2017 ACC/AHA/AAPA/ABC/ACPM/AGS/APhA/ASH/ASPC/NMA/PCNA Guideline for the Prevention, Detection, Evaluation, and Management of High Blood Pressure in Adults

Executive Summary

A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines

WRITING COMMITTEE MEMBERS

Paul K. Whelton, MB, MD, MSc, FAHA, *Chair* Robert M. Carey, MD, FAHA, *Vice Chair*

Wilbert S. Aronow, MD, FACC, FAHA* Donald E. Casey, Jr, MD, MPH, MBA, FAHA† Karen J. Collins, MBA‡ Cheryl Dennison Himmelfarb, RN, ANP, PhD, FAHA§ Sondra M. DePalma, MHS, PA-C, CLS, AACC Samuel Gidding, MD, FACC, FAHA¶ Kenneth A. Jamerson, MD# Daniel W. Jones, MD, FAHA† Eric J. MacLaughlin, PharmD** Paul Muntner, PhD, FAHA† Bruce Ovbiagele, MD, MSc, MAS, MBA, FAHA⁺ Sidney C. Smith, Jr, MD, MACC, FAHA⁺⁺ Crystal C. Spencer, JD[‡] Randall S. Stafford, MD, PhD^{‡‡} Sandra J. Taler, MD, FAHA§§ Randal J. Thomas, MD, MS, FACC, FAHA || || Kim A. Williams, Sr, MD, MACC, FAHA⁺ Jeff D. Williamson, MD, MHS¶¶ Jackson T. Wright, Jr, MD, PhD, FAHA##

ACC/AHA TASK FORCE MEMBERS

Glenn N. Levine, MD, FACC, FAHA, Chair Patrick T. O'Gara, MD, FAHA, MACC, Chair-Elect Jonathan L. Halperin, MD, FACC, FAHA, Immediate Past Chair Sana M. Al-Khatib, MD, MHS, FACC, FAHA Federico Gentile, MD, FACC Joshua A. Beckman, MD, MS, FAHA Samuel Gidding, MD, FAHA*** Zachary D. Goldberger, MD, MS, FACC, FAHA Kim K. Birtcher, MS, PharmD, AACC Biykem Bozkurt, MD, PhD, FACC, FAHA*** Mark A. Hlatky, MD, FACC, FAHA Ralph G. Brindis, MD, MPH, MACC*** John Ikonomidis, MD, PhD, FAHA Joaquin E. Cigarroa, MD, FACC José A. Joglar, MD, FACC, FAHA Lesley H. Curtis, PhD, FAHA*** Laura Mauri, MD, MSc, FAHA Susan J. Pressler, PhD, RN, FAHA*** Anita Deswal, MD, MPH, FACC, FAHA Lee A. Fleisher, MD, FACC, FAHA Barbara Riegel, PhD, RN, FAHA

Duminda N. Wijeysundera, MD, PhD

*American Society for Preventive Cardiology Representative. †ACC/AHA Representative. ‡Lay Volunteer/Patient Representative. §Preventive Cardiovascular Nurses Association Representative. || American Academy of Physician Assistants Representative. ¶Task Force Liaison. #Association of Black Cardiologists Representative. **American Pharmacists Association Representative. †*ACC/AHA Prevention Subcommittee Liaison. ‡‡American College of Preventive Medicine Representative. §§American Society of Hypertension Representative. || || Task Force on

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Performance Measures Liaison. ¶¶American Geriatrics Society Representative. ##National Medical Association Representative. ***Former Task Force member; current member during the writing effort.

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Preamble

Since 1980, the American College of Cardiology (ACC) and American Heart Association (AHA) have translated scientific evidence into clinical practice guidelines (guidelines) with recommendations to improve cardiovascular health. In 2013, the National Heart, Lung, and Blood Institute (NHLBI) Advisory Council recommended that the NHLBI focus specifically on reviewing the highest-quality evidence and partner with other organizations to develop recommendations (1, 2). Accordingly, the ACC and AHA collaborated with the NHLBI and stakeholder and professional organizations to complete and publish 4 guidelines (on assessment of cardiovascular risk, lifestyle modifications to reduce cardiovascular risk, management of blood cholesterol in adults, and management of overweight and obesity in adults) to make them available to the widest possible constituency. In 2014, the ACC and AHA, in partnership with several other professional societies, initiated a guideline on the prevention, detection, evaluation, and management of high blood pressure (BP) in adults. Under the management of the ACC/AHA Task Force, a Prevention Subcommittee was appointed to help guide development of the suite of guidelines on prevention of cardiovascular disease (CVD). These guidelines, which are based on systematic methods to evaluate and classify evidence, provide a cornerstone for quality cardiovascular care. The ACC and AHA sponsor the development and publication of guidelines without commercial support, and members of each organization volunteer their time to the writing and review efforts. Guidelines are official policy of the ACC and AHA.

Intended Use

Practice guidelines provide recommendations applicable to patients with or at risk of developing CVD. The focus is on medical practice in the United States, but guidelines developed in collaboration with other organizations can have a global impact. Although guidelines may be used to inform regulatory or payer decisions, they are intended to improve patients' quality of care and align with patients' interests. Guidelines are intended to define practices meeting the needs of patients in most, but not all, circumstances and should not replace clinical judgment.

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Clinical Implementation

Management in accordance with guideline recommendations is effective only when followed by both practitioners and patients. Adherence to recommendations can be enhanced by shared decision making between clinicians and patients, with patient engagement in selecting interventions on the basis of individual values, preferences, and associated conditions and comorbidities.

Methodology and Modernization

The ACC/AHA Task Force on Clinical Practice Guidelines (Task Force) continuously reviews, updates, and modifies guideline methodology on the basis of published standards from organizations, including the Institute of Medicine (3, 4), and on the basis of internal reevaluation. Similarly, the presentation and delivery of guidelines are reevaluated and modified on the basis of evolving technologies and other factors to facilitate optimal dissemination of information to healthcare professionals at the point of care.

Toward this goal, this guideline continues the introduction of an evolved format of presenting guideline recommendations and associated text called the "modular knowledge chunk format." Each modular "chunk" includes a table of related recommendations, a brief synopsis, recommendation-specific supportive text, and when appropriate, flow diagrams or additional tables. References are provided within the modular chunk itself to facilitate quick review. Additionally, this format will facilitate seamless updating of guidelines with focused updates as new evidence is published, as well as content tagging for rapid electronic retrieval of related recommendations; therefore, the present document represents a transitional format that best suits the text as written. Future guidelines will fully implement this format, including provisions for limiting the amount of text in a guideline.

Recognizing the importance of cost–value considerations in certain guidelines, when appropriate and feasible, an analysis of the value of a drug, device, or intervention may be performed in accordance with the ACC/AHA methodology (5).

To ensure that guideline recommendations remain current, new data are reviewed on an ongoing basis, with full guideline revisions commissioned in approximately 6-year cycles. Publication of new, potentially practice-changing study results that are relevant to an existing or new drug, device, or management strategy will prompt evaluation by the Task Force, in consultation with the relevant guideline writing committee, to determine whether a focused update should be commissioned. For additional information and policies regarding guideline development, we encourage readers to consult the ACC/AHA guideline methodology manual (6) and other methodology articles (7-10).

Selection of Writing Committee Members

The Task Force strives to avoid bias by selecting experts from a broad array of backgrounds. Writing committee members represent different geographic regions, sexes, ethnicities, races, intellectual perspectives/biases, and scopes of clinical practice. The Task Force may also invite organizations and professional societies with related interests and expertise to participate as partners, collaborators, or endorsers.

Relationships With Industry and Other Entities

The ACC and AHA have rigorous policies and methods to ensure that guidelines are developed without bias or improper influence. The complete relationships with industry and other entities (RWI) policy can be found <u>at http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/relationships-with-industry-policy</u>. Appendix 1 of the present document lists writing committee members' relevant RWI. For the purposes of full transparency, writing committee members' comprehensive disclosure information is available online (<u>http://hyper.ahajournals.org/lookup/suppl/doi:10.1161/HYP.00000000000066/-/DC1</u>). Comprehensive disclosure information for the Task Force is available at <u>http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/guidelines-and-clinical-documents/guidelines-and-clinical-documents/guidelines-and-clinical-documents/guidelines-and-clinical-documents/guidelines-and-clinical-documents/guidelines-and-clinical-documents/guidelines-and-clinical-documents/guidelines-and-documents-task-forces.</u>

Evidence Review and Evidence Review Committees

In developing recommendations, the writing committee uses evidence-based methodologies that are based on all available data (6-9). Literature searches focus on randomized controlled trials (RCTs) but also include registries, nonrandomized comparative and descriptive studies, case series, cohort studies, systematic reviews, and expert opinion. Only key references are cited.

An independent evidence review committee (ERC) is commissioned when there are 1 or more questions deemed of utmost clinical importance that merit formal systematic review. The systematic review will determine which patients are most likely to benefit from a drug, device, or treatment strategy and to what degree. Criteria for commissioning an ERC and formal systematic review include: a) the absence of a current authoritative systematic review, b) the feasibility of defining the benefit and risk in a time frame consistent with the writing of a guideline, c) the relevance to a substantial number of patients, and d) the likelihood that the findings can be translated into actionable recommendations. ERC members may include methodologists, epidemiologists, healthcare providers, and biostatisticians. The recommendations developed by the writing committee on the basis of the systematic review are marked with "SR".

Guideline-Directed Management and Therapy

The term guideline-directed management and therapy (GDMT) encompasses clinical evaluation, diagnostic testing, and pharmacological and procedural treatments. For these and all recommended drug treatment regimens, the reader should confirm the dosage by reviewing product insert material and evaluate the treatment regimen for contraindications and interactions. The recommendations are limited to drugs, devices, and treatments approved for clinical use in the United States.

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Class of Recommendation and Level of Evidence

The Class of Recommendation (COR) indicates the strength of the recommendation, encompassing the estimated magnitude and certainty of benefit in proportion to risk. The Level of Evidence (LOE) rates the quality of scientific evidence that supports the intervention on the basis of the type, quantity, and consistency of data from clinical trials and other sources (Table 1) (6-8).

The reader is encouraged to consult the full-text guideline (11) for additional guidance and details about hypertension, since the executive summary contains mainly the recommendations

Glenn N. Levine, MD, FACC, FAHA Chair, ACC/AHA Task Force on Clinical Practice Guidelines

> American Heart Association

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Table 1. Applying Class of Recommendation and Level of Evidence to Clinical Strategies, Interventions, Treatments, or Diagnostic Testing in Patient Care* (Updated August 2015)

CLASS (STRENGTH) OF RECOMMENDATION

Suggested phrases for writing recommendations:

Is recommended

CLASS I (STRONG)

- Is indicated/useful/effective/beneficial
- Should be performed/administered/other
- Comparative-Effectiveness Phrasest:
- Treatment/strategy A is recommended/indicated in preference to treatment B
- Treatment A should be chosen over treatment B

Suggested phrases for writing recommendations:

- Is reasonable
- Can be useful/effective/beneficial
- Comparative-Effectiveness Phrases†:
- Treatment/strategy A is probably recommended/indicated in preference to treatment B
- It is reasonable to choose treatment A over treatment B

CLASS IIb (WEAK)

Suggested phrase ing recommendations:

- May/might be
- May/might be
- Usefulness/effe s is unknown/unclear/uncertain or not well esta

CLASS III: No Bene RATE)

Benefit = Risk

Risk > Benefit

- Suggested phrase ng recommendations:
- Is not recommended
- Is not indicated/useful/effective/beneficial
- Should not be performed/administered/other

CLASS III: Harm (STRONG)

- Suggested phrases for writing recommendations:
- Potentially harmful
- Causes harm
- Associated with excess morbidity/mortality
- Should not be performed/administered/other

LEVEL (QUALITY) OF EVIDENCE[±]

LEVEL A

- · High-quality evidencet from more than 1 RCT
- Meta-analyses of high-quality RCTs
- One or more RCTs corroborated by high-quality registry studies

LEVEL B-R

- Moderate-quality evidence[‡] from 1 or more RCTs
- Meta-analyses of moderate-quality RCTs

LEVEL B-NR

(Nonrandomized)

(Randomized)

- Moderate-quality evidencet from 1 or more well-designed, well-executed nonrandomized studies, observational studies, or registry studies
- Meta-analyses of such studies

- Randomized or nonrandomized observational or registry studies with limitations of design or execution
- Meta-analyses of such studies
- Physiological or mechanistic studies in human subjects

Consensus of expert opinion based on clinical experience

COR and LOE are determined independently (any COR may be paired with any LOE).

A recommendation with LOE C does not imply that the recommendation is weak. Many important clinical questions addressed in guidelines do not lend themselves to clinical trials. Although RCTs are unavailable, there may be a very clear clinical consensus that a particular test or therapy is useful or effective.

- * The outcome or result of the intervention should be specified (an improved clinical outcome or increased diagnostic accuracy or incremental prognostic information).
- † For comparative-effectiveness recommendations (COR I and IIa; LOE A and B only), studies that support the use of comparator verbs should involve direct comparisons of the treatments or strategies being evaluated.
- the method of assessing quality is evolving, including the application of standardized, widely used, and preferably validated evidence grading tools; and for systematic reviews, the incorporation of an Evidence Review Committee.
- COR indicates Class of Recommendation; EO, expert opinion; LD, limited data; LOE, Level of Evidence; NR, nonrandomized; R, randomized; and RCT, randomized controlled trial.

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1. Introduction

In 2013, the National Heart, Lung, and Blood Institute (NHLBI) Advisory Council recommended that the NHLBI focus specifically on reviewing the highest-quality evidence and partner with other organizations to develop recommendations (1, 2). Accordingly, the ACC and AHA collaborated with the NHLBI and stakeholder and professional organizations to complete and publish 4 guidelines (on assessment of cardiovascular risk, lifestyle modifications to reduce cardiovascular risk, management of blood cholesterol in adults, and management of overweight and obesity in adults) to make them available to the widest possible constituency. In 2014, the ACC and AHA in partnership with several other professional societies initiated a guideline on the prevention, detection, evaluation and management of high blood pressure in adults. Under the management of the ACC/AHA Task Force, a Prevention Subcommittee was appointed to help guide development of the suite of guidelines on prevention of cardiovascular disease.

As early as the 1920s, and subsequently in the 1959 Build and Blood Pressure Study (3) of almost 5 million adults insured between 1934 and 1954, a strong direct relationship was noted between level of BP and risk of clinical complications and death. In the 1960s, these findings were confirmed in a series of reports from the Framingham Heart Study (4). The 1967 and 1970 Veterans Administration Cooperative Study Group reports ushered in the era of effective treatment for high BP (5, 6). The first comprehensive guideline for detection, evaluation, and management of high BP was published in 1977, under the sponsorship of the NHLBI (7). In subsequent years, a series of Joint National Committee (JNC) BP guidelines were published to assist the practice community and improve prevention, awareness, treatment, and control of high BP (7-9). The present guideline updates prior JNC reports.

1.1. Methodology and Evidence Review

An extensive evidence review, which included literature derived from research involving human subjects, published in English, and indexed in MEDLINE (through PubMed), EMBASE, the Cochrane Library, the Agency for Healthcare Research and Quality, and other selected databases relevant to this guideline, was conducted between February and August 2015. Key search words included but were not limited to the following: adherence; aerobic; alcohol intake; ambulatory care; antihypertensive: agents, drug, medication, therapy; beta adrenergic blockers; blood pressure: arterial, control, determination, devices, goal, high, improve, measurement, monitoring, ambulatory; calcium channel blockers; diet; diuretic agent; drug therapy; heart failure: diastolic, systolic; hypertension: white coat, masked, ambulatory, isolated ambulatory, isolated clinic, diagnosis, reverse white coat, prevention, therapy, treatment, control; intervention; lifestyle: measures, modification; office visits; patient outcome; performance measures; physical activity; potassium intake; protein intake; renin inhibitor; risk reduction: behavior, counseling; screening; sphygmomanometers; spironolactone; therapy; treatment: adherence, compliance, efficacy, outcome, protocol, regimen; weight. Additional relevant studies published through June 2016, during the guideline writing process, were also considered by the writing committee and added to the evidence tables when appropriate. The final evidence tables included the Online Data Supplement in (http://hyper.ahajournals.org/lookup/suppl/doi:10.1161/HYP.000000000000066/-/DC2) summarize the evidence used by the writing committee to formulate recommendations.

As noted in the preamble, an independent ERC was commissioned to perform a formal systematic review of 4 critical clinical questions related to hypertension (Table 2), the results of which were considered by the writing committee for incorporation into this guideline. Concurrent with this process, writing committee members evaluated other published data relevant to the guideline. The findings of the ERC and the writing committee members were formally presented and discussed, and then guideline recommendations were developed. The systematic review report, "Systematic Review for the 2017 ACC/AHA/AAPA/ABC/ACPM/AGS/APhA/ASH/ASPC/NMA/PCNA Guideline for the Prevention, Detection,

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Evaluation, and Management of High Blood Pressure in Adults," is published in conjunction with this guideline (10), and its respective data supplements are available online (http://hyper.ahajournals.org/lookup/suppl/doi:10.1161/HYP.0000000000000067/-/DC2). No writing committee member reported a RWI. Drs. Whelton, Wright and Williamson had leadership roles in SPRINT (Systolic Blood Pressure Intervention Trial). Dr. Carey chaired committee discussions in which the SPRINT results were considered.

Question		Section
Number	Question	Number
1	Is there evidence that self-directed monitoring of BP and/or ambulatory BP monitoring are superior to office-based measurement of BP by a healthcare worker for 1) preventing adverse outcomes for which high BP is a risk factor and 2) achieving better BP control?	4.2
2	What is the optimal target for BP lowering during antihypertensive therapy in adults?	8.1.5 9.3 9.6
3	In adults with hypertension, do various antihypertensive drug classes differ in their comparative benefits and harms?	8.1.6 8.2
4	In adults with hypertension, does initiating treatment with antihypertensive pharmacological monotherapy versus initiating treatment with 2 drugs (including fixed- dose combination therapy), either of which may be followed by the addition of sequential drugs, differ in comparative benefits and/or harms on specific health outcomes?	An8.1.6.1 Heart Association

BP indicates blood pressure.

1.2. Organization of the Writing Committee

The writing committee consisted of clinicians, cardiologists, epidemiologists, internists, an endocrinologist, a geriatrician, a nephrologist, a neurologist, a nurse, a pharmacist, a physician assistant, and 2 lay/patient representatives. It included representatives from the ACC, AHA, American Academy of Physician Assistants (AAPA), Association of Black Cardiologists (ABC), American College of Preventive Medicine (ACPM), American Geriatrics Society (AGS), American Pharmacists Association (APhA), American Society of Hypertension (ASH), American Society for Preventive Cardiology (ASPC), National Medical Association (NMA), and Preventive Cardiovascular Nurses Association (PCNA).

1.3. Document Review and Approval

This document was reviewed by 2 official reviewers nominated by the ACC and AHA; 1 reviewer each from the AAPA, ABC, ACPM, AGS, APhA, ASH, ASPC NMA, and PCNA; and 38 individual content reviewers. Reviewers' RWI information was distributed to the writing committee and is published in this document (Appendix 2).

This document was approved for publication by the governing bodies of the ACC, AHA, AAPA, ABC, ACPM, AGS, APhA, ASH, ASPC, NMA, and PCNA.

1.4. Scope of the Guideline

The present guideline is intended to be a resource for the clinical and public health practice communities. It is designed to be comprehensive but succinct and practical in providing guidance for prevention, detection, evaluation, and management of high BP. It is an update of the NHLBI publication, "The Seventh Report of the Joint National Committee on Prevention, Detection, Evaluation and Treatment of High Blood Pressure" (JNC

7) (9). It incorporates new information from studies of office-based BP-related risk of CVD, ambulatory blood pressure monitoring (ABPM), home blood pressure monitoring (HBPM), telemedicine, and various other areas. This guideline does not address the use of BP-lowering medications for the purposes of prevention of recurrent CVD events in patients with stable ischemic heart disease (SIHD) or chronic heart failure (HF) in the absence of hypertension; these topics are the focus of other ACC/AHA guidelines (11, 12). In developing the present guideline, the writing committee reviewed prior published guidelines, evidence reviews, and related statements. Table 3 contains a list of publications and statements deemed pertinent to this writing effort and is intended for use as a resource, thus obviating the need to repeat existing guideline recommendations.

Title	Organization	Publication Year	
Guidelines			
Lower-extremity peripheral artery	AHA/ACC	2016 (13)	
disease			
Management of primary	Endocrine Society	2016 (14)	
aldosteronism: case detection,			
diagnosis, and treatment			
Stable ischemic heart disease	ACC/AHA/AATS/PCNA/SCAI/STS	2014 (15)*2012 (11)	
Pheochromocytoma and	Endocrine Society	2014 (16)	
paraganglioma		American	
Atrial fibrillation	AHA/ACC/HRS	2014 (17)	
Valvular heart disease	ACC/AHA	2017 (18)	
Assessment of cardiovascular risk	ACC/AHA	2013 (19)	
Hypertension in pregnancy	ACOG	2013 (20)	
Heart failure	ACC/AHA	2017 (21)	
T 740	012t 010	2013 (12)	
Lifestyle management to reduce	AHA/ACC	2013 (22)	
cardiovascular risk			
Management of arterial	ESH/ESC	2013 (23)	
hypertension			
Management of overweight and	AHA/ACC/TOS	2013 (24)	
obesity in adults			
ST-elevation myocardial infarction	ACC/AHA	2013 (25)	
Treatment of blood cholesterol to	ACC/AHA	2013 (26)	
reduce atherosclerotic			
cardiovascular risk in adults			
Cardiovascular diseases during	ESC	2011 (27)	
pregnancy			
Effectiveness-based guidelines for	AHA/ACC	2011 (28)	
the prevention of cardiovascular			
disease in women			
Secondary prevention and risk-	AHA/ACC	2011 (29)	
reduction therapy for patients with			
coronary and other atherosclerotic			
vascular disease			
Assessment of cardiovascular risk in	ACC/AHA	2010 (30)	
asymptomatic adults			
Thoracic aortic disease	ACC/AHA/AATS/ACR/ASA/SCA/SCAI/SIR/	2010 (31)	
	STS/SVM		
Diagnosis, evaluation, and	NHLBI	2004 (32)	
treatment of high blood pressure in			
children and adolescents			

Table 3. Associated Guidelines and Statements

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Statements		
Salt sensitivity of blood pressure	AHA	2016 (33)
Cardiovascular team-based care and	ACC	2015 (34)
the role of advanced practice		
providers		
Treatment of hypertension in	AHA/ACC/ASH	2015 (35)
patients with coronary artery		
disease		
Ambulatory blood pressure	AHA	2014 (36)
monitoring in children and		
adolescents		
An effective approach to high blood	AHA/ACC/CDC	2014 (37)
pressure control		
Ambulatory blood pressure	ESH	2013 (38)
monitoring		
Performance measures for adults	ACC/AHA/AMA-PCPI	2011 (39)
with coronary artery disease and		
hypertension		
Interventions to promote physical	AHA	2010 (40)
activity and dietary lifestyle changes		American
for cardiovascular risk factor		Gib Heart
reduction in adults		Association
Resistant hypertension: diagnosis,	AHA	2008 (41)
evaluation, and treatment		

*The full-text SIHD guideline is from 2012 (11). A focused update was published in 2014 (15).

AATS indicates American Association for Thoracic Surgery; ACC, American College of Cardiology; ACOG, American College of Obstetricians and Gynecologists; ACR, American College of Radiology; AHA, American Heart Association; AMA, American Medical Association; ASA, American Stroke Association; ASH, American Society of Hypertension; CDC, Centers for Disease Control and Prevention; ESC, European Society of Cardiology; ESH, European Society of Hypertension; HRS, Heart Rhythm Society; NHLBI, National Heart, Lung, and Blood Institute; PCNA, Preventive Cardiovascular Nurses Association; PCPI, Physician Consortium for Performance Improvement; SCA, Society of Cardiovascular Anesthesiologists; SCAI, Society for Cardiovascular Angiography and Interventions; SIHD, stable ischemic heart disease; SIR, Society of Interventional Radiology; STS, Society of Thoracic Surgeons; SVM, Society for Vascular Medicine; and TOS, The Obesity Society.

1.5. Abbreviations and Acronyms

Abbreviation/Acronym	Meaning/Phrase
ABPM	ambulatory blood pressure monitoring
ACE	angiotensin-converting enzyme
AF	atrial fibrillation
ARB	angiotensin receptor blocker
BP	blood pressure
ССВ	calcium channel blocker
CHD	coronary heart disease
CKD	chronic kidney disease
СРАР	continuous positive airway pressure
CVD	cardiovascular disease
DBP	diastolic blood pressure
DM	diabetes mellitus
ECG	electrocardiogram

ESRD	end-stage renal disease
GDMT	guideline-directed management and therapy
GFR	glomerular filtration rate
HBPM	home blood pressure monitoring
EHR	electronic health record
HF	heart failure
HF <i>p</i> EF	heart failure with preserved ejection fraction
HF <i>r</i> EF	heart failure with reduced ejection fraction
ICH	intracerebral hemorrhage
JNC	Joint National Commission
LV	left ventricular
LVH	left ventricular hypertrophy
MI	myocardial infarction
MRI	magnetic resonance imaging
PAD	peripheral artery disease
RAS	renin-angiotensin system
RCT	randomized controlled trial
SBP	systolic blood pressure
SIHD	stable ischemic heart disease
TIA	transient ischemic attack

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2. BP and CVD Risk

2.1. Observational Relationship

Observational studies have demonstrated graded associations between higher systolic blood pressure (SBP) and diastolic blood pressure (DBP) and increased CVD risk (1, 2). In a meta-analysis of 61 prospective studies, the risk of CVD increased in a log-linear fashion from SBP levels <115 mm Hg to >180 mm Hg and from DBP levels <75 mm Hg to >105 mm Hg (1). In that analysis, 20 mm Hg higher SBP and 10 mm Hg higher DBP were each associated with a doubling in the risk of death from stroke, heart disease, or other vascular disease. In a separate observational study including >1 million adult patients ≥30 years of age, higher SBP and DBP were associated with increased risk of CVD incidence and angina, myocardial infarction (MI), HF, stroke, peripheral artery disease (PAD), and abdominal aortic aneurysm, each evaluated separately (2). An increased risk of CVD associated with higher SBP and DBP has been reported across a broad age spectrum, from 30 years to >80

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years of age. Although the relative risk of incident CVD associated with higher SBP and DBP is smaller at older ages, the corresponding high BP–related increase in absolute risk is larger in older persons (≥65 years) given the higher absolute risk of CVD at an older age (1).

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2.2. BP Components

Epidemiological studies have evaluated associations of SBP and DBP, as well as derived components of BP measurements (including pulse pressure, mean BP, and mid-BP), with CVD outcomes (Table 4). When considered separately, higher levels of both SBP and DBP have been associated with increased CVD risk (1, 2). Higher SBP has consistently been associated with increased CVD risk after adjustment for, or within strata of, DBP (3-5). In contrast, after consideration of SBP through adjustment or stratification, DBP has not been consistently associated with CVD risk (6, 7). Although pulse pressure and mid-BP have been associated with increased CVD risk independent of SBP and DBP in some studies, SBP (especially) and DBP are prioritized in the present document because of the robust evidence base for these measures in both observational studies and clinical trials and because of their ease of measurement in practice settings (8-11).

Table 4. BP Measurement Definitions

BP Measurement	Definition
SBP	First Korotkoff sound*
DBP	Fifth Korotkoff sound*
Pulse pressure	SBP minus DBP
Mean arterial pressure	DBP plus one third pulse pressure ⁺
Mid-BP	Sum of SBP and DBP, divided by 2

*See Section 4 for a description of Korotkoff sounds.

[†]Calculation assumes normal heart rate.

BP indicates blood pressure; DBP, diastolic blood pressure; and SBP, systolic blood pressure.

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2.3. Population Risk

In 2010, high BP was the leading cause of death and disability-adjusted life years worldwide (1, 2). In the United States, hypertension (see Section 3.1 for definition) accounted for more CVD deaths than any other modifiable CVD risk factor and was second only to cigarette smoking as a preventable cause of death for any reason (3). In a follow-up study of 23,272 U.S. NHANES (National Health and Nutrition Examination Survey) participants, >50% of deaths from coronary heart disease (CHD) and stroke occurred among individuals with hypertension (4). Because of the high prevalence of hypertension and its associated increased risk of CHD, stroke, and end-stage renal disease (ESRD), the population-attributable risk of these outcomes associated with hypertension is high (4, 5). In the population-based ARIC (Atherosclerosis Risk in Communities) study, 25% of the cardiovascular events (CHD, coronary revascularization, stroke, or HF) were attributable to hypertension. In the Northern Manhattan study, the percentage of events attributable to hypertension was higher in women (32%) than in men (19%) and higher in blacks (36%) than in whites (21%) (6). In 2012, hypertension was the second leading assigned cause of ESRD, behind diabetes mellitus (DM), and accounted for 34% of incident ESRD cases in the U.S. population (7).

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2.4. Coexistence of Hypertension and Related Chronic Conditions

Recommendation for Coexistence of Hypertension and Related Chronic Conditions			
Refere	References that support the recommendation are summarized in Online Data Supplement 1.		
COR	LOE	Recommendation	
		1. Screening for and management of other modifiable CVD risk factors are	
l I	B-NR	recommended in adults with hypertension (1, 2).	

Table 5. CVD Risk Factors Common in Patients With Hypertension

Modifiable Risk Factors*	Relatively Fixed Risk Factors [†]
Current cigarette smoking, secondhand smoking	• CKD
Diabetes mellitus	Family history
Dyslipidemia/hypercholesterolemia	Increased age
 Overweight/obesity 	 Low socioeconomic/educational status
Physical inactivity/low fitness	Male sex
Unhealthy diet	Obstructive sleep apnea
,	Psychosocial stress

*Factors that can be changed and, if changed, may reduce CVD risk.

⁺Factors that are difficult to change (CKD, low socioeconomic/educational status, obstructive sleep apnea (3)), cannot be changed (family history, increased age, male sex), or, if changed through the use of current intervention techniques, may not reduce CVD risk (psychosocial stress) (3).

CKD indicates chronic kidney disease; and CVD, cardiovascular disease.

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3. Classification of BP

3.1. Definition of High BP

Recommendation for Definition of High BP			
References that support the recommendation are summarized in Online Data Supplement 2.			
COR	LOE	Recommendation	
I	B-NR	1. BP should be categorized as normal, elevated, or stage 1 or 2 hypertension to prevent and treat high BP (Table 6) (1-20).	

BP Category	SBP		DBP
Normal	<120 mm Hg	and	<80 mm Hg
Elevated	120–129 mm Hg	and	<80 mm Hg
Hypertension			
Stage 1	130–139 mm Hg	or	80–89 mm Hg
Stage 2	≥140 mm Hg	or	≥90 mm Hg

Table 6. Categories of BP in Adults*

*Individuals with SBP and DBP in 2 categories should be designated to the higher BP category. BP indicates blood pressure (based on an average of ≥2 careful readings obtained on ≥2 occasions, as detailed in Section 4); DBP, diastolic blood pressure; and SBP systolic blood pressure.

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3.2. Lifetime Risk of Hypertension

Observational studies have documented a relatively high incidence of hypertension over periods of 5 to 10 years of follow-up (1, 2). Thus, there is a much higher long-term population burden of hypertension as BP progressively increases with age. Several studies have estimated the long-term cumulative incidence of developing hypertension (3, 4). In an analysis of 1132 white male medical students (mean age: approximately 23 years at baseline) in the Johns Hopkins Precursors study, 0.3%, 6.5%, and 37% developed hypertension at age 25, 45, and 65 years, respectively (5). In MESA (Multi-Ethnic Study of Atherosclerosis), the percentage of the population developing hypertension over their lifetimes was higher for African Americans and Hispanics than for whites and Asians (3). For adults 45 years of age without hypertension, the 40-year risk of developing hypertension was 93% for African-American, 92% for Hispanic, 86% for white, and 84% for Chinese adults (3). In the Framingham Heart Study, approximately 90% of adults free of hypertension at age 55 or 65 years developed hypertension during their lifetimes (4). All of these estimates were based on use of the 140/90– mm Hg cutpoint for recognition of hypertension and would have been higher had the 130/80–mm Hg cutpoint been used.



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- 2. Parikh NI, Pencina MJ, Wang TJ, et al. A risk score for predicting near-term incidence of hypertension: the Framingham Heart Study. Ann Intern Med. 2008;148:102-10.
- 3. Carson AP, Howard G, Burke GL, et al. Ethnic differences in hypertension incidence among middle-aged and older adults: the multi-ethnic study of atherosclerosis. Hypertension. 2011;57:1101-7.
- 4. Vasan RS, Beiser A, Seshadri S, et al. Residual lifetime risk for developing hypertension in middle-aged women and men: the Framingham Heart Study. JAMA. 2002;287:1003-10.
- 5. Shihab HM, Meoni LA, Chu AY, et al. Body mass index and risk of incident hypertension over the life course: the Johns Hopkins Precursors Study. Circulation. 2012;126:2983-9.

3.3. Prevalence of High BP

	SBP/DBP ≥130/80 mm Hg or Self- Reported Antihypertensive Medication†		SBP/DBP ≥140/90 mm Hg or Self- Reported Antihypertensive Medication‡	
Overall, crude	46%		32%	
	Men (n=4717)	Women (n=4906)	Men (n=4717)	Women (n=4906)
Overall, age-sex	48%	43%	31%	32%
adjusted				
Age group, y				
20–44	30%	19%	11%	10%
45–54	50%	44%	33%	27%
55–64	70%	63%	53%	52%
65–74	77%	75%	64%	63%
75+	79%	85%	71%	78%
Race-ethnicity§				
Non-Hispanic white	47%	41%	31%	30%
Non-Hispanic black	59%	56%	42%	46%
Non-Hispanic Asian	45%	36%	29%	27% American
Hispanic	44%	42%	27%	32% Heart

Table 7. Prevalence of I	lypertension Based on 2 SBF	<pre>/DBP Thresholds*†</pre>

The prevalence estimates have been rounded to the nearest full percentage.

*130/80 and 140/90 mm Hg in 9623 participants (≥20 years of age) in NHANES 2011–2014.

⁺BP cutpoints for definition of hypertension in the present guideline.

‡BP cutpoints for definition of hypertension in JNC 7.

§Adjusted to the 2010 age-sex distribution of the U.S. adult population.

BP indicates blood pressure; DBP, diastolic blood pressure; NHANES, National Health and Nutrition Examination Survey; and SBP, systolic blood pressure.

4. Measurement of BP

4.1. Accurate Measurement of BP in the Office

Recommendation for Accurate Measurement of BP in the Office				
COR	LOE	Recommendation		
I	C-EO	 For diagnosis and management of high BP, proper methods are recommended for accurate measurement and documentation of BP (Table 8). 		

Key Steps for Proper BP	Specific Instructions
Measurements	
Step 1: Properly prepare the patient	 Have the patient relax, sitting in a chair (feet on floor, back supported) for >5 min. The patient should avoid caffeine, exercise, and smoking for at least 30 min before measurement. Ensure patient has emptied his/her bladder. Neither the patient nor the observer should talk during the rest period or during the measurement. Remove all clothing covering the location of cuff placement. Measurements made while the patient is sitting or lying on an examining table do not fulfill these criteria.
Step 2: Use proper technique for BP measurements	 Use a BP measurement device that has been validated, and ensure that the device is calibrated periodically.* Support the patient's arm (e.g., resting on a desk). Position the middle of the cuff on the patient's upper arm at the level of the right atrium (the midpoint of the sternum). Use the correct cuff size, such that the bladder encircles 80% of the arm, and note if a larger- or smaller-than-normal cuff size is used (Table 9). Either the stethoscope diaphragm or bell may be used for auscultatory readings (3, 4).
Step 3: Take the proper measurements needed for diagnosis and treatment of elevated BP/hypertension	 At the first visit, record BP in both arms. Use the arm that gives the higher reading for subsequent readings. Separate repeated measurements by 1–2 min. For auscultatory determinations, use a palpated estimate of radial pulse obliteration pressure to estimate SBP. Inflate the cuff 20–30 mm Hg above this level for an auscultatory determination of the BP level. For auscultatory readings, deflate the cuff pressure 2 mm Hg per second, and listen for Korotkoff sounds.
Step 4: Properly document accurate BP readings	 Record SBP and DBP. If using the auscultatory technique, record SBP and DBP as onset of the first Korotkoff sound and disappearance of all Korotkoff sounds, respectively, using the nearest even number. Note the time of most recent BP medication taken before measurements.
Step 5: Average the readings	Use an average of ≥ 2 readings obtained on ≥ 2 occasions to estimate the individual's level of BP.
Step 6: Provide BP readings to patient *See Section 4.2 for additional g	Provide patients the SBP/DBP readings both verbally and in writing.

Table 8. Checklist for Accurate Measurement of BP (1, 2)

*See Section 4.2 for additional guidance.

BP indicates blood pressure; DBP, diastolic blood pressure; and SBP, systolic blood pressure.

Adapted with permission from Mancia et al. (1) (Oxford University Press), Pickering et al. (5) (American Heart Association, Inc.), and Weir et al. (2) (American College of Physicians, Inc.).

Table 9. Selection Criteria for BP Cuff Size for Measurement of BP in Adults

Arm Circumference	Usual Cuff Size
22–26 cm	Small adult
27–34 cm	Adult
35–44 cm	Large adult
45–52 cm	Adult thigh

Adapted with permission from Pickering et al. (5) (American Heart Association, Inc.). BP indicates blood pressure.

References

- 1. Mancia G, Fagard R, Narkiewicz K, et al. 2013 ESH/ESC guidelines for the management of arterial hypertension: the Task Force for the Management of Arterial Hypertension of the European Society of Hypertension (ESH) and of the European Society of Cardiology (ESC). Eur Heart J. 2013;34:2159-219.
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- 4. Kantola I, Vesalainen R, Kangassalo K, et al. Bell or diaphragm in the measurement of blood pressure? J Hypertens. 2005;23:499-503.
- Pickering TG, Hall JE, Appel LJ, et al. Recommendations for blood pressure measurement in humans and experimental animals: part 1: blood pressure measurement in humans: a statement for professionals from the Subcommittee of Professional and Public Education of the American Heart Association Council on High Blood Pressure Research. Circulation. 2005;111:697-716.

4.2. Out-of-Office and Self-Monitoring of BP

Recommendation for Out-of-Office and Self-Monitoring of BP			
References that support the recommendation are summarized in Online Data Supplement 3 and			
Systematic Review Report. Association			
COR	LOE	Recommendation	
I.	A ^{SR}	1. Out-of-office BP measurements are recommended to confirm the diagnosis of hypertension (Table 11) and for titration of BP-lowering medication, in conjunction with telehealth counseling or clinical interventions (1-4).	

SR indicates systematic review.

Table 10. Procedures for Use of HBPM (5-7)

Patient training should occur under medical supervision, including:

- Information about hypertension
- Selection of equipment
- Acknowledgment that individual BP readings may vary substantially
- Interpretation of results

Devices:

• Verify use of automated validated devices. Use of auscultatory devices (mercury, aneroid, or other) is not generally useful for HBPM because patients rarely master the technique required for measurement of BP with auscultatory devices.

- Monitors with provision for storage of readings in memory are preferred.
- Verify use of appropriate cuff size to fit the arm (Table 9).
- Verify that left/right inter-arm differences are insignificant. If differences are significant, instruct patient to measure BPs in the arm with higher readings.

Instructions on HBPM procedures:

- Remain still:
 - Avoid smoking, caffeinated beverages, or exercise within 30 min before BP measurements.
 - Ensure ≥5 min of quiet rest before BP measurements.
- Sit correctly:

- Sit with back straight and supported (on a straight-backed dining chair, for example, rather than a sofa).
- Sit with feet flat on the floor and legs uncrossed.
- Keep arm supported on a flat surface (such as a table), with the upper arm at heart level.
- Bottom of the cuff should be placed directly above the antecubital fossa (bend of the elbow).
- Take multiple readings:
 - Take at least 2 readings 1 min apart in morning before taking medications and in evening before supper. Optimally, measure and record BP daily. Ideally, obtain weekly BP readings beginning 2 weeks after a change in the treatment regimen and during the week before a clinic visit.
- Record all readings accurately:
 - Monitors with built-in memory should be brought to all clinic appointments.
 - BP should be based on an average of readings on ≥ 2 occasions for clinical decision making.

The information above may be reinforced with videos available online:

http://www.heart.org/HEARTORG/Conditions/HighBloodPressure/SymptomsDiagnosisMonitoringofHighBloodPressure/Home-Blood-Pressure-Monitoring_UCM_301874_Article.jsp#.WcQNfLKGMnM

See Table 11 for HBPM targets.

BP indicates blood pressure; and HBPM, home blood pressure monitoring.

Table 11. Corresponding Values of SBP/DBP for Clinic, HBPM, Daytime, Nighttime, and 24-Hour ABPM Measurements

Association					
Clinic	НВРМ	Daytime ABPM	Nighttime ABPM	24-Hour ABPM	
120/80	120/80	120/80	100/65	115/75	
130/80	130/80	130/80	110/65	125/75	
140/90	135/85	135/85	120/70	130/80	
160/100	145/90	145/90	140/85	145/90	

ABPM indicates ambulatory blood pressure monitoring; BP, blood pressure; DBP diastolic blood pressure; HBPM, home blood pressure monitoring; and SBP, systolic blood pressure.

- 1. Uhlig K, Balk EM, Patel K, et al. Self-Measured Blood Pressure Monitoring: Comparative Effectiveness. Rockville, MD: Agency for Healthcare Research and Quality (U.S.); 2012.
- 2. Margolis KL, Asche SE, Bergdall AR, et al. Effect of home blood pressure telemonitoring and pharmacist management on blood pressure control: a cluster randomized clinical trial. JAMA. 2013;310:46-56.
- 3. McManus RJ, Mant J, Haque MS, et al. Effect of self-monitoring and medication self-titration on systolic blood pressure in hypertensive patients at high risk of cardiovascular disease: the TASMIN-SR randomized clinical trial. JAMA. 2014;312:799-808.
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4.3. Ambulatory BP Monitoring

4.4. Masked and White Coat Hypertension

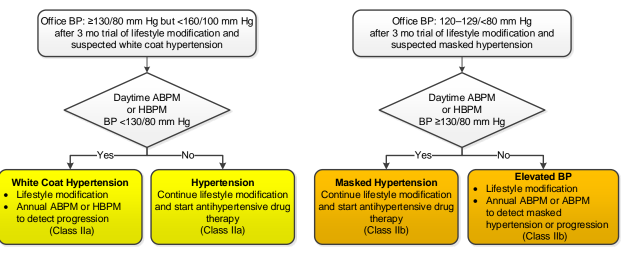
	Recommendations for Masked and White Coat Hypertension				
Reference	References that support recommendations are summarized in Online Data Supplements 4, 5, and 6.				
COR	LOE	Recommendation			
lla	B-NR	 In adults with an untreated SBP greater than 130 mm Hg but less than 160 mm Hg or DBP greater than 80 mm Hg but less than 100 mm Hg, it is reasonable to screen for the presence of white coat hypertension by using either daytime ABPM or HBPM before diagnosis of hypertension (1-8). 			
lla	C-LD	2. In adults with white coat hypertension, periodic monitoring with either ABPM or HBPM is reasonable to detect transition to sustained hypertension (2, 5, 7).			
lla	C- LD	3. In adults being treated for hypertension with office BP readings not at goal and HBPM readings suggestive of a significant white coat effect, confirmation by ABPM can be useful (9, 10).			
lla	B-NR	4. In adults with untreated office BPs that are consistently between 120 mm Hg and 129 mm Hg for SBP or between 75 mm Hg and 79 mm Hg for DBP, screening for masked hypertension with HBPM (or ABPM) is reasonable (3, 4, 6, 8, 11).			
llb	C-LD	5. In adults on multiple-drug therapies for hypertension and office BPs within 10 mm Hg above goal, it may be reasonable to screen for white coat effect with HBPM (or ABPM) (3, 7, 12).			
llb	C-EO	6. It may be reasonable to screen for masked uncontrolled hypertension with HBPM in adults being treated for hypertension and office readings at goal, in the presence of target organ damage or increased overall CVD risk.			
llb	C-EO	7. In adults being treated for hypertension with elevated HBPM readings suggestive of masked uncontrolled hypertension, confirmation of the diagnosis by ABPM might be reasonable before intensification of antihypertensive drug treatment.			

Table 12. BP Patterns Based on Office and Out-of-Office Measurements
--

	Office/Clinic/Healthcare Setting	Home/Nonhealthcare/ABPM Setting
Normotensive	No hypertension	No hypertension
Sustained hypertension	Hypertension	Hypertension
Masked hypertension	No hypertension	Hypertension
White coat hypertension	Hypertension	No hypertension

ABPM indicates ambulatory blood pressure monitoring; and BP, blood pressure.



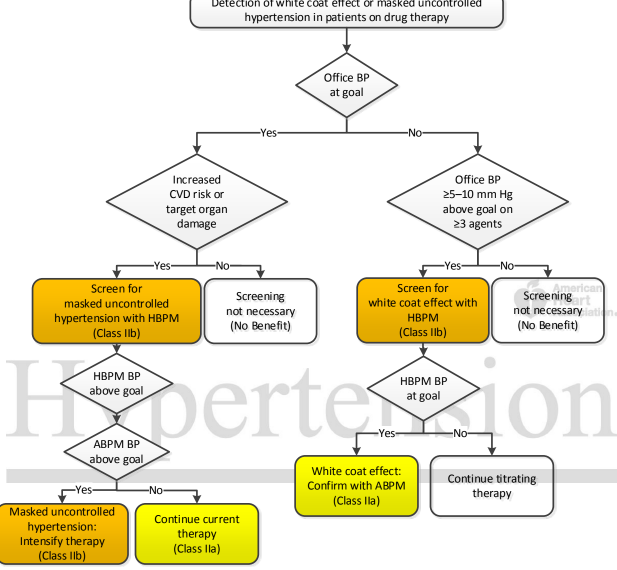


Colors correspond to Class of Recommendation in Table 1.

ABPM indicates ambulatory blood pressure monitoring; BP, blood pressure; and HBPM, home blood pressure monitoring.



Figure 2. Detection of White Coat Effect or Masked Uncontrolled Hypertension in Patients on Drug Therapy
Detection of white coat effect or masked uncontrolled



Colors correspond to Class of Recommendation in Table 1.

See Section 8 for treatment options.

ABPM indicates ambulatory blood pressure monitoring; BP, blood pressure; CVD, cardiovascular disease; and HBPM, home blood pressure monitoring.

- 1. Pickering TG, James GD, Boddie C, et al. How common is white coat hypertension? JAMA. 1988;259:225-8.
- 2. Piper MA, Evans CV, Burda BU, et al. Diagnostic and predictive accuracy of blood pressure screening methods with consideration of rescreening intervals: a systematic review for the U.S. Preventive Services Task Force. Ann Intern Med. 2015;162:192-204.
- 3. Ohkubo T, Kikuya M, Metoki H, et al. Prognosis of "masked" hypertension and "white-coat" hypertension detected by 24-h ambulatory blood pressure monitoring 10-year follow-up from the Ohasama study. J Am Coll Cardiol. 2005;46:508-15.

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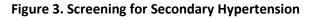
- 4. Fagard RH, Cornelissen VA. Incidence of cardiovascular events in white-coat, masked and sustained hypertension versus true normotension: a meta-analysis. J Hypertens. 2007;25:2193-8.
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- 9. Viera AJ, Hinderliter AL, Kshirsagar AV, et al. Reproducibility of masked hypertension in adults with untreated borderline office blood pressure: comparison of ambulatory and home monitoring. Am. J Hypertens. 2010;23:1190-7.
- 10. Viera AJ, Lin FC, Tuttle LA, et al. Reproducibility of masked hypertension among adults 30 years or older. Blood Press Monit. 2014;19:208-15.
- 11. Stergiou GS, Asayama K, Thijs L, et al. Prognosis of white-coat and masked hypertension: International Database of HOme blood pressure in relation to Cardiovascular Outcome. Hypertension. 2014;63:675-82.
- 12. Tomiyama M, Horio T, Yoshii M, et al. Masked hypertension and target organ damage in treated hypertensive patients. Am J Hypertens. 2006; 19:880-6.

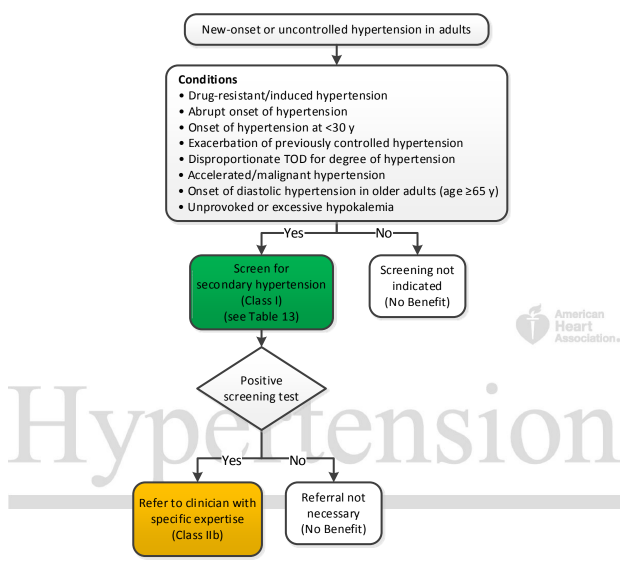


5. Causes of Hypertension

5.1. Secondary Forms of Hypertension

	Recommendations for Secondary Forms of Hypertension						
COR	LOE	Recommendations					
I	C-EO	1. Screening for specific form(s) of secondary hypertension is recommended when the clinical indications and physical examination findings listed in Table 13 are present or in adults with resistant hypertension.					
llb	C-EO	2. If an adult with sustained hypertension screens positive for a form of secondary hypertension, referral to a physician with expertise in that form of hypertension may be reasonable for diagnostic confirmation and treatment.					





Colors correspond to Class of Recommendation in Table 1.

TOD indicates target organ damage (e.g., cerebrovascular disease, hypertensive retinopathy, left ventricular hypertrophy, left ventricular dysfunction, heart failure, coronary artery disease, chronic kidney disease, albuminuria, peripheral artery disease).

	Prevalence	Physical Clinical Indications Examination		Screening Tests	Additional/ Confirmatory Tests
Common causes					
Renal parenchymal	1%–2%	Urinary tract infections;	Abdominal	Renal	Tests to
disease (1, 2)		obstruction, hematuria;	mass	ultrasound	evaluate cause
		urinary frequency and	(polycystic		of renal
		nocturia; analgesic	kidney		disease
		abuse; family history of			

		polycystic kidney	disease); skin		
		disease; elevated	pallor		
			panoi		
		serum creatinine;			
		abnormal urinalysis			B ¹
Renovascular	5%-34%*	Resistant hypertension;	Abdominal	Renal	Bilateral
disease (3)		hypertension of abrupt	systolic-	Duplex	selective renal
		onset or worsening or	diastolic bruit;	Doppler	intra-arterial
		increasingly difficult to	bruits over	ultrasound;	angiography
		control; flash	other arteries	MRA;	
		pulmonary edema	(carotid –	abdominal	
		(atherosclerotic); early-	atherosclerotic	СТ	
		onset hypertension,	or		
		especially in women	fibromuscular		
		(fibromuscular	dysplasia),		
		hyperplasia)	femoral		
Primary	8%-20%†	Resistant hypertension;	Arrhythmias	Plasma	Oral sodium
aldosteronism (4,		hypertension with	(with	aldosterone	loading test
5)		hypokalemia	hypokalemia);	/renin ratio	(with 24-h
		(spontaneous or	especially	under	urineleart
		diuretic induced);	atrial	standardize	aldosterone)
		hypertension and	fibrillation	d conditions	or IV saline
		muscle cramps or		(correction	infusion test
		weakness;		of	with plasma
	710	hypertension and		hypokalemia	aldosterone at
		incidentally discovered		and	4 h of infusion
		adrenal mass;		withdrawal	Adrenal CT
		hypertension and		of	scan,
		obstructive sleep		aldosterone	adrenal vein
		apnea; hypertension		antagonists	sampling.
		and family history of		for 4–6 wk)	BB-
		early-onset			
		hypertension or stroke			
		hypertension of stroke			
Obstructive sleep	25%-50%	Resistant hypertension;	Obesity,	Berlin	Polysomnogra
apnea (6)‡	2370 3070	snoring; fitful sleep;	Mallampati	Questionnai	phy
aprica (0)+		breathing pauses	class III–IV;	re (7);	pily
		during sleep; daytime	loss of normal	Epworth	
			nocturnal BP	-	
		sleepiness	fall	Sleepiness	
			Idli	Score (8); overnight	
				oximetry	
Drug or alashal	20/ 40/	Sodium containing	Fine tremer	-	Rosponso to
Drug or alcohol	2%–4%	Sodium-containing	Fine tremor,	Urinary drug	Response to
induced (9)§		antacids; caffeine;	tachycardia,	screen (illicit	withdrawal of
		nicotine (smoking);	sweating	drugs)	suspected
		alcohol; NSAIDs; oral	(cocaine,		agent
		contraceptives;	ephedrine,		

		cyclosporine or tacrolimus; sympathomimetics (decongestants, anorectics); cocaine,	MAO inhibitors); acute abdominal pain (cocaine)		
		amphetamines and other illicit drugs; neuropsychiatric agents; erythropoiesis- stimulating agents;			
		clonidine withdrawal; herbal agents (Ma Huang, ephedra)			
Uncommon causes					
Pheochromocytom a/paraganglioma (10)	0.1%-0.6%	Resistant hypertension; paroxysmal hypertension or crisis superimposed on	Skin stigmata of neurofibromat osis (café-au-	24-h urinary fractionated metanephri nes or	CT or MRI scan of abdomen/pelv is
		sustained hypertension; "spells," BP lability, headache, sweating, palpitations,	lait spots; neurofibromas); Orthostatic	plasma metanephri nes under standard	American Heart Association
Hy	/p	pallor; positive family history of pheochromocytoma/ paraganglioma; adrenal incidentaloma	hypotension	conditions (supine position with indwelling	31
				IV cannula)	
Cushing's	<0.1%	Rapid weight gain,	Central	Overnight 1-	24-h urinary
syndrome (11)		especially with central distribution; proximal	obesity, "moon" face,	mg dexamethas	free cortisol excretion
		muscle weakness; depression; hyperglycemia	dorsal and supraclavicular fat pads, wide	one suppression test	(preferably multiple); midnight
			(1-cm) violaceous		salivary cortisol
			striae, hirsutism		
Hypothyroidism (9)	<1%	Dry skin; cold intolerance; constipation; hoarseness; weight gain	Delayed ankle reflex; periorbital puffiness; coarse skin;	Thyroid- stimulating hormone; free thyroxine	None
			cold skin; slow movement; goiter		

1 h	-10/		Lidles for a	Thumpiel	Dediesst
Hyperthyroidism	<1%	Warm, moist skin; heat	Lid lag; fine	Thyroid-	Radioactive
(9)		intolerance;	tremor of the	stimulating	iodine uptake
		nervousness;	outstretched	hormone;	and scan
		tremulousness;	hands; warm,	free	
		insomnia; weight loss;	moist skin	thyroxine	
		diarrhea; proximal			
		muscle weakness			
Aortic coarctation	0.1%	Young patient with	BP higher in	Echocardiog	Thoracic and
(undiagnosed or		hypertension (<30 y of	upper	ram	abdominal CT
repaired) (12)		age)	extremities		angiogram or
			than in lower		MRA
			extremities;		
			absent		
			femoral		
			pulses;		
			continuous		
			murmur over		
			patient's back,		
			chest, or		American Heart
					Association
			abdominal		
			bruit; left		
			thoracotomy	•	
			scar		
		ATTA	(postoperative		
)		
Primary	Rare	Hypercalcemia	Usually none	Serum	Serum
hyperparathyroidis				calcium	parathyroid
m (13)					hormone
Congenital adrenal	Rare	Hypertension and	Signs of	Hypertensio	11-beta-OH:
hyperplasia (14)		hypokalemia;	virilization (11-	n and	elevated
		virilization (11-beta-	beta-OH) or	hypokalemia	deoxycorticost
		hydroxylase deficiency	incomplete	with low or	erone (DOC),
		[11-beta-OH]);	masculinizatio	normal	11-
		incomplete	n (17-alpha-	aldosterone	deoxycortisol,
		masculinization in	OH)	and renin	and
		males and primary	,		androgens17-
		amenorrhea in females			alpha-OH;
		(17-alpha-hydroxylase			decreased
		deficiency [17-alpha-			androgens and
		OH])			estrogen;
		Unjj			_
					elevated
					deoxycorticost
					erone and
					corticosterone

Mineralocorticoid	Rare	Early-onset	Arrhythmias	Low	Urinary
excess syndromes		hypertension; resistant	(with	aldosterone	cortisol
other than primary		hypertension;	hypokalemia)	and renin	metabolites;
aldosteronism (14)		hypokalemia or			genetic testing
		hyperkalemia			
Acromegaly (15)	Rare	Acral features,	Acral features;	Serum	Elevated age-
		enlarging shoe, glove,	large hands	growth	and sex-
		or hat size; headache,	and feet;	hormone ≥1	matched IGF-1
		visual disturbances;	frontal bossing	ng/mL	level; MRI scan
		diabetes mellitus		during oral	of the
				glucose load	pituitary

*Depending on the clinical situation (hypertension alone, 5%; hypertension starting dialysis, 22%; hypertension and peripheral vascular disease, 28%; hypertension in the elderly with congestive heart failure, 34%).

*8% in general population with hypertension; up to 20% in patients with resistant hypertension.
 *Although obstructive sleep apnea is listed as a cause of secondary hypertension, RCTs on the effects of continuous positive airway pressure on lowering BP in patients with hypertension have produced mixed results (see Section 5.4.4 for details).

§For a list of frequently used drugs causing hypertension and accompanying evidence, see Table 14. BP indicates blood pressure; CT, computed tomography; DOC, 11-deoxycorticosterone; IGF-1, insulin-like growth factor-1; IV, intravenous; MAO, monamine oxidase; MRI, magnetic resonance imaging; MRA, magnetic resonance arteriography; NSAIDs, nonsteroidal anti-inflammatory drugs; OH, hydroxylase; and RCT, randomized clinical trial.

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5.1.1. Drugs and Other Substances With Potential to Impair BP Control

Agent	Possible Management Strategy
Alcohol	 Limit alcohol to ≤1 drink daily for women and ≤2 drinks for men (1)
Amphetamines (e.g., amphetamine,	Discontinue or decrease dose (2)
methylphenidate dexmethylphenidate,	Consider behavioral therapies for ADHD (3)
dextroamphetamine)	
Antidepressants (e.g., MAOIs, SNRIs, TCAs)	 Consider alternative agents (e.g., SSRIs) depending on part indication
	Avoid tyramine-containing foods with MAOIs
Atypical antipsychotics (e.g., clozapine,	Discontinue or limit use when possible
olanzapine)	Consider behavior therapy where appropriate
	Recommend lifestyle modification (see Section 6.2)
	Consider alternative agents associated with lower risk of
	weight gain, diabetes mellitus, and dyslipidemia (e.g.,
	aripiprazole, ziprasidone) (4, 5)
Caffeine	 Generally limit caffeine intake to <300 mg/d
	Avoid use in patients with uncontrolled hypertension
	Coffee use in patients with hypertension is associated with
	acute increases in BP; long-term use is not associated with
	increased BP or CVD (6)
Decongestants (e.g., phenylephrine,	Use for shortest duration possible, and avoid in severe or
pseudoephedrine)	uncontrolled hypertension
	Consider alternative therapies (e.g., nasal saline, intranasal
	corticosteroids, antihistamines) as appropriate
Herbal supplements (e.g., Ma Huang	Avoid use
[ephedra], St. John's wort [with MAO	
inhibitors, yohimbine])	
Immunosuppressants (e.g., cyclosporine)	Consider converting to tacrolimus, which may be associated
	with fewer effects on BP (7-9)
Over a second base	
Oral contraceptives	• Use low-dose (e.g., 20–30 mcg ethinyl estradiol) agents (10) or
Urai contraceptives	• Use low-dose (e.g., 20–30 mcg ethinyl estradiol) agents (10) or a progestin-only form of contraception, or consider
Urai contraceptives	
Urai contraceptives	a progestin-only form of contraception, or consider

Table 14. Frequently Used Medications and Other Substances That May Cause Elevated BP*

NSAIDs	٠	Avoid systemic NSAIDs when possible
	•	Consider alternative analgesics (e.g., acetaminophen,
		tramadol, topical NSAIDs), depending on indication and risk
Recreational drugs (e.g., "bath salts"	٠	Discontinue or avoid use
[MDPV], cocaine, methamphetamine,		
etc.)		
Systemic corticosteroids (e.g.,	٠	Avoid or limit use when possible
dexamethasone, fludrocortisone,	•	Consider alternative modes of administration (e.g., inhaled,
methylprednisolone, prednisone,		topical) when feasible
prednisolone)		
Angiogenesis inhibitor (e.g., bevacizumab)	٠	Initiate or intensify antihypertensive therapy
and tyrosine kinase inhibitors (e.g.,		
sunitinib, sorafenif)		

*List is not all inclusive.

ADHD indicates attention-deficit/hyperactivity disorder; BP, blood pressure; CVD, cardiovascular disease; IUD, intrauterine device; MAOI, monoamine-oxidase inhibitors; MDPV, methylenedioxypyrovalerone; NSAIDs, nonsteroidal antiinflammatory drugs; SNRI, serotonin norepinephrine reuptake inhibitor; SSRI, selective serotonin reuptake inhibitor; and TCA, tricyclic antidepressant.

American Heart Association

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5.1.2. Primary Aldosteronism

	Recommendations for Primary Aldosteronism				
COR	LOE	Recommendations			
I	C-EO	1. In adults with hypertension, screening for primary aldosteronism is recommended in the presence of any of the following concurrent conditions: resistant hypertension, hypokalemia (spontaneous or substantial, if diuretic induced), incidentally discovered adrenal mass, family history of early-onset hypertension, or stroke at a young age (<40 years).			
I.	C-LD	2. Use of the plasma aldosterone: renin activity ratio is recommended when adults are screened for primary aldosteronism (1).			
I	C-EO	3. In adults with hypertension and a positive screening test for primary aldosteronism, referral to a hypertension specialist or endocrinologist is recommended for further evaluation and treatment.			

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5.1.3. Renal Artery Stenosis

	Recommendations for Renal Artery Stenosis				
COR	LOE Recommendations				
I.	Α	1. Medical therapy is recommended for adults with atherosclerotic renal artery stenosis (1, 2).			
IIb	C-EO	2. In adults with renal artery stenosis for whom medical management has failed (refractory hypertension, worsening renal function, and/or intractable HF) and those with nonatherosclerotic disease, including fibromuscular dysplasia, it may be reasonable to refer the patient for consideration of revascularization (percutaneous renal artery angioplasty and/or stent placement).			

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5.1.4. Obstructive Sleep Apnea

Recommendation for Obstructive Sleep Apnea				
COR	COR LOE Recommendations			
lib	B-R	1. In adults with hypertension and obstructive sleep apnea, the effectiveness of continuous positive airway pressure (CPAP) to reduce BP is not well established (1-5).		

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		Recommendations for Nonpharmacological Interventions			
Refer	References that support recommendations are summarized in Online Data Supplements 9-21.				
COR	LOE	Recommendations			
1	A	1. Weight loss is recommended to reduce BP in adults with elevated BP or hypertension who are overweight or obese (1-4).			
I	А	2. A heart-healthy diet, such as the DASH (Dietary Approaches to Stop Hypertension) diet, that facilitates achieving a desirable weight is recommended for adults with elevated BP or hypertension (5-7).			
I.	А	3. Sodium reduction is recommended for adults with elevated BP or hypertension (8-12).			
I	A	4. Potassium supplementation, preferably in dietary modification, is recommended for adults with elevated BP or hypertension, unless contraindicated by the presence of CKD or use of drugs that reduce potassium excretion (13-17).			
I.	А	5. Increased physical activity with a structured exercise program is recommended for adults with elevated BP or hypertension (3, 4, 12, 18-22).			
I	А	6. Adult men and women with elevated BP or hypertension who currently consume alcohol should be advised to drink no more than 2 and 1 standard drinks* per day, respectively (23-28).			

6. Nonpharmacological Interventions

*In the United States, 1 "standard" drink contains roughly 14 g of pure alcohol, which is typically found in 12 oz of regular beer (usually about 5% alcohol), 5 oz of wine (usually about 12% alcohol), and 1.5 oz of distilled spirits (usually about 40% alcohol) (29).

	Nonpharmacological	Dose	Approximate Impact on SBP			
	Intervention		Hypertension	Normotension	Reference	
Weight loss	Weight/body fat	Best goal is ideal	-5 mm Hg	-2/3 mm Hg	(1)	
0	0, ,	body weight, but aim	0	, 0		
		for at least a 1-kg				
		reduction in body				
		weight for most				
		adults who are				
		overweight. Expect				
		about 1 mm Hg for				
		every 1-kg reduction				
		in body weight.				
Healthy	DASH dietary	Consume a diet rich	-11 mm Hg	-3 mm Hg	(6, 7)	
diet ,	pattern	in fruits, vegetables,	0	U		
	•	whole grains, and				
		low-fat dairy				
		products, with				
		reduced content of				
		saturated and total			d	
		fat.			American Heart	
Reduced	Dietary sodium	Optimal goal is <1500	-5/6 mm Hg	-2/3 mm Hg	(9, 10) sociation	
intake of		mg/d, but aim for at				
dietary		least a 1000-mg/d				
sodium		reduction in most				
		adults.				
Enhanced	Dietary potassium	Aim for 3500–5000	-4/5 mm Hg	-2 mm Hg	(13)	
intake of		mg/d, preferably by				
dietary		consumption of a diet				
potassium		rich in potassium.				
Physical	Aerobic	• 90–150 min/wk	-5/8 mm Hg	-2/4 mm Hg	(18, 22)	
activity		● 65%–75% heart				
		rate reserve				
	Dynamic resistance	● 90–150 min/wk	-4 mm Hg	-2 mm Hg	(18)	
		● 50%–80% 1 rep				
		maximum				
		 6 exercises, 3 				
		sets/exercise, 10				
		repetitions/set				
	Isometric resistance	 4 × 2 min (hand 	-5 mm Hg	-4 mm Hg	(19, 30)	
		grip), 1 min rest				
		between exercises,				
		30%–40% maximum				
		voluntary				
		contraction, 3				
		sessions/wk				
		● 8–10 wk				
Moderation	Alcohol consumption	In individuals who	-4 mm Hg	-3 mm Hg	(22-24)	
in alcohol		drink alcohol, reduce				
intake		alcohol† to:				
		 Men: ≤2 drinks 				
		daily				

Table 15. Best Proven Nonpharmacological Interventions for Prevention and Treatment of Hypertension*

● Women: ≤1 drink		
daily		

*Type, dose, and expected impact on BP in adults with a normal BP and with hypertension. DASH indicates Dietary Approaches to Stop Hypertension; and SBP, systolic blood pressure. Resources:

Your Guide to Lowering Your Blood Pressure With DASH—How Do I Make the DASH? Available at: https://www.nhlbi.nih.gov/health/resources/heart/hbp-dash-how-to. Accessed September 15, 2017. (31) Top 10 Dash Diet Tips. Available at: http://dashdiet.org/dash_diet_tips.asp. Accessed September 15, 2017. (32) †In the United States, one "standard" drink contains roughly 14 g of pure alcohol, which is typically found in 12 oz of regular beer (usually about 5% alcohol), 5 oz of wine (usually about 12% alcohol), and 1.5 oz of distilled spirits (usually about 40% alcohol) (29).

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7. Patient Evaluation

Table 16. Historical Features Favoring Hypertension Cause

Primary Hypertension	Secondary Hypertension
Gradual increase in BP, with slow rate	BP lability, episodic pallor and dizziness (pheochromocytoma)
of rise in BP	Snoring, hypersomnolence (obstructive sleep apnea)
Lifestyle factors that favor higher BP	Prostatism (chronic kidney disease due to post-renal urinary
(e.g., weight gain, high-sodium diet,	tract obstruction)
decreased physical activity, job change	Muscle cramps, weakness (hypokalemia from primary
entailing increased travel, excessive	aldosteronism or secondary aldosteronism due to
consumption of alcohol)	renovascular disease)
Family history of hypertension	Weight loss, palpitations, heat intolerance (hyperthyroidism)
	Edema, fatigue, frequent urination (kidney disease or failure)
	History of coarctation repair (residual hypertension associated
	with coarctation)
	Central obesity, facial rounding, easy bruisability (Cushing's
	syndrome)

•	•	Medication or substance use (e.g., alcohol, NSAIDS, cocaine,
		amphetamines)
	•	Absence of family history of hypertension
DD indicates blood pressure, and NSAIDs, peneto		idal anti inflommatany druga

BP indicates blood pressure; and NSAIDs, nonsteroidal anti-inflammatory drugs.

7.1. Laboratory Tests and Other Diagnostic Procedures

Table 17. Basic and Optional Laboratory	Tests for Primary Hypertension
---	--------------------------------

Fasting blood glucose*
Complete blood count
Lipid profile
Serum creatinine with eGFR*
Serum sodium, potassium, calcium*
Thyroid-stimulating hormone
Urinalysis
Electrocardiogram
Echocardiogram
Uric acid
Urinary albumin to creatinine ratio

*May be included in a comprehensive metabolic panel. eGFR indicates estimated glomerular filtration rate.

8. Treatment of High BP

8.1. Pharmacological Treatment

8.1.1. Initiation of Pharmacological BP Treatment in the Context of Overall CVD Risk

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For any specific difference in BP, the relative risk of CVD is constant across groups that differ in absolute risk of atherosclerotic CVD (1-4), albeit with some evidence of lesser relative risk but greater excess risk in older than in younger adults (5-8). Thus, there are more potentially preventable CVD events attributable to elevated BP in individuals with higher than with lower risk of CVD and in older than in younger adults. The relative risk reduction for CVD prevention with use of BP-lowering medications is fairly constant for groups that differ in CVD risk across a wide range of estimated absolute risk (9, 10) and across groups defined by sex, age, body mass index, and the presence or absence of DM, AF, and CKD (5, 11-21). As a consequence, the absolute CVD risk reduction attributable to BP lowering is greater at greater absolute levels of CVD risk (9, 10, 12, 15-19, 22, 23). Put another way, for a given magnitude of BP reduction due to antihypertensive medications, fewer individuals at high CVD risk would need to be treated to prevent a CVD event (i.e., lower number needed to treat) than those at low CVD risk.

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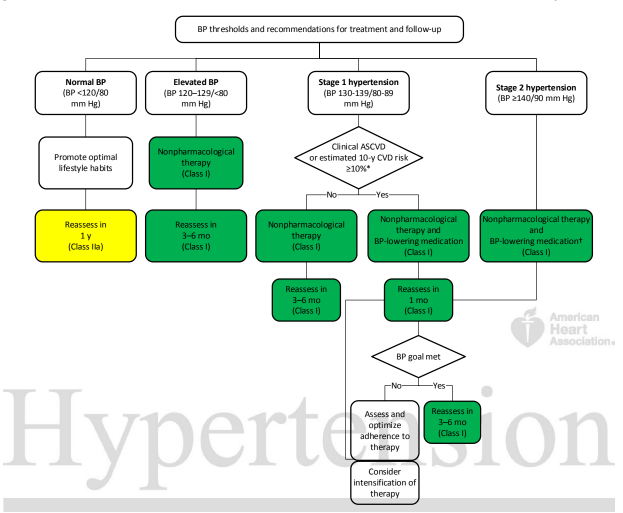
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8.1.2. BP Treatment Threshold and the Use of CVD Risk Estimation to Guide Drug Treatment of Hypertension

Recomm	Recommendations for BP Treatment Threshold and Use of Risk Estimation* to Guide Drug					
	Treatment of Hypertension					
Refe	References that support recommendations are summarized in Online Data Supplement 23.					
COR	COR LOE Recommendations					
I	SBP: A DBP: C-EO	1. Use of BP-lowering medications is recommended for secondary prevention of recurrent CVD events in patients with clinical CVD and an average SBP of 130 mm Hg or higher or an average DBP of 80 mm Hg or higher, and for primary prevention in adults with an estimated 10-year atherosclerotic cardiovascular disease (ASCVD) risk of 10% or higher and an average SBP 130 mm Hg or higher or an average DBP 80 mm Hg or higher (1-9).				
I	C-LD	 Use of BP-lowering medication is recommended for primary prevention of CVD in adults with no history of CVD and with an estimated 10-year ASCVD risk <10% and an SBP of 140 mm Hg or higher or a DBP of 90 mm Hg or higher (3, 10-13). 				

*ACC/AHA Pooled Cohort Equations (http://tools.acc.org/ASCVD-Risk-Estimator/) (13a) to estimate 10-year risk of atherosclerotic CVD. ASCVD was defined as a first CHD death, non-fatal MI or fatal or non-fatal stroke.







Colors correspond to Class of Recommendation in Table 1.

*Using the ACC/AHA Pooled Cohort Equations (14). Note that patients with DM or CKD are automatically placed in the high-risk category. For initiation of RAS inhibitor or diuretic therapy, assess blood tests for electrolytes and renal function 2 to 4 weeks after initiating therapy.

[†]Consider initiation of pharmacological therapy for stage 2 hypertension with 2 antihypertensive agents of different classes. Patients with stage 2 hypertension and BP \geq 160/100 mm Hg should be promptly treated, carefully monitored, and subject to upward medication dose adjustment as necessary to control BP. Reassessment includes BP measurement, detection of orthostatic hypotension in selected patients (e.g., older or with postural symptoms), identification of white coat hypertension or a white coat effect, documentation of adherence, monitoring of the response to therapy, reinforcement of the importance of adherence, reinforcement of the importance of treatment, and assistance with treatment to achieve BP target.

ACC indicates American College of Cardiology; AHA, American Heart Association; ASCVD, atherosclerotic cardiovascular disease; BP, blood pressure; CKD, chronic kidney disease; DM, diabetes mellitus; and RAS, renin-angiotensin system.

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8.1.3. Follow-Up After Initial BP Evaluation

Recommendations for Follow-Up After Initial BP Elevation						
Refe	References that support recommendations are summarized in Online Data Supplement 24.					
COR	LOE	Recommendations				
I	B-R	 Adults with an elevated BP or stage 1 hypertension who have an estimated 10-year ASCVD risk less than 10% should be managed with nonpharmacological therapy and have a repeat BP evaluation within 3 to 6 months (1, 2). 				
I	B-R	2. Adults with stage 1 hypertension who have an estimated 10-year ASCVD risk of 10% or higher should be managed initially with a combination of nonpharmacological and antihypertensive drug therapy and have a repeat BP evaluation in 1 month (1, 2).				
I	B-R	3. Adults with stage 2 hypertension should be evaluated by or referred to a primary care provider within 1 month of the initial diagnosis, have a combination of nonpharmacological and antihypertensive drug therapy (with 2 agents of different classes) initiated, and have a repeat BP evaluation in 1 month (1, 2).				
I	B-R	 For adults with a very high average BP (e.g., SBP ≥180 mm Hg or DBP ≥110 mm Hg), evaluation followed by prompt antihypertensive drug treatment is recommended (1, 2). 				
lla	C-EO	5. For adults with a normal BP, repeat evaluation every year is reasonable.				

References

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8.1.4. General Principles of Drug Therapy

Refe	Recommendation for General Principle of Drug Therapy References that support recommendations are summarized in Online Data Supplement 25.				
COR	LOE	Recommendation			
III: Harm	Α	6. Simultaneous use of an ACE inhibitor, ARB, and/or renin inhibitor is potentially harmful and is not recommended to treat adults with hypertension (1-3).			

Table 18. Oral Antihypertensive Drugs

Class	Drug	Usual Dose, Range (mg/day)*	Daily Frequency	Comments	
Primary agents	Primary agents				
	Chlorthalidone	12.5–25	1		
	Hydrochlorothiazide	25–50	1		

Thiazide or	Indapamide	1.25-2.5	1	•	Chlorthalidone is preferred on the basis of
thiazide-type	Metolazone	2.5–10	1	1	prolonged half-life and proven trial reduction
diuretics	Wietoldzone				of CVD.
					Monitor for hyponatremia and hypokalemia,
				-	uric acid and calcium levels.
					Use with caution in patients with history of
					acute gout unless patient is on uric acid–
					lowering therapy.
					lowering therapy.
ACE	Benazepril	10-40	1 or 2	•	Do not use in combination with ARBs or direct
inhibitors	Captopril	12.5-150	2 or 3		renin inhibitor.
	Enalapril	5–40	1 or 2	٠	There is an increased risk of hyperkalemia,
	Fosinopril	10–40	1		especially in patients with CKD or in those on K ⁺
	Lisinopril	10–40	1		supplements or K⁺-sparing drugs.
	Moexipril	7.5–30	1 or 2	•	There is a risk of acute renal failure in patients
	Perindopril	4–16	1		with severe bilateral renal artery stenosis.
	Quinapril	10-80	1 or 2	•	Do not use if patient has history of angioedema
	Ramipril	2.5–10	1 or 2		with ACE inhibitors.
	Trandolapril	1–4	1	•	Avoid in pregnancy.
ARBs	Azilsartan	40–80	1	٠	Do not use in combination with ACE inhibitors
	Candesartan	8–32	1		or direct renin inhibitor. Heart
	Eprosartan	600–800	1 or 2	٠	There is an increased risk of hyperkalemia in
	Irbesartan	150-300	1		CKD or in those on K ⁺ supplements or K ⁺ -
	Losartan	50–100	1 or 2		sparing drugs.
	Olmesartan	20–40	1	٠	There is a risk of acute renal failure in patients
	Telmisartan	20-80	1		with severe bilateral renal artery stenosis.
	Valsartan	80–320	1	•	Do not use if patient has history of angioedema
					with ARBs. Patients with a history of
					angioedema with an ACE inhibitor can receive
					an ARB beginning 6 weeks after ACE inhibitor is
					discontinued.
				•	Avoid in pregnancy.
CCB—	Amlodipine	2.5–10	1	•	Avoid use in patients with HFrEF; amlodipine or
dihydropyridi	Felodipine	5–10	1	_	felodipine may be used if required.
nes	Isradipine	5–10	2	•	They are associated with dose-related pedal
	Nicardipine SR	5–20	1	_	edema, which is more common in women than
	Nifedipine LA	60–120	1	_	men.
	Nisoldipine	30–90	1		
CCB—	Diltiazem SR	180–360	2	٠	Avoid routine use with beta blockers because
nondihydrop	Diltiazem ER	120–480	1		of increased risk of bradycardia and heart
yridines	Verapamil IR	40-80	3		block.
	Verapamil SR	120–480	1 or 2	•	Do not use in patients with HF <i>r</i> EF.
	Verapamil-delayed	100–480	1 (in the	•	There are drug interactions with diltiazem and
	onset ER (various		evening)		verapamil (CYP3A4 major substrate and
	forms)				moderate inhibitor).
Secondary age					
Diuretics—	Bumetanide	0.5-4	2	•	These are preferred diuretics in patients with
loop	Furosemide	20-80	2		symptomatic HF. They are preferred over thiazides in patients with moderate-to-severe
	Torsemide	5–10	1		
	Amilorida	E 10	1 05 2	\vdash	CKD (e.g., GFR <30 mL/min).
	Amiloride	5–10	1 or 2	<u> </u>	

Diuretics— potassium sparing	Triamterene	50–100	1 or 2	 These are monotherapy agents and minimally effective antihypertensive agents. Combination therapy of potassium-sparing diuretic with a thiazide can be considered in patients with hypokalemia on thiazide monotherapy. Avoid in patients with significant CKD (e.g., GFR (45 ml (min))
Diuretics— aldosterone antagonists	Eplerenone Spironolactone	50–100 25–100	12	 <45 mL/min). These are preferred agents in primary aldosteronism and resistant hypertension. Spironolactone is associated with greater risk of gynecomastia and impotence as compared with eplerenone. This is common add-on therapy in resistant hypertension. Avoid use with K⁺ supplements, other K⁺- sparing diuretics, or significant renal dysfunction. Eplerenone often requires twice-daily dosing for adequate BP lowering.
Beta blockers— cardioselectiv e Beta blockers— cardioselectiv	Atenolol Betaxolol Bisoprolol Metoprolol tartrate Metoprolol succinate Nebivolol	25–100 5–20 2.5–10 100–400 50–200 5–40	12 1 1 2 1 1	 Beta blockers are not recommended as first- line agents unless the patient has IHD or HF. These are preferred in patients with bronchospastic airway disease requiring a beta blocker. Bisoprolol and metoprolol succinate are preferred in patients with HFrEF. Avoid abrupt cessation. Nebivolol induces nitric oxide-induced vasodilation. Avoid abrupt cessation.
e and vasodilatory Beta blockers— noncardiosel ective	Nadolol Propranolol IR Propranolol LA	40–120 160–480 80–320	1 2 1	 Avoid in patients with reactive airways disease. Avoid abrupt cessation.
Beta blockers— intrinsic sympathomi metic activity	Acebutolol Carteolol Penbutolol Pindolol	200-800 2.5-10 10-40 10-60	2 1 1 2	 Generally avoid, especially in patients with IHD or HF. Avoid abrupt cessation.
Beta blockers— combined alpha- and beta- receptor	Carvedilol Carvedilol phosphate Labetalol	12.5–50 20–80 200–800	2 1 2	 Carvedilol is preferred in patients with HFrEF. Avoid abrupt cessation.
Direct renin inhibitor	Aliskiren	150–300	1	 Do not use in combination with ACE inhibitors or ARBs. Aliskiren is very long acting.

				 There is an increased risk of hyperkalemia in CKD or in those on K⁺ supplements or K⁺-sparing drugs. Aliskiren may cause acute renal failure in patients with severe bilateral renal artery stenosis. Avoid in pregnancy.
Alpha-1	Doxazosin	1-8	1	These are associated with orthostatic
blockers	Prazosin	2–20	2 or 3	hypotension, especially in older adults.
	Terazosin	1–20	1 or 2	 They may be considered as second-line agent in patients with concomitant BPH.
Central	Clonidine oral	0.1-0.8	2	• These are generally reserved as last-line
alpha₁-	Clonidine patch	0.1–0.3	1 weekly	because of significant CNS adverse effects,
agonist and	Methyldopa	250-1000	2	especially in older adults.
other centrally acting drugs	Guanfacine	0.5–2	1	 Avoid abrupt discontinuation of clonidine, which may induce hypertensive crisis; clonidine must be tapered to avoid rebound hypertension.
Direct	Hydralazine	250-200	2 or 3	• These are associated with sodium and water
vasodilators	Minoxidil	5–100	1 -3	retention and reflex tachycardia; use with a
ТТ				 diuretic and beta blocker. Hydralazine is associated with drug-induced lupus-like syndrome at higher doses. Minoxidil is associated with hirsutism and requires a loop diuretic. Minoxidil can induce pericardial effusion.

*Dosages may vary from those listed in the FDA approved labeling (available at https://dailymed.nlm.nih.gov/dailymed/.

ACE indicates angiotensin-converting enzyme; ARB, angiotensin receptor blocker; BP, blood pressure; BPH, benign prostatic hyperplasia; CCB, calcium channel blocker; CKD, chronic kidney disease; CNS, central nervous system; CVD, cardiovascular disease; ER, extended release; GFR, glomerular filtration rate; HF, heart failure; HF*r*EF, heart failure with reduced ejection fraction; IHD, ischemic heart disease; IR, immediate release; LA, long-acting; and SR, sustained release.

From Chobanian et al. JNC 7. (4)

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8.1.5. BP Goal for Patients With Hypertension

	Recommendations for BP Goal for Patients With Hypertension				
Refere	References that support recommendations are summarized in Online Data Supplement 26 and				
		Systematic Review Report.			
COR	LOE	Recommendations			
	SBP:	1. For adults with confirmed hypertension and known CVD or 10-year ASCVD			
	B-R ^{SR}	event risk of 10% or higher (see Section 8.1.2), a BP target of less than 130/80			
•	DBP:	mm Hg is recommended (1-5).			
	C-EO				
	SBP:	2. For adults with confirmed hypertension, without additional markers of			
llb	B-NR	increased CVD risk, a BP target of less than 130/80 mm Hg may be			
	DBP:	reasonable (6-9).			
	C-EO				

SR indicates systematic review.

References

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8.1.6. Choice of Initial Medication

	Recommendation for Choice of Initial Medication				
Referer	References that support the recommendation are summarized in Online Data Supplement 27 and				
	Systematic Review Report.				
COR	LOE	Recommendation			
I.	A ^{SR}	1. For initiation of antihypertensive drug therapy, first-line agents include thiazide diuretics, CCBs, and ACE inhibitors or ARBs. (1, 2)			

SR indicates systematic review.

References

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8.1.6.1. Choice of Initial Monotherapy Versus Initial Combination Drug Therapy

Reco	Recommendations for Choice of Initial Monotherapy Versus Initial Combination Drug				
	Therapy*				
COR	LOE	Recommendation			
I	C-EO	1. Initiation of antihypertensive drug therapy with 2 first-line agents of different classes, either as separate agents or in a fixed-dose combination, is recommended in adults with stage 2 hypertension and an average BP more than 20/10 mm Hg above their BP target.			
lla	C-EO	2. Initiation of antihypertensive drug therapy with a single antihypertensive drug is reasonable in adults with stage 1 hypertension and BP goal <130/80 mm Hg with dosage titration and sequential addition of other agents to achieve the BP target.			

*Fixed-dose combination antihypertensive medications are listed in Online Data Supplement D.

8.2. Follow-Up of BP During Antihypertensive Drug Therapy

Appropriate follow-up and monitoring enable assessment of adherence (see Section 12.1) and response to therapy, help identify adverse responses to therapy and target organ damage, and allow assessment of progress toward treatment goals. High-quality RCTs have successfully and safely developed strategies for follow-up, monitoring, and reassessment from which recommendations can be made (Figure 4) (1, 2). A systematic approach to out-of-office BP assessment is an essential part of follow-up and monitoring of BP, to assess response to therapy; check for evidence of white coat hypertension, white coat effect, masked hypertension, or masked uncontrolled hypertension; and help achieve BP targets (see Sections 4 and 12).

References

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8.2.1. Follow-Up After Initiating Antihypertensive Drug Therapy

Re	Recommendation for Follow-Up After Initiating Antihypertensive Drug Therapy				
Refer	References that support the recommendation are summarized in Online Data Supplement 28.				
COR	LOE	Recommendation			
I	B-R	1. Adults initiating a new or adjusted drug regimen for hypertension should have a follow-up evaluation of adherence and response to treatment at monthly intervals until control is achieved (1-3).			

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References

- Ambrosius WT, Sink KM, Foy CG, et al. The design and rationale of a multicenter clinical trial comparing two strategies for control of systolic blood pressure: the Systolic Blood Pressure Intervention Trial (SPRINT). Clin Trials. 2014;11:532-46.
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8.3.2. Monitoring Strategies to Improve Control of BP in Patients on Drug Therapy for High BP

Recommendation f	Recommendation for Monitoring Strategies to Improve Control of BP in Patients on Drug				
Therapy for High BP					
References that support the recommendation are summarized in Online Data Supplement 29.					

COR	LOE	Recommendation
I	A	1. Follow-up and monitoring after initiation of drug therapy for hypertension control should include systematic strategies to help improve BP, including use of HBPM, team-based care, and telehealth strategies (1-6).

References

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9. Hypertension in Patients With Comorbidities

Certain comorbidities may affect clinical decision-making in hypertension. These include ischemic heart disease, HF with reduced ejection fraction (HFrEF), HFpEF, CKD (including renal transplantation), cerebrovascular disease, AF, PAD, DM, and metabolic syndrome (1). As noted in Section 8.1.2, this guideline generally recommends use of BP-lowering medications for secondary prevention of CVD in patients with clinical CVD (CHD, HF, and stroke) and an average BP \geq 130/80 mm Hg and for primary prevention of CVD in adults with an estimated 10-year ASCVD risk of \geq 10% and an average SBP \geq 130 mm Hg or an average DBP \geq 80 mm Hg. Although we recommend use of the ACC/AHA Pooled Cohort Equations (http://tools.acc.org/ASCVD-Risk-Estimator/) to estimate 10-year risk of ASCVD to establish the BP threshold for treatment, the vast majority of adults with a co-morbidity are likely to have a 10-year risk of ASCVD that exceeds 10%. In some

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instances, clinical trial confirmation of treatment in patients with comorbidities is limited to a target BP of 140/90 mm Hg. In addition, the selection of medications for use in treating high BP in patients with CVD is guided by their use for other compelling indications (e.g., beta blockers after MI, ACE inhibitors for HF*r*EF), as discussed in specific guidelines for the clinical condition (2-4). The present guideline does not address the recommendations for treatment of hypertension occurring with acute coronary syndromes.

References

- Aronow WS, Fleg JL, Pepine CJ, et al. ACCF/AHA 2011 expert consensus document on hypertension in the elderly: a report of the American College of Cardiology Foundation Task Force on Clinical Expert Consensus documents developed in collaboration with the American Academy of Neurology, American Geriatrics Society, American Society for Preventive Cardiology, American Society of Hypertension, American Society of Nephrology, Association of Black Cardiologists, and European Society of Hypertension. Circulation. 2011;123:2434-506.
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- 3. Fihn SD, Blankenship JC, Alexander KP, et al. 2014 ACC/AHA/AATS/PCNA/SCAI/STS focused update of the guideline for the diagnosis and management of patients with stable ischemic heart disease: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines, and the American Association for Thoracic Surgery, Preventive Cardiovascular Nurses Association, Society for Cardiovascular Angiography and Interventions, and Society of Thoracic Surgeons. Circulation. 2014;130:1749-67.
- 4. Gerhard-Herman MD, Gornik HL, Barrett C, et al. 2016 AHA/ACC guideline on the management of patients with lower extremity peripheral artery disease: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines. Circulation. 2017;135:e726-79.

Recom	Recommendations for Treatment of Hypertension in Patients With Stable Ischemic Heart					
	Disease (SIHD)					
Refere	References that support recommendations are summarized in Online Data Supplements 30-32.					
COR	LOE	Recommendations				
	SBP:	1. In adults with SIHD and hypertension, a BP target of less than 130/80 mm				
	B-R	Hg is recommended (1-5).				
I.	DBP:					
	C-EO					
	SBP:	2. Adults with SIHD and hypertension (BP ≥130/80 mm Hg) should be treated				
	B-R	with medications (e.g., GDMT (6) beta blockers, ACE inhibitors, or ARBs) for				
		compelling indications (e.g., previous MI, stable angina) as first-line therapy,				
•	DBP:	with the addition of other drugs (e.g., dihydropyridine CCBs, thiazide				
	C-EO	diuretics, and/or mineralocorticoid receptor antagonists) as needed to				
		further control hypertension (7-10).				
		3. In adults with SIHD with angina and persistent uncontrolled hypertension,				
1	B-NR	the addition of dihydropyridine CCBs to GDMT (6) beta blockers is				
		recommended (8, 11, 12).				
		4. In adults who have had a MI or acute coronary syndrome, it is reasonable to				
lla	B-NR	continue GDMT (6) beta blockers beyond 3 years as long-term therapy for				
		hypertension (13, 14).				
		5. Beta blockers and/or CCBs might be considered to control hypertension in				
llb	C-EO	patients with CAD (without HFrEF) who had an MI more than 3 years ago				
		and have angina.				

9.1. Stable Ischemic Heart Disease

Figure 5 is an algorithm on management of hypertension in patients with SIHD.

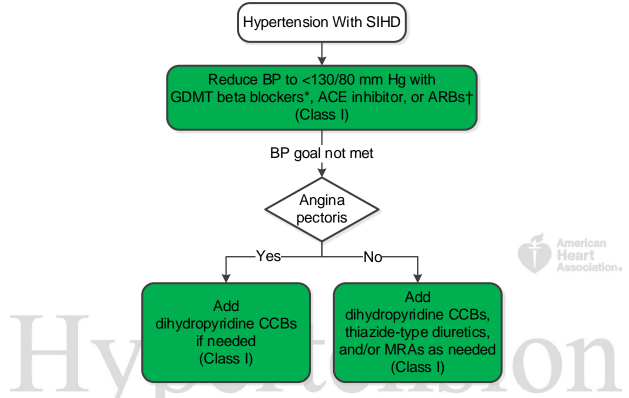


Figure 5. Management of Hypertension in Patients With SIHD

Colors correspond to Class of Recommendation in Table 1.

*GDMT beta blockers for BP control or relief of angina include carvedilol, metoprolol tartrate, metoprolol succinate, nadolol, bisoprolol, propranolol, and timolol. Avoid beta blockers with intrinsic sympathomimetic activity. The beta blocker atenolol should not be used because it is less effective than placebo in reducing cardiovascular events. †If needed for BP control.

ACE indicates angiotensin-converting enzyme; ARB, angiotensin receptor blocker; BP, blood pressure; CCB, calcium channel blocker; GDMT, guideline-directed management and therapy; and SIHD, stable ischemic heart disease.

- 1. Wright JT Jr, Williamson JD, Whelton PK, et al. A randomized trial of intensive versus standard blood-pressure control. SPRINT Research Group. N Engl J Med. 2015;373:2103-16.
- 2. Bundy JD, Li C, Stuchlik P, et al. Systolic blood pressure reduction and risk of cardiovascular disease and mortality: a systematic review and network meta-analysis. JAMA Cardiol. 2017;2:775-81.
- 3. Leenen FH, Nwachuku CE, Black HR, et al. Clinical events in high-risk hypertensive patients randomly assigned to calcium channel blocker versus angiotensin-converting enzyme inhibitor in the antihypertensive and lipid-lowering treatment to prevent heart attack trial. Hypertension 2006;48:374-84.
- 4. Zanchetti A, Julius S, Kjeldsen S, et al. Outcomes in subgroups of hypertensive patients treated with regimens based on valsartan and amlodipine: An analysis of findings from the VALUE trial. J Hypertens. 2006;24:2163-8.
- 5. Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial Collaborative Research Group. Diuretic versus alpha-blocker as first-step antihypertensive therapy: final results from the Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT). Hypertension. 2003;42:239-46.

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- 6. Fihn SD, Blankenship JC, Alexander KP, et al. 2014 ACC/AHA/AATS/PCNA/SCAI/STS focused update of the Guideline for the diagnosis and management of patients with stable ischemic heart disease: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines, and the American Association for Thoracic Surgery, Preventive Cardiovascular Nurses Association, Society for Cardiovascular Angiography and Interventions, and Society of Thoracic Surgeons. Circulation. 2014;130:1749-67.
- Fox KM, EURopean trial On reduction of cardiac events with Perindopril in stable coronary Artery disease Investigators. Efficacy of perindopril in reduction of cardiovascular events among patients with stable coronary artery disease: randomised, double-blind, placebo-controlled, multicentre trial (the EUROPA study). Lancet. 2003;362:782-8.
- Law MR, Morris JK, Wald NJ. Use of blood pressure lowering drugs in the prevention of cardiovascular disease: meta-analysis of 147 randomised trials in the context of expectations from prospective epidemiological studies. BMJ. 2009;338:b1665.
- 9. Pfeffer MA, Braunwald E, Moye LA, et al. Effect of captopril on mortality and morbidity in patients with left ventricular dysfunction after myocardial infarction. Results of the survival and ventricular enlargement trial. The SAVE Investigators. N Engl J Med. 1992;327:669-77.
- Yusuf S, Sleight P, Pogue J, et al. Effects of an angiotensin-converting-enzyme inhibitor, ramipril, on cardiovascular events in high-risk patients. The Heart Outcomes Prevention Evaluation Study Investigators. N Engl J Med. 2000;342:145-53.
- 11. Leon MB, Rosing DR, Bonow RO, et al. Clinical efficacy of verapamil alone and combined with propranolol in treating patients with chronic stable angina pectoris. Am J Cardiol. 1981;48:131-9.
- 12. Staessen JA, Fagard R, Thijs L, et al. Randomised double-blind comparison of placebo and active treatment for older patients with isolated systolic hypertension. The Systolic Hypertension in Europe (Syst-Eur) Trial Investigators. Lancet. 1997;350:757-64.
- 13. Freemantle N, Cleland J, Young P, et al. beta Blockade after myocardial infarction: systematic review and meta regression analysis. BMJ. 1999;318:1730-7.
- de Peuter OR, Lussana F, Peters RJG, et al. A systematic review of selective and non-selective beta blockers for prevention of vascular events in patients with acute coronary syndrome or heart failure. Neth J Med. 2009;67:284-94.

9.2. Heart Failure

Recommendation for Prevention of HF in Adults With Hypertension				
Refer	References that support the recommendation are summarized in Online Data Supplement 33.			
COR	COR LOE Recommendation			
	SBP:	1. In adults at increased risk of HF, the optimal BP in those with hypertension		
I	B-R	should be less than 130/80 mm Hg (1-3).		
	DBP:			
	C-EO			

- 1. Lv J, Ehteshami P, Sarnak MJ, et al. Effects of intensive blood pressure lowering on the progression of chronic kidney disease: a systematic review and meta-analysis. CMAJ. 2013;185:949-57.
- Thomopoulos C, Parati G, Zanchetti A. Effects of blood pressure lowering on outcome incidence in hypertension: 7. Effects of more vs. less intensive blood pressure lowering and different achieved blood pressure levels - updated overview and meta-analyses of randomized trials. J Hypertens. 2016;34:613-22.
- 3. Xie X, Atkins E, Lv J, et al. Effects of intensive blood pressure lowering on cardiovascular and renal outcomes: updated systematic review and meta-analysis. Lancet. 2015;387:435-43.

Recommendations for Treatment of Hypertension in Patients With HFrEF			
References that support recommendations are summarized in Online Data Supplement 34.			
COR	LOE	Recommendation	
I	C-EO	1. Adults with HFrEF and hypertension should be prescribed GDMT (2) titrated to attain a BP of less than 130/80 mm Hg.	
III: No Benefit	B-R	2. Nondihydropyridine CCBs are not recommended in the treatment of hypertension in adults with HF <i>r</i> EF (1).	

9.2.1. Heart Failure With Reduced Ejection Fraction

Reference

1. Goldstein RE, Boccuzzi SJ, Cruess D, et al. Diltiazem increases late-onset congestive heart failure in postinfarction patients with early reduction in ejection fraction. The Adverse Experience Committee; and the Multicenter Diltiazem Postinfarction Research Group. Circulation. 1991;83:52-60.

American

9.2.2. Heart Failure With Preserved Ejection Fraction

COR LOE Recommendations I C-EO 1. In adults with HFpEF who present with symptoms of volume over diuretics should be prescribed to control hypertension. I C-EO 2. Adults with HFpEF and persistent hypertension after management		Recommendations for Treatment of Hypertension in Patients With HFpEF			
IC-EO1. In adults with HFpEF who present with symptoms of volume over diuretics should be prescribed to control hypertension.2.Adults with HFpEF and persistent hypertension after management	Referen	References that support recommendations are summarized in Online Data Supplements 35 and 36.			
C-EO diuretics should be prescribed to control hypertension. 2. Adults with HFpEF and persistent hypertension after management	COR LOE Recommendations		Recommendations		
	I	C-EO	1. In adults with HFpEF who present with symptoms of volume overload, diuretics should be prescribed to control hypertension.		
blockers titrated to attain SBP of less than 130 mm Hg (1-6).	I	C-LD	2. Adults with HFpEF and persistent hypertension after management of volume overload should be prescribed ACE inhibitors or ARBs and beta blockers titrated to attain SBP of less than 130 mm Hg (1-6).		

- Pfeffer MA, Claggett B, Assmann SF, et al. Regional variation in patients and outcomes in the Treatment of Preserved Cardiac Function Heart Failure With an Aldosterone Antagonist (TOPCAT) Trial. Circulation. 2015;131:34-42.
- Yancy CW, Jessup M, Bozkurt B, et al. 2013 ACCF/AHA guideline for the management of heart failure: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. Circulation. 2013;128:e240-327.
- Aronow WS, Ahn C, Kronzon I. Effect of propranolol versus no propranolol on total mortality plus nonfatal myocardial infarction in older patients with prior myocardial infarction, congestive heart failure, and left ventricular ejection fraction > or = 40% treated with diuretics plus angiotensin-converting enzyme inhibitors. Am J Cardiol. 1997;80:207-9.
- 4. van Veldhuisen DJ, Cohen-Solal A, Bohm M, et al. Beta-blockade with nebivolol in elderly heart failure patients with impaired and preserved left ventricular ejection fraction: Data From SENIORS (Study of Effects of Nebivolol Intervention on Outcomes and Rehospitalization in Seniors With Heart Failure). J Am Coll Cardiol. 2009;53:2150-8.
- 5. Yusuf S, Pfeffer MA, Swedberg K, et al. Effects of candesartan in patients with chronic heart failure and preserved left-ventricular ejection fraction: the CHARM-Preserved Trial. Lancet. 2003;362:777-81.
- 6. Massie BM, Carson PE, McMurray JJ, et al. Irbesartan in patients with heart failure and preserved ejection fraction. N Engl J Med. 2008;359:2456-67.

9.3. Chronic Kidney Disease

Recommendations for Treatment of Hypertension in Patients With CKD References that support recommendations are summarized in Online Data Supplements 37 and 38			
and Systematic Review Report.			
LOE	Recommendations		
SBP:	1. Adults with hypertension and CKD should be treated to a BP goal of less than 130/80 mm Hg (1-6).		
DBP:	130/80 mm ng (1-0).		
C-EO			
B-R	 In adults with hypertension and CKD (stage 3 or higher or stage 1 or 2 with albuminuria [≥300 mg/d, or ≥300 mg/g albumin-to-creatinine ratio or the equivalent in the first morning void]), treatment with an ACE inhibitor is reasonable to slow kidney disease progression (3, 7-12). 		
C-EO	3. In adults with hypertension and CKD (stage 3 or higher or stage 1 or 2 with albuminuria [≥300 mg/d, or ≥300 mg/g albumin-to-creatinine ratio in the first morning void]) (7, 8), treatment with an ARB may be reasonable if an ACE inhibitor is not tolerated.		
	that su LOE SBP: 3-R ^{SR} DBP: C-EO B-R		

Figure 6 is an algorithm on management of hypertension in patients with CKD.



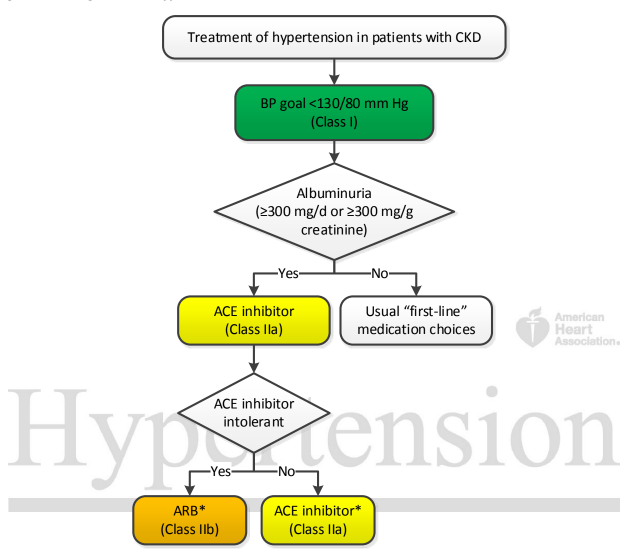


Figure 6. Management of Hypertension in Patients With CKD

Colors correspond to Class of Recommendation in Table 1.

*CKD stage 3 or higher or stage 1 or 2 with albuminuria ≥300 mg/d or ≥300 mg/g creatinine. ACE indicates angiotensin-converting enzyme; ARB, angiotensin receptor blocker; BP blood pressure; and CKD, chronic kidney disease.

- Klahr S, Levey AS, Beck GJ, et al. The effects of dietary protein restriction and blood-pressure control on the progression of chronic renal disease. Modification of Diet in Renal Disease Study Group. N Engl J Med. 1994;330:877-84.
- 2. Ruggenenti P, Perna A, Loriga G, et al. Blood-pressure control for renoprotection in patients with non-diabetic chronic renal disease (REIN-2): multicentre, randomised controlled trial. Lancet. 2005;365:939-46.
- 3. Wright JT Jr, Bakris G, Greene T, et al. Effect of blood pressure lowering and antihypertensive drug class on progression of hypertensive kidney disease: results from the AASK trial. JAMA. 2002;288:2421-31.
- 4. Upadhyay A, Earley A, Haynes SM, et al. Systematic review: blood pressure target in chronic kidney disease and proteinuria as an effect modifier. Ann Intern Med. 2011;154:541-8.
- 5. Lv J, Ehteshami P, Sarnak MJ, et al. Effects of intensive blood pressure lowering on the progression of chronic kidney disease: a systematic review and meta-analysis. CMAJ. 2013;185:949-57.

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- 6. Jafar TH, Stark PC, Schmid CH, et al. Progression of chronic kidney disease: the role of blood pressure control, proteinuria, and angiotensin-converting enzyme inhibition: a patient-level meta-analysis. Ann Intern Med. 2003;139:244-52.
- Lambers Heerspink HJ, Brantsma AH, de Zeeuw D, et al. Albuminuria assessed from first-morning-void urine samples versus 24-hour urine collections as a predictor of cardiovascular morbidity and mortality. Am J Epidemiol. 2008;168:897-905.
- 8. Lambers Heerspink HJ, Gansevoort RT, Brenner BM, et al. Comparison of different measures of urinary protein excretion for prediction of renal events. J Am Soc Nephrol. 2010;21:1355-60.
- 9. Contreras G, Greene T, Agodoa LY, et al. Blood pressure control, drug therapy, and kidney disease. Hypertension. 2005;46:44-50.
- 10. Esnault VL, Brown EA, Apetrei E, et al. The effects of amlodipine and enalapril on renal function in adults with hypertension and nondiabetic nephropathies: a 3-year, randomized, multicenter, double-blind, placebo-controlled study. Clin Ther. 2008;30:482-98.
- 11. Marin R, Ruilope LM, Aljama P, et al. A random comparison of fosinopril and nifedipine GITS in patients with primary renal disease. J Hypertens. 2001;19:1871-6.
- 12. Giatras I, Lau J, Levey AS. Effect of angiotensin-converting enzyme inhibitors on the progression of nondiabetic renal disease: a meta-analysis of randomized trials. Angiotensin-Converting-Enzyme Inhibition and Progressive Renal Disease Study Group. Ann Intern Med. 1997;127:337-45.

9.3.1. Hypertension After Renal Transplantation

Re	Recommendations for Treatment of Hypertension After Renal Transplantation				
Referen	References that support recommendations are summarized in Online Data Supplements 39 and 40.				
COR	LOE	Recommendations			
	SBP:	1. After kidney transplantation, it is reasonable to treat patients with			
lle	B-NR	hypertension to a BP goal of less than 130/80 mm Hg (1).			
lla	DBP:				
	C-EO				
		2. After kidney transplantation, it is reasonable to treat patients with			
lla	B-R	hypertension with a calcium antagonist on the basis of improved GFR and			
		kidney survival (2).			

Heart

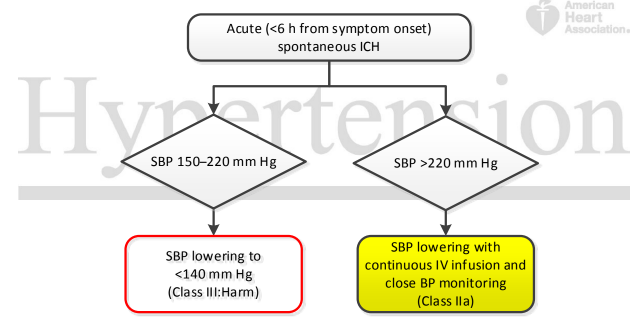
- 1. Wright JT Jr, Williamson JD, Whelton PK, et al. A randomized trial of intensive versus standard blood-pressure control. SPRINT Research Group. N Engl J Med. 2015;373:2103-16.
- 2. Cross NB, Webster AC, Masson P, et al. Antihypertensive treatment for kidney transplant recipients. Cochrane Database Syst Rev. 2009;CD003598.

9.4. Cerebrovascular Disease

9.4.1. Acute Intracerebral Hemorrhage

Recomm	Recommendations for Management of Hypertension in Patients With Acute Intracerebral			
	Hemorrhage (ICH)			
Refe	erences the	at support recommendations are summarized in Online Data Supplement 41.		
COR	LOE	Recommendations		
lla	C-EO	1. In adults with ICH who present with SBP greater than 220 mm Hg, it is reasonable to use continuous intravenous drug infusion (Table 19) and close BP monitoring to lower SBP.		
III: Harm	А	2. Immediate lowering of SBP (Table 19) to less than 140 mm Hg in adults with spontaneous ICH who present within 6 hours of the acute event and have an SBP between 150 mm Hg and 220 mm Hg is not of benefit to reduce death or severe disability and can be potentially harmful (1, 2).		

Figure 7. Management of Hypertension in Patients With Acute ICH



Colors correspond to Class of Recommendation in Table 1.

BP indicates blood pressure; ICH, intracerebral hemorrhage; IV, intravenous; and SBP, systolic blood pressure.

- 1. Anderson CS, Heeley E, Huang Y, et al. Rapid blood-pressure lowering in patients with acute intracerebral hemorrhage. N Engl J Med. 2013;368:2355-65.
- 2. Qureshi AI, Palesch YY, Barsan WG, et al. Intensive blood-pressure lowering in patients with acute cerebral hemorrhage. N Engl J Med. 2016;375:1033-43.

9.4.2. Acute Ischemic Stroke

	Recommendations for Management of Hypertension in Patients With Acute Ischemic Stroke			
Refe	erences th	at support recommendations are summarized in Online Data Supplement 42.		
COR	LOE	Recommendations		
I	B-NR	1. Adults with acute ischemic stroke and elevated BP who are eligible for treatment with intravenous tissue plasminogen activator should have their BP slowly lowered to less than 185/110 mm Hg before thrombolytic therapy is initiated (1, 2).		
I	B-NR	 In adults with an acute ischemic stroke, BP should be less than 185/110 mm Hg before administration of intravenous tissue plasminogen activator and should be maintained below 180/105 mm Hg for at least the first 24 hours after initiating drug therapy (3). 		
lla	B-NR	3. Starting or restarting antihypertensive therapy during hospitalization in patients with BP greater than 140/90 mm Hg who are neurologically stable is safe and reasonable to improve long-term BP control, unless contraindicated (4, 5).		
lib	C-EO	4. In patients with BP of 220/120 mm Hg or higher who did not receive intravenous alteplase or endovascular treatment and have no comorbid conditions requiring acute antihypertensive treatment, the benefit or initiating or reinitiating treatment of hypertension within the first 48 to 72 hours is uncertain. It might be reasonable to lower BP by 15% during the first 24 hours after onset of stroke.		
III: No Benefit	A	5. In patients with BP less than 220/120 mm Hg who did not receive intravenous thrombolysis or endovascular treatment and do not have a comorbid condition requiring acute antihypertensive treatment, initiating or reinitiating treatment of hypertension within the first 48 to 72 hours after an acute ischemic stroke is not effective to prevent death or dependency (4- 9).		

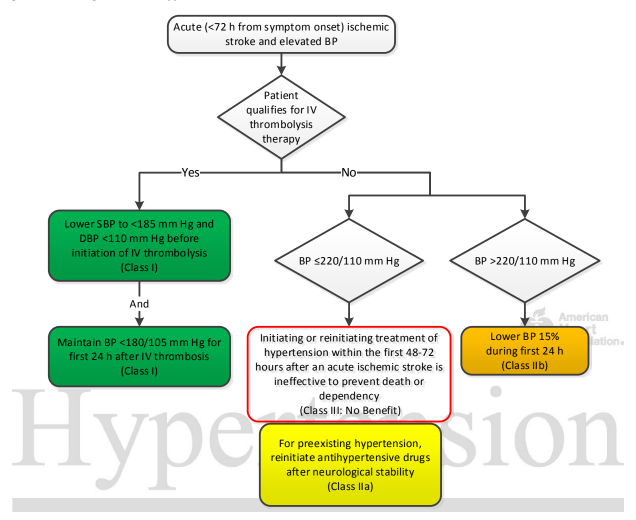


Figure 8. Management of Hypertension in Patients With Acute Ischemic Stroke

Colors correspond to Class of Recommendation in Table 1.

BP indicates blood pressure; DBP, diastolic blood pressure; IV, intravenous; and SBP, systolic blood pressure.

- 1. National Institute of Neurological Disorders and Stroke rt-PA Stroke Study Group. Tissue plasminogen activator for acute ischemic stroke. N Engl J Med. 1995;333:1581-7.
- 2. Hacke W, Kaste M, Bluhmki E, et al. Thrombolysis with alteplase 3 to 4.5 hours after acute ischemic stroke. N Engl J Med. 2008;359:1317-29.
- 3. Ahmed N, Wahlgren N, Brainin M, et al. Relationship of blood pressure, antihypertensive therapy, and outcome in ischemic stroke treated with intravenous thrombolysis: retrospective analysis from Safe Implementation of Thrombolysis in Stroke-International Stroke Thrombolysis Register (SITS-ISTR). Stroke. 2009;40:2442-9.
- 4. Robinson TG, Potter JF, Ford GA, et al. Effects of antihypertensive treatment after acute stroke in the Continue or Stop Post-Stroke Antihypertensives Collaborative Study (COSSACS): a prospective, randomised, open, blinded-endpoint trial. Lancet Neurol. 2010;9:767-75.
- 5. He J, Zhang Y, Xu T, et al. Effects of immediate blood pressure reduction on death and major disability in patients with acute ischemic stroke: the CATIS randomized clinical trial. JAMA. 2014;311:479-89.
- 6. Wang H, Tang Y, Rong X, et al. Effects of early blood pressure lowering on early and long-term outcomes after acute stroke: an updated meta-analysis. PLoS ONE. 2014;9:e97917.

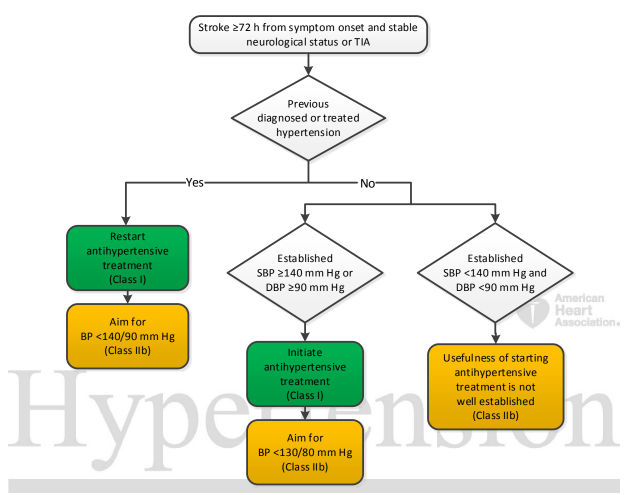
- Zhao R, Liu F-D, Wang S, et al. Blood pressure reduction in the acute phase of an ischemic stroke does not improve short- or long-term dependency or mortality: a meta-analysis of current literature. Medicine (Baltimore). 2015;94:e896.
- 8. Bath PM, Krishnan K. Interventions for deliberately altering blood pressure in acute stroke. Cochrane Database Syst Rev. 2014;10:CD000039.
- 9. Sandset EC, Bath PMW, Boysen G, et al. The angiotensin-receptor blocker candesartan for treatment of acute stroke (SCAST): a randomised, placebo-controlled, double-blind trial. Lancet. 2011;377:741-50.

9.4.3. Secondary Stroke Prevention

	Recommendations for Treatment of Hypertension for Secondary Stroke Prevention References that support recommendations are summarized in Online Data Supplements 43 and 44.		
COR	LOE	Recommendations	
I	A	1. Adults with previously treated hypertension who experience a stroke or transient ischemic attack (TIA) should be restarted on antihypertensive treatment after the first few days of the index event to reduce the risk of recurrent stroke and other vascular events (1-3).	
I	A	2. For adults who experience a stroke or TIA, treatment with a thiazide diuretic, ACE inhibitor, or ARB, or combination treatment consisting of a thiazide diuretic plus ACE inhibitor, is useful (1, 3-5).	
I	B-R	3. Adults not previously treated for hypertension who experience a stroke of TIA and have an established BP of 140/90 mm Hg or higher should be prescribed antihypertensive treatment a few days after the index event to reduce the risk of recurrent stroke and other vascular events (1-3).	
I	B-NR	 For adults who experience a stroke or TIA, selection of specific drugs should be individualized on the basis of patient comorbidities and agen pharmacological class (6). 	
llb	B-R	5. For adults who experience a stroke or TIA, a BP goal of less than 130/80 mm Hg may be reasonable (6, 7).	
llb	B-R	 For adults with a lacunar stroke, a target SBP goal of less than 130 mm H_g may be reasonable (8). 	
llb	C-LD	 In adults previously untreated for hypertension who experience an ischemic stroke or TIA and have a SBP less than 140 mm Hg and a DBP less than 90 mm Hg, the usefulness of initiating antihypertensive treatment is not wel established (9). 	

Figure 9 is an algorithm on management of hypertension in patients with a previous history of stroke (secondary stroke prevention).

Figure 9. Management of Hypertension in Patients With a Previous History of Stroke (Secondary Stroke Prevention)



Colors correspond to Class of Recommendation in Table 1.

DBP indicates diastolic blood pressure; SBP, systolic blood pressure; and TIA, transient ischemic attack.

- 1. Liu L, Wang Z, Gong L, et al. Blood pressure reduction for the secondary prevention of stroke: a Chinese trial and a systematic review of the literature. Hypertens Res. 2009;32:1032-40.
- 2. Lakhan SE, Sapko MT. Blood pressure lowering treatment for preventing stroke recurrence: a systematic review and meta-analysis. Int Arch Med. 2009;2:30.
- 3. PROGRESS Collaborative Group. Randomised trial of a perindopril-based blood-pressure-lowering regimen among 6,105 individuals with previous stroke or transient ischaemic attack. Lancet. 2001;358:1033-41.
- 4. PATS Collaborating Group. Post-stroke antihypertensive treatment study. A preliminary result. Chin Med J. 1995;108:710-7.
- 5. Lee M, Saver JL, Hong K-S, et al. Renin-angiotensin system modulators modestly reduce vascular risk in persons with prior stroke. Stroke. 2012;43:113-9.
- 6. Wang W-T, You L-K, Chiang C-E, et al. Comparative effectiveness of blood pressure-lowering drugs in patients who have already suffered from stroke: traditional and Bayesian network meta-analysis of randomized trials. Medicine (Baltimore). 2016;95:e3302.
- 7. Katsanos AH, Filippatou A, Manios E, et al. Blood pressure reduction and secondary stroke prevention: a systematic review and metaregression analysis of randomized clinical trials. Hypertension. 2017;69:171-9.

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- 8. Benavente OR, Coffey CS, Conwit R, et al. Blood-pressure targets in patients with recent lacunar stroke: the SPS3 randomised trial. Lancet. 2013;382:507-15.
- 9. Arima H, Chalmers J, Woodward M, et al. Lower target blood pressures are safe and effective for the prevention of recurrent stroke: the PROGRESS trial. J Hypertens. 2006;24:1201-8.

9.5. Peripheral Arterial Disease

Recommendation for Treatment of Hypertension in Patients With PAD			
Refer	References that support the recommendation are summarized in Online Data Supplement 45.		
COR	LOE	Recommendation	
I.	B-NR	1. Adults with hypertension and PAD should be treated similarly to patients with hypertension without PAD (1-4).	

References

- 1. Ostergren J, Sleight P, Dagenais G, et al. Impact of ramipril in patients with evidence of clinical or subclinical peripheral arterial disease. Eur Heart J. 2004;25:17-24.
- 2. Thompson AM, Hu T, Eshelbrenner CL, et al. Antihypertensive treatment and secondary prevention of cardiovascular disease events among persons without hypertension: a meta-analysis. JAMA. 2011;305:913-22.
- Bavry AA, Anderson RD, Gong Y, et al. Outcomes among hypertensive patients with concomitant peripheral and coronary artery disease: findings from the INternational VErapamil-SR/Trandolapril STudy. Hypertension. 2010;55:48-53.
- 4. Zanchetti A, Julius S, Kjeldsen S, et al. Outcomes in subgroups of hypertensive patients treated with regimens based on valsartan and amlodipine: an analysis of findings from the VALUE trial. J Hypertens. 2006;24:2163-8.

	Recommendations for Treatment of Hypertension in Patients With DM		
Referen	ces that su	upport recommendations are summarized in Online Data Supplements 46 and 47	
		and Systematic Review Report.	
COR	LOE	Recommendations	
	SBP:	1. In adults with DM and hypertension, antihypertensive drug treatment	
	B-R ^{SR}	should be initiated at a BP of 130/80 mm Hg or higher with a treatment goal	
1	DBP:	of less than 130/80 mm Hg (1-8).	
	C-EO		
I.	A ^{sr}	2. In adults with DM and hypertension, all first-line classes of antihypertensive agents (i.e., diuretics, ACE inhibitors, ARBs, and CCBs) are useful and effective (1, 9, 10).	
llb	B-NR	3. In adults with DM and hypertension, ACE inhibitors or ARBs may be considered in the presence of albuminuria (11, 12).	

9.6. Diabetes Mellitus

SR indicates systematic review.

- 1. Emdin CA, Rahimi K, Neal B, et al. Blood pressure lowering in type 2 diabetes: a systematic review and metaanalysis. JAMA. 2015;313:603-15.
- 2. Arguedas JA, Leiva V, Wright JM. Blood pressure targets for hypertension in people with diabetes mellitus. Cochrane Database Syst Rev. 2013;10:CD008277.

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- 3. Cushman WC, Evans GW, Byington RP, et al. Effects of intensive blood-pressure control in type 2 diabetes mellitus. ACCORD Study. N Engl J Med. 2010;362:1575-85.
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- 11. Palmer SC, Mavridis D, Navarese E, et al. Comparative efficacy and safety of blood pressure-lowering agents in adults with diabetes and kidney disease: a network meta-analysis. Lancet. 2015;385:2047-56.
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9.7. Metabolic Syndrome

Metabolic syndrome is a state of metabolic dysregulation characterized by visceral fat accumulation, insulin resistance, hyperinsulinemia, and hyperlipidemia, as well as predisposition to type 2 DM, hypertension, and atherosclerotic CVD (1-3). According to data from the NHANES III and NHANES 1999–2006 (1, 4), the prevalence of metabolic syndrome in the United States was 34.2% in 2006 and has likely increased substantially since that time. The metabolic syndrome is linked to several other disorders, including nonalcoholic steatohepatitis, polycystic ovary syndrome, certain cancers, CKD, Alzheimer's disease, Cushing's syndrome, lipodystrophy, and hyperalimentation (5, 6).

Lifestyle modification, with an emphasis on improving insulin sensitivity by means of dietary modification, weight reduction, and exercise, is the foundation of treatment of the metabolic syndrome. The optimal antihypertensive drug therapy for patients with hypertension in the setting of the metabolic syndrome has not been clearly defined (1). Although caution exists with regard to the use of thiazide diuretics in this population because of their ability to increase insulin resistance, dyslipidemia, and hyperuricemia and to accelerate conversion to overt DM, no data are currently available demonstrating deterioration in cardiovascular or renal outcomes in patients treated with these agents (1). Indeed, as shown in follow-up of ALLHAT, chlorthalidone use was associated with only a small increase in fasting glucose levels (1.5–4.0 mg/dL), and this increase did not translate into increased CVD risk at a later date (7-10). In addition, in post hoc analysis of the nearly two thirds of participants in ALLHAT that met criteria for the metabolic syndrome, chlorthalidone was unsurpassed in reducing CVD and renal outcomes compared with lisinopril, amlodipine, or doxazosin (9, 11). Similarly, high-dose ARB therapy reduces arterial stiffness in patients with hypertension with the metabolic syndrome, but no outcomes data are available from patients in which this form of treatment was used (12). Use of traditional beta blockers may lead to dyslipidemia or deterioration of glucose tolerance, and ability to lose weight (2). In several large clinical trials, the risk of developing DM as a result of traditional betablocker therapy was 15% to 29% (2). However, the newer vasodilating beta blockers (e.g., labetalol, carvedilol, nebivolol) have shown neutral or favorable effects on metabolic profiles compared with the traditional beta

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blockers (13). Trials using vasodilator beta blockers have not been performed to demonstrate effects on CVD outcomes.

References

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- 5. Chen J, Muntner P, Hamm LL, et al. The metabolic syndrome and chronic kidney disease in U.S. adults. Ann Intern Med. 2004;140:167-74.
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- 9. Wright JT Jr, Harris-Haywood S, Pressel S, et al. Clinical outcomes by race in hypertensive patients with and without the metabolic syndrome: Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT). Arch Intern Med. 2008;168:207-17.
- 10. Wright JT Jr, Probstfield JL, Cushman WC, et al. ALLHAT findings revisited in the context of subsequent analyses, other trials, and meta-analyses. Arch Intern Med. 2009;169:832-42.
- 11. Black HR, Davis B, Barzilay J, et al. Metabolic and clinical outcomes in nondiabetic individuals with the metabolic syndrome assigned to chlorthalidone, amlodipine, or lisinopril as initial treatment for hypertension: a report from the Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT). Diabetes Care. 2008;31:353-60.
- 12. Laurent S, Boutouyrie P, Vascular Mechanism Collaboration. Dose-dependent arterial destiffening and inward remodeling after olmesartan in hypertensives with metabolic syndrome. Hypertension. 2014;64:709-16.
- 13. Reisin E, Owen J. Treatment: special conditions. Metabolic syndrome: obesity and the hypertension connection. J Am Soc Hypertens. 2015;9:156-9.

9.8. Atrial Fibrillation

	Recommendation for Treatment of Hypertension in Patients With AF			
Refer	References that support the recommendation are summarized in Online Data Supplement 48.			
COR	LOE	Recommendation		
lla	B-R	1. Treatment of hypertension with an ARB can be useful for prevention of recurrence of AF (1, 2).		

- 1. Healey JS, Baranchuk A, Crystal E, et al. Prevention of atrial fibrillation with angiotensin-converting enzyme inhibitors and angiotensin receptor blockers: a meta-analysis. J Am Coll Cardiol. 2005;45:1832-9.
- 2. Zhao D, Wang Z-M, Wang L-S. Prevention of atrial fibrillation with renin-angiotensin system inhibitors on essential hypertensive patients: a meta-analysis of randomized controlled trials. J Biomed Res. 2015;29:475-85.

9.9. Valvular Heart Disease

	Recommendations for Treatment of Hypertension in Patients With Valvular Heart Disease		
Reference	ces that su	pport recommendations are summarized in Online Data Supplements 49 and 50.	
COR	LOE	Recommendation	
I	B-NR	1. In adults with asymptomatic aortic stenosis, hypertension should be treated with pharmacotherapy, starting at a low dose and gradually titrating upward as needed (1-4).	
lla	C-LD	2. In patients with chronic aortic insufficiency, treatment of systolic hypertension with agents that do not slow the heart rate (i.e., avoid beta blockers) is reasonable (5, 6).	

References

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- 2. Eleid MF, Nishimura RA, Sorajja P, et al. Systemic hypertension in low-gradient severe aortic stenosis with preserved ejection fraction. Circulation. 2013;128:1349-53.
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- 5. Scognamiglio R, Rahimtoola SH, Fasoli G, et al. Nifedipine in asymptomatic patients with severe aortic regurgitation and normal left ventricular function. N Engl J Med. 1994;331:689-94.
- 6. Evangelista A, Tornos P, Sambola A, et al. Long-term vasodilator therapy in patients with severe aortic regurgitation. N Engl J Med. 2005;353:1342-9.

9.10. Aortic Disease

Recommendation for Management of Hypertension in Patients With Aortic Disease				
COR	LOE	Recommendation		
I	C-EO	1. Beta blockers are recommended as the preferred antihypertensive agents in patients with hypertension and thoracic aortic disease (1, 2).		

- 1. Genoni M, Paul M, Jenni R, et al. Chronic beta-blocker therapy improves outcome and reduces treatment costs in chronic type B aortic dissection. Eur J Cardiothorac Surg. 2001;19:606-10.
- 2. Suzuki T, Isselbacher EM, Nienaber CA, et al. Type-selective benefits of medications in treatment of acute aortic dissection (from the International Registry of Acute Aortic Dissection [IRAD]). Am J Cardiol. 2012;109:122-7.

10. Special Patient Groups

Special attention is needed for specific patient subgroups.

10.1.1 Racial and Ethnic Differences in Treatment

	Recommendations for Race and Ethnicity				
Refe	References that support recommendations are summarized in Online Data Supplement 51.				
COR	LOE	Recommendations			
I	B-R	1. In black adults with hypertension but without HF or CKD, including those with DM, initial antihypertensive treatment should include a thiazide-type diuretic or CCB (1-4).			
I	C-LD	2. Two or more antihypertensive medications are recommended to achieve a BP target of less than 130/80 mm Hg in most adults with hypertension, especially in black adults with hypertension (5-7).			

References

- 1. Leenen FH, Nwachuku CE, Black HR, et al. Clinical events in high-risk hypertensive patients randomly assigned to calcium channel blocker versus angiotensin-converting enzyme inhibitor in the antihypertensive and lipid-lowering treatment to prevent heart attack trial. Hypertension 2006;48:374-84.
- 2. Wright JT Jr, Probstfield JL, Cushman WC, et al. ALLHAT findings revisited in the context of subsequent analyses, other trials, and meta-analyses. Arch Intern Med. 2009;169:832-42.
- 3. Wright JT Jr, Dunn JK, Cutler JA, et al. Outcomes in hypertensive black and nonblack patients treated with chlorthalidone, amlodipine, and lisinopril. JAMA. 2005;293:1595-608.
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10.2. Sex-Related Issues

The prevalence of hypertension is lower in women than in men until about the fifth decade but is higher later in life (1). Other than special recommendations for management of hypertension during pregnancy, there is no evidence that the BP threshold for initiating drug treatment, the treatment target, the choice of initial antihypertensive medication, or the combination of medications for lowering BP differs for women versus men (2, 3).

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- 3. Turnbull F, Woodward M, Neal B, et al. Do men and women respond differently to blood pressure-lowering treatment? Results of prospectively designed overviews of randomized trials. Eur Heart J. 2008;29:2669-80.

10.2.1. Women

A potential limitation of RCTs, including SPRINT, is that they are not specifically powered to determine the value of intensive SBP reduction in subgroups, including women in the case of SPRINT. However, in prespecified analyses, there was no evidence of an interaction between sex and treatment effect. Furthermore, no significant differences in CVD outcomes were observed between men and women in a large meta-analysis that included 31 RCTs with about 100,000 men and 90,000 women with hypertension (1 Some have called for conduct of a SPRINT-like trial with sufficient power to assess the effects of intensive SBP reduction in women {Wenger, 2016 #9131). Some have called for a SPRINT-like trial with sufficient power to assess the effects of intensive SBP reduction in women (2). In meta-analyses, there was no convincing evidence that different antihypertensive drug classes exerted sex-related differences in BP lowering or provided distinct CVD protection (1). Calcium antagonists offered slightly greater benefits for stroke prevention than did ACE inhibitors for women than for men, whereas calcium antagonists reduced all-cause deaths compared with placebo in men but not in women. However, these sex-related differences might have been due to chance because of the large number of statistical comparisons that were performed. The Heart Attack Trial and Hypertension Care Computing Project reported that beta blockers were associated with reduced mortality in men but not in women, but this finding was likely due to the low event rates in women (3). Similarly, in the open-label Second Australian National BP study, a significant reduction in CVD events was demonstrated in men but not in women with ACE inhibitors versus diuretics (4).

Adverse effects of antihypertensive therapy were noted twice as often in women as in men in the TOMHS study (5). A higher incidence of ACE inhibitor–induced cough and of edema with calcium antagonists was observed in women than in men (6). Women were more likely to experience hypokalemia and hyponatremia and less likely to experience gout with diuretics (7). Hypertension in pregnancy has special requirements (see Section 10.2.2).

- 1. Turnbull F, Woodward M, Neal B, et al. Do men and women respond differently to blood pressure-lowering treatment? Results of prospectively designed overviews of randomized trials. Eur Heart J. 2008;29:2669-80.
- 2. Wenger NK, Ferdinand KC, Bairey Merz CN, et al. Women, hypertension, and the Systolic Blood Pressure Intervention Trial. Am J Med. 2016;129:1030-6.
- 3. Fletcher A, Beevers DG, Bulpitt C, et al. Beta adrenoceptor blockade is associated with increased survival in male but not female hypertensive patients: a report from the DHSS Hypertension Care Computing Project (DHCCP). J Hum Hypertens. 1988;2:219-27.
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- Lewis CE, Grandits A, Flack J, et al. Efficacy and tolerance of antihypertensive treatment in men and women with stage 1 diastolic hypertension. Results of the Treatment of Mild Hypertension Study. Arch Intern Med. 1996;156:377-85.
- 6. Kloner RA, Sowers JR, DiBona GF, et al. Sex- and age-related antihypertensive effects of amlodipine. The Amlodipine Cardiovascular Community Trial Study Group. Am J Cardiol. 1996;77:713-22.
- 7. Igho Pemu P, Ofili E. Hypertension in women: part I. J Clin Hypertens (Greenwich). 2008;10:406-10.

	Recommendations for Treatment of Hypertension in Pregnancy				
Referer	References that support recommendations are summarized in Online Data Supplement 53.				
COR	LOE	Recommendations			
I	C-LD	1. Women with hypertension who become pregnant, or are planning to become pregnant, should be transitioned to methyldopa, nifedipine, and/or labetalol (1) during pregnancy (2-6).			
III: Harm	C-LD	2. Women with hypertension who become pregnant should not be treated with ACE inhibitors, ARBs, or direct renin inhibitors (4-6).			

References

- 1. James PR, Nelson-Piercy C. Management of hypertension before, during, and after pregnancy. Heart. 2004;90:1499-504.
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- 6. Ferrer RL, Sibai BM, Mulrow CD, et al. Management of mild chronic hypertension during pregnancy: a review. Obstet Gynecol. 2000;96:849-60.

10.3. Age-Related Issues

10.3.1. Older Persons

	Recommendations for Treatment of Hypertension in Older Persons					
Refe	References that support recommendations are summarized in Online Data Supplement 54.					
COR	LOE	Recommendations				
I	A	 Treatment of hypertension with a SBP treatment goal of less than 130 mm Hg is recommended for noninstitutionalized ambulatory community- dwelling adults (≥65 years of age) with an average SBP of 130 mm Hg or higher (1). 				
lla	C-EO	 For older adults (≥65 years of age) with hypertension and a high burden of comorbidity and limited life expectancy, clinical judgment, patient preference, and a team-based approach to assess risk/benefit is reasonable for decisions regarding intensity of BP lowering and choice of antihypertensive drugs. 				

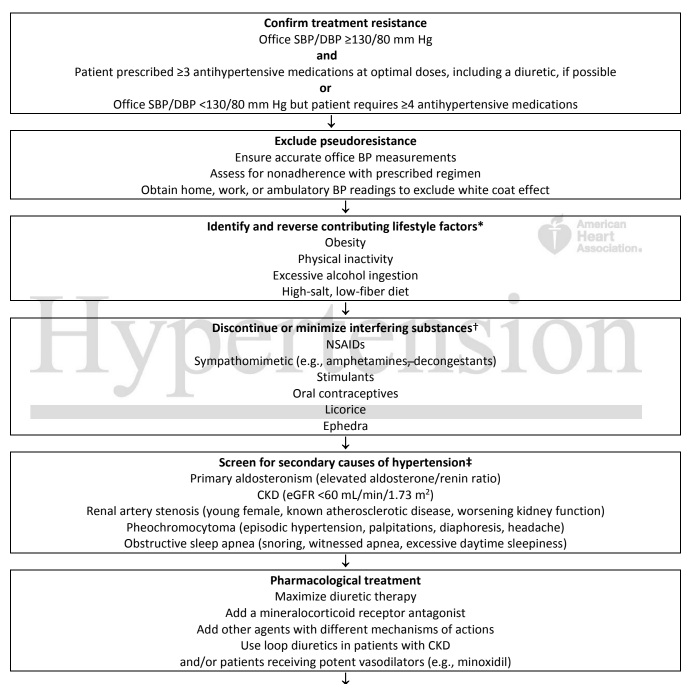
Reference

1. Williamson JD, Supiano MA, Applegate WB, et al. Intensive vs standard blood pressure control and cardiovascular disease outcomes in adults aged ≥75 years: a randomized clinical trial. JAMA. 2016;315:2673-82.

11. Other Considerations

11.1. Resistant Hypertension

Figure 10. Resistant Hypertension: Diagnosis, Evaluation, and Treatment



Refer to specialist

Refer to appropriate specialist for known or suspected secondary cause(s) of hypertension Refer to hypertension specialist if BP remains uncontrolled after 6 mo of treatment

*See additional details in Section 6, Nonpharmacological Intervention.

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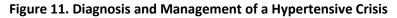
*See Section 5.4.1 and Table 14 for complete list of drugs that elevate BP.
*See Section 5.4 and Table 13 for secondary hypertension.
BP indicates blood pressure; CKD, chronic kidney disease; DBP, diastolic blood pressure; eGFR, estimated glomerular filtration rate; NSAIDs, nonsteroidal anti-inflammatory drugs; and SBP, systolic blood pressure.
Adapted with permission from Calhoun et al. (1) (American Heart Association, Inc.).

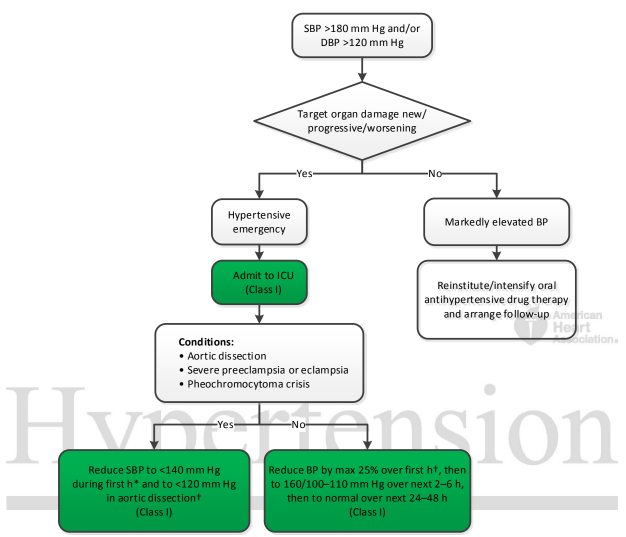
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1. Calhoun DA, Jones D, Textor S, et al. Resistant hypertension: diagnosis, evaluation, and treatment: a scientific statement from the American Heart Association Professional Education Committee of the Council for High Blood Pressure Research. Hypertension. 2008;51:1403-19.

11.2. Hypertensive Crises—Emergencies and Urgencies

Recommendations for Hypertensive Crises and Emergencies				
Refe	References that support recommendations are summarized in Online Data Supplement 55.			
COR	LOE	Recommendations		
I	B-NR	 In adults with a hypertensive emergency, admission to an intensive care unit is recommended for continuous monitoring of BP and target organ damage and for parenteral administration of an appropriate agent (Tables 19 and 20) (1, 2). 		
I	C-EO	2. For adults with a compelling condition (i.e., aortic dissection, severe preeclampsia or eclampsia, or pheochromocytoma crisis), SBP should be reduced to less than 140 mm Hg during the first hour and to less than 120 mm Hg in aortic dissection.		
I	C-EO	3. For adults without a compelling condition, SBP should be reduced by no more than 25% within the first hour; then, if stable, to 160/100 mm Hg within the next 2 to 6 hours; and then cautiously to normal during the following 24 to 48 hours.		





Colors correspond to Class of Recommendation in Table 1.

*Use drug(s) specified in Table 19.

+If other comorbidities are present, select a drug specified in Table 20.

BP indicates blood pressure; DBP, diastolic blood pressure; ICU, intensive care unit; and SBP, systolic blood pressure.

Class	Drug(s)	Usual Dose Range	Comments
CCB-	Nicardipine	Initial 5 mg/h,	Contraindicated in advanced aortic
dihydropyridines		increasing every 5 min by 2.5 mg/h to maximum 15 mg/h.	stenosis; no dose adjustment needed for elderly.
	Clevidipine	Initial 1–2 mg/h, doubling every 90 s until BP approaches target, then increasing by less than double every 5–10 min; maximum dose 32 mg/h; maximum duration 72 h.	Contraindicated in patients with soybean, soy product, egg, and egg product allergy and in patients with defective lipid metabolism (e.g., pathological hyperlipidemia, lipoid nephrosis or acute pancreatitis). Use low-end dose range for elderly patients.
Vasodilators— Nitric-oxide dependent	Sodium nitroprusside	Initial 0.3–0.5 mcg/kg/min; increase in increments of 0.5 mcg/kg/min to achieve BP target; maximum dose 10 mcg/kg/min; duration of treatment as short as possible. For infusion rates ≥4–10 mcg/kg/min or duration >30 min, thiosulfate can be coadministered to prevent cyanide toxicity.	Intra-arterial BP monitoring recommended to prevent "overshoot." Lower dosing adjustment required for elderly. Tachyphylaxis common with extended use. Cyanide toxicity with prolonged use can result in irreversible neurological changes and cardiac arrest.
Нy	Nitroglycerin	Initial 5 mcg/min; increase in increments of 5 mcg/min every 3–5 min to a maximum of 20 mcg/min.	Use only in patients with acute coronary syndrome and/or acute pulmonary edema. Do not use in volume-depleted patients.
Vasodilators— direct	Hydralazine	Initial 10 mg via slow IV infusion (maximum initial dose 20 mg); repeat every 4–6 h as needed.	BP begins to decrease within 10–30 min, and the fall lasts 2–4 h. Unpredictability of response and prolonged duration of action do not make hydralazine a desirable first-line agent for acute treatment in most patients.
Adrenergic blockers—beta1 receptor selective antagonist	Esmolol	Loading dose 500–1000 mcg/kg/min over 1 min followed by a 50- mcg/kg/min infusion. For additional dosing, the bolus dose is repeated and the infusion increased in 50- mcg/kg/min increments as needed to a maximum of 200 mcg/kg/min.	Contraindicated in patients with concurrent beta-blocker therapy, bradycardia, or decompensated HF. Monitor for bradycardia. May worsen HF. Higher doses may block beta ₂ receptors and impact lung function in reactive airway disease.
Adrenergic blockers— combined alpha ₁ and nonselective	Labetalol	Initial 0.3–1.0-mg/kg dose (maximum 20 mg) slow IV injection every 10 min or 0.4–1.0-mg/kg/h IV infusion up to 3 mg/kg/h. Adjust	Contraindicated in reactive airways disease or chronic obstructive pulmonary disease. Especially useful in hyperadrenergic syndromes. May worsen HF and should not be given in

beta receptor antagonist		rate up to total cumulative dose of 300 mg. This dose	patients with second- or third-degree heart block or bradycardia.
-		can be repeated every 4–6 h.	
Adrenergic blockers— nonselective alpha receptor antagonist	Phentolamine	IV bolus dose 5 mg. Additional bolus doses every 10 min as needed to lower BP to target.	Used in hypertensive emergencies induced by catecholamine excess (pheochromocytoma, interactions between monamine oxidase inhibitors and other drugs or food, cocaine toxicity, amphetamine overdose, or clonidine withdrawal).
Dopamine ₁ - receptor selective agonist	Fenoldopam	Initial 0.1–0.3 mcg/kg/min; may be increased in increments of 0.05–0.1 mcg/kg/min every 15 min until target BP is reached. Maximum infusion rate 1.6 mcg/kg/min.	Contraindicated in patients at risk of increased intraocular pressure (glaucoma) or intracranial pressure and those with sulfite allergy.
ACE inhibitor	Enalaprilat	Initial 1.25 mg over a 5-min period. Doses can be increased up to 5 mg every 6 h as needed to achieve BP target.	Contraindicated in pregnancy and should not be used in acute MI or bilateral renal artery stenosis. Mainly useful in hypertensive emergencies associated with high plasma renin activity. Dose not easily adjusted.
	7116	Prte 1	Relatively slow onset of action (15 min) and unpredictability of BP response. re; IV, intravenous; and MI, myocardial

BP indicates blood pressure; CCB, calcium channel blocker; HF, heart failure; IV, intravenous; and MI, myocardial infarction.

Table 20. Intravenous Antihypertensive Drugs for Treatment of Hypertensive Emergencies in Patients With
Selected Comorbidities

Comorbidity	Preferred	Comments	
-	Drug(s)*		
Acute aortic dissection	Esmolol labetalol	Requires rapid lowering of SBP to ≤120 mm Hg.	
		Beta blockade should precede vasodilator (e.g., nicardipine or	
		nitroprusside) administration, if needed for BP control or to	
		prevent reflex tachycardia or inotropic effect; SBP ≤120 mm Hg	
		should be achieved within 20 min.	
Acute pulmonary edema	Clevidipine,	Beta blockers contraindicated.	
	nitroglycerin		
	nitroprusside		
Acute coronary syndromes	Esmolol ⁺	Nitrates given in the presence of PDE-5 inhibitors may induce	
	labetalol	profound hypotension. Contraindications to beta blockers	
	nicardipine	include moderate-to-severe LV failure with pulmonary edema,	
	nitroglycerin†	bradycardia (<60 bpm), hypotension (SBP <100 mm Hg), poor	
		peripheral perfusion, second- or third-degree heart block, and	
	Chaudalia in a	reactive airways disease.	
Acute renal failure	Clevidipine	N/A	
	fenoldopam nicardipine		
Eclampsia or preeclampsia	Hydralazine	Requires rapid BP lowering.	
	labetalol	ACE inhibitors, ARBs, renin inhibitors, and nitroprusside	
	nicardipine	contraindicated.	
Perioperative hypertension	Clevidipine	Intraoperative hypertension is most frequently seen during	
(BP \ge 160/90 mm Hg or SBP	esmolol	anesthesia induction and airway manipulation.	
elevation $\geq 20\%$ of the	nicardipine,		
preoperative value that	nitroglycerin		
persists for >15 min)	0,		
Acute sympathetic discharge	Clevidipine	Requires rapid lowering of BP.	
or catecholamine excess	nicardipine		
states (e.g.,	phentolamine		
pheochromocytoma, post-			
carotid endarterectomy			
status)			
Acute ICH	Section 9.4.1	Section 9.4.1	
Acute ischemic stroke	Section 9.4.2	Section 9.4.2	

*Agents are listed in alphabetical order, not in order of preference.

[†]Agent of choice for acute coronary syndromes.

ACE indicates angiotensin-converting enzyme; ARB, angiotensin receptor blocker; BP, blood pressure; bpm, beats per minute; ICH, intracerebral hemorrhage; LV, left ventricular; PDE-5, phosphodiesterase type-5; and SBP, systolic blood pressure.

References

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- 2. Peacock WF, Chandra A, Char D, et al. Clevidipine in acute heart failure: results of the A Study of Blood Pressure Control in Acute Heart Failure--A Pilot Study (PRONTO). Am Heart J. 2014;167:529-36.

11.3. Cognitive Decline and Dementia

	Recommendation for Prevention of Cognitive Decline and Dementia			
Refer	References that support the recommendation are summarized in Online Data Supplement 56.			
COR	LOE	Recommendation		
lla	B-R	1. In adults with hypertension, BP lowering is reasonable to prevent cognitive decline and dementia (1-6).		

References

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SIDD

Recommendations for Treatment of Hypertension in Patients Undergoing Surgical				
	Procedures			
Referen	ces that su	pport recommendations are summarized in Online Data Supplements 57 and 58.		
COR	LOE	Recommendations		
		Preoperative		
I.	B-NR	 In patients with hypertension undergoing major surgery who have been on beta blockers chronically, beta blockers should be continued (1-7). 		
lla	C-EO	2. In patients with hypertension undergoing planned elective major surgery, it is reasonable to continue medical therapy for hypertension until surgery.		
llb	B-NR	3. In patients with hypertension undergoing major surgery, discontinuation of ACE inhibitors or ARBs perioperatively may be considered (8-10).		
llb	C-LD	4. In patients with planned elective major surgery and SBP of 180 mm Hg or higher or DBP of 110 mm Hg or higher, deferring surgery may be considered (11, 12).		
III: Harm	B-NR	5. For patients undergoing surgery, abrupt preoperative discontinuation of beta blockers or clonidine is potentially harmful (2, 13).		
III: Harm	B-NR	6. Beta blockers should not be started on the day of surgery in beta blocker- naïve patients (14).		

11.4. Patients Undergoing Surgical Procedures

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Intraoperative				
I	C-EO	7. Patients with intraoperative hypertension should be managed with intravenous medications (Table 19) until such time as oral medications can be resumed.		

References

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- 12. Howell SJ, Sear JW, Foex P. Hypertension, hypertensive heart disease and perioperative cardiac risk. Br J Anaesth. 2004;92:570-83.
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- 14. Devereaux PJ, Yang H, Yusuf S, et al. Effects of extended-release metoprolol succinate in patients undergoing noncardiac surgery (POISE trial): a randomised controlled trial. Lancet. 2008;371:1839-47.

12. Strategies to Improve Hypertension Treatment and Control

12.1.1. Antihypertensive Medication Adherence Strategies

	Recommendations for Antihypertensive Medication Adherence Strategies			
Reference	ces that su	upport recommendations are summarized in Online Data Supplements 59 and 60.		
COR	LOE	Recommendations		
I	B-R	 In adults with hypertension, dosing of antihypertensive medication once daily rather than multiple times daily is beneficial to improve adherence (1- 3). 		
lla	B-NR	2. Use of combination pills rather than free individual components can be useful to improve adherence to antihypertensive therapy (4-7).		

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Available fixed-dose combination drug therapy is listed in Online Data Supplement D.

References

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- 7. Yang W, Chang J, Kahler KH, et al. Evaluation of compliance and health care utilization in patients treated with single pill vs. free combination antihypertensives. Curr Med Res Opin. 2010;26:2065-76.

12.1.2. Strategies to Promote Lifestyle Modification

1.2. 00	4109.00	American
	Reco	mmendation for Strategies to Promote Lifestyle Modification
Refer	ences that	support the recommendation are summarized in Online Data Supplement 61.
COR	LOE	Recommendations
I	C-EO	1. Effective behavioral and motivational strategies to achieve a health lifestyle (i.e., tobacco cessation, weight loss, moderation in alcohol intake increased physical activity, reduced sodium intake, and consumption of healthy diet) are recommended for adults with hypertension (1, 2).

References

- 1. Artinian NT, Fletcher GF, Mozaffarian D, et al. Interventions to promote physical activity and dietary lifestyle changes for cardiovascular risk factor reduction in adults: a scientific statement from the American Heart Association. Circulation. 2010;122:406-41.
- Eckel RH, Jakicic JM, Ard JD, et al. 2013 AHA/ACC guideline on lifestyle management to reduce cardiovascular risk: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. Circulation. 2014;129(suppl 2):S76-99.

12.2. Structured, Team-Based Care Interventions for Hypertension Control

Recomm	Recommendation for Structured, Team-Based Care Interventions for Hypertension Control				
Refer	References that support the recommendation are summarized in Online Data Supplement 62.				
COR	LOE	Recommendations			
I.	Α	1. A team-based care approach is recommended for adults with hypertension (1-7).			

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References

- 1. Carter BL, Rogers M, Daly J, et al. The potency of team-based care interventions for hypertension: a meta-analysis. Arch Intern Med. 2009;169:1748-55.
- 2. Clark CE, Smith LF, Taylor RS, et al. Nurse led interventions to improve control of blood pressure in people with hypertension: systematic review and meta-analysis. BMJ. 2010;341:c3995.
- 3. Proia KK, Thota AB, Njie GJ, et al. Team-based care and improved blood pressure control: a community guide systematic review. Am J Prev Med. 2014;47:86-99.
- 4. Santschi V, Chiolero A, Colosimo AL, et al. Improving blood pressure control through pharmacist interventions: a meta-analysis of randomized controlled trials. J Am Heart Assoc. 2014;3:e000718.
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12.3. Health Information Technology–Based Strategies to Promote Hypertension Control

12.3.1. EHR and Patient Registries



	Recommendations for EHR and Patient Registries					
Refe	References that support recommendations are summarized in Online Data Supplement 63.					
COR	LOE	Recommendations				
I.	B-NR	1. Use of the EHR and patient registries is beneficial for identification of patients with undiagnosed or undertreated hypertension (1-3).				
I.	B-NR	2. Use of the EHR and patient registries is beneficial for guiding quality improvement efforts designed to improve hypertension control (1-3).				

References

- 1. Rakotz MK, Ewigman BG, Sarav M, et al. A technology-based quality innovation to identify undiagnosed hypertension among active primary care patients. Ann Fam Med. 2014;12:352-8.
- Borden WB, Maddox TM, Tang F, et al. Impact of the 2014 expert panel recommendations for management of high blood pressure on contemporary cardiovascular practice: insights from the NCDR PINNACLE registry. J Am Coll Cardiol. 2014;64:2196-203.
- 3. Jaffe MG, Lee GA, Young JD, et al. Improved blood pressure control associated with a large-scale hypertension program. JAMA. 2013;310:699-705.

12.3.2. Telehealth Interventions to Improve Hypertension Control

Recommendation for Telehealth Interventions to Improve Hypertension Control				
Refer	References that support the recommendation are summarized in Online Data Supplement 64.			
COR	LOE	Recommendations		
lla	Α	1. Telehealth strategies can be useful adjuncts to interventions shown to reduce BP for adults with hypertension (1-5).		

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References

- 1. Omboni S, Gazzola T, Carabelli G, et al. Clinical usefulness and cost effectiveness of home blood pressure telemonitoring: meta-analysis of randomized controlled studies. J Hypertens. 2013;31:455-67; discussion 467-8.
- 2. Verberk WJ, Kessels AGH, Thien T. Telecare is a valuable tool for hypertension management, a systematic review and meta-analysis. Blood Press Monit. 2011;16:149-55.
- 3. Agarwal R, Bills JE, Hecht TJW, et al. Role of home blood pressure monitoring in overcoming therapeutic inertia and improving hypertension control: a systematic review and meta-analysis. Hypertension. 2011;57:29-38.
- 4. Liu S, Dunford SD, Leung YW, et al. Reducing blood pressure with Internet-based interventions: a meta-analysis. Can J Cardiol. 2013;29:613-21.
- 5. Burke LE, Ma J, Azar KMJ, et al. Current science on consumer use of mobile health for cardiovascular disease prevention: a scientific statement from the American Heart Association. Circulation. 2015;132:1157-213.

12.4. Improving Quality of Care for Patients With Hypertension

12.4.1. Performance Measures

Recommendation for Performance Measures					
Refer	References that support the recommendation are summarized in Online Data Supplement 65.				
COR	LOE	Recommendations American			
lla	B-NR	 Use of performance measures in combination with other quality improvement strategies at patient-, provider-, and system-based levels is reasonable to facilitate optimal hypertension control (1-3). 			

References

- 1. Svetkey LP, Pollak KI, Yancy WS Jr, et al. Hypertension improvement project: randomized trial of quality improvement for physicians and lifestyle modification for patients. Hypertension. 2009;54:1226-33.
- 2. de Lusignan S, Gallagher H, Jones S, et al. Audit-based education lowers systolic blood pressure in chronic kidney disease: the Quality Improvement in CKD (QICKD) trial results. Kidney Int. 2013;84:609-20.
- 3. Jaffe MG, Lee GA, Young JD, et al. Improved blood pressure control associated with a large-scale hypertension program. JAMA. 2013;310:699-705.

12.4.2. Quality Improvement Strategies

	Recommendation for Quality Improvement Strategies			
Reference	References that support the recommendation are summarized in Online Data Supplements 66 and 67.			
COR	LOE	Recommendations		
lla	B-R	1. Use of quality improvement strategies at the health system, provider, and patient levels to improve identification and control of hypertension can be effective (1-8).		

References

- 1. Walsh JME, McDonald KM, Shojania KG, et al. Quality improvement strategies for hypertension management: a systematic review. Med Care. 2006;44:646-57.
- 2. Carter BL, Rogers M, Daly J, et al. The potency of team-based care interventions for hypertension: a meta-analysis. Arch Intern Med. 2009;169:1748-55.
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- 8. Agarwal R, Bills JE, Hecht TJW, et al. Role of home blood pressure monitoring in overcoming therapeutic inertia and improving hypertension control: a systematic review and meta-analysis. Hypertension. 2011;57:29-38.

12.5. Financial Incentives

	Recommendations for Financial Incentives					
Refe	References that support recommendations are summarized in Online Data Supplement 68.					
COR	LOE	Recommendations				
lla	B-R	1. Financial incentives paid to providers can be useful in achieving improvements in treatment and management of patient populations with hypertension (1-3).				
lla	B-NR	2. Health system financing strategies (e.g., insurance coverage and copayment benefit design) can be useful in facilitating improved medication adherence and BP control in patients with hypertension (4).				

References

- 1. Hysong SJ, Simpson K, Pietz K, et al. Financial incentives and physician commitment to guideline-recommended hypertension management. Am J Manag Care. 2012;18:e378-91.
- 2. Petersen LA, Simpson K, Pietz K, et al. Effects of individual physician-level and practice-level financial incentives on hypertension care: a randomized trial. JAMA. 2013;310:1042-50.
- 3. Karunaratne K, Stevens P, Irving J, et al. The impact of pay for performance on the control of blood pressure in people with chronic kidney disease stage 3-5. Nephrol Dial Transplant. 2013;28:2107-16.
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13. The Plan of Care for Hypertension

Table 21

	Recommendation for the Plan of Care for Hypertension					
COR	LOE	Recommendation				
I	C-EO	 Every adult with hypertension should have a clear, detailed, and current evidence-based plan of care that ensures the achievement of treatment and self-management goals, encourages effective management of comorbid conditions, prompts timely follow-up with the healthcare team, and adheres to CVD GDMT (Table 22). 				

Clinician's Sequential Flow Chart for the Management of Hypertension						
Measure office BP accurately	Section 4					
Detect white coat hypertension or masked	Section 4					
hypertension by using ABPM and HBPM						
Evaluate for secondary hypertension	Section 5					
Identify target organ damage	Sections 5 and 7					
Introduce lifestyle interventions	Section 6					
Identify and discuss treatment goals	Sections 7 and 8					
Use ASCVD risk estimation to guide BP threshold for	Section 8.1.2					
drug therapy						
Align treatment options with comorbidities	Section 9					
Account for age, race, ethnicity, sex, and special	Sections 10 and 11					
circumstances in antihypertensive treatment						
Initiate antihypertensive pharmacological therapy	Section 8					
Insure appropriate follow-up	Section 8					
Use team-based care	Section 12					
Connect patient to clinician via telehealth	Section 12					
Detect and reverse nonadherence	Section 12					
Detect white coat effect or masked uncontrolled	Section 4					
hypertension	Association					
Use health information technology for remote	Section 12					
monitoring and self-monitoring of BP						

Table 21. Clinician's Sequential Flow Chart for the Management of Hypertension

ABPM indicates ambulatory blood pressure monitoring; ASCVD, atherosclerotic cardiovascular disease; BP, blood pressure; and HBPM, home blood pressure monitoring.

13.1. Health Literacy

Communicating alternative behaviors that support self-management of healthy BP in addition to medication adherence is important. This should be done both verbally and in writing. Today, mobile phones have a recording option. For patients with mobile phones, the phone can be used to inform patients and family members of medical instructions after the doctor's visit as an additional level of communication. Inclusion of a family member or friend that can help interpret and encourage self-management treatment goals is suggested when appropriate. Examples of needed communication for alternative behaviors include a specific regimen relating to physical activity; a specific sodium-reduced meal plan indicating selections for breakfast, lunch, and dinner; lifestyle recommendations relating to sleep, rest, and relaxation; and finally, suggestions and alternatives to environmental barriers, such as barriers that prevent healthy food shopping or limit reliable transportation to and from appointments with health providers and pharmacy visits.

13.2. Access to Health Insurance and Medication Assistance Plans

Health insurance and medication plan assistance for patients is especially important to improving access to and affordability of medical care and BP medications. Learning how the patient financially supports and budgets for his or her medical care and medications offers the opportunity to share additional insight relating to cost reductions, including restructured payment plans. Ideally, this would improve the patient's compliance with medication adherence and treatment goals.

13.3. Social and Community Services

Health care can be strengthened through local partnerships. Hypertensive patients, particularly patients with lower incomes, have more opportunity to achieve treatment goals with the assistance of strong local partnerships. In patients with low socioeconomic status or patients who are challenged by social situations, integration of social and community services offers complementary reinforcement of clinically identified treatment goals. Social and community services are helpful when explicitly related to medical care. However, additional financial support and financial services are incredibly beneficial to patients, some of whom may choose to skip a doctor's appointment to pay a residential utility bill.



	Associated Section(s) of Guideline			
Plan of Care	and Other Reference(s)			
Pharmacological and nonpharmacological treatments				
Medication selection (initial and ongoing)	Section 8.1			
Monitoring for adverse effects and adherence	Sections 8.3.1, 8.3.2, 12.1.1			
Nonpharmacological interventions	Sections 6, 12.1.2 (1)			
• Diet				
Exercise				
Weight loss if overweight				
 Moderate alcohol consumption 				
Management of common comorbidities and conditions				
Ischemic heart disease	Section 9.1 (2, 3)			
Heart failure	Section 9.2 (4)			
Reduced ejection fraction				
Preserved ejection fraction				
Diabetes mellitus	Section 9.6 (5)			
Chronic kidney disease	Section 9.3			
Cerebrovascular disease	Section 9.4			
Peripheral arterial disease	Section 9.5			
Atrial fibrillation	Section 9.8 Association			
Valvular heart disease	Section 9.9			
Left ventricular hypertrophy	Section 7.3			
Thoracic aortic disease	Section 9.10			
Patient and family education	1			
Achieving BP control and self-monitoring	Sections 4.2, 8.2			
Risk assessment and prognosis	Section 8.1.2			
Sexual activity and dysfunction	Section 11.4			
Special patient groups				
Pregnancy	Section 10.2.2			
Older persons	Section 10.3.1			
Children and adolescents	Section 10.3.2			
Metabolic syndrome	Section 9.7			
Possible secondary causes of hypertension	Section 5.4			
Resistant hypertension	Section 11.1			
Patients with hypertension undergoing surgery	Section 11.5			
Renal transplantation	Section 9.3.1			
Psychosocial factors				
Sex-specific issues	Section 10.2			
Culturally sensitive issues (race and ethnicity)	Section 10.2			
Resource constraints	Section 12.5			
Clinician follow-up, monitoring, and care coordination	Section 12.5			
Follow-up visits	Sections 8 1 3 8 3 1 8 3 7			
Team-based care	Sections 8.1.3, 8.3.1, 8.3.2 Section 12.2			
Electronic health record				
Health information technology tools for remote and self-monitoring	Section 12.3.1			
	Section 12.3.2			
Socioeconomic and cultural factors	Section 12.1.2			
Health literacy	Section 13.1.3			
Access to health insurance and medication assistance plans	Section 13.1.3			
Social services	Section 13.1.3			

Community services	Section 13.1.3
BP indicates blood pressure.	

References

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- 3. Fihn SD, Gardin JM, Abrams J, et al. 2012 ACCF/AHA/ACP/AATS/PCNA/SCAI/STS guideline for the diagnosis and management of patients with stable ischemic heart disease: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines, and the American College of Physicians, American Association for Thoracic Surgery, Preventive Cardiovascular Nurses Association, Society for Cardiovascular Angiography and Interventions, and Society of Thoracic Surgeons. Circulation. 2012;126:e354-471.
- 4. Yancy CW, Jessup M, Bozkurt B, et al. 2013 ACCF/AHA guideline for the management of heart failure: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. Circulation. 2013;128:e240-327.
- 5. Standards of Medical Care in Diabetes--2016: Summary of Revisions. Diabetes Care. 2016;39(suppl 1):S4-5.

Association

14. Summary of BP Thresholds and Goals for Pharmacological Therapy

 Table 23. BP Thresholds for and Goals of Pharmacological Therapy in Patients With Hypertension

 According to Clinical Conditions

	BP Threshold, mm	
Clinical Condition(s)	Hg	BP Goal, mm Hg
General		
Clinical CVD or 10-year ASCVD risk ≥10%	≥130/80	<130/80
No clinical CVD and 10-year ASCVD risk <10%	≥140/90	<130/80
Older persons (≥65 years of age; noninstitutionalized,	≥130 (SBP)	<130 (SBP)
ambulatory, community-living adults)		
Specific comorbidities		
Diabetes mellitus	≥130/80	<130/80
Chronic kidney disease	≥130/80	<130/80
Chronic kidney disease after renal transplantation	≥130/80	<130/80
Heart failure	≥130/80	<130/80
Stable ischemic heart disease	≥130/80	<130/80
Secondary stroke prevention	≥140/90	<130/80
Secondary stroke prevention (lacunar)	≥130/80	<130/80
Peripheral arterial disease	≥130/80	<130/80

ASCVD indicates atherosclerotic cardiovascular disease; BP, blood pressure; CVD, cardiovascular disease; and SBP, systolic blood pressure.

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American College of Cardiology

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American College of Cardiology/American Heart Association

Katherine A. Sheehan, PhD, Director, Guideline Strategy and Operations Abdul R. Abdullah, MD, Science and Medicine Advisor Naira Tahir, MPH, Associate Guideline Advisor

American Heart Association

John J. Warner, MD, President Nancy Brown, Chief Executive Officer Rose Marie Robertson, MD, FAHA, Chief Science and Medicine Officer Gayle R. Whitman, PhD, RN, FAHA, FAAN, Senior Vice President, Office of Science Operations Jody Hundley, Production Manager, Scientific Publications, Office of Science Operations

Key Words: ACC/AHA Clinical Practice Guidelines, blood pressure; hypertension; ambulatory care; antihypertensive agents; behavior modification; risk reduction; treatment adherence; treatment outcomes; Systems of care, hypertension emergency, secondary hypertension, blood pressure, measurement, diabetes, chronic kidney disease, resistant hypertension, nonpharmacologic treatment, lifestyle measures

Appendix 1. Author Relationships With Industry and Other Entities (Relevant)—2017 ACC/AHA/AAPA/ABC/ACPM/AGS/APhA/ASH/ASPC/NMA/PCNA Guideline for the Prevention, Detection, Evaluation, and Management of High Blood Pressure in Adults (October 2017)

			Speakers	Ownership/ Partnership/		Institutional, Organizational, or Other	Expert	
Committee Member	Employment	Consultant	Bureau	Principal	Personal Research	Financial Benefit	Witness	Salary
Paul K. Whelton <i>(Chair)</i>	Tulane University School of Hygiene and Tropical Medicine—Show Chwan Professor of Global Public Health	None	None	None	None	None	None American Heart Associat	None
Robert M. Carey (Vice Chair)	University of Virginia—Dean Emeritus and University Professor, Department of Medicine	None	None	None	None	None	None	None
Wilbert S. Aronow	Westchester Medical Center and New York Medical College— Professor of Medicine	None	None	None	None	None	None	None
Donald E. Casey, Jr	Thomas Jefferson College of Population Health—Adjunct Faculty; Alvarez & Marsal Ipo4health— Principal and Founder	None	None	None	None	None	None	None
Karen J. Collins	Collins Collaboration— President	None	None	None	None	None	None	None

Cheryl Dennison	John Hopkins	None	None	None	None	None	None	None
Himmelfarb	University—							
	Professor of							
	Nursing and							
	Medicine,							
	Institute for							
	Clinical and							
	Translational							
	Research							
Sondra M DePalma	PinnacleHealth	None	None	None	None	None	None	None
	CardioVascular							
	Institute—							
	Physician					C.	America	
	Assistant;						Heart	-
	American						Associat	on.
	Academy of PAs—						P10000101	COTTO
	Director,							
	Regulatory and							
	Professional							
	Practice							
Samuel Gidding	Alfred I. Dupont	None	None	None	None	None	None	None
	Hospital for							
	Children—Chief,							
	Division of							
	Pediatric							
	Cardiology,							
	Nemours Cardiac							
	Center	line .						
David C. Goff, Jr*	Colorado School	None	None	None	None	None	None	None
	of Public Health—							
	Professor and							
	Dean,							
	Department of							
	Epidemiology							

Kenneth A. Jamerson	University of Michigan Health System— Professor of Internal Medicine and Frederick G.L. Huetwell Collegiate Professor of Cardiovascular Medicine	None	None	None	None	None	None	None
Daniel W. Jones	University of Mississippi Medical Center— Professor of Medicine and Physiology; Metabolic Diseases and Nutrition— University	None	None	None	None	None	None American Heart Associat	None
	Sanderson Chair in Obesity Mississippi Center for Obesity Research— Director, Clinical and Population Science				ns	10)1	
Eric J. MacLaughlin	Texas Tech University Health Sciences Center— Professor and Chair, Department of Pharmacy Practice, School of Pharmacy	None	None	None	None	None	None	None

Paul Muntner	University of	None	None	None	None	None	None	None
	Alabama at							
	Birmingham—							
	Professor,							
	Department of							
	Epidemiology							
Bruce Ovbiagele	Medical	None	None	None	None	None	None	None
	University of							
	South Carolina—							
	Pihl Professor and							
	Chairman of							
	Neurology							
Sidney C. Smith, Jr	University of	None	None	None	None	None	Nonenerica	None
	North Carolina at						Heart	
	Chapel Hill—						Associat	on
	Professor of						10000000	
	Medicine; Center							
	for Cardiovascular							
	Science and							
	Medicine—							
	Director							-
Crystal C. Spencer	Spencer Law,	None	None	None	None	None	None	None
	PA—Attorney at							
	Law							
Randall S. Stafford	Stanford	None	None	None	None	None	None	None
	Prevention							
	Research							
	Center—Professor	-						
	of Medicine;							
	Program on							
	Prevention							
	Outcomes—							
	Director							
Sandra J. Taler	Mayo Clinic—	None	None	None	None	None	None	None
	Professor of							
	Medicine, College							
	of Medicine							
Randal J. Thomas	Mayo Clinic—	None	None	None	None	None	None	None
	Medical Director,							
	Cardiac							
	Rehabilitation							
	Program							

Kim A. Williams, Sr	Rush University Medical Center— James B. Herrick Professor; Division of Cardiology—Chief	None	None	None	None	None	None	None
Jeff D. Williamson	Wake Forest Baptist Medical Center—Professor of Internal Medicine; Section on Gerontology and Geriatric Medicine—Chief	None	None	None	None	None	None American	None
Jackson T. Wright, Jr	Case Western Reserve University— Professor of Medicine; William T. Dahms MD Clinical Research	None	None	None	None	None	None	None
Н	Unit—Program Director; University Hospitals Case Medical Center— Director, Clinical Hypertension Program				ns	10)1	

This table represents the relationships of committee members with industry and other entities (RWI) that are considered relevant to this document. Although most ACC/AHA guideline writing committees are constituted such that no more than half the members may have relevant RWI for 1 year before and during development of the guideline, rules for the prevention guidelines require that no members have relevant RWI from 1 year before appointment until 1 year after publication of the guideline. Members' RWI were reviewed and updated at all meetings and conference calls of the writing committee during the document development period. The complete ACC/AHA policy on RWI is available at http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/relationships-with-industry-policy.

We gratefully acknowledge the contributions of Dr. Lawrence Appel, who served as a member of the Writing Committee from November 2014 to September 2015.

*Dr. David C. Goff resigned from the writing committee in December 2016 because of a change in employment before the recommendations were balloted. The writing committee thanks him for his contributions, which were extremely beneficial to the development of the draft.

AAPA indicates American Academy of Physician Assistants; ACC, American College of Cardiology; ACPM, American College of Preventive Medicine; AGS, American Geriatrics Society; AHA, American Heart Association; APhA, American Pharmacists Association; ASH, American Society of Hypertension; ASPC, American Society for Preventive Cardiology; ABC, Association of Black Cardiologists; NMA, National Medical Association; and PCNA, Preventive Cardiovascular Nurses Association.

Appendix 2. Reviewer Relationships With Industry and Other Entities (Comprehensive)—2017 ACC/AHA/AAPA/ABC/ACPM/AGS/APhA/ASH/ASPC/NMA/PCNA Guideline for the Prevention, Detection, Evaluation, and Management of High Blood Pressure in Adults (October 2017)

Reviewer	Representati on	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness	Salary
Kim K. Birtcher	Official Reviewer— TFPG Lead Reviewer	University of Houston College of Pharmacy— Clinical Professor, Department of Pharmacy Practice and Translational Research	• Jones & Bartlett Learning	None	None	None	Accreditation Council for Clinical Lipidology†	None Ameri Heat Assoc	
Roger Blumenthal	Official Reviewer— Prevention Subcommitte e	Johns Hopkins Hospital— Kenneth Jay Pollin Professor of Cardiology; Ciccarone Center for the Prevention of Heart Disease— Director	None	None	None	None	None	None	None

Anna	Official	University of	None	None	None	None	None	None	None
Dominiczak	Reviewer—	Glasgow—	None	None	None	None	None	None	None
DOMINICZAK	AHA	Regius							
	ALIA	Professor of							
		Medicine;							
		Vice-Principal							
		and Head of							
		College of							
		Medical,							
		Veterinary							
		and Life							
		Sciences							
Carlos M.	Official	Wake Forest	None	None	None	None	None	Nono	None
Ferrario	Reviewer—	School of	None	None	None	NOTE	None	None Ameri	NOTE
Ferrario	AHA	Medicine—						Hear	T
	ALIA	Professor, of						Assoc	iation.
		Physiology							
		and							
		Pharmacolog							
		y; Hypertension							
		and Vascular							
		Disease							
		Center—							
		Director		2		1			
Eugene	Official	University of	 RubiconMD* 	None	None	Amgen	None	• Third	None
Yang	Reviewer—	Washington	Regeneron*	None	None	 Angen Inc.* 	None	• miru party,	None
Tang	ACC-BOG	School of	• Regeneron					CAD,	
	ACC-BOG	Medicine—				Gilead		2016*	
		Associate				Sciences, Inc.		2010	
		Clinical				(DSMB)*			
		Professor of				(DSIVIB)			
		Medicine;							
		UW Medicine							
		Eastside							
		Specialty							
		Center—							
		Medical							
		Director							
		Director					1		

Robert Jay	Organizationa	Massachusett	None	None	None	None	None	 Defendan 	None
Amrien	l Reviewer—	s General						t, aortic	
	AAPA	Hospital—						dissection	
		Clinical						, 2016*	
		Physician						-	
		Assistant,							
		Chelsea							
		Health							
		Center;							
		Bryant							
		University—							
		Physician							
		Assistant						Ameri	can
		Program						Heal	+
Greg	Organizationa	Montana	None	None	None	None	American Academy of	None Assoc	None
Holzman	l Reviewer—	Department					Family Medicine ⁺	- russe	P (1944) 10 (1947) 1 (1940)
	ACPM	of Public					 American College of 		
		Health and					Preventive Medicine ⁺		
		Human							
		Services—							
		State Medical							
		Officer							
Martha	Organizationa	University of	None	None	None	None	• REATA (spouse)*	None	None
Gulati	l Reviewer—	Arizona							
	ASPC	College of							
		Medicine—							
		Professor of							
		Medicine;	-						
		Chief,							
		Division of							
		Cardiology;							
		University							
		Medicine							
		Cardiovascula							
		r Institute in							
		Phoenix—							
		Physician							
		Executive							
		Director,							
		Banner							

Wallace Johnson	Organizationa I Reviewer— NMA	University of Maryland Medical Center— Assistant Professor of	None	None	None	Amgen†	None	None	None
		Medicine							
Nancy Houston Miller	Organizationa l Reviewer— PCNA	The Lifecare Company— Associate Director	 Moving Analytics* 	None	None	None	None	None	None
Aldo J. Peixoto	Organizationa I Reviewer— ASH	Yale University School of Medicine— Professor of Medicine (Nephrology); Associate	• Lundbeck Inc.	None	None	• Bayer Healthca re Pharmac euticals†	Bayer Healthcare Pharmaceuticals	None Ameri Heat Assoc	
		Chair for Ambulatory Services Operations and Quality, Department of Internal Medicine; Clinical Chief,	þe	1	te	1	S 1(D1	1
		Section of Nephrology							
Carlos Rodriguez	Organizationa I Reviewer— ABC	Wake Forest University— Professor, Epidemiology and Prevention	• Amgen Inc.	None	None	None	None	None	None

Joseph	Organizationa	University of	None	None	None	None	National Lipid	• Defendan	None
Saseen	l Reviewer—	Colorado					Association ⁺	t, statin	
	APhA	Anschutz						use, 2016	
		Medical							
		Campus—							
		Vice-Chair,							
		Department							
		of Clinical							
		Pharmacy,							
		Skaggs School							
		of Pharmacy							
		and							
		Pharmaceutic						Ameri	000
		al Sciences						Hoal	et in the second
Mark	Organizationa	University of	None	None	None	None	American Geriatrics	None Assoc	None
Supiano	l Reviewer—	Utah School					Society [†]	Assoc	ARTOTA
	AGS	of Medicine—					Division Chief [†]		
		D. Keith					McGraw-Hill Medical		
		Barnes, MD,							
		and Dottie							
		Barnes							
		Presidential							
		Endowed							
		Chair in							
		Medicine;		1		1			
		Chief,							
		Division of							
		Geriatrics; VA							
		Salt Lake City							
		Geriatric							
		Research—							
		Director,							
		Education,							
		and Clinical							
		Center;							
		University of							
		Utah Center							
		on Aging							
		Executive—							
		Director							
		Director					1		l

Sana M. Al-	Content	Duke Clinical	None	None	None	 AHRQ* 	• Elsevier*	• Third	None
Khatib	Reviewer—	Research				• FDA*	• NIH, NHLBI	party,	
	ACC/AHA	Institute—				 PCORI* 		implantab	
	Task Force on	Professor of				• VA		le	
	Clinical	Medicine				Health		cardiverte	
	Practice					System		r	
	Guidelines					(DSMB)		defibrillat	
						(ors, 2017	
George	Content	University of	None	None	None	 AbbVie, 	None	None	None
Bakris	Reviewer	Chicago				Inc.			
		Medicine—				 Janssen, 			
		Professor of				Bayer,		-	
		Medicine;				Relypsa		Ameri	can
		Director,						Hear	rt
		Hypertensive						Assoc	iation.
		Diseases Unit							
Jan Basile	Content	Medical	None	 Amgen 	None	• Eli Lilly	None	None	None
	Reviewer	University of		Inc.		and			
		South		 Arbor 		Compan			
		Carolina—		 Janssen 		У			
		Professor of		Pharmace		NHLBI			
		Medicine,		uticals,					
		Seinsheimer		Inc					
		Cardiovascula				2			
		r Health							
		Program;							
		Ralph H							
		Johnson VA							
		Medical							
		Center—							
		Internist							
Joshua A.	Content	Vanderbilt	 AstraZeneca* 	None	• EMX†	 Bristol 	Vascular Interventional	None	• 2015-
Beckman	Reviewer—	University	 Merck* 		 JanaCare[†] 	Myers	Advances*		Defendant;
	ACC/AHA	Medical	 SANOFI* 			Squibb*			Venous
	Task Force on	Center:							thromboemb
	Clinical	Director,							olism*
	Practice	Cardiovascula							
	Guidelines	r Fellowship							
		Program,							

John	Content	University of	• CVRx	None	None	• CVRx*	None	None	None
Bisognano	Reviewer	Rochester				● NIH*			
		Medical							
		Center—							
		Cardiologist							
Biykem	Content	Baylor	None	None	None	 Novartis 	None	None	None
Bozkurt	Reviewer—	College of				Corporat			
	ACC/AHA	Medicine—				ion			
	Task Force on	Medical Care							
	Clinical	Line							
	Practice	Executive,							
	Guidelines	Cardiology							
		Chief, Gordon						Amer	can
		Cain						and the second se	-6
		Chair,						Hear	1. Lotton
		Professor of						Assoc	iation.
		Medicine,							
		Debakey							
David	Content	University of	Novartis	None	None	MEDTRO	None	None	None
Calhoun	Reviewer	Alabama,	Valencia	None	Hone	NIC*		None	None
camban		Birmingham	Technologies*			ReCor			
		School of	reennoiogies			Medical*			
		Medicine—				Wiedlear			
		Professor,							
		Department		1		1			
		of							
		Cardiovascula							
	(r Disease							

Les auties E	Contont	0.0000	Naza	Neze	Nega			. Defenden	Nana
Joaquin E.	Content	Oregon	None	None	None	• NIH	ACC/AHA Taskforce on	Defendan	None
Cigarroa	Reviewer—	Health and					Clinical Practice	t, CAD,	
	ACC/AHA	Science					Guidelines ⁺	2011+	
	Task Force on	University—					 AHA, Board of Directors, 	Defendan	
	Clinical	Clinical					Western	t, sudden	
	Practice	Professor of					Affiliate ⁺	death/CA	
	Guidelines	Medicine					 American Stroke 	D, 2010†	
							Association,		
							Cryptogenic Stroke		
							Initiative Advisory		
							Committee ⁺		
							 Catheterization and 	_	
							Cardiovascular	C Ameri	can
							Intervention ⁺	Hear	rt -
							SCAI Quality	Assoc	iation.
							Interventional Council ⁺		
William	Content	Memphis VA	None	None	None	• Lilly	 Novartis Corporation⁺ 	None	None
Cushman	Reviewer	Medical					• Takeda†		
		Center—							
		Chief,			1				
		Preventive							
		Medicine							
		Section;							
		University of							
		Tennessee							
		College of							
		Medicine—							
		Professor,							
		Medicine,							
		Preventive							
		Medicine,							
		and							
		Physiology		1	1	1			

Anita	Content	Baylor	None	None	None	• NIH *	• bAurora Health Care	None	None
Deswal	Reviewer	College of	NOTE	None	None		Inc.	None	None
—ACC/AHA		Medicine							
	Task Force on	Associate					American Heart		
	Clinical	Professor of					Association ⁺		
							AHA Committee on		
	Practice	Medicine,					Heart Failure and		
	Guidelines						Transplantation –		
							Chair†		
							 Heart Failure Society of America⁺ 		
Dave Dixon	Content	Virginia	None	None	None	None	None	None	None
	Reviewer—	Commonweal						d Amari	0.010
	Cardiovascula	th University						Ameri	can
	r Team	School of						Hear	1
		Pharmacy—						Assoc	iation.
		Associate							
		Professor	0.01/#						
Ross	Content	Winnipeg	• GSK*	None	None	None	None	None	None
Feldman	Reviewer	Regional	 Servier* 						
		Health	 Valeant 						
		Authority—	Pharmaceutica						
		Medical	ls						
		Director,	International*	_					
		Cardiac		2		1			
		Sciences							
		Program;							
		University of							
		Manitoba—							
		Professor of							
	-	Medicine							
Keith	Content	Tulane	 Amgen Inc.* 	None	None	None	 Novartis 	None	None
Ferdinand	Reviewer	University	 Boehringer 						
		School of	Ingelheim*						
		Medicine—	● Eli Lilly*						
		Professor of	 Sanofi- 						
		Clinical	Aventis*						
		Medicine	 Novartis 						
			 Quantum 						
			Genomics						
			Sanofi-						

Stephan	Content	University of	None	None	None	None	• University of	None	None
Fihn	Reviewer	Washington	None	None	None	None	Washington	None	None
	Reviewei	—Professor					Washington		
		of Medicine,							
	Heath								
		Services;							
		Division							
		Head,							
		General							
		Internal							
		Medicine;							
		Director,							
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		n; VA Puget							
		Sound Health							
		Care							
		System—							
		General		1		1			
		Internist							
Lawrence	Content	National	None	None	None	None	● NIH*	None	None
Fine	Reviewer	Heart, Lung							
		and Blood							
		Institute—							
		Chief, Clinical							
		Applications							
		and							
		Prevention							
		Branch,							
		Division of							
		Prevention							
		and							
		Population							
		Sciences							

John Flack	Content Reviewer	Southern Illinois University School of Medicine— Chair and Professor Department of Internal Medicine; Chief, Hypertension Specialty Services	 Regeneron* NuSirt 	None	None	 Bayer Healthca re Pharmac euticals[†] GSK[†] 	 American Journal of Hypertension* CardioRenal Medicine† International Journal of Hypertension† Southern Illinois University Department of Medicine* 	None	None
Joseph Flynn	Content Reviewer	Seattle Children's Hospital— Chief of the Division of Nephrology; University of Washington School of Medicine— Professor of Pediatrics	• Ultragenyx, Inc. (DSMB)	None	None	None	• UpToDate, Springer*	None Assoc	None
Federico Gentile	Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	Centro Cardiologico	None	None	None	None	None	None	None
Joel Handler	Content Reviewer	Kaiser Permanente — Physician; National Kaiser Permanente Hypertension — Clinical Leader	None	None	None	None	None	None	None

Hani Jneid	Content	Baylor	None	None	None	None	None	None	None
	Reviewer —	College of							
	ACC/AHA	Medicine—							
	Task Force on	Associate							
	Clinical Data	Professor of							
	Standards	Medicine,							
		MEDVAMC							
José A.	Content	UT	None	None	None	None	None	None	None
Joglar	Reviewer—	Southwestern							
	ACC/AHA	Medical							
	Task Force on	Center—							
	Clinical	Professor of						-	
	Practice	Internal						Amer	can
	Guidelines	Medicine;						TD Hear	rt -
		Cardiovascula							iation
		r Clinical							
		Research							
		Center—							
		Director							
Amit Khera	Content	University of	None	None	None	None	None	None	None
	Reviewer	Texas							
		Southwestern							
		Medical							
		Center—				1			
		Assistant							
		Professor of							
	2	Medicine							

	Contout	Davidan	News	News	News	News	Nexe		News
Glenn N.	Content	Baylor	None	None	None	None	None	• Defendan	None
Levine	Reviewer—	College of						t,	
ACC/AHA	Medicine-						catherizat		
	Task Force on	Professor of						ion	
	Clinical	Medicine;						laborator	
	Practice	Director,						У	
	Guidelines	Cardiac Care						procedure	
		Unit						, 2016	
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								death,	
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Giuseppe	Content	University of	Boehringer	None	None	None	 Novartis* 	None	None
Mancia	Reviewer	Milan-	Ingelheim*						
		Bicocca—	• CVRx						
		Professor of	 Ferrer 						
		Medicine;	 MEDTRONIC 						
		Chairman,	 Menarini 						
		Department	International*						
		of Clinical	 Recordati 						
		Medicine,	Servier						
		Prevention	International*						
		and Applied	Actavis						
		Biotechnologi	- / (CCUVIS						

Andrew	Content	Cardiovascula	None	None	None	Novartis	Bristol-Myers Squibb	None	None
Geriatric	Reviewer—	r Associates—				Corporat	Company		
	Cardiologist				ion†	• Janssen			
	Cardiology					• Pfizer	Pharmaceuticals, Inc.		
	Section					Inc†	• NIH		
Pamela	Content	Seinsheimer	Amgen Inc.	None	None	 Amgen 	None	None	None
Morris	Reviewer—	Cardiovascula	 AstraZeneca 			Inc.			
	Prevention	r Health	 Sanofi 						
	Council, Chair	Program—	Regeneron						
		Director;	-0						
		Women's							
		Heart Care							
		Medical						Ameri	can
		University of						TD Hear	rt -
		South							iation.
		Carolina—						-	
		Co-Director							
Martin	Content	Sunnybrook	 Ideal Life Inc* 	None	None	None	None	None	None
Myers	Reviewer	Health							
		Sciences							
		Centre—							
		Affiliate							
		Scientist;							
		University of							
		Toronto-							
		Professor,							
	2	Cardiology							
Rick	Content	Mayo Clinic	None	None	None	None	None	None	None
Nishimura	Reviewer	College of							
		Medicine—							
		Judd and							
		Mary Morris							
		Leighton							
		Professor of							
		Medicine;							
		Mayo Clinic—							
		Division of							
		Cardiovascula							
		r Diseases							

Patrick T.	Content	Harvard	None	None	None	None	MEDTRONIC	None	None
O'Gara	Reviewer—	Medical					• NIH*		
	ACC/AHA	School—							
	Task Force on	Professor of							
	Clinical	Medicine;							
	Practice	Brigham and							
	Guidelines	Women's							
		Hospital—							
		Director,							
		Strategic							
		Planning,							
		Cardiovascula							
		r Division						Ameri	can
Suzanne	Content	University of	Actelion	None	None	AstraZen	• NIH/NHLBI,	None	None
Oparil	Reviewer	, Alabama at	Lundbeck			eca	• Takeda		iation.
		Birmingham	 Novo Nordisk, 			(Duke	WHF/ESH/EPH	- A3500	ACIDITI
			Inc.			Universit	, 20.1, 21.1.		
		Distinguished	inc.			y)*			
		Professor of				• Bayer			
		Medicine;				Healthca			
		Professor of				re			
		Cell,				Pharmac			
		Development				euticals,			
		al and				Inc.*			
		Integrative		1		Novartis			
		Biology,				*			
		Division of				● NIH*			
		Cardiology							
Carl Pepine	Content	Shands	None	None	None	Capricor,	None	None	None
	Reviewer—	Hospital at				Inc.			
	CV Disease in	University of				• NIH			
	Women	Florida—				Cytori			
	Committee	Professor of				Therape			
		Medicine,				utics,			
		Chief of				Inc.			
		Cardiovascula				 Sanofi- 			
		r Medicine				Aventis			
						InVentiv			
						e Health			
						Clinical.			
						LLC			

Mahboob	Content	Case Western	None	None	None	None	None	None	None
Rahman	Reviewer	Reserve							
naman	nemener	University							
		School of							
		Medicine—							
		Professor of							
		Medicine							
Vankata	Content	UT	None	None	None	None	None	None	None
Ram	Reviewer	Southwestern							
		Medical							
		Center;							
		Apollo							
		Institute for						💪 Ameri	can
		Blood						Hear	1
		Pressure						Assoc	iation.
		Clinics						- russe	10000
Barbara	Content	University of	None	None	None	• Co-	 Novartis Corp † 	None	None
Riegel	Reviewer—	Pennsylvania				Investiga			
	ACC/AHA	School of				tor-			
	Task Force on	Nursing-				mentor†			
	Clinical	Professor				• Co-			
	Practice	6 7 6				investiga			
	Guidelines					tor NIH			
						• NIH			
						grant			
						PCORI			
Edward	Content	National	Medical	None	None	None	 American Society of 	None	None
Roccella	Reviewer	Heart, Lung,	University of				Hypertension ⁺		
		and Blood	South Carolina				 Consortium for 		
		Institute—					Southeast Hypertension		
		Coordinator,					Control ⁺		
		National High					 Consortium Southeast 		
		Blood					Hypertension Control		
		Pressure					 Inter American Society 		
		Education					of Hypertension ⁺		
		Program							

Ernesto	Content	Jewish	Novartis	Novartis	None	 Servier* 	CME Medical Grand	None	None
Schiffrin	Reviewer	General	Servier			Canadian	Rounds		
		Hospital—				Institute			
		Physician-in-				s for			
		Chief, Chief of				Health			
		the				Research			
		Department				*			
		of Medicine							
		and Director							
		of the							
		Cardiovascula							
		r Prevention							
		Centre;						Amer	can
		McGill						Hea	
		University—						Asso	iation
		Professor,						1.3501	ACT COLLO
		Department							
		of Medicine,							
		Division of							
		Experimental							
		Medicine							
Raymond	Content	University of	MEDTRONIC	None	None	• NIH*	• ASN	None	None
Townsend	Reviewer	Pennsylvania					UpToDate		
		School of							
		Medicine—							
		Professor of							
		Medicine;							
		Director,							
		Hypertension							
		Section,							
		Department							
		of Internal							
		Medicine /							
		Renal;							
		Institute for							
		Translational							
		Medicine and							
		Therapeutics							
		-Member							

Michael	Content	SUNY	Ablative	 Menarini* 	None	None	None	None	None
Weber	Reviewer	Downstate	Solutions*	Merck &					
		College of	 Allergan, Inc 	Co., Inc.*					
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Paul K. Whelton, Robert M. Carey, Wilbert S. Aronow, Donald E. Casey, Jr, Karen J. Collins, Cheryl Dennison Himmelfarb, Sondra M. DePalma, Samuel Gidding, Kenneth A. Jamerson, Daniel W. Jones, Eric J. MacLaughlin, Paul Muntner, Bruce Ovbiagele, Sidney C. Smith, Jr, Crystal C. Spencer, Randall S. Stafford, Sandra J. Taler, Randal J. Thomas, Kim A. Williams, Sr, Jeff D. Williamson and Jackson T. Wright, Jr

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Search Terms:

An extensive evidence review, which included literature derived from research involving human subjects, published in English, and indexed in MEDLINE (through PubMed), EMBASE, the Cochrane Library, the Agency for Healthcare Research and Quality, and other selected databases relevant to this guideline, was conducted between February and August 2015. Key search words included but were not limited to the following: *adherence; aerobic; alcohol intake; ambulatory care; antihypertensive: agents, drug, medication, therapy; beta adrenergic blockers; blood pressure: arterial, control, determination, devises, goal, high, improve, measurement, monitoring, ambulatory; calcium channel blockers; diet; diuretic agent; drug therapy; heart failure: diastolic, systolic; hypertension: white coat, masked, ambulatory, isolated ambulatory, isolated clinic, diagnosis, reverse white coat, prevention, therapy, treatment, control; intervention; lifestyle: measures, modification; office visits; patient outcome; performance measures; physical activity; potassium intake; protein intake; renin inhibitor; risk reduction: behavior, counseling; screening; sphygmomanometers; spironolactone; therapy; treatment: adherence, compliance, efficacy, outcome, protocol, regimen; weight. Additional relevant studies published through June 2016, during the guideline writing process, were also considered by the writing committee, and added to the evidence tables when appropriate.*

Abbreviations:

1°, primary; 2°, secondary; AASK, African American Study of Kidney Disease and Hypertension; ABI, ankle-brachial index; ABCD, Appropriate Blood Pressure Control in Diabetes; ABPM, ambulatory blood pressure monitoring; ACCESS, Acute Candesartan Cilexetil Evaluation in Stroke Survivors; ACCOMPLISH, Avoiding Cardiovascular Events Through Combination Therapy in Patients Living With Systolic Hypertension; ACCORD, Action to Control Cardiovascular Risk in Diabetes; ACE, angiotensin-converting enzyme; ACEI, angiotensin-converting enzyme inhibitor; ACS, acute coronary syndrome; ADVANCE, Action in Diabetes and Vascular Disease; AF, atrial fibrillation; AFL, atrial flutter; AHR, adjusted hazard ratio; AIPRD, Angiotensin-Converting Enzyme Inhibition in Progressive Renal Disease; ALLHAT, Antihypertensive Lipid Lowering Treatment to Prevent Heart Attack Trial; AMI, acute myocardial infarction; ARB, angiotensin-receptor blocker; ARIC, Atherosclerosis Risk in Communities; ASCOT, Anglo-Scandinavian Cardiac Outcomes Trial; BB, beta blocker; BMI, body mass index; BP, blood pressure; BPLTTC, Blood Pressure Lowering Treatment Trialists' Collaboration; bpm, beats per minute; BUN, blood urea nitrogen; CABG, coronary artery bypass graft; CAD, coronary artery disease; CATIS, China Antihypertensive Trial in Acute Ischemic Stroke; CCB, calcium-channel blocker; CCU, coronary care unit; CHD, coronary heart disease; CHF, congestive heart failure; CHHIPS, Controlling Hypertension and Hypotension Immediately Post-Stroke; CI, confidence interval; CKD, chronic kidney disease; COMFORT, Combination Pill of Losartan Potassium and Hydrochlorothiazide for Improvement of Mediation Compliance Trial; COSSACS, the Continue or Stop Post-Stroke Antihypertensives Collaborative Study; CPAP, continuous positive airway pressure; Cr, creatinine; CrCL, creatinine clearance; CRP, c-reactive protein; CR/XL, metoprolol controlled release/extended release; CT, computed tomography; CV, cardiovascular; CVD, cardiovascular disease; DASH, Dietary Approaches to Stop Hypertension; DBP, diastolic blood pressure; DM, diabetes mellitus; DM-1, diabetes mellitus type-1; DM-2, diabetes mellitus type-2; ECG, electrocardiogram; ED, emergency department; EF, ejection fraction; eGFR, estimated glomerular filtration rate; ESKD, end-stage kidney disease; ESRD, end-stage renal disease; FC, functional class; FDC, fixed dose combination; FEVER, Felodipine EVent Reduction; GITS, gastrointestinal therapeutic system; GFR, glomerular filtration rate; HBPM, home blood pressure monitoring; HCTZ, hydrochlorthiazide; HDL, high-density lipoprotein; HDL-C, high-density lipoprotein cholesterol; HEDIS, Healthcare Effectiveness Data and Information Set; HF, heart failure; HFrEF, reduced ejection fraction; HFpEF, heart failure with preserved ejection fraction; HIV, human immunodeficiency virus; HR, hazard ratio; HTN, hypertension; ICD, implantable cardioverter-defibrillator; ICH, intracerebral hemorrhage; IDACO, International Database of Ambulatory Blood Pressure in relation to Cardiovascular Outcome; IHD, ischemic heart disease; IMT, intimal media thickness; INDANA, Individual Data Analysis of Antihypertensive drug intervention trials; INTERACT2, the second Intensive Blood Pressure Reduction in Acute Cerebral Hemorrhage Trial; INVEST, International Verapamil-Trandolapril Study; INWEST, the Intravenous Nimodipine West European Stroke Trial; IQI, interquartile interval; IQR, interquartile range; IRR, incident rate ratio; ISDN, isosorbide dinitrate; IV, intravenous; JNC-7, 7th Report of the Joint National Committee; KPNC, Kaiser Permanente Northern California; LDL, low-density lipoprotein; LGSAS, low-gradient severe aortic stenosis; LIFE, Losartan Intervention For Endpoint Reduction in Hypertension; LVEF, left ventricular ejection fraction; LVH, left ventricular hypertrophy; LVMI; left ventricular mass index; MACCE, major adverse cardiac and cerebrovascular events; MACE, major adverse cardiac events; MAP, mean arterial pressure; MD, mean difference; MDPIT, Multicenter Dilitiazem Postinfarction Research Group; MDRD, Modification of Diet in Renal Disease; MERIT, Metoprolol CR/XL Randomised Intervention Trial; MESA, Multi-Ethnic Study of Atherosclerosis; MH, masked hypertension; MI, myocardial infarction; MOSES, The Morbidity and Mortality After Stroke, Eprosartan Compared With Nitrendipine for Secondary Prevention; MPR, medication possession ratio; MRFIT, Multiple Risk Factor Intervention Trial; MRI, magnetic resonance imaging; N/A, not available; NCQA, National Committee for Quality Assurance; NEMESIS, North East Melbourne Stroke Incidence Study; NHANES, National Health and Nutrition Examination Surveys; NIH, National Institute of Health; NNT, number needed to treat; NR, not relative;

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NS, nonsignificant; NSAID, nonsteroidal anti-inflammatory drug; NUTRICODE, Nutrition and Chronic Diseases Expert Group; NYHA, New York Heart Association; ONTARGET, Ongoing Telmisartan Alone and in Combination With Ramipril Global Endpoint Trial; OR, odds ratio; OSA, obstructive sleep apnea; P4P, pay for performance; PA, pulmonary artery; PAD, peripheral artery disease; PAMELA, Pressione Arteriose Monitorate E Loro Associazioni; PCP, primary care provider; periop, perioperative; PREDIMED, Prevention with a Mediterranean Diet; PRISMA, Preferred Reporting Items for Systematic Reviews and Meta-Analyses; PROBE, Prospective, randomized, open, blinded endpoint; PROGRESS, The perindopril protection against recurrent stroke study; PRONTO, Prospective Optical Coherence Tomography Imaging of Patients with endovascular Age-Related Macular Degeneration Treated with Intraocular Ranibizumab; pt, patient; PTCA, percutaneous transluminal coronary angioplasty; PVD, peripheral vascular disease; QI, quality improvement; RAAS, renin angiotensin aldosterone system; RCT, randomized controlled trial; REIN-2, Blood Pressure Control for Renoprotection in Patients with Non-diabetic Renal Disease; RH, relative hazard; ROADMAP, Randomized Olmesartan and Diabetes Microalbuminuria Prevention; RR, relative risk; Rx, medical prescription; SAE, severe adverse event; SBP, systolic blood pressure; SCOPE-AS, Symptomatic Cardiac Obstruction – Pilot Study of Enalapril in Aortic Stenosis; SD, standard deviation; SE, stress echocardiography; SH, sustained hypertension; SHEP, Summer Health Enrichment; SITS-ISTR, Safe Implementation of Thrombolysis in Stroke-International Stroke Thrombolysis Register; SKIPOGH, Swiss Kidney Project on Genes in Hypertension; SPC, single pill combination; SPRINT, Systolic Blood Pressure Intervention Trial; Syst-Eur, Systolic Hypertension in Europe; t-PA, tissue plasminogen activator; TIA, transient ischemic attack; TOHP, Trials of Hypertension Prevention; TOMHS, Treatment of Mild Hypertension Study; TONE, Trial of Nonpharmacologic Intervention in the Elderly; TOPCAT, Treatment of Preserved Cardiac Function Heart Failure With Aldosterone Antagonist; TR, target range; UA, unstable angina; U.K., United Kingdom; UKPDS, United Kingdom Prospective Diabetes Study; U.S., United States; VA, Veterans Affairs; VA Coop; Veterans Administration Cooperative Study Group on Antihypertensive Agents; VA NEPHRON-D, Veterans Affairs Nephropathy in Diabetes; VALIANT, Valsartan in Acute Myocardial Infarction Trial; VALUE, Valsartan Antihypertensive Long-term Use Evaluation; WCH, white coat hypertension; and WPW; Wolff-Parkinson-White syndrome.

Data Supplement 1. Coexist	ence of Hypertension	and Related Chronic	Conditions (Section 2.4)
Data Supplement 1. COEXIST	ence of hypertension	and Related Children	

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR RR; & 95% CI)	Summary/Conclusion Comment(s)
Wilson PW, et al., 1999 (1) <u>10335688</u>	Study type: Nonrandomized Size: 2,406 men, 2,569 women (1,759 men, 1,818 women with follow-up)	Inclusion criteria: Men and women 18–74 y and free of CHD at baseline, from the Framingham Offspring Study Exclusion criteria: N/A	<u>1° endpoint</u> : Total CHD (first occurrence of angina, UA, MI, and coronary death), Hard CHD (first MI and coronary death) <u>Results:</u> Presence of ≥3 risk factors was associated with a 2.39 times greater risk of CHD in men (95% CI: 1.56–3.36; p<0.001) and a 5.90 increased risk of CHD in women (95% CI: 2.54–13.73; p<0.001)	• CVD risk factors infrequently occur in isolation (only 28%–30% of the time); presence of ≥3 risk factors occurred 17% of the time in both men and women; presence of ≥3 risk factors associated with high risk of CHD and coronary death (attributable risk of 20% in men and 48% in women)
Berry JD, et al., 2012 (2) <u>22276822</u>	Study type: Nonrandomized Size: 257,384 black and white men and women, including 67,890 pts (from 17 meta-analysis) and 189,494 pts (from MRFIT)	Inclusion criteria: Meta- analysis of 18 cohort studies Exclusion criteria: N/A	<u>1° endpoint</u> : Fatal CHD, nonfatal MI, fatal or nonfatal stroke <u>Results:</u> Participants with optimal RF profile (total cholesterol <180 mg/dL, untreated BP <120 mm Hg systolic, and <80 mm Hg diastolic, nondiabetic, nonsmoker) compared to participants with ≥2 risk factors had lower risk of CVD through the age of 80 y (4.7% vs. 29.6% for men, 6.4% vs. 20.5% for women), lower lifetime risk of fatal heart disease and nonfatal MI (3.6% vs. 37.5% for men, <1% vs. 18.3% for women), and lower lifetime risk of fatal or nonfatal stroke (2.3% vs. 8.3% for men, 5.3% vs. 10.7% for women)	• Increased burden of 80 risk factors associated with higher lifetime risk of CVD

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; and CI; & 95% CI)	Summary/Conclusion Comment(s)
Lewington S, et al., 2002 <u>12493255</u>	Study type: Meta-analysis of 61 observational cohort studies	Inclusion criteria: Men and women with no history of previous CVD and record of key study variables. Exclusion criteria: Prior CVD	1° endpoint: Cause-specific mortality Results: 958,074 persons followed for a mean of 12 y to death (12.7 million person-y at risk. Number of deaths attributed to: -Stroke: 11960 -IHD: 34,283 -Other vascular:10092 -Non-vascular: 60797 Above a SBP ≥115 mm Hg and DBP ≥75 mm Hg, there was a progressive rise in vascular death with progressively high BP with no evidence of a J-curve (approximately doubling of stroke and IHD mortality for a 20 mm Hg higher level of SBP or 10 mm Hg higher level of DBP, in those 40–69 y). With progressively higher age, the BP-related proportional risk of vascular mortality was somewhat reduced but the corresponding absolute risk was much higher.	• In adults aged 40–89 y, usual BP is strongly related to vascular (and overall) mortality, without evidence of a threshold down to at least an SBP/DBP of 115/75 mm Hg.
Rapsomaniki E, et al., 2014 <u>24881994</u>	Study type: Observational cohort study Size: 1.25 million patients, in 225 primary care practices in the UK, followed for a median of 5.2 y using electronic medical records.	Inclusion criteria: Men and women ≥30 y, with no previous diagnosis of CVD, who had been registered at their practices for ≥1 year. Exclusion criteria: N/A	<u>1° endpoint</u> : 12 acute and chronic CVD outcomes <u>Results:</u> 83,098 initial CVD events recorded. Within each of 3 age groups (30–59, 60–79, and ≥80 y), the lowest risk for CVD was in those with a SBP 90–114 mm Hg and DBP 60–74 mm Hg. There was a direct relationship between level of BP and most CVD outcomes, with no evidence of J-curve, with the strongest relationship for SBP and stroke and weakest for abdominal aneurysm.	• Despite modern treatments, the lifetime burden of BP- related CVD was substantial.
Wilson PW, et al., 1999 (1) <u>10335688</u>	Study type: Nonrandomized	Inclusion criteria: Men and women 18– 74 y and free of CHD at baseline, from the Framingham Offspring Study	<u>1° endpoint</u> : Total CHD (first occurrence of angina, UA, MI, and coronary death), Hard CHD (first MI and coronary death)	• CVD risk factors infrequently occur in isolation (only 28%–30% of the time)

Data Supplement 2. Definition of High BP (Section 3.1)
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	Size: 2,406 men, 2,569 women (1,759 men, 1,818 women with follow-up)	Exclusion criteria: N/A	Results: Presence of ≥3 risk factors was associated with a 2.39 times greater risk of CHD in men (95% CI: 1.56–3.36; p<0.001) and a 5.90 increased risk of CHD in women (95% CI: 2.54– 13.73; p<0.001)	 Presence of ≥3 risk factors occurred 17% of the time in both men and women Presence of ≥3 risk factors associated with high risk of CHD and coronary death (attributable risk of 20% in men and 48% in women)
Guo X, et al., 2013 (3) <u>23634212</u>	<u>Study type</u> : Meta- analysis of nonrandomized studies <u>Size</u> : 870,678 pts	Inclusion criteria: Studies reporting adjusted risk for CVD or mortality with pre- HTN Exclusion criteria: N/A	<u>1° endpoint</u> : CVD and all-cause mortality <u>Results:</u> SBP/DBP 120–129/80–84 mm Hg compared to <120/80 mm Hg: • All-cause mortality: RR: 0.91; 95% CI: 0.81– 1.02) • CVD mortality: RR: 1.10 (95% CI: 0.92, 1.30) SBP/DBP 130–139/85–89 mm Hg compared to <120/80 mm Hg: • All-cause mortality: 1.00; 95% CI: 0.95–1.06 • CVD mortality: DD: 1.2(+0.5%) CI: 1.12, 1.41	 SBP/DBP of 120–129/80– 84 mm Hg associated with increased risk for all-cause or CVD mortality. SBP/DBP of 130–139/85– 89 mm Hg associated with an increased risk for CVD mortality.
Guo X, et al., 2013 (4) <u>24234576</u>	<u>Study type</u> : Meta- analysis of nonrandomized studies <u>Size</u> : 1,010,858 pts	Inclusion criteria: Studies reporting adjusted risk for fatal and nonfatal stroke, CHD, MI and total CVD events with pre- HTN, 120–129/80–84 mm Hg or 130– 139/85–89 mm Hg Exclusion criteria: N/A	 CVD mortality: RR: 1.26; 95% CI: 1.13–1.41 <u>endpoint</u>: Fatal and nonfatal stroke, CHD, MI and total CVD events <u>Results:</u> SBP/DBP 120-129/80-84 mm Hg compared to <120/80 mm Hg: CVD RR: 1.24; 95% CI: 1.10–1.39 MI RR: 1.43; 95% CI: 1.10–1.86 Stroke: RR: 1.35; 95% CI: 1.10–1.66 SBP/DBP 130–139/85–89 mm Hg compared to <120/80 mm Hg: CVD RR: 1.56; 95% CI: 1.36–1.78 MI RR: 1.99; 95% CI: 1.69–2.24 	• Compared to pts with SBP/DBP<120/80 mm Hg, the RR for CVD, MI and stroke were larger for pts with SBP/DBP of 130–139/85–89 mm Hg vs. SBP/DBP of 120– 129/80–84 mm Hg.
Huang Y, et al., 2013 (5) <u>23915102</u>	Study type: Meta- analysis of nonrandomized studies Size: 468,561 pts from 18 prospective cohort studies	Inclusion criteria: Studies reporting risk for CVD, CHD and stroke, with 120– 139/80–89 mm Hg, 120–129/80–84 mm Hg or130–139/85–89 mm Hg Adults ≥18 y BP evaluated at baseline	<u>1° endpoint</u> : CVD, CHD, and stroke <u>Results:</u> Comparing SBP/DBP 120–129/80–84 mm Hg to <120/80 mm Hg:	• Compared to pts with SBP/DBP <120/80 mm Hg, the RR for CVD was larger for pts with SBP/DBP of 130– 139/85–89 mm Hg vs.

		≥2 y follow-up for outcomes Results reported with adjustment <u>Exclusion criteria</u> : N/A	Comparing SBP/DBP RR: 130–139/85–89 mm Hg to <120/80 mm Hg: • CVD RR: 1.63; 95% CI: 1.47–1.80; p value comparing these risk ratios=0.02 • The RR comparing CHD and stroke by levels of SBP/DBP: 130–139/85–89 mm Hg and SBP/DBP of 120–129/80–84 mm Hg vs. <120/80 mm Hg were not reported.	SBP/DBP of 120–129/80–84 mm Hg
Huang Y, et al., 2014 (6) <u>24074825</u>	Study type: Meta- analysis of nonrandomized studies Size: 1,003,793 pts were derived from 6 prospective cohort studies	Inclusion criteria: Studies reporting adjusted risk for ESRD with 120–139/80– 89 mm Hg, 120–129/80–84 mm Hg or130–139/85–89 mm Hg Adults ≥18 y BP evaluated at baseline ≥ 1 y follow-up for ESRD Results reported with adjustment Exclusion criteria: 1) enrollment depended on having a condition or risk factor, 2) the study reported only age- and sex-adjusted RRs, and 3) data were derived from the same cohort or from a 2° analysis	<u>1° endpoint</u> : ESRD <u>Results:</u> Comparing SBP/DBP 120–129/80–84 mm Hg to <120/80 mm Hg: • ESRD RR: 1.44; 95% CI: 1.19–1.74 Comparing SBP/DBP 130–139/85–89 mm Hg to <120/80 mm Hg: • ESRD RR: 2.02; 95% CI: 1.70–2.40; • p value comparing these risk ratios=0.01	• Compared to pts with SBP/DBP <120/80 mm Hg, the RR for ESRD was larger for pts with SBP/DBP of 130– 139/85–89 mm Hg vs. SBP/DBP of 120–129/80–84 mm Hg
Huang Y, et al., 2013 (7) <u>24623843</u>	<u>Study type</u> : Meta- analysis of nonrandomized studies <u>Size</u> : 762,393 pts from 19 prospective cohort studies	Inclusion criteria: Studies reporting adjusted risk for stroke with 120–139/80– 89 mm Hg, 120–129/80-84 mm Hg or130– 139/85–89 mm Hg • Adults ≥18 y • BP evaluated at baseline • ≥1 y follow-up for stroke • Results reported with adjustment Exclusion criteria: • Enrollment depended on having a specific risk factor condition (e.g., DM or other baseline chronic diseases) • The RR was unadjusted or only adjusted for age and sex • Data were derived from the same cohort or meta-analysis of other cohort studies.	<u>1° endpoint</u> : Stroke <u>Results:</u> Comparing SBP/DBP 120–129/80–84 mm Hg to <120/80 mm Hg: • Stroke: RR: 1.44; 95% CI: 1.27–1.63 Comparing SBP/DBP 130–139/85–89 mm Hg to <120/80 mm Hg: • Stroke: RR: 1.95; 95% CI: 1.73–2.21 • p value comparing these risk ratios ≤0.001	• Compared to pts with SBP/DBP <120/80 mm Hg, the RR for stroke was larger for pts with SBP/DBP of 130– 139/85–89 mm Hg vs. SBP/DBP of 120–129/80–84 mm Hg

Huang Y, et al., 2014 (8) <u>24439976</u>	Study type: Meta- analysis of nonrandomized studies Size: 1,129,098 pts from 20 prospective cohort studies	Inclusion criteria: • Studies reporting adjusted risk for all-cause/CVD mortality with 120–139/80–89 mm Hg, 120-129/80–84 mm Hg or 130–139/85–89 mm Hg • Adults ≥18 y • BP evaluated at baseline • ≥2 y follow-up for mortality • Results reported with adjustment Exclusion criteria: • Enrollment depended on having a specific risk factor condition (e.g., DM or other baseline chronic diseases) • The RR was unadjusted or only adjusted for age and sex • Data were derived from the same cohort or meta-analysis of other cohort studies.	<u>1° endpoint</u> : All-cause and CVD mortality <u>Results:</u> Comparing SBP/DBP 120–129/80-84 mm Hg to <120/80 mm Hg: • All-cause mortality RR: 0.96; 95% CI: 0.85–1.08 • CVD mortality RR: 1.08; 95% CI: 0.98–1.18 Comparing SBP/DBP 130-139/85-89 mm Hg to <120/80 mm Hg: • All-cause mortality RR: 1.03; 95% CI: 0.95–1.12 • CVD mortality RR: 1.28; 95% CI: 1.16–1.41 • p value comparing these risk ratios: • All-cause mortality p=0.33 • CVD mortality p=0.01	 Compared to pts with SBP/DBP <120/80 mm Hg, the RR for CVD mortality was larger for pts with SBP/DBP of 130–139/85-89 mm Hg vs. SBP/DBP of 120–129/80-84 mm Hg. The RR for not all-cause mortality was similar for these 2 BP levels.
Huang Y, et al., 2015 (9) <u>25699996</u>	Study type: Meta- analysis of nonrandomized studies Size: 591,664 pts from 17 prospective cohort studies	Inclusion criteria: • Studies reporting adjusted risk for CHD with 120–139/80–89 mm Hg, 120–129/80– 84 mm Hg or130–139/85–89 mm Hg • Adults ≥18 y • BP evaluated at baseline • Results reported with adjustment Exclusion criteria: • Enrollment depended on having a specific risk factor condition (e.g., DM or other baseline chronic diseases) • The RR was unadjusted or only adjusted for age and sex • Data were derived from the same cohort or meta-analysis of other cohort studies.	<u>1° endpoint</u> : CHD <u>Results:</u> Comparing SBP/DBP 120–129/80–84 mm Hg to <120/80 mm Hg: • CHD RR: 1.27; 95% CI: 1.07–1.50 Comparing SBP/DBP 130-139/85-89 mm Hg to <120/80 mm Hg: • CHD RR: 1.58; 95% CI: 1.24–2.02 • p value comparing these RR: 0.15	 Compared to pts with SBP/DBP<120/80 mm Hg, the RR for CHD was larger for pts with SBP/DBP of 130– 139/85–89 mm Hg vs. SBP/DBP of 120-129/80–84 mm Hg. However, this difference was not statistically significant.
Lee M, et al., 2011 (10) <u>21956722</u>	Study type: Meta- analysis of nonrandomized studies	 Inclusion criteria: Studies reporting adjusted risk for stroke with 120–139/80–89 mm Hg, 120–129/80–84 mm Hg or130–139/85–89 mm Hg Adults ≥18 y 	<u>1° endpoint</u> : Incident stroke <u>Results:</u> Comparing SBP/DBP 120–129/80–84 mm Hg to <120/80 mm Hg: • Stroke RR: 1.22; 95% CI: 0.95–1.57	• Compared to pts with SBP/DBP <120/80 mm Hg, the RR for stroke was larger for pts with SBP/DBP of 130– 139/85–89 mm Hg vs.

	Size: 518,520 pts from 18	BP evaluated at baseline	Comparing SBP/DBP 130–139/85–89 mm Hg to	SBP/DBP of 120-129/80-84
	prospective cohort studies	Results reported with adjustment	<120/80 mm Hg:	mm Hg
			• Stroke RR: 1.79; 95% CI: 1.49–2.16	
		Exclusion criteria:		
		Cross-sectional, case-control or		
		retrospective cohort		
		• The RR was unadjusted or only adjusted		
		for age and sex • 95% CI not reported		
		 Data were derived from the same cohort 		
		or meta-analysis of other cohort studies		
		Results from trial of antihypertensive		
		medication		
Shen L, et al.,	Study type: Meta-	Inclusion criteria:	<u>1° endpoint</u> : CHD	 Compared to pts with
2013 (11)	analysis of	 Studies reporting adjusted risk for CHD 		SBP/DBP <120/80 mm Hg,
<u>23608614</u>	nonrandomized studies	with 120–139/80–89 mm Hg, 120–129/80–	Results: Comparing SBP/DBP 120–129/80–84	the RR for CHD was larger for
	Size: $0.24, 106$ ptc from 10	84 mm Hg or 130–139/85–89 mm Hg	mm Hg to <120/80 mm Hg:	pts with SBP/DBP of 130–
	Size: 934,106 pts from 18 prospective cohort studies	BP evaluated at baseline	• CHD RR: 1.16; 95% CI: 0.96–1.42	139/85–89 mm Hg vs. SBP/DBP of 120–129/80–84
	prospective conort studies	 95% CI was reported 	Comparing SBP/DBP 130–139/85–89 mm Hg to <pre><pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre></pre><pre><td>SBP/DBP 01 120-129/80-84 mm Hg</td></pre></pre>	SBP/DBP 01 120-129/80-84 mm Hg
		Exclusion criteria: N/A	• CHD RR: 1.53; 95% CI: 1.19–1.97)	mining
Wang S, et al.,	Study type: Meta-	Inclusion criteria:	1° endpoint: CVD, CVD mortality, all-cause	Compared to pts with
2013 (12)	analysis of	Prospective cohort studies reporting risk	mortality	SBP/DBP<120/80 mm Hq, RR
<u>23932039</u>	nonrandomized studies	for outcomes with 120–139/80–89 mm Hg	,	for CVD and CVD mortality
		Pts free of CVD at baseline,	Results: Comparing SBP/DBP 120-129/80-84	were larger for pts with
	Size: 396,200 pts from 13	 Follow-up ≥5 y 	mm Hg to <120/80 mm Hg:	SBP/DBP of 130–139/85–89
	prospective cohort studies	 Adjusted results reported 	• CVD RR: 1.41; 95% CI: 1.25–1.59	mm Hg vs. SBP/DBP of 120-
		 95% CI was reported 	• CVD mortality RR: 1.18; 95% CI: 0.98–1.42	129/80–84 mm Hg.
			• All-cause mortality RR: 0.99; 95% CI: 0.88–1.13	No difference in all-cause mortality was present across
		Exclusion criteria: N/A	Comparing SBP/DBP 130–139/85–89 mm Hg to	mortality was present across BP levels.
			<120/80 mm Hg: CVD RR: 1.74; 95% CI: 1.51–2.01 	וסעסו.
			• CVD RR: 1.74; 95% CI: 1.51–2.01 • CVD mortality RR: 1.33; 95% CI: 1.13–1.58	
			 All-cause mortality RR: 1.02; 95% CI: 0.97–1.08 	
Cushman WC, et	Study type: 2° analysis of	Inclusion criteria: Men and women ≥55 y	1° endpoint: Achieving SBP/DBP<140/90 mm	• BP control (<140/90 mm
al., 2002 (13)	an RCT	with HTN and 1 additional CHD risk factor	Hg, use of ≥2 drug classes	Hg) can be achieved in most
<u>12461301</u>				pts ≥2 or more drug classes
	Size: 33,357 pts in the	Exclusion criteria: Pts randomized to		are often required.
	ALLHAT	doxazosin.		

			Results: SBP/DBP control was achieved by 66% at 5 y of follow-up and 63% of pts were on ≥2 drug classes.	
Dalhof B, et al., 2002 (14) <u>11937178</u>	<u>Study type</u> : RCT <u>Size</u> : 9,193 pts 55–80 y in the Losartan Intervention For Endpoint reduction in HTN	Inclusion criteria: Men and women with ECG signs of LVH. Trough sitting SBP 160–200 mm Hg or DBP 95–115 mm Hg after 1–2 wk of placebo. Exclusion criteria: 2° HTN, MI/stroke within 6 mo, angina, HF or LVEF <40%.	<u>1° endpoint</u> : Following a titration schedule to reach a target SBP/DBP<140/90 mm Hg <u>Results:</u> Mean SBP/DBP at baseline was 174/98 mm Hg. Over 90% of pts required ≥2 drug classes during follow-up.	• Pts with a mean SBP/DBP of 160–200/95–115 mm Hg will need ≥2 classes of antihypertensive medication to achieve SBP/DBP <140/90 mm Hg.
Wald DS, et. al., 2009 (15) <u>19272490</u>	Study type: Meta- analysis of RCT Size: 10,968 pts in 42 trials of factorial designs comparing monotherapy, combination therapy and placebo.	Inclusion criteria: Randomized placebo- controlled trials comparing 2 of 4 (thiazides, BB s, ACEIs, and CCB) drug classes. Exclusion criteria: Trials <2 wk duration, no placebo group, nonrandomized order of treatment.	<u>1° endpoint</u> : Mean BP reduction. <u>Results:</u> Combination therapy vs. monotherapy produced larger SBP reductions: • Thiazide alone (7.3 mm Hg) • Thiazide+second drug class (14.6 mm Hg) • BB alone (9.3 mm Hg) • BB +second drug class (18.9 mm Hg) • ACE-inhibitor alone (6.8 mm Hg) • ACE-inhibitor+second drug class (13.9 mm Hg) • CCB alone (8.4 mm Hg) • CCB +second drug class (14.3 mm Hg)	• Combination therapy results in substantially larger SBP and DBP reductions compared with monotherapy, even after dose titration.
Lewington S, et al., 2002 (16) <u>12493255</u>	Aim: To describe the age- specific relevance of BP to cause-specific mortality Study type: Meta- analysis of cohort studies Size: 61 prospective studies with 12.7 million person-y of observation, 56,000 vascular deaths in 40–89 y.	Inclusion criteria: Collaboration was sought from the investigators of all prospective observational studies in which data on BP, blood cholesterol, date of birth (or age), and sex had been recorded at a baseline screening visit, and in which cause and date of death (or age at death) had been routinely sought for all screens during more than 5,000 person-y of follow- up (see appendix A). Relevant studies were identified through computer searches of Medline and Embase, by hand- searches of meeting abstracts, and by extensive discussions with investigators. Exclusion criteria: To minimize the effects of reverse causality (whereby	 1° endpoint: Not completely clear, but for our purposes, stroke and IHD death would be co-1°. Also looked at other vascular deaths. HRs for stroke mortality for a 20 mm Hg lower SBP by age-group 40–49: 0.36 (95% CI: 0.32–0.40) 50–59: 0.38 (95% CI: 0.35–0.40) 60–69: 0.43 (95% CI: 0.41–0.45) 70–79: 0.50 (95% CI: 0.48–0.52) 80–89: 0.67 (95% CI: 0.63–0.71) HRs for IHD mortality for a 20 mm Hg lower SBP by age-group 40–49: 0.49 (95% CI: 0.45–0.53) 50–59: 0.50 (95% CI: 0.49–0.52) 60–69: 0.54 (95% CI: 0.49–0.52) 60–69: 0.54 (95% CI: 0.53–0.55) 70–79: 0.60 (95% CI: 0.58–0.61) 	• Throughout middle and old age, usual BP is strongly and directly related to vascular (and overall) mortality, without any evidence of a threshold down to at least 115/75 mm Hg.

		established disease could change the usual BP), studies were excluded if they had selected pts on the basis of a positive history of stroke or heart disease, and individuals from contributing studies were excluded from the present analyses if they had such a history recorded at baseline.	80–89: 0.67 (95% CI: 0.64–0.70) • HRs for other vascular mortality for a 20 mm Hg lower SBP by age-group 40–49: 0.43 (95% CI: 0.38–0.48) 50–59: 0.50 (95% CI: 0.47–0.54) 60–69: 0.53 (95% CI: 0.51–0.56) 70–79: 0.64 (95% CI: 0.61–0.67) 80–89: 0.70 (95% CI: 0.65–0.75) • Similar results for DBP also in figure 1. • Similar results for men and women separately for stroke, figure 3, and IHD, figure 5.	
Ettehad D, et al., 2016 (17) <u>26724178</u>	Aim: This systematic review and meta-analysis aims to combine data from all published large-scale BP-lowering trials to quantify the effects of BP reduction on CV outcomes and death across various baseline BP levels, major comorbidities, and different pharmacological interventions. <u>Study type:</u> Meta- analysis of RCTs <u>Size:</u> 123 studies with 613,815 pts	 Inclusion criteria: RCTs of BP-lowering treatment that included a minimum of 1,000 pt-y of follow-up in each study arm. No trials were excluded because of presence of baseline comorbidities, and trials of antihypertensive drugs for indications other than HTN were eligible. Eligible studies fell into 3 categories: 1st, random allocation of pts to a BP-lowering drug or placebo; 2nd, random allocation of pts to different BP-lowering drugs; and third, random allocation of pts to different BP-lowering targets. Exclusion criteria: 		

Law MR, et al., 2009 (18) <u>19454737</u>	Study type: Meta- analysis of use of BP- lowering drugs in prevention of CVD from 147 randomized trials Size: Of 147 randomized trials of 464,000 pts, 37 trials of BBs in CAD included 38,892 pts, and 37 trials of other antihypertensive drugs in CAD included 85,395 pts	Inclusion criteria: The database search used Medline (1966 to Dec. 2007) to identify randomized trials of BP-lowering drugs in which CAD events or strokes were recorded. The search also included the Cochrane Collaboration and Web of Science databases and the citations in trials and previous meta-analyses and review articles. Exclusion criteria: Trials were excluded if there were <5 CAD events and strokes or if treatment duration was <6 mo.	 CVD- 0.74 (95% CI: 0.67–0.83) Total deaths CVD+ 0.90 (95% CI: 0.83–0.98) CVD- 0.84 (95% CI: 0.75–0.93) Other outcomes similarly in figure 5 In appendix, in general, benefits for CVD prevention seen in groups with and without baseline CHD, Stroke, DM, CKD and HF when examined separately, but no absolute risks provided to enable estimation of how far down the absolute risk curve these findings have been demonstrated. Some evidence of BB inferiority to other med classes in figure 6. Did not report absolute risks so do not know lower level of risk in treated populations. 1° endpoint: CAD events; stroke Results: In 37 trials of pts with a history of CAD, BB reduced CAD events 29% (95% CI: 22%– 34%). In 27 trials in which BBs were used after acute MI, BB reduced CAD events 31% (95% CI: 24%–38%), and in 11 trials in which BB were used after long-term CAD, BB insignificantly reduced CAD events 13%. In 7 trials, BB reduced stroke 17% (95% CI: 1%–30%). CAD events were reduced 14% (95% CI: 2%–25%) in 11 trials of thiazide diuretics, 17% (95% CI: 11%–22%) in 21 trials of ACEIs, insignificantly 14% in 4 trials of angiotensin receptor blockers, and 15% (95% CI: 8%–22%) in 22 trials of CCB. Stroke was reduced 38% (95% CI: 28%–47%) in 10 trials of thiazide diuretics, 22% (95% CI: 25%–42%) in 9 trials of ACEI, and 34% (95% CI: 25%–42%) in 9 trials of CCB. 	 Lack of individual pt data, which would have allowed a more reliable assessment of treatment effects in different pt groups. Interpretation: Lowering of BP into what has been regarded the normotensive range should therefore be routinely considered for the prevention of CVD among those deemed to be of sufficient absolute risk. With the exception of the extra protective effect of BB given shortly after a MI and the minor additional effect of CCBs in preventing stroke, all the classes of BP-lowering drugs have a similar effect in reducing CAD events and stroke for a given reduction in BP.
Sundstrom J, et al., 2015 (19) <u>25531552</u>	Aim: To investigate whether pharmacologic BP reduction prevents CV events and deaths in pts with grade 1 HTN.	Inclusion criteria: RCTs of at least 1 y duration; pts ≥18 y, at least 80% of whom had grade 1 HTN and no previous CVD (MI, angina pectoris, CABG, PCI, stroke, TIA, carotid surgery, peripheral arterial	<u>1° endpoint</u> : Total major CV events, comprising stroke (nonfatal stroke or death from cerebrovascular disease), coronary events (nonfatal MI or death from CHD, including sudden death), HF (causing death or resulting in	• BP-lowering therapy is likely to prevent stroke and death in pts with uncomplicated grade 1 HTN.

	<u>Study type:</u> Meta- analysis of RCTs <u>Size:</u> 10 RTCs with 15,266 pts	surgery, intermittent claudication, or renal failure); and compared an antihypertensive drug provided as monotherapy or a stepped-care algorithm vs. placebo or another control regimen. Exclusion criteria: Excluded trials did not contribute an event for any of the outcomes of interest.	hospitalization), or CV death; OR: 0.86 (95% CI: 0.74–1.01) Other endpoints: Each of the above outcomes independently; and total deaths. CHD 0.91 (95% CI: 0.74–1.12) Stroke 0.72 (95% CI: 0.55–0.99) HF 0.80 (95% CI: 0.57–1.12) CVD deaths 0.75 (95% CI: 0.57–0.98) Total deaths 0.78 (95% CI: 0.67–0.92) Only the first event for a pt was used for the analysis of each outcome, but a pt who had >1 outcome type could contribute to more than 1 analysis. They also tabulated overall withdrawals and withdrawals due to adverse events.	• 5 y risks in BPLTTC control groups CVD events 7.4%, CVD deaths 3.1%
Thomopoulos C, et al., 2014 (20) <u>25259547</u>	Aim: Investigating whether all grades of HTN benefit from BP-lowering treatment and which are the target BP levels to maximize outcome reduction. Study type: Meta- analysis of RCTs Size: 32 RCTs with 104,359 pts	Inclusion criteria: Intentional BP-lowering comparing active drug treatment with placebo, or less active treatment (intentional BP-lowering trials), or comparison of an active drug with placebo over baseline antihypertensive treatment, resulting in a BP difference of at least 2 mm Hg in either SBP or DBP (nonintentional BP-lowering trials); enrolling of hypertensive individuals only or a high proportion (at least 40%) of them. Exclusion criteria: N/A	 <u>1° endpoint</u>: As some trials were done on low-risk pts, others on higher risk pts, no evaluation of absolute risk-reduction was made. However, a 2° analysis was done including trials or trial subgroups with mean baseline SBP/DBP values in grade 1 range and a low-to-moderate risk (<5% CV deaths in 10 y in controls): FEVER stratum with baseline SBP below the median (<153 mm Hg); HTN Detection and Follow-up Program stratum with baseline DBP 90–94 mm Hg and no CVD; OSLO (e17); TOMHS (e28) and USPHS. Risks of stroke, CHD, the composite of stroke and CHD, and all-cause death were significantly reduced by BP-lowering in these low-to-moderate risk pts (control group: average CV mortality 4.5% in10 y) with a moderate BP elevation (average SBP/DBP 145.5/91 mm Hg) at randomization. Standardized RR associated with 10/5 reduction in BP: stroke 0.33 (95% CI: 0.11–0.98) CHD 0.68 (95% CI: 0.48–0.95) 	 Meta-analyses favor BP-lowering treatment even in grade 1 HTN at low-to-moderate risk, and lowering SBP/DBP to <140/90 mm Hg. Achieving <130/80 mm Hg appears safe, but only adds further reduction in stroke.

CVD death 0.57 (95% CI: 0.32–1.02) total death 0.53 (95% 0.35–0.80)	
Compared outcomes of achieved on study SBP	
<130 vs. ≥130	
Standardized Risk ratio associated with 10/5	
reduction in BP: stroke 0.68 (95% CI: 0.57, 0.83)	
CHD 0.87 (95% CI: 0.76, 1.00)	
HF 0.92 (95% CI: 0.47, 1.77)	
CVD 0.81 (95% CI: 0.67, 1.00)	
CVD death 0.88 (95% CI: 0.77, 1.01) total death	
0.88 (95% CI: 0.77, 0.99)	
Outcomes of achieved on study SBP 130-139	
vs. ≥140	
Standardized RR associated with 10/5 reduction	
in BP: stroke 0.63 (95% CI: 0.52, 0.77)	
CHD 0.77 (95% CI: 0.72, 0.77)	
HF 0.76 (95% CI: 0.47, 1.25)	
CVD 0.74 (95% CI: 0.62, 0.88)	
CVD 0.74 (75 % Cl. 0.02, 0.00) CVD death 0.81 (95% Cl. 0.67, 0.97) total death	
0.87 (95% CI: 0.75, 1.00)	
Similar pattern of results for on treatment DBP.	
	BP-lowering,
	\sim <130 mm Hg,
	reater vascular
	than standard
targets or different BP changes from ESKD, and adverse events. Progression of regimens.	
	sk pts, there are
	cenefits from more
	P-lowering,
	or those with SPB
	Hg at baseline.
	absolute benefits of
	P-lowering in high-
	uals are large.
Other endpoints:	uais ale lalye.
<u>Comparator</u> : MI RR: 0.87 (95% CI: 0.76–1.00) p=0.042 Limitation	ç.
	<u>s.</u> ndividual pt data,
	Id have allowed a
	ble assessment of
levels in the less intensive BP-lowering Total deaths RR: 0.91 (95% CI: 0.81–1.03)	no assessment of

regimen group were 140/81 mm Hg, compared with 133/76 mm Hg in the more intensive treatment group.	Other results:• Benefit for CVD not different by baseline SBP120–139: 0.89 (95% CI: 0.76–1.05)140–160: 0.83 (95% CI: 0.68–1.00)>160: 0.89 (95% CI: 0.73–1.09)p-heterogeneity: 0.60• Benefit for CVD not different for more intensiveand less intensive targets in intensive group<140 or <150 mm Hg: 0.76 (95% CI: 0.60–0.97)<120– <130 mm Hg: 0.91 (95% CI: 0.84–1.00; p-hetero: 0.06)• Absolute benefits were proportional to absoluterisk.• For trials in which all pts had vascular disease,renal disease, or DM at baseline, the averagecontrol group rate of major vascular events was2.9% per y compared with 0.9% per y in othertrials, and the numbers needed to treat were 94(95% CI: 44–782) in these trials vs. 186 (95% CI:107–708) in all other trials.	treatment effects in different pt groups. • Interpretation: Supports treating pt with and without CVD at threshold of 130 to <130. Supports treating at threshold of about 130 even down to a CVD event rate of 0.9% per y.
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Data Supplement 3. Out-of-Office and Self-Monitoring of BP (Section 4.2)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population (N)	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Pickering TG, et al., 1988 (22) <u>3336140</u>	Study type: • Observational Cohort • 24-h ABPM <134/90 • Systematic review • Office vs. ABPM or HBPM Size: 292 pts	N/A	<u>1° endpoint</u> : WCH=21%	• Multiple methodologies used to define MH. Prevalence 8.5%–16.6% (general population), 14.7%–30.4% (nonelevated clinic population)

Uhlig K, et al., 2012 (23) <u>22439158</u>	Study type: • Systematic review • Self-monitoring vs. usual care vs. self- monitoring+support	N/A	<u>1° endpoint</u> : Change in clinic SBP/DBP	 Self-monitoring vs. usual care resulted in lower SBP/DBP (-3.1/-2.0 mm Hg) at 6 mo Self-monitoring + support vs. usual care resulted in lower SBP/DBP SBP/DBP -3.4 8.9/-1.94.4 mm Hg up to 12 mo. Self-monitoring may confer a small benefit for BP control.
McManus RJ, et al., 2014 (24) <u>25157723</u>	Study type: • RCT • Self-monitoring with self- titration vs. usual care. Size: 552 pts	Inclusion criteria: SBP/DBP ≥130/85 mm Hg	<u>1° endpoint</u> : Change in SBP/DBP at 12 mo	• Self-monitoring with self-titration was associated with SBP and DBP differences of 9.2 mm Hg and 3.4 mm Hg, respectively.
Margolis KL, et al., 2013 (25) <u>23821088</u>	Aim: Assess impact of follow-up and monitoring system including home BP tele-monitoring and pharmacist case management on BP control in pts treated for HTN <u>Study type:</u> Cluster RCT <u>Size</u> : 450 pts	Inclusion criteria: Pts from 16 clinics in integrated health system in Minneapolis, MN	222 pts randomized to 8 usual care clinics and 228 randomized to 8 intervention clinics Intervention included 12 mo of home BP tele-monitoring and pharmacist case management, with 6 mo of follow-up afterward	 Intervention group achieved better BP control compared to usual care during 12 mo of intervention and persisting during 6 mo of follow-up SBP was <140/90 in 57.2% (95% CI: 44.8%, 68.7%) of intervention pts at 6 and 12 mo vs. 30% (95% CI: 23.2%, 37.8%) in usual care (p=0.001) Combination of home BP tele-monitoring and pharmacist case management helped control HTN better than usual care at 6, 12, and 18 mo.24
Margolis KL, et al., 2013 (25) <u>23821088</u>	Study type: • RCT • Home BP telemonitoring with pharmacist case management vs. usual care. <u>Size</u> : 450 pts	Inclusion criteria: Uncontrolled BP	<u>1° endpoint</u> : SBP/DBP <140/90 mm Hg (<130/80 mm Hg in DM or CKD) at 6 and 12 mo. <u>2° endpoint</u> : Change in BP, pt satisfaction, and BP control at 18 mo (6 mo after intervention stopped).	 Telemonitoring resulted in better BP control (57% vs. 30%) at 6 and 12 mo and larger SBP declines at 6, 12, and 18 mo. Some aspects of pt satisfaction (e.g., clinicians listening carefully) improved with telemonitoring.
McManus RJ, et al., 2014 (24) <u>25157723</u>	Study type: • RCT • Self-monitoring with self- titration vs. usual care. Size: 552 pts	Inclusion criteria: SBP/DBP ≥130/85 mm Hg	<u>1° endpoint</u> : Change in SBP/DBP at 12 mo	• Self-monitoring with self-titration was associated with SBP and DBP differences of 9.2 mm Hg and 3.4 mm Hg, respectively.

Siu AL, et al., 2015 <u>26458123</u>	Study type: U.S. Preventive Services Task Force commissioned systematic review and meta-analysis of office and out of office BP relationships for diagnostic accuracy of diagnosing high BP after an initial office-based classification of high BP.	Inclusion criteria: • Adults ≥18 y. • 24 studies based on "confirmation" by means of ABPM and 6 by means of HPBM.	 <u>1° endpoint</u>: ABPM or HBPM conformation of office-based diagnosis of high BP. CVD risk-relationships for ABPM, HBPM and office-based BPs also reviewed. ABPM was recommended as the best method to confirm an office-based diagnosis of high BP, with HBPM an acceptable alternative, based on "over diagnosis" of high BP with office BP measurements (White coat hypertension) and stronger relationships between out of office BP measurements (especially ABPM) with vascular events. 	• Screen for high BP in adults ≥18 y and confirm office-based high BP using out of office BP measurements 9preferably ABPM).
Uhlig K, et al., 2012 (23) <u>22439158</u>	Study type: • Systematic review • Self-monitoring vs. usual care vs. self- monitoring+support	N/A	<u>1° endpoint</u> : Change in clinic SBP/DBP	 Self-monitoring vs. usual care resulted in lower SBP/DBP (-3.1/-2.0 mm Hg) at 6 mo Self-monitoring + support vs. usual care resulted in lower SBP/DBP SBP/DBP -3.4 8.9/-1.94.4 mm Hg up to 12 mo. Self-monitoring may confer a small benefit for BP control.
Yi SS, et al., 2015 (26) <u>25737487</u>	Study type: • RCT • Self-monitoring of BP vs. usual care. <u>Size</u> : 900 pts	N/A	 <u>1° endpoint</u>: Change in clinic SBP/DBP and HTN control (SBP/DBP <140/90 mm Hg) Decline in SBP at 9 mo was 14.7 mm Hg and 14.1 mm Hg in the intervention and usual care groups (p=0.70); HTN was controlled in 38.9% and 39.1% in the intervention and control groups (p=0.91) 	• Self-monitoring of BP by itself does not improve BP above usual care.
Agarwal R, et. al., 2011 (27) <u>21115879</u>	Study type: • Systematic review	N/A	<u>1° endpoint</u> : • Change in clinic SBP/DBP and MAP	• Self-monitoring is associated with a reduction in BP. This effect is larger when accompanied by telemonitoring.

	Self-monitoring vs. usual care vs. self- monitoring+telemonitoring <u>Size</u> : 9,446 pts		• Mean reduction in SBP, DBP and MAP with home monitoring was 2.63 mm Hg (95% CI: 4.24– 1.02), 1.68 (95% CI: 2.58–0.79), 4.0 (95% CI: 1.79–6.22). The effect for SBP was larger when accompanied by telemonitoring (3.20; 95% CI: 4.66–1.73 vs. 1.26; 95% CI: 2.20–0.31).	
Fagard RH, et. al., 2007 (28) <u>17921809</u>	Study type: • Systematic review • MH and WCH vs. sustained normotension Size: 11,502 pts	N/A	<u>1° endpoint</u> : CVD events. The adjusted HR for CVD events was 1.12 (95% CI: 0.84–1.50) for WCH vs. sustained normotension (p=0.59) and 2.00 (95% CI: 1.58– 2.52) for MH vs. sustained normotension (p<0.001)	 MH is associated with increased CVD risk but WCH is not associated with increased risk.

Data Supplement 4. White Coat Hypertension (Section 4.4)

Study Acronym; Author; Year Published	Study Type/Design; Definitions	Patient Population (N)	HBPM (%)	Daytime ABPM (%)	24-h ABPM (%)	Results/Comments
Viera AJ, et al., 2010 (29) <u>20671718</u>	Office BP ×3 Duplicate measures of: 24-h ABPM >130/80 Daytime ABPM>135/85 HBPM >135/85	 50 pts Untreated Borderline HTN and BP >110/70 and <160/110 	• MH=43/35	• MH=54/53	• MH=51/45	 For MH diagnosis 95% agreement daytime and 24-h ABPM Only 47%–53% agreement between HBPM and either daytime or 24-h ABPM
Viera AJ, et al., 2014 (30) <u>24842491</u>	Office BP ×3 Duplicate measures of: 24-h ABPM >130/80 Daytime ABPM >135/85 HBPM >135/85	 420 pts Untreated Borderline HTN and BP >120/80 and <149/95 	• MH=15–17	• MH=43-44	• MH=48-50	 For MH Diagnosis 92%–94% agreement daytime and 24-h ABPM 70% agreement between HBPM and either daytime K=0.3–0.36
Bayo B, et al., 2006 (31) <u>16534404</u>	Office BP ×3HBPM ×3 d	190 untreated ptsSpanishBorderline	• WCH=35 (95% CI: 28-42)		• WCH=42 (95% CI: 34, 48)	• Compared to ABPM, HBPM pulse pressure variation: 59% negative predictive value: 69%

Asayama K, et al., 2015 (32) <u>25135185</u>	Obs (IDACO) database CV outcomes risk by WCH, MH, NTN ABPM measured: Office BP ×2 >140/90 (office) >130/80 (24-h ABPM) >135/85 (daytime ABPM) >120/70 (nighttime ABPM)	8,237 untreated pts	N/A	• WCH=9.1 • MH=13.4	• WCH=10.7 • MH=9.7	 Overlap from daytime to 24-h ABPM: WCH=86% MH=61%
Conen D, et al., 2014 (33) <u>25185130</u>	 Obs 13 IDACO Cohorts Office ×2 Awake ABPM >135/85 24-h ABP >130/80 Analyzed by decade in y 	• 7,506 untreated pts	 WCH=2.2% age 18–30, increasing to 19.5% in both sexes age >70 y MH=inverted U distribution (13% and 11% in 18–30 y 18% and 20% in those 30–50 y Increased prevalence in men 		 WCH=3.0 in age 18–30 increasing to 19.1% both sexes age >70 y MH=inverted U distribution (12% and 9% in youngest and oldest, 19% and 17% in those 30– 50 y Increase prevalence in men 	• Similar prevalence using either 24-h or awake ABPM
Nasothimiou EG, et al., 2012 (34) 22357523	 Office BP ×3 × >140/90 HBPM >135/85 Daytime ABPM >135/85 	• 613 pts (66% untreated, 34% treated)	● WCH=15% ● MH=15%	• WCH=14% • MH=16%	N/A	 WCH: 89% agreement daytime ABPM and HBPM, kappa=0.79 MH: 88% agreement, kappa=0.56
Coll de TG, et al., 2011 (35) <u>21183853</u>	Office ×2 >140/90 Daytime ABPM >135/85 HBPM >135/85	403 untreated pts	• WCH=24%	• WCH=8.1%	N/A	N/A
Stergiou GS, et al., 2005 (36) <u>15925734</u>	 Office ×3 ×2 >140/90 HBPM ≥135/85 awake ● ABPM ≥135/85 	438 untreated/ treated pts	• MH=12% • WCH=16%	● MH=14% ● WCH=15%	N/A	 No difference in proportions of pts Dx with MH or WCH by HBPM or awake ABPM No difference between treated and untreated. However, only 44% overlap for MH, but 90%–95% if 5 mm Hg zone of uncertainty added.

Sega R, et al., 2001 (37) <u>11560854</u>	Population-based PAMELA Study	• 2,051 pts	● WCH=12% ● MH=9%	• WCH=12% • MH=9%	N/A	 70% agreement between ABPM and HBPM for WCH and 57% for MH
	 Office ×3 >140/90 HBPM >132/83 ABPM >125/79 LVMI by echo 					

Data Supplement 5. White Coat Hypertension (Prevalence) (Section 4.4)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Vinyoles E et al., 2008 (38) <u>18300853</u>	Study type: • Cross-sectional, comparative multicenter descriptive study Size: 6,176 pts	N/A	<u>1° endpoint</u> : WCH=21%	 Multiple methodologies used to define MH. Prevalence 8.5%–16.6% (general population), 14.7%–30.4% (nonelevated clinic population)
Pickering TG, et al., 1988 (22) <u>3336140</u>	Study type: • Observational cohort • 24-h ABPM <134/90 • Systematic review • Office vs. ABPM or HBPM Size: 292 pts	N/A	<u>1° endpoint</u> : WCH=21%	 Multiple methodologies used to define MH. Prevalence 8.5%–16.6% (general population), 14.7%–30.4% (nonelevated clinic population)
Piper MA, et al., 2015 (39) 25531400	Study type: • Systematic review • Office vs. ABPM or HBPM	N/A	<u>1° endpoint</u> : ●WCH=5–35% (ABPM) ● WCH conversion to SH ~1%–5% y	 Prevalence of WCH sufficiently high to require ABPM confirmation of SH in those with elevated clinic BP
Asayama K, et al., 2014 (32) <u>25135185</u>	Study type: • Observational (IDACO) database • ABPM measured: • Office BP ×2 • >140/90 (office) • >130/80 (24-h ABPM) • >135/85 (daytime ABPM) • >120/70 (nighttime ABPM)	Inclusion criteria: Untreated, >18 y	<u>1° endpoint</u> : • WCH=6.3%-12.5% • MH=9.7%-19.6%	• Variable prevalence of both WCH and MH based on method of defining
	<u>Size:</u> 8,237			

Conen D, et al., 2014 (33) <u>25185130</u>	Study type: • Observational • 13 IDACO cohorts • Office ×2 • Awake ABPM >135/85 • 24-h ABP >130/80 • Analyzed by decade in y Size: 7,506 pts	Inclusion criteria: >18 y, untreated	 <u>1° endpoint</u>: WCH=2.2% age 18–30 y, increasing to 19.5% both sexes age >70 y MH=inverted U distribution (13% and 11% in youngest and oldest, 18% and 20% in those 30–50 y) Increase prevalence in males 	 Increase in WCH prevalence with increasing age in both sexes Peak MH prevalence age 30–50 y with drop at age extremes. Greater prevalence of MH in males. Similar prevalence when 24-h vs. awake ABPM used
Alwan H, et al., 2014 (40) <u>24663506</u>	Study type: Observational SKIPOGH Office BP ×4 Daytime ABPM Office >140/90 Daytime >135/85	Inclusion criteria: >18 y, untreated	<u>1° endpoint:</u> • WCH=2.6% • MH=15.8%	• Pts with pre-HTN had 7 times higher rate of MH
Stergiou GS, et al., 2014 (41) <u>24420553</u>	Study type: • Observational • 5 IDACO cohort Studies • Office ×2 >140/90 • Home >135/85 • Median 8.3-y follow-up Size: 5,007 pts	Inclusion criteria: >18 y, untreated	<u>1° endpoint</u> : Long-term follow-up for CVD events	• WCH=13.8% • MH=8.1%
Pierdomenico SD, et al., 2011 (42) <u>20847724</u>	Study type: Meta-analysis of observational cohort studies (8 WCH, 5 MH) 24-h ABPM >130/80 Daytime >135/85 Size: 7,961 pts	Inclusion criteria: >18 y, untreated	<u>1° endpoint:</u> Long-term follow-up for CVD events	• WCH=16.1% • MH=5.8%
Hansen TW, et al., 2007 (43) <u>17620947</u>	Study type: • 4 observational studies • Office <140/90 • 24-h ABPM >135/85 Size: 7,030 pts	78% untreated	Study endpoints: • F/NF CVD • Median follow-up =9.5 y 1° Results: • Adj HR vs. NTN • WCH=1.22 (CI: 0.96–1.53), p=0.09 • MH=1.62 (CI: 1.35–1.96), p<0.001	N/A

	● SH=1.80 (CI: 1.59–2.03), p<0.001	

Data Supplement 6. White Coat Hypertension (Correlation with Clinical Outcomes) (Section 4.4)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Study Endpoints and Length of Follow-up	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Summary/Conclusions/ Comment
NICE 2011 (44) <u>22855971</u>	Study type: • Systematic Review • 3 Meta-analyses • 11 observational studies "best method" comparison of office vs. HBPM or ABPM that best predicted (i.e., statistically significant predictors and higher HR values) clinical outcomes (after adjustment for covariates in multivariate analyses)	 Home vs. office (n=7,685) ABPM vs. office (n=33,158) Home vs. ABPM vs. Office (n=2,442) 	• Outcomes of interest: mortality, stroke, MI, HF, DM, vascular procedures, hospitalization for angina, and other MACCE	 For predicting clinical outcomes: ABPM vs. office (9 studies): ABPM superior to office (8 studies) No difference between ABPM and office (1 study) HBPM vs. office (3 studies): HBPM superior to office (2 studies) No difference between HBPM and office (1 study) HBPM vs. ABPM vs. office (2 studies): HBPM similar to ABPM and both superior to office (1 study) No difference between HBPM, ABPM and office (1 study) 	• Overall recommendation for ABPM to confirm HTN diagnosis (HBPM recommended if ABPM not practical)
Pierdomenico SD, et al., 2011 (42) <u>20847724</u>	Study type: Meta-analysis (8 studies) • NTN vs. WCH or MH based mostly on daytime ABPM <135/85 Size: 7,961	Inclusion criteria: Untreated	 Follow-up 3.2–12.8 y Composite CVD 	 WCH vs. NTN: OR: 0.96; 95% CI: 0.65–1.42 MH vs. NTN: OR: 2.09; 95% CI: 1.55–2.81 SH vs. NTN: OR: 2.59; 95% CI: 2.00–3.35 	N/A
Asayama K, et al., 2014 (32) <u>25135185</u>	Study type: Observational (IDACO) database • CV outcomes risk by WCH, MH, NTN • ABPM measured: Office BP ×2 >140/90 (office)	Inclusion criteria: >18 y, untreated	 F/NF CVD/stroke, 729 CV events Follow-up 10.6 y 	 WCH adjusted HR: 1.2; 95% CI: 0.93– 1.54; p=0.16 MH adjusted HR: 1.81; 95% CI: 1.41– 2.32; p<0.0001 SH adjusted HR: 2.31; 95% CI: 1.91– 2.80; p<0.0001 	N/A

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	(24-h ABPM) >130/80 (daytime ABPM) >135/85 (nighttime ABPM) >120/70 Size: 8,237				
Verdecchia P, et al., 2005 (45) <u>15596572</u>	Study type: Population- based (4 international cohorts) • Office ×3 >140/90 • Awake ABPM >130/80 Size: 5,955	• 26% NTN	StrokeFollow-up 5.4 y	 WCH adjusted HR: 1.15; 95% CI: 0.61–2.16; p=0.66 SH adjusted HR: 2.01; 95% CI: 1.31–3.08; p<0.001 	• Stroke not increased in WCH but tended to approach systolic HTN risk 6 y after baseline ABPM.
Hansen TW, et al., 2007 (43) <u>17620947</u>	Study type: Observational 4 studies • Office <140/90 • 24-h ABPM >135/85 Size: 7,030	78% untreated	 F/NF CVD Median follow-up=9.5 y 	 WCH adjusted HR: 1.22 (95% CI: 0.96, 1.53), p=0.09 MH adjusted HR: 1.62; 95% CI: 1.35–1.96; p<0.001 SH adjusted HR: 1.80; 95% CI: 1.59–2.03; p<0.001 	N/A
Fagard RH, et al., 2007 (28) <u>17921809</u>	Study type: Meta-analysis 7 studies • Office <140/90 • 24-h ABPM or HBPM Size: 11,502	Treated and untreated	 F or F/NF CVD Follow-up 3.2–12.3 y (mean=8 y) 	 WCH adjusted HR: 1.12 (95% CI: 0.84– 1.50), p=0.59 MH adjusted HR: 2.0; 95% CI: 1.58– 2.52; p<0.001 Systolic HTN adjusted HR: 2.28; 95% CI: 1.87–2.78; p<0.001 	N/A
Mancia G, et al., 2013 (46) <u>23716584</u>	Study type: Observational PAMELA Study • Office ×3<140/90 • HBPM>135/85 and-24-h • ABPM>130/80 Size: 2,051	• 22% treated	 CV and all- cause mortality Follow-up 16 y 	 CV mortality in WCH adjusted HR: 2.04 (95% CI: 0.87–4.78), p=0.10 All-cause mortality in WCH adjusted HR: 1.50; 95% CI: 1.03–2.18; p=0.03 	 Trend but insignificant increase in CV mortality and significant increase in total mortality in WCH Risk of developing systolic HTN greater in those with WCH
Tomiyama M, et al., 2006 (47) <u>16942927</u>	Study type: Cross-sectional study assessing target organ damage by BP control status. Control: Office <140/90, daytime <135/85. Size: 332	Treated pts	 LVMI, carotid IMT, UAE Cross-sectional 	LVMI, carotid IMT and UAE increased in masked uncontrolled HTN compared to controlled HTN. LVMI and UAE increased in SH	• SH and masked uncontrolled HTN but not WCE associated with increased target organ damage

Ohkubo T, et al., 2005 (48) <u>16053966</u>	Study type: Observational cohort • Office ×2 >140/90 • Awake ABPM >135/85 Size: 1,332	 Untreated (70%) Treated (30%) 	 CVD mortality/stroke Follow-up 10 y 	 WCH RH: 1.28; 95% CI: 0.76–2.14); p=0.4 MH RH: 2.13; 95% CI:1.38–3.29; p<0.001 SH RH: 2.26; 95% CI:1.77–4.54; p<0.0001 	 Similar results treated and untreated, males, and females
Tientcheu D, et al., 2015 (49) <u>26564592</u>	Study type: Observational cohort • Home readings ×5 ×2 visits taken by research staff • Office readings ×5 Size: 3,027	 Dallas Heart Study 54% African American 30%–39% treated 	 Clinical CVD incl TIA, UA 	 WCH adj HR: 2.09; 95% CI: 1.05–4.15; p=0.035 MH adj HR: 2.03; 95% CI: 1.36–3.03; p<0.001 SH adj HR: 3.12; 95% CI: 2.13–4.56; p<0.001 	Higher CVD with SH, MH and WCH (African Americans only). CVD risk not increased in whites with WCH

Data Supplement 7. Renal Artery Stenosis (Section 5.4.3)

Study Acronym (if applicable) Author Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Lawes CM, et al., 2003 (50) <u>12658016</u>	<u>Study type:</u> Meta- analysis of RCTs of BP drugs recording CHD events and strokes <u>Size</u> : 464,000 pts	N/A	• CHD RR or 46% Stroke 64%	• All classes of BP meds confer benefit while BB confer greater benefit in those with CAD
Riaz IB, et al., 2014 (51) <u>25145333</u>	Study type: 540 studies and 7 RCTs Size: 2,139 pts	N/A	• Incidence of nonfatal MI 6.74% in both the stenting and medical therapy groups: OR: 0.99; 95% CI: 0.70–1.43; p=0.99, incidence of renal events in stenting population was found to be 19.58% vs. 20.53% in medical therapy OR: 0.95; 95% CI: 0.76–1.18; p=0.62.	 BP effect, CV accident not specifically reported
Cooper CJ, et al., 2014 (52) <u>24245566</u>	Study type: Residential treatment center medical therapy with or without renal stent Size: 947 pts	Inclusion criteria: Atherosclerotic renal artery stenosis	• Composite endpoint of death from CV or renal causes, MI, stroke, hospitalization for congestive HF, progressive renal insufficiency, or the need for renal-replacement therapy. 35.1% and 35.8%, respectively; HR with stenting: 0.94; 95% CI: 0.76–1.17; p=0.58 Difference in SBP favoring the stent group: -2.3 mm Hg; 95% CI: -4.4– -0.2; p=0.03.	N/A

Xie X, et al., 2015 (21) <u>26559744</u>	Study type: MA of RTC that randomly assigned individuals to different target BP levels Size: 44,989 pts	• 19 trials	 Achieved BP 133/76 mm Hg (intensive) 140/81 (less intense) Major CV events: 14%; 95% CI: 4%–22% MI: 13%; 95% CI: 0%–24% Stroke: 22%; 95% CI: 10%–32% Albuminuria: 10%; 95% CI: 3%–16% Retinopathy progression: 19%; 95% CI: 0%–34%. More intensive had no effects on HF: 15%; 95% CI: -11%–34% CV death: 9%; 95% CI: -11%–26% Total mortality: 9%; 95% CI: -3%–19% ESKD: 10%; 95% CI: -6%–23% 	• More intensive approach reduced major CV events (stroke and MI) except heat failure, CVD, ESRD, and total mortality.
Brunström M, et al., 2016 (53) <u>26920333</u>	Study type: Meta- analysis of levels of BP control in DM hypertensives. Size: 73,738 pts	• 49 trials (most pts with DM-2)	Baseline SBP >150 <u>RR for</u> • All death: 0.89; 95% CI:0.80–0.99 • CVD: 0.75; 95% CI: 0.57–0.99 • MI: 0.74; 95% CI: 0.63–0.87 • Stroke: 0.77; 95% CI: 0.65–0.91 • ESRD: 0.82; 95% CI: 0.71–0.94 Baseline SBP140–150 <u>RR of</u> • Death: 0.87; 95% CI: 0.78–0.98) • MI: 0.84; 95% CI: 0.76–0.9 • HF: 0.80; 95% CI: 0.66–0.97 If baseline SBP,140 mm Hg, however, further treatment increased the risk of CV mortality (1.15; 95% CI: 1.00–1.32	• BP lowering reduces major CV events in DM. Caution for initiating treatment in diabetics with SBP <140/90
Ettehad D, et al., 2015 (17) <u>26724178</u>	Study type: Meta- analysis of large RTCs of antihypertensive treatment <u>Size</u> : 613,815 pts	• 123 studies	Every 10 mm Hg reduction in SBP RR: • Major CV events: 0.80; 95% CI: 0.77–0.83 • CHD: 0.83; 95% CI: 0.78–0.88 • Stroke: 0.73; 95% CI: 0.68–0.77), HF (0.72, 0.67–0.78 • All-cause mortality: 0.87; 95% CI: 0.87; 0.84–0.91 • ESRD: 0.95; 0.84–1.07	• BP lowering reduces CV risk across various baseline BP levels and comorbidities. Suggest lowering SBP <130 mm Hg and BP-lowering treatment to pts with a history of CVD, CHD, stroke, DM, HF, and CKD.

Thomopolous C, et al., 2016 (54) <u>26848994</u>	Study type: Meta- analysis of RTCs of more vs. less intense BP control	• 16 trials (52,235 pts) compared more vs. less intense treatment 34 (138,127 pts) active vs. placebo	More intense BP • Stroke RR: 0.71; 95% CI: 0.60–0.84) • CHD RR: 0.80; 95% CI: 0.68–0.95) • Major CV events RR: 0.75; 95% CI: 0.68–0.85 • CV mortality RR: 0.79; 95% CI: 0.63–0.97 Stratification of SBP cutoffs (150,140 and 130 mm Hg) showed that a SBP/DBP difference of 10/5 mm Hg across each cutoff reduced risk of all outcomes	Intensive BP reduction improves CV outcomes compared to less intense Achieved BP <130/80 may be associated with CV benefit.
Julius S, et al., 2006 (55) <u>16537662</u>	<u>Study type:</u> RCT in pre- HTN; 16 mg candesartan vs. placebo <u>Size</u> : 809 pts	• 58% men	• During the first 2 y, HTN developed in 154 (40.4%) pts in the placebo group compared with only 53 (13.6%) of those in the candesartan group, for RR: 66.3% (p<0.0001). After 4 y, HTN developed in 240 (63.0%) in the placebo group vs. only 208 (53.2%) in the candesartan group RR: 15.6% (p<0.0069).	• 2/3 of those with pre-HTN develop HTN within 4 y. Candesartan interrupts the onset and reduced by 15.6%
Ference BA, et al., 2014 (56) <u>24591335</u>	Study type: Evaluated the effect of 12 polymorphisms (associated with BP) on the odds of CHD and compared it with the effect of lower SBP observed in both prospective cohort studies and BP-lowering randomized trials Size: 199,477 pts	63 studies	• 12 polymorphisms were associated with a 0.32 mm Hg lower SBP (p=1.79×10 ⁻⁷) and a 0.093-mm Hg/decade slower age-related rise in SBP (p=3.05×10 ⁻⁵). The effect of long-term exposure to lower SBP on CHD mediated by these polymorphisms was 2-fold greater than that observed in prospective cohort studies (p=0.006) and 3-fold greater than that observed in short-term BP treatment trials (p=0.001).	• SBP may be causally associated with the rate of rise in SBP with age and has a cumulative effect on the risk of CHD.

Data Supplement 8. RCTs Comparing Obstructive Sleep Apnea (Section 5.4.4)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Barb F, et al., 2010 (57)	<u>Aim</u> : Assess the effect on BP of 1 y of	Inclusion criteria: Pts with	Intervention: CPAP	1° endpoint: Decrease in BP	Limitations: Not blinded; both groups consisted of pts with severe sleep-
<u>20007932</u>	treatment with CPAP in nonsleepy pts with HTN and OSA.	HTN (on medications or ≥140/90) and	Comparator: Conservative treatment	Results: At 12 mo, CPAP decreased SBP by 1.89 mm Hg (95% CI: 3.90–0.11 mm Hg; p=0.065) and DBP 2.19 mm Hg	apnea.

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	Study type: RCT Size: 359 pts; 12 mo of follow-up	apnea-hypopnea index >19.	(dietary counseling and sleep hygiene advice).	(95% CI: 3.46– -0.93 mm Hg: p=0.001). The most significant reduction in BP was in pts who used CPAP for more than 5.6 h/night.	Conclusions: CPAP induced a significant reduction in BP, albeit small, in hypertensive pts with OSA.
Martinez-Garcia MA, et al., 2013 (58) <u>24327037</u>	Aim: Assess the effect of CPAP on BP in pts with OSA and resistant hypertension. Study type: RCT Size: 194 pts; 3 mo follow-up	Inclusion criteria: Pts with resistant hypertension and OSA.	Intervention: CPAP Comparator: No therapy	 <u>1° endpoint</u>: Change in 24-h ABPM from baseline to 12 wk. <u>Results</u>: When the changes in BP were compared between groups by intent to treat, the CPAP group achieved a greater decrease in 24-h mean BP (3.1 mm Hg (95% Cl: 0.6, 5.6); p=0.02) and 24-h DBP (3.2 mm Hg (95% Cl: 1.0, 5.4; p=0.005) but not in 24-h SBP (3.1 mm Hg (95% Cl: -0.6–6.7; p=0.10) compared to control. There was also a greater nocturnal BP dipping pattern in CPAP treated pts than control (35.9% vs. 21.6%; adjusted OR: 2.4; Cl: 1.2–5.1; p=0.02). There was a significant positive correlation between h of CPAP use and the decrease in mean 24-h BP (r=0.29; Cl) 	Limitations: Did not use sham CPAP as placebo; open-label; short follow-up. Conclusions: Among pts with resistant hypertension and OSA, CPAP treatment for 12 wk compared with control resulted in a decrease in 24-h mean and DBP and improvement in nocturnal pressure pattern.
Lozano L, et al., 2010 (59) <u>20577130</u>	Aim: Assess effect of CPAP on pts with OSA and resistant hypertension. Study type: RCT Size: 96 pts; 3 mo of follow-up	Inclusion criteria: Pts with resistant hypertension and OSA.	Intervention: CPAP + conventional drug treatment Comparator: Conventional drug treatment alone	0.006), SBP (r=0.25; p=0.02) and DBP (r=0.30; p=0.005). <u>1° endpoint</u> : Decrease in 24-h ABPM from baseline to 12 wk. <u>Results:</u> Pts with ABPM confirmed resistant hypertension treated with CPAP, unlike those treated with conventional therapy, showed a decrease in 24-h DBP (-4.9±6.4 vs. 0.1±7.3 mm Hg; p=0.027). Pts who used CPAP >5.8 h showed a greater reduction in daytime DBP (-6.12 mm Hg; 95% Cl: -1.45–10.82; p=0.004), 24-h DBP (-6.98 mm Hg; 95% Cl: -1.86– 12.1; p=0.009) and 24-h SBP (-9.71 mm Hg; 95% Cl: -0.20– -19.22; p=0.46).	Limitations: Small study; only 3 mo follow-up; lack of sham control. Conclusions: CPAP as a complement to usual treatment improved mean 24-h DBP in pts with OSA and ABPM- confirmed resistant hypertension.

Muxfeldt ES, et al., 2015 (60) <u>25601933</u>	Aim: Evaluate the effect of CPAP on pts with resistant hypertension and OSA. <u>Study type</u> : RCT <u>Size:</u> 434 pts; 6 mo of follow-up	Inclusion criteria: Pts with resistant hypertension and OSA	Intervention: CPAP + conventional antihypertensive therapy Comparator: Antihypertensive therapy alone. Conventional antihypertensive therapy included spironolactone.	 <u>1° endpoint</u>: BP reduction at 6 mo via ABPM <u>Results:</u> On an intention-to-treat analysis, there was no significant difference in any BP change, neither in nocturnal BP fall, between CPAP and control groups. The best effect of CPAP was on night-time SBP in per-protocol analysis, with greater reduction of 4.7 mm Hg (95% Cl: -1.6%–5.8%; p=0.25, in comparison with the control group. Median use of CPAP was 4.8 h. 	Limitations: Nonblinded design; per protocol analysis underpowered to show the prespecified outcome of 6–7 mm Hg SBP differences between CPAP and control groups. Conclusions: CPAP had no significant effect on clinic or ambulatory BP in pts with resistant hypertension and moderately severe to severe OSA. However, in the specific subgroup of pts with uncontrolled ambulatory BP, CPAP may modestly reduce night-time SBP and improve the nocturnal BP fall pattern. The reason for lack of BP reduction in the overall study may have been due to excellent control of BP with median 5 medications, including spironolactone, in the majority of pts.
Pedrosa RP, et al., 2013 (61) <u>23598607</u>	Aim: Evaluate the effect of CPAP on pts with resistant hypertension and OSA. <u>Study type</u> : RCT with <u>Size:</u> 40 pts; 6 mo follow-up	Inclusion <u>criteria:</u> Pts with resistant hypertension and OSA	Intervention: CPAP + conventional antihypertensive therapy (n=20) Comparator: Antihypertensive therapy alone (n=20).	<u>1° endpoint</u> : BP reduction at 6 mo by ABPM. <u>Results</u> : BP was 162±4/97±2 mm Hg prior to randomization. CPAP was used for 6 h/night. Compared with the control group, awake SBP/DBP decreased significantly in the CPAP group (- 6.5±3.3/-4.5±1.9 vs. +3.1±3.3/2.1±2/7 mm Hg; p<0.05). BP changes were significant only when pts were awake but not at night by ABPM.	Limitations: Small; but strength was rigorous exclusion of pts who were nonadherent; control arm did not undergo placebo treatment; nonblinded. Conclusions: Treatment of OSA with CPAP significantly reduces daytime BP in pts with resistant hypertension.

Data Supplement 9. RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Dietary Fiber Intake) (Section 6.2)

Study Acronym;	Aim of Study;	Patient Population	Study Intervention (#	Endpoint Results	Relevant 2° Endpoint (if any);
Author:	Study Type;		patients) /	(Absolute Event Rates,	Study Limitations;
Year Published			Study Comparator (# patients)	P value; OR or RR; & 95% CI)	Adverse Events

Whelton SP, et al., 2005 (62) <u>15716684</u>	Aim: Study the effect of dietary fiber intake on BP Study type: Systematic review and meta- analysis Size: • 21 RCTs (25 comparisons) with 1,477 pts • 20 of the RCTs were conducted in nonhypertensive persons • 13 double-blind; 3 single blind and 9 open label	Inclusion criteria: • RCT • ≥16 y • English language publication before Feb. 2004 • No concurrent interventions Exclusion criteria: Missing key data	Intervention: Fiber supplementation, either as a pill (8 trials), cereal/fruit/veg (15 trials), Pectin (1 trial), Guar gum (1 trial) <u>Comparator</u> : Placebo or no fiber supplementation	<u>1° endpoint</u> : In a pooled analysis of the overall group (hypertensive and normotensive persons), the mean for change in SBP was -1.15 mm Hg; 95% CI: -2.68–0.39 mm Hg and for DBP was -1.65 mm Hg; 95% CI: -2.70– -0.61 mm Hg. In the subgroup of 20 trials conducted in nonhypertensives, the mean change in SBP was -0.14 mm Hg; 95% CI: -1.10–0.86 mm Hg. In the subgroup of 5 trials conducted in hypertensives, the mean change in BP was -5.95 mm Hg; 95% CI: -9.50– -2.40) mm Hg.	 This is the most detailed and comprehensive review of the topic. It provides limited evidence, overall, that fiber supplementation results in a significant in BP and suggests no evidence in support of an effect in normotensives.
Streppel MT, et al., 2005 (63) <u>15668359</u>	Aim: Study the effect of fiber supplementation on BP Study type: Systematic review and meta- analysis Size: • 23 RCTs (25 comparisons) in 1,404 pts • Mean duration=9 wk • Mean age=42 y • 16 double-blind, with 14 (67%) of the 21 comparisons conducted in normotensive pts • 3 trials based on plant protein and 4 trials based on animal protein	Inclusion criteria • Human RCT • BP 1° or 2° outcome • Publications between January 1966–January 2003 Exclusion criteria: • Inadequate reporting of the data • Concurrent intervention	Intervention: Fiber supplementation (average dose=11.5 g/d); soluble fiber in 11 trials, insoluble fiber in 7 trials, and a mixture in the remaining trials Comparator: Placebo or no fiber supplementation	<u>1° endpoint</u>: In the overall group (hypertensive and normotensive pts), a pooled analysis identified a MD for change in SBP of -1.13 mm Hg; 95% CI: -2.49–0.23. In a subgroup of 17 trials conducted in "nonhypertensives" (mean baseline BP<140/90 mm Hg or <50% receiving antihypertensive medication), the mean treatment effect was -0.23 mm Hg; 95% CI: -1.43–0.98 in univariate analysis and -1.00 mm Hg; 95% CI: - 1.94– -0.06 mm Hg in multivariate analysis that adjusted for age, sex, study design, duration of intervention, and fiber dose. The corresponding effects in 8 trials conducted in hypertensives were -4.53 mm Hg; 95% CI: -6.69– -2.38 mm Hg; and -2.42 mm Hg; 95% CI: -5.28–0.45 mm Hg. Safety endpoint: N/A	• Findings consistent with experience in the meta-analysis by Whelton et al.

Evans CE, et al., 2011 (64) <u>25668347</u>	Aim: Study the effect of fiber supplementation on BP Study type: Systematic review and meta- analysis Size: 28 trials met the inclusion criteria and reported fiber intake and SBP and/or DBP. 18 trials were included in a meta-analysis.	Inclusion criteria • RCTs, in humans of at least 6 wk duration • Fiber isolate or fiber-rich diet against a control or placebo • Published between 1 January 1990 and 1 December 2013. Exclusion criteria: N/A	Intervention: Fiber supplementation (average dose =11.5 g/d) -soluble fiber in 11 trials, insoluble fiber in 7 trials, and a mixture in the remaining trials <u>Comparator</u> : Placebo or no fiber supplementation	<u>1° endpoint</u> : Studies were categorized into 1 of 12 fiber-type categories. The pooled estimates for all fiber types were -0.9 mm Hg (95% CI: -2.5–0.6 mm Hg) and -0.7 mm Hg (95% CI: -1.9–0.5 mm Hg) for SBP and DBP, respectively. The median difference in total fiber was 6g. Analyses of specific fiber types concluded that diets rich in beta-glucans reduce SBP by 2.9 mm Hg (95% CI: 0.9, 4.9 mm Hg) and DBP by 1.5 mm Hg (95% CI: 0.2–2.7 mm Hg) for a median difference in beta-glucans of 4 g. Heterogeneity for individual fiber types was generally low	 Higher consumption of beta- glucan fiber is associated with lower SBP and DBP. The results of this review are consistent with recommendations to increase consumption of foods rich in dietary fiber, but some additional emphasis on sources of beta-glucans, such as oats and barley, may be warranted.

Data Supplement 10. RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Fish Oil) (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Campbell F, et al., 2012 (65) <u>22345681</u>	<u>Aim</u> : Study the effect of fish oil supplementation on BP <u>Study type</u> : Systematic review and meta-analysis <u>Size</u> : • 17 RCTs (25 comparisons) with 1,524 pts. • 9 trials were conducted in normotensives (1,049	Inclusion criteria: • RCT • English language publication before January 2011 • Duration ≥8 wk Exclusion criteria: N/A	Intervention: Fish oil given in capsule form, with doses varying from 0.8–13.33 g/d. <u>Comparator</u> : Placebo (usually corn oil, olive oil, or safflower oil).	<u>1° endpoint</u> : In a pooled analysis of the 8 trials conducted in hypertensive pts, the mean for change in SBP was - 2.56 mm Hg; 95% CI: -4.53– -0.58 mm Hg. The corresponding SBP change for the 9 trials conducted in normotensives was -0.50 mm Hg; 95% CI: -1.44– 0.45.	 This is the most recent of many that have been published. Previous meta-analyses have been conducted by Appel et al (1993), Morris et al. (1993), Geleijnse et al (2002) and Dickinson et al. (2006). In general, the findings have been fairly consistent in demonstrating a relatively small (2 3/4 mm Hg SBP) but significant effect, with most of this being attributable to the results in trials conducted in hypertensive pts.

Rodriguez- Leyva D, et al., 2013 (66) <u>24126178</u>	pts with mean age of 47 y). Follow-up varied 2–26 wk. <u>Aim:</u> Study the effect of flaxseed on BP in hypertensive pts <u>Study type</u> : RCT <u>Size</u> : 110 pts with PAD	Inclusion criteria: • >40 y • PAD for >6 mo, ABI <0.9 Exclusion criteria: Inability to walk, bowel disease, moderate to severe renal failure, life expectancy <2 y with high cardiac risk, allergy to any of the study products, pts who plan to undergo surgery during the course of the trial, and no more	Intervention: Pts given 1 food item per day for 6 mo, containing either 30 g of milled flax seed or placebo. Flaxseed contains omega-3 fatty acids, lignans, and fiber. Comparator: Placebo	<u>1° endpoint</u> : SBP and DBP consistently decreased in the flaxseed group over the course of the study. After 6 mo, SBP in the flaxseed group dropped significantly to 136±22 mm Hg (p=0.04). On the contrary, in the placebo group, SBP rose slightly to 146±21 mm Hg. After 6 mo of intervention, DBP in the flaxseed group fell to 72±11 mm Hg (p=0.004), whereas DBP in the placebo group remained the same (79±10 mm Hg).	• Based on this 1 RCT, flaxseed appeared to have a significant BP lowering effect
		of the trial, and no more than 2 fish meals per wk			

Data Supplement 11. RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Potassium Supplementation to Placebo or Usual Diet) (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Whelton PK, et al., 1997 (67) <u>9168293</u>	Aim: Study the effect of potassium supplementation on BP Study type: Systematic review and meta-analysis Size:	Inclusion criteria: • Human RCT • Without HTN • Potassium supplementation vs. control • No concurrent interventions <u>Exclusion criteria</u> : Missing key data	Intervention: Potassium supplementation in 1,049 pts (potassium chloride tabs in 10 RCTs with 618 pts and diet in 2 RCT with 431 pts) <u>Comparator</u> : No potassium supplementation	 <u>1° endpoint</u>: Significant reduction in BP. Overall (hypertensives and normotensives), mean: 3.11 mm Hg; 95% CI: -4.321.91 mm Hg. In the 12 trials conducted in normotensives, mean: -1.8 mm Hg; 95% CI: -2.90.6 mm Hg for SBP and -1.0 mm Hg; 95% CI: -2.1-0.0 for DBP 	 This is the most comprehensive presentation of the effects of potassium on BP, including experience in normotensives. Significant reduction in SBP overall and in the subgroups with and without HTN. In a subsequent meta-analysis of 23 trials, Geleijnse JM, Kok FJ, and Grobbee DE (J Hum Hypertens. 2003;17:471-480) reported a similar effect of potassium on SBP in both hypertensives and nonhypertensives (mean of -3.2 and -1.4 mm Hg, respectively).

	 Overall, 33 RCT (n=2,609) 2 RCTs (n=1,049) in normotensives 		(placebo in 10 RCT and usual diet in 2 RCT)	• In the 20 trials conducted in hypertensives, mean: -4.4 mm Hg; 95% CI: -6.6– -2.2 for SBP and -2.5 mm Hg; 95% CI: -4.9– -0.1 for DBP <u>Safety endpoint</u> : N/A	 The 1 RCT conducted in African-Americans (n=87) identified a mean treatment effect size of -6.9 mm Hg; 95% Cl: -9.34.4 for SBP (p<0.001) and -2.5 mm Hg; 95% Cl: -4.30.8 for DBP (p=0.004). In the entire cohort (trials conducted in pts with HTN and normotension), net changes in SBP and DBP were directly related to level of urinary sodium excretion during the trial.
Aburto NJ, et al., 2013 (68) <u>23558164</u>	Aim: Study the effect of potassium supplementation on BP Study type: Systematic review and meta-analysis Size: 21 RCTs (n=1,892); 16 in pts with HTN (n=818) and 3 RCTs in pts without HTN (n=757)	Inclusion criteria: • RCT in humans • Duration ≥4 wk • 24-h collections of urinary potassium • No concomitant interventions Exclusion criteria: Pts who were acutely ill, HIV positive, hospitalized, or had impaired urinary excretion of potassium	Intervention: Potassium supplementation in 20 trials, supplements plus diet/education in 1 trial, and diet/education alone in 2 trials. Comparator: No potassium supplementation (placebo or usual diet)	 <u>1° endpoint</u>: Overall change in SBP=- 5.93; 95% CI: -10.151.70. After removing outlier trials, the change was -3.49 mm Hg; 95% CI: -5.151.82 mm Hg. In 16 trials conducted in hypertensives, change in SBP was -5.32 mm Hg; 95% CI: -7.203.43. In the 3 trials conducted in persons without HTN, change in SBP was 0.09 mm Hg; 95% CI: -0.77-0.95. Safety endpoint: N/A 	 1 trial (TOHP Phase I) incorrectly entered twice so only 2 trials really available. However, this does not change overall finding. The negative results for normotensives in this meta-analysis (and difference with the findings by Whelton et al) probably reflects the requirement for a duration of ≥4 wk and the fact that few trials of this duration have been conducted in normotensives.
Geleijnse JM, et al., 2003 (69) <u>12821954</u>	Aim: Study the effect of potassium supplementation on BP Study type: Systematic review and meta- regression analysis Size: 27 RCTs; 19 in pts with HTN and 11 RCTs in pts without HTN	Inclusion criteria: • RCT in adults • Published after 1966 • Duration ≥2 wk • No concomitant interventions Exclusion criteria: • Disease • Outlier results (1 trial)	Intervention: Potassium supplementation Comparator: No potassium supplementation	 <u>1° endpoint</u>: Overall change in SBP=- 2.42; 95% CI: -3.751.08 In the 19 trials conducted in hypertensives, change in SBP was -3.51 mm Hg; 95% CI: -5.311.72 In the 3 trials conducted in persons without HTN, change in SBP was 0.97 mm Hg; 95% CI: -3.07-1.14 <u>Safety endpoint</u>: N/A 	 Imputation for missing data In addition to the treatment effect difference by presence/absence of HTN, there was a trend toward a larger treatment effect in older age (≥45 y), and to a lesser extent higher baseline urinary Na (>150 mmol/24 h) and greater increase in urinary K (>44 mmol/24 h)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Rebholz CM, et al., 2012 (70) 23035142	Aim: Study the effect of protein intake on BP Study type: Systematic review and meta- analysis Size: • 40 RCTs (44 comparisons) with 3,277 pts • 32 comparisons of protein vs. carbohydrate • 12 comparisons of vegetable vs. animal protein • 35 of the RCTs were conducted in normotensive persons (28 with SBP in the prehypertensive range)	Inclusion criteria: • RCT in humans • ≥18 y • Publication between January 1,1950 and April 1, 2011 • No concurrent interventions • No more than 10% difference in calories, sodium, potassium, fiber between the treatment arms • Duration ≥1 wk Exclusion criteria: Missing key data	Intervention: • Protein intake • 1st meta-analysis: any source of protein, with a median protein supplementation dose of 40 g/d (20–66 g/d) • 2 nd meta-analysis: specifically vegetable or animal protein Comparator: • 1st meta-analysis: carbohydrate • 2 nd meta-analysis: vegetable or animal protein	 1º endpoint: 1st meta-analysis There was a fairly consistent trend for a small BP lowering effect of protein compared to carbohydrate intake (86% of the trials). In a pooled analysis of the overall group (hypertensive and normotensive persons), the mean for change in SBP was -1.76 (95% CI: - 2.331.20). In a subgroup of 15 trials in which none of the participants were receiving antihypertensive medication, the mean change in SBP was -1.95 (95% CI: -2.621.29). 2nd meta-analysis For the comparison of vegetable vs. animal protein, there was no evidence of a difference in BP. In a pooled analysis of the overall group (hypertensive and normotensive pts) the mean change in SBP was -0.10 (95% CI: -2.31-2.11) mm Hg. In a subgroup of 8 trials in which none of the pts were receiving antihypertensive medication, the mean change in SBP was -0.55 (95% CI: -3.06-1.96). 1° Safety endpoint: N/A 	 This is the most detailed and comprehensive review of the topic. It provides strong evidence that protein supplementation results in a significant but modest reduction in BP and suggests that the effect size is similar following supplementation with protein from vegetables or animals.
Tielemans SM, et al., 2013 (71) <u>23514841</u>	Aim: Study the effect of protein intake on BP	Inclusion criteria • RCTs, in "generally healthy adults"	Intervention: Protein intake	1° endpoint: • At baseline, the mean for age and SBP were 50 (range: 31–74) and 128	• Findings consistent with experience in the meta- analysis by Rebholz et al.

Data Supplement 12. RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Protein Intake on BP) (Section 6.2)

	Study type: Systematic review and meta- analysis Size: 16 RCT (210 comparisons) of protein vs. carbohydrate in 1,449 pts, with 14 (67%) of the 21 comparisons conducted in normotensive pts. -3 trials based on plant protein and 4 trials based on animal protein	 Publications between January 1966–January 2012 <u>Exclusion criteria</u>: Inadequate reporting of the data Concurrent intervention 	<u>Comparator</u> : Carbohydrate intake	 (range: 112–144). During the trials, the MD in protein intake was 48 g/d (range: 26–74 g/d). In the overall group (hypertensive and normotensive participants), a pooled analysis of comparisons from 14 trials (1,208 pts) identified a MD for change in SBP of -2.11 (95% CI: -2.8– -1.37) for protein vs. carbohydrate. In 3 RCTs that employed plant protein (327 pts), the mean treatment effect was -1.95 (95% CI: -3.21– -0.69) and in 4 RCTs that employed animal protein (574 pts), the corresponding difference was -2.20 (95% CI: -3.36– -1.03). Safety endpoint: N/A 	
Dong JY, et al., 2013 (72) <u>23829939</u>	Aim: Study the effect of protein intake on BP in DM-2 Study type: Systematic review and meta- analysis Size: 9 RCTs with 418 pts	Inclusion criteria: • RCTs in adults with DM-2 • Publications up to August 2012 • High protein diet intervention and ≥5% difference in dietary protein intake between intervention and control groups • Trial duration ≥4 wk Exclusion criteria: Inadequate reporting of key data	Intervention: High protein diet intervention and ≥5% difference in dietary protein intake between intervention and control groups Comparator: N/A	<u>1° endpoint</u> : Pooled experience in the 14 trials identified a nonsignificant reduction in mean SBP of -3.10 (95% CI: -4.63– -1.56). <u>Safety endpoint</u> : N/A	 Heterogeneous group of open label trials with a range of duration from 4–24 wk (median of 12 wk). In addition to DM-2, all of the participants were overweight or obese. The quality of the trials varied, drop-out rates ranged from 0%–0%, and only 1 trial was analyzed using an intent to treat approach.
Dong JY, et al., 2013 (73) <u>23823502</u>	Aim: Study the effect of probiotic fermented milk on BP. Study type: Systematic review and meta- analysis. All but 1	Inclusion criteria: • RCTs • Placebo controlled • Published prior to March 2012 Exclusion criteria:	Intervention: Probiotic fermented milk (100– 450 g/d) <u>Comparator</u> : Not specified but all of the trials reported to be	<u>1° endpoint</u> : Pooled experience in the 9 trials identified a nonsignificant reduction in mean SBP of -3.59 (95% CI: -7.58–0.40). <u>Safety endpoint</u> : N/A	• The most recent of several meta-analyses conducted by different groups of investigators that have reported a similar effect size following administration of lactopeptides, especially the

use a parallel design. Antihypertensive drug	 Intervention with enzymatically hydrolysed milk Cointervention 	placebo controlled. However, 2 were single blind and 1 was open label.	 lactotripeptides Valine- Proline-Proline and Isolucine- Proline-Proline. These findings may have special relevance for countries, like Japan, where consumption of fermented milk products is common.
<u>Size</u> : 14 RCTs with 702 pts (median size=40).			

Data Supplement 13. RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Sodium Reduction to Placebo or Usual Diet) (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
NUTRICODE Mozaffarian D, et al., 2014 (74) <u>25119608</u>	Aim: Study the effect of sodium reduction on BP and CVD mortality Study type: Meta- regression analysis Size: 103 RCTs (107 comparisons) with 6,970 pts; 38 of the 107 comparisons were conducted in normotensive pts	Inclusion criteria: RCT in 2 previous Cochrane meta-analyses Exclusion criteria: • Duration <1 wk • Mean 24-h collections or estimates of urinary sodium reduced <20 mmol in the intervention group compared to control • Concomitant interventions	Intervention: Sodium reduction Comparator: No sodium reduction	 <u>1° endpoint</u>: Strong evidence for a linear relationship between reduction in sodium intake and lower levels of SBP throughout the entire distribution of sodium studied, with larger reductions in older persons, blacks (compared to whites) and hypertensives (compared to normotensives). For a white, normotensive population at age 50 y, each reduction of 100 mmol/d (2.3 g/d) in dietary sodium lowered SBP by a mean: 3.74 (95% CI: 5.18–2.29). Modeling based on global estimates of sodium intake, effect of sodium reduction on BP, and effect of BP reduction on CVD mortality attributed 1.65 million CVD deaths annually due sodium intake >2 g/d. this would represent 9.5% (95% CI: 6.4–12.8) of all CVD mortality. Estimates were not 	 RCT meta-regression analysis that provides evidence for BP lowering following a reduction in dietary sodium intake, overall and in normotensive persons, with a more pronounced effect in those who were older, black and had a higher starting level of BP. These findings are consistent with other reports. The modeling analysis suggested sodium reduction would yield important population health benefits but did not specify the magnitude of the potential benefit for pts within the normal BP range.

Aburto NJ, et al., 2013 (68) 23558164	<u>Aim</u> : Study the effect of sodium reduction on BP <u>Study type</u> : Systematic review and meta-analysis <u>Size</u> : Overall study included 36 trials (49 comparisons) conducted in 6,736 pts. Of these, 3,263 were nonhypertensive. The results in normotensives in this table are based on experience in 7 RCTs conducted in 3,067 normotensive pts.	Inclusion criteria: • RCT in humans • Trial duration ≥4 wk • 24-h urinary sodium ≥40 mmol/d less in treatment compared to control group • No concurrent interventions • Not acutely ill Exclusion criteria: Lack of above	Intervention: Sodium reduction Comparator: No sodium reduction	provided separately for hypertensive and normotensive persons. <u>1° Safety endpoint</u> : N/A <u>1° endpoint</u> : In pooled analysis, the overall change in SBP was -3.39 (95% CI: -4.31– - 2.46) mm Hg. In the pts with HTN, the change was -4.06 (95% CI: -5.15– -2.96). In the normotensives, the change was -1.38 (95% CI: -2.74–0.02). <u>Safety endpoint</u> : In the small number of relevant trials, there was no significant effect of sodium reduction on lipid levels (Total cholesterol, LDL-cholesterol, HDL- cholesterol, triglyceride levels; 11 trials) or on plasma (7 trials) or urinary catecholamine levels (2 trials). Experience in 4 trials (3 which could not be included in the meta-analysis) suggested a beneficial effect of sodium reduction on urinary protein excretion.	• Study inclusion/exclusion criteria designed to yield a group of trials that would provide results that have relevance for clinical practice and public health. In this context, reduced sodium intake resulted in a statistically significant but small reduction in SBP.
He FJ, et al., 2013 (75) <u>22437256</u>	Aim: Study the effect of sodium reduction on BP Study type: Systematic review, meta-analysis and meta-regression analysis Size: Overall study included 34 trials (37 comparisons) conducted in 3,230	Inclusion criteria: • RCTs • Healthy adults ≥18 y • Trial duration ≥4 wk • Sodium intake only difference between treatment and control group • 24-h urine sodium ≥40 mmol less in treatment compared to control Exclusion criteria: Lack of above	Intervention: Sodium reduction Comparator: No sodium reduction	 <u>1° endpoint</u>: In an overall pooled analysis, the change for SBP was -4.18 (95% CI: -5.183.18) mm Hg. In the trials of persons with HTN, the mean change was -5.39 (95% CI: -6.624.15) mm Hg. In the trials conducted in normotensives, the change in SBP was -2.42 (95% CI: -3.561.29) mm Hg. In meta-regression analysis, change in 24-h urinary sodium was significantly associated with reduction in SBP (4.3 mm Hg for a 100 mmol reduction in 24-h urinary sodium). 	 Study inclusion/exclusion criteria designed to yield a group of trials that would provide results that have relevance for clinical practice and public health. In this context, reduced sodium intake resulted in a significant and potentially important reduction in SBP. The meta-regression results were consistent with a dose- response relationship in normotensive pts

	pts. 12 of the RCTs (14 comparisons) were conducted in 2,240 normotensive pts.			Safety endpoint: In the small number of relevant trials (which included both hypertensive and normotensive pts) that provided safety endpoint measurements (4–14 trials), there was no change in total, LDL- or HDL- cholesterol, or triglyceride levels. There were small significant increases in plasma renin activity, aldosterone, and noradrenaline levels but these were consistent with expected physiologic responses to sodium reduction.	
Graudal NA, et al., 2012 (76) <u>22068710</u>	Aim: Study the effect of sodium reduction on BP Systematic review and meta-analysis Size: Overall study included 167 trials. Of these, 71 RCTs were conducted in 5,577 normotensive pts, with the following characteristics: Median age: 27 y (13–67 y) Median trial duration: 7 d (4– 1,100 d) 5,292 Whites (71 studies) 268 Blacks (7 studies) 215 Asians (3 studies)	Inclusion criteria: • RCTs • 24-h collections or estimates from ≥8 h collections of urinary sodium excretion Exclusion criteria: Systematic studies in unhealthy pts with diseases other than HTN	Intervention: Sodium reduction Comparator: No sodium reduction	 <u>1° endpoint</u>: The overall effect of sodium reduction was not presented. A forest plot of 71 comparisons (from 61 trials) in the 4,919 normotensive whites assigned to sodium reduction compared to usual sodium intake identified a trend towards lower SBP in 50 (70%), no difference in 8 (11%), and higher SBP in 13 (19%). In a pooled analysis, sodium reduction compared to usual sodium intake in the normotensives yielded the following MDs in SBP: Whites: -1.27 (95% CI: -1.88– -0.66) Blacks: -4.02 (95% CI: -7.37– -0.68) Asians: -1.27 (95% CI: -3.07– -0.54) A corresponding analysis in the hypertensives yielded the following MDs in SBP: Whites: -5.48 (95% CI: -6.53– -4.43) Blacks: -6.44 (95% CI: -8.85– -4.03) Asians: -10.21 (95% CI: -16.98– -3.44) 	 Heterogeneous group of trials that included many small studies of short duration in young persons. Overall finding of lower BP in those assigned to a reduced intake of dietary sodium, with an apparently greater effect in Blacks compared to Whites and Asians. The hormone changes in this meta-analysis likely reflect a physiologic response to sodium reduction, especially in studies of short duration and rapid changes in sodium intake. The increases in total cholesterol and triglyceride levels were not noted in the meta-analyses conducted by Aburto et al. and He et al.

DASH-Sodium Trial Sacks FM, et al., 2001 (77) <u>11136953</u>	Aim: Study the effect of sodium reduction on BP Study type: Randomized, controlled crossover trial Size: Overall study based on 412 pts, of whom 243 were normotensive	Inclusion criteria: Adults ≥22 y Exclusion criteria: Taking antihypertensive medication, heart disease, renal disease, poorly controlled hyperlipidemia or DM, DM requiring insulin, special dietary requirements, >14 drinks/wk	Intervention: Feeding study in which pts were randomized to a DASH or control diet at 3 levels of assigned dietary sodium intake (High=210 mmol/d; Intermediate=100 mmol/d; Low=50 mmol/d) <u>Comparator</u> : Each pt served as their own control (crossover design)	comparisons in both hypertensive and normotensive participants) that provided safety endpoint measurements, significant increases in the standard MD for plasma renin activity (70 trials), aldosterone (51 trials), noradrenaline (31 trials), adrenaline (14 trials), and weighted MD for total cholesterol (24 trials), and triglyceride (18 trials) levels. There was no significant effect of sodium reduction on LDL-cholesterol (15 trials) and HDL-cholesterol (17 trials). <u>1° endpoint</u> : • Reduced sodium intake resulted in a significant reduction in SBP, with a greater reduction during assignment to the Low compared to the Intermediate sodium intake diet. At every level of sodium intake, the achieved reduction in SBP was greater on the control group compared to the DASH diet and for Blacks compared to other pts. • Reducing sodium intake from the high to intermediate level decreased SBP by 2.1 mm Hg (p<0.001) during the control diet and 1.3 mm Hg (p=0.03) during the DASH diet. • Reducing sodium intake from the intermediate low level decreased SBP by a further 4.6 mm Hg (p<0.001) during the control diet and 1.7 mm Hg (p<0.01) during the DASH diet. Safety endpoint : N/A 1° endpoint :	 This trial provides the best (direct) evidence for a dose- response treatment relationship between sodium intake and level of BP. It also suggests the relative effect of reduced sodium intake is greater in persons with a typical U.S. diet but the combination of sodium reduction and consumption of a DASH-type diet results in a lower level of BP than can be achieved with either dietary modification on its own. Consistent with other trials and meta-analyses, it suggests the effect of a reduced sodium intake is greater in Blacks compared to others, especially for those consuming a typical U.S. diet. This was the largest trial of
(Sodium component) Kumanyika SK, et al., 2005 (78) <u>15372064</u>	of sodium reduction on BP and prevention of HTN. Study type: Randomized,	 Healthy community- dwelling adults 30–54 y BMI between 110% and 165% of desirable body weight 	Behavior change intervention aimed at studying the effects of modest (25%–30%) reductions in dietary sodium intake during	<u>Change in SBP</u> Compared to usual care, the sodium reduction group experienced a significant mean reduction of 51 mmol for 24-h urinary excretion and -2.9 (SD: 0.5) mm Hg (p<0.001) in SBP at 6 mo (-5.1 mm Hg in	 on this was the largest that of sodium reduction in HTN prevention and also provides the longest duration of follow. The assumptions for a main effects factorial analysis (independence of the

	controlled factorial trial. <u>Size</u> : 2,382 pts, of whom 594 were randomized to sodium reduction (alone) and 596 were randomized to usual care.	 Not taking BP-lowering medication Mean SBP <140 mm Hg and DBP 83–89 mm Hg Exclusion criteria: Taking antihypertensive medication, Heart disease, renal disease, poorly controlled hyperlipidemia or DM, DM requiring insulin, special dietary requirements, >14 drinks/wk. 	up to 48 mo (minimum 36 mo) of follow-up. <u>Comparator</u> : Usual care group	 the sodium reduction group and -2.2 mm Hg in the usual care group). A progressive reduction in effect size for urinary sodium excretion and BP was noted over time, with mean for SBP at 18, 36 mo and termination of -2.0 (SD: 0.5) mm Hg (p<0.001), -1.2 (SD: 0.5) mm Hg (p=0.02), and -1.0 (SD: 0.5) mm Hg (p=0.5). <u>Prevention of HTN</u> At 6 mo of follow-up the incidence of new onset HTN was 39% lower in the pts randomized to reduced dietary sodium intake compared to the usual care group (p=0.04). During more prolonged follow-up, the effect size decreased but remained significant after 48 mo of follow-up (14% reduction; p=0.04). Overall, the incidence of HTN was reduced by 18% (p=0.048). <u>Safety endpoint</u>: N/A 	interventions) were not demonstrated. Given this finding, the most reliable analysis of this trial was comparison of the experience in each active intervention group with the usual care group. This results in a reduction in statistical power. • Consistent with the pattern in the proceeding TOHP I trial sodium reduction reduced BP and the incidence of HTN but the effect sizes for sodium reduction and BP as well as the difficulty of maintaining the intervention in highly motivated and extensively counselled participants underscores the difficulty of achieving sodium reduction in the general population without changes in food processing and restaurant/fast food preparation practices.
TOHP Phase I 1992 (79) <u>1586398</u>	Aim: Study the effect of sodium reduction on BP and prevention of HTN Study type: Randomized, controlled factorial trial. Size: Overall, 2,182 adults, with the 327 assigned to sodium reduction compared	Inclusion criteria: • Community-dwelling adults 30–54 y • Not on antihypertensive medication • DBP 80-89 mm Hg • Healthy Exclusion criteria: • Disease • Inability to comply with the protocol	Intervention: Behavior change intervention Comparator: Usual care	<u>1° endpoint</u> : Change in DBP <u>2° endpoint</u> : Change in SBP <u>Safety endpoint</u> : CVD events, symptoms and general and well being	 Significantly lower DBP (0.9 mm Hg; p<0.05) and SBP (1.7 mm Hg; p<0.01) in the sodium reduction group compared to usual care Few CVD events No difference in symptoms Significant improvement in general well-being at 6 and 18 mo (p<0.05)

Cook NR, et al., 2007 (80)	to 417 usual care controls <u>Aim</u> : Study the effect of sodium reduction	Inclusion criteria: Assigned to dietary	Intervention: Behavior change	<u>1° endpoint:</u> • 200 CVD events and 77 deaths during	Dietary sodium reduction, previously shown to reduce BP
<u>17449506</u>	on CVD morbidity and mortality. <u>Study type:</u> 10–15 y post-trial follow-up of TOHP I and TOPH II pts that took advantage of the randomized trial design. Vital status was obtained for 100% of the pts and information on morbidity was obtained from 2,415 (77%) of the pts. <u>Size</u> : 744 TOHP Phase I and 2,382 TOHP Phase II pts	sodium reduction or control in TOHP Phase I or TOHP Phase II. <u>Exclusion criteria</u> : None	intervention aimed at studying the effects of modest (25%–30%) reductions in dietary sodium intake during TOHP Phase I or TOHP Phase II. <u>Comparator</u> : No sodium reduction intervention.	 follow-up Kaplan-Meier plots identified trends toward less morbidity and mortality in those who had been randomized to sodium reduction compared to usual care, with a consistent pattern for the TOHP I and TOHP II participants Risk of a CVD event was 30% lower (RR: 0.70; 95% CI: 0.53–0.94; p=0.018) among those randomized to sodium reduction compared to usual care, after adjustment for trial, clinic, age, race, sex, baseline weight and sodium excretion RR for total mortality was 0.80 (95% CI: 0.51–1.26). 	and prevent HTN in the TOHP I and TOHP II trials, appeared to reduce CVD events during extended post-trial follow-up of the pts from these 2 trials.

Data Supplement 14. RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Stress Reduction) (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Canter PH, et al., 2004 (81) <u>15480084</u>	<u>Aim</u> : Study the effect of transcendental meditation on BP <u>Study type</u> : Systematic review <u>Size</u> :	Inclusion criteria: • RCT in humans • Publication in any language until May 2004 • No concurrent interventions	Intervention: • Use of transcendental meditation techniques as taught by Maharishi Mahesh Yogi • Practiced on a regular basis over an extended period	<u>1° endpoint</u> : Statistically significant reduction in SBP reported in 3 of 5 trials that provided such information. <u>1° Safety endpoint</u> : N/A	 Only a handful of RCTs available from the large number of publications on this topic. Trials had methodological weaknesses and were subject to potential bias due to the affiliation of authors to the transcendental meditation organization.

 6 RCTs with wide range of pts: young to elderly; healthy volunteers to Blacks with HTN. HTN: 2 trials High normal BP: 2 trials Normotensive: 1 trial Not stated: 1 trial Sample sizes ranging from 34–156 pts Follow-up from 2 mo–1 y 	<u>Exclusion criteria</u> : N/A	Comparator: No treatment, sham, alternative treatment	 A few trials reported small reductions in SBP but clinical relevance of findings is unclear. Most of the trials were underpowered and could have missed a significant finding. The authors concluded that "there is at present insufficient good quality information to conclude whether or not transcendental meditation has a cumulative positive effect on BP"
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Data Supplement 15. RCTs and Meta-analyses Studying the Effect of Nonpharmacologic Interventions on BP (Dietary Patterns) (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Appel LJ, et al., 1997 (82) <u>9099655</u>	Aim: Study the effect of dietary patterns on BP Study type: • Multicenter RCT • 3 arm parallel design • 3 wk pre- randomization run-in phase • Feeding study with 8 wk of intervention Size: 459 adults, mean age 44 y. (326 normotensive)	Inclusion criteria: • Adults ≥22 y • SBP<160 mm Hg and DBP 80–95 mm Hg • No antihypertensive medication Exclusion criteria: • CVD event within 6 mo • Poorly controlled DM or hyperlipidemia • BMI ≥35 • Pregnancy or lactation • Chronic illness that would interfere with participation • Unwillingness to stop taking vitamins, mineral supplements, Ca++ antacids	Intervention: • Diet high in fruits and vegetables • "Combination" diet high in fruits, vegetables, low-fat dairy products, and reduced total fat, saturated fat and cholesterol. <u>Comparator</u> : Usual U.S. diet	 <u>1° endpoint</u>: Compared to the control diet, both intervention diets reduced BP, with an overall mean (95% Cl) reduction of: Fruits and Veg. Diet: SBP: -2.8 (95% Cl: -4.7– -0.9) DBP: -1.1 (95% Cl: -2.4– -0.3) Combination Diet: SBP: -5.5 (95% Cl: -7.4– -3.7) DBP: -3.0 (95% Cl: -4.3– -1.6) The BP changes in the subgroup with HTN were: Fruits and Veg. Diet: SBP: -7.2 (-11.4, -3.0) DBP: -2.8 (-5.4, -0.3) Combination Diet: SBP: -5.5 (-8.2, -2.7) 	 This trial was the first of several to document the value of the combination diet (later renamed the DASH diet). The BP reductions noted with the DASH (combination) diet were substantial and well maintained. Generalizability was limited due to the nature of the intervention (feeding study) and the relatively short period of intervention experience (8 wk)

		 Consuming ≥14 		The corresponding changes in the	
		alcoholic drinks with		subgroup of normotensives were:	
		 Renal insufficiency 		 Fruits and Veg. Diet: 	
		_		SBP: -0.8 (-2.7, 1.1)	
				DBP: -0.3 (-1.9, 1.3)	
				 Combination Diet: 	
				SBP: -3.5 (-5.3, -1.6)	
				DBP: -2.1 (-3.6, -0.5)	
				1° Safety endpoint: Infrequent and	
				similar occurrence of	
				gastrointestinal symptoms in each	
				group	
Sacks FM, et al.,	Aim: Study the effect	Inclusion criteria:	Intervention: 3 levels of	<u>1° endpoint:</u>	This trial provided additional
2001 (77)	of different levels of	● Adults ≥22 y	dietary sodium while	 At each level of sodium intake, 	documentation of the effectiveness of a
<u>11136953</u>	sodium intake on BP	 Average SBP between 	consuming a DASH or	SBP and DBP were lower during	DASH diet in lowering BP in
	during consumption of	120–159 mm Hg and	usual U.S. diet. The	consumption of the DASH diet	normotensives (and hypertensives) and
	a DASH or usual U.S.	average DBP between	target sodium intake	compared to the usual U.S. diet, the	the complementary benefit of
	diet	80–95 mm Hg	levels for a daily energy intake of 2,100 kcal	difference being greatest with high	consuming a reduced intake of sodium.
	Study type:	No use of	were:	sodium intake and lowest with low	
	Multicenter RCT with	antihypertensive	High: 150 mmol (3,450	sodium intake, with the mean SBP difference between the DASH and	
	2 parallel diet arms	medication	mg)/d	usual US diets during high,	
	(DASH diet or usual	Exclusion criteria:	Intermediate: 100 mmol	intermediate and low sodium intake	
	U.S. diet)	Heart disease, renal	(2,300 mg)/d	being -5.9 (95% CI: -8.0– -3.7), -5.0	
	• Within each arm,	insufficiency, poorly	Low: 50 mmol (1,150	(95% CI: -7.6– -2.5), and -2.2 (95%	
	randomized cross-over	controlled hyperlipidemia	mg)/d	CI: $-4.40.1$). The corresponding	
	trial with 3 periods	or DM, DM requiring	3,	differences for DBP were -2.9 (95%	
	testing different levels	insulin, special dietary	The mean achieved	Cl: -4.3– -1.5), -2.5 (95% Cl: -4.1– -	
	of sodium intake (no	requirements, >14	levels of sodium during	0.8), and -1.0 (95% CI: -2.5, 0.4).	
	washout)	alcoholic drinks /wk.	the high, intermediate	• In both the DASH and usual U.S.	
			and low sodium periods	diet arms, SBP and DBP were	
	Size: 412, with 59%		were 144, 107 and 67	significantly lower during	
	(243) being		mmol/d in the DASH	intermediate compared to high	
	normotensive		diet group and 141, 106,	sodium intake, and during low	
			and 64 mmol/d in the	compared to intermediate sodium	
			usual U.S. diet group.	intake, with the decrement being	
			Comporator: Soo	greater for the latter change.	
			Comparator: See description above	 In comparison to consumption of a 	
			description above	usual U.S. diet at the high level of	

PREMIER Appel LJ, et al., 2003 (83) 12709466	Aim: Study the effect of 2 behavioral interventions, aimed at dietary change, on BP Study type: • Multicenter RCT with 3 parallel arms: • Established • Established plus DASH diet • Advice only Size: 810 adults, with 62% (506) normotensive. At baseline mean age	Inclusion criteria: • Adults ≥25y • Average SBP between 120–159 mm Hg and average DBP between 80–95 mm Hg • No use of antihypertensive medication • BMI between 18.5 and 45 kg/m ² Exclusion criteria: • Regular use of drugs that affect BP • Target organ damage or	Intervention: • Structured behavioral interventions that used an identical format (4 individual and 14 group sessions) to facilitate adoption of "established" dietary recommendations for reduction in BP or "established" plus the DASH diet. The "established" dietary recommendations used in PREMIER were a) weight loss in overweight participants	sodium intake, the normotensive group consuming the DASH diet at the low level of sodium intake had a mean SBP difference of 7.1 mm Hg (p<0.001). <u>1° Safety endpoint:</u> Participants tended to report less symptoms during periods of reduced sodium intake, with a statistically significant reduction in reports of headache (p<0.05) consistent with prior experience in the TONE trial. <u>1° endpoint</u> • Compared to control (advice only), SBP and DBP were significantly reduced with both active interventions but there was no significant difference in the effect size between the 2 active intervention groups. This was true for both the normotensive and hypertensive pts, with the effect size being larger in the hypertensive group. In the normotensives, the MD for change in SBP was identical for the "established" compared to "established plus DASH Diet" groups: -3.1 (95% CI: -5.1– -1.1) mm Ha	 This was an interesting trial which employed a behavior change approach to implement both active interventions. The investigators goal was to determine the additive value of the DASH Diet in persons already following key elements of conventional (established) recommendations for nonpharmacologic intervention to lower BP. The intervention approach in this trial was less effective in achieving weight loss and reduction in dietary sodium compared to the corresponding experience in the TOHP and TONE trials and the DASH Diet effects on intermediate variables (such as fruit and
	810 adults, with 62% (506) normotensive. At baseline, mean age, BMI and SBP/DBP	that affect BP • Target organ damage or DM • Use of weight-loss meds	in PREMIER were a)	"established plus DASH Diet"	experience in the TOHP and TONE
	were 50 y, 33 kg/m ² , and 135/85 mm Hg, respectively.	 Hx CVD event HF, angina, cancer, within 2 y Consumption of >21 	activity, reduced alcohol intake in pts consuming alcohol.	for the "established" intervention group and -2.0 (95% CI: -3.4– -0.6) for the "established intervention plus	studies. • Despite the modest intervention effects, both SBP and DBP were
	Duration: 6 mo, with observations at 3 and 6 mo.	 Ochsamption of 221 alcoholic drinks /wk Pregnancy, planned pregnancy, lactation 	• Compared to experience in the advice only (control) group, there was only modest achievement of	 DASH Diet) group. Overall, the incidence of HTN was lowest and the percent with optimal BP was highest in the "established plus DASH" diet but the incidence of 	significantly reduced with the conventional intervention approach (in normotensives as well as overall) and addition of the DASH diet did not have a

Appel LJ, et al.,	Aim: Compare effects	Inclusion criteria:	intervention goals in the "established" group, with a MDs of 3.8 kg (8.4 lbs) for body weight, 11.6 mmol (267 mg)/d) for urinary sodium excretion, no change in physical activity (but better fitness), and no change in alcohol consumption (but very low alcohol consumption at baseline). • Weight loss was somewhat greater in the "established" plus DASH diet group, with a MD of 4.8 kg (10.6 lbs) for body weight. This group also manifested expected effects of the DASH diet, with significantly higher urinary potassium and phosphorous levels, greater consumption of fruits and vegetables, dietary calcium, dairy products, and a lower consumption of total fat and saturated fat. <u>Comparator:</u> Advice only Intervention:	HTN was significantly less and the percent with optimal BP was higher in both active intervention groups compared to advice only. The difference between the 2 active intervention groups was not significant. In the normotensives, there was a nonsignificant trend towards less HTN and a significantly higher percent with optimal BP in both active intervention groups compared to advice only, with no significant difference for percent with optimal BP in the 2 active intervention groups. <u>1° Safety endpoint</u> : N/A	 significant effect on reduction of SBP or DBP. There were some nonsignificant trends for slightly lower BP, less HTN, and more optimal BP in the "established plus DASH Diet" group compared to "established" group. The authors also cited use of the DASH Diet as a means to beneficially influence CVD risk factors in addition to BP. This clinical trial demonstrated that
2005 (84) <u>16287956</u>	of 3 diets, each with a reduced intake of saturated fats, on BP and serum lipids	 Adults ≥30 y Average SBP between 120–159 mm Hg and 	High protein with reduced fat/saturated fat content	Compared with the high carbohydrate diet, the high protein diet:	substituting either protein or monounsaturated fat in place of carbohydrate resulted in a small

	Study type: • 2 center RCT • 3 period crossover design • Each 8 wk period was separated by a 2– 4 wk wash-out phase Size: 161–164 included in analyses (191 pts randomized). 132 (80.5%) of the 164 included in the BP analyses were normotensive. Mean age and BMI were 54 y and 30.2 kg/m ² , respectively.	average DBP between 80–95 mm Hg • No use of antihypertensive medication <u>Exclusion criteria:</u> DM, CVD (current or H/O), LDL cholesterol >220 mg/dL, fasting triglycerides >750 mg/dL, weight >350 lb., taking that effect BP or lipids, unwillingness to stop vitamin/mineral supplements, >14 alcoholic drinks/wk.	High unsaturated fats (predominantly monounsaturated fat) with low saturated fat content <u>Comparator:</u> High carbohydrate with reduced fat/saturated fat content	 Reduced SBP by -1.4 mm Hg (p=0.002) overall and by -0.9 mm Hg (p=0.047) in the normotensives Reduced LDL cholesterol by 3.3 mg/dL (p=0.01) overall and by -2.1 mg/dL (p=0.14) in the normotensives Reduced HDL-C by -1.3 mg/dL (p=0.02) overall Reduced serum Triglycerides by - 15.7 mg/dL (p<0.001) overall Compared with the high carbohydrate diet, the high unsaturated fat diet: Reduced SBP by -1.3 mm Hg (p=0.005) overall and by -0.9 (p=0.06) in the normotensives Reduced LDL cholesterol by -1.5 mg/dL (p=0.01) and by -2.1 (p=0.14) in the normotensives Increased HDL-C by 1.1 mg/dL (p=0.03) overall 	reduction in SBP and improvement in lipid profile.
Bazzano LA, et al., 2014 (85) <u>25178568</u>	<u>Aim:</u> Compare the effects of a low- carbohydrate and a low-fat diet on body weight and CVD risk factors (including BP) <u>Study type</u> : Single center parallel arm RCT that compared the 2 diets over 12 mo of intervention. <u>Size:</u> 148 pts, with a mean age of 46.8 y at	Inclusion criteria: • 22–75 y • BMI: 30–45 kg/m ² Exclusion criteria: • CVD • DM-2 • Kidney disease • Use of prescription weight loss meds/surgery • Weight loss >6.8 kg during prior 6 mo	Intervention: • Low-carbohydrate diet, with digestible carbohydrate (total carbohydrate minus total fiber) <40 g/d • Behavioral counselling that employed a mix of 20 individual and group meetings <u>Comparator:</u> • Low fat diet, with <30% of daily energy	 Reduced serum Triglycerides by - 9.6 (p=0.02) overall <u>1° endpoint:</u> Compared to the low-fat diet group, the low-carbohydrate diet group had a mean decrease at 12 mo of: Body weight: -3.5 (95% CI: -5.6 1.4) kg Fat mass: -1.5 (95% CI: -2.60.4) HDL-C: 7.0 (11.0-3.0) mg/dL Ratio total/HDL-C: -0.44 (95% CI: - 0.710.16) Sr. triglyceride: -14.1 (95% CI: - 27.40.8) mg/dL 	 This clinical trial provides 1 of the longest follow-up experiences related to the topic. It suggests low carbohydrate diets may be somewhat better than traditional low fat diets in achievement of weight loss, improvement of lipid profile, inflammation, and CHD risk. Although the BP differences were not significant, there was a consistent trend toward lower BPs in the low-carbohydrate diet group.

	baseline. Mean SBP/DBP at baseline were 124.9/79.4 and 120.3/77.5 mm Hg in the low-fat and low- carbohydrate groups, respectively. The corresponding BMIs were 97.9 and 96.3 kg/m ² . All 148 pts were included in the analysis (intention to treat)		intake from fat (<7% from saturated fat) • Behavioral counselling that used identical format to that employed in the low carbohydrate group	 At 3, 6, and 12 mo, BP tended to be lower in the low-carbohydrate group but none of the differences in SBP or DBP were significant. CRP was reduced in both diet groups but to a significantly greater extent in the low-carbohydrate group. At 6 and 12 mo pts in the low carbohydrate group experienced a significant improvement in their 10-y Framingham CHD risk score. In contrast, there was no change in Framingham CHD risk in the low-fat diet group. <u>1° Safety endpoint:</u> No serious side effects noted 	
Nordmann AJ, et al., 2006 (86) <u>16476868</u>	Aim: Compare effects of low-carbohydrate and low-fat diets on weight loss and CVD risk factors Study type: • Systematic review and meta-analysis • Cochrane Collaboration strategy Size: 5 trials (447 pts)	Inclusion criteria: • RCT • Adults ≥16 y • Low-carbohydrate diet and low-fat diet interventions • BMI ≥25 kg/m ² • Follow-up ≥6 m Exclusion criteria: • Cross-over or sequential design • Missing data	Intervention: Low- carbohydrate diet: maximum of 60 g/d carbohydrate Comparator: Low-fat diet: maximum of 30% energy from fat	 <u>1° endpoint:</u> At 6 mo, the low-carbohydrate diet pts, compared to the low-fat diet participants, had a mean reduction in body weight that was greater by -3.3 (95% CI: -5.3 - 1.4) kg, and a more favorable profile for HDL-cholesterol and triglyceride levels. In contrast, the profile for total-cholesterol and HDL-cholesterol and HDL-cholesterol was more favorable in those assigned to a low-fat diet. The profile for SBP tended to be better in the low carbohydrate diet pts but the differences were not significant: MD at 6 mo: -2.4 (95% CI: -4.9–0.1) mm Hg. 1° Safety endpoint: N/A 	 This systematic review/meta-analysis tends to suggest low-carbohydrate diets are somewhat more effective in reducing body weight compared to the traditionally recommended low-fat diets. Although the BP differences were not significant they would probably have reached a conventional level of significance had subsequent clinical trials (including the Bazzano et al. trial) been included in the analysis.

Nordmann AJ, et al., 2011 (87) <u>21854893</u>	Aim: Compare effects of Mediterranean and low-fat diets on weight loss and CVD risk factors Study type: • Systematic review and meta-analysis • Cochrane Collaboration strategy Size: 6 trials (2,650 pts)	Inclusion criteria: • RCT • Intent to treat analysis • Overweight/obese with at least 1 additional CVD risk factor • Follow-up ≥6 mo Exclusion criteria: N/A	Intervention: Mediterranean diet: moderate fat intake (main sources olive oil and nuts), rich in vegetables, and low in red meat. Comparator: Low fat diet: ≤30% of energy intake from fat	 <u>1° endpoint:</u> Compared to the lowfat diet, the Mediterranean diet resulted in MDs of: Body weight: -2.2 (95% Cl: -3.9 – -0.6) kg BMI: -0.6 (95% Cl: -1.0 – -0.1) kg/m² SBP: -1.7 (95% Cl: -3.3 – -0.05) mm Hg DBP: -1.5 (95% Cl: -2.1 – -0.8) Fasting Plasma Glucose: -3.8 (95% Cl: -7.0 – -0.6) mg/dL Total-Cholesterol.: -7.4 (95% Cl: -10.3 – -4.4) CRP: -1.0 (95% Cl: -1.5 – -0.5) <u>1° Safety endpoint:</u> N/A 	 Overall, this study suggests the Mediterranean diet compared to the traditional low fat diet results in greater weight loss, a better CVD risk factor profile (including better BP control), and less inflammation. The number of eligible trials was small and the study samples were heterogeneous (2 2° and 4 1° prevention trials).
Yokoyama Y, et al., 2014 (88) <u>24566947</u>	Aim: Compare the effects of vegetarian and omnivorous diets on BP Study type: Systematic review and meta-analysis Size: • 7 trials (n=311). • 6 were RCT (n=198) • 4 parallel and 3 cross-over designs • All were open • Follow-up ≥6 wk (mean=15.7 wk) • Mean age=44.5 y	Inclusion criteria: • Adults ≥20 y • English language publications between Jan 1946-Nov 2013 Exclusion criteria: • Twin pt studies • Multiple interventions • Only categorical BP results	Intervention: • Lacto-ovo in 4 trials • Lacto in 1 trial • Vegan in 2 trials Comparator: Omnivorous diet in all trials	 <u>1° endpoint:</u> Compared to the omnivorous diet, the vegetarian diet resulted in MDs of: SBP: -4.8 (95% CI: -6.6– -3.1) mm Hg DBP: -2.2 (95% CI: -3.5– -1.0) SBP was lower in the vegetarian diet group in 5 of the 7 trials (significant in 3) and DBP was lower in 6 of the 7 trials (significant in 2). <u>1° Safety endpoint:</u> N/A 	 Overall, this meta-analysis of clinical trials suggested BP was lower in those who consumed a vegetarian diet compared to their counterparts who consumed an omnivorous diet. However, the trials were generally small, heterogeneous in their design and conduct, and of questionable quality. Even greater reductions in SBP and DBP were noted in a MA of 32 observational studies.
PREDIMED Toledo E, et al., 2013 (89) <u>24050803</u>	Aim: Compare the effects of a Mediterranean and lower-fat diet on BP	Inclusion criteria: • Adults, men 5,580 y, women 60–80 y • Free from CVD	Intervention: Pts assigned to a control group or to 1 of 2 Mediterranean diets.	<u>1° endpoint:</u> The percentage of pts with controlled BP increased in all 3 intervention groups (p-value for within-group changes: p<0.001). Pts	• Both the traditional Mediterranean diet and a low-fat diet exerted beneficial effects on BP and could be part of advice to pts for controlling BP.

	DM or at least 3 major	The control group	allocated to either of the 2	 However, lower values of DBP were
Study type: RCT,	CVD risk factors (smoking,	received education on	Mediterranean diet groups had	noted in the 2 groups following the
single-blinded, in	HTN, elevated LDL	following a low-fat diet,	significantly lower DBP than the pts	Mediterranean diet with extra virgin
Spanish primary	cholesterol, low HDL,	while the groups on	in the control group (-1.53 mm Hg	olive oil or with nuts than in the control
healthcare centers	overweight/obese, family	Mediterranean diets	(95% CI: -2.01– -1.04) for the	group.
	history of early CHD)	received nutritional	Mediterranean diet supplemented	
<u>Size</u> : 7,447 men (55–		education and also free	with extra virgin olive oil, and -0.65	
80 y) and women (60–	Exclusion criteria: Do	foods; either extra virgin	mm Hg (95% CI: -1.15– -0.15) mm	
80 y) at high risk for	not meet criteria listed	olive oil, or nuts.	Hg for the Mediterranean diet	
CVD.	above		supplemented with nuts). No	
		Comparator: Lower fat	between-group differences in	
		diet	changes of SBP were seen	

Data Supplement 16. RCTs and Meta-analysis RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Alcohol Reduction) (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Xin X, et al., 2001 (90) <u>11711507</u>	Aim: Study the effect of alcohol reduction on BP Study type: Systematic review and meta-analysis Size: • 15 RCTs (25 comparisons) with 2,234 pts. • 6 trials were conducted in normotensives (269 pts with a mean age ranging from 26.5– 45.5 y). Average	Inclusion criteria: • RCT in humans • Publication between 1966-1999 • Duration ≥1 wk • Only pts regularly consuming alcohol • Only difference between the comparison groups was alcohol intake <u>Exclusion criteria</u> : Comparison of different doses of alcohol intake	Intervention: Reduction in alcohol consumption. In most trials this was achieved by randomization to "light" alcohol but some RCT were based on a behavioral intervention aimed at reducing the number of drinks consumed. <u>Comparator</u> : Usual consumption of alcohol	 <u>1° endpoint</u>: Overall, alcohol reduction was associated with a significant reduction in mean SBP of -3.31 (95% Cl: -4.102.52) and DBP of -2.04 (95% Cl: -2.581.49). In the subgroup of 7 RCTs in persons with HTN, the mean changes in SBP and DBP were -3.9 (95% Cl: -5.042.76) and -2.41 (95% Cl: -3.021.57). In the subgroup of 6 RCTs in normotensives the corresponding changes in SBP and DBP were -3.5 (95% Cl: -4.612.51) and -1.80 (95% Cl: -3.030.58). 	 This is the most recent meta-analysis of this topic. Although this meta-analysis reports % reduction in alcohol intake, most trials aimed at reducing the number of alcoholic drinks consumed achieved a reduction of about 3 drinks/d. The intervention results were consistent with the relationship alcohol and BP in observational epidemiology – about a 1 mm Hg higher SBP per alcoholic drink consumed. In observational studies, type of alcohol does not seem to matter and at lower levels of alcohol consumption (<1 standard size alcoholic drink per day in women and <2 in men) there does not

	consumption of alcohol at baseline was not reported. Follow-up varied from 1–18 wk			 In a meta-regression analysis, a dose-response was noted between % reduction in alcohol consumption and mean reduction in BP. <u>1° Safety endpoint</u>: N/A 	 seem to be an important biological effect of alcohol on BP. The relationship between alcohol consumption and BP is predictable and consistent in observational and RCT studies. However, the relationship between alcohol consumption and CVD is more complex as alcohol is associated with an apparently beneficial effect on CVD risk, possibly mediated by an increase in HDL-cholesterol. Pregnant women, pts with HTN and those at risk of a drinking problem should not drink alcohol. Established light drinkers (<2 standard drinks/d in men and <1/d in women) who are normotensive are in a favorable risk category for CVD.
Stewart SH, et al., 2008 (91) <u>18821872</u>	Aim: Study the effect of reduced alcohol intake on BP. Study type: Randomized, controlled factorial trial. Size: 1,383 pts.	Inclusion criteria: • Alcohol dependence. • 4—21 d of abstinence. • Men: >21 drinks/wk; Women >14 drinks/wk. • At least 2 heavy drinking days within a consecutive 30-d period during 90 d prior to baseline. Exclusion criteria: • Other substance abuse. • Psychiatric disorder requiring medication. • Unstable medical condition	Intervention: Pharmacotherapy (naltrexone, acamprosate, or both) and counseling strategies (behavioral and/or medical management). Comparator: Placebo.	 Change in BP: Based on up to 5 repeated measures of BP over 16 wk. Data modeled to estimate change in BP over time. For pts with higher than average baseline SBP (>132 mm Hg), SBP declined by an average of 12 mm Hg (149—137) in the intervention arm compared to placebo, with a corresponding decline in DBP of 8 mm Hg. For those with a baseline SBP ≤132 mm Hg there was no change in SBP (120—121 mm Hg) or DBP. Safety endpoint: N/A 	 This trial was designed to evaluate interventions for treatment of alcohol dependence. BP measurements were not standardized. About 20% of the observations were missing and assumed to be random.
Dickenson HO, et al., 2006 (92) <u>16508562</u>	<u>Aim</u> : Study effectiveness of lifestyle	Inclusion criteria: • Only parallel trials	Intervention: Lifestyle change aimed at reduced consumption of alcohol	<u>1° endpoint</u> : -Net reduction (95% CI): SBP -3.8 (-6.1— -1.4)	Relatively small number of trialsLimited details provided

Wallace P, et al.,	interventions, including reduced alcohol intake, for treatment of HTN. <u>Study type</u> : 1 of 10 meta-analyses. <u>Size</u> : 4 trials which collectively studied 305 pts <u>Aim</u> : Study	 SBP ≥140 mm Hg and/or DBP ≥85 mm Hg ≥8 wk duration BP outcome Exclusion criteria: 2° HTN or renal disease Pregnant women Change in BP meds during trial Inclusion criteria: 	Comparator: Usual care Usual care	DBP -3.2 (-5.0— -1.4) <u>Safety endpoint</u> : N/A <u>Endpoints</u> :	 The goal was to blind those
1988 (93) <u>3052668</u>	effectiveness of general practitioner advice to reduce heavy drinking. <u>Study type</u> : • RCT <u>Size</u> : 909 adults (641 men and 268 women)	Heavy drinking during wk prior to screening interview. <u>Exclusion criteria</u> : None mentioned	counselling aimed at reduced consumption of alcohol. <u>Comparator</u> : Usual care	 1° outcome was reduction in percent with heavy consumption of alcohol (mean net change=46%). Liver enzymes and BP also measured at 6 and 12 mo. Pretreatment SBP/DBP=133.5/79.9 mm Hg. Net reduction SBP=-2.12 (95% CI: -4.190.00) Safety endpoint: N/A	 conducting the outcome assessment to treatment assignment but by 6 mo assignment was known in 20-30% of the participants. A reduction in SBP was noted despite use of a modest intervention.
Lang T, et al., 1995 (94) <u>8596098</u>	<u>Aim</u> : Worksite study of reduced alcohol intake effect on BP in heavy drinkers with HTN. <u>Study type</u> : RCT <u>Size</u> : 14 site physicians; 129 adults (95% men)	Inclusion criteria: • Heavy drinking (documented by history and liver enzyme elevation). • HTN (SBP/DBP >140/90 mm Hg) Exclusion criteria: • 2° HTN • Severe liver disease • Planned move/retirement.	Intervention: Physician and worker counselling aimed at reduced consumption of alcohol. Comparator: Usual care. Duration: Follow-up visits at 1, 3, 6, and 18 mo.	Endpoints: • Baseline SBP/DBP=162.5/98.0. Although all of the workers had HTN, only about 20% were being treated with antihypertensive medications at baseline. • At 1 y, the net change in SBP=-5.5 (p<0.05). When 5 sites with <5 workers/site were excluded, the net change in SBP=-7.3 mm Hg (p<0.01). • At 2 y, the net change in SBP=-6.6 (p<0.05).	 Behavioral intervention state of the art for its time Careful measurements of BP using Hawksley RZ sphygmomanometer. Main analyses do not seem to have accounted for cluster design.

				Safety endpoint: N/A	
Roerecke M, et al., 2017 Lancet Public Health. 2017;2:e108-120.	Aim: Study the effect of reduced alcohol intake on BP. Systematic review and meta-analysis. Size: 36 RCT with 2865 participants. Design: • 15 parallel-arm trials • 21 crossover trials Setting: • 13 in hypertension • 13 in normotension • 12 HTN and NT • Only 3 trials presented data for women.	Inclusion criteria: • RCT in adult humans • Publication on or before July 13, 2016. • Full text articles. • Change in alcohol intake for ≥1 wk	Intervention: Reduction in alcohol consumption. Strategy varied from controlled inpatient administration to randomization to "light" alcohol to pragmatic primary care trials with counselling to reduce alcohol intake. Duration: Follow-up from 1 wk to 2 y (median 4 wk).	 1° endpoint: Overall, alcohol reduction was associated with a significant reduction in mean SBP of -3.31 (95% CI: -4.102.52) and DBP of -2.04 (95% CI: -2.581.49). In the subgroup of 7 RCTs in persons with HTN, the mean changes in SBP and DBP were SBP: -3.13 (95% CI: -3.932.32) DBP: -2.00 (95% CI: -2.651.35). In meta-regression analysis, there was a strong relationship between the extent of BP reduction and change in BP, with no reduction in BP for those consuming 2 or less drinks at baseline but increasing reductions in BP for those with progressively higher intakes of alcohol at baseline. For instance, in those consuming ≥6 drinks/day and reducing their alcohol intake by approximately 50%, the estimated reduction in SBP and DBP were: SBP: -5.5 (95% CI: -6.704.30) DBP: -3.97 (95% CI: -4.703.25). Similar patterns of the effect of baseline alcohol intake on treatment effect were noted for a variety of subgroups. 	N/A

Data Supplement 17. RCTs and Meta-analysis RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Calcium Supplementation)	
(Section 6.2)	

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Van Mierlo LA, et al., 2006 (95) <u>16673011</u>	Aim: Study the effect of calcium supplementation on BP Study type: Systematic review and meta-analysis Size: • 40 RCTs with 2,492 pts. • 27 RCTs in pts <140/90 mm Hg (n=1,728) • Follow-up varied from 3–208 wk (median=8.5 wk) • Age range 11–77 y (mean=43.7 y)	Inclusion criteria: • RCT in humans • Publication between 1996 and 2003 • Nonpregnant normotensive pts or hypertensive pts • Only difference between the comparison groups was magnesium intake • Follow-up ≥2 wk Exclusion criteria: Study pts having renal disease or hyperparathyroidism	Intervention: Increased calcium intake, with a range from 355–2,000 mg/d (mean=1,200 mg/d; median=1,055 mg/d), primarily as a gluconate or carbonate salt. Comparator: Placebo or usual intake – 32 double-blind.	1° endpoint:• Overall, increased calcium intake was associated with a significant reduction in mean SBP of -1.86 (95% CI: -2.910.81) and DBP of -0.99 (95% CI: -1.61- -0.37).• The reduction was slightly less but still significant in the subset of 32 double-blind trials, with a mean SBP of -1.67 (95% CI: - 2.870.47) and DBP of -0.93 (95% CIL -1.640.22).• There was no significant difference between the effect size in those with a baseline BP ≥ or<140/90 mm Hg. • The mean change in SBP and DBP for those with a baseline BP≥140/90 mm Hg (23 comparisons) was -2.17 (95% CI: - 3.780.55) and -0.95 (95% CI: - 1.890.01), respectively. • The mean in SBP and DBP for those with a baseline BP <140/90 mm Hg was -1.67 (95% CI: -3.01- -0.27) and -1.02 (95% CI: -1.85- 0.19) mm Hg, respectively.• The authors reported slightly larger effect sizes in those with a lower initial calcium intake, in trials that employed a dietary	 This is the most recent SR/MA on this topic to include RCT conducted in both normotensive and hypertensive pts. The authors interpreted their results as being consistent with a beneficial effect of calcium supplementation on BP, with about a 2 mm Hg reduction in SBP for a 1 g increase in calcium intake. This is slighter larger effect size than noted in several earlier meta-analyses. A subsequent Cochrane Collaboration meta-analysis was confined to 13 RCT in 485 adults (≥18 y) with HTN studied for ≥8 wk (Dickinson HO et al. Cochrane Database of Systematic Reviews. 2006; CD004639). The authors noted a significant reduction in mean of -2.5 (95% CI: -4.5– -0.6) for SBP but a more modest insignificant change of -0.8 (95% CI: -2.1– 0.4) for DBP. Due to the poor quality of the RCT and heterogeneity of the results, the authors concluded the reduction in SBP was likely an artifact due to bias. Although not included in most metaanalyses, calcium supplementation has been effective as a treatment in pregnant women at risk for pre-eclampsia. Several of the meta-analyses (including the 1 by van Mierlo et al) have suggested a bigger effect size in persons with a lower intake of calcium at baseline and in trials that utilized a dietary intervention.

	intervention (compared to a supplement), and in the 4 trials conducted in Asians.	• Most of the trials were of short duration and did not (have the capacity) report on potential adverse effects such renal stones.
	<u>1° Safety endpoint</u> : N/A	 In addition to being small, several trials were of uncertain quality. Overall, RCT experience provides limited and inconsistent evidence from trials of variable quality in support of calcium supplementation for prevention (or treatment) of HTN. Better evidence supports the role of calcium supplements, in conjunction with vitamin D, in strengthening bone density.

Data Supplement 18. RCTs and Meta-analyses RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Physical Activity) (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Whelton SP, et al., 2002 (96) <u>11926784</u>	Aim: Study the effect of aerobic exercise on BP Study type: Systematic review and meta-analysis Size: 38 reports (54 comparisons) with 2,419 pts; 27 of the comparisons were conducted in normotensive pts	Inclusion criteria: • English language publication between 1966–2001 • RCT in adults ≥18 y • Duration ≥2 wk • No concurrent interventions Exclusion criteria: Missing BP data	Intervention: Aerobic exercise <u>Comparator</u> : No exercise prescribed	 <u>1° endpoint</u>: For the overall group, a pooled analysis of experience in 53 trials identified a mean net change in SBP of - 3.84 (95% Cl: -4.97– -2.72). In subgroup analysis, the effect was noted in different ethnic groups, in trials that employed different designs, durations, and sample sizes, in trials with obese, overweight or normal weight pts, and in trials that employed different types, intensity levels, and duration of aerobic exercise. In the subgroup of 15 trials in hypertensives, the mean net change in SBP was -4.94 (95% Cl: -7.17– -2.70). 	 This meta-analysis provides the most comprehensive analysis of the effect of aerobic exercise on BP and provides strong evidence in support of aerobic exercise as an intervention to lower BP in normotensives. Recognizing this, many of the trials were small and of short duration.

Cornelissen VA, et al., 2013 (97) <u>23525435</u>	Aim: Study the effectof different types ofphysical activity on BPDynamic aerobicenduranceResistance trainingDynamicStatic (Isometric)Study type:Systematic review andmeta-analysisSize: Overall, 93studies (>5,000 pts)59 DynamicResistanceTraining studies5 Combined DynamicResistanceTraining studies5 Combined DynamicAerobic andResistance training4 Static (Isometric)Resistance12 Differentinterventions within 1trialAim: Study the effect	Inclusion criteria: • Parallel arm RCTs • Adults≥18 y • Peer reviewed journals up to February 2012 • Trial duration ≥4 wk Exclusion criteria: Inadequate reporting of the data	Intervention: Physical activity Comparator: No prescription of physical activity	 In the subgroup of 27 trials conducted in normotensives, the mean net change in SBP was -4.04 (95% CI: -5.32– -2.75). <u>1° Safety endpoint</u>: N/A <u>1° endpoint</u>: Overall (trials in hypertensives and normotensive), pooled experience identified a significant reduction in BP with all forms of physical activity (aerobic and both forms of resistance training), with mean reductions in SBP of -3.5 mm Hg following aerobic endurance training, - 1.8 mm Hg following dynamic resistance training, and -10.9 mm Hg following static (isometric) resistance training (p<0.001 for the difference between the effect size following static [isometric] and other forms of physical activity). In subgroup analysis, dynamic aerobic endurance and dynamic resistance training resulted in mean SBP changes of -2.1 (95% CI: -3.3 – 0.83) and -4.3 (95% CI: -7.7 – 0.90), respectively, in the pts with pre-HTN and smaller, nonsignificant reductions in the remaining pts with a normal BP. <u>Safety endpoint</u>: N/A 	 Most recent in a series of progressively updated publications from Dr. Cornelissen and her colleagues. The findings suggest a beneficial effect of all forms of physical activity on BP, with a disproportionately large effect of resistance training on BP. Many of the available RCTs have been small, of short duration, and of uncertain quality. Suggests resistance training is
2013 (98) <u>23541664</u>	of resistance exercise on BP Study type: Systematic review and meta-analysis	 RCTs in adults (≥18 y) BP-lowering 1° outcome 	Dynamic resistance training but overall reporting of the details was poor.	(hypertensive and normotensive pts) identified a small, nonsignificant reduction in mean SBP of -1.03 (95% CI: -3.44–0.39). The corresponding finding	effective in lowering BP and was the basis for recommending this intervention in the Canadian HTN Education Program recommendations.

	Size: 9 RCTs (11 intervention groups and 14 comparisons) conducted in 452 pts. 10 (71%) of the 14 comparisons were conducted in normotensives	 Trial duration ≥4 wk Resistance training only intervention Exclusion criteria: Handgrip/isometric exercise 	<u>Comparator</u> : No resistance training but not detailed in this article	for DBP was -2.19 (95% CI: -3.87– - 0.51). <u>Safety endpoint</u> : N/A	• The discrepancy in effect size between this meta-analysis and the 1 conducted by Cornelisson et al may have been due to the more restrictive requirement by Rossi et al that change in BP be the 1° outcome.
Garcia-Hermosa A, et al., 2013 (99) <u>23786645</u>	Aim: Study the effect of exercise on BP in obese children. Systematic review and meta-analysis. Size: 9 RCTs (410 pts).	Inclusion criteria: • Children ≤14 y with obesity • RCT • Duration ≥8 wk • 1° outcome: change in BP <u>Exclusion criteria</u> : Concomitant intervention	Intervention: Physical activity, principally aerobic exercise. Comparator: No physical exercise, nutrition, education, or dietary restriction intervention	<u>1° endpoint</u> : Change in SBP: In pooled analysis, mean change in SBP was -0.4 (95% CI: -0.66– -0.24). <u>Safety endpoint</u> : N/A	 This meta-analysis focused specifically on the effect of physical activity on BP in children with obesity. Although it is not stated explicitly, it seems likely that all of the participants were normotensive and not receiving medication that could influence level of BP. The findings are consistent with other meta-analyses of the effect of physical activity on BP. Only limited information regarding study details is provided in this publication. The interventions were heterogeneous in type, duration, and quality.
Carlson DJ, et al., 2014 (100) <u>24582191</u>	Aim: Study the effect of physical activity on BP in children with obesity. Study type: Systematic review and meta-analysis. Size: 9 RCTs (223 pts: 127 intervention and 96 controls): 6 were conducted in normotensives.	Inclusion criteria: • Adults ≥18 y • RCT, including cross-over trials. • Duration ≥4 wk • Published in a peer reviewed journal between January 1, 1966 and July 31, 2013 Exclusion criteria: Studies that employed any intervention other	Intervention: Pure isometric exercise. Comparator: Use of a control group was a requirement but no additional specific information provided.	 <u>1° endpoint</u>: In the overall pooled analysis (hypertensive and normotensive trials), mean change in SBP was -6.77 (95% CI: -7.935.62) mm Hg. In the subgroup of 3 trials with hypertensive pts (all on antihypertensive medication), the mean change in SBP was -4.31 (95% CI: -6.422.21) mm Hg. In the subgroup of 6 trials with normotensive pts, the mean change in SBP was -7.83 (95% CI: -9.216.45) mm Hg. 	 This study provides information regarding the effect of pure isometric exercise interventions on BP in adults. The BP reductions reported in this meta-analysis are surprisingly large but the overall effect pattern is quite consistent with other meta-analyses of isometric exercise.

Cornelissen VA, et al., 2011 (101) <u>21896934</u>	Aim: Study the effect of resistance training on BP. Study type: Meta-	than pure isometric exercise (e.g., dynamic resistance) Inclusion criteria: • Adults ≥18 y • RCT, including cross-over trials. • Duration ≥4 wk	Intervention: Resistance training, including isometric and dynamic modalities.	<u>Safety endpoint</u> : N/A <u>1° endpoint</u> : Resistance training induced a significant SBP/DBP reduction in 28 normotensive or prehypertensive study groups of -3.9 (-6.4, -1.2)/-3.9 (- 5.6, -2.2] mm Hg). In the 5 hypertensive	 This meta-analysis supports the BP-lowering potential of dynamic resistance training and isometric handgrip training. Results further suggest that
	analysis <u>Size</u> : 28 randomized, controlled trials, involving 33 study groups and 1,012 pts.	Published in a peer reviewed journal up to June 2010 <u>Exclusion criteria</u> : Interventions other than pure isometric exercise (e.g. dynamic resistance)	<u>Comparator</u> : Use of a control group was a requirement but no additional specific information provided.	study groups, the change in mean SBP/DBP was -4.1 (95% CI: -0.63–1.4)/- 1.5 (95% CI: -3.4–0.40) mm Hg. When the study groups were divided according to the mode of training, isometric handgrip training in 3 groups resulted in a larger decrease in SBP/DBP (-13.5 [95% CI: -16.5– -10.5]/-6.1[95% CI: -8.3– -3.9] mm Hg) than dynamic resistance training in 30 groups (-2.8 [95% CI: -4.3– -1.3]/-2.7 [95% CI: -3.8– -1.7] mm Hg). Safety endpoint: N/A	 isometric handgrip training may be more effective for reducing BP than dynamic resistance training. However, given the small amount of isometric studies available, additional studies are warranted to confirm this finding.

Data Supplement 19. RCTs and Meta-analysis RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Magnesium Supplementation) (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Kass L, et al., 2012 (102) <u>22318649</u>	<u>Aim</u> : Study the effect of magnesium supplementation on BP <u>Study type</u> : Systematic review and meta-analysis	Inclusion criteria: • RCT in humans • Parallel or cross- over design • Publication before July 2010 • Adults >18 y • Only difference between the	Intervention: Increased magnesium intake, with a range in elemental magnesium of 120 to 973 mg/d and a mean of 410 mg/d. Comparator: Placebo or usual intake	<u>1° endpoint:</u> • Overall, increased magnesium intake was associated with a small nonsignificant reduction in mean SBP of -0.32 (95% CI: - 0.41– -0.23) and DBP of -0.36 (95% CI: -0.44– -0.27).	 This is the most recent systematic review/meta-analysis on this topic. The authors interpreted their results as being consistent with a beneficial effect of magnesium supplementation on BP. However, this interpretation seems at odds with the data. In an earlier meta-analysis of 20 RCT (6 in normotensives) by Jee Systolic

Size: 22 RCTs (23 comparisons) with 1,173 pts. Data for RCTs conducted in normotensive pts were not presented. However, most RCTs were conducted in normotensives and only 6 of the RCTs included some (or all) pts who were being treated with antihypertensive medication. Overall mean age was -50 y. Follow-up varied from 3–24 wk, with a mean of 11.3 wk.	comparison groups was magnesium intake Exclusion criteria: Comparison of different doses of alcohol intake		 Forest plots revealed considerable heterogeneity in effect size. The authors reported slightly larger effect sizes in subgroup analysis of cross-over RCT and RCT that employed a dose of magnesium >370 mg/d. <u>1° Safety endpoint</u>: N/A 	 HTN et al (Am J Hyperts. 2002;15:691-696) magnesium supplementation resulted in small mean NS reductions of -0.6 (95% CI: -2.2–1.0) mm Hg in SBP and -0.8 (95% CI: -1.9–0.4) in DBP. In meta-regression analysis, there was an apparent dose-response with SBP and DBP reductions of -4.3 (95% CI: -6.3–2.2) and -2.3 (95% CI: -4.9–0) mm Hg for each 10 mmol/d higher level of magnesium intake. A Cochrane systematic review/meta-analysis of magnesium supplementation for treatment of HTN in adults (Dickinson HO et al. Cochrane Database Systematic Review 2006: CD 004640) included 12 RCT (n=545) with follow-up of 8–26 wk. Overall, mean SBP and DBP were reduced by -1.3 (95% CI: -4.0–1.5) and -2.2 (95% CI: -3.4–0.9) mm Hg, respectively. The authors noted the studies were of poor quality, with considerable heterogeneity, and felt the results were likely biased. Some authors have suggested there may be a greater BP effect when the intervention is by means of diet change but there is insufficient RCT evidence to support this position. Magnesium sulfate is the drug of choice for prevention of seizures in the pre-eclamptic woman, or prevention of recurrence of seizures in the eclamptic woman, as demonstrated in RCT and a 2010 Cochrane review (Duley L et al. Cochrane Database of Systematic Reviews. CD000127, 2010). Overall, RCT experience provides insufficient evidence to recommend oral
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		magnesium supplementation as a means to prevent (or treat) HTN.

Data Supplement 20. RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Weight Loss) (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; and 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Neter JE, et al., 2003 (103) <u>12975389</u>	<u>Aim</u> : Study the effect of weight loss on BP <u>Study type</u> : Systematic review and meta-analysis <u>Size</u> : 25 RCTs (34 comparisons) with 4,874 pts; 17 of the comparisons were conducted in normotensive pts	Inclusion criteria: • RCT in humans • English language publication between 1966– 2002 • Nonpharmacologic intervention Exclusion criteria: • Duration <8 wk • Missing data • Objective not weight loss • Concomitant intervention(s)	Intervention: Weight loss (calorie reduction, physical activity, or combination of both) Comparator: No weight loss prescription	 <u>1° endpoint</u>: For the overall group, mean baseline body weight was 88.3 kg and mean change in body weight following the application of the weight loss intervention was -5.1 (95% CI: -6.034.25) kg. This represents a mean percent change of -5.8%. There was strong evidence for a BP lowering effect of weight loss on BP, overall and in normotensive subgroup. In the normotensive group, the mean for change in SBP was 4.08 (95% CI: -6.012.16). Overall, a 1 kg reduction in body weight was associated with a mean change in SBP of -1.05 (95% CI: -1.430.66) mm Hg. 	 Substantial evidence for a reduction in BP, overall and in normotensives. With the exception of the mean (95% Cl) changes in BP, this paper provides limited data for the normotensive group
Ho M, et al., 2012 (104) <u>23166346</u>	<u>Aim</u> : Study the effect of lifestyle weight loss interventions in obese/overweight children on weight	Inclusion criteria: • RCTs, in obese/overweight children and adolescents ≤18 y	Intervention: Lifestyle weight loss program with a dietary component <u>Comparator</u> : No treatment, usual care or	<u>1° endpoint</u> : Pooled experience in the 7 RCTs with BP experience identified a significant reduction in mean SBP of - 3.40 (95% CI: -5.19– -1.61). The pooled SBP MD was -3.72 (95% CI: -4.74– - 2.69) in the 3 RCTs with a duration >1 y	• Findings in children are consistent with experience in adult normotensives and with experience in hypertensive pts.

	change and cardio- metabolic risk factors <u>Study type</u> : Systematic review and meta-analysis <u>Size</u> : • Overall, 38 studies • 33 included in various meta-analyses • Effect on SBP studied in 7 RCTs that included 554 pts	 English language publications between 1975– 2010 Trial duration ≥2 mo Exclusion criteria: Studies that targeted prevention/weight maintenance Drug trials Trials in persons with an eating disorder Inadequate reporting of the data 	written education materials	<u>Safety endpoint</u> : N/A	Considerable heterogeneity in the data
Cai L, et al., 2014 (105) <u>24552832</u>	Aim: Study the effect of childhood obesity prevention programs on BP Study type: Systematic review and meta-analysis Size: Overall study included 23 studies (28 comparisons) conducted in 18,925 pts.	Inclusion criteria: • RCTs, quasi-experimental studies, and natural experiments in humans • Children and adolescents 2–18 y • Conducted in a developed country • English language publications • Trial duration ≥1 y (≥6 mo for school-based intervention studies) Exclusion criteria: • Studies that only targeted obese/overweight children or those with a medical condition • Inadequate reporting of the data	Intervention: • Weight loss • 15 school-based • 12 some combination of school, home and/or community-based • 1 child care <u>Comparator</u> : No weight loss	<u>1º endpoint</u> : Pooled experience in 19 studies (20 comparisons) identified a small but significant reduction in mean SBP of -1.65 (95% CI: -2.56– -0.71). The effect size was greater in studies that employed an intervention that combined diet and physical activity (mean change in SBP of -2.11 mm Hg). <u>Safety endpoint</u> : N/A	 Study included a mix of RCTs (13), quasi- experimental studies (9), and natural experiments (1). Included studies conducted over several decades (1985–2012). A significant reduction in BP was only noted in the studies conducted between 2000–2009: mean change in SBP of -3.73 (95% CI: - 5.372.09) Findings of a BP reduction in childhood consistent with evidence from the publications by Neter and Ho.
TOHP, Phase II Hypertension Prevention Collaborative Research Group,	<u>Aim</u> : Study the effect of weight loss on BP and prevention of HTN.	Inclusion criteria: • Healthy community-dwelling adults 30–54 y	Intervention: Behavior change intervention (combination of diet change and physical activity) aimed at	1° endpoint: Change in SBP • Compared to usual care, the weight loss group experienced a significant mean reduction of -4.5 kg in body	• Largest trial of weight loss in prevention of HTN and also provides the longest duration of follow-up

1997 (106) <u>9080920</u>	Study type: Randomized, controlled factorial trial. Size: 2,382 pts, of whom 1,192 were randomized to a weight loss intervention and 1,190 were randomized to a no weight loss intervention.	 BMI between 110% and 165% of desirable body weight Not taking BP-lowering medication Mean SBP <140 mm Hg and DBP 83-89 mm Hg Exclusion criteria: Taking antihypertensive medication Heart disease, renal disease, poorly controlled hyperlipidemia or DM, DM requiring insulin, special dietary requirements >14 drinks/wk 	studying the effects of a modest reduction in body weight during up to 48 mo (minimum 36 mo) of follow-up. <u>Comparator</u> : Usual care group	 weight and -3.7 (SD: 0.5; p<0.001) mm Hg in SBP at 6 mo (-6.0 mm Hg in the weight loss group and -2.2 mm Hg in the usual care group). A progressive reduction in the effect sizes for body weight and BP was noted over time, with mean for SBP at 18, 36 mo and termination of -1.8 (SD: 0.5; p<0.001), -1.3 (SD: 0.5; p=0.01), and - 1.1 (SD: 0.5; p=0.04). <u>Prevention of HTN</u> At 6 mo of follow-up the incidence of new onset HTN was 42% lower in the participants randomized to weight loss compared to the usual care group (p=0.02). During more prolonged follow-up, the effect size decreased but remained borderline significant after 48 mo of follow-up (13% reduction; p=0.06). Overall, the incidence of HTN was reduced by 21% (p=0.02). <u>Safety endpoint</u>: N/A 	 The assumptions for a main effects factorial analysis (independence of the interventions) were not demonstrated. Given this finding, the most reliable analysis of this trial was comparison of the experience in each active intervention group with the usual care group. This results in a reduction in statistical power. Consistent with the pattern in the proceeding TOHP I trial weight loss reduced BP and the incidence of HTN but the effect sizes for weight loss and BP as well as the difficulty of maintaining the intervention in highly motivated and extensively counselled participants underscores the difficulty of achieving and maintaining ideal body weight in the general population by means of lifestyle change.
TOHP, Phase I 1992 (79) <u>1586398</u>	<u>Aim</u> : Study the effect of weight loss on BP and prevention of HTN <u>Study type</u> : Randomized, controlled factorial trial. <u>Size</u> : Overall, 2,182 adults, with the 308	Inclusion criteria: • Community- dwelling adults 30–54 y • Not on antihypertensive medication • DBP 80-89 mm Hg • Healthy Exclusion criteria: • Disease	Intervention: Behavior change intervention (combination of diet change and physical activity) Comparator: Usual care	<u>1° endpoint</u> : Change in DBP <u>2° endpoint: Change in SBP <u>Safety endpoint</u>: CVD events, symptoms and general and well being</u>	 Significantly lower DBP (2.3 mm Hg; p<0.01) and SBP (2.9 mm Hg; p<0.01) in the weight loss group compared to usual care Few CVD events No difference in symptoms Significant improvement in general well-being at 6 and 18 mo (p<0.05)

	assigned to weight loss compared to 256 usual care controls	 Inability to comply with the protocol 			
TONE Whelton PK, et al., 1998 (107) <u>9515998</u>	Aim: Study the effect of weight loss on BP and need for antihypertensive drug therapy Study type: RCT, factorial design Size: 585 (obese) participants	Inclusion criteria: • Community-dwelling adults 60–80 y • SBP <145 mm Hg and DBP	Intervention: Behavior change intervention (combination of diet change and physical activity) Comparator: Usual care, with similar level of contact compared to active intervention group	 <u>1° endpoint</u>: Recurrence of HTN following withdrawal of antihypertensive medication (or CVD event) <u>2° endpoint</u>: BP (while still on antihypertensive medication prior to tapering of medication) <u>Safety endpoint</u>: CVD events, symptoms (including headaches), dietary composition 	 Significant reduction in SBP prior to withdrawal of antihypertensive medication (mean±SE=-4.0±1.3 mm Hg) 1° outcome significantly less common in weight loss group compared to usual care – Rel. HR: 0.70; 95% CI, 0.57–0.87; p<0.001 No overt evidence for adverse effects of intervention

Data Supplement 21. RCTs and Systematic Reviews for RCTs Studying the Effect of Nonpharmacologic Interventions on BP (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
TOHP, Phase II (Weight Loss component) 1997 (1) <u>9080920</u>	<u>Aim</u> : Study the effect of weight loss on BP and prevention of HTN. <u>Study type</u> : Randomized, controlled factorial trial.	Inclusion criteria: • Healthy community- dwelling adults 30–54 y • BMI between 110% and 165% of desirable body weight • Not taking BP-lowering medication • Mean SBP <140 mm Hg and DBP 83-89 mm Hg	Intervention: Behavior change intervention (combination of diet change and physical activity) aimed at studying the effects of a modest reduction in body weight during up to 48 mo (minimum 36 mo) of follow-up.	 <u>1° endpoint</u>: <u>Change in SBP</u> Compared to usual care, the weight loss group experienced a significant mean (standard error) reduction of -4.5 kg in body weight and -3.7 (0.5) (p<0.001) mm Hg in SBP at 6 mo (-6.0 mm Hg in the weight loss group and -2.2 mm Hg in the usual care group). A progressive reduction in the effect sizes for body weight and BP 	 This was the largest trial of weight loss in prevention of HTN and also provides the longest duration of follow-up The assumptions for a main effects factorial analysis (independence of the interventions) were not demonstrated. Given this finding, the most reliable analysis of this trial was comparison of the experience in each active intervention group with the usual care group. This results in a reduction in statistical power.

	Size: 2,382 pts, of whom 1,192 were randomized to weight loss and 1,190 were randomized to no weight loss intervention	Exclusion criteria: • Taking antihypertensive medication • Heart disease, renal disease, poorly controlled hyperlipidemia or DM, DM requiring insulin, special dietary requirements • >14 drinks/wk.	<u>Comparator</u> : Usual care group	 was noted over time, with mean (SD) for SBP at 18, 36 mo and termination of -1.8 (0.5) (p<0.001), - 1.3 (0.5) (p=0.01), and -1.1 (0.5) (p=0.04). <u>Prevention of</u> HTN At 6 mo of follow-up the incidence of new onset HTN was 42% lower in the participants randomized to weight loss compared to the usual care group (p=0.02). During more prolonged follow-up, the effect size decreased but remained borderline significant after 48 mo of follow-up (13% reduction; p=0.06). Overall, the incidence of HTN was reduced by 21% (p=0.02). Safety endpoint: N/A 	• Consistent with the pattern in the proceeding TOHP I trial weight loss reduced BP and the incidence of HTN but the effect sizes for weight loss and BP as well as the difficulty of maintaining the intervention in highly motivated and extensively counselled participants underscores the difficulty of achieving and maintaining ideal body weight in the general population by means of lifestyle change.
TONE (Weight Loss component) Whelton PK, et al., 1998 (3) <u>9515998</u>	<u>Aim</u> : Study the effect of weight loss on BP and need for antihypertensive drug therapy <u>Study type</u> : RCT, factorial design <u>Size</u> : 585 (obese) participants	Inclusion criteria: • Community-dwelling adults 60-80 y • SBP <145 mm Hg and DBP <85 mm Hg on 1 antihypertensive medication Exclusion criteria: • Heart attack or stroke within 6 mo • Current angina, HF, insulin-dependent DM • Inability to comply with protocol	Intervention: Behavior change intervention (combination of diet change and physical activity) Comparator: Usual care, with similar level of contact compared to active intervention group	1° endpoint: Recurrence of HTN following withdrawal of antihypertensive medication (or CVD event) 2° endpoint: BP (while still on antihypertensive medication prior to tapering of medication) Safety endpoint: CVD events, symptoms (including headaches), dietary composition	 Significant reduction in SBP prior to withdrawal of antihypertensive medication (mean±standard error=-4.0±1.3 mm Hg) 1° outcome significantly less common in weight loss group compared to usual care – Rel. HR: 0.70; 95% CI: 0.57– 0.87; p<0.001 No overt evidence for adverse effects of intervention
TOHP, Phase I (Weight Loss component) 1992 (4) <u>1586398</u>	<u>Aim</u> : Study the effect of weight loss on BP and prevention of HTN	 Inclusion criteria: Community-dwelling adults 30–54 y 	Intervention: Behavior change intervention (combination of diet change and physical activity)	<u>1° endpoint</u> : Change in DBP <u>2° endpoint</u> : Change in SBP	 Significantly lower DBP (2.3 mm Hg; p<0.01) and SBP (2.9 mm Hg; p<0.01) in the weight loss group compared to usual care Few CVD events

<u>Study type</u> : Randomized, controlled factorial trial.	 Not on antihypertensive medication DBP 80-89 mm Hg Healthy 	Comparator: Usual care	Safety endpoint: CVD events, symptoms and general and well being	 No difference in symptoms Significant improvement in general well- being at 6 and 18 mo
Size: Overall, 2,182 adults, with the 308 assigned to weight loss compared to 256 usual care controls	Exclusion criteria: • Disease • Inability to comply with the protocol			

Data Supplement 22. Observational Studies of CV Target Organ Damage Including LVH (Section 7.2)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
LIFE Devereux RB, et al., 2004 (108) <u>15547162</u>	Study type: Sub- study of pts with HTN and ECG LVH Size: 941	Inclusion criteria: • 55–80 y • BP 160–200/95–115 mm Hg • No MI or stroke within 6 mo • Had echo • Did not require treatment with BB, ACE or AT-1 antagonist for other reasons Intervention: Treatment to BP of 140/90 mm Hg beginning with pts randomized to losartan or atenolol	 <u>1° endpoint</u>: Change in LV mass assessed by echo and change in BP in relation to CVD events <u>Results:</u> Composite endpoint of CV death, MI, or stroke reached in 104 in 4.8 y of follow-up Reduction in 1° endpoint per SD reduction in LV mass independent of BP change OR: 0.74 (95% CI: 0.6–0.91; p=0.003) Reductions for each composite endpoint component and total mortality were also significant; results independent of change in ECG LVH 	Reduction in LV mass by echo independently related to CVD outcomes
CARDIA Armstrong AC, et al., 2014 (109) <u>24507735</u>	Study type: Observational study of population-based cohorts	Inclusion criteria: African American and white men and women stratified by education (above/below high school) 18– 30 y at study start and followed for over 20 y; previously healthy	 <u>1° endpoint</u>: Composite of hard CVD events <u>Results:</u> LV mass indexed to body surface area or to height predicted CV events independently of the Framingham risk score (HR: 1.21; 95% Cl: 1.05–1.39; p<0.007) 	 LV mass measured at age 18– 30 y leads to modest risk reclassification later in life Low number of events limits generalizability

	<u>Size</u> : 3,980		• Net reclassification improvement for LVM/height was 0.13 (p<0.01) and for LVM/BSA was 0.11 (p=0.02).	
ARIC Okwuosa TM, et al., 2015 (110) <u>25497261</u>	Study type: Observational study of population-based cohorts <u>Size</u> : 14,489	Inclusion criteria: African American and white men and women population-based cohort mean age 54.7 ± 5.7 y at study start and followed for over 25 y; previously healthy	 <u>1° endpoint</u>: Pooled cohort CV events and 10-y Framingham CVD events <u>Results:</u> 792 (5.5%) 10-y Pooled Cohort CV events and 690 (4.8%) 10-y Framingham CHD events. LVH was associated with CVD events (HR: 1.62; 95% CI: 1.38–1.90) and CHD events (HR: 1.56; 95% CIL 1.32–1.86. LVH by ECG did not significantly reclassify or improve C statistic compared with Framingham risk score (C statistics 0.767/0.719; net reclassification index =0.001 [p=not significant]), compared with (C statistics 0.770/0.718), respectively. 	ECG LVH does not improve risk reclassification
MESA Zalawadiya SK, et al., 2015 (111) <u>24699336</u>	Study type: Observational study of population-based cohorts Size: 4,921	Inclusion criteria: Multi-ethnic cohort of men and women followed for a mean follow-up of 4.5 y	<u>1° endpoint</u> : Hard CVD endpoints <u>Results:</u> MRI calculated LVH (indexed to BSA or height; >95 th percentile) predicted hard CVD events (LVH-BSA: HR: 2.36; 95% CI: 1.37–4.04; p=0.002; LVH-height [1.7]: HR: 1.95; 95% CI: 1.17– 3.26; p=0.01). but did not improve risk reclassification beyond conventional risk factors	• Though LVH predicted events it did not improve risk reclassification

Data Supplement 23. RCTs on Use of Risk Estimation to Guide Treatment of Hypertension (Section 8.1.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% Cl)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events Summary
Sundstrom J, et al., 2014 (112) <u>25131978</u>	<u>Aim</u> : We aimed to investigate whether the benefits of BP- lowering drugs are proportional to baseline CV risk, to	Inclusion criteria: BPLTTC: trials were eligible if they met the original inclusion criteria specified in the protocol, 11 and were part of the subset of studies that randomly allocated	Intervention: BP-lowering meds Comparator: Placebo or less intensive treatment	<u>1° endpoint</u> : • Total major CV events, consisting of stroke (nonfatal stroke or death from cerebrovascular disease), CHD (nonfatal MI or death from CHD	Summary: • Lowering BP provides similar relative protection at all levels of baseline CV risk, but progressively greater absolute risk reductions as baseline risk

	establish whether absolute risk could be used to inform treatment decisions for BP-lowering therapy, as is recommended for lipid-lowering therapy. <u>Study type</u> : Meta- analysis of RCTs <u>Size</u> : 11 trials and 26 randomized groups with 67,475 pts (51,917 pts data available for the calculation of the risk equations)	pts to either a BP-lowering drug or placebo, or to a more intensive or less intensive BP regimen. Trials had to have a minimum of 1,000 pt-y of planned follow-up in each randomized group, and should not have presented their main results before the protocol was finalized in July, 1995. <u>Exclusion criteria</u> : Not stated		 including sudden death), HF (resulting in death or admission to hospital), or CV morbidity. The mean estimated baseline levels of 5-y CV risk for each of the 4 risk groups were 6.0% (SD: 2–0), 12.1% (1–5), 17.7% (1–7), and 26.8% (5–4). In each consecutive higher risk group, BP-lowering treatment reduced the risk of CV events relatively by 18% (95% CI: 7–27), 15% (95% CI: 4–25), 13% (95% CI: 2–22), and 15% (95% CI: 5–24), respectively (p=0.30 for trend) in each group with BP-lowering treatment for 5 y would prevent 14 (95% CI: 8–21), 20 (95% CI: 8–31), 24 (95% CI: 8–40), and 38 (95% CI: 16–61) CV events, respectively (p=0.04 for trend). 	increases. These results support the use of predicted baseline CVD risk equations to inform BP-lowering treatment decisions. • Lowest risk group had >83% with a risk that exceeds 4%.
Sundstrom J, et al., 2015 (19) <u>25531552</u>	<u>Aim:</u> To investigate whether pharmacologic BP reduction prevents CV events and deaths in pts with grade 1 HTN. <u>Study type:</u> Meta- analysis of RCTs <u>Size:</u> 10 RTCs with 15,266 pts	Inclusion criteria: RCTs of at least 1 y duration; pts ≥18 y, at least 80% of whom had grade 1 HTN and no previous CVD (MI, angina pectoris, CABG, PCI, stroke, TIA, carotid surgery, peripheral arterial surgery, intermittent claudication, or renal failure); and compared an antihypertensive drug provided as monotherapy or a stepped-care algorithm vs. placebo or another control regimen. Exclusion criteria: Excluded trials did not contribute an event	Intervention: BP-lowering meds Comparator: • Placebo or less intensive treatment • The difference in average achieved BP between the active and control groups was 3.6/2.4 mm Hg in the BPLTTC (Appendix Table 2, available at www.annals.org) but is unknown for the other contributing trial subgroups.	 <u>1° endpoint</u>: Total major CV events, comprising stroke (nonfatal stroke or death from cerebrovascular disease), coronary events (nonfatal MI or death from CHD, including sudden death), HF (causing death or resulting in hospitalization), or CV death; OR: 0.86 (95% CI: 0.74–1.01) <u>Other endpoints:</u> Each of the above outcomes independently; and total deaths. CHD 0.91 (95% CI: 0.74–1.12) Stroke 0.72 (95% CI: 0.55–0.99) HF 0.80 (95% CI: 0.57–1.12) CVD deaths 0.75 (95% CI: 0.57– 0.98) 	Summary: • BP-lowering therapy is likely to prevent stroke and death in pts with uncomplicated grade 1 HTN. • 5 y risks in BPLTTC control groups CVD events 7.4% CVD deaths 3.1%

Thompson AM, et al., 2011 (113) 21364140	Aim: To evaluate the effect of antihypertensive treatment on 2° prevention of CVD events and all-cause mortality among pts without clinically defined HTN. <u>Study type:</u> Meta- analysis of RCTs <u>Size:</u> 25 RCTs with 64,162 pts	for any of the outcomes of interest. Inclusion criteria: Studies were eligible for inclusion if they were RCTs of antihypertensive treatment among pts with BP <140 mm Hg systolic or <90 mm Hg diastolic for the prevention of CVD events (fatal or nonfatal stroke, fatal or nonfatal MI, CHF, or CVD mortality). Exclusion criteria: Studies were excluded if CVD events were not reported by HTN status in studies that included pts with and without HTN; the study population did not include pts with BP in the normal or prehypertensive ranges; the study population did not include pts with preexisting CVD or CVD equivalents, such as diabetes; antihypertensive treatment was not part of the intervention; treatment allocation was not random; a measure of	Intervention: BP-lowering meds, the majority were studies of ACEI, next most common were BBs. Comparator: Placebo or active comparator	 Total deaths 0.78 (95% CI: 0.67– 0.92) Only the first event for a pt was used for the analysis of each outcome, but a pt who had >1 outcome type could contribute to more than 1 analysis. They also tabulated overall withdrawals and withdrawals due to adverse events. <u>P endpoint</u>: Composite CVD (fatal or nonfatal stroke, fatal or nonfatal MI, CHF, or CVD mortality): CVD RR: 0.85 (95% CI: 0.80– 0.90), absolute risk reduction: 27.1/1,000. This implies that a 2.7% absolute risk reduction reflects a 15% RR reduction, so the baseline risk for CVD would have been about 18%, but the follow-up interval is unclear. <u>Other endpoints:</u> Stroke RR: 0.77 (95% CI: 0.61, 0.98) MI RR: 0.80 (95% CI: 0.65, 0.77) CVD death RR: 0.83 (95% CI: 0.69, 0.99) Total deaths RR: 0.87 (95% CI: 0.80, 0.95) <u>Other results:</u> Table 4 shows similar results for CVD from studies 	Summary: Among pts with clinical history of CVD but without HTN, antihypertensive treatment was associated with decreased risk of stroke, CHF, composite CVD events, and all-cause mortality. Limitations: • Difference in achieved BP was not reported. • Average baseline SBP not reported. No information on the entry levels of BP other than not hypertensive. Difficult to use to establish a treatment threshold or goal. • Many of these studies were designed to try to demonstrate specific drug benefits rather than BP-lowering benefits. Can we attribute the benefits to BP- lowering? We know these pts did not have HTN but we do not know the lower limit of the
		intervention; treatment allocation		Other results: Table 4 shows similar results for CVD from studies of pts with CAD vs. other, HF vs. other, and DM vs. non-DM. Similar results from studies of ACEI vs. other. These results support the	did not have HTN but we do

		intervention and control groups other than antihypertensive treatment.		conclusion that the effect is not a drug effect, but is a BP-lowering effect, and that the effect is seen in people with CVD broadly defined, not just in HF pts.	establish a treatment initiation threshold or goal.
Xie X, et al., 2015 (21) <u>26559744</u>	Aim: To assess the efficacy and safety of intensive BP-lowering strategies. Study type: Meta- analysis of RCTs Size: 19 RCTs with 44,989 pts	Inclusion criteria: RCTs with at least 6 mo follow-up that randomly assigned pts to more intensive vs. less intensive BP- lowering treatment, with different BP targets or different BP changes from baseline. Reference lists from identified trials and review articles were manually scanned to identify any other relevant studies. Exclusion criteria: N/A	Intervention: BP-lowering meds Comparator: • Less intensive treatment • BP difference 6.8/3.5 • The mean follow-up BP levels in the less intensive BP-lowering regimen group were 140/81 mm Hg, compared with 133/76 mm Hg in the more intensive treatment group.	 <u>1° endpoint</u>: CVD, other major CV events, defined as a MI, stroke, HF, or CV death, separately and combined; nonvascular and all-cause mortality; ESKD, and adverse events. Progression of albuminuria (defined as new onset of micro-albuminuria/macro-albuminuria or a change from micro-albuminuria) and retinopathy (retinopathy progression of 2 or more steps) were also recorded for trials that were done in pts with DM CVD RR: 0.86 (95% CI: 0.78–0.96) <u>Other endpoints:</u> MI RR: 0.87 (95% CI: 0.76–1.00; p=0.042) Stroke RR: 0.78 (95% CI: 0.68–0.90) HF RR: 0.85 (95% CI: 0.66–1.11) CVD death RR: 0.91 (95% CI: 0.74–1.11) Total deaths RR: 0.91 (95% CI: 0.81–1.03) <u>Other results:</u> Benefit for CVD not different by baseline SBP 120–139: 0.89 (95% CI: 0.78–1.00) > 160: 0.89 (95% CI: 0.73–1.09) 	Summary: Intensive BP- lowering, including to <130 mm Hg, provided greater vascular protection than standard regimens. In high-risk pts, there are additional benefits from more intensive BP- lowering, including for those with SPB <140 mm Hg at baseline. The net absolute benefits of intensive BP- lowering in high-risk individuals are large. Limitations: • Lack of individual pt data, which would have allowed a more reliable assessment of treatment effects in different pt groups. • Interpretation: Supports treating pt with and without CVD at threshold of 130 to <130. Supports treating at threshold of about 130 even down to a CVD event rate of 0.9% per y.

				 p-heterogeneity: 0.60 Benefit for CVD not different for more intensive and less intensive targets in intensive group <140 or <150 mm Hg: 0.76 (95% Cl: 0.60–0.97) <120– <130 mm Hg: 0.91 (95% Cl: 0.84–1.00) p-hetero: 0.06 Absolute benefits were proportional to absolute risk. For trials in which all pts had vascular disease, renal disease, or DM at baseline, the average control group rate of major vascular events was 2.9% per y compared with 0.9% per y in other trials, and the numbers needed to treat were 94 (95% Cl: 44–782) in these trials vs. 186 (95% Cl: 107– 708) in all other trials. Increase in severe hypotension: 0.3% vs. 0.1% per person y OR: 2.68 (95% Cl: 1.21–5.89) 	
Ettehad D, et al., 2015 (17) <u>26724178</u>	<u>Aim:</u> This systematic review and meta- analysis aims to combine data from all published large-scale BP-lowering trials to quantify the effects of BP reduction on CV outcomes and death across various baseline BP levels, major comorbidities, and different pharmacological interventions.	Inclusion criteria: • RCTs of BP-lowering treatment that included a minimum of 1,000 pt-y of follow- up in each study arm. No trials were excluded because of presence of baseline comorbidities, and trials of antihypertensive drugs for indications other than HTN were eligible. • Eligible studies fell into 3 categories: 1st, random allocation of pts to a BP-lowering drug or placebo; 2nd, random allocation of pts to different BP-	Intervention: BP-lowering meds <u>Comparator</u> : Placebo, active comparator or less intensive treatment	1° endpoint:• CVD.• Major CVD events, CHD, stroke, HF, renal failure, and all-cause mortality.• Standardized RR for 10 mm Hg difference in SBP• CVD RR: 0.80 (95% CI: 0.77– 0.83)Other endpoints: • CHD RR: 0.83 (95% CI: 0.78– 0.88) • Stroke RR: 0.73 (95% CI: 0.68– 0.77)	Summary: • BP-lowering significantly reduces vascular risk across various baseline BP levels and comorbidities. Our results provide strong support for lowering BP to SBP <130 mm Hg and providing BP-lowering treatment to individuals with a history of CVD, CHD, stroke, DM, HF, and CKD. • In stratified analyses, we saw no strong evidence that proportional effects were diminished in trials that included people with lower

	Study type: Meta-	lowering drugs; and third,		• HF RR: 0.72 (95% CI: 0.67–0.78)	baseline SBP (<130 mm Hg),
	analysis of RCTs	random allocation of pts to		• Total deaths RR: 0.87 (95% CI:	and major CV events were
		different BP-lowering targets.		• 10/al deallis KK. 0.07 (95% Cl. 0.84–0.91)	clearly reduced in high-risk pts
	Size: 123 studies with	different Dr Howening targets.		0.84–0.91)	with various baseline
	613,815 pts	Exclusion criteria: <1,000 pt-y			comorbidities. Both of these
	013,013 μις	of follow-up in each treatment		Other results:	
				Benefit for CVD and other	major findings—the efficacy of
		group.		endpoints not different by baseline	BP-lowering below 130 mm Hg
				SBP, including <130 mm Hg fig 4	and the similar proportional
				in paper	effects in high risk
				CVD: 0.63; 95% CI: 0.50–0.80;	populations—are consistent
				p=0.22	with and extend the findings of
				CHD: 0.55; 95% CI: 0.42–0.72;	the SPRINT trial.
				p=0.93	
				Stroke: 0.65; 95% CI: 0.27–1.57;	Limitations:
				p=0.38	 Lack of individual pt data,
				HF: 0.83; 95% CI: 0.41–1.70;	which would have allowed a
				p=0.27	more reliable assessment of
				Total deaths: 0.53; 95% CI: 0.37-	treatment effects in different pt
				0.76; p=0.79	groups.
				 More precision around estimates 	 Interpretation: Lowering of
				of benefits in SBP 130–139 at	BP into what has been
				baseline, fig 4 in paper	regarded the normotensive
				Results similar in trials of people	range should therefore be
				with and without CVD at baseline	routinely considered for the
				figure 5	prevention of CVD among
				CVD+ 0.77 (95% CI: 0.71–0.81)	those deemed to be of
				CVD- 0.74 (95% CI: 0.67–0.83)	sufficient absolute risk.
				Total deaths	
				CVD+ 0.90 (95% CI: 0.83–0.98)	
				CVD- 0.84 (95% CI: 0.75–0.93)	
				Other outcomes similarly in figure 5	
				• In appendix, in general, benefits	
				for CVD prevention seen in groups	
				with and without baseline CHD,	
				Stroke, DM, CKD and HF when	
				examined separately, but no	
				absolute risks provided to enable	
				estimation of how far down the	
				absolute risk curve these findings	
				have been demonstrated.	
L				חמיה מכרוו מכוווטווזנו מנכע.	

SPRINT Wright JT Jr, et al., 2015 (114) 26551272	Aim: To test the effectiveness of a goal SBP<120 mm Hg vs. a goal SBP<140 mm Hg for the prevention of CVD in pts with SBP≥130 mm Hg at baseline. Study type: RCT Size: 9361 pts followed median of 3.26 y.	Inclusion criteria: SBP≥130 mm Hg, with upper limit varying as number of pre-trial BP- lowering meds increased. age ≥50 y Presence of at least 1 of the following: • Clinical or subclinical CVD • CKD stage ≥3 • Age≥75 • Framingham General CVD risk≥15% in 10 y Exclusion criteria: DM, history of stroke, ESRD (eGFR <20)	Intervention: Intensive BP- lowering treatment to goal SBP <120 mm Hg Comparison: • Standard BP-lowering treatment to goal SBP<140 mm Hg • Net treatment difference ~3 drugs (2.8) on average vs. 2 drugs (1.8) on average • During the trial, mean SBP was 121.5 vs. 134.6.	 Some evidence of BB inferiority to other med classes in figure 6. Did not report absolute risks so do not know lower level of risk in treated populations. 1° <u>endpoint:</u> CVD (MI, ACS, stroke, HF, CVD death) HR: 0.75 (95% CI: 0.64, 0.89) <u>Other endpoints:</u> Total deaths HR: 0.73 (95% CI: 0.60-0.90) 1° or death HR: 0.78 (95% CI: 0.67-0.90) Components of 1° composite mostly consistent in direction other than ACS – no difference. <u>CKD outcomes:</u> 1° in CKD pts: reduction in GFR of ≥50% or ESRD HR: 0.89 (95% CI: 0.42, 1.87) Incident albuminuria HR: 0.72 (95% 0.48, 1.07) In pts without CKD: reduction in GFR ≥30% and to <60 HR: 3.49 (95% CI: 2.44-5.10) 	 Summary: More intensive SBP lowering to a goal of <120 mm Hg with achieved mean of approximately 121 mm Hg resulted in less CVD and lower total mortality over 3.26 y in comparison with a goal SBP <140 mm Hg and achieved SBP of ~135 mm Hg. There were small increases in some expected SAEs. Perhaps unexpected, a sizable increase in reduced eGFR in the non-CKD group and AKI/ARF overall was observed in the intensive group. While of uncertain etiology and significance, there is speculation this could be an acute hemodynamic effect, especially given the findings reagending alwaming.
				 Incident albuminuria HR: 0.72 (95% 0.48, 1.07) In pts without CKD: reduction in GFR ≥30% and to <60 	uncertain etiology and significance, there is speculation this could be an acute hemodynamic effect,
				 (95% CI: 0.63–1.04) <u>Adverse events:</u> SAEs: 1.04; p=0.25 Significant absolute increases seen in intensive group for hypotension (1%), syncope (0.6%), electrolyte abnormality (0.8%), acute kidney injury/acute renal failure (1.6%) over the study period. 	Limitations: Few pts were untreated at baseline ~9%, so SPRINT provides little if any insight at present regarding BP-lowering medication initiation for untreated people with SBP 130–139.

Lawes MR, et al., 2009 (115) <u>16222626</u>	Aim: • To determine the quantitative efficacy of different classes of BP-lowering drugs in preventing CHD and stroke, and who should receive treatment. • 5 questions encapsulate this uncertainty. 1st, do BBs have a special effect over and above lowering BP in preventing CHD events in people with a history of CHD? 2 nd , does the effect of BP- lowering drugs in preventing CHD and stroke differ in people with and without a history of CVD (i.e., is there a different effect in 2° and 1° preventing CHD and stroke? 4 th , should the use of BP-lowering	Inclusion criteria: The database search (by MRL) used Medline (1966 to December 2007; any language) to identify randomized trials of BP-lowering drugs in which CHD events or strokes were recorded (irrespective of whether BP reduction was considered the mechanism of action). Search terms were "antihypertensive agents" or "HTN" or "diuretics, thiazide" or "adrenergic beta- antagonists" or "angiotensin- converting enzyme inhibitors" or "receptors, angiotensin/antagonists & inhibitors" or "tetrazoles" or "CCB s" or "vasodilator agents" or the names of all BP-lowering drugs listed in the British National Formulary as keywords or text words. Limits were Medline publication type "clinical trial" or "controlled clinical trial" or "RCT" or "meta-analysis". We also searched the Cochrane Collaboration and Web of Science databases and the citations in trials and previous meta-analysis and review articles.	Intervention: BP-lowering medications Comparison: Placebo or less intensive treatment	 1.7% fewer pts had orthostatic hypotension in intensive group; p=0.01. <u>1° endpoint:</u> CHD and stroke co-1° Standardized to a 10/5 mm Hg BP reduction Overall CHD: 0.78 (95% CI: 0.73–0.83) Stroke: 0.59 (95% CI: 0.52–0.67) In absence of vascular disease CHD: 0.79 (95% CI: 0.72–0.86) Stroke: 0.54 (95% CI: 0.72–0.86) Stroke: 0.54 (95% CI: 0.45–0.65) History of CHD CHD: 0.76 (95% CI: 0.68–0.86) Stroke: 0.65 (95% CI: 0.63–0.80) History of stroke CHD: 0.79 (95% CI: 0.62–1.00) Stroke: 0.66 (95% CI: 0.56–0.79) No big drug class effects except more benefit for BBs shortly after MI. Treatment benefits seen down to pre-treatment SBP of 110–119 mm Hg for CHD events RR: 0.78 (95% CI: 0.63–0.96) and 130–139 mm Hg for stroke RR: 0.75 (95% CI: 0.63–0.89) 	Summary: The effect of BP- lowering drugs in reducing the risk of disease is entirely or largely due to BP reduction, with 1 main exception, a special extra effect of BBs in people who have had a recent MI The proportional reduction in CHD events and stroke for a given reduction in BP, an approximate halving in risk for each 10 mm Hg diastolic reduction, is the same in people with and without a history of vascular disease and in people without high BP as well as in those with high BP There is benefit in lowering BP in anyone at sufficient CV risk whatever their BP, so avoiding the need to measure BP routinely. Limitation: • Most of the pts without HTN were in the trials of people with pre-existing CVD; hence, most of the results of BP lowering in people with SBP<140 are in people with CVD. • No absolute risks or benefits provided. Not possible to estimate how far down the risk
	use of BP-lowering drugs be limited to people with high BP and not given to those at high risk of CVD	Exclusion criteria: We excluded nonrandomized trials and trials in which treated groups but not control groups			estimate how far down the risk curve these results apply. <u>Interpretation:</u> This MA provides stronger support for

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who have a lower BP?	had other interventions as well			treating at levels <140 for
A corollary is whether	as BP reduction, such as			people with CVD than for
BP should be reduced	cholesterol reduction. We			people without CVD.
to a limited extent	excluded trials in pts with			
only, a treat to target	chronic renal failure because			
approach. Although	these pts typically have high BP			
cohort	and high rates of CVD and their			
(prospective\observati	response to standard BP-			
onal) studies do not	lowering therapy may differ from			
show a lower BP limit	other people. We also excluded			
below which risk	trials in which fewer than 5 CHD			
ceases to decline ("the	events and strokes were			
lower the better"), this	recorded or the duration of			
has not been shown in	treatment was less than 6 mo,			
randomized trials	as these data would contribute			
across a wide range of	little to the overall results and			
BP.	substantially increase the			
Finally, what is the	complexity of the analyses.			
quantitative effect of	RCTs were otherwise included			
taking ≥1 BP-lowering	irrespective of pt age, disease			
drugs in lowering BP	status, BP before treatment, or			
and preventing CHD	use of other drugs.			
events and stroke				
according to dose,				
pretreatment BP, and				
age? To date no such				
quantitative summary				
of effect, taking				
account of these				
determining factors,				
has been made.				
Study type: Meta-				
analysis of RCTs				
,				
Size: 147 RCTs of BP-				
lowering meds and				
CHD events (22,000)				
and stroke (12,000).				
		1		

Lewington S,	Aim: To describe the	Inclusion criteria: Collaboration	Intervention: N/A	1° endpoint:	Summary: Throughout middle
et al., 2002	age-specific relevance	was sought from the		 Not completely clear, but for our 	and old age, usual BP is
(16)	of BP to cause-specific	investigators of all prospective	Comparator: N/A	purposes, stroke and IHD death	strongly and directly related to
12493255	mortality	observational studies in which		would be co-1°. Also looked at	vascular (and overall) mortality,
	5	data on BP, blood cholesterol,	• The exposures of interest	other vascular deaths.	without any evidence of a
	Study type: Meta-	date of birth (or age), and sex	were the level of SBP and	HRs for stroke mortality for a 20	threshold down to at least
	analysis of cohort	had been recorded at a baseline	DBP and age-group.	mm Hg lower SBP by age-group	115/75 mm Hg.
	studies	screening visit, and in which		40–49: 0.36 (95% CI: 0.32–0.40)	3
		cause and date of death (or age		50–59: 0.38 (95% CI: 0.35–0.40)	
	Size: 61 prospective	at death) had been routinely		60–69: 0.43 (95% CI: 0.41–0.45)	
	studies with 12.7	sought for all screens during		70–79: 0.50 (95% CI: 0.48–0.52)	
	million person-y of	more than 5,000 person-y of		80–89: 0.67 (95% CI: 0.63–0.71)	
	observation, 56,000	follow-up (see appendix A;		• HRs for IHD mortality for a 20	
	vascular deaths in 40-	http://image.thelancet.com/extra		mm Hg lower SBP by age-group	
	89 y.	s/01art8300webappendixA.pdf).		40–49: 0.49 (95% CI: 0.45–0.53)	
		Relevant studies were identified		50–59: 0.50 (95% CI: 0.49–0.52)	
		through computer searches of		60–69: 0.54 (95% CI: 0.53–0.55)	
		Medline and Embase, by hand-		70–79: 0.60 (95% CI: 0.58–0.61)	
		searches of meeting abstracts,		80–89: 0.67 (95% CI: 0.64–0.70)	
		and by extensive discussions		HRs for other vascular mortality	
		with investigators.		for a 20 mm Hg lower SBP by age-	
				group	
		Exclusion criteria: To minimize		40–49: 0.43 (95% CI: 0.38–0.48)	
		the effects of reverse causality		50–59: 0.50 (95% CI: 0.47–0.54)	
		(whereby established disease		60–69: 0.53 (95% CI: 0.51–0.56)	
		could change the usual BP),		70–79: 0.64 (95% CI: 0.61–0.67)	
		studies were excluded if they		80–89: 0.70 (95% CI: 0.65–0.75)	
		had selected pts on the basis of		 Similar results for DBP also in 	
		a positive history of stroke or		figure 1.	
		heart disease, and individuals		 Similar results for men and 	
		from contributing studies were		women separately for stroke, figure	
		excluded from the present		3, and IHD, figure 5.	
		analyses if they had such a			
		history recorded at baseline.			
Thomopoulos	Aim: Investigating	Inclusion criteria: Intentional	Intervention/Comparator:	<u>1° endpoint</u> :	Summary: Meta-analyses
C, et al.,	whether all grades of	BP-lowering comparing active	Criteria of eligibility were	 As some trials were done on low- 	favor BP-lowering treatment
2014 (20)	HTN benefit from BP-	drug treatment with placebo, or	intentional BP-lowering	risk pts, others on higher risk pts,	even in grade 1 HTN at low-to-
<u>25259547</u>	lowering treatment and	less active treatment (intentional	comparing active drug	no evaluation of absolute risk-	moderate risk, and lowering
	which are the target	BP-lowering trials), or	treatment with placebo, or	reduction was made. However, a	SBP/DBP to <140/90 mm Hg.
		comparison of an active drug	less active treatment	2° analysis was done including	Achieving <130/80 mm Hg

BP levels to maximize	with placebo over baseline	(intentional BP-lowering	trials or trial subgroups with mean	appears safe, but only adds
outcome reduction.	antihypertensive treatment,	trials), or comparison of an	baseline SBP/DBP values in grade	further reduction in stroke.
	resulting in a BP difference of at	active drug with placebo	1 range and a low-to-moderate risk	
Study type: Meta-	least 2 mm Hg in either SBP or	over baseline	(<5% CV deaths in 10 y in	
analysis of RCTs	DBP (nonintentional BP-lowering	antihypertensive treatment,	controls): FEVER stratum with	
	trials); enrolling of hypertensive	resulting in a BP difference	baseline SBP below the median	
Size: 32 RCTs with	individuals only or a high	of at least 2 mm Hg in	(<153 mm Hg) (e7); HTN Detection	
104,359 pts	proportion (at least 40%) of	either SBP or DBP	and Follow-up Program stratum	
	them.	(nonintentional BP-lowering	with baseline DBP 90–94 mm Hg	
		trials); enrolling of	and no CVD (e9); OSLO (e17);	
	Exclusion criteria: N/A	hypertensive individuals	TOMHS (e28) and USPHS (e29).	
		only or a high proportion (at	Risks of stroke, CHD, the	
		least 40%) of them. Other	composite of stroke and CHD, and	
		inclusion criteria can be	all-cause death were significantly	
		found in the preceding	reduced by BP-lowering in these	
		paper. 51 trials were found	low-to-moderate risk pts (control	
		eligible either for assessing	group: average CV mortality 4.5%	
		BP-lowering effects in	in10 y) with a moderate BP	
		different HTN grades or for	elevation (average SBP/DBP	
		assessing the effects of	145.5/91 mm Hg) at randomization.	
		achieving different BP	Standardized risk ratio associated	
		levels	with 10/5 reduction in BP: stroke	
			0.33 (95% CI: 0.11–0.98) CHD 0.68 (95% CI: 0.48–0.95)	
			CVD death 0.57 (95% CI: 0.32–	
			1.02) total death 0.53 (95% 0.35–	
			0.80)	
			Compared outcomes of achieved	
			• Compared outcomes of achieved on study SBP <130 vs. ≥130	
			Standardized Risk ratio associated	
			with 10/5 reduction in BP: stroke	
			0.68 (95% CI: 0.57, 0.83)	
			CHD 0.87 (95% CI: 0.76, 1.00)	
			HF 0.92 (95% CI: 0.47, 1.77)	
			CVD 0.81 (95% CI: 0.67, 1.00)	
			CVD death 0.88 (95% CI: 0.77,	
			1.01) total death 0.88 (95% CI:	
			0.77, 0.99)	
			 Outcomes of achieved on study 	
			SBP 130–139 vs. ≥140	

Lonn EM, et al., 2016 (116) <u>27041480</u>	<u>Aim:</u> To assess efficacy of fixed-dose antihypertensive therapy in adults with intermediate CVD risk. <u>Study type</u> : Double- blind, placebo- controlled RCT, factorial design <u>Size</u> : 12,705 pts	Inclusion criteria: Men ≥55 y and women ≥60 y at intermediate risk for CVD. No BP restrictions. Exclusion criteria: • Known CVD • Indications or contraindications to study meds • Mod/advanced CKD • Symptomatic hypotension	Intervention: FDC of ARB (candesartan 16 mg/d) and diuretic (hydrochlorothiazide 12.5 mg/d) or placebo Follow-up: Median=5.6 y	Standardized Risk ratio associated with 10/5 reduction in BP: stroke 0.63 (95% CI: 0.52–0.77) CHD 0.77 (95% CI: 0.70–0.86) HF 0.76 (95% CI: 0.47–1.25) CVD 0.74 (95% CI: 0.62–0.88) CVD death 0.81 (95% CI: 0.67– 0.97) total death 0.87 (95% CI: 0.75–1.00) • Similar pattern of results for on treatment DBP. <u>1° endpoint</u> : 1 co-1° CVD composite outcomes • CVD mortality, nonfatal MI, nonfatal stroke • Above plus cardiac arrest, HF, revascularization	Summary: • SBP/DBP reduction of 6.0/3.0 mm Hg • No difference in treatment effect • 1st co-1° 0.93 (0.79–1.10) • 2nd co-1° 0.95 (0.81–1.11) • Suggestion of a subgroup effect in tertile with the highest baseline BP and increased CVD risk.
Neaton JD et al., 1993 (117) <u>8336373</u>	Aim: To compare 6 antihypertensive drugs (representing different drug classes) Study type: Double- blind, placebo- controlled RCT Size: 902 pts with stage 1 HTN	 Inclusion criteria: Men and women 45–69 y Not taking antihypertensive medications, with DBP 90–99 mm Hg Taking 1 antihypertensive medication, with DBP <95 mm Hg and between 85–99 mm Hg after withdrawal of BP medications 	Intervention: Treatment (number): Once daily (AM): • Placebo (234) • Chlorthalidone 15 mg/d (136) • Acebutolol 400 mg/d (132) • Doxazosin 2 mg/d (134) • Amlodipine 5 mg/d (131) • Enalapril 5 mg/d (135) <u>Follow-up</u> : Median=4.4 y	<u>1º endpoint</u> : BP, QoL, side effects, chemistries, ECG, clinical events	Summary: • Drugs (plus diet) more effective compared to placebo (plus diet) for control of BP. • Minimal differences between drug regimens

Van Dieren S, et al., 2012 (118) <u>22677192</u>	Aim: To assess differences in treatment effects of a fixed combination of perindopril- indapamide on major clinical outcomes in pts with type 2 DM across subgroups of CV risk. Study type: RCT Size: 11,140 pts with DM-2, from the ADVANCE trial	Inclusion criteria: DM-2, aged ≥55 y, with a history of major macrovascular or microvascular disease, or at least 1 other risk factor for vascular disease Exclusion criteria: A definite indication for, or contraindication to, any of the study treatments, a definite indication for long-term insulin treatment or were participating in any other clinical trial.	Intervention: Perindopril- indapamide or matching placebo	 <u>1° endpoint</u>: The Framingham equation was used to calculate 5-y CVD risk and to divide participants into 2 risk groups, moderate-to-high risk (<25% and no history of macrovascular disease), very high risk (>25% and/or history of macrovascular disease). Endpoints were macrovascular and microvascular events. 	<u>Summary</u> : Relative effects of BP-lowering with perindopril- indapamide on CV outcomes were similar across risk groups whilst absolute effects trended to be greater in the high-risk group.
Montgomery AA, et al., 2003 (119) <u>12923409</u>	Aim: To estimate the effectiveness and cost-effectiveness of BP-lowering treatment over a lifetime. Study type: Markov decision analysis model comparing treatment and nontreatment of HTN. Size: Hypothetical cohorts for 20 different strata of sex, age (30– 79 y, in 10-y bands), and CV risk (low and high)	Inclusion criteria: We created models for 20 different strata of sex, age (age 30–70 y in 10-y bands), and 2 risk profiles (designated as 'low' and 'high' risk). These example risk profiles represent the extremes of absolute CV risk, based on data from the Health Survey for England and using a Framingham risk function. We recognize that the risk of most individuals seen in primary care will be somewhere between the examples presented here. The data included were as follows: age- and sex-specific mean SBP of untreated individuals with SBP>0.160 mm Hg were used for both high-risk and low-risk profiles. In addition, low-risk profile was defined as nonsmoker, 10th percentile total cholesterol 90th percentile HDL	Intervention: Treatment and nontreatment of HTN.	<u>1º endpoint</u> : Life expectancy, and incremental cost: effectiveness ratios for treatment and nontreatment strategies	 Probabilities of clinical events were obtained from published literature. <u>Summary</u>: Incremental cost per quality-adjusted life y among low-risk groups ranged from £1,030 to £3,304. Cost-effectiveness results for low-risk pts were sensitive to the utility of receiving antihypertensive treatment. Treatment of highrisk individuals was highly cost-effective, such that it was the dominant strategy in the oldest age group, and resulted in incremental costs per quality-adjusted life y ranging from £34–£265 in younger age groups. Policy decisions about which pts to treat depend on whether a life-expectancy or cost-

		cholesterol, no DM, and no LVH, and high-risk profile was defined as smoker, 90th percentile total cholesterol, 10th percentile HDL cholesterol, DM, and LVH. <u>Exclusion criteria</u> : N/A			effectiveness perspective is taken. Treatment increases life expectancy in all strata of age, sex, and CV risk. However, younger individuals stand to gain proportionately more from BP treatment than do the elderly. In terms of cost- effectiveness, pts at high risk of CVD are a highly cost- effective group to treat. In pts at lower risk of CVD, consideration should be given to issues of pt preference and cost.
Kassai B, et al., 2005 (120) <u>17315403</u>	Aim: Consideration of absolute risk has been recommended for making decisions concerning preventive treatment in HTN. Aim to estimate the benefit of antihypertensive therapy over a life- time. Study type: Meta- analysis on individual data in HTN and specific cause of death from national statistics. Disease-free survival curves until all pts have died were built using the "life-table" method. The treatment effect estimated from INDANA was applied to this curve to obtain the disease-free	Inclusion criteria: To estimate the rate of cv and non-CV deaths in a hypothetical U.S. population of untreated hypertensive pts, we used the following procedure: age-specific death rates in the U.S. general population were obtained from national vital statistics (1994), and in untreated hypertensive population they were obtained from the control groups of the INDANA database. This latter group represents a unique cohort of 14 942 untreated or placebo-treated hypertensive pts, 26–96 y with an average follow-up of 5 y <u>Exclusion criteria</u> : N/A	Intervention: The gain in life expectancy without stroke, CHD, and CV events was estimated from the area between the 2 survival curves of treated and control groups. The relative gain in life expectancy was defined as the ratio of gain in life expectancy to life expectancy.	1° endpoint: Stroke and CHD co-1° Results: CHD Age ABb RGLEe Y RRa (%) NNTc GLEd (%) 40 0.86 0.3 333 20 4.1 50 0.88 1.0 100 17 4.3 60 0.90 1.9 53 13 3.4 70 0.91 3.9 26 10 5.4 Stroke Age ABb RGLEe Y RRa (%) NNTc GLEd (%) 40 0.80 0.4 250 32 5.9 50 0.84 1.0 100 26 5.7 60 0.86 2.3 44 21 7.1 70 0.87 5.7 18 17 9.1 a RR at 10 y y b Absolute benefit at 10 y y c NNT to avoid 1 event. d Gain in life expectancy in mo without events.	Summary: Absolute gains in life expectancy are likely to be greater for younger, lower risk people with HTN than for older, higher risk people with HTN. However, the NNT to prevent an event will likely be greater especially in the short term in younger, lower risk people. This modeling analysis provides support for treating younger, lower risk individuals with HTN, but relies on the assumption that the relative benefits of treatments observed in short-term trials of higher risk individuals applies over a longer term to lower risk individuals.

Czernichow S et al., 2011 (121) 20881867	survival curve of the life-long treated population. Gains in event-free life expectancy were estimated from survival curves. A sensitivity analysis was performed to assess the impact of possible death misclassifications. <u>Size</u> : 6 RCTs, ~30,000 pts <u>Aim:</u> The objective of this systematic review and meta-analysis was to compare the relative reductions in risk achieved with different starting levels of BP (and treatment regimens). <u>Study type:</u> Meta- analysis of RCTs <u>Size:</u> 32 trials with 201,566 pts (20,079 1° outcome events)	Inclusion criteria: RCTs of BP- lowering (drug vs. control or less intensive treatment) or different classes of drug therapy that included a minimum of 1,000 pt- y of follow-up in each study arm. Exclusion criteria: <1,000 pt-y of follow-up in each treatment group.	Intervention: BP-lowering meds Comparator: Placebo, active comparator or less intensive treatment	 e Relative gain in life expectancy without events. <u>1° endpoint:</u> Major CVD events (stroke, CHD, and HF. No evidence of differences in the ratio of risk across varying levels of baseline BP (with all classes of BP-lowering medications). 	Summary: • Effectiveness of BP-lowering regiments in reducing RR of major CVD events does not seem to be influenced by starting level of BP. Limitations: • The majority of the participants studied were at high risk for CVD. • Information pertaining to the effect of treatment on absolute risk was not presented in this manuscript.
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Data Supplement 24. Follow-Up After Initial BP Evaluation (Section 8.1.3)

Study Acronym;	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if any);
Author;	Study Type;		(# patients) /	(Absolute Event Rates,	Study Limitations;
Year Published	Study Size (N)		Study Comparator	P value; OR or RR; & 95%	Adverse Events;
			(# patients)	CI)	Summary

Ambrosius WT, et al., 2014 (122) <u>24902920</u>	Aim: To describe the study design of the SPRINT Study type: SPRINT RCT	Inclusion criteria: Adults ≥50 y, average SBP_≥130 mm Hg and evidence of CVD, CKD, or 10-y Framingham risk score ≥15%, or ≥75 y	Intervention: 9,361 pts randomized to 2 treatment groups: • Standard treatment group, SBP target <140 mm Hg • Intensive treatment group: SBP target <120 mm Hg.	<u>1° endpoint</u> : MI, ACS, stroke, HF, or CVD death.	Relevant 2° endpoint: All-cause mortality, decline in kidney function or development of ESRD, incident dementia, decline in cognitive function, and small-vessel cerebral ischemic diseaseSummary: This paper describes the protocol followed in the SPRINT trial that was successful in helping participants to attain and maintain BP targets in the study groups. Once treated, participants had follow-up visits to assessment BP control monthly until BP was at target. Medications were titrated and added as per protocol, when target BP was not attained.
Cushman WC, et al., 2007 (123) <u>17599425</u>	Aim: To describe the study design of the BP trial of the ACCORD Trial Study type: Description of study design and protocol for the ACCORD RCT	Inclusion criteria: Adults with a diagnosis of DM-2 for at least 3 mo and at high risk for CVD events, who meet the following BP criteria: (1) SBP 130–160 mm Hg and taking 0–3 antihypertensive medications; (2) SBP 161–170 and on 0–2 antihypertensive medications; or (3) SBP 171-180 and taking 0-1 antihypertensive medication. Other entry criteria included spot urine sample <2+, protein–Cr ratio <700 mg protein/1 g Cr, or 24-h protein excretion <1.0 g/24 h.	Intervention: • Unmasked, open- label, factorial design, randomized trial with a sample size of 4,733 pts • Pts randomized to intensive SBP control (<120 mm Hg) or standard control (<140 mm Hg)	<u>1° endpoint</u> : Major CVD event (nonfatal MI or stroke, or CV death)	Relevant 2° endpoint: Expanded macrovascular outcome (1° outcome plus coronary revascularization or HF hospitalization), total mortality, each of the separate components of the 1° outcome, HF death or hospitalization, and composite microvascular disease outcome (kidney and eye disease). Summary: This paper describes the protocol followed in the ACCORD trial that was successful in helping participants to attain and maintain BP targets in the study groups. Once treated, participants had follow-up visits to assessment BP control monthly until BP was at target. Medications were titrated and added as per protocol, when target BP was not attained.

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% Cl)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
VA NEPHRON-D Fried LF, et al., 2013 (124) 24206457	Aim: Assess the efficacy of combination of an ACEI and an ARB vs. ARB monotherapy in reducing the progression of proteinuric diabetic nephropathy Study type: Multicenter, double- blind, RCT at 32 VA Medical Centers Size: 1448 pts	Inclusion criteria: Pts with type 2 DM, a urinary albumin-to-creatinine ratio of ≥300, and an eGFR 30.0–89.9 mL/min/1.73 m ² Exclusion criteria: • Subjects with known nondiabetic kidney disease • Serum K+ >5.5 mmol/L • Current treatment with sodium polystyrene sulfonate • Inability to stop prescribed medication that increases the risk of hyperkalemia	Intervention: Losartan 100 mg daily plus lisinopril 10–40 mg daily (n=724) <u>Comparator</u> : Losartan 100 mg daily plus placebo (n=724)	<u>1° endpoint</u> : After a median follow-up of 2.2 y, the study was stopped early due to safety concerns. There was no difference in the 1° outcome of first occurrence of change in eGFR (decrease of ≥30 mL/min/1.73 m ² if initial GFR was ≥60 mL/min/1.73 m ² or a decline of ≥50% if initial eGFR was <60 mL/min/1.73 m ²), ESRD, or death (HR with combination therapy: 0.88; 95% CI: 0.70–1.12; p=0.30). <u>Safety endpoint</u> : Combination therapy increased the risk of hyperkalemia (HR: 2.8; 95% CI: 1.8–4.2; p<0.001) and acute kidney injury (HR: 1.7; 95% CI: 1.3–2.2; p<0.001).	2° endpoint: There was no difference in the 2° endpoint of first occurrence of change in eGFR or ESRD (HR: 0.78; 95% CI: 0.58–1.05; p=0.10). There were no differences between combination therapy or losartan monotherapy for the endpoints of ESRD, death, composite of MI, HF, or stroke, MI, CHF, and stroke (p>0.05 for all). <u>Summary</u> : Combination therapy of losartan plus lisinopril did not improve renal outcomes compared to losartan alone, and was associated with greater risk of acute kidney injury and hyperkalemia.
ALTITUDE Parving HH, et al., 2012 (125) <u>23121378</u>	<u>Aim</u> : Determine if addition of aliskiren as an adjunct to an ACEI or ARB reduces the risk of CV and renal events in pts with type 2 DM	Inclusion criteria: • ≥35 y with type 2 DM • On ACEI or ARB • At least 1 of the following: persistent macroalbuminuria (urine microalbumin to creatinine ratio ≥200 mg/g) and eGFR ≥30 mL/min/1.73 m ² , persistent microalbuminuria (≥20 mg/g and <200 mg/g) and a mean eGFR ≥30 and <60	Intervention: Aliskiren 300 mg daily added to conventional treatment with an ACEI or ARB (n=4,274) Comparator: Placebo (n=4,287)	<u>1° endpoint</u> : After a median follow-up of 32.9 mo the study was stopped early. There was no difference in the 1° composite outcome death from CV causes or first occurrence of cardiac arrest with resuscitation; nonfatal MI; nonfatal stroke;	 <u>2° endpoint</u>: There was no difference between aliskiren and placebo for the individual components of the composite 1° outcome (all p>0.05) other than cardiac arrest with resuscitation, which was increased significantly with aliskiren (HR: 2.40; 95% CI: 1.05–5.48; p=0.04).

Data Supplement 25. RCTs for General Principles of Drug Therapy (Combination Therapies that Inhibit the RAAS) (Section 8.1.4)

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	Study type: Doubled-blind, multicenter RCT Size: 8561	mL/min/1.73 m ² , or history of CVD (e.g., MI, stroke, HF, or CAD) and a mean eGFR \geq 30 and <60 mL/min/1.73 m ² Exclusion criteria: • Serum K+ >5.0 mmol/L • Type 1 DM • Unstable serum Cr • CV history (NYHA Class III or IV, SBP \geq 170 mm Hg or DBP \geq 110 mm Hg or SBP \geq 135 and <170 mm Hg or DBP \geq 82 and <100 mm Hg with at least 3 agents, 2 nd or third degree heart block, renal artery stenosis • Surgical or medical conditions (malignancy in last 5 y, <2 y life expectancy, renal transplant or immunosuppressive therapy, drug/alcohol abuse, hypersensitivity/allergy/contraindication to study drugs, pregnancy) • Concomitant treatment with \geq 2 agents blocking RAAS or K+-sparing diuretics.		unplanned hospitalization for HF; ESRD; death attributable to kidney failure or need for renal- replacement therapy with no dialysis or transplantation available or initiated; or doubling of the baseline serum Cr between aliskiren or placebo (HR: 1.08; 95% CI: 0.98–1.20; p=0.12). <u>Safety endpoint</u> : The combination of aliskiren added to an ACEI or an ARB was associated with greater risk of hyperkalemia and hypotension (11.2% vs. 7.2% and 12.8% vs. 8.3%; p<0.001 for both, respectively).	 There was no differences in CV composite outcome, renal composite outcome, or death from any cause (p>0.05 for all) <u>Summary</u>: Aliskiren added to background treatment of an ACEI or ARB did not decrease CV or renal outcomes, and was associated with increased risk of cardiac arrest with resuscitation, hyperkalemia, and hypotension.
ONTARGET Yusuf S, et al., 2008 (126) <u>18378520</u>	Aim: Evaluate whether use of an ARB was noninferior to ACEI, and whether the combination was superior to ACE alone in the prevention of vascular events in pts with CVD or DM but not HF. Study type: Multi- center, double-blind, RCT	Inclusion criteria: • ≥55 y • Coronary, peripheral, or cerebrovascular disease or DM with end-organ damage Exclusion criteria: • Inability to discontinue ACEI or ARB • Known hypersensitivity or intolerance to ACEI or ARB • Selected CVDs (congestive HF, hemodynamically significant valvular or outflow tract obstruction, constrictive pericarditis, complex congenital heart disease, syncopal episodes of unknown etiology <3 mo, planned cardiac surgery	Intervention: Ramipril 10 mg daily (n=8,576) Comparator: • Telmisartan 80 mg daily (n=8,542) • Combination of telmisartan and ramipril (n=8,502)	1° endpoint: After a median follow-up of 56 mo, there was no difference between ramipril vs. telmisartan or combination therapy vs. ramipril in the 1° composite outcome of death from CV causes, MI, stroke, or hospitalization for HF (RR: 1.01; 95% CI: 0.94– 1.09 and RR: 0.99; 95% CI: 0.92–1.07, respectively) Safety endpoint: • Combination therapy was associated with greater risk of hyperkalemia than	 <u>2° endpoint</u>: There was no difference in composite of death from CV causes, MI, or stroke in the ramipril vs. telmisartan groups RR: 0.99; 95% CI: 0.9–1.07); p=0.001 or ramipril vs. combination RR: 1.00; 95% CI: 0.93–1.09 There were no differences between ramipril vs. telmisartan or ramipril vs. combination therapy in 2° outcomes including MI, stroke, hospitalization for HF, death from CV causes, or death from any cause (p>0.05 for all).

<u>Size</u> : 25,620	or PTCA <3 mo, uncontrolled HTN on treatment [e.g., BP >160/100 mm Hg], heart transplant recipient, stroke due to subarachnoid hemorrhage) • Other conditions (significant renal artery disease, hepatic dysfunction, uncorrected volume or sodium depletion, 1° hyperaldosteronism, hereditary fructose intolerance, other major noncardiac illness or expected to reduce life expectancy or significant disability interfere with study participation, simultaneously taking another experimental drug, unable to	ramipril monotherapy (480 pts vs. 283 pts; p<0.001) • Hypotensive symptoms were cited as reason for permanent discontinuing more in telmisartan vs. ramipril (RR: 1.54; p<0.001) and combination therapy vs. ramipril monotherapy (RR: 2.75; p<0.001) • Renal impairment was more common in combination therapy vs. ramipril monotherapy vs.	<u>Summary</u> : Combination therapy with telmisartan and ramipril did not decrease the risk of CV events in pts at high risk compared to monotherapy with ramipril. In addition, combination therapy was associated with increased risk of hypotension, hyperkalemia, and renal impairment.
	another experimental drug, unable to provide written informed consent).	ramipril monotherapy RR: 1.33; 95% CI: 1.2–1.44	

Data Supplement 26. BP Goal for Patients with Hypertension (Section 8.1.5)

Study Acronym (if applicable) Author Year Published	Study Type/Design; Study Size (N)	Patient Population	Study Intervention (# patients) Study Comparator (# patients)	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Lawes CM, et al., 2003 (50) <u>12658016</u>	Study type: Meta- analysis of RCTs of BP drugs recording CHD events and strokes Size: 464,000 pts	N/A	N/A	• CHD RR or 46% Stroke 64%	• All classes of BP meds confer benefit while BB confer greater benefit in those with CAD
LV J, et al., 2013 (127) <u>23798459</u>	Study type: MA of RTC that randomly assigned individuals to different target BP levels Size: 15 trials including a total of 37,348 pts	N/A	N/A	 7.5/4.5 mm Hg BP difference. Intensive BP lowering achieved. <u>RR for</u> Major CV events: 11%; 95% CI: 1%–21%) MI: 13%; 95% CI: 0%– 25% 	More intensive strategy for BP control reduced cardio- renal endpoint

				 Stroke: 24%; 95% CI: 8%-37% ESRD: 11%; 95% CI: 3%-18% Albuminuria: 10%; 95% CI: 4%-16% Retinopathy 19%; 95% CI: 0%-34% p=0.051 	
Xie X, et al., 2015 (21) <u>26559744</u>	Study type: MA of RTC that randomly assigned individuals to different target BP levels <u>Size</u> : 19 trials (n=44,989)	N/A	N/A	Achieved BP 133/76 mm Hg (intensive) 140/81 (less intense) • Major CV events: 14%; 95% CI: 4%–22% • MI: 13%; 95% CI: 0%– 24% • Stroke: 22%; 95% CI: 10%–32% • Albuminuria: 10%; 95% CI: 3%–16% • Retinopathy progression: 19%; 95% CI: 0%–34%. • More intensive had no effects on HF: 15%; 95% CI: -11%–34% • CV death: 9%; 95% CI: - 11%–26% • Total mortality: 9%; 95% CI: -3%–19% • ESKD: 10%; 95% CI: - 6%–23%	More intensive approach reduced major CV events (stroke and MI) except heat failure, CVD, ESRD, and total mortality.

Verdecchia P et al., 2016 <u>27456518</u>	Study type: Cumulative meta- analysis of RCTs to study benefit of more vs. less intensive BP lowering <u>Size</u> : 18 trials (n=53,405)	N/A	N/A	 Stroke, MI, HF, CVD mortality, and all-cause mortality Difference in achieved SBP/DBP=7.6/4.5 mm Hg For stroke and MI the cumulative Z score crossed the efficacy boundary after addition of the SPRINT results For CVD mortality and HF, the cumulative Z curve crossed the conventional significance boundary (but not the sequential monitoring boundary) For all-cause mortality, the cumulative Z curve did not reside in the futility are but did not cross the conventional significance boundary 	• The results strongly supported the benefit of intensive BP reduction for prevention of stroke and MI and suggested benefit for prevention of CVD mortality and HF
Bangalore S, et al., 2017 <u>28109971</u>	Study type: Network meta- analysis in which the authors attempted to compare the benefits and adverse effects resulting from intensive reduction in SBP Size: 17 trials (n=55,163)	N/A	N/A	 There was a significant reduction in stroke (RR: 0.54) and MI (RR: 0.68) The point estimate favored all-cause mortality, CVD mortality and HF but the results did not achieve significance SBP targets <120 and <130 mm Hg ranked #1 and #2 as the most efficacious Serious adverse effects were more common at a lower SBP (120 vs. 150 or 140 mm Hg) 	• Overall, the beneficial effects of treatment were consistent with other reports. The cluster plots of treatment benefit vs. risk are difficult to interpret due to limitations of the available data base and the authors' decision to weight treatment benefits and potential adverse effects equally.

				• Cluster plots for combined efficacy and safety suggested a SBP <130 mm Hg as the optimal target for SBP reduction during treatment	
Bundy JD, et al., 2017 <u>28564682</u>	Study type: Systematic review and network meta- analysis to assess the benefits of intensive SBP reduction during treatment of hypertension Size: 42 trials (n=144,220)	N/A	N/A	• In general, there were linear associations between achieved SPB and risk of CVD and all- cause mortality, with the lowest risk at a SBP of 120–124 mm Hg.	• This was by far the largest and best powered meta- analysis to assess the relationship between SBP reduction and major outcomes during treatment of hypertension. The findings provided strong evidence for the "lower is better" approach to treatment in patients with a high SBP who are at high risk for CVD.
Lawes CMM, et al., 2002 <u>16222626</u>	Study type: Review of observational reports and randomized controlled trials	N/A	N/A	 The relative benefits of BP lowering for CHD prevention likely to be consistent across a wide range of different populations Likely to be considerable benefit for BP lowering beyond traditional thresholds, especially in those at high risk for CVD BP lowering is likely to be more important than choice of initial agent A large majority of patients being treated for 	• Strongly supports lower BPs during BP treatment, especially in those at high risk of CVD

	hypertension have suboptimal BPs. Initiatives to lower their BP further are essential	

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (include Absolute Event Rates, P value; OR or RR; and 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Xie X, et al., 2015 (21) <u>26559744</u>	Aim: To assess the efficacy and safety of intensive BP- lowering strategies. Study type: Meta- analysis of RCTs Size: 19 RCTs with 44,989 pts	Inclusion criteria: RCTs with at least 6 mo follow-up that randomly assigned pts to more intensive vs. less intensive BP-lowering treatment, with different BP targets or different BP changes from baseline. Reference lists from identified trials and review articles were manually scanned to identify any other relevant studies. <u>Exclusion criteria:</u> N/A	Intervention: BP-lowering meds Comparator: • Less intensive treatment • BP difference 6.8/3.5 • The mean follow-up BP levels in the less intensive BP- lowering regimen group were 140/81 mm Hg, compared with 133/76 mm Hg in the more intensive treatment group.	 <u>1° endpoint</u>: CVD, other major CV events, defined as a MI, stroke, HF, or CV death, separately and combined; nonvascular and all-cause mortality; ESKD, and adverse events. Progression of albuminuria (defined as new onset of micro-albuminuria/macro-albuminuria or a change from micro-albuminuria to macro-albuminuria) and retinopathy (retinopathy progression of 2 or more steps) were also recorded for trials that were done in pts with DM CVD RR: 0.86 (95% CI: 0.78–0.96) 	Summary: Intensive BP- lowering, including to <130 mm Hg, provided greater vascular protection than standard regimens. In high- risk pts, there are additional benefits from more intensive BP-lowering, including for those with SPB <140 mm Hg at baseline. The net absolute benefits of intensive BP- lowering in high-risk individuals are large. Limitations: • Lack of individual pt data, which would have allowed a more reliable assessment of

	Other endpoints: • MI RR: 0.87 (95% CI: 0.76– 1.00) p=0.042 • Stroke RR: 0.78 (95% CI: 0.68–0.90) • HF RR: 0.85 (95% CI: 0.66– 1.11) • CVD death RR: 0.91 (95% CI: 0.74–1.11) • Total deaths RR: 0.91 (95% CI: 0.81–1.03)	treatment effects in different pt groups. • Interpretation: Supports treating pt with and without CVD at threshold of 130 to <130. Supports treating at threshold of about 130 even down to a CVD event rate of 0.9% per y.
	Other results:• Benefit for CVD not differentby baseline SBP120–139: 0.89 (95% CI: 0.76–1.05)140–160: 0.83 (95% CI: 0.68–1.00)>160: 0.89 (95% CI: 0.73–1.09)p-heterogeneity: 0.60• Benefit for CVD not differentfor more intensive and lessintensive targets in intensivegroup<140 or <150 mm Hg: 0.76	
	control group rate of major vascular events was 2.9% per y compared with 0.9% per y in other trials, and the numbers needed to treat were 94 (95%	

				 CI: 44–782) in these trials vs. 186 (95% CI: 107–708) in all other trials. Increase in severe hypotension: 0.3% vs. 0.1% per person y OR: 2.68 (95% CI: 1.21–5.89) 	
Julius S, et al., 2006 (55) <u>16537662</u>	<u>Study type</u> : RCT in pre-HTN16 mg candesartan vs. placebo <u>Size</u> : 809 pts	• 58% men	N/A	• During the first 2 y, HTN developed in 154 (40.4%) pts in the placebo group compared with only 53 (13.6%) of those in the candesartan group, for a RR of 66.3% (p<0.0001). After 4 y, HTN developed in 240 (63.0%) in the placebo group vs. only 208 (53.2%) in the candesartan group RR 15.6% (p<0.0069).	• 2/3 of those with pre-HTN develop HTN within 4 y. Candesartan interrupts the onset and reduced by 15.6%
Lawes CM, et al., 2003 (50) <u>12658016</u>	Study type: Meta- analysis of RCTs of BP drugs recording CHD events and strokes Size: 464,000 pts	N/A	N/A	CHD RR or 46% Stroke 64%	All classes of BP meds confer benefit while BB confer greater benefit in those with CAD
Lonn EM, et al., 2016 (116) <u>27041480</u>	Aim: To assess efficacy of fixed-dose antihypertensive therapy in adults with intermediate CVD risk. <u>Study type</u> : Double- blind, placebo- controlled RCT, factorial design	Inclusion criteria: Men ≥55 y and women ≥60 y at intermediate risk for CVD. No BP restrictions. Exclusion criteria: • Known CVD • Indications or contraindications to study meds • Mod/advanced CKD • Symptomatic hypotension	Intervention: FDC of ARB (candesartan 16 mg/d) and diuretic (hydrochlorothiazide 12.5 mg/d) or placebo Follow-up: Median=5.6 y	 <u>1° endpoint</u>: 1 co-1° CVD composite outcomes CVD mortality, nonfatal MI, nonfatal stroke Above plus cardiac arrest, HF, revascularization 	Summary: • SBP/DBP reduction of 6.0/3.0 mm Hg • No difference in treatment effect • 1st co-1° 0.93 (0.79–1.10) • 2nd co-1° 0.95 (0.81–1.11)

	<u>Size</u> : 12,705 pts				• Suggestion of a subgroup effect in tertile with the highest baseline BP and increased CVD risk.
Neaton JD, et al., 1993 (117) <u>8336373</u>	Aim: To compare 6 antihypertensive drugs (representing different drug classes) Study type: Double- blind, placebo- controlled RCT Size: 902 pts with stage 1 HTN	Inclusion criteria: • Men and women 45–69 y • Not taking antihypertensive medications, with DBP 90–99 mm Hg • Taking 1 antihypertensive medication, with DBP <95 mm Hg and between 85–99 mm Hg after withdrawal of BP medications	Intervention: Treatment (number): Once daily (AM): • Placebo (234) • Chlorthalidone 15 mg/d (136) • Acebutolol 400 mg/d (132) • Doxazosin 2 mg/d (134) • Amlodipine 5 mg/d (131) • Enalapril 5 mg/d (135) <u>Follow-up</u> : Median=4.4 y	<u>1º endpoint</u> : BP, QoL, side effects, chemistries, ECG, clinical events	Summary: • Drugs (plus diet) more effective compared to placebo (plus diet) for control of BP. • Minimal differences between drug regimens

Data Supplement 27. Choice of Initial Medication (Section 8.1.6)

Study Acronym Author Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Psaty BM, et al., 2003 <u>12759325</u>	Study type: Network meta- analysis to compare value of different first-line antihypertensive drugs in prevention of major CVD and all-cause mortality Size: 42 trials (n=192,478)	N/A	 For all outcomes, low-dose diuretics were better than placebo None of the other first-line agents (β-blockers, ACEI, CCBs, α-receptor blockers and ARBs) were superior to low-dose diuretics For several outcomes, low-dose diuretics were superior to other agents 	• Low-dose diuretics were identified as the most effective first-line treatment for prevention of CVD and all-cause mortality during treatment of hypertension	N/A

Brunström M, et al., 2016 (53) 26920333	Study type: Meta- analysis of levels of BP control in DM hypertensives. Size: 73,738 pts	• 49 trials (most pts with DM-2)	Baseline SBP >150 <u>RR for</u> • All death: 0.89; 95% CI:0.80– 0.99 • CVD: 0.75; 95% CI: 0.57– 0.99 • MI: 0.74; 95% CI: 0.63–0.87 • Stroke: 0.77; 95% CI: 0.65– 0.91 • ESRD: 0.82; 95% CI: 0.71– 0.94 Baseline SBP140–150 <u>RR of</u> • Death: 0.87; 95% CI: 0.78– 0.98) • MI: 0.84; 95% CI: 0.76–0.9 • HF: 0.80; 95% CI: 0.76–0.9 The seline SBP,140 mm Hg, 100– 1.32	• BP lowering reduces major CV events in DM. Caution for initiating treatment in diabetics with SBP <140/90	N/A
Ettehad D, et al., 2015 (17) <u>26724178</u>	Study type: Meta- analysis of large RTCs of antihypertensive treatment Size: 123 studies (613,815 pts)	N/A	Every 10 mm Hg reduction in SBP RR: • Major CV events: 0.80; 95% CI: 0.77–0.83 • CHD: 0.83; 95% CI: 0.78– 0.88 • Stroke: 0.73; 95% CI: 0.68– 0.77), HF (0.72, 0.67–0.78 • All-cause mortality: 0.87; 95% CI: 0.87; 0.84–0.91 • ESRD: 0.95; 0.84–1.07	• BP lowering reduces CV risk across various baseline BP levels and comorbidities. Suggest lowering SBP <130 mm Hg and BP-lowering treatment to pts with a history of CVD, CHD, stroke, DM, HF, and CKD.	N/A
Thomopolous C, et al., 2016 (54) <u>26848994</u>	Study type: Meta- analysis of RTCs of more vs. less intense BP control	• 16 trials (52,235 pts) compared more vs. less intense treatment 34 (138,127 pts) active vs. placebo	More intense BP • Stroke RR: 0.71; 95% CI: 0.60–0.84) • CHD RR: 0.80; 95% CI: 0.68–0.95)	 Intensive BP reduction improves CV outcomes compared to less intense Achieved BP <130/80 may be associated with CV benefit. 	N/A

			 Major CV events RR: 0.75; 95% CI: 0.68–0.85 CV mortality RR: 0.79; 95% CI: 0.63–0.97 Stratification of SBP cutoffs (150,140 and 130 mm Hg) showed that a SBP/DBP difference of 10/5 mm Hg across each cutoff reduced risk of all outcomes 		
Julius S, et al., 2006 (55) <u>16537662</u>	Study type: RCT in pre-HTN 16 mg candesartan vs. placebo Size: 809 pts	• 58% men	• During the first 2 y, HTN developed in 154 (40.4%) pts in the placebo group compared with only 53 (13.6%) of those in the candesartan group, for a RR of 66.3% (p<0.0001). After 4 y, HTN developed in 240 (63.0%) in the placebo group vs. only 208 (53.2%) in the candesartan group RR 15.6% (p<0.0069).	• 2/3 of those with pre-HTN develop HTN within 4 y. Candesartan interrupts the onset and reduced by 15.6%	N/A
Ference BA, et al., 2014 (56) <u>24591335</u>	Study type: Evaluated the effect of 12 polymorphisms (associated with BP) on the odds of CHD and compared it with the effect of lower SBP observed in both prospective cohort studies and BP-lowering randomized trials Size: 199,477 pts in 63 studies	N/A	•12 polymorphisms were associated with a 0.32 mm Hg lower SBP (p=1.79×10 ⁻⁷) and a 0.093-mm Hg/decade slower age-related rise in SBP (p=3.05×10 ⁻⁵). The effect of long-term exposure to lower SBP on CHD mediated by these polymorphisms was 2- fold greater than that observed in prospective cohort studies (p=0.006) and 3-fold greater than that observed in short- term BP treatment trials (p=0.001).	• SBP may be causally associated with the rate of rise in SBP with age and has a cumulative effect on the risk of CHD.	N/A

Data Supplement 28.	Follow-Up After Initiatin	a Antihypertensive Drug	Therapy (Section 8.3.1)
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Study Acronym Author Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Ambrosius WT, et al., 2014 (122) <u>24902920</u>	<u>Aim</u> : To describe the study design of the SPRINT trial <u>Study type</u> : description of study design and protocol for the SPRINT RCT	Inclusion criteria: Adults ≥50 y, average SBP ≥130 mm Hg and evidence of CVD, CKD, or 10-y Framingham risk score ≥15%, or age ≥75 y	Intervention: 9361 participants randomized to 2 treatment groups: (1) Standard treatment group, SBP target <140 mm Hg, and (2) Intensive treatment group: SBP target <120 mm Hg.	<u>1° endpoint</u> : MI, ACS, stroke, HF, or CVD death.	Relevant 2° endpoint: All- cause mortality, decline in kidney function or development of ESRD, incident dementia, decline in cognitive function, and small- vessel cerebral ischemic disease
					Summary: This paper describes the protocol followed in the SPRINT trial that was successful in helping participants to attain and maintain BP targets in the study groups. Once treated, participants had follow-up visits to assessment BP control monthly until BP was at target. Medications were titrated and added as per protocol, when target BP was not attained.
Cushman WC, et al., 2007 (123) <u>17599425</u>	<u>Aim</u> : To describe the study design of the BP trial of the ACCORD trial. <u>Study type</u> : description of study design and protocol	Inclusion criteria: Adults with a diagnosis of type 2 DM for at least 3 mo and at high risk for CVD events, who meet the following BP criteria: (1) SBP 130–160 mm Hg and taking 0–3 antihypertensive medications; (2) SBP 161–170 and on 0–2 antihypertensive	Intervention: • Unmasked, open-label, factorial design, randomized trial with a sample size of 4,733 pts • Patients were randomized to intensive SBP control (<120 mm Hg) or standard control (<140 mm Hg)	<u>1° endpoint</u> : Major CVD event (nonfatal MI or stroke, or CV death)	Relevant 2° endpoint: Expanded macrovascular outcome (1° outcome plus coronary revascularization or HF hospitalization), total mortality, each of the separate components of the 1° outcome, HF death or hospitalization, and

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	for the ACCORD RCT	medications; or (3) SBP 171– 180 and taking 0–1 antihypertensive medication. Other entry criteria included spot urine sample <2+, protein–Cr ratio <700 mg protein/1 g creatinine, or 24-h protein excretion <1.0 g/24 h.			composite microvascular disease outcome (kidney and eye disease). <u>Summary:</u> This paper describes the protocol followed in the ACCORD trial that was successful in helping pts to attain and maintain BP targets in the study groups. Once treated, pts had follow-up visits to assessment BP control monthly until BP was at target. Medications were titrated and added as per protocol, when target BP was not attained.
Xu W, et al., 2015 (128) <u>25655523</u>	Aim: Retrospective assessment of the impact of follow-up intervals and treatment intensification thresholds on CVD events Study type: Retrospective cohort Size: 88,756 adult pts with HTN from The Health Improvement Network database	Inclusion criteria: Primary care practices in the U.K., 1986–2010.	N/A	 Median follow-up of 37.4 mo after the treatment strategy assessment period 9,985 (11.3%) pts had an acute CV event or died. No difference in risk of the outcome with systolic intensification thresholds 130– 150 mm Hg, but HR: 1.21 for thresholds >150 mm Hg Outcome risk increased progressively from the lowest (0–1.4 mo) to the highest 5th of time to medication intensification (HR: 1.12; 95% CI: 1.05–1.20; p=0.009) for intensification between 1.4 and 4.7 mo after detection of elevated BP). The highest fifth of time to follow-up (>2.7 mo) was also associated with increased outcome risk HR: 	 Increased risk of acute CVD event or death with: Systolic intensification thresholds >150 mm Hg Delays of >1.4 mo before medication intensification after SBP elevation Delays of >2.7 mo before BP follow-up after antihypertensive medication intensification Timely medical management and follow-up impacts outcomes in the treatment of pts with HTN. Retrospective study, but still sheds important light on the impact of follow-up actions

				1.18; 95% CI: 1.11–1.25; p<0.001	
Birtwhistle RV, et al., 2004 (129) <u>14726370</u>	Aim: Assess impact of follow-up intervals on BP control in stable, treated pts with HTN Study type: RCT Size: 609 pts, 30– 74 y with essential HTN, on drug treatment, with HTN controlled for ≥3 mo prior to entry into study.	Inclusion criteria: 50 family practices in southeastern Ontario, Canada.	• 302 pts randomized to follow- up every 3 mo, 307 randomized to follow-up every 6 mo.	 Pts in both groups visited doctor more frequently than their assigned interval. Mean BP was similar in the groups, as was control of HTN. Pt satisfaction and adherence to treatment were similar in the groups. About 20% of pts in each group had BPs that were out of control during the study. 	 Study addresses follow-up interval for pts with treated, stable, and controlled HTN. No difference in BP control or pt satisfaction between 3 and 6 mo follow-up groups. May be helpful with recommendations for pts with treated, stable HTN.

Data Supplement 29. Monitoring Strategies to Improve Control of BP in Patients on Drug Therapy for High BP (Section 8.3.2)

Study Acronym Author Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (include Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Brennan T, et al., 2010 (130) <u>20415618</u>	Aim: Assess impact of follow-up and monitoring system including home BP monitoring and telephonic nurse case management on BP control in pts treated for HTN <u>Study type:</u> RCT <u>Size</u> : 638 African American pts with high BP from a national health maintenance organization plan	Inclusion criteria: HTN	Intervention: Intervention group received telephonic nurse case management, pt education materials, lifestyle counseling, and a home BP monitor <u>Comparator</u> : Control group received a home BP monitor only	• Intervention group achieved lower SBP (123.6 vs. 126.7 mm Hg, p=0.03) and was 50% more likely than the control group to achieve BP control OR: 1.50; 95% CI: 0.997–2.27; p=0.052	Combination of home BP monitoring and nurse case management controlled HTN better than home BP alone

Bosworth, et al., 2009 (131) <u>19920269</u>	Aim: Assess impact of telephone follow-up intervention and/or home BP monitoring on BP control in pts with treated HTN Study type: RCT Size: 636 pts were randomized; 475 pts completed the trial, including 24-mo follow-up period.	Inclusion criteria: Pts with HTN, from 2 university- affiliated primary care clinics.	• 636 pts randomized to usual care or 1 of 3 intervention groups: (1) Nurse-administered telephone intervention targeting HTN -related behaviors, (2) home BP monitoring 3 times weekly, and (3) both interventions	 475 pts (75%) completed the 24-mo BP follow-up. At 24 mo, improvements in the proportion of pts with BP control relative to the usual care group were 4.3% (95% Cl: -4.5%, 12.9%) in the behavioral intervention group, 7.6% (95% Cl: -1.9%, 17.0%) in the home BP monitoring group, and 11.0% (95% Cl: 1.9%, 19.8%) in the combined intervention group. Relative to usual care, the 24-mo difference in SBP was 0.6 mm Hg (95% Cl: -2.2, 3.4 mm Hg) for the behavioral intervention group, -0.6 mm Hg (95% Cl: -3.6, 2.3 mm Hg) for the BP monitoring group, and -3.9 mm Hg (95% Cl: -6.9 – 0.9 mm Hg) for the combined intervention group; patterns were similar for DBP 	• Home BP monitoring and tailored behavioral telephone intervention improved BP control, SBP, and DBP at 24 mo relative to usual care. Combined therapy was significantly better than either therapy alone.
Bosworth, et al., 2011 (132) <u>21747013</u>	<u>Aim</u> : Assess impact of telephone follow-up interventions on BP control in pts with treated HTN <u>Study type</u> : RCT <u>Size</u> : Of 1551 eligible pts, 593 randomized	Inclusion criteria: Primary care clinics at a VA Medical Center	• 593 pts randomized to either usual care or to 1 of 3 telephone follow-up groups: (1) nurse-administered behavioral management, (2) nurse- and physician- administered medication management, or (3) a combination of both	 1° endpoint: BP control measured every 6 mo for 18 mo Behavioral management and medication management alone showed significant improvements at 12 mo-12.8% (95% Cl: 1.6%, 24.1%) and 12.5% (95% Cl: 1.3%, 23.6%), respectively-but not at 18 mo. In subgroup analyses, among those with poor baseline BP control, SBP decreased in the combined intervention group by 14.8 mm Hg (95% Cl: -21.8– -7.8 mm Hg) at 12 mo and 8.0 mm Hg (95% Cl: -15.5– -0.5 mm Hg) at 18 mo, relative to usual care. 	 Telephone-based case management for high BP control effectively lowers BP for up to 1 y, but then BP control slackens. Interventions had the most impact on pts with worst BP control at study entry. Study carried out in the Veteran's Administration outpatient practice; unclear if results would apply to other practice settings.

Green BB, et al., 2008 (133) <u>18577730</u>	Aim: Assess impact of follow-up and monitoring system including home BP monitoring, Internet-based BP management tool, and pharmacist care management on BP control in pts treated for HTN <u>Study type:</u> Cluster RCT <u>Size</u> : 778 pts from 16 clinics in integrated group practice in Washington state.	Inclusion criteria: Uncontrolled HTN and Internet access	 2 intervention groups: one with home BP monitoring and Internet tool, and the other with home BP monitoring, Internet tool, and pharmacist care management Compare to usual care 1 y follow-up 	 Intervention group with all components achieved better BP control vs. usual care 56% (95% Cl: 49%–62%) or combination intervention group achieved BP control vs. usual care (p<0.001) and intervention with only home BP monitor and Internet tool (p<0.001) 	• Combination of home BP monitoring, Internet-based BP management tools, and pharmacist case management helped control HTN better than usual care and better than BP monitoring and Internet-based tool alone.
Heisler M, et al., 2012 (134) <u>22570370</u>	Aim: Assess impact of follow-up pharmacist care management system on BP control in pts treated for HTN Study type: Cluster RCT Size: 1797 intervention and 2303 control pts from 16 primary care clinics at 5 medical centers (3 VA and 2 Kaiser Permanente)	Inclusion criteria: Uncontrolled HTN and Internet access	 14-mo intervention period BP 6 mo prior to and 6 mo after intervention period were compared in intervention and control groups 	 Mean SBP was 2.4 mm Hg lower (95% CI: -3.4– -1.5), p<0.001 in the intervention group immediately after the intervention period, compared to the control group BP decrease was the same in the intervention and control groups (9 mm Hg). 	 Pharmacist care management system in a "real world" setting was more effective than usual care in lowering BP in the short-term, but in the longer-term follow-up did not differ significantly from usual care. This study is one of very few studies to show no significant longer term impact of a care management system on BP control in pts with HTN.
Margolis KL, et al., 2013 (25) <u>23821088</u>	Aim: Assess impact of follow-up and monitoring system including home BP tele-monitoring and pharmacist case management on BP control in pts treated for HTN <u>Study type:</u> Cluster RCT <u>Size</u> : 450 pts from 16 clinics in integrated health system in Minneapolis, MN	Inclusion criteria: Uncontrolled HTN	 222 pts randomized to 8 usual care clinics and 228 randomized to 8 intervention clinics Intervention included 12 mo of home BP tele- monitoring and pharmacist case management, with 6 mo of follow-up afterward 	 Intervention group achieved better BP control compared to usual care during 12 mo of intervention and persisting during 6 mo of follow-up SBP was <140/90 in 57.2% (95% CI: 44.8%, 68.7%) of intervention pts at 6 and 12 mo vs. 30% (95% CI: 23.2%, 37.8%) in usual care (p=0.001) 	• Combination of home BP tele- monitoring and pharmacist case management helped control HTN better than usual care at 6, 12, and 18 mo

Data Supplement 30, RC	s Comparing Stable Ischem	ic Heart Disease (Section 9.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
INVEST Bangalore S, et al., 2014 (135) 25145522	Aim: To investigate optimal BP in pts ≥60 y with CAD and SBP >150 mm Hg treated with antihypertensive drugs Study type: Post- hoc analysis of PROBE trial (INVEST study— atenolol/HCTZ or verapamil- SR/trandolapril) Size: 8,354 pts	Inclusion criteria: Pts ≥60 y with CAD and SBP >150 mm Hg treated with antihypertensive therapy Exclusion criteria: N/A	Intervention: • 4,787 pts (57%) achieved SBP<140 mm Hg (group 1) • SBP achieved was <140 mm Hg (group 1) Comparator: • 1,747 pts (21%) achieved SBP of 140– 149 mm Hg (group 2); 1,820 pts (22%) achieved SBP ≥150 mm Hg (group 3) • SBP achieved was 140–149 mm Hg (group 2) and 150 mm Hg or higher (group 3)	<u>1° endpoint</u> : All-cause death, nonfatal MI, or nonfatal stroke. Multiple propensity score-adjusted 1° outcome showed that compared with group 1, the risk of 1° outcome adjusted HR: 1.12 (95% CI: 0.95– 1.32; p=0.19); for group 2 adjusted HR: 1.85 (95% CI: 1.59, 2.14), p<0.0001; for group 3 adjusted HR: 1.64 (95% CI: 1.40, 1.93), p<.0001 <u>1° Safety endpoint</u> : No significant difference between the 3 groups	Relevant 2° endpoint: Multiple propensity score- adjusted analysis:• Compared with group 1, no significant difference in all-cause mortality in group 2 but increased all-cause mortality in group 3 (HR: 1.64; 95% Cl: 1.40–1.93; p<0.0001).

					from antihypertensive treatment. No adverse events were reported.
					<u>Summary</u> : The optimal SBP in pts ≥60 y with CAD and SBP >150 mm Hg treated with antihypertensive therapy was <140 mm Hg
Law MR, et al., 2009 (18) <u>19454737</u>	Study type: Meta- analysis of use of BP-lowering drugs in prevention of CVD from 147 randomized trials Size: Of 147 randomized trials of 464,000 pts, 37 trials of BBs in CAD included 38,892 pts, and 37 trials of other antihypertensive drugs in CAD included 85,395 pts	Inclusion criteria: The database search used Medline (1966 to Dec. 2007) to identify randomized trials of BP- lowering drugs in which CAD events or strokes were recorded. The search also included the Cochrane Collaboration and Web of Science databases and the citations in trials and previous meta-analyses and review articles. Exclusion criteria: Trials were excluded if there were <5 CAD events and strokes or if treatment duration was <6 mo.	N/A	1° endpoint: CAD events; stroke Results: In 37 trials of pts with a history of CAD, BBs reduced CAD events 29% (95% CI: 22%, 34%). In 27 trials in which BBs were used after acute MI, BBs reduced CAD events 31% (95% CI: 24%–38%), and in 11 trials in which BBs were used after long-term CAD, BBs insignificantly reduced CAD events 13%. In 7 trials, BBs reduced stroke 17% (95% CI: 1%– 30%). CAD events were reduced 14% (95% CI: 2%–25%) in 11 trials of thiazide diuretics, 17% (95% CI: 11%–22%) in 21 trials of ACEIs, insignificantly 14% in 4 trials of angiotensin receptor blockers, and 15% (95% CI: 8%–22%) in 22 trials of CCBs. Stroke was reduced 38% (95% CI: 28%–47%) in 10 trials of thiazide diuretics, 22% (95% CI: 8%–34%) in 13	With the exception of the extra protective effect of BBs given shortly after a MI and the minor additional effect of CCBs in preventing stroke, all the classes of BP-lowering drugs have a similar effect in reducing CAD events and stroke for a given reduction in BP.
				trials of ACEIs, and 34%	

				(95% CI: 25%–42%) in 9 trials of CCBs.	
HOPE Yusuf S, et al., 2000 (136) <u>10639539</u>	Aim:To investigate effect of ACE-I (Ramipril 10 mg) on CV events in high risk pts. over 5y with a mean entry BP of 139/79 mm Hg in both groupsStudy type:RCT, 2×2 factorial designSize:9,297	Inclusion criteria: Pts ≥55 y with history of CAD, stroke, PVD or DM with either HTN, elevated total cholesterol, low LDL cholesterol, smoking, or micro albuminuria. Exclusion criteria: HF, <0.40 EF, on ACE-I or Vitamin E, uncontrolled HTN /overt nephropathy, Had MI or stroke <4 wk	Intervention: Ramipril (10 mg) (4,645) Comparator: Placebo (4,652)	<u>1° endpoint</u> : Composite of MI, stroke, or mortality from CV causes. <u>Results:</u> Endpoint reduction Ramipril group vs. Placebo (14% vs. 17.8%; RR: 0.78; CI: 0.70–0.86; p<0.001)	 Death from cardiac causes reduced (6.1% vs. 8.1%; p<0.001) Death from MI reduced (9.9% vs. 12.3%; p<0.001) Death from any cause (10.4 % vs. 12.2%; p=0.005)
SAVE Pfeffer M., et al., 1992 (137) <u>1386652</u>	Aim: To assess if captopril decrease morbidity and mortality in pts with LV dysfunction after MI. <u>Study type</u> : RCT <u>Size</u> : 2,231	Inclusion criteria: Pts (21–80 y) surviving 3 d after MI, EF≤40%. Exclusion criteria: Pts not randomized within 16 d after MI, contra. to ACE-I use, Serum Cr. >2.5 mg/dL, severe comorbidities, unstable infarction, need for revascularization	Intervention: Captopril (titrated doses) (115) <u>Comparator</u> : Placebo (1116)	<u>1° endpoint and results</u> : All-cause mortality: 20% vs. 25%, RR: 19%; 95% CI: 3%–32%; p=0.019 <u>Other endpoints</u> : Fatal and nonfatal major CV events were reduced in the captopril group.	 Captopril vs. Placebo group BP at 1 y (125±18 / 77±10 mm Hg for placebo vs. 119±18/74±10 mm Hg for captopril; p<0.001) Dizziness, alteration in taste, cough and diarrhea were reported significantly more in the captopril group Ventricular size on Echo studies was independent predictor of adverse CV outcomes
EUROPA Fox KM, et al., 2003 (138) <u>13678872</u>	<u>Aim</u> : To investigate efficacy of perindopril in CV events in pts with stable CAD. <u>Study type</u> : RCT	Inclusion criteria: Pts ≥18 y (women) with CAD >mo before screening, revascularization >6 mo before screening, ≥70% narrowing of major	Intervention: Perindopril (6,110) Comparator: Placebo (6,108)	<u>1º endpoint</u> : Composite of CV death, nonfatal MI, cardiac arrest with successful CPR <u>Results:</u> RR 20%; 95% CI: 9%–29; p=0.0003	• Perindopril resulted reduction in all these outcomes: composite of total mortality, nonfatal MI, hospital admission for UA, and cardiac arrest with successful CPR; CV mortality and nonfatal MI, the individual components these outcomes and revascularization, stroke, and admission for HF

MERIT-HF Goldstein S, et al., 1999 (139) 10526701	Size: 12,218 pts <u>Aim</u> : To investigate if metoprolol (CR/XL) once daily with std. treatment lowers mortality in pts with HF <i>r</i> EF <u>Study type</u> : RCT <u>Size</u> : 3,991 pts	coronary artery. Men with history of chest pain, positive ECG, echo or nuclear test $\frac{Exclusion criteria:}{echo or nuclear test}$ $\frac{Exclusion criteria:}{echo or nuclear test}$ $\frac{Exclusion criteria:}{echo or nuclear test}$ $\frac{Exclusion criteria:}{echo or nuclear test}$ $\frac{Inclusion criteria:}{entry, Stable clinical}$ $\frac{Exclusion criteria:}{echo or nuclear test}$	Intervention: Metoprolol CR/XL (1,990) <u>Comparator</u> : Placebo (2,001)	1° endpoint: All-cause mortality in the intent to treat Results: 145 vs. 217 deaths [11.0 %], RR: 0.66 (95% Cl: 0.53–0.81; p=0.0009) or adjusted for interim analyses p=0.0062.	 Fewer sudden deaths in the metoprolol group (p=0.0002) Lesser deaths from HF<i>r</i>EF in the metoprolol group (p=0.002) Metoprolol improved survival and was well tolerated
		run-in phase, EF ≤0.40.		interim analyses	

Packer M, et al., 2001 (140) <u>11386263</u>	Aim: To assess survival in severe chronic HF pts by the use of carvedilol. Study type: RCT Size: 2,289 pts	Inclusion criteria: HF pts with dyspnea/exertion for 2 mo at least and left EF<25% despite treatment clinically euvolemic; allowed on digitalis, nitrates, hydralazine, spironolactone, or	Intervention: Carvedilol (1,156) Comparator: Placebo (1,133)	1° endpoint: • Death from any cause 130 vs. 190 deaths RR: 35%; 95% CI: 19%–48%; p=0.00013 • Combined risk of death/hospitalization (24% lower risk in the carvedilol; (95% CI: 13%–33%; p<0.001	 Study stopped early (1.3-y follow-up) due to benefit on survival Long-term treatment is very valuable. Not all the pts with severe HF were allowed in the study
		amiodarone. Hospitalized pts with no acute illness. <u>Exclusion criteria</u> : HF due to uncorrected prim. valvular disease or		Safety endpoint: Lesser pts in carvedilol group required permanent discontinuation because of adverse events or for reasons other than death	
		reversible cardiomyopathy cardiac transplant pts., coronary revasc. <2 mo, acute MI or stroke, ventricular tachycardia, on alpha		(p=0.02)	
		blocker or CCB or on antiarrythmics class I <4 wk, SBP <85 mm Hg, serum Cr >2.8 mg/dL, change in body weight >1.5 kg during screening.			
CAPRICORN Dargie HJ, et al., 2001 (141) <u>11356434</u>	<u>Aim</u> : To investigate outcomes after carvedilol after MI in pts with LV dysfunction.	Inclusion criteria: Pts ≥18 y, MI within 3–21 d of entry, LVEF≤40%, concurrent ACEI stable dose for at least 24 h, HF pts treated and	Intervention: Carvedilol (975) Comparator: Placebo (984)	<u>1° endpoint</u> : All-cause mortality or hospital admissions for CV issues <u>Results</u> : 12% vs. 15%; RR: 23%; 95% CI: 0.60–	 CV mortality, nonfatal MI reduced in the carvedilol group No difference between groups SCD and admission due to HF
	<u>Study type</u> : RCT <u>Size</u> : 1,959 pts	controlled with ACEI and diuretics but not inotropes.		0.98; p=0.03 No difference between groups for death or CV hospital admissions	

MERIT-HF HTN Herlitz J, et al., 2002 (142) <u>11862577</u>	<u>Aim</u> : To assess metoprolol CR/XL influence on mortality and hospitalizations in HF and HTN pts. <u>Study type</u> : RCT <u>Size</u> : 1,747 pts	Exclusion criteria: SBP<90 mm Hg, uncontrolled HTN, bradycardia, insulin- dependent DM, BBs not for HF, Beta-2 agonists and steroids Inclusion criteria: Same as above MERIT- HF, 1999 study (HTN subgroup) Exclusion criteria: Same as above MERIT- HF	Intervention: Metoprolol CR/XL (871) <u>Comparator</u> : Placebo (876)	<u>1° endpoint</u> : Total mortality <u>Results</u> : RR: 0.61; 95% Cl: 0.44–0.84; p=0.0022	 Total mortality reduction was driven by reduction in the SCD and death from worsening HF 12.5% pts had earlier discontinuation due to any cause. Lesser no. of pts in the metoprolol group (n=21) discontinued due to worsening HF The mean reduction in BP (adjusted) was 1.7 mm Hg in the metoprolol group vs. 4.8 mm Hg in placebo group (p=0.0001)
CIBIS-II 1999 (143) <u>10023943</u>	<u>Aim</u> : To determine efficacy of bisoprolol in reducing mortality in chronic HF. <u>Study type</u> : RCT <u>Size</u> : 2,647 pts	Inclusion criteria: 18– 80 y, LVEF≤35%, dyspnea, orthopnea, fatigue, NYHA class III- IV Exclusion criteria: Uncontrolled HTN, MI, UA <3 mo revascularization. treatment, heart transplant, AV block <1 degree, SBP <100 mm Hg, renal failure, reversible obstructive lung disease	Intervention: Bisoprolol (1,327) Comparator: Placebo (1,320)	<u>1° endpoint</u> : All-cause mortality <u>Results</u> : 11.8% vs. 17.3% deaths with a RR: 0.66; 95% CI: 0.54–0.81; p<0.0001	 The trial stopped early due to benefit. Bisoprolol group had significantly fewer SCDs. Mean age was 61 y so more data on elderly pts is needed
Elkayam U, et al., 1990 (144) <u>2242521</u>	Aim: To assess comparative efficacy and safety of nifedipine and ISDN alone and the combination for treating for chronic CHF.	Inclusion criteria: 18– 75 y HF pts, NYHA class II and III, LVEF<40%, clinically stable, maintenance dose of Digitalis and diuretics.	Intervention: Nifedipine (21), ISDN (20), Nifedipine+ISDN (23) Comparator: Placebo	Endpoints and Results: <u>HF-worsening</u> : 9 in Nifedipine group vs. 3 in ISDN group (p<0.09); and 21 in nifedipine-ISDN group (p<0.001 vs. nifedipine, p<0.0001 vs. ISDN)	 In clinical deterioration nifedipine pts (8) vs. rest of the pts (No difference in LVEF or VO² max) Although all the 3 drug regimens improved exercise capacity, nifedipine treatment alone or in combination resulted in clinical deterioration and worsening of CHF

The Multicenter Dilitiazem Postinfarction Research Group 1988 (145) <u>2899840</u>	Study type: RCT with a crossover design Size: 28 pts Aim: To assess dilitiazem effect on recurrent infarction and death after acute MI Study type: RCT Size: 2,466 pts Size: 2,466 pts	Exclusion criteria: Pregnancy, nursing, history of MI <1 mo before entry, valvular disease, Angina, significant pulmonary, hepatic, renal and hematologic disease, unable to walk on the treadmill, noncompliance Inclusion criteria: 25–75 y admitted to CCU, MI with enzyme confirmation. Exclusion criteria: • Cardiogenic shock, • Symptomatic hypotension, • PH with right HF, • 2nd/3rd degree heart block, • HR <50 bpm, • Contraceptives, • WPW syndrome, • CCBs, • Severe comorbidities or • Cardiac surgery Inclusion criteria:	Intervention: Dilitiazem 240 mg (1,234) Comparator: Placebo (1,232)	Clinical deterioration discontinuation: Nifedipine 29% vs. ISDN group 5% (p<0.05) DBP: Nifedipine alone or combination with ISDN (reduction, p<0.05) 1º endpoints and results: • Total mortality: identical in both groups • Cardiac death and nonfatal MI: 11% fewer in dilitiazem but difference was NS	No combined benefit from dilitiazem on mortality or cardiac events
Goldstein RE, et al., 1991 (146) <u>1984898</u>	Aim: To determine in dilitiazem increases late onset CHF in post-MI pts with early decline in EF. <u>Study type</u> : RCT <u>Size</u> : 2,466 pts	Same as above Exclusion criteria: Same as above	Dilitiazem 240 mg (1,234) <u>Comparator</u> : Placebo (1,232)	<u>1° endpoint and results</u> : Same as above <u>Follow-up Results</u> : Pts with BL EF<0.40, late CHF in Dilitizam group (21%) vs. Placebo (12%) [p=0.004].	 Life table analysis confirmed increased frequency of late CHF in pts taking dilitiazem (p=0.0017) Dilitiazem related CHF exclusively associated with systolic LVD with or without BBs

Freemantle N, et al., 1999 (147) <u>10381708</u>	Aim: To evaluate BBs effectiveness for short-term treatment and long-term 2° prevention in acute MI. Study type: Meta- analysis of RCTs Size: 54,234 pts (82 RCTs)	Inclusion criteria: RCTs with treatment lasting >1 d and with follow-ups on clinical effectiveness in pts with MI Exclusion criteria: Cross-over RCTs	Intervention: BBs (mostly propranolol, timolol, metoprolol) Comparator: Controls (placebo/other treatment)	 <u>1° endpoint</u>: All-cause mortality <u>Results</u>: Long-term trials RR reduction: 23% (95% CI: 15%–31%) Short-term trials RR reduction: 4%; 95% CI: -8%–5% 	 Meta-regression in long-term trials indicated a near significant trend for decreased benefit in drugs with ISA. NS in withdrawal between BBs of different cardio selectivity.
de Peuter OR, et al., 2009 (148) <u>19841485</u>	Aim: To determine influence of beta-2 blockade in addition to beta-1 blockade for preventing vascular events in pts with ACS or HF. Study type: Meta- analysis of RCTs Size: 34,360 pts (33 RCTs)	Inclusion criteria: • RCTs comparing Beta-1 blockers vs. BBs 1 + 2 directly (5) • RCTs comparing Beta-1 blockers vs. Beta 1 + 2 blockers with a control group (28) Exclusion criteria: Studies not pre- specifying total mortality and vascular event as outcomes <3 mo follow- up, duplicate data, sub studies.	Intervention: Beta-1 blockers Comparator: BBs 1+2 with or without control group	<u>1° endpoint</u> : Total mortality, vascular events. <u>Results</u> : <u>ACS Population:</u> 1 study with different BBs underpowered to detect difference. Beta-1 vs. Placebo NS reduced mortality or vascular events <u>HF population:</u> Beta 1 + 2 blockers vs. Beta 1 blockers decreased mortality RR: 0.86; 95% CI: 0.78–0.94 Beta 1 and Beta 1+2 decreased total mortality. Only Beta 1+2 blockers reduced vascular events.	 Supplementary beta 2 blockade may be more beneficial. Indirect comparisons and heterogeneity among studies
Leon MB, et al., 1981 (149) <u>7246435</u>	<u>Aim</u> : To evaluate effectiveness of verapamil as a single agent and in combination with propranolol in pts with stable AP.	Inclusion criteria: Symptomatic angina pectoris pts, 1) not sufficiently controlled on BBs and nitrates and noncardiac	Intervention: Propranolol, verapamil, Combination of propranolol and verapamil	<u>Results</u>: Large dose verapamil significantly lowered BP. Propranolol and verapamil combined (at best dose) further lowered BP, improved	 HR and pressure-rate product lowered significantly on combination therapy PR interval increased on combination treatment Regarding antianginal properties, verapamil seemed to be more effective than propranolol.

		effects from propranolol	Comparator: Placebo	exercise time by 4.7 ± 0.7	
	Study type: RCT	hindering treatment	<u>oomparator</u> i nacobo	min (p<0.001)	
	(triple crossover)	2) who could stay 4 wk		N /	
		in hospital			
	<u>Size</u> : 11 pts				
		Exclusion criteria: LVD			
		with CHF or LVEF<30%			
		at rest and <25% for			
		exercise, HR<50 b/min,			
		≥first degree heart block			
Staessen JA, et	Aim: To determine if	Inclusion criteria: Pts	Intervention: Active	1° endpoint: Fatal and	 All fatal and nonfatal cardiac endpoints (with sudden
al., 1997 (150)	active treatment	≥60 y, sitting SPB 160–	treatment (2,398)	nonfatal strokes	death) decreased in the active treatment group
<u>9297994</u>	reduces	219 mm Hg, sitting DBP		combined.	(p=0.03)
	complications from	95 mm Hg, and	Comparator: Placebo		 Cardiac mortality was lower in active treatment (-
	isolated systolic HTN	standing SBP ≥140 mm	(2,297)	Results: 13.7 vs. 7.9	27%; p=0.07). All-cause mortality was not different.
	in the elderly.	Hg.		endpoints/ 1,000 pts-y (42% reduction; p=0.003)	 Nitrendipine used for active arm.
	Study type: RCT	Exclusion criteria:			
		Systolic HTN 2nd to a			
	<u>Size</u> : 4,965 pts	disorder, retinal			
		hemorrhages/papillede			
		ma, CHF, aneurysms,			
		serum Cr ≥180 µmol/L,			
		history of nosebleed,			
		stroke, MI <1 y,			
		dementia, substance			
		abuse, severe			
		comorbidities			
Wright JT, et al.,	Aim: To compare in	Inclusion criteria:	Intervention: 4,678	<u>1º endpoint</u> :	• At 3.26-y median follow-up, compared with standard
2015 (114)	pts with a SBP of	9,361 pts, mean 67.9 y	pts were randomized	• At 1 y, the mean SBP	BP treatment, intensive BP treatment reduced all-
<u>26551272</u>	130–180 mm Hg and an increased CV risk	(28.2% ≥75 y; 35.6% women; 57.7% non-	to intensive BP treatment	was 121.4 mm Hg with	cause mortality 27% (p=0.003), HF 38% (p=0.002),
	but without DM the	Hispanic white; 31.5%	ueauneni	intensive treatment (mean	CV mortality 43% (p=0.005), and the 1° composite outcome or death 23% (p < 0.001)
	effect of a target SBP	African American;	Comparator: 4,683	number of	outcome or death 22% (p<0.001)
	of <140 mm Hg vs. a	10.5% Hispanic) with a	pts were randomized	antihypertensive drugs was 2.8) and 136.2 mm	• Intensive BP treatment reduced the 1° composite
	target SBP of <120	SBP of 130–180 mm Hq	to standard BP	Hg with standard	endpoint 33% (14% to 49%) in pts aged 75 y and older and 20% (0% to 36%) in pts 50–74 y
	mm Hg on the 1°	and an increased CV	treatment	treatment (mean number	Serious adverse events were similar in both
	composite outcome	risk but without DM,		of antihypertensive drugs	• Serious adverse events were similar in both treatment groups. However, intensive BP treatment
	of MI, other ACSs,	history of stroke,		was 1.8)	caused more hypotension (2.4% vs. 1.4%; p=0.001),
		symptomatic HF within		was 1.0j	more syncope (2.3% vs. 1.7% ; p=0.05), more
	l			1	more syncope (2.3% vs. 1.7%, p=0.03), more

	stroke, HF, or CV death	past 6 mo, LVEF <35%, and eGFR <20 mL/min/1.73 mm ² ; CVD was present in 20.1%, and the Framingham 10-y CVD risk score was ≥15% in 61.3% of pts		• At 3.26-y median follow- up, the 1° composite outcome was reduced 25% (p<0.001) by intensive BP treatment	electrolyte abnormality (3.1% vs. 2.3%; p=0.02), and more acute kidney injury or acute renal failure (4.1% vs. 2.5%; p<0.001). The incidence of bradycardia, injurious falls, and orthostatic hypotension with dizziness was similar in both treatment groups
ALLHAT Collaborative Research Group, 2003 <u>12925554</u>	Aim: In a follow-up analysis, to compare diuretic vs. alpha- blocker as first step treatment of hypertension.	Inclusion criteria: Men and women ≥ 55 y with BP ≥140/90 mm Hg or on medications for hypertension with at least one additional risk factor for coronary heart disease.	Intervention: 15,255 patients were randomized to chlorthalidone and 9,061 to doxazosin and followed for 3.2 y.	Primary endpoint: Combined fatal coronary heart disease or non-fatal MI, analyzed by intention to treat.	 There was no difference in primary outcome between the arms (RR: 1.02; 95% CI: 0.94–1.13). However, the doxazosin arm compared with the chlorthalidone arm had a higher risk for stroke (RR: 1.26; 95% CI: 1.10–1.46) and combined cardiovascular disease (RR: 1.20; 95% CI: 1.13–1.27). The findings confirmed the superiority of diuretic-based over alpha blocker based antihypertensive treatment in the prevention of cardiovascular disease.
Zanchetti A, et al., 2006 <u>17053536</u>	Aim: To provide additional analyses of the primary endpoint in the VALUE trial, including sex, age, race, geographic region, smoking status, type 2 diabetes, total cholesterol, left ventricular hypertrophy, proteinuria, serum creatinine, history of coronary heart disease, stroke or transient ischemic attack and history of peripheral artery disease.	Inclusion criteria: The 15,245 patients participating in VALUE were divided into subgroups according to baseline characteristics.	Statistical analysis: Subgroup interaction analyses were conducted by the Cox proportion hazard model. Within each subgroup, treatment effects were assessed by hazard ratios and 95% CIs.		 For cardiac morbidity and mortality, the only significant subgroup by treatment interaction was of sex (p=0.016) with HR indicating a relative excess of cardiac events in women but not in men, but SBP differences in favor of amlodipine were greater in women. In the VALUE cohort, in no subgroup of patients were there differences in the incidence of the composite cardiac endpoint with valsartan and amlodipine treatment despite greater BP reduction in the amlodipine group.

Leenen FHH, et al., 2006 <u>16864749</u>	<u>Aim</u> : To compare the long-term relative safety and outcomes of ACE inhibitor- and CCB-based regimens in older hypertensive individuals in ALLHAT.	Inclusion criteria: men and women age ≥55 y with untreated (BP 140– 180/90–110 mm Hg) or treated hypertension (BP ≤160/100 mm Hg on ≤2 antihypertensive drugs) with ≥ 1 additional risk factor for coronary heart disease.	Intervention: Patients (were randomized to amlodipine (9,048) or Lisinopril (9,054).	Primary outcome: Combined fatal coronary heart disease or non-fatal MI, analyzed by intention to treat. Follow-up: 4.9 y	 Risk of coronary heart disease was similar between amlodipine and Lisinopril For stroke, combined cardiovascular disease, gastrointestinal bleeding and angioedema, risks are higher with Lisinopril compared to amlodipine. For heart failure, risks are higher with amlodipine compared to Lisinopril.
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Data Supplement 31. Meta-analyses of ischemic heart disease (Section 9.1)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR; and CI)	Summary/Conclusion Comment(s)
Bundy JD, et al., 2017 <u>28564682</u>	Study type: Network meta- analysis Size: 144,220 patients in 42 RCTs.	Inclusion criteria: • Random allocation into an antihypertensive medication, control or treatment target • Allocation to antihypertensive Antihypertensive treatment was independent of other treatment regimens • ≥100 patients in each treatment group • Trial duration ≥ 6 mo • One or more events for each treatment group reported • Minimum 5 mm Hg difference in SBP level between the 2 treatment groups • Outcomes included major CVD, stroke, CHD, CVD mortality or all- cause mortality	• There were linear associations between mean achieved SBP and risk of cardiovascular disease and mortality, with the lowest risk at 120 to 124 mm Hg. Randomized groups with a mean achieved SBP of 120 to 124 mm Hg had a hazard ratio (HR) for major cardiovascular disease of 0.71 (95% CI: 0.60–0.83) compared with randomized groups with a mean achieved SBP of 130 to 134 mm Hg, an HR of 0.58 (95% CI: 0.48–0.72) compared with those with a mean achieved SBP of 140 to 144 mm Hg, an HR of 0.46 (95% CI: 0.34–0.63) compared with those with a mean achieved SBP of 150 to 154 mm Hg, and an HR of 0.36 (95% CI: 0.26–0.51) compared with those with a mean achieved SBP of 160 mm Hg or more.	• This study suggests that reducing SBP to levels below currently recommended targets significantly reduces the risk of cardiovascular disease and all- cause mortality and strongly support more intensive control of SBP among adults with hypertension.

Data Supplement 22 Neprendemized Trials	Obconvational Studios and/or Dog	istrias of Isabamia Haart Disaasa (Saction 0.1)
Data Supplement 52. Normanuomizeu mais	, Observational Studies, and/or Rey	istries of Ischemic Heart Disease (Section 9.1)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; and 95% CI)	Summary/Conclusion Comment(s)
PROVE IT-TIMI 22 Bangalore S, et al., 2010 (151) 21060068	Study type: Nonrandomized trial of optimal BP after ACS Size: 4,162 pts	Inclusion criteria: Pts with acute MI or high-risk UA within 10 d randomized to pravastatin or atorvastatin and to gatifloxacin or placebo treated with standard medical and interventional treatment for ACS Exclusion criteria: N/A	<u>1° endpoint</u> : Composite of all-cause death, MI, UA requiring rehospitalization, revascularization after 30 d, and stroke with a mean follow-up of 24 mo <u>Results:</u> The relationship between SBP and DBP followed a J- or U-shaped curve association with the 1° outcome with increased events rates at both low and high BP values. A nonlinear Cox proportional hazards model showed a nadir of 136/85 mm Hg (range 130–140/80–90 mm Hg) at which the incidence of 1° outcome was lowest. There was a relatively flat curve for SBP of 110–130 mm Hg and for DBP of 70–90 mm Hg, suggesting a BP <110/70 mm Hg may be dangerous.	• After an ACS, a J- or U-shaped association existed between BP and the incidence of new CV events. The lowest incidence of CV events occurred with a BP of 130–140/80–90 mm Hg and a relatively flat curve for SBP of 110–130 mm Hg and of DBP of 70–90 mm Hg, suggesting a BP <110/70 mm Hg may be dangerous.
Law MR, et al., 2009 (18) <u>19454737</u>	Study type: Meta- analysis of use of BP-lowering drugs in prevention of CVD from 147 randomized trials Size: Of 147 randomized trials of 464,000 pts, 37 trials of BBs in CAD included 38,892 pts, and 37 trials of other antihypertensive drugs in CAD included 85,395 pts	Inclusion criteria: The database search used Medline (1966 to Dec. 2007) to identify randomized trials of BP-lowering drugs in which CAD events or strokes were recorded. The search also included the Cochrane Collaboration and Web of Science databases and the citations in trials and previous meta-analyses and review articles. Exclusion criteria: Trials were excluded if there were <5 CAD events and strokes or if treatment duration was <6 mo.	<u>1° endpoint</u> : CAD events; stroke <u>Results:</u> In 37 trials of pts with a history of CAD, BBs reduced CAD events 29% (95% CI: 22%, 34%). In 27 trials in which BBs were used after acute MI, BBs reduced CAD events 31% (95% CI: 24%–38%), and in 11 trials in which BBs were used after long-term CAD, BBs insignificantly reduced CAD events 13%. In 7 trials, BBs reduced stroke 17% (95% CI: 1%–30%). CAD events were reduced 14% (95% CI: 2%–25%) in 11 trials of thiazide diuretics, 17% (95% CI: 11%– 22%) in 21 trials of ACEIs, insignificantly 14% in 4 trials of angiotensin receptor blockers, and 15% (95% CI: 8%–22%) in 22 trials of CCBs. Stroke was reduced 38% (95% CI: 28%–47%) in 10 trials of thiazide diuretics, 22% (95% CI: 8%–34%) in 13 trials of ACEIs, and 34% (95% CI: 25%–42%) in 9 trials of CCBs.	• With the exception of the extra protective effect of BBs given shortly after a MI and the minor additional effect of CCBs in preventing stroke, all the classes of BP-lowering drugs have a similar effect in reducing CAD events and stroke for a given reduction in BP.

Study Acronym (if applicable) Author Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; and 95% CI)	Summary/Conclusion Comment(s)
LV J, et al., 2013 (127) <u>23798459</u>	Study type: MA of RTC that randomly assigned individuals to different target BP levels Size: 37,348 pts	•15 trials	 7.5/4.5 mm Hg BP difference. Intensive BP lowering achieved. <u>RR for</u> Major CV events: 11%; 95% CI: 1%–21%) MI: 13%; 95% CI: 0%–25% Stroke: 24%; 95% CI: 8%–37% ESRD: 11%; 95% CI: 3%–18% Albuminuria: 10%; 95% CI: 4%–16% Retinopathy 19%; 95% CI: 0%–34% p=0.051 	More intensive strategy for BP control reduced cardio-renal endpoint
Xie X, et al., 2015 (21) <u>26559744</u>	Study type: MA of RTC that randomly assigned individuals to different target BP levels Size: 44,989 pts	• 19 trials	Achieved BP 133/76 mm Hg (intensive) 140/81 (less intense) • Major CV events: 14%; 95% CI: 4%–22% • MI: 13%; 95% CI: 0%–24% • Stroke: 22%; 95% CI: 10%–32% • Albuminuria: 10%; 95% CI: 3%–16% • Retinopathy progression: 19%; 95% CI: 0%–34%. • More intensive had no effects on HF: 15%; 95% CI: -11%–34% • CV death: 9%; 95% CI: -11%–26% • Total mortality: 9%; 95% CI: -3%–19% • ESKD: 10%; 95% CI: -6%–23%	• More intensive approach reduced major CV events (stroke and MI) except heat failure, CVD, ESRD, and total mortality.
Thomopolous C, et al., 2016 (54) <u>26848994</u>	Study type: Meta- analysis of RTCs of more vs. less intense BP control	• 16 trials (52,235 pts) compared more vs. less intense treatment 34 (138,127 pts) active vs. placebo	More intense BP • Stroke RR: 0.71; 95% CI: 0.60–0.84) • CHD RR: 0.80; 95% CI: 0.68–0.95) • Major CV events RR: 0.75; 95% CI: 0.68–0.85 • CV mortality RR: 0.79; 95% CI: 0.63–0.97 Stratification of SBP cutoffs (150,140 and 130 mm Hg) showed that a SBP/DBP difference of 10/5 mm Hg across each cutoff reduced risk of all outcomes	• Intensive BP reduction improves CV outcomes compared to less intense Achieved BP <130/80 may be associated with CV benefit.

Data Supplement 33. RCTs Comparing Heart Failure (Section 9.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% Cl)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Herlitz J, et al., 2002 (142) <u>11862577</u>	Aim: To see effect of metoprolol vs. placebo on mortality and hospitalizations among pts with history of HTN and HF with reduced LVEF Study type: RCT Size: 1,747 pts	Inclusion criteria: NYHA class II–IV HF with LVEF ≤40% within 3 mo of enrollment; supine resting HR ≥68 bpm; stable clinical condition Exclusion criteria: Acute MI or UA within 28 d of randomization; indication or contraindication for treatment with BBs or drugs with beta-blocking properties; poor compliance; CABG surgery or PTCA in past 4 mo	Intervention: • Administration of metoprolol • 871 pts randomized to metoprolol <u>Comparator</u> : • Administration of placebo • 876 pts randomized to placebo	 <u>1° endpoint</u>: At 1-y follow-up, compared with placebo, metoprolol reduced all-cause mortality 39% (95% Cl: 16%–56%; p=0.002) and all-cause mortality or all-cause hospitalization 24% (95% Cl: 11%–35%; p=0.0007) <u>1° Safety endpoint</u>: Early permanent cessation of drug was 12.5% for metoprolol and 15.9% for placebo (p=0.048); 21 pts on metoprolol and 35 pts on placebo had early cessation because of worsening 	Relevant 2° endpoint: At 1-y follow- up, compared with placebo, metoprolol reduced CV death 41% (95% CI: 17%–57%; p=0.002), death from HF: 51% (95% CI: 1%–75%; p=0.042), sudden cardiac death 49% (95% CI: 21%–67%; p=0.002), all- cause mortality or HF hospitalization 28% (95% CI: 11%–42%; p=0.002), and cardiac death or nonfatal acute MI 44% (95% CI: 23%–60%; p=0.0003)Study limitations and adverse events: Early permanent cessation of drug was 12.5% for metoprolol and 15.9% for placebo (p=0.048); 21 pts on M and 35 pts on placebo had early cessation because of worsening HF; all-cause withdrawals were 22% less with metoprolol; (p=0.048); adverse events were 28% less with metoprolol (p=0.026); worsening HF was 41% less with metoprolol (p=0.056)Summary: In an RCT of pts with HF with reduced EF and a history of HTN, compared with placebo, metoprolol succinate reduced all- cause mortality and all-cause mortality or all-cause hospitalization
Packer M, et al., 2001 (140) <u>11386263</u>	Aim: To assess survival in severe	Inclusion criteria: HF pts with dyspnea/exertion for 2 mo at least and left EF<25% despite	Intervention: Carvedilol (1,156)	 <u>1° endpoint</u>: Death from any cause 130 vs. 190 deaths (RR: 35%; 	Study stopped early (1.3 y follow- up) due to benefit on survival

Data Supplement 34. RCTs Comparing HF*r*EF (Section 9.2.1)

CAPRICORN Dargie HJ, et al.,	chronic HF pts by the use of carvedilol. <u>Study type</u> : RCT <u>Size</u> : 2,289 pts <u>Aim</u> : To investigate outcomes after	treatment clinically euvolemic; allowed on digitalis, nitrates, hydralazine, spironolactone, or amiodarone. Hospitalized pts with no acute illness. Exclusion criteria: HF due to uncorrected prim. valvular disease or reversible cardiomyopathy, cardiac transplant pts., coronary revasc. <2 mo, acute MI or stroke, ventricular tachycardia, on alpha blocker or CCB or on antiarrythmics class I <4 wk, SBP <85 mm Hg, serum Cr >2.8 mg/dL, change in body weight >1.5 kg during screening. Inclusion criteria: Pts ≥18 y, MI within 3–21 d of entry, LVEF	Comparator: Placebo (1,133) Intervention: Carvedilol (975)	 95% CI: 19%–48%; p=0.00013) Combined risk of death/hospitalization (24% lower risk in the carvedilol; 95% CI: 13%–33%; p<0.001) <u>Safety endpoint</u>: Lesser pts in carvedilol group required permanent discontinuation because of adverse events or for reasons other than death (p=0.02) <u>1° endpoint</u>: All-cause mortality or hospital 	 Long-term treatment is very valuable. Not all the pts with severe HF were allowed in the study Output of the study CV mortality, nonfatal MI reduced in the carvedilol group
2001 (141) <u>11356434</u>	carvedilol after MI in pts with LV dysfunction. <u>Study type</u> : RCT <u>Size</u> : 1,959 pts	 ≤40%, concurrent ACEI stable dose for at least 24 h, HF pts treated and controlled with ACEI and diuretics but not inotropes. Exclusion criteria: SBP <90 mm Hg, uncontrolled HTN, bradycardia, insulin-dependent DM, BBs not for HF, Beta-2 agonists, and steroids 	<u>Comparator</u> : Placebo (984)	admissions for CV issues <u>Results</u> : 12% vs. 15%; RR: 23% (95% CI: 0.60–0.98; p=0.03) No difference between groups for death or CV hospital admissions	 No difference between groups sudden death and admission due to HF
Elkayam U, et al., 1990 (144) <u>2242521</u>	Aim: To assess comparative efficacy and safety of nifedipine and ISDN alone and the combination for treating for chronic CHF.	Inclusion criteria: 18–75 y old HF pts, NYHA class II and III, LVEF<40%, clinically stable, maintenance dose of Digitalis and diuretics. Exclusion criteria: Pregnancy, nursing, history of MI <1 mo before entry, valvular disease, angina, significant pulmonary,	Intervention: Nifedipine (21), ISDN (20), Nifedipine+ISDN (23) Comparator: Placebo	Endpoints and Results: • HF-worsening: 9 in Nifedipine group vs. 3 in ISDN group (p<0.09); and 21 in nifedipine-ISDN group (p<0.001 vs. nifedipine, p<0.0001 vs. ISDN) • Clinical deterioration discontinuation:	 In clinical deterioration nifedipine pts (8) vs. rest of the pts (No difference in LVEF or VO² max.) Although all the 3 drug regimens improved exercise capacity, nifedipine treatment alone or in combination resulted in clinical deterioration and worsening of CHF

MDPIT Goldstein RE, et al., 1991 (146) <u>1984898</u>	Study type: Crossover RCT Size: 28 pts Aim: To determine if dilitiazem increases late onset CHF in post-MI pts with early decline in EF. Study type: RCT Size: 2,466 pts	hepatic, renal and hematologic disease., unable to walk on the treadmill, noncompliance Inclusion criteria: 18–75 y HF pts, NYHA class II and III, LVEF <40%, clinically stable, maintenance dose of digitalis and diuretics. Exclusion criteria: Pregnancy, nursing, history of MI <1 mo before entry, valvular disease, Angina, significant pulmonary, hepatic, renal and hematologic disease., unable to walk on the treadmill, noncompliance	Intervention: Dilitiazem 240 mg (1,234) Comparator: Placebo (1,232)	Nifedipine 29% vs. ISDN group 5% (p<0.05) • DBP: Nifedipine alone or combination with ISDN (reduction, p<0.05) 1° endpoint and results: • HF-worsening: 9 in Nifedipine group vs. 3 in ISDN group (p<0.09); and 21 in nifedipine-ISDN group (p<0.001 vs. nifedipine, p<0.0001 vs. ISDN) • Clinical deterioration discontinuation: Nifedipine 29% vs. ISDN group 5% (p<0.05) • DBP: Nifedipine alone or combination with ISDN (reduction, p<0.05) Follow-up Results: Pts with BL EF<0.40, late CHF in Dilitizam group (21%) vs.	 Life table analysis confirmed increased frequency of late CHF in pts taking dilitiazem (p=0.0017) Dilitiazem related CHF exclusively associated with systolic LVD with or without BB s
Cohn JN, et al., 2001 (152) <u>11759645</u>	<u>Aim</u> : To determine the effect of valsartan vs. placebo on mortality plus morbidity in pts with HF <i>r</i> EF	Inclusion criteria: 5,010 pts, mean age 63 y, with NYHA class II-IV HF <i>r</i> EF	Intervention/Compar ator: 5,010 pts on standard therapy for HF were randomized to valsartan or placebo	Placebo (12%) p=0.004. <u>1° endpoint and results</u> : • At 23-mo follow-up, mortality was similar in pts treated with valsartan or placebo • The combined endpoint of mortality plus morbidity was reduced 13.2% (p=0.009) by valsartan because of a lower rate of HF hospitalization for HF (13.8% vs. 18.2%; p<0.001)	• Treatment with valsartan resulted in improvements in NYHA class, LVEF, signs and symptoms of HF, and quality of life compared with placebo (p<0.01).
SOLVD Investigators, 1991 (153) <u>2057034</u>	Aim: To determine the effect of enalapril vs. placebo on mortality and on mortality plus	Inclusion criteria: 2,569 pts, mean age 61 y, with HF <i>r</i> EF (90% with NYHA class II and III HF)	Intervention/Compar ator: 2,569 pts on standard therapy for	1° endpoint and results: At 41.4-mo follow-up, compared with placebo, enalapril	• At 41.4-mo follow-up, compared with placebo, enalapril reduced mortality by 16% (p=0.0036)

1002 (15.1)	hospitalization for HF in pts with HF <i>r</i> EF		HF were randomized to enalapril or placebo	reduced mortality or hospitalization for worsening HF by 26% (p<0.0001)	
1993 (154) <u>8104270</u>	<u>Aim</u> : To determine the effect of ramipril vs. placebo on mortality in pts with HF <i>r</i> EF	Inclusion criteria: 2,006 pts, mean age 65 y, with HFrEF after MI and without NYHA class0HF	Intervention/Compar ator: 2,006 pts were randomized to ramipril or placebo	<u>1° endpoint and results</u> : At 15-mo mean follow-up, compared with placebo, ramipril reduced all-cause mortality 27% (p=0.002)	• Analysis of prespecified 2° outcomes showed that ramipril reduced the first validated outcome (death, severe/resistant HF, MI, or stroke) by 19% (p=0.008).
Garg R, et al., 1995 (155) <u>7654275</u>	<u>Aim</u> : A meta-analysis was performed to determine the effect of ACEIs vs. placebo on mortality and on mortality plus hospitalization for HF in pts with HF <i>r</i> EF	Inclusion criteria: The meta- analysis included 32 trials of 7,105 pts with HF <i>r</i> EF treated with ACEIs vs. placebo	Intervention/Compar ator: In 25 trials, pts were treated with digoxin and/or diuretics, 4 trials only used diuretics, 1 trial used only digoxin, and 2 trials used no background therapy	<u>1° endpoint and results</u> : Compared with placebo, ACEIs reduced all-cause mortality 23% (p<0.001) and all-cause mortality or hospitalization for HF 35% (p<0.001).	• The reduction in mortality was primarily due to a 31% (17%–42%) reduction in death from progressive HF.
Pfeffer MA, et al., 2003 (156) <u>14610160</u>	<u>Aim</u> : To determine the effect of valsartan, captopril, or both on mortality in pts with MI complicated by HF, LV dysfunction, or both	Inclusion criteria: 14,703 pts, mean age 65 y, with MI complicated by HF, LV dysfunction, or both	Intervention: 4,909 pts were randomized to valsartan, 4,909 pts were randomized to captopril <u>Comparator</u> : 4,885 pts were randomized to valsartan plus captopril.	<u>1° endpoint and results</u> : At 24.7-mo median follow-up, mortality was similar in the 3 treatment groups.	• The incidence of adverse events causing discontinuation of drug was 5.8% with valsartan, 7.7% with captopril, and 9.0 % with valsartan plus captopril (p<0.05 comparing valsartan with captopril and valsartan plus captopril with captopril).
Maggioni AP, et al., 2002 (157) <u>12392830</u>	Aim: A subgroup analysis of the Val- HeFT study was performed to determine the effect of valsartan vs. placebo on mortality and on mortality plus morbidity in pts with HF <i>r</i> EF not receiving ACEIs	Inclusion criteria: 366 pts, mean age 67 y, with HF <i>r</i> EF not receiving ACEIs	Intervention/Compar ator: 185 pts were randomized to valsartan and 181 pts were randomized to placebo	<u>1° endpoint and results</u> : Compared with placebo, valsartan reduced mortality 33% (p=0.017) and mortality plus morbidity 44% (p<0.001).	• Compared with placebo, valsartan reduced first hospital admission for HF 53% (p=0.0006).

Granger CB, et al., 2003 (158) <u>13678870</u>	<u>Aim</u> : To determine the effect of candesartan vs. placebo on mortality in pts with HF <i>r</i> EF intolerant to ACEIs	Inclusion criteria: 2,028 pts, mean age 67 y, with HF <i>r</i> EF intolerant to ACEIs	Intervention/Compar ator: 1,013 pts were randomized to candesartan and 1,015 pts were randomized to placebo	<u>1° endpoint and results</u> : At 33.7-mo median follow-up, compared with placebo, the 1° endpoint of CV death or hospital admission for HF was reduced 30% by candesartan (p<0.0001).	• Compared with placebo, candesartan reduced CV death, hospital admission for HF, MI, stroke, or coronary revascularization 24% (p<0.0001).
Pitt B, et al., 2003 (159) <u>12668699</u>	<u>Aim</u> : To determine the effect of eplerenone vs. placebo on mortality and on CV death or hospitalization for CV events in pts with MI complicated by HF <i>r</i> EF	Inclusion criteria: 6,632 pts, mean age 64 y, with HF <i>r</i> EF after MI	Intervention/Compar ator: 3,313 pts were randomized to eplerenone and 3,319 pts were randomized to placebo	<u>1° endpoint and results:</u> At 16-mo mean follow-up, eplerenone reduced mortality 15% (p=0.008) and CV death or hospitalization for CV events 17% (p=0.005).	• Compared with placebo, eplerenone reduced death from any cause or any hospitalization 8% (p=0.02) and sudden cardiac death 21% (p=0.03), reduced hypokalemia from 13.1% to 8.4% (p<0.001), and increased serious hyperkalemia from 3.9%–5.5% (p=0.002).
Taylor AL, et al., 2004 (160) <u>15533851</u>	<u>Aim</u> : To determine the effect of ISDN plus hydralazine vs. placebo on mortality, first hospitalization for HF, and change in quality of life in black pts with HF <i>r</i> EF	Inclusion criteria: 1,050 African American pts, mean age 57 y, with HF <i>r</i> EF and NYHA class III or IV HF.	Intervention/Compar ator: 518 pts were randomized to ISDN plus hydralazine and 532 pts were randomized to placebo	<u>1° endpoint and results:</u> At 10-mo mean follow-up, compared with placebo, the mean 1° endpoint of mortality, first hospitalization for HF, and change in quality of life was reduced by ISDN plus hydralazine (p=0.01).	 Compared with placebo, ISDN plus hydralazine reduced mortality from 10.2%–6.2% (p=0.02) causing cessation of the study. Compared with placebo, ISDN plus hydralazine reduced all-cause mortality 43% (first hospitalization for HF 33% (p=0.001), and improved quality of life (p=0.02).
The Multicenter Dilitiazem Postinfarction Research Group, 1988 (145) <u>2899840</u>	Aim: To assess dilitiazem effect on recurrent infarction and death after acute MI <u>Study type</u> : RCT <u>Size</u> : 2,466 pts	Inclusion criteria: 25–75 y admitted to CCU, MI with enzyme confirmation. Exclusion criteria: Cardiogenic shock, symptomatic hypotension, PH with right HF, 2nd/3rd degree heart block, HR <50 bpm, contraceptives, WPW syndrome, CCBs, severe comorbidities or cardiac surgery	Intervention: Dilitiazem 240 mg (1,234) Comparator: Placebo (1,232)	 <u>1° endpoints and results</u>: Total mortality: identical in both groups Cardiac death and nonfatal MI: 11% fewer in dilitiazem but difference was NS 	No combined benefit from dilitiazem on mortality or cardiac events
ONTARGET Investigators, et al., 2008 (126) <u>18378520</u>	<u>Aim</u> : Evaluate whether use of an ARB was noninferior to ACEI, and whether the combination was	Inclusion criteria: • ≥55 y • Coronary, peripheral, or cerebrovascular disease or DM	Intervention: Ramipril 10 mg daily (n=8,576) Comparator:	<u>1° endpoint</u> : After a median follow-up of 56 mo, there was no difference between ramipril vs. telmisartan or combination therapy vs. ramipril in the 1°	• Telmisartan was equivalent to ramipril in pts with vascular disease or high-risk DM and was associated with less angioedema. The combination of the 2 drugs was associated with more

superior to ACE alone	with end-organ damage	• Telmisartan 80 mg	composite outcome of death	adverse events without an increase in
in the prevention of		daily (n=8,542)	from CV causes, MI, stroke,	benefit
vascular events in pts	Exclusion criteria:	 Combination of 	or hospitalization for HF (RR:	
with CVD or DM but	Inability to discontinue ACEI or	telmisartan and	1.01; 95% CI: 0.94–1.09 and	
not HF.	ARB	ramipril (n=8,502)	RR: 0.99; 95% CI: 0.92–1.07,	
	Known hypersensitivity or		respectively)	
Study type: Multi-	intolerance to ACEI or ARB			
center, double-blind,	Selected CVDs (congestive		Safety endpoint:	
RCT	HF, hemodynamically significant		Combination therapy was	
	valvular or outflow tract		associated with greater risk of	
Size: 25,620 pts	obstruction, constrictive		hyperkalemia than ramipril	
	pericarditis, complex congenital		monotherapy (480 pts vs. 283	
	heart disease, syncopal		pts; p<0.001)	
	episodes of unknown etiology <3		Hypotensive symptoms	
	mo, planned cardiac surgery or		were cited as reason for	
	PTCA <3 mo, uncontrolled HTN		permanent discontinuing more	
	on treatment [e.g., BP >160/100		in telmisartan vs. ramipril (RR:	
	mm Hg], heart transplant		1.54; p<0.001) and	
	recipient, stroke due to		combination therapy vs.	
	subarachnoid hemorrhage)		ramipril monotherapy (RR:	
	Other conditions (significant		2.75; p<0.001)	
	renal artery disease, hepatic		Renal impairment was more	
	dysfunction, uncorrected volume		common in combination	
	or sodium depletion, 1°		therapy vs. ramipril	
	hyperaldosteronism, hereditary		monotherapy (RR: 1.33; 95%	
	fructose intolerance, other major		CI: 1.22–1.44)	
	noncardiac illness or expected to			
	reduce life expectancy or			
	significant disability interfere with			
	study participation,			
	simultaneously taking another			
	experimental drug, unable to			
	provide written informed			
	consent).			

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
TOPCAT Pfeffer MA, et al., 2015 (161) 25406305	Aim: To investigate variation in pts and outcome in TOPCAT between pts from the Americas vs. Russia/Georgia Study type: Post- hoc analysis of prospective, double- blind, RCT Size: 3,445 pts	Inclusion criteria: NYHA class II–IV HF with LVEF ≤40% within 3 mo of enrollment; supine resting heart rate ≥68 bpm; stable clinical condition Exclusion criteria: Acute MI or UA within 28 d of randomization; indication or contraindication for treatment with BBs or drugs with beta- blocking properties; poor compliance; CABG surgery or PTCA in past 4 mo	Intervention: • Americas 886 on spironolactone • Russia/Georgia 836 on spironolactone • Spironolactone 15–45 mg daily <u>Comparator</u> : • Americas 881 on placebo • Russia/Georgia 842 on placebo • Placebo	 <u>1º endpoint</u>: Composite of CV death, aborted cardiac arrest, or HF hospitalization at 3.3 y follow-up was: Americas: 27.3% for spironolactone and 31.8% for placebo HR: 0.82; 95% CI: 0.69– 0.98; p=0.026; Russia/Georgia 9.3% for spironolactone and 8.4% for placebo HR: 1.10; 95% CI: 0.79– 1.51; p=0.58 <u>1º Safety endpoint:</u> Doubling of serum creatinine: Americas: 17.8% for spironolactone and 11.6% for placebo HR: 1.60; 95% CI: 1.25–2.05; p<0.001 Russia/Georgia 2.0% for S and 2.1% for p HR: 0.95; 95% CI: 0.49– 1.85; p=0.89 Creatinine >3.0 mg/dL Americas 9.8% for spironolactone and 9.1% for placebo HR: 1.10; 95% CI: 0.81–1.49; p=0.55 Russia/Georgia 0.2% for spironolactone and 0.4% for placebo HR: 0.5; 95% CI: 0.09–2.75; p=0.43 Hyperkalemia (potassium >5.5 mmol/L) Americas 25.2% for spironolactone and 8.9% for placebo OR: 3.46; 95% CI: 2.62–4.56; p<0.001 	Relevant 2° endpoint: CV mortality: Americas 10.8% for spironolactone and 14.4% for placebo HR: 0.74; 95% Cl 0.57–0.97; p=0.027; Russia/Georgia 7.7% for spironolactone and 5.8% for placebo HR: 1.31; 95% Cl: 0.91–1.90; p=0.15. Aborted cardiac arrest: NS between groups. HF hospitalization: 20.8% for spironolactone and 24.5% for placebo HR: 0.82; 95% Cl: 0.67–0.99; p=0.042; Russia/Georgia 2.6% for spironolactone and 3.4% for placebo HR: 0.76; 95% Cl: 0.44–1.32; p=0.327; Recurrent HF: 361 events for spironolactone and 438 events for placebo (IRR: 0.75; 95% Cl: 0.58– 0.96; p=0.024) Russia/Georgia 33 events for spironolactone and 37 events for placebo (IRR: 0.83; 95% Cl: 0.42–1.62; p=0.58) All-cause mortality: NS between groups in Americas and Russia/Georgia. All-cause hospitalization: NS between groups in Americas and Russia/Georgia. MI: NS between groups; Stroke: NS between groupsStudy limitations and adverse events: The pts enrolled in Russia/Georgia in the TOPCAT trial did not demonstrate either the expected morbidity and mortality associated with symptomatic HF or

Data Supplement 35. RCTs Comparing HF*p*EF (Section 9.2.2)

Aronow WS, et	<u>Aim</u> : To determine	Inclusion criteria:	Intervention: 79 pts were	 Russia/Georgia 11.8% for spironolactone and 9.4% for placebo OR: 1.30; 95% CI: 0.95–1.77; p=0.10 Hypokalemia (potassium <3.5 mmol/L) Americas 15.2% for spironolactone and 26.2% for placebo) 0.51 (95% CI: 0.40–0.64; p<0.001) Russia/Georgia 17.2% for S and 19.4% for p OR: 0.87 (95% CI: 0.68–1.11; p=0.26) 	most pharmacological responses to spironolactone <u>Summary</u> : In pts with HF with preserved EF, spironolactone reduced the 1° endpoint of composite of CV death, aborted cardiac arrest, or HF hospitalization in the Americas group but not in the Russia/Georgia group. The pts enrolled in the Russia/Georgia group did not demonstrate either the expected morbidity and mortality associated with symptomatic HF with preserved EF or most pharmacological responses to spironolactone <u>Relevant 2° endpoint:</u> At 1-y follow-
al., 1997 (162) <u>9230162</u>	effect of propranolol vs. no propranolol on mortality plus nonfatal MI in pts with prior MI and HF <i>p</i> EF	Pts ≥62 y with MI and LVEF ≥40% and HF NYHA class II or III treated with diuretics and ACEIs for 2 mo	randomized to treatment with propranolol <u>Comparator</u> : 79 pts were randomized to no propranolol. • All pts continued diuretic and ACEI therapy.	up, multivariate Cox regression analysis showed that compared with no propranolol, propranolol reduced mortality 35% (p=0.03) and mortality plus nonfatal MI 37% (p=0.018)	up, LVEF was increased by propranolol from 57% to 63% (p<0.001) and LV mass was decreased by propranolol from 312 grams to 278 grams (p=0.001) Propranolol was stopped because of adverse effects in 11 of 79 pts (14%)
Kostis JB, et al., 1997 (163) <u>9218667</u>	<u>Aim</u> : To determine the effect of antihypertensive drug therapy vs. placebo in prevention of HF in pts with isolated systolic HTN	Inclusion criteria: Pts ≥60 y with isolated systolic HTN in the SHEP program	Intervention/Comparator: 4,736 pts were randomized to antihypertensive drug therapy or placebo	<u>1° endpoint</u> : At 4.5-y follow-up, fatal or nonfatal HF was reduced 49% (p<0.001) by antihypertensive drug therapy (NNT to prevent 1 event =48)	<u>Relevant 2° endpoint:</u> CV mortality and nonfatal hospitalized HF was reduced 30% (p=0.002) by antihypertensive drug therapy
Beckett NS, et al., 2008 (164) <u>18378519</u>	<u>Aim</u> : To determine the effect of antihypertensive drug therapy on fatal or nonfatal stroke in pts ≥80 y	Inclusion criteria: Pts ≥80 y with a SBP≥160 mm Hg	Intervention/Comparator: 3,845 pts were randomized to antihypertensive drug therapy or placebo	<u>1° endpoint</u> : The 1° endpoint of fatal or nonfatal stroke was reduced 30% (p=0.06) by antihypertensive drug therapy	Relevant 2° endpoint: Antihypertensive drug therapy reduced HF 64% (p<0.001) all-cause mortality 21% (p=0.02), and CV death 23% (p=0.06)

Van Veldhuisen DJ, et al., 2009 (165) <u>19497441</u>	<u>Aim</u> : To determine the effect of nebivolol vs. placebo in pts with HF <i>r</i> EF and HF <i>p</i> EF	Inclusion criteria: Pts ≥70 y, history of HF, and HF <i>r</i> EF or HF <i>p</i> EF	Intervention/Comparator: 1,359 pts with a history of HF <i>r</i> EF and 752 pts with a history of HF <i>p</i> EF were randomized to nebivolol or to placebo	<u>1° endpoint</u> : At 21-mo follow-up, the 1° endpoint of all-cause mortality or CV hospitalization was reduced by nebivolol 14% (95% CI: 0.72– 1.04) in pts with HF <i>r</i> EF and 19% (95% CI: 0.63, 1.04) in pts with HF <i>p</i> EF	Relevant 2° endpoint: HR for reduction of all-cause mortality by nebivolol: 0.84 (95% CI: 0.86–1.08) for HF <i>r</i> EF and 0.91 (95% CI: 0.62–1.33) for HF <i>p</i> EF
Yusef S, et al., 2003 (166) <u>13678871</u>	<u>Aim</u> : To determine the effects of candesartan vs. placebo in pts with HF <i>p</i> EF	Inclusion criteria: 3,032 pts, mean age 67 y, with HF <i>p</i> EF and NYHA class II-IV HF	Intervention/Comparator: 3,032 pts were randomized to candesartan or placebo	<u>1° endpoint</u> : At 36.6 m follow-up, the 1° outcome of CV death or hospitalization for HF was reduced 11% (p=0.118) by candesartan	Relevant 2° endpoint: Hospitalization was reduced 16% (p=0.047) by candesartan
Massie BM, et al., 2008 (167) <u>19001508</u>	<u>Aim</u> : To determine the effect of irbesartan vs. placebo on all- cause mortality or hospitalization for a CV cause in pts with HF <i>p</i> EF	Inclusion criteria: Pts 60 y and older with HF <i>p</i> EF and NYHA class II, III, or IV HF	Intervention/Comparator 4,128 pts were randomized to irbesartan or placebo	<u>1° endpoint</u> : At 49.5-mo follow-up, the 1° outcome of all-cause mortality or hospitalization for CV cause was reduced 5% by irbesartan (p=0.35)	<u>Relevant 2° endpoint:</u> Irbesartan did not significantly reduce the 2° outcomes of death from HF or hospitalization for HF, death from any cause and from CV causes, and quality of life
Piller LB, et al., 2011 (168) <u>21969009</u>	<u>Aim</u> : To determine mortality rates in pts who developed HF in ALLHAT	Inclusion criteria: 1,761 pts, mean age 70 y, developed HF during ALLHAT	Intervention/Comparator At 8.9-y mean follow-up, 1,348 of 1,761 pts (77%) with HF died	<u>1° endpoint</u> : Post-HF all-cause mortality was similar for pts treated with chlorthalidone, amlodipine, and lisinopril. 10-y adjusted rates for mortality were 86% for amlodipine, 87% for lisinopril, and 83% for chlorthalidone	<u>Relevant 2° endpoint:</u> All-cause mortality rates were similar for those with HF <i>r</i> EF (84%) and for those with HF <i>p</i> EF (81%) with no significant differences by randomized treatment arm
Law MR, et al., 2009 (18) <u>19454737</u>	Study type: Meta- analysis of use of BP-lowering drugs in prevention of CVD from 147 randomized trials Size: Of 147 randomized trials of 464,000 pts, 37 trials of BBs in CAD included 38,892 pts, and 37 trials of	Inclusion criteria: The database search used Medline (1966- Dec. 2007 in any language) to identify randomized trials of BP-lowering drugs in which CAD events or strokes were recorded. The search also included the Cochrane Collaboration and	<u>1° endpoint</u> : CAD events; stroke <u>Results:</u> In 37 trials of pts with a history of CAD, BBs reduced CAD events 29% (95% CI: 22%–34%). In 27 trials in which BBs were used after acute MI, BBs reduced CAD events 31% (95% CI: 24%–38%), and in 11 trials in which BBs were used after long-term CAD, BBs	• With the exception of the extra protective effect of BBs given shortly after a MI and the minor additional effect of CCBs in preventing stroke, all the classes of BP-lowering drugs have a similar effect in reducing CAD events and stroke for a given reduction in BP.	N/A

drugs	vpertensive in CAD led 85,395 pts databases and the citations in trials and previous meta- analyses and review articles.	atabases and the itations in trials and revious meta- nalyses and reviewevents 13%. In 7 trials, B reduced stroke 17% (95% 1%–30%). CAD events v reduced 14% (95% CI: 2 25%) in 11 trials of thiazi diuretics, 17% (95% CI:	Bs 6 CI: rere %– de	
	5	rticles. 25%) in 11 trials of thiazi diuretics, 17% (95% CI: 11%–22%) in 21 trials of ACEIs, insignificantly 14%	de	
	events and strokes or if treatment duration was <6 mo.	r if treatment (95% CI: 8%–22%) in 22	s 3%- de %-	
		and 34% (95% CI: 25%– 42%) in 9 trials of CCBs.		

Data Supplement 36. Nonrandomized Trials, Observational Studies, and/or Registries of HFpEF (Section 9.2.2)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Law MR, et al.,	Study type: Meta-	Inclusion criteria: The database	1° endpoint: CAD events; stroke	 With the exception of
2009 (18)	analysis of use of BP-	search used Medline (1966–Dec.		the extra protective
<u>19454737</u>	lowering drugs in	2007 in any language) to identify	Results: In 37 trials of pts with a history of CAD, BBs reduced	effect of BBs given
	prevention of CVD from	randomized trials of BP-lowering	CAD events 29% (95% CI: 22%, 34%). In 27 trials in which BBs	shortly after a MI and
	147 randomized trials	drugs in which CAD events or	were used after acute MI, BBs reduced CAD events 31% (95%	the minor additional
		strokes were recorded. The	CI: 24%, 38%), and in 11 trials in which BBs were used after	effect of CCBs in
	Size: Of 147 randomized	search also included the	long-term CAD, BBs insignificantly reduced CAD events 13%.	preventing stroke, all
	trials of 464,000 pts, 37	Cochrane Collaboration and Web	In 7 trials, BBs reduced stroke 17% (95% CI: 1%–30%). CAD	the classes of BP-
	trials of BBs in CAD	of Science databases and the	events were reduced 14% (95% CI: 2%–25%) in 11 trials of	lowering drugs have a
	included 38,892 pts, and	citations in trials and previous	thiazide diuretics, 17% (95% CI: 11%–22%) in 21 trials of	similar effect in
	37 trials of other	meta-analyses and review	ACEIs, insignificantly 14% in 4 trials of angiotensin receptor	reducing CAD events
	antihypertensive drugs in	articles.	blockers, and 15% (95% CI: 8%–22%) in 22 trials of CCBs.	and stroke for a given
	CAD included 85,395 pts		Stroke was reduced 38% (95% CI: 28%–47%) in 10 trials of	reduction in BP.

	thiazide diuretics, 22% (95% CI: 8%–34%) in 13 trials of ACEIs, and 34% (95% CI: 25%–42%) in 9 trials of CCBs.	
duration was <6 mo.		

Data Supplement 37. RCTs Comparing CKD (Section 9.3)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
MDRD Klahr S, et al., 1994 (169) <u>8114857</u>	Aim: To determine whether restricted protein intake or tighter HTN control would delay progression of CKD Study type: Randomized management to low or usual BP goal and usual, low or very low protein intake Size: • Total n=840 Study 1 n=585 Study 2 n=255 • Mean follow-up 2.2 y • Mean MAP, mm Hg (SD): Study 1: 98 (11) Study 2: 98 (11) • Mean SBP, mm Hg (SD): Study 1: 131 (18) Study 2: 133 (18)	Inclusion criteria: Adults 18–70 y, with renal insufficiency (serum Cr 1.2–7.0 mg/dL in women and 1.4–7.0 mg/dL in men or CrCl <70 mL/min per 1.73 m ²) and MAP≤125 mm Hg (normotensives included) Exclusion criteria: Pregnancy, body weight <80% or >160% of standard, DM requiring insulin, urine protein >10 g/d, history of renal transplant, chronic medical conditions, doubts regarding compliance.	Intervention:• Study 1 includedsubjects with GFR 25–55mL/min 1.73 m² (n=585);• Study 2 includedsubjects with GFR 13–24mL/min 1.73 m² (n=255)• Low MAP goal ≤92 mmHg for those 18–60 y;≤98 for those ≥61 y• Usual: MAP goal ≤107m Hg for those 18–60;MAP ≤113 for subjects≥61• 2 studies:Study 1: above BP goalsplus usual or low proteindiet (1.3 or 0.58 g proteinper kg of body weight/d)Study 2: above BP goalsplus low or very lowprotein diet (0.58 or 0.28g per kg/d)• Between groupdifference in MAP, mmHg 4.7; p<0.001	1° endpoint: Rate of decline in GFR, mL/min (95% Cl) • Study 1 From baseline to 4 mo Low: 3.4; 95% Cl: 2.6–4.1 Usual: 1.9; 95% Cl: 1.1–2.7 p=0.010 4 mo to study end, Low: 2.8; 95% Cl: 2.2–3.3 Usual: 3.9; 95% Cl: 3.3–4.5 p=0.006 Baseline to 3 y, Low: 10.7; 95% Cl: 9.1–12.4 Usual: 12.3; 95% Cl: 9.1–12.4 Usual: 12.3; 95% Cl: 10.6–14.0 p=0.18 • Study 2 From baseline to end of study, Low: 3.7; 95% Cl: 3.1–4.3 Usual: 4.2; 95% Cl: 3.6–4.9 p=0.28 ESRD or death: • Study 2 RR for low vs. usual: 0.85; 95% Cl: 0.60– 1.22 p=NR	 Limitations: Drug therapy was not randomized. Recommended ACEI ± diuretic then CCB and others. More subjects in the low BP goal groups received ACEIs (48%, 51% also reported elsewhere) compared to the usual BP goal group (28%, 32% also reported e/w) (not noted in 1° manuscript but reported in Peterson JC, et al., 1995 (170)). 1.9% study 1, 1.2% study 2 lost to follow-up. Rate of GFR decline was slower than expected in the control groups and was not constant. Summary: No significant benefits overall from either low protein or lower BP target. There was a significant interaction between baseline urinary protein excretion and BP interventions (p=0.01) indicating that low BP was of benefit to subjects with >1 g proteinuria with slower progression of loss of GFR

REIN-2	Mean DBP, mm Hg (SD): Study 1: 81 (10) Study 2: 81 (10)	Inclusion criteria:	<u>Comparator</u> : By BP and protein intake goals Intervention:		Limitations: The study was
REIN-2 Ruggeneti P, et al., 2005 (171) <u>15766995</u>	Aim: To determine whether intensive BP control will achieve further renoprotection (delayed progression to ESRD) compared to standard BP control in pts with chronic nephropathies Study type: Multicenter RCT of pts all placed on ACEI (ramipril) at maximum dose tolerated to achieve DBP <90 then assigned to conventional or intensified BP control. Add-on drug was dihydropyridine felodipine 5–10 mg/d Size: 335 (median time 19 mo)	 Adults, age 18–70 y, with nondiabetic nephropathy, persistent proteinuria (urinary protein excretion >1 g/24 h for ≥3 mo) and not on ACEIs in previous 6 wk Pts with proteinuria 1–3 g/24 h included if CrCI <70 mL/min/1.73 m² For overall population, mean SBP, mm Hg (SD): Intensive: 137.0 (16.7) Conventional: 136.4 (17.0) For overall population, mean DBP, mm Hg (SD): Intensive: 84.3 (9.0) Conventional: 83.9 (10.4) Exclusion criteria: Urinary tract infection, CHF class III–IV, treatment with corticosteroids, NSAIDs, immunosuppression, acute MI or stroke in prior 6 mo, severe uncontrolled HTN, 	 Intervention: Intensive: BP goal <130/80 mm Hg Conventional: DBP goal <90 mm Hg, irrespective of SBP For baseline proteinuria subgroups, result BP values NR For the overall population, achieved BP, mm Hg (SD) Intensive: 129.6/79.5 (10.9/5.3) Conventional: 133.7/82.3 (12.6/7.1) p=0.0019/<0.0001 For the overall population, change in BP, mm Hg Intensive: -7.4/-4.8 Conventional: -2.7/-1.6 p=NR For the overall population, BP difference between groups, mm Hg 4.1/2.8 p=NR 	1° endpoint• Time to ESRD; over 36 mo follow-up, median 19 mo1° outcome: ESRD in pts with baseline proteinuria 1–3 g/24 h HR (95% CI): 1.06 (95% CI: 0.51–2.20) p=0.89• ESRD in pts with baseline proteinuria >3 g/24 h HR (95% CI): 1.09 (95% CI: 0.55–2.19) p=0.81• 23% of intensive and 20% of conventional control groups progressed to ESRD.• Median rate of GFR decline, mL/min/1.73 m²/mo (IQR) in pts with baseline proteinuria <3 g/24: Intensive: 0.18 (95% CI: 0.03–0.49) Conventional: 0.21 (95% CI: -0.03–0.40) p=0.89• Median rate of GFR decline, mL/min/1.73 m/mo (IQR) in pts with baseline proteinuria ≥3 g/24: Intensive: 0.51; 95% CI: 0.16–1.05 Conventional: 0.39; 95% CI: 0.030.98 p=0.39	Limitations: The study was stopped at the 1 st interim analysis for futility. Median time 19 mo Summary: In pts with non-DM proteinuric nephropathies receiving background ACEI therapy, no additional benefits from further BP reduction by felodipine could be shown. Dihydropyridine CCBs do not offer additional renoprotection to ACEIs or ARBs.

		suspicion for	Comparator: By BP		
		renovascular disease,	goals		
		obstructive uropathy,	5		
		DM-1, collagen vascular			
		disease, cancer, elevated			
		aspartate transaminase,			
		chronic cough, history of			
		allergy or poor tolerance			
		to study meds, alcohol			
		abuse, pregnancy,			
		breastfeeding, ineffective			
		contraception.			
AASK	Aim: To compare the	Inclusion criteria:	Intervention:	<u>1° endpoint:</u>	Limitations:
Wright JT, et al.,	effects of 2 levels of	Adult African-	• Low: MAP goal ≤92	• 1° outcome: difference in mean slopes,	Based on DSMD
2002 (172)	BP and 3	Americans, 18–70 y, with	mm Hg	acute GFR slope, mL/min/1.73 m ² /3 mo	recommendation, amlodipine arm
<u>12435255</u>	antihypertensive drug classes on GFR	HTN (DBP ≥95) and	Usual: MAP goal 102-	(SE):	halted early and those pts
	decline in HTN	GFR of 20–65	107 mm Hg	• 1.82 (0.54) in low BP group	switched to open label Rx,
		mL/min/1.73 m ² , no DM	 Initial treatment with a 	p<0.001	continued study schedule and
	Study type:	• At entry: mean MAP,	B Blocker (metoprolol),	• 1° outcome: difference in mean slopes,	same BP goals
	Randomized 3×2	mm Hg:	and ACEI (ramipril) or a	chronic GFR slope, mL/min/1.73 m ² /y	<u>j</u>
	factorial trial	Low: 115 (27)	dihydropyridine	(SE): 0.21 (0.22)	Summary:
	Measured GFR with	Usual: 113 (15)	(amlodipine) with open	p=0.33 NS	No difference in GFR decline
	iothalamate	 Mean SBP, mm Hg 	label agents added to	• Difference in mean slopes, total GFR	with lower BP goal and no
	Iotrialamate	(SD):	achieve BP goals	slope, mL/min/1.73 m^2/y (SE):	difference in composite clinical
	Size: 1,094	Low:152 (25)	 Study duration: 	-0.25 (0.22)	endpoints
	<u>5120</u> . 1,074	Usual: 149 (23)	3–6.4 у	p=0.24	Average rate of GFR decline 2
		Mean DBP, mm Hg:	BP similar across drug	Main 2° clinical composite outcome:	mL/min/y is similar or slower than
		Low: 96 (15)	groups except 2 mm Hg lower in amlodipine	GFR event, ESRD, or death,	previous reports
		Usual: 95 (14)	group	% risk reduction (95% CI): 2 (95% CI: -	• There was a trend favoring the
		. ,	Mean from 3 mo to	22–21)	lower BP goal in subjects with
		Exclusion criteria:	study end	p=0.85	higher baseline proteinuria and
		DBP<95, history of DM,	• MAP, mm Hg (SD)	• GFR event or ESRD,	the opposite trend for those
		Urinary protein/creatinine	• MAP, IIIII Hy (SD) Low: 95.8 (8)	% Risk Reduction: -2; 95% CI: -31–20;	without proteinuria
		ratio >2.5, accelerated or	Usual: 104 (7)	p=0.87	Ramipril treatment group had
		malignant HTN, non-BP	• SBP/DBP, mm Hg (SD)	• ESRD or death,	slower progression compared
		related cause of CKD,	• SBP/DBP, IIIII Hg (SD) Low: 128/78 (12/8)	% risk reduction: 12; 95% CI: -13–32;	with metoprolol and amlodipine
		serious systemic disease,	Usual: 141/85 (12/7)	p=0.31	combined, less evident between
		clinical CHF, specific	· · ·	• ESRD alone,	ramipril and metoprolol
		indication or	MAP change, mm Hg	% risk reduction: 6; 95% CI: -29–31;	
		contraindication for a	Low: -20	p=0.72	
L				p-0.72	

		study drug or procedure	Usual: -9 • SBP/DBP change, mm Hg Low: -24/-8 Usual: -18/-10 • Achieved mean BP difference between groups, mm Hg MAP: 11 SBP: 16 DBP: 8 <u>Comparator</u> : N/A	 2° outcome: urine protein excretion <u>Safety endpoint</u>: Acute and chronic rate of change in GFR (slope): NS for chronic and total slope in subgroup analyses by baseline proteinuria strata Acute slope: p=0.08 for interaction Total slope: p=0.04 for interaction Chronic slope: p=0.16 for interaction Clinical composite outcome: includes reduction in GFR by 50% or by 25 mL/min/m², ESRD, death, NS in subgroup analyses by baseline proteinuria strata; p=0.007 for interaction For above outcomes, trends favored the lower BP goal over the usual goal in participants with higher baseline proteinuria and opposite trends in participants with little or no proteinuria Within each drug group, risk reductions for any 2° clinical outcome of the low vs. usual BP goal were not significantly different between pts with baseline urine protein to creatinine ratio ≤0.22 (p=NS) 	
Contreras G, et al., 2005 (173) <u>15897360</u>	Aim: Within AASK to examine the effect of BP intervention separately in the 3 drug treatment groups Study type: • Randomized 3×2 factorial trial • Measured GFR with iothalamate	Inclusion criteria: • Adult African Americans, ages 18–70, with HTN (DBP ≥95) and GFR of 20–65 mL/min/1.73 m ² , no DM Mean MAP, mm Hg: Low, Amlodipine: 115.3 (18.3) Usual, Amlodipine: 112.7 (14.7)	Intervention: • Analysis by initial drug treatment group • Low, Amlodipine: MAP goal ≤92 mm Hg, Amlodipine (5–10 mg/d) Usual, Amlodipine: MAP goal 102–107 mm Hg, Amlodipine (5–10 mg/d) • Low, Metoprolol: MAP goal ≤92 mm Hg, Matamatal (50, 200	 <u>1° endpoint</u>: GFR event, ESRD, or death prior to dialysis, Amlodipine, Low vs. Usual Goal RR: 32%; 95% Cl: -14–60; p=0.14 Metoprolol, Low vs. Usual Goal RR: 4%; 95% Cl: -39–33; p=0.84 Ramipril, Low vs. Usual Goal RR: -8%; 95% Cl: -93–15; p=0.24 p for interaction=0.17 GFR event or ESRD, Amlodipine, Low vs. Usual Goal RR: 26% (000000000000000000000000000000000000	Limitations: Post-hoc analysis, effects on GFR may have been obscured by early rise and later fall with amlodipine, follow-up only 3–6.4 y, many comparisons so risk for type I error, unable to test ACEI – DHP CCB combination. Summary: • BP effect was similar among
	<u>Size</u> : 1,094	Low, Metoprolol: 114.5 (17.5)	Metoprolol (50–200 mg/d)	26%; 95% CI: -33–58; p=0.32	drug groups for GFR slope and main clinical composite.

Llaval Matagnalal 110.4		Matamalal Lawren Havel Carl DD	
Usual, Metoprolol: 112.4	Usual, Metoprolol: MAP	Metoprolol, Low vs. Usual Goal RR:	BP effect differed among drug
(14.1)	goal 102–107 mm Hg,	7%; 95% CI: -42–39; p=0.74	groups for composite of ESRD or
Low, Ramipril: 115.2	Metoprolol (50–200	Ramipril, Low vs. Usual Goal RR:	death and ESRD alone.
(15.2)	mg/d)	-42%; 95% CI: -126–11; p=0.14	 Higher event rates for
Usual, Ramipril: 114.0	 Low, Ramipril: MAP 	p for interaction=0.20	amlodipine and usual BP goal
(16.7)	goal ≤92 mm Hg,	 ESRD or death prior to dialysis, 	compared with other groups.
 Mean SBP, mm Hg: 	Ramipril (2.5–10 mg/d)	Amlodipine, Low vs. Usual Goal RR:	 Low BP goal associated with
Low, Amlodipine: 152.2	Usual, Ramipril: MAP	51%; 95% CI: 13–73; p=0.016	reduced risk of ESRD or death
(28.2)	goal 102–107 mm Hg,	 Metoprolol, Low vs. Usual Goal RR: 	and ESRD for amlodipine but not
Usual, Amlodipine: 147.7	Ramipril (2.5–10 mg/d)	11%; 95% CI: -40–44; p=0.61	for other drug groups (in the
(21.9)	 Note: Amlodipine arms 	Ramipril, Low vs. Usual Goal RR:	absence of ACEI treatment).
Low, Metoprolol: 152.0	terminated 1 y early	-32%; 95% CI: -114–18; p=0.26	
(25.7)	Achieved MAP	p for interaction=0.035	
Usual, Metoprolol: 147.7	difference between	 Death alone (prior to dialysis), 	
(21.4)	groups, mm Hg	Amlodipine, Low vs. Usual Goal RR:	
Low, Ramipril: 151.0	Amlodipine, Low vs.	48%; 95% CI: -59–83; p=0.25	
(22.5)	Usual:12.89	• Metoprolol, Low vs. Usual Goal RR: -1;	
Usual, Ramipril: 150.9	Metoprolol, Low vs.	95% CI: -110-5; p=0.97	
(24.1)	Usual: 11.11	Ramipril, Low vs. Usual Goal RR:	
Mean DBP, mm Hg:	Ramipril, Low vs. Usual:	21%; 95% CI: -92–67; p=0.61; p for	
Low, Amlodipine: 96.55	10.12	interaction=0.61	
(15.1)	p=NR		
Usual, Amlodipine: 94.87	 Achieved SBP 	Safety endpoint:	
(12.9)	difference between	• ESRD alone, Amlodipine, Low vs.	
Low, Metoprolol: 95.45	groups, mm Hg	Usual Goal: RR: 54%; 95% CI: 8–77;	
(15.4)	Amlodipine, Low vs.	p=0.028	
Usual, Metoprolol: 94.47	Usual: 18.4	Metoprolol, Low vs. Usual Goal RR:	
(12.5)	Metoprolol, Low vs.	11%; 95% CI: -60–50; p=0.70	
Low, Ramipril: 96.90	Usual: 15.4	• Ramipril, Low vs. Usual Goal RR:	
(13.6)	Ramipril, Low vs. Usual:	-65%; 95% CI: -195–8; p=0.09; p for	
Usual, Ramipril: 95.12	12.6	interaction=0.021	
(15.3)	p=NR	 Death alone (prior to dialysis), 	
(10.0)		Amlodipine, Low vs. Usual Goal: RR:	
Exclusion criteria:	 Achieved DBP 	48%; 95% CI: -59–83; p=0.25	
DBP<95, history of DM,	difference between	 Metoprolol, Low vs. Usual Goal: RR: - 	
Urinary protein/creatinine	groups, mm Hg	■ Metoproloi, Low Vs. Osuai Goal. RR 1; 95% CI: -110–5; p=0.97	
ratio >2.5, accelerated or	Amlodipine, Low vs.		
malignant HTN, non-BP	Usual: 10.14	• Ramipril, Low vs. Usual Goal RR:	
related cause of CKD,	Metoprolol, Low vs.	21%; 95% CI: -92–67; p=0.61; p for	
serious systemic	Usual: 8.86	interaction=0.61	
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		disease, clinical CHF,	Ramipril, Low vs. Usual:	 Proteinuria within each drug group, 	
		specific indication or	8.96	risk reductions for any 2° clinical	
		contraindication for a	p=NR	outcome of the low vs. usual BP goal	
		study drug or procedure		were not significantly different between	
			Comparator: N/A	pts with baseline urine protein to	
				creatinine ratio ≤0.22 and >0.22 (p=NS)	
Norris K, et al.,	Aim: Compared effect	Inclusion criteria:	Intervention:	1° endpoint:	Limitations:
2006 (174)	of treatment on CV	Adult African	 Achieved SBP/DBP, 	Number of deaths before ESRD, n of	 Limited power, only 202 CV
17059993	event rate during mean	Americans, 18–70 y, with	mm Hg (SD)	events	events – low incidence. CV
	follow-up of 4.1 y by	HTN (DBP ≥95) and	Low: 128/78	Low: 38	outcomes were 2° endpoints of
	drug class and level of	GFR of 20–65	Usual: 141/85	Usual: 47; p=NR	high priority (prespecified).
	BP control.	mL/min/1.73 m ² , no DM	p=NR	Major CAD events, n of events (rate	 >50% had a history of heart
	Determined baseline	Mean MAP, mm Hg:	• SBP/DBP change, mm	per person-y)	disease at entry, 40% with LVH
	factors that predict CV	114 (16)	Hg	Low: 19 (0.008)	by ECG. 1/3 smokers, almost
	outcomes	Mean SBP, mm Hg:	Low: -23/-19	Usual: 23 (0.010); p=NS	50% had income <15K.
	outcomes	150 (24)	Usual: -8/-9	• Stroke events, number of events (rate	
	Study type:	Mean DBP,	p=NR	per person-y)	Summary:
	Randomized 3×2	mm Hg: 96	Achieved mean BP	Low: 26 (0.011)	CV outcome rate was not
	factorial trial	(14)	difference between	Usual: 29 (0.013); p=NS	related to randomized
	Measured GFR with	(14)			
	iothalamate	Evolucion oritorio, N/A	groups, mm Hg SBP: 15	• HF events, n of events (rate per	interventions, either drug or BP
	IUIIIaiaiilaite	Exclusion criteria: N/A	DBP: 10	person-y)	target. ● 7 baseline risk factors were
	Sizo.			Low: 27 (0.012)	
	<u>Size</u> : 1,094		p=NR	Usual: 23 (0.010)	independently associated with
	1,094			p=NS	increased risk for CV composite
			Comparator: N/A	CV composite outcome, n of events	outcome in multivariable analyses
				(rate per person-y)	after controlling for age, sex,
				Low: 71 (0.032)	baseline GFR, baseline
				Usual: 78 (0.035); p=NS	proteinuria: PP, duration of HTN,
				 Composite outcome or ESRD, n of 	protein/creatinine ratio, urine
				events (rate per person-y)	sodium-potassium ratio and
				Low: 143 (0.064)	annual income <15,000.
				Usual: 159 (0.072)	
				p=NS	
				Overall rate of CV events, n of events	
				(rate per person-y)	
				Low: 108 (0.048)	
				Usual: 94 (0.042); p=NS	
				• CV death, n of events (rate per	
				person-y)	
				1 57	
				Low: 16 (0.007)	

				Usual: 15 (0.006); p=NS	
Amlodipine Versus	Aim:	Inclusion criteria:	Intervention:	1° endpoint: Change in GFR from	Summary:
Enalapril in Renal	To compare GFR	• 18–80 y	• Amlodipine: 5–10 mg/d	baseline to final assessment	 No difference in GFR change
Failure (AVER trial)	decline in nondiabetic,	• CrCl 20–60	 Enalapril: 5–20 mg/d 		or serum creatinine at trial end
Esnault VL, et al.,	nonnephrotic adults	mL/min/1.73 m ²	Therapy initiated with	<u>2° Outcome:</u> Clinical composite of renal	Last observation: mean change in
2008 (175)	with HTN and	(Cockcroft-Gault)	amlodipine 5 mg/d or	replacement therapy, discontinuation	GFR, mL/min/1.73 m ²
18405787	estimated CrCl 20–60	Nondiabetic	enalapril 5 mg/d. Drugs	due to deterioration of renal function,	Amlodipine -4.92, Enalapril -3.98;
	mL/min/1.73 m ² when	 Enrollment confirmed 	up-titrated to amlodipine	50% decrease in GFR, doubling of	p=NS
	randomized to a CCB	at end of 4-wk placebo	10 mg/d or enalapril 20	serum Cr, hospitalization for transient	• Last observation: mean change
	(amlodipine, 5–10	run-in if sitting DBP	mg/d at wk 8 and 12 if	renal failure. "Other 2º outcome	in Serum Cr from baseline (mg/d)
	mg/d) or an ACEI	between 90 and 119 mm	DBP >90 mm Hg.	measures" included: changes in serum	Amlodipine +0.57, Enalapril
	(enalapril, 5–20 mg/d).	Нд	After 18 wk, if maximal	Cr, sitting DBP and SBP, heart rate,	+0.47; p=NS
		 Mean SBP, mm Hg 	tolerated dose of study	total and HDL cholesterol, 24-h urinary	 No difference in composite 2°
	Study type: RCT	(SD):	drug did not decrease BP	protein excretion, ambulatory BP	endpoints.
		Amlodipine: 165.1 (15.4)	to target, add on anti-	monitoring, and safety measures.	 Mean BP (mm Hg): baseline to
	<u>Size</u> :	Enalapril: 165.2 (16.6)	HTN treatments were the	Composite Outcomes: 2º clinical	last observation
	Amlodipine: 132	 Mean DBP, mm Hg 	following: atenolol (50-	composite	Amlodipine 164.8/101.8 to
	Enalapril: 131	(SD):	100 mg/d), loop diuretics		140.1/85.4, delta -24.7/16.4
		Amlodipine: 102.0 (6.7)	(furosemide 20–500	Safety endpoint: Proteinuria subgroup,	Enalapril 165.0/102.5 to
		Enalapril: 102.5 (7.1)	mg/d or torsemide, 5–	>1 g/d: protein excretion rate decreased	140.3/86.4, delta -24.7/16.1
		 Mean serum Cr, mg/dL 	200 mg/d), alpha	significantly in pts taking enalapril plus	
		(SD):	blockers (prazosin, 2.4–5	diuretic (median -270 mg/d; p<0.001)	
		Amlodipine: 2.00 (0.8)	mg/d or doxazosin, 1–16	but not in pts taking amlodipine plus	
		Enalapril: 2.05 (0.7)	mg/d) and centrally	diuretic (-25 mg/d) at last obs	
			acting drugs (rilmenidine		
		Exclusion criteria:	(1–2 mg/d or		
		 Nephrotic proteinuria 	methyldopa, 250–500		
		• 2° or malignant HTN	mg/d).		
		(DBP >120 mm Hg)			
		 A major CV event 	 BP goal: 		
		within 3 mo	Amlodipine: <130/85 mm		
		 Angina pectoris 	Hg		
		Congestive heart	Enalapril: <130/85 mm		
		disease (NYHA II-IV)	Hg		
		Uncontrolled	Duration of treatment:		
		arrhythmias	Median follow-up 2.93 y		
		II-III AV block	in amlodipine group; 2.95		
		Need for serious	y in enalapril group		
		steroids, NSAIDS or			
		cytotoxic drugs			

ESPIRAL Marin R, et al., 2001 (176) <u>11593109</u>	Aim: To investigate in a random comparison the capacity of an angiotensin converting enzyme inhibitor (fosinopril), and that of a long-acting dihydropiridine (nifedipine GITS) to modify the decay in renal function in pts with primary renal disease, exhibiting a progressive increase in serum Cr during the previous 2 y. Study type: Randomized open label trial	 Women of child-bearing potential not using appropriate contraceptives Any disease that could limit the ability of pts to comply with protocol requirements Inclusion criteria: 18–75 y Serum Cr between 1.5 and 5 mg/dL (133–442 µmol/l) HTN defined as BP >140/90 mm Hg or by the use of antihypertensive agent(s) Proven progression of chronic renal failure in the previous 2 y, defined by increase by >25% or >0.5 mg/dL (44.2 µmol/l) in serum Cr Mean SBP, mm Hg (SD): Nifedipine GITS: 157.5 (20) 	Intervention: • Nifedipine GITS: 30– 60 mg QD • Fosinopril: 10–30 mg QD • Drugs added in step- wise fashion to achieve BP goal. • Step 1: Randomized drug • Step 2: Furosemide (up to 100 mg) • Step 3: Atenolol (up to 100 mg) • Step 4: Doxazosin (up to 12 mg) • BP goal: Nifedipine GITS: <140/90 mm Hg	 <u>1° endpoint:</u> 1° Outcome: Time elapsed until serum Cr values doubled, or the need to enter a dialysis program 2° Outcome: CV events (including MI, stroke, angina, and death), proteinuria evolution and serum Cr <u>Safety endpoint</u>: N/A 	 Limitations: SBP was 4–6 mm Hg lower with ACEI which may have impacted improved outcomes. Still positive effects remained from fosinopril after adjusted for BP levels. Sodium restriction may have favored the ACEI group. Summary: Renal survival was significantly better if fosinopril used as first agent, unrelated to the primary renal disease. Proteinuria decreased by 57% in the fosinopril group and increased by 7% in the nifedipine GITS group while BP control did net differ between tractment
	renal function in pts	chronic renal failure in	 Step 2: Furosemide 		
	disease, exhibiting a	by increase by >25% or	 Step 3: Atenolol (up to 		better if fosinopril used as first
	serum Cr during the	in serum Cr	 Step 4: Doxazosin (up 		renal disease.
		(SD):	● BP goal:		in the fosinopril group and
	Randomized open	(20)			GITS group while BP control did
	label trial	Fosinopril: 155 (17) ● Mean DBP, mm Hg	Fosinopril: <140/90 mm Hg		not differ between treatment groups for DBP.
	<u>Size</u> : 241 Nifedipine GITS: 112 Fosinopril: 129	(SD): Nifedipine GITS: 96 (11) Fosinopril: 96 (8)	• Duration of treatment: mean follow-up NR; authors report minimum follow-up of 3 y and this		 3-y follow-up Doubling of serum Cr or entering dialysis N (%) Nifedipine GITS 40 (36%)
		Exclusion criteria: • DM • Previous recent history	is when most outcome measures reported		Fosinopril 27 (21%) OR: 0.47 (0.26–0.84); p=0.01 • Decrease in SBP, mm Hg (SD)
		of CVD (stroke, MI, or HF)			Nifedipine GITS 14.0 (22.5) Fosinopril 19.8 (19.6), p NR
		 Taking concomitant medications that could 			Decrease in DBP, mm Hg (SD) Nifedipine GITS 14.9 (11.8)

Parving HH, et al., of dual blockade of Pts with HTN, 18–85 y, losartan then aliskiren or Ratio of albumin to creatinine at 6 mo regarding function, survival, CV	Bakris GL, et al., 2010 (177) 20170948	Aim: To examine the effect of initial antihypertensive therapy with benazepril plus amlodipine compared to benazepril plus hydrochlorothiazide on progression of CKD Study type: RCT, forced drug titration Size: • Overall benazepril plus amlodipine n=5,744 benazepril plus hydrochlorothiazide n=5,762 • Pts with CKD benazepril plus amlodipine n=561 benazepril plus amlodipine n=561 benazepril plus hydrochlorothiazide n=532 • Pts without CKD benazepril plus amlodipine n=5,171 benazepril plus amlodipine n=5,171 benazepril plus amlodipine n=5,171 benazepril plus amlodipine n=5,171 benazepril plus amlodipine n=5,171 benazepril plus hydrochlorothiazide n=5,218 Aim: Compare effects of dual blockade of	Inclusion criteria: • Males or females ≥55 y, with HTN, high CV risk (history of coronary events, MI, revascularization, stroke, CKD, PAD, LVH, DM) • Entry BP for pts with CKD benazepril plus amlodipine: 145.1/78.6 (20.2/11.2) benazepril plus hydrochlorothiazide: 145.0/78.1 (20.5/10.7) • Rate of DM same in CKD and non-CKD pts (58.9% vs. 60.5%; p=0.302 Exclusion criteria: N/A Inclusion criteria: Pts with HTN, 18–85 y,	Intervention: • Initial antihypertensive therapy with benazepril plus amlodipine compared to benazepril plus hydrochlorothiazide • BP after dose adjustment benazepril plus amlodipine: 131.6/73.3 (18.2/10.3 SD), 4119 (75%) controlled • Benazepril plus hydrochlorothiazide: 132.5/74.4 (17.9/11.2 SD), 3963 (72%) controlled p<0.0013 Target <140/90 and <130/80 for DM or CKD Comparator: N/A Intervention: All on losartan then aliskiren or	 <u>1° endpoint:</u> Overall: time to first event of composite CV morbidity and mortality Progression of CKD, a prespecified endpoint, was defined as doubling of serum creatinine concentration or ESRD (estimated glomerular filtration rate <15 mL/min/1·73 m² or need for dialysis). All randomized pts were included in the intention-to-treat analysis. There were 113 (2.0% x 0%) events of CKD progression in the benazepril plus amlodipine group compared with 215 (3.7% x 7%) in the benazepril plus hydrochlorothiazide group HR: 0.52, (95% CI: 0.41–0.65), p<0.0001 2° endpoints: CKD plus death, change in albuminuria, change in eGFR Subset with more advanced CKD analyzed for rate of progression Safety endpoint: N/A 	Limitations: • Trial terminated early (mean follow-up 2.9 y [SD 0.4]) because of superior efficacy of benazepril plus amlodipine compared with benazepril plus hydrochlorothiazide with 20% lower CV risk. • Very small proportion of study population had albuminuria above 33.9 mg/mmol combined with early trial termination to reduce renal events. • Funded by Novartis. <u>Summary:</u> • Initial antihypertensive treatment with benazepril plus amlodipine slowed progression of nephropathy to a greater extent compared to benazepril plus hydrochlorothiazide. <u>Limitations:</u> No renal endpoints regarding function, survival, CV
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2008 (178) <u>18525041</u>	RAAS by aliskiren 300 mg/d added to maximal dose losartan 100 mg/d and optimal HTN therapy <u>Study type:</u> RCT, double-blinded, duration was 6 mo <u>Size:</u> 805 entered open label, 599 randomized, 524 completed.	and DM-2 and nephropathy (early morning alb/creat >300 mg/g or >200 mg/g in on RAAS blocker already <u>Exclusion criteria:</u> Non-DM kidney disease, >3,500 mg/g alb/ Cr ratio, eGFR, 30 mL/min/BSA, chronic urinary tract infections, baseline serum potassium >5.1, severe HTN, major CVD in prior 6 mo	placebo added <u>Comparator</u> : All on losartan, aliskiren or placebo added	 2°: decline in eGFR, development of renal dysfunction (serum creatinine >176.8 micromol/l (2.0 mg/dL) <u>Safety endpoint:</u> Hyperkalemia 5% in aliskiren group, 5.7% in placebo group but more frequent individual elevations >5.5 in aliskiren group 	events, BP 2/1 mm Hg lower in aliskiren group; supported by Novartis <u>Summary</u> : • Outcome was degree of albuminuria. Aliskiren reduced urinary alb/creat ratio by 20% (95% Cl 9–30; p<0.001) • From post hoc analysis: Antiproteinuric effects consistent across CKD stages (19%, 22%, and 18% for stages 3, 2, and 1). For CKD 3, renal dysfunction more frequent in placebo group (29.3 vs. 13.6%; p=0.032) • No differences in deaths or acute renal failure by treatment group (0.7% in both)
VA NEPHRON-D Fried LF, et al., 2010 (124) 20728887	<u>Aim</u> : To test the efficacy of the combination of losartan with lisinopril as compared with standard treatment with losartan alone in slowing the progression of proteinuric diabetic kidney disease <u>Study type:</u> RCT, multi-center, double- blind <u>Size</u> : 1448 were randomized	Inclusion criteria: Pts without adverse events on full dose losartan DM-2, eGFR 30–89.9 mL/min/1.73 m² by 4 variable MDRD formula, urinary albumin/creatinine ratio of ≥300 in a random sample Exclusion criteria: Known non-DM kidney disease, serum potassium >5.5 mmol/L, current treatment with sodium polystyrene sulfonate or inability to stop prescribed medications increasing risk of hyperkalemia.	Intervention:● Pts with DM-2 already taking losartan 100 mg/d with albumin to creatinine ratio of ≥300 were randomized to either lisinopril 10–40 mg/d or placebo.● 132 1° endpoints in the combination therapy group; No benefit to mortality or CV events. Combination therapy increase risk of hyperkalemia 6.3 events/100 person-y vs. 2.6 events/100 person-y vs. 2.6 events/100 person-y (p<0.001) and acute kidney injury 12.2 vs. 6.7 events/100 person-y (p<0.001)	<u>1° endpoint</u> : First occurrence of a change in eGFR (a decline of ≥30 mL/min/1.73 m² if initial GFR ≥60 or a decline of ≥50% if initial eGFR <60, ESRD or death 2° endpoint: First occurrence of decline in eGFR or ESRD <u>Safety endpoint</u> : mortality, hyperkalemia, acute kidney injury	<u>Summary</u> : Study stopped early due to safety concerns. Combination of ACEI and ARB was associated with increased risk of adverse events among pts with diabetic nephropathy

		Comparator: 152	
		primary endpoints in	
		monotherapy group	

Data Supplement 38. Nonrandomized Trials, Observational Studies, and/or Registries of CKD (Section 9.3)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Upadhyay A, et al., 2011 (179) <u>21403055</u>	Aim: To summarize trials comparing lower vs. higher BP targets in pts with CKD; focus on proteinuria as an effect modifier Study type: Systematic review Size: 2,272	Inclusion criteria: >50 pts/group, 1 y follow-up, outcomes of death, kidney failure, CV events, change in kidney function, number of antihypertensive agents, adverse events. 3 trials (MDRD, AASK, REIN-2; 8 reports)	<u>Results:</u> Overall trials did not show that BP target of <125/75–130/80 is more beneficial than a target of <140/90. Lower quality evidence suggests a low target may be beneficial in subgroups with proteinuria >300–1,000/d	Limitations: No pts with DM-1 included. Duration (mean follow- up 2–4 y) may be too short to detect differences in clinically important outcomes. Reporting of adverse events not uniform. Summary: Available evidence is inconclusive but does not prove a BP target <130/80 improves clinical outcomes more than a target of <140/90 in adults with CKD.
Lv, et al., 2013 (127) <u>23798459</u>	Aim: To assess the renal and CV effects of intensive BP lowering in people with CKD Study type: Systematic review Size: 9,287 pts with CKD and 1,264 kidney failure events	 Inclusion criteria: Randomized trials of pts with CKD assigned to different target BP that reported kidney failure and CV events. 11 trials on 9,287 pts with CKD and 1,264 kidney failure events (doubling of serum creatinine, 50% decline in GFR or ESKD) Included AASK, REIN-2, MDRD, Wuhl (children), Toto, Schrier plus 5 trials with CKD subgroups, also included the late nonrandomized follow-up studies for AASK and MDRD BP targets varied substantially between trials. 2 trials targeted mean BP <92 mm Hg for the intensive treatment arm, and 107 mm Hg in the standard treatment arm. 1 trial aimed for BP<130/80 mm Hg vs. a DBP of 90 mm Hg, 1 study targeted <120/80 mm Hg vs. 	<u>Results:</u> Compared with standard regimens, more intensive BP lowering reduced risk of composite endpoint HR: 0.82; 95% CI: 0.68–0.98, and ESKD HR: 0.79; 95% CI: 0.67–0.93. Effect was modified by proteinuria (p=0.006) and markers of trial quality. Intensive BP lowering reduced the risk of kidney failure HR: 0.73; 95% CI: 0.62–0.86 but not in pts without proteinuria at baseline HR: 1.12; 95% CI: 0.67–1.87. No clear effect on CV events or death.	 <u>Limitations</u>: All trials used open label, in 2 pts were blinded, substantial variability in design quality. There was substantial variability in BP targets by MAP, systolic and DBP or only DBP. Most trials did not include pts with diabetic kidney disease <u>Summary:</u> Renal outcomes: 7 trials (N=5,308) recorded a total of 1,264 kidney failure events. A -7.7 mm Hg difference in SBP and a -4.9 mm Hg difference in DBP seen between treatment arms. Overall, a more intensive regimen reduced risk of composite kidney failure events by 17% HR: 0.82; 95% CI: 0.68–0.98, reduced the risk of ESKD alone by 18% (pooled HR for composite outcomes: 0.79; 95% CI: 0.67–0.93). Intensive BP lowering had no effect on kidney failure in pts who did not have proteinuria (3 trials involving 1,218 pts HR: 1.12; 95% CI: 0.67–1.87), but it did reduce the risk of progressive kidney failure by 27% (5 trials involving 1,703 pts HR: 0.73; 95% CI: 0.62–0.86 in people who did have proteinuria at baseline.

		135–140/85–90 mm Hg, and 4 studies had DBP<75–80 mm Hg vs. from 80–90 mm Hg. A trial involving pediatric pts targeted a 24-h mean BP <the 50th<br="">percentile, compared with the 50th to 95th percentiles in the control group. 2 trials had more liberal targets for intensive treatment (<140–150 mm Hg systolic, 85 mm Hg diastolic)</the>		 CV outcomes: major CV events reported in 5 trials (472 CV events in 5,308 pts with CKD). Intensive BP lowering did not reduce risk of CV events in pts with CKD, but the CIs remained wide RR: 1.09; 95% CI: 0.83–1.42. 6 trials reported stroke outcomes (197 events in 5,411 pts), 5 trials reported MI (138 events in 4,317 pts), and 5 trials reported HF (118 events in 5,308 pts). They saw no clear effect of intensive treatment on any of these vascular outcomes. Death: 10 trials involving 6,788 participants reported 846 deaths. There was no clear effect of intensive BP lowering on risk of all-cause death RR: 0.94; 95% CI: 0.84–1.05) or CV death RR: 1.20; 95% CI: 0.82–1.75
2003 (180) 12965979 excr with prog durin antil ther. with prog durin antil ther. with prog durin antil ther. with prog durin antil ther. with prog durin antil ther. mod second ther. Stud in pt prog non dise	ihypertensive rapy with and hout ACEIs. <u>udy type:</u> 11 RCTs ots with edominantly ndiabetic kidney ease <u>re:</u> 1,860 pooled in evel meta- alysis; mean ration of follow-up	 Inclusion criteria: Pt-level meta-analysis using data from the AIPRD Study Group database to assess relationships among pts with nondiabetic kidney disease across a wide range of urine protein excretion values during antihypertensive therapy with and without ACEIs. The AIPRD Study Group database included 1,860 pts with nondiabetic kidney disease enrolled in 11 RCTs of ACEIs to slow the progression of kidney disease. The database contained information on BP, urine protein excretion, serum creatinine, and onset of kidney failure during 22,610 visits. Included only randomized trials (with a minimum 1 y follow-up) that compared the effects of antihypertensive regimens that included ACEIs with the effects of regimens that did not include ACEIs. HTN or decreased kidney function was required for entry into all studies. Exclusion criteria: Common to all studies: acute kidney failure, treatment with immunosuppressive meds, clinically significant chronic HF, obstructive uropathy, renal artery stenosis, active systemic disease, DM-1, history of transplantation, history of allergy to 	<u>1° endpoint</u> : Progression of CKD defined as doubling of serum creatinine or onset of kidney failure <u>Results</u> : Kidney disease progression documented in 311 pts, 124 (13.2%) in the ACEI group and 187 (20.5%) in the control group (p=0.001). 176 (9.5%) developed kidney failure: 70 (7.4%) in the ACEI group and 106 (11.6%) in the control group (p=0.002). SBP of 110–129 mm Hg and urine protein excretion <2.0 g/d were associated with lowest risk for kidney disease progression. ACEI beneficial after adjustment for BP and urine protein excretion (RR: 0.67; 95% CI: 0.53–0.84). The increased risk for kidney progression at higher SBP levels was greater in pts with urine protein excretion >1.0 g/d (p<0.006).	Limitations: Studies included were not designed to assess the effect of lowering BP and urine protein excretion on kidney disease progression. <u>Conclusions:</u> Although reverse causation cannot be excluded with certainty, SBP goal between 110 and 129 mm Hg may be beneficial in pts with urine protein excretion >1.0 g/d. SBP <110 mm Hg may be associated with higher risk for kidney disease progression.

		ACEIs, and pregnancy.		
Giatras I, et al., 1997 (181) <u>9273824</u>	<u>Aim:</u> To use meta- analysis to assess effects if ACEIs on development of ESRD in nondiabetic pts <u>Study type:</u> Meta- analysis <u>Size:</u> 1,594 pts from 10 studies	Inclusion criteria: All randomized studies comparing ACEIs with other antihypertensive agents, with at least 1 y of follow-up Exclusion criteria: Studies of diabetic renal disease and renal transplants were excluded.	Results:• Among 806 pts receiving ACEIs, 52 (6.4%) developed ESRD and 17 (2.1%) died.• In 788 controls, 72 (9.1%) developed ESRD and 12 (1.5%) died. The pooled RR were 0.70; 95% CI: 0.51–0.97 for ESRD and 1.24; CI: 0.55– 2.83 for death.• The decreases in weighted mean systolic and DBPs during follow-up were 4.9 and 1.2 mm Hg greater, respectively, in the pts who received ACEIs.	 <u>Limitations:</u> Included studies through 5/1996, published (7) and nonpublished (3) study results. Did not require that pts have HTN or renal insufficiency at baseline. Did not report results by severity of proteinuria related to the diseases included many of which are not characterized by proteinuria. <u>Summary</u>: ACEIs are more effective than other antihypertensive agents in reducing the development of end-stage nondiabetic renal disease, and they do not increase mortality. It could not be determined whether this beneficial effect is due to the greater decline in BP or to other effects of ACE inhibition.
ONTARGET Investigators, et al., 2008 (126) <u>18378520</u>	Aim: Evaluate whether use of an ARB was noninferior to ACEI, and whether the combination was superior to ACE alone in the prevention of vascular events in pts with CVD or DM but not HF. Study type: Multi- center, double-blind, RCT Size: 25,620 pts	Inclusion criteria: • ≥55 y • Coronary, peripheral, or cerebrovascular disease or DM with end- organ damage Exclusion criteria: • Inability to discontinue ACEI or ARB • Known hypersensitivity or intolerance to ACEI or ARB • Selected CVDs (congestive HF, hemodynamically significant valvular or outflow tract obstruction, constrictive pericarditis, complex congenital heart disease, syncopal episodes of unknown etiology <3 mo, planned cardiac surgery or PTCA <3 mo, uncontrolled HTN on treatment [e.g., BP >160/100 mm Hg], heart transplant recipient, stroke due to subarachnoid hemorrhage) • Other conditions (significant renal artery disease, hepatic dysfunction, uncorrected volume or sodium depletion,	Intervention: Ramipril 10 mg daily (n=8,576) Comparator: • Telmisartan 80 mg daily (n=8,542) • Combination of telmisartan and ramipril (n=8,502)	 <u>1° endpoint</u>: After a median follow-up of 56 mo, there was no difference between ramipril vs. telmisartan or combination therapy vs. ramipril in the 1° composite outcome of death from CV causes, MI, stroke, or hospitalization for HF RR: 1.01; 95% CI: 0.94–1.09 and RR: 0.99; 95% CI: 0.92–1.07, respectively <u>Safety endpoint</u>: Combination therapy was associated with greater risk of hyperkalemia than ramipril monotherapy (480 pts vs. 283 pts; p<0.001) Hypotensive symptoms were cited as reason for permanent discontinuing more in telmisartan vs. ramipril RR: 1.54; p<0.001 and combination therapy vs. ramipril monotherapy RR: 2.75; p<0.001 Renal impairment was more common in combination therapy vs. ramipril monotherapy RR: 1.33; 95% CI: 1.22–1.44).

VALIANT White HD, et al., 2005 (182) <u>16301343</u>	Aim: Evaluate whether use of an ARB or the combination of an ACEI and an ARB was superior to a proven effective dose of an ACEI after AMI in pts with HF and/or LVEF <40%. Study type: Multi- center, double-blind, RCT Size: 14,703 pts	1° hyperaldosteronism, hereditary fructose intolerance, other major noncardiac illness or expected to reduce life expectancy or significant disability interfere with study participation, simultaneously taking another experimental drug, unable to provide written informed consent). Inclusion criteria: • ≥18 y • Between 12 h and 10 d after AMI • Clinical or radiological signs of HF and/or evidence of depressed LV systolic function with EF<40% or reduced echo wall motion index Exclusion criteria: • Cardiogenic shock • Serum creatinine >2.5 mg/dL • Known hypersensitivity or intolerance to ACEI or ARB • SBP<100 mm Hg • Known or suspected bilateral renal artery stenosis • Stroke or TIA within previous 3 mo • Refractory ventricular arrhythmia • Refractory angina • Right ventricular MI • Mitral stenosis, mitral regurgitation, aortic stenosis, aortic regurgitation, aortic stenosis, aortic regurgitation, aortic stenosis, aortic regurgitation of hemodynamic significance • Obstructive cardiomyopathy • Previous major organ transplant • Conditions likely to lead to poor adherence	Intervention: Valsartan 160 mg bid Comparator: • Captopril 50 mg tid • Combination of captopril 50 mg tid and valsartan 160 mg bid • Analyzed by prespecified age groups of <65 y (n=6988) 65–74 y (n=4555) 75–84 y (n=2777) ≥85 y (n=383)	 <u>1° endpoint</u>: All-cause mortality <u>2° endpoint</u>: Composite of CV mortality or emergency treatment or hospitalization for new or worsening HF, reinfarction, stroke, and resuscitated cardiac arrest On 3-y multivariable analysis, each 10-y age increase was associated with HR: 1.49; 95% CI: 1.43–1.56); p<0.0001 for mortality and an OR: 1.38; 95% CI: 1.31–1.46; p<0.0001 for readmission with HF. Similar but slightly smaller trend for composite endpoint, higher mainly in the oldest group. Valsartan was at least as effective as captopril in reducing mortality and other adverse outcomes in all age groups and combination therapy with both agents added no incremental benefit. Combination therapy increased the incidence of adverse effects leading to discontinuation in all age groups Safety endpoint: Adverse events associated with captopril and valsartan were more common in the elderly and in pts receiving combination therapy. Renal dysfunction was more common with older age and combination therapy.
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Data Supplement 39. RCTs Comparing Hypertension after Renal Transplantation (Section 9.3.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Midtvedt K, et al., 2001 (183) <u>11468543</u>	Aim: To compare the effect of an ACEI (lisinopril) with a CCB (controlled release nifedipine) in the treatment of post- transplant HTN focusing on changes in LVH. Study type: prospective RCT Size:154 pts 123 completed 1 y good quality echo data for 116 at 2 and 12 mo post treatment	Inclusion criteria: All RTx pts with HTN by DBP ≥95 in first 3 wk after transplant Exclusion criteria: Normotensive, isolated systolic HTN, refusal, requirement of ACEI for HF.	Intervention: Renal transplant recipients with HTN (DBP ≥95 mm Hg) in the first 3 wk after Transplant were randomized to double-blind nifedipine CR 30 mg or lisinopril 10 mg daily. Comparator: 2 treatment arms	<u>1° endpoint</u> : BP controlled in both groups (mean 140 \pm 16/87 \pm 8 with nifedipine, 136 \pm 17/85 \pm 8 with lisinopril, NS). LV mass reduced by 15% (p<0.001) in both groups (from 153 \pm 43 to 131 \pm 38 g/m ² with nifedipine and from 142 \pm 35 to 121 \pm 34 g/m ² with lisinopril) with no difference between groups at baseline or at follow-up.	Summary: In renal transplant pts with HTN with well-controlled BP, there is regression of LV mass after renal transplantation which is observed to be similar in pts treated with lisinopril or nifedipine.
Midtvedt K, et al., 2001 (184) <u>11740389</u>	Aim: To examine whether graft function as determined by GFR was better maintained with a CCB (controlled release nifedipine) as compared to an ACEI (lisinopril) in hypertensive renal transplant recipients treated with cyclosporine. Study type: Prospective RCT Size:154 pts • 123 completed 1 y good quality echo data for 116 at 2 and 12 mo post-Transplant • 64 recruited to complete a 2nd y	Inclusion criteria: All renal transplant pts with HTN by DBP ≥95 in first 3 wk after transplant Exclusion criteria: Normotensive, isolated systolic HTN, refusal, requirement of ACEI for HF.	Intervention: Renal transplant pts with HTN (DBP ≥95 mm Hg) in the first 3 wk after transplant were randomized to double-blind nifedipine CR 30 mg or lisinopril 10 mg daily. <u>Comparator</u> : 2 treatment arms	<u>1° endpoint:</u> • GFR baseline at 3–5 wk after entry, and at 1 and 2 y • Nifedipine: baseline GFR 46 mL/min, at 1 y 56 • Lisinopril: baseline GFR 43, at 1 y 44 • delta N vs. L: 9.6 at 1 y (95% CI: 5.5–13.7 mL/min; p=0.0001), 10.3 at 2 y (95% CI: 4.0–16.6 mL/min; p=0.0017) • Baseline GFR similar, change in GFR significant after 1 y and remained statistically significant after 2 y	Summary: Both nifedipine and lisinopril were safe and effective in treatment of HTN in renal transplant pts treated with cyclosporine. Pts receiving nifedipine but not lisinopril had improved renal function over 2 y.

Suwelack B, et al., 2000 (185) <u>11009288</u>	Aim: To compare the structural and functional cardiac changes of quinapril vs. atenolol administered to hypertensive kidney transplant recipients Study type: Prospective RCT Size: 31 cyclosporine treated stable function recipients with HTN 6–12 wk after transplant	Inclusion criteria: Cyclosporine-based immunosuppression, stable graft function with serum creatinine <2.5 mg/dL. Exclusion criteria: Pts with severe aortic or mitral regurgitation or with heart rates >100 beats/min	Intervention: • Cyclosporine treated stable function pts with HTN 6–12 wk after transplant randomized to double-blinded quinapril or atenolol to target DBP<90. • Echo within 24 h of first dose and at 24 mo • Stepwise increase in dose, could then add furosemide 40–80 mg/d, third-line CCB <u>Comparator</u> : 2 treatment arms	 <u>1° endpoint</u>: BP was lower in the atenolol group, delta 10.7 ± 3.4 mm Hg vs. 4.5 ± 2.9 mm Hg with quinapril E/A ratio (impaired relaxation) increased (improved) only in quinapril group (+0.11; p<0.05) and decreased by 0.03 (p>0.05 vs. start of treatment) in the atenolol group. Difference in E/A ratio alterations was significant (p<0.05). LV mass index decreased only in quinapril group (p<0.05) from entry to 24 mo. 	 Summary: In hypertensive renal allograft recipients, quinapril in contrast to atenolol provided a sufficient reduction in LVH and a concomitant improvement in LV diastolic cardiac relaxation and these effects occurred independently from BP reduction. While the conclusion was that quinapril showed a benefit not seen with atenolol, the actual numbers are very close (14.1 ± 10.1 atenolol, 15.8 ± 7.7 quinapril). BP reduction was twice as great in the atenolol group as in the quinapril group. Arterial BP did not correlate with cardiac mass reduction.
Paoletti E, et al., 2007 (186) <u>17591533</u>	<u>Aim</u> : To assess the effectiveness of ACEIs in regressing LVH persisting after renal transplantation during an 18-mo observation period. To assess the impact of cyclosporine vs. tacrolimus in affecting LVH outcome. <u>Study type</u> : Prospective RCT <u>Size</u> : 70 renal transplant recipients at 3–6 mo after transplant.	Inclusion criteria: • Renal transplant pts with serum creatinine <2.5 mg/dL, urine protein excretion not exceeding 1 g/d and with persistent LVH at 3–6 mo after transplant. • Previously randomized to either cyclosporine or tacrolimus immunosuppression. • All were pts of deceased donor transplants. Exclusion criteria:	Intervention: • RCT Lisinopril (n=36) vs. placebo (n=34), also used other agents to treat HTN • Endpoint LVMI at 18 mo • Echo at 3–6 mo and at 18 mo <u>Comparator</u> : Treatment vs. placebo	 <u>1° endpoint</u>: Change in LV mass index at 18 mo. BP decreased in both groups (p=NS, between group differences SBP -1.7 ± 3.3 mm Hg; 95% Cl: -4.8–8.2; and DBP 0.3 ± 2.2 mm Hg; 95% Cl: -4.8–4.1). LVMI regressed more in ACEI group (-9.1 ± 13.3 g/m 2.7; p<0.001) but only in those on cyclosporine immunosuppression. Interaction of LVMI effect and 	Summary: LVMI regressed more in ACEI group but only in those on cyclosporine immunosuppression. Interaction of LVMI effect and cyclosporine in post hoc analysis.

	Aim . To toot the office ou of the	 No DM, HF, severe valvular disease, previous renal artery stenosis blocking agents, acute rejections in prior 3 mo or significant renal artery stenosis. Pts receiving a preemptive 2nd transplant or a living donor transplant were excluded. 	Intervention	cyclosporine in post hoc analysis. • 74/104 had LVMI above normal. • Change in LVMI ACEIs vs. controls p<0.001 Number of meds comparable • Number using CCB/BBs/diuretic/others was 17/21/2/9 for ACEI, 24/26/3/15 controls	Summary Study stopped
VA NEPHRON-D Fried LF, et al., 2010 (124) 20728887	<u>Aim</u> : To test the efficacy of the combination of losartan with lisinopril as compared with standard treatment with losartan alone in slowing the progression of proteinuric diabetic kidney disease <u>Study type:</u> RCT, multi-center, double-blind <u>Size:</u> 1,448 were randomized	Inclusion criteria: Pts without adverse events on full dose losartan DM-2, eGFR 30–89.9 mL/min/1.73 m² by 4 variable MDRD formula, urinary albumin/creatinine ratio of ≥300 in a random sample Exclusion criteria: Known nondiabetic kidney disease, serum potassium >5.5 mmol/L, current treatment with sodium polystyrene sulfonate or inability to stop prescribed medications increasing risk of hyperkalemia.	 Intervention: Pts with DM-2 already taking losartan 100 mg/d with albumin to creatinine ratio of ≥300 were randomized to either lisinopril 10–40 mg/d or placebo. 132 1° endpoints in the combination therapy group No benefit to mortality or CV events. Combination therapy increase risk of hyperkalemia 6.3 events/100 person-y vs. 2.6 events/100 person-y (p<0.001) and acute kidney injury 12.2 vs. 6.7 events/100 person-y (p<0.001) Comparator: 152 1° endpoints in monotherapy group 	<u>1° endpoint</u> : First occurrence of a change in eGFR (a decline of ≥30 mL/min/1.73 m ² if initial GFR ≥60 or a decline of ≥50% if initial eGFR <60, ESRD or death <u>2° endpoint</u> : First occurrence of decline in eGFR or ESRD <u>Safety endpoint</u> : Mortality, hyperkalemia, acute kidney injury	Summary: Study stopped early due to safety concerns. Combination of ACEI and ARB was associated with increased risk of adverse events among pts with diabetic nephropathy

Data Supplement 40. Nonrandomized Trials, Observational Studies, and/or Registries for Hypertension after Renal Transplantation (Section 9.3.1)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Cross NB, et al., 2009 (187) <u>19588343</u>	Study type: Comparative assessment by drug class using RCTs and quasi-RCTs lasting at least 2 wk in kidney transplant pts Size: • 60 studies, 3,802 pts, most taking cyclosporine based immunosuppression • 29 studies (n=2,262) compared CCB to placebo, 10 (n=445) ACEI to placebo, 7 (n=405) CCB to ACEI	Inclusion criteria: 21 studies for HTN, 6 for erythrocytosis, 2 CAN, 2 LVH, 30 not specified Exclusion criteria: N/A	<u>1° endpoint</u> : To assess comparative effects of antihypertensive agents in kidney transplant pts <u>Results:</u> Used random effects meta- analysis, risk ratios for dichotomous outcomes and MD for continuous outcomes, both with 95% CI. Stratified analyses and meta- regression to investigate heterogeneity.	 CCBs vs. placebo or no treatment had strongest results: improved GFR MD: 4.45 mL min (95% CI: 2.22–6.68), reduced graft loss RR: 0.75, (95% CI: 0.57–0.99). ACEI vs. placebo inconclusive for GFR MD: -8.07 mL/min (95% CI: -18.57–2.43) and variable for graft loss. Compared to CCB, ACEI decreased GFR MD: -11.48 mL/min; 95% CI: -5.75– -7.21), proteinuria MD: -0.28 g/24 h (95% CI: -0.47– -0.10), also reduced hemoglobin MD: -12.96 g/L (95% CI: -5.72– -10.21) and increased hyperkalemia RR: 3.74 (95% CI: 1.89– 7.43). Graft loss data were inconclusive. CCB may be preferred as first line for HTN after kidney transplant. ACEI may have some detrimental effects. There were not enough studies with other agents.
Jennings DL, et al., 2008 (188) <u>18094340</u>	Study type: Literature review Size: 5 studies with 3 reporting safety endpoints and 2 reporting clinical efficacy endpoints	Inclusion criteria: Studies using either ACEI or ARB initiated within the first 12 wk after renal transplant	<u>1° endpoint</u> : Safety or efficacy <u>Results:</u> • No significant increase in serum creatinine or potassium after up to 9 mo Rx • Early initiation of ACEI may be more effective than BB in reducing LVH and proteinuria after 24 mo treatment	<u>Conclusion:</u> Reasonable to consider RAAS inhibitors as first-line treatment in pts with HTN and compelling indications i.e., DM, HF in first 12 wk after renal transplant.
Ninomiya T, et al., 2013 (189) <u>24092942</u>	<u>Aim:</u> To define CV effects of lowering BP in pts with CKD <u>Study type:</u>	Inclusion criteria: Had to meet 1 of the following criteria: Pts randomized to a BP-lowering drug/regimen or a control group (placebo or less intensive BP lowering regimen) or pts randomized	<u>Results:</u> Compared with placebo, BP lowering regimens reduced the risk of major CV events by about a sixth per 5 mm Hg reduction in SBP in individuals with (HR: 0.83; 95% CI	Limitations: • Limited numbers with CKD and most were stage 3a: • There were 121,995 pts (80%) with eGFR ≥60 mL/min/1.73 m ² (mean eGFR 81 (SD 17)

	 Meta-analysis of RCTs Individual pt data available for 23 trials, with summary data from another 3. Meta- analysis was performed according to baseline kidney function. <u>Size:</u> 26 trials (152,290 pts), including 30,295 pts with reduced eGFR, defined as eGFR <60 mL/min/1.73 m². 	between regimens based on different classes of drugs to lower BP. Trials required to have at least 1,000 pt-y of planned follow-up in each randomized arm and not to have presented or published their main results before finalization of the overview protocol in July 1995. <u>Exclusion criteria:</u> Trials prior to July 1995.	0.76–0.90) and without reduced eGFR (HR: 0.83; 95% CI: 0.79– 0.88), with no evidence for any difference in effect (p=1.00 for homogeneity). The results were similar irrespective of whether BP was reduced by regimens based on ACEIs, calcium antagonists, or diuretics/BBs. There was no evidence that the effects of different drug classes on major CV events varied between pts with different eGFR (all p>0.60 for homogeneity).	 mL/min/1.73 m²) and 30,295 pts (20%) with eGFR <60 mL/min/1.73 m² (mean 52 (SD 7) mL/min/1.73 m²) at baseline (table 4¹). Only 439 pts (0.3%) had eGFR <30 mL/min/1.73 m² at baseline. Limited numbers had proteinuria, present in 2,500 (7%) of 37161 pts with data available. Summary: These analyses provided compelling evidence for the CV benefits of reduction in BP in pts with stage 1–3 CKD. The proportional reductions in risk of major CV events were similar in pts with and without evidence of CKD, however those with CKD stood to gain larger absolute benefits because their baseline risk was much higher. BP-lowering is an effective strategy for preventing CV events among pts with moderately reduced eGFR. There is little evidence from these overviews to support the preferential choice of particular drug classes for the prevention of CV events in CKD.
ONTARGET Investigators, et al., 2008 (126) <u>18378520</u>	<u>Aim</u> : Evaluate whether use of an ARB was noninferior to ACEI, and whether the combination was superior to ACE alone in the prevention of vascular events in pts with CVD or DM but not HF. <u>Study type</u> : Multi- center, double-blind, RCT <u>Size</u> : 25,620	Inclusion criteria: • ≥55 y • Coronary, peripheral, or cerebrovascular disease or DM with end-organ damage Exclusion criteria: • Inability to discontinue ACEI or ARB • Known hypersensitivity or intolerance to ACEI or ARB • Selected CVDs (congestive HF, hemodynamically significant valvular or outflow tract obstruction, constrictive pericarditis, complex congenital heart disease, syncopal episodes of unknown etiology <3 mo, planned cardiac surgery or PTCA <3 mo, uncontrolled HTN on treatment [e.g., BP >160/100 mm Hg], heart transplant recipient,	Intervention: Ramipril 10 mg daily (n=8,576) <u>Comparator</u> : • Telmisartan 80 mg daily (n=8,542) • Combination of telmisartan and ramipril (n=8,502)	 <u>1° endpoint</u>: After a median follow-up of 56 mo, no difference between ramipril vs. telmisartan or combination therapy vs. ramipril in the 1° composite outcome of death from CV causes, MI, stroke, or hospitalization for HF RR: 1.01 (95% CI: 0.94–1.09) and RR: 0.99 (95% CI: 0.92–1.07), respectively. <u>Safety endpoint</u>: Combination therapy was associated with greater risk of hyperkalemia than ramipril monotherapy (480 pts vs. 283 pts; p<0.001) Hypotensive symptoms were cited as reason for permanent discontinuing more in telmisartan vs. ramipril RR: 1.54, p<0.001; and combination therapy vs. ramipril monotherapy RR: 2.75, p<0.001 Renal impairment was more common in combination therapy vs. ramipril monotherapy RR: 1.33; 95% CI: 1.22–1.44

VALIANT White HD, et al., 2005 (182) <u>16301343</u>	Aim: Evaluate whether use of an ARB or the combination of an ACEI and an ARB was superior to a proven effective dose of an ACEI after AMI in pts with HF and/or LVEF <40%. Study type: Multi- center, double-blind, RCT Size: 14,703	stroke due to subarachnoid hemorrhage) • Other conditions (significant renal artery disease, hepatic dysfunction, uncorrected volume or sodium depletion, 1° hyperaldosteronism, hereditary fructose intolerance, other major noncardiac illness or expected to reduce life expectancy or significant disability interfere with study participation, simultaneously taking another experimental drug, unable to provide written informed consent). Inclusion criteria: • ≥18 y • Between 12 h and 10 d after AMI • Clinical or radiological signs of HF and/or evidence of depressed LV systolic function with EF<40% or reduced echo wall motion index Exclusion criteria: • Cardiogenic shock • Serum creatinine >2.5 mg/dL • Known hypersensitivity or intolerance to ACEI or ARB • SBP<100 mm Hg • Known or suspected bilateral renal artery stenosis • Stroke or TIA within previous 3 mo • Refractory ventricular arrhythmia • Refractory angina • Right ventricular MI	Intervention: Valsartan 160 mg bid Comparator: • Captopril 50 mg tid • Combination of captopril 50 mg tid and valsartan 160 mg bid • Analyzed by prespecified age groups of <65 (n=6,988) 65 to 74 (n=4,555) 75 to 84 (n=2,777) ≥85 y (n=383)	 <u>1° endpoint</u>: All-cause mortality <u>2° endpoint</u>: Composite of CV mortality or emergency treatment or hospitalization for new or worsening HF, reinfarction, stroke, and resuscitated cardiac arrest On 3-y multivariable analysis, each 10-y increase was associated with HR: 1.49 (95% CI: 1.43–1.56), p<0.0001 for mortality and OR: 1.38 (95% CI: 1.31–1.46; p<0.0001) for readmission with HF. Similar but slightly smaller trend for composite endpoint, higher mainly in the oldest group. Valsartan was at least as effective as captopril in reducing mortality and other adverse outcomes in all age groups and combination therapy with both agents added no incremental benefit. Combination therapy increased the incidence of adverse effects leading to discontinuation in all age groups
		 Mitral stenosis, mitral regurgitation, aortic stenosis, aortic regurgitation of hemodynamic significance Obstructive cardiomyopathy Previous major organ transplant 		 Safety endpoint: Adverse events associated with captopril and valsartan were more common in the elderly and in pts receiving combination therapy. Renal dysfunction was more common with older age and combination therapy.

		Conditions likely to I adherence	ead to poor		
SPRINT Senior Williamson JD, et al., 2016 (190) <u>27195814</u>	Aim: Intensive SBP goal <120 mm Hg) vs. standard (SBP goal <140) Study type: RCT Size: 2,636; 30% met criteria for being classified as ambulatory frail Mean follow-up:3.1 y	Inclusion criteria: Men and women age 75+; mean age 79.8 y; 38% women; 17% black, 74% Caucasian Exclusion criteria: Nursing home residents; prevalent DM, stroke, Class III/IV HF, dementia	Intervention: Medications and dietary advice to achieve SBP of <120 mm Hg Comparator: Medications and dietary advice to achieve SBP of <140 mm Hg Achieved SBP: Intensive= 123.4 mm Hg Standard= 134.8 mm Hg	 <u>1° endpoint</u>: Composite CVD outcome (AMI, non-MI ACS, Stroke, HF, CVD death. <u>Results</u>: 102 events in the intensive treatment group vs. 148 events in the standard treatment group; HR: 0.66; 95% CI: 0.51–0.85 and all-cause mortality (73 deaths vs. 107 deaths, respectively; HR: 0.67; 95% CI: 0.49–0.91. No difference in falls, orthostatic hypotension, or overall SAEs. NNT for 1° outcome=27 and NNT for all-cause mortality=41 	Limitations: Does not apply to nursing home pts of those with dementia or advance <u>Conclusions:</u> Intensive SBP is safe and effective for lowering CVD events and total mortality in adults ≥75 y

Data Supplement 41. RCTs Comparing Acute Intracerebral Hemorrhage Outcomes (Section 9.4.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
INTERACT2 Anderson CS, et al., 2013 (191) <u>23713578</u>	Aim: To assess whether rapid lowering of elevated BP would improve the outcome in pts with ICH. Study type: Phase III RCT Study size: 2,839 pts	Inclusion criteria: Pts with spontaneous ICH within the previous 6 h with elevated SBP	Design: Intensive treatment to lower BP (with a target systolic level of <140 mm Hg within 1 h) vs. guideline- recommended treatment (with a target SBP <180 mm Hg) among pts with SBP between 150 and 220 mm using agents of the physician's choosing.	 <u>1° outcome</u>: Death or major disability (score of 3 to 6 on the modified Rankin scale) at 90 d. <u>Pre-specified 2° outcome</u>: Ordinal analysis of the modified Rankin score. <u>Key findings</u>: Among the 2,794 pts for whom the 1° outcome could be determined, 719 of 1,382 participants (52.0%) receiving 	Summary: • In pts with ICH, intensive lowering of BP did not result in a significant reduction in the rate of death or severe disability. • However, there may be improved functional outcomes with intensive lowering of BP. • INTERACT-2 is so far the largest (and only phase 3) RCT evaluating efficacy of intensive BP lowering.

				 intensive treatment, vs. 785 of 1,412 (55.6%) receiving guideline-recommended treatment, had a 1° outcome event; intensive treatment OR: 0.87; 95% CI: 0.75–1.01; p=0.06. The ordinal analysis showed significantly lower modified Rankin scores with intensive treatment. OR for greater disability: 0.87; 95% CI: 0.77–1.00; p=0.04. Mortality was 11.9% in the group receiving intensive treatment and 12.0% in the group receiving guideline-recommended treatment. Nonfatal serious adverse events occurred in 23.3% and 23.6% of the pts in the 2 groups, respectively. 	 No clear relationship between outcome and time from onset of ICH to commencing treatment and no significant effect of intensive BP- lowering treatment on hematoma growth. Of note, only1 third of pts achieved the target SBP level within 1 h (half achieved the target by 6 h), and most (75%) presented with mild to moderate size (<20 mL) hematomas.
ATACH-1 2010 (192) <u>19770736</u>	Aim: To determine the feasibility and acute (i.e., within 72 h) safety of 3 levels of SBP reduction in subjects with supratentorial ICH treated within 6 h after symptom onset. Study type: Phase I, dose- escalation, multicenter prospective study. Study size: 60	Inclusion criteria: Pts with ICH with elevated SBP ≥170 mm Hg who presented to the ED within 6 h of symptom onset.	 Design: IV nicardipine to reduce SBP to a target of: <u>#1</u>: 170–200 mm Hg in the first cohort of pts <u>#2</u>: 140–170 mm Hg in the 2nd cohort <u>#3</u>: 110–140 mm Hg in the third cohort. Each subject was followed-up for 3 mo to preliminarily assess mortality and the clinical outcomes. A total of 18, 20, and 22 pts were enrolled in the respective 3 tiers of SBP treatment goals. 	 <u>1° outcome</u>: Treatment feasibility (achieving and maintaining the SBP goals for 18–24 h) <u>2° outcomes</u>: <u>#1</u>: Neurologic deterioration within 24 h; <u>#2</u>: Serious adverse events within 72 h. <u>Key findings</u>: Overall, 9 of 60 pts had treatment failures (all in the last tier). A total of 7 subjects with neurologic deterioration were observed: 1 (6%), 2 (10%), and 4 (18%) in tier 1, 2, and 3, respectively. Serious adverse events were observed in1 subject (5%) in tier 2 and in 3 subjects (14%) in tier 3. However, the safety stopping rule was not activated in any of the tiers. 3 (17%), 2 (10%), and 5 (23%) subjects in tiers1, 2, and 3, respectively, died within 3 mo 	Summary: • Observed proportions of neurologic deterioration and serious adverse events were below the prespecified safety thresholds, and the 3-mo mortality rate was lower than expected in all SBP tiers.
INTERACT-1	Aim: To assess the safety and efficiency of	Inclusion criteria: Pts with	Design: Early intensive lowering of BP (target SBP	1° outcome: Proportional change in hematoma volume at 24 h.	Summary: Early intensive BP- lowering treatment is clinically

Anderson CS, et al.,	this treatment, as a	acute	140 mm Hg; n=203) vs.		feasible, well tolerated, and might
2008 (193)	run-in phase to a larger	spontaneous	standard guideline-based	2° outcomes: Measurements of	reduce hematoma growth in ICH.
18396107	trial.	ICH diagnosed	management of BP (target	hematoma volume.	reduce nematoria growth in fort.
10070107		by CT within 6 h	SBP 180 mm Hg; $n=201$).		
	Study type:	of onset,	22. 100 min 19, 11 201).	Safety and clinical outcomes: Assessed	
	Randomized pilot trial	elevated SBP		for up to 90 d.	
		(150–220 mm			
	Study size: 404	Hg), and no		Key findings:	
	<u></u>	definite		 Mean hematoma volumes were smaller 	
		indication or		in the guideline group (12.7 mL, SD 11.6)	
		contraindication		than in the intensive group (14.2 mL, SD	
		to treatment		14.5).	
				• From randomization to 1 h, mean SBP	
				was 153 mm Hg in the intensive group and	
				167 mm Hg in the guideline group	
				(difference 13.3 mm Hg (95% CI: 8.9–17.6)	
				mm Hg; p<0.0001); from 1 h to 24 h, BP	
				was 146 mm Hg in the intensive group and	
				157 mm Hg in the guideline group (10.8	
				mm Hg; 95% CI: 7.7–13.9 mm Hg;	
				p<0.0001).	
				 Mean proportional hematoma growth 	
				was 36.3% in the guideline group and	
				13.7% in the intensive group (difference	
				22.6%; 95% CI: 0.6%–44.5%; p=0.04) at	
				24 h.	
				 After adjustment for initial hematoma 	
				volume and time from onset to CT, median	
				hematoma growth differed between the	
				groups with p=0.06; the absolute	
				difference in volume between groups was	
				1.7 mL (95% CI: -0.5–3.9; p=0.13). RR of	
				hematoma growth ≥33% or ≥12.5 mL was	
				36% lower (95% CI: 0%–59%; p=0.05) in	
				the intensive group than in the guideline	
				group. Adjusted RR: 8% (95% CI: -1.0%-	
				17%; p=0.05).	
				 Intensive BP-lowering treatment did not 	
				alter the risks of adverse events or 2°	
				clinical outcomes at 90 d.	

Tsivgoulis G, et al., 2014 (194) <u>25239836</u>	Aim: To evaluate the safety and efficacy of intensive BP reduction in pts with acute-onset ICH Study type: Systematic review and meta-analysis of RCTs. Study size: 4 eligible studies, including a total of 3,315 pts	Inclusion criteria: Pts with acute ICH randomized to either intensive or guideline BP- reduction protocols.	• Intensive early BP lowering after acute ICH onset compared with guideline-based treatment	 Key findings: Death rates similar between pts randomized to intensive BP-lowering treatment and those receiving guideline BP-lowering treatment OR: 1.01; 95% CI: 0.83–1.23; p=0.914 Intensive BP-lowering treatment associated with strong trend towards lower 3-mo death or dependency vs. guideline treatment OR: 0.87; 95% CI: 0.76–1.01; p=0.062. Intensive BP reduction was also associated with a greater attenuation of absolute hematoma growth at 24 h (standardized MD± standard error: -0.110 ± 0.053; p=0.038). 	 Summary: Intensive BP management in pts with acute ICH is safe. Intensively treated ICH pts tended to have more favorable 3-mo functional outcome. Intensive BP reduction associated with a greater attenuation of absolute hematoma growth at 24 h. Starting antihypertensive treatment in the initial 5–10 d after ICH may have a different outcome from that seen after an ischemic stroke because of 2° edema formation and hemodynamic changes
ATACH2 Qureshi Al, et al., 2016 27276234	<u>Aim</u> : To determine the relative efficacy of intensive vs. standard antihypertensive treatment that was initiated within 4.5 H after symptom onset and continued for the next 24 H in patients with spontaneous supratentorial intracerebral hemorrhage <u>Study type</u> : Phase III RCT <u>Study size</u> : 1,000 pts	Inclusion criteria: Pts with spontaneous ICH (volume, <60 cm3) and a Glasgow Coma Scale (GCS) score of 5 or more	Design: Intravenous nicardipine administered within 4.5 H after symptom onset and continued for the next 24 H to lower BP	 <u>1° outcome</u>: Moderately severe or severe disability or who had died (modified Rankin scale score, 4 to 6) at 3 months <u>Key findings</u>: Among 1,000 participants with a mean (±SD) systolic BP of 200.6±27.0 mm Hg at baseline, 500 were assigned to intensive treatment and 500 to standard treatment. Enrollment was stopped because of futility Death or disability occurred in 38.7% of patients in the intensive-treatment group and 37.7% in the standard-treatment group. RR: 1.04; 95%CI: 0.85–1.27. Serious adverse events occurring within 72 H after randomization were reported in 1.6% of the patients in the intensive-treatment group. Renal adverse events within 7 d after randomization were significantly higher in the intensive-treatment group than in the 	Summary: Treatment of patients with spontaneous ICH to achieve a target systolic BP of 110 to 139 mm Hg did not result in a lower rate of death or disability compared to conventional reduction to a target of 140–179 mm Hg. Furthermore, there was more than twice the frequency of renal adverse events in the more intensively treated arm within a week of treatment initiation.

	standard-treatment group (9.0% vs. 4.0%,	
	p=0.002).	

Data Supplement 42. RCTs Comparing Acute Ischemic Stroke Outcomes (Section 9.4.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
COSSACS Robinson TG, et al., 2010 20621562	Aim: Assess the efficacy and safety of continuing or stopping pre- existing antihypertensive drugs in patients with acute stroke Study type: RCT Size: 763	Inclusion criteria: Acute ischemic stroke (or ICH) within previous 48 h Exclusion criteria: • Impaired level of consciousness • Unable to swallow • Hypertensive emergency • BP >200/120 mm Hg • Premorbid disability • Intravenous alteplase	Intervention: Continue previous antihypertensive medication/s (n=379) <u>Comparator</u> : Stop previous antihypertensive medication/s (n=384)	<u>1° endpoint</u> : Death or major disability (mRS 3–6) at 14 d: RR: 0.86 (95% CI: 0.65–1.14; p=0.3) <u>Safety endpoint</u> : Adverse events, minor and serious: p>0.05 for all	Relevant 2° endpoint• 2-wk NIHSS: p=0.46 and 2-wkBarthel Index: p=0.30• 2-wk BP: significantly lower in the continue arm (mean difference of - 13 mm Hg in SBP and -8 mm Hg in DBP) p<0.0001

CATIS He J, et al., 2014 24240777	Aim: Evaluate whether immediate blood pressure reduction in	Inclusion criteria: • Age >22 y • Acute ischemic stroke within previous	Intervention: Antihypertensive medication to maintain BP <140/90 for the first	<u>1° endpoint</u> : Death or major disability (mRS 3–6) at 14 d: OR: 1.0 (95% CI: 0.88–1.14; p=0.98)	Relevant 2° endpoint • Death or major disability (mRS 3– 5) at 90 d: OR: 0.99 (95% CI: 0.86– 1.15; p=0.93)
	patients with acute ischemic stroke would reduce death and major disability at 14 d or hospital discharge Study type: RCT Size: 4071	24 h Exclusion criteria: • Impaired level of consciousness • Hypertensive emergency • BP >220/120 • Atrial fibrillation • Intravenous alteplase	<u>Comparator</u> : No antihypertensive medication for the first wk (n=2033)	Safety endpoint: • Vascular disease events p=0.28 • Recurrent stroke p=0.07	 Lower blood pressure at 14 d (mean difference of -8.6 mm Hg in SBP and -3.9 mm Hg in DBP; p<0.001) and at 90 d (mean difference of -2.9 mm Hg in SBP and -1.4 mm Hg in DBP; p<0.001) in the active arm <u>Study limitations</u> Antihypertensive regimen was not standardized
					Summary/conclusions • Early treatment of hypertension was safe but ineffective to prevent death or dependency • Early initiation of anti- hypertensives was associated with better BP control at 2 wk
Wang H, et al., 2014 (195) <u>24853087</u>	Aim: To assess the effects of early BP lowering on early and long-term outcomes after acute stroke. Study type: Systematic review and meta-analysis of RCTs. Study size: 17 trials (n=13,236 pts)	Inclusion criteria: Prospective RCTs of pts ≥18 y with acute ischemic or hemorrhagic stroke; intervention compared with placebo was initiated within 7 d of stroke onset; intervention aimed to lower BP or intervention achieved BP reduction;1 or more functional outcomes reported, such as death or dependency.	• Early BP lowering after acute stroke onset compared with placebo	 <u>1° outcomes</u>: Early (within 30 d) and long-term (from 3–12 mo). <u>Key findings</u>: Early BP lowering after acute stroke onset associated with more death within 30 d compared with placebo RR: 1.34; 95% CI: 1.02–1.74; p=0.03. Early BP lowering after acute stroke onset not associated with early neurological deterioration, early death within 7 d, long-term death, early and long-term dependency, early and long-term stroke recurrence, long-term MI and long-term CVE. 	Summary: Results do not support early BP lowering after acute stroke. Early BP lowering may be associated with greater risk of death within 30 d after acute stroke.

Zhao R, et al., 2015 (196) 26061309	Aim: To determine whether lowering BP during the acute phase of an ischemic stroke improves short- and long-term outcomes. Systematic review and meta-analysis of RCTs. Study size: 22 RCTs	Exclusion criteria: Studies with the pts of subarachnoid hemorrhage, studies without available full- text or relevant data, studies about ongoing trials and those written in languages other than English. Inclusion criteria: Pts with acute stroke (ischemic or hemorrhagic) treated with an antihypertensive agent or placebo. Groups: Treatment groups were n=5,672 (range, 6–2,308), and in the control groups was 5,416 (range, 6– 2033). Follow-up: Ranged from 5 d–12 mo	• Early BP lowering after acute stroke onset compared with placebo	 <u>1° outcomes</u>: Change in SBP and DBP after treatment and short- and long-term dependency and mortality rates. <u>Key findings</u>: Treatment groups had a greater decrease in BP than control groups, and this effect was seen with different classes of antihypertensive drugs. Short-term and long-term dependency rates were similar between treatment and control groups (short-term dependency: pooled OR: 1.041; 95% CI: 0.936–1.159; p=0.457; long-term dependency: pooled OR: 1.013; 95% CI: 0.915–1.120; p=0.806). Short-term or long-term mortality was similar between the treatment and control groups (short-term mortality: pooled OR: 1.020 (95% CI: 0.749–1.388; p=0.902); long-term mortality: pooled OR: 1.039 (95% CI: 0.883–1.222; p=0.644). 	Summary: Antihypertensive agents effectively reduce BP during the acute phase of an ischemic stroke, but seem to confer no benefit with regard to short- and long-term dependency and mortality.
Ahmed N, et al., 2000 (197) <u>10835440</u>	<u>Aim</u> : To investigate outcome in INWEST subgroups with increasing levels of BP reduction.	Inclusion criteria: Pts with a diagnosis of ischemic stroke in the carotid artery territory within 24 h.	Interventions: • Nimodipine as IV infusion of 1 mg/h for 5 d followed by oral dose of 120 mg daily for a total treatment period of 21 d (n=101)	 <u>1° outcomes</u>: Neurological outcome per the Orgogozo scale and functional outcome per the Barthel scale at d 21 <u>Key findings</u>: Nimodipine treatment resulted in a significant reduction in BP from baseline vs. placebo during the first few d. 	Summary: • DBP, but not SBP, reduction was associated with neurological worsening after the IV high-dose nimodipine after acute stroke. • For low-dose nimodipine, the results were inconclusive.

	Study type: Post- hoc analysis of RCT Size: 265		 Nimodipine as IV infusion of 2 mg/h for 5 d followed by oral dose of 120 mg daily for a total treatment period of 21 d (n=94) <u>Comparator</u>: Placebo (n=100) 	 A significant correlation between DBP reduction and worsening of the neurological score was found for the high-dose group (beta=0.49; p=0.048). Pts with a DBP reduction of ≥20% in the high-dose group had a significantly increased adjusted OR for death or dependency (n/N=25/26, OR: 10.16; 95% CI: 1.02–101.74) and death alone (n/N=9/26, OR: 4.336; 95% CI: 1.131–16.619) vs. all placebo pts (n/N=62/92 and 14/92, respectively). No correlation between SBP change and outcome. 	
Bath PM, et al., 2014 (198) <u>25353321</u>	Aim: To assess the clinical effectiveness of altering BP in pts with acute stroke, and the effect of different vasoactive drugs on BP in acute stroke. Update of previously published Cochrane reviews (1997, 2001, and 2008). Study type: Meta- analysis of RCTs of interventions that aimed to alter BP vs. control in pts within 1 wk of acute ischemic or hemorrhagic stroke.	Inclusion criteria: RCTs of interventions that aimed to alter BP compared with control in pts with 1 wk of acute ischemic or hemorrhagic stroke	• BP lowering after acute stroke onset compared with placebo	 <u>1° outcome</u>: Functional outcome <u>Key findings</u>: At 24 h after randomization #1: Oral ACEIs reduced SBP MD: -8 mm Hg (95% CI: -17–1) and DBP MD: -3 mm Hg (95% CI: -9–2), sublingual ACEIs reduced SBP MD: - 12.00 mm Hg (95% CI: -26–2) and DBP MD: - 2 (95% CI: -10–6). Oral angiotensin receptor antagonists reduced SBP MD: -1 mm Hg (95% CI: -3–2) and DBP MD: -1 mm Hg (95% CI: -3–2) and DBP MD: -1 mm Hg (95% CI: -3–1). Oral BBs reduced SBP MD: -14 mm Hg (95% CI: -27– -1) and DBP MD: -1 mm Hg (95% CI: -27– -1) and DBP MD: -1 mm Hg (95% CI: -9–7), IV BBs reduced SBP MD: -5 mm Hg (95% CI: -18–8) and DBP MD: -5 mm Hg (95% CI: -13–3). Oral CCBs reduced SBP MD: -13 mm Hg (95% CI: -43–17) and DBP MD: -13 mm Hg (95% CI: -43–17) and DBP MD: -6 mm Hg (95% CI: -31–6). Nitric oxide donors reduced SBP MD: -12 mm Hg (95% CI: -19– -5) and DBP MD: -3 (95% CI: -4– -2). 	 Summary: No current evidence showing that lowering BP during the acute phase of stroke improves functional outcome. It seems reasonable to withhold BP-lowering drugs until pts are medically and neurologically stable, after which drugs can then be reintroduced. CCBs, ACEI, angiotensin receptor antagonists, BBs and nitric oxide donors each lower BP in acute stroke while phenylephrine appears to increase BP.

	Size: 26 trials involving 17,011 pts (8,497 pts were assigned active therapy and 8,514 pts received placebo/control). Not all trials contributed to each outcome.			 Phenylephrine, nonsignificantly increased SBP MD: 21 mm Hg (95% CI: -13–55) and DBP MD: 1 mm Hg (95% CI: -15–16). BP lowering did not reduce death or dependency either by drug class OR: 0.98 (95% CI: 0.92–1.05), stroke type OR: 0.98 (95% CI: 0.92–1.05) or time to treatment OR: 0.98 (95% CI: 0.92–1.05). Treatment within 6 h of stroke appeared effective in reducing death or dependency OR: 0.86 (95% CI: 0.76–0.99) but not death OR: 0.70 (95% CI: 0.38–1.26) by trial end. While death or dependency did not differ between pts who continued pre-stroke antihypertensive treatment vs. those who stopped it temporarily (worse outcome with continuing treatment OR: 1.06; 95% CI: 0.91–1.24), disability scores at the end of the trial were worse in pts randomized to continue treatment_(Barthel Index MD: -3.2 (95% CI: -5.8– -0.6). 	
SITS-ISTR Ahmed N, et al., 2009 (199) <u>19461022</u>	Aim: To determine the association of BP and antihypertensive therapy with clinical outcomes after thrombolysis for acute ischemic stroke Study type: Retrospective analysis of prospectively maintained thrombolysis registry.	Inclusion criteria: • Pts with acute ischemic stroke treated with IV rtPA • BP values were recorded at baseline, 2 h, and 24 h after thrombolysis. <u>Categories</u> : By history of HTN and antihypertensive therapy within 7 d after thrombolysis: • Group 1, HTN treated with antihypertensives (n=5,612)	Various categories of HTN treatments	 <u>1° outcomes</u>: Symptomatic (National Institutes of Health Stroke Scale score deterioration ≥4) ICH Type 2, mortality, and independence at (modified Rankin Score 0 to 2) 3 mo. <u>Key findings</u>: High SBP 2–24 h after thrombolysis as a continuous variable was associated with worse outcome (p<0.001) and as a categorical variable had a linear association with symptomatic hemorrhage and a U- shaped association with mortality and independence with <u>SBP 141–150 mm Hg</u> <u>associated with most favorable outcomes</u>. No difference in symptomatic hemorrhage OR: 1.09 (95% CI: 0.83–1.51; p=0.58) and independence OR: 1.03 (95% CI: 0.93–1.10; p=0.80) but lower mortality OR: 0.82 (95% 	 Summary: Strong association of high SBP after thrombolysis with poor outcome. Higher BPs during the initial 24 h were associated with greater risk of ICH in a linear fashion. U-shaped relation found between BP during initial 24 h and death or dependency at 3 mo, with best outcomes associated with SBP of 141–150 mm Hg.

	<u>Study size</u> : 11,080 pts from 2002–2006.	 Group 2, HTN withholding antihypertensives (n=1,573) Group 3, without history of HTN treated with antihypertensives (n=995) Group 4, without history of HTN not treated with antihypertensives (n=2,632). 		CI: 0.73–0.92; p=0.0007) for Group 1 vs. Group 4. • Group 2 had a higher symptomatic hemorrhage (OR: 1.86; 95% CI: 1.34–2.68; p=0.0004) and mortality (OR: 1.62; 95% CI: 1.41–1.85; p<0.0001) and lower independence (OR: 0.89; 95% CI: 0.80– 0.99; p=0.04) vs. with Group 4. Group 3 had similar results as Group 1.	
ACCESS Schrader J, et al., 2003 (200) <u>12817109</u>	Aim: To assess safety of modest BP reduction by candesartan in early treatment of stroke; and provide an estimate of the number of cases required to perform a larger phase III efficacy study. Study type: Prospective, double-blind, RCT; multicenter phase II study. Size: 342 pts	Inclusion criteria: Motor deficit, a cerebral CT scan excluding ICH, and necessity to treat HTN per prevailing recommendation Exclusion criteria: >85 y, disorders in consciousness preventing acquisition of consent, occlusion or >70% stenosis of the internal carotid artery, malignant HTN, manifest cardiac failure, high-grade aortic or mitral stenosis, UA pectoris, or contraindications against candesartan.	Design: 4 mg candesartan daily or placebo on d 1. On d 2, dosage was increased to 8 or 16 mg candesartan or placebo if BP >60 mm Hg SPB or 100 mm Hg DBP. Treatment was targeted to a 10%–15% BP reduction within 24 h.	<u>1° outcome</u> : Trial was stopped prematurely when 342 pts (339 valid) had been randomized because of an imbalance in endpoints. <u>Key findings</u> : Cumulative 12 mo mortality and the number of vascular events differed significantly in favor of the candesartan cilexetil group (OR: 0.475; 95% CI: 0.252– 0.895).	Summary: Early antihypertensive therapy with candesartan might be a safe therapeutic option in acute stroke, but study sample size very small.
SCAST Sandset EC, et al., 2011 (201) <u>21316752</u>	<u>Aim</u> : To examine whether careful BP-lowering treatment with the candesartan is	Inclusion criteria: Pts >18 y with acute stroke (ischemic or hemorrhagic) and SBP of ≥140 mm Hg were	Design: Pts randomized to candesartan (n=1,017) or placebo (1,012) (1:1) for 7 d, with doses	<u>1° effect variables</u> : Composite of vascular death, MI, or stroke during the first 6 mo; and functional outcome at 6 mo, as measured by the modified Rankin Scale.	Relevant 2° endpoint:• Similar effects for all prespecified2° endpoints.• During follow-up, 9 (1%) pts on candesartan and 5 (<1%) on

	beneficial in pts with acute stroke and raised BP. <u>Study type:</u> Double-blind RCT <u>Study size</u> : 2,029 pts	included within 30 h of symptom onset.	increasing from 4 mg on d 1–16 mg on d 3–7.	 Data for status at 6 mo were available for 2,004 pts (99%; 1,000 candesartan, 1,004 placebo). <u>Key findings:</u> BPs significantly lower in pts allocated candesartan vs. placebo (mean 147/82 mm Hg [SD 23/14] in the candesartan group on d 7 vs. 152/84 mm Hg [22/14] in the placebo group; p<0.0001). Risk of the composite vascular endpoint did not differ between treatment groups (candesartan, 120 events, vs. placebo, 111 events; adjusted HR: 1.09; 95% CI: 0.84–1.41; p=0.52. Analysis of functional outcome suggested a higher risk of poor outcome in the candesartan group (adjusted OR: 1.17; 95% CI: 1.00–1.38; p=0.048. 	placebo had symptomatic hypotension, and renal failure was reported for 18 (2%) pts taking candesartan and 13 (1%) allocated placebo. Summary: Careful BP-lowering treatment with candesartan was not beneficial in pts with acute stroke and raised BP. Indeed, there was the suggestion of a harmful effect.
CATIS He J, et al., 2014 (202) <u>24240777</u>	Aim: To evaluate whether immediate BP reduction in pts with acute ischemic stroke would reduce death and major disability at 14 d or hospital discharge. Single-blind, blinded end-points RCT. Study size: 4,071 pts	Inclusion criteria: Pts with nonthrombolysed ischemic stroke within 48 h of onset and elevated SBP	Design: Pts (n=2,038) randomized to antihypertensive treatment (aimed at lowering SBP by 10% to 25% within first 24 h, achieving BP <140/90 mm Hg within 7 d, and maintaining this level during hospitalization) vs. to discontinue all antihypertensive medications (control) during hospitalization (n=2,033).	 <u>1° outcome</u>: Combination of death and major disability (modified Rankin Scale score ≥3) at 14 d or hospital discharge. <u>Key findings</u>: Mean SBP was reduced from 166.7 mm Hg to 144.7 mm Hg (-12.7%) within 24 h in the antihypertensive treatment group and from 165.6 mm Hg to 152.9 mm Hg (-7.2%) in the control group within 24 h after randomization (difference, -5.5% (95% CI: -4.96.1%); absolute difference, -9.1 mm Hg (95% CI: -10.28.1), p<0.001). 1° outcome did not differ between treatment groups (OR: 1.00; 95% CI: 0.88-1.14; p=0.98) at 14 d or hospital discharge. BP at 14 d and 90 d: significantly lower in the active arm (mean difference of -2.9 mm Hg in systolic BP and -1.4 mm Hg in diastolic BP) 	 <u>Relevant 2° endpoint:</u> Death and major disability at 3-mo posttreatment follow-up did not differ between treatment groups (500 events [antihypertensive treatment] vs. 502 events [control]; OR: 0.99; 95% CI: 0.86–1.15; p=0.93). <u>Summary</u>: Among pts with acute ischemic stroke, BP reduction with antihypertensive medications, vs. absence of hypertensive medication, did not reduce the likelihood of death and major disability at 14 d or hospital discharge. Early initiation of antihypertensives was associated with better BP control at 2 wk

COSSACS	Aim: To assess	Inclusion criteria: Pts	Design: Continue	19 outcomer Death or dependency at 2 wk	Summary
Robinson TG, et al.,	the efficacy and	>18 y taking	(n=379) or stop (n=384)	1° outcome: Death or dependency at 2 wk.	Summary: • Continuation of antihypertensive
2010 (203)	safety of	antihypertensive drugs	pre-existing	Key findings:	drugs did not reduce 2-wk death or
20621562	continuing or	enrolled within 48 h of	antihypertensive drugs	• 72 of 379 pts in the continue group and 82	dependency, CV event rate, or
20021002	stopping pre-	stroke and last dose of	for 2 wk.	of 384 pts in the stop group reached the 1°	mortality at 6 mo
	existing	antihypertensive drug.		endpoint RR: 0.86; 95% CI: 0.65–1.14;	Early reinitiation of
	antihypertensive	antinypertensive urug.		p=0.3.	antihypertensives was associated
	drugs in pts who			 Difference in SBP at 2 wk between the 	with better BP control at 2 wk
	recently had a			continue group and the stop group was 13	 Lower BP levels in those who
	stroke.			mm Hg (95% CI: 10–17) and the difference	continued antihypertensive
				in DBP was 8 mm Hg (6–10; difference	treatment after acute mild stroke
	Study type:			between groups; p<0.0001).	were not associated with an
	Multicenter,			 No substantial differences were observed 	increase in adverse events.
	prospective,			between groups in rates of serious adverse	 Of note, COSSACS was likely
	randomized, open,			events, 6-mo mortality, or major CV events.	underpowered due to early
	blinded-endpoint				termination of the trial.
	trial.				
	Study size:				
	763 pts		.		
CHHIPS	Aim: To assess	Inclusion criteria: Pts	Design:	<u>1° outcome:</u> Death or dependency at 2 wk.	Summary:
Potter JF, et al.,	feasibility, safety,	with cerebral infarction	Within 36 h of		Labetalol and lisinopril are
2009 (204) 19058760	and effects of 2	or cerebral	symptom onset: #1: Oral labetalol,	Key findings:	effective antihypertensive drugs in acute stroke that do not raise risk of
19030700	regimens for lowering BP in pts	hemorrhage who were hypertensive SBP	lisinopril vs. placebo if	• 1° outcome occurred in 61% (69) of the particular $E0\%$ (25) of the placebe group	serious adverse events.
	who with acute	>160 mm Hg)	they were	active vs. 59% (35) of the placebo group (RR: 1.03; 95% CI: 0.80–1.33; p=0.82)	 Early lowering of BP with lisinopril
	stroke.	>100 min rig)	nondysphagic;	 No evidence of early neurological 	and labetalol after acute stroke may
	50000		#2: IV labetalol,	deterioration with active treatment (RR: 1.22;	be a promising approach to lower
	Study type:		sublingual lisinopril, or	95% CI: 0.33–4.54; p=0.76) despite greater	mortality and disability.
	Double-blind pilot		placebo if they had	drop in SBP within the first 24 h in this group	 However, pilot nature and very
	trial.		dysphagia.	vs. placebo (21 [17–25] mm Hg vs. 11 [5–17]	small sample size limit
			 Labetalol (n=58), 	mm Hg; p=0.004).	generalizability.
	Study size: 179		lisinopril (n=58), or	 No rise in serious adverse events with 	S
	pts		placebo (n=63).	active treatment (RR: 0.91; 95% CI: 0.69-	
			 Doses were titrated 	1.12; p=0.50) but 3-mo mortality was halved	
			up if target BP was not	(9.7% vs. 20.3%; HR: 0.40; 95% CI: 0.2–1.0;	
			reached.	p=0.05).	
Bath PM, et al.,	Aim: To assess	Inclusion criteria: Pts	Design:	1° outcome: Function, assessed with the	Summary:
2015 (205)	outcomes after	admitted to hospital	 7 d of transdermal 	modified Rankin Scale at 90 d	 In pts with acute stroke and high
25465108	stroke in pts given	with an acute ischemic	glyceryl trinitrate (5 mg		BP transdermal glyceryl trinitrate

	drugs to lower their BP. Study type: Multicenter, randomized partial- factorial trial Study size: 4,011 pts	or hemorrhagic stroke and raised SBP (140– 220 mm Hg)	 per d), started within 48 h of stroke onset vs. No glyceryl trinitrate (control group). Pts taking antihypertensive drugs before index stroke randomly assigned to continue vs. stop taking these drugs. 	 Key findings: Mean BP was 167 (SD: 19) mm Hg/90 (13) mm Hg at baseline (median 26 h (16– 37) after stroke onset), and was significantly reduced on d 1 in 2,000 pts allocated to glyceryl trinitrate vs. 2,011 controls (difference -7.0 (95% CI: -8.5– -5.6) mm Hg/- 3.5 [-4.4– -2.6] mm Hg; both p<0.0001), and on d 7 in 1,053 pts allocated to continue antihypertensive drugs compared with 1,044 pts randomized to stop them (difference: -9.5 (95% CI: -11.8– -7.2) mm Hg/-5.0 [-6.4– -3.7] mm Hg; both p<0.0001). D-90 functional outcome did not differ in either treatment comparison-glyceryl trinitrate vs. no glyceryl trinitrate (OR: 1.01; 95% CI 0.91–1.13; p=0.83), and with continue vs. stop antihypertensive drugs (OR: 1.05; 95% CI: 0.90–1.22; p=0.55). 	 lowered BP with acceptable safety but did not improve functional outcome. Continuing prestroke antihypertensive drugs in acute stroke pts in the first few d did not confer benefit.
ATACH-1 2010 (192) <u>19770736</u>	Aim: To determine the feasibility and acute (i.e., within 72 h) safety of 3 levels of SBP reduction in subjects with supratentorial ICH treated within 6 h after symptom onset. Study type: Phase I, dose- escalation, multicenter prospective study. Study size: 60	Inclusion criteria: Pts with ICH with elevated SBP ≥170 mm Hg who presented to the ED within 6 h of symptom onset.	 Design: IV nicardipine to reduce SBP to a target of: #1: 170-200 mm Hg in the first cohort of pts #2: 140-170 mm Hg in the 2nd cohort #3: 110-140 mm Hg in the third cohort. Each subject was followed-up for 3 mo to preliminarily assess mortality and the clinical outcomes. A total of 18, 20, and 22 pts were enrolled in the respective 3 tiers of SBP treatment goals. 	 <u>1° outcome</u>: Treatment feasibility (achieving and maintaining the SBP goals for 18–24 h) <u>2° outcomes</u>: <u>#1</u>: Neurologic deterioration within 24 h; <u>#2</u>: Serious adverse events within 72 h. <u>Key findings</u>: Overall, 9 of 60 pts had treatment failures (all in the last tier). A total of 7 subjects with neurologic deterioration were observed: 1 (6%), 2 (10%), and 4 (18%) in tier 1, 2, and 3, respectively. Serious adverse events were observed in1 subject (5%) in tier 2 and in 3 subjects (14%) in tier 3. However, the safety stopping rule was not activated in any of the tiers. 3 (17%), 2 (10%), and 5 (23%) subjects in tiers1, 2, and 3, respectively, died within 3 mo 	 Summary: Observed proportions of neurologic deterioration and serious adverse events were below the prespecified safety thresholds, and the 3-mo mortality rate was lower than expected in all SBP tiers. Results formed the basis of an ongoing larger randomized trial (ATACH-2) addressing the efficacy of SBP reduction in pts with ICH.

INTERACT-1 Anderson CS, et al., 2008 (193) <u>18396107</u>	Aim: To assess the safety and efficiency of this treatment, as a run-in phase to a larger trial. Study type: Randomized pilot trial Study size: 404	Inclusion criteria: Pts with acute spontaneous ICH diagnosed by CT within 6 h of onset, elevated SBP (150- 220 mm Hg), and no definite indication or contraindication to treatment	Design: Early intensive lowering of BP (target SBP 140 mm Hg; n=203) vs. standard guideline-based management of BP (target SBP 180 mm Hg; n=201).	<u>1° outcome</u> : Proportional change in hematoma volume at 24 h. <u>2° outcomes</u> : Measurements of hematoma volume. <u>Safety and clinical outcomes</u> : Assessed for up to 90 d. <u>Key findings</u> : • Mean hematoma volumes were smaller in the guideline group (12.7 mL, SD 11.6) than in the intensive group (14.2 mL, SD 14.5). • From randomization to 1 h, mean SBP was 153 mm Hg in the intensive group and 167 mm Hg in the guideline group (difference 13.3 mm Hg (95% CI: 8.9–17.6) mm Hg; p<0.0001); from 1 h to 24 h, BP was 146 mm Hg in the intensive group and 157 mm Hg in the guideline group (10.8 mm Hg; 95% CI: 7.7–13.9 mm Hg; p<0.0001). • Mean proportional hematoma growth was 36.3% in the guideline group and 13.7% in the intensive group (difference 22.6%; 95% CI: 0.6%–44.5%; p=0.04) at 24 h. • After adjustment for initial hematoma volume and time from onset to CT, median hematoma growth differed between the groups with p=0.06; the absolute difference in volume between groups was 1.7 mL (95% CI: -0.5–3.9; p=0.13). RR of hematoma growth ≥33% or ≥12.5 mL was 36% lower (95% CI: 0%–59%; p=0.05) in the intensive group than in the guideline group. Adjusted RR: 8% (95% CI: -1.0%–17%; p=0.05). • Intensive BP-lowering treatment did not alter the risks of adverse events or 2° clinical outcomes at 90 d.	Summary: Early intensive BP- lowering treatment is clinically feasible, well tolerated, and appears to reduce hematoma growth in ICH.
Hack W, et al., 2008 (206)	Aim: To assess the efficacy and	Inclusion criteria: Pts 18–80 y, who had	<u>Design</u> :	<u>1° outcome</u> : Disability at 90 d, dichotomized as a favorable outcome (a score of 0 or 1 on	<u>Summary</u> : Compared with placebo, IV alteplase administered between 3

<u>18815396</u>	safety of alteplase administered between 3 and 4.5 h after the onset of a stroke. <u>Study type</u> : RCT <u>Study size</u> : 821 pts	received a clinical diagnosis of acute ischemic stroke, and were able to receive the study drug within 3–4 h after the onset of symptoms. Exclusion criteria: SBP >185 mm Hg or DBP >110 mm Hg or aggressive treatment (IV medication) necessary to reduce BP to these limits	 Eligible pts were randomly assigned 1:1 to receive 0.9 mg of alteplase per kg, administered IV (with an upper limit of 90 mg), or placebo. 418 pts were assigned to receive alteplase and 403 pts were assigned to receive placebo 	 the modified Rankin scale, which has a range of 0–6, with 0 indicating no symptoms at all and 6 indicating death) or an unfavorable outcome (a score of 2–6 on the modified Rankin scale). <u>2° outcome:</u> global outcome analysis of 4 neurologic and disability scores combined. <u>Safety outcomes:</u> death, symptomatic intracranial hemorrhage, and other serious adverse events. <u>Key findings:</u> More pts had a favorable outcome with alteplase than with placebo (52.4% vs. 45.2%; OR: 1.34; 95% CI: 1.02–1.76; p=0.04. Incidence of ICH was higher with alteplase than with placebo (for any ICH, 27.0% vs. 17.6%; p=0.001; for symptomatic intracranial hemorrhage, 2.4% vs. 0.2%; p=0.08). Mortality did not differ significantly between the alteplase and placebo groups (7.7% and 8.4%, respectively; p=0.68). No significant difference in the rate of other serious adverse events. 	and 4.5 h after the onset of symptoms significantly improved clinical outcomes in pts with acute ischemic stroke; alteplase was more frequently associated with symptomatic ICH.
NINDS rt-PA Stroke Study Group, 1995 (207) <u>7477192</u>	Aim: To assess the difference in clinical efficacy between IV t-PA and placebo among pts with an acute ischemic stroke Study type: Double-blind RCT	Inclusion criteria: Pts with an ischemic stroke with a clearly defined time of onset (<3 h), a deficit measurable on the NIH stroke scale, and a base-line CT scan of the brain that showed no evidence of ICH. Exclusion criteria:	Design: RCT with acute ischemic stroke pts randomized to t-PA vs. placebo	 <u>1° outcome</u>: Clinical outcome at 3 mo, according to scores on the Barthel index, modified Rankin scale, Glasgow outcome scale, and NIH stroke scale: <u>Key findings</u>: As compared with pts given placebo, pts treated with t-PA were at least 30% more likely to have minimal or no disability at 3 mo on the assessment scales. Symptomatic ICH within 36 h after the onset of stroke occurred in 6.4% of pts given 	Summary : Despite an increased incidence of symptomatic ICH, treatment with IV t-PA within 3 h of the onset of ischemic stroke improved clinical outcome at 3 mo

Γ	Study size: 624	SBP >185 mm Hg or	t-PA but only 0.6% of pts given placebo
	pts	DBP >110 mm Hg	(p<0.001).
		Ŭ	 Mortality at 3 mo was 17% in the t-PA
			group and 2% in the placebo group (p=0.30).

Data Supplement 43. RCTs Comparing Secondary Stroke Prevention (Section 9.4.3)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Post-stroke Antihypertensive Treatment Study (PATS) 1995 (208) <u>8575241</u>	Aim: To assess whether lowering BP prevents the recurrence of stroke in Chinese pts with history of cerebrovascular disease <u>Study type</u> : Double- blind RCT <u>Size</u> : 5,665 pts	Inclusion criteria: Pts with history of stroke or TIA <u>Exclusion</u> criteria: N/A	Intervention: Indapamide 2.5 mg daily (n=2,840 pts) <u>Comparator</u> : Placebo (n=2,825 pts)	<u>1° outcome</u> : Recurrence of fatal or nonfatal stroke. <u>Key findings</u> : Average SBP/DBP at randomization was 153.8/92.8 mm Hg. At median follow-up (2 y), BP was 6.8/3.3 mm Hg lower in pts on active treatment. 143 pts on indapamide vs. 219 pts on placebo had recurrent strokes (HR: 0.69; 95% CI: 0.54–0.89; p<0.001).	 <u>2° outcome</u>: Major fatal and nonfatal CV events In addition, 199 pts on indapamide and 258 pts on placebo had a CV event (HR: 0.75; 95% CI: 0.89–0.62; p=0.002). 2,825 pts received a placebo and 2,840 pts received. <u>Summary</u>: For pts with a history of stroke or TIA, BP reduction of 5/2 mm Hg with 2.5 mg of indapamide lowered the first incidence of fatal and nonfatal stroke by 29%, with 3-y absolute benefit of 29 events per 1,000 pts.
PROGRESS 2001 (209) <u>11589932</u>	<u>Aim</u> : To determine effects of a BP- lowering regimen in hypertensive and nonhypertensive pts with a history of stroke or TIA. <u>Study type</u> : Double- blind, placebo- controlled trial <u>Size</u> : 6,105	Inclusion criteria: Pts with history of stroke (evidence of an acute disturbance of focal neurological function with symptoms lasting more than 24 h and	Intervention: Active treatment comprised a flexible regimen based on the ACEI perindopril (4 mg daily), with addition of diuretic indapamide at discretion of treating physicians (n=3,051) <u>Comparator</u> : Placebo (n=3,054)	 <u>1° outcome</u>: Total stroke (fatal or nonfatal) <u>Key findings</u>: Over 4 y of follow-up, active treatment reduced BP by 9/4 mm Hg. 307 (10%) pts assigned active treatment suffered a stroke, vs. 420 (14%) assigned placebo (RR reduction: 28% (95% CI: 17, 38), p<0.0001). Combination therapy with perindopril plus indapamide reduced BP by 12/5 mm Hg and stroke risk by 43% (95% CI: 30%–54%). Single-drug therapy reduced 	Relevant 2° endpoint:Active treatment also reduced the risk of total major vascular events (26% [16–34]).There were similar reductions in the risk of stroke in hypertensive and nonhypertensive subgroups (all p<0.01).

		thought to be due to ICH or ischemia) or TIA within the previous 5 y. <u>Exclusion</u> <u>criteria</u> : N/A • Pts clinically stable for at least 2 wk after their most recent vascular event before entry to the study.		BP by 5/3 mm Hg and produced no discernable reduction in the risk of stroke.	reductions and larger risk reductions than single drug therapy with perindopril alone. • This trial showed the benefits of BP lowering in both hypertensive pts. However, based on older definitions, presence of baseline HTN in the trial was defined as ≥160/90 mm Hg.
MOSES Schrader J, et al., 2005 (210) <u>15879332</u>	Aim: To assess among hypertensive stroke pts, whether for the same level of BP control, eprosartan would be more effective than nitrendipine in reducing cerebrovascular and CV morbidity and mortality. Study type: PROBE design Size: 1,405	Inclusion <u>criteria</u> : High- risk hypertensives with cerebral event during the last 24 mo (proven by cerebral CT scan or nuclear magnetic resonance) <u>Exclusion</u> <u>criteria</u> : Internal carotid artery occlusion or stenosis >70%, manifest HF (NYHA grade III–IV), age >85 y at the time of	Intervention: Eprosartan 600 mg (n=681) Comparator: Nitrendipine 10 mg (n=671)	<u>1° endpoint</u> : Composite of total mortality and all CV and cerebrovascular events, including all recurrent events. <u>Key findings</u> : BP reduced to comparable extent without significant differences between 2 groups during study period (150.7/84 mm Hg vs. 152.0/87.2 mm Hg with eprosartan and nitrendipine therapy to 137.5/80.8 mm Hg and 136.0/80.2 mm Hg, respectively). 75.5% reached values <140/90 mm Hg with eprosartan regimen and 77.7% with nitrendipine. During follow-up, 461 1° events occurred: 206 eprosartan and 255 nitrendipine (IDR: 0.79; 95% CI: 0.66– 0.96; p=0.014.	 <u>Relevant 2° endpoint:</u> CV events were: 77 eprosartan and 101 nitrendipine (IDR: 0.75; 95% CI: 0.55–1.02; p=0.06); cerebrovascular events: 102 eprosartan and134 nitrendipine (IDR: 0.75; 95% CI: 0.58–0.97; p=0.03). <u>Summary</u>: The combined 1° endpoint was significantly lower in the eprosartan group. However, it was a reduction in TIAs that accounted for most of the benefit in cerebrovascular events, with no significant difference in ischemic strokes. Also a more traditional analysis of time to first cerebrovascular event did not show a benefit of eprosartan.

PROFESS Yusuf S, et al., 2008 (211) <u>18753639</u> SPS-3	Aim: To evaluate the effects of therapy with an ARB, telmisartan, initiated early after a stroke Study type: Double- blind RCT Size: 20,332 pts	the cerebrovascula r event, pts treated with anticoagulants for a cardiac arrhythmia, high-grade aortic or mitral valve stenosis, or UA pectoris. Inclusion criteria: Pts ≥55 y with an ischemic stroke <90 d before randomization Exclusion criteria: 1° hemorrhagic stroke, severe disability after the qualifying stroke	Intervention: Telmisartan 80 mg daily (n=10,146) Comparator: Placebo (n=10,186) Intervention: SBP	<u>1° endpoint</u> : Recurrent stroke <u>Key findings</u> : During mean follow-up of 2.5 y, mean BP was 3.8/2.0 mm Hg lower in telmisartan group vs. placebo group. 880 pts (8.7%) in telmisartan group vs. 934 pts (9.2%) in placebo group had a subsequent stroke (HR: 0.95; 95% CI: 0.86–1.04; p=0.23). 1° outcome: All stroke (including	Relevant 2° endpoint: Major CV events (death from CV causes, recurrent stroke, MI, or new or worsening HF) occurred in 1,367 pts (13.5%) in telmisartan group vs. 1,463 pts (14.4%) in placebo group (HR: 0.94; 95% CI: 0.87–1.01; p=0.11). Summary: • Therapy with telmisartan initiated soon after ischemic stroke and continued for 2.5 y did not significantly lower Rate of recurrent stroke, or major CV events. • Impact of treatment with telmisartan may have been affected by the high rate of discontinuation of treatment medication because of hypotensive symptoms, syncope, diarrhea, and nausea experienced in the telmisartan arm and the more aggressive treatment with other standard antihypertensive therapies in the placebo arm. Thus, adverse side effects from treatment medication adherence after stroke. 2° outcomes: No difference between target
Benavente OR, et al., 2013 (212) <u>23726159</u>	effects of different BP targets on rate of recurrent stroke in pts	<u>criteria</u> : Pts with recent, MRI-defined symptomatic	target of 130–149 mm Hg (n=1,519)	ischemic strokes and intracranial hemorrhages).	<u>2° outcomes</u> : No unerence between larger groups in disabling or fatal stroke 0.81, (95% Cl: 0.53–1.23; p=0.32) or composite outcome of MI or vascular death 0.84 (95% Cl: 0.68–1.04; p=0.32). However,

W	vith recent lacunar	lacunar	Comparator: SBP	 After 1 y, mean SBP was 138 mm Hg 	hemorrhagic stroke occurred in 16 pts
	troke.	infarctions.	target of <130 mm Hg (n=1,501)	(95% CI: 137–139) in the higher-target group and 127 mm Hg (95% CI: 126–	assigned to the higher-target group (0.29% per y) vs. 6 assigned to the lower-target
R. Ia	Study type: Randomized open- abel trial Size: 3,020 pts	Exclusion criteria: Pts with cortical strokes, cardioembolic disease, or carotid stenosis were excluded.	ng (n= 1,55 i)	 glodp and 127 mining (70% of 120 128) in the lower-target group. Recurrent stroke was observed in 152 pts assigned to higher-target group (2.8% per y) vs. 125 assigned to the lower- target group (2.3% per y; HR: 0.81; 95% Cl: 0.64–1.03). 	group (0.11% per y; HR: 0.37 (95% CI: 0.15– 0.95). Serious complications of hypotension were observed in 15 pts assigned to the higher-target group (0.26% per y) and 23 assigned to the lower-target group (0.40% per y; HR: 1.53; 95% CI: 0.80–2.93). <u>Summary</u> : Use of a SBP target of less than 130 mm Hg was not significantly better than a target of 130–149 mm Hg for preventing any recurrent stroke. However, the lower target appeared to confer benefit for prevention of hemorrhagic stroke.

Data Supplement 44. Nonrandomized Trials, Observational Studies, and/or Registries of Secondary Stroke Prevention (Section 9.4.3)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Rashid P, et al., 2003 (213) <u>14576382</u>	Study type: Meta- analysis of RCTs Size: 7 RCTs	Inclusion criteria: Pts with a history of ischemic stroke, TIA, or ICH Exclusion criteria: N/A	<u>1° outcome</u> : Recurrent stroke <u>Key findings</u> : Antihypertensive drug therapy associated with a 24% reduction in recurrent stroke risk (RR: 0.76; 95% CI: 0.63–0.92) Recurrent stroke risk reduction seen in both hypertensive and normotensive (as defined by the respective trials) pts and linked to magnitude of reduction in SBP	2° outcomes: Nonfatal stroke OR: 0.79 (95% CI: 0.65–0.95), MI OR: 0.79 (95% CI: 0.63, 0.98), and total vascular events OR: 0.79 (95% CI: 0.66– 0.95). No effect seen on vascular or all-cause mortality. ACEIs and diuretics separately, and particularly together, reduced vascular events, while beta-receptor antagonists had no discernable effect. Summary: Use of antihypertensive agents to lower BP for the prevention of vascular events in pts with previous stroke or TIA is efficacious.
Lakhan SE, et al., 2009 (214) <u>19843330</u>	<u>Aim</u> : To examine the role of BP reduction using antihypertensive	Inclusion criteria: Pts with a history of ischemic stroke, TIA, or ICH Exclusion criteria: N/A	<u>1° outcome</u> : Recurrent stroke BP-lowering agents reduced recurrent stroke OR: 0.71 (95% CI: 0.59–0.86; p=0.0004) and	<u>2°outcomes</u> : BP-lowering agents did not affect the rate of MI or all-cause mortality.

	agents to prevent recurrent stroke. Systematic review and meta-analysis Size: 10 RCTs		CV events OR: 0.69 (95% CI: 0.57–0.85; p=0.0004) in pts with a prior stroke or TIA.	<u>Summary</u> : BP lowering agents reduced the occurrence of subsequent stroke and CV events. Rate of MI and all-cause mortality was unchanged.
Liu L, et al., 2009 (215) <u>19798097</u>	Aim: To examine role of BP reduction using antihypertensive agents to prevent recurrent stroke. Study type: Systematic review and meta-analysis Size: 10 RCTs	Inclusion criteria: Pts with a history of ischemic stroke, TIA, or ICH Followed up 2 to 5 y. Exclusion criteria: N/A	<u>1° outcome</u> : Recurrent stroke <u>Key findings</u> : Antihypertensive drugs associated with significant reduction in recurrent strokes (RR: 0.78; 95% CI: 0.68– 0.90). Impact of antihypertensive treatment after ischemic stroke was similar in a restricted group of subjects with HTN and when all subjects, including those with and without HTN, were included. Pooled OR: 0.63 (95% CI: 0.54–0.73; p<0.0001) for trials involving diuretics as a component of therapy and 0.93 (95% CI: 0.87– 1.01; p=0.086) for trials in which treatment included renin system inhibitors (p<0.0001 for heterogeneity).	<u>2° outcomes</u> : Significant reduction in recurrent stroke seen with diuretics (alone or in combination with ACEIs) but not with renal artery stenosis inhibitors, BBs, or CCBs used alone; however, statistical power was limited, particularly for the assessment of BBs and CCBs. <u>Summary</u> : In conclusion, BP lowering by indapamide treatment reduced the recurrence of stroke and the incidence of CV events in Chinese pts with cerebrovascular disease. Whether prevention of stroke recurrence depends on drug class, degree of BP lowering or both requires further investigation.
Lee M, et al., 2012 (216) <u>21796663</u>	Aim: To compare impact of achieving tight vs. usual SBP control on stroke prevention <u>Study size</u> : 11 studies with 42,572 pts and 794 stroke events.	Inclusion criteria: (1) Achieved SBP<130 mm Hg in an active treatment group and SBP 130 to 39 mm Hg in a comparator group by trial; (2) trial duration at least 6 mo; (3) total pts and number of stroke events reported separately for active treatment and comparator groups. Exclusion criteria: (1) Nonrandomized trials; (2) trials in which either the	 <u>1° outcome</u>: Association of future stroke risk and achieved level of different SBP (intensive vs. usual) <u>Key findings</u>: Final SBPs, weighted for trial size, were a mean of 126.5 mm Hg in the intensive treatment arms and 132.6 mm Hg in the conventional arms (mean SBP reduction, 6.1 mm Hg). In subgroup analyses, those with established (symptomatic) CVD at entry did not experience stroke risk reduction with tight control (0.92; 95% CI: 0.83–1.03). 	Summary: Achieving an SBP <130 mm Hg vs. 130–139 mm Hg appears to provide additional stroke protection only among pts with risk factors but no established CVD.

		comparator or the active therapy		
		group received additional treatment		
		that other group did not; (3)		
		majority of participants had		
		ESRD;		
		(4) <10 stroke events in a trial,		
		because stroke was not a major		
		endpoint; (5) SBP not		
		significantly different between		
		active and comparator groups at		
		trial end; (6) Achieved SBP<130		
	Aim. To susheets	mm Hg in a comparator group.	10 automa Malanua da constructor for t	Summony Treatment with an ACEL or ADD has a
Lee M, et al., 2012 (217)	Aim: To_evaluate whether use of	Inclusion criteria: (1) RCT design; (2) pts had a history of	<u>1° outcome</u> : Major vascular event (nonfatal stroke, nonfatal MI, or death from CV causes)	Summary: Treatment with an ACEI or ARB has a clear but rather modest effect on reducing vascular
22052520	ACEIs or ARB	stroke or TIA; (3) active	or stroke (ischemic or hemorrhagic)	risk in persons with prior stroke.
22032320	reduces future	treatment consisted of ACEIs or	of stroke (ischernic of hemorragic)	
	vascular events in	ARBs; (4) follow-up duration at	Key findings: Use of ACEIs or ARBs in	
	persons with prior	least 6 mo; (5) total pts and	persons with prior stroke was associated with	
	stroke.	number of future major vascular	lower risks of future major vascular events RR:	
	Size: 8 RCTs with	events and/or recurrent stroke	0.91 (95% CI: 0.87–0.97; p=0.001); NNT=71	
	29,667 pts	were reported separately for	and recurrent stroke RR: 0.93 (95% CI: 0.86-	
		active treatment and comparator	0.99; p=0.03); NNT=143.	
		groups.		
		Exclusion criteria: (1)		
		mandatory ACEI or ARB use in		
		control groups; (2) study		
		purpose was to examine efficacy		
		of ACEIs or ARBs in pts with		
		acute stroke		
Arima H, et al.,	<u>Aims</u> :	Inclusion criteria: Pts with	1° outcome: Total stroke (fatal or nonfatal)	Summary:
2006 (218)	<u>#1</u> : To investigate	history of cerebrovascular event		• These analyses provide no evidence of a J-curve
<u>16685221</u>	the effects of	(stroke or TIA) within the	Key findings:	relationship between BP level and stroke risk
	randomized	previous 5 y	• Smaller BP differences between active vs.	among pts with cerebrovascular disease. However,
	treatment on	Crowney	placebo groups (p<0.0001) and corresponding	ischemic stroke, TIA, and hemorrhagic pts were all
	recurrent stroke by baseline BP levels	Groups: Defined by baseline BP of <120,	lesser	enrolled and within 5 y of the index event suggesting that these pts were generally
	#2: To investigate	120–139, 140–159, and 160 mm	risk reductions (p trend=0.05) with lower baseline BPs	neurologically stable and not acknowledging the
	association	Hg or greater	Association of stroke incidence with achieved	neuroiogically stable and not acknowledging the
	433001411011	ing of greater	Association of stroke incluence with achieved	

	between achieved follow-up BP levels and recurrent stroke risk. <u>Study type:</u> Post-hoc analysis of PROGRESS trial. <u>Size</u> : 6,105 pts		follow-up SBP level was strong and continuous with no evidence of a J-curve in the range of achieved follow-up SBP from 112–168 mm Hg (p trend <0.0001 RR of study treatment on the discontinuation of randomized treatment increased progressively across the subgroups with lower baseline SBP levels at entry (p trend=0.04), but there was no corresponding difference in effects of randomized treatment on the risks of death or hospital admission (both p trend >0.2) or hypotension, renal dysfunction, electrolyte disturbance, hip fracture, or depression between pts with different levels of baseline BP at baseline (all p trend >0.1) • Minor side-effects were progressively more common at lower BP levels	differences in pathophysiologic mechanism between stroke types. • First analysis showed that the effectiveness of antihypertensive treatment for 2° stroke prevention diminished as baseline BP declined (relative RRs were 39%, 31%, 14%, and 0%, respectively, in the groups defined previously). This trend of decreasing effect was despite successful reduction of mean SBP in each active-treatment group compared with placebo (11.1, 9.2, 7.6, and 7.4 mm Hg reductions, respectively, in the groups defined previously). Also of note, 40% of pts with a baseline BP<140 mm Hg were taking antihypertensive therapy at baseline.
White CL, et al., 2015 (219) <u>25850462</u>	Aim: To determine safety and tolerability of lowering BP in older adults with lacunar stroke Study type: Post- hoc analysis of randomized trial Study Size: 494 pts	Inclusion criteria: Pts with lacunar stroke ≥75 y	 (p homogeneity=0.04). <u>1° outcome</u>: Rates of side effects related to lowering SBP <u>2° outcome</u>: Stroke recurrence and death from vascular causes <u>Key findings</u>: Older pts achieved SBP levels similar to younger pts (mean SBP of 125 mm Hg in lower SBP target group and 137 mm Hg in higher target group) 3.5 y of follow-up 21% reported dizziness and 15% reported lightheadedness when standing; only significant difference between younger and older groups was unsteadiness when standing (23% vs. 32%, p<0.001). No difference in recurrent stroke by target SBP level among the older subjects (HR: 1.01; 95% CI: 0.59–1.73), but the 	Summary: Pts ≥75 y with a recent lacunar stroke who achieved a lower SBP target (<130 mm Hg) were significantly more likely to report unsteadiness on standing than their younger counterparts. Lower SBP was not related to a decrease in recurrent stroke risk in elderly pts with lacunar stroke but there was a potential protective advantage from vascular death.

Ovbiagele B, et al., 2011 (220) <u>22089721</u>	<u>Aim</u> : To assess the association of maintaining low-normal vs. high-	Inclusion criteria: Pts 55 y or older with an ischemic stroke <90 d before randomization	lower target SBP group in older pts was linked to a significant reduction in vascular death (HR: 0.42; 95% CI: 0.18–0.98; p=0.049). <u>1° outcome</u> : First recurrence of stroke of any type 2° outcome : Composite of stroke, MI, or death	<u>Relevant 2° endpoint:</u> Compared with pts in the high-normal SBP group, the risk of 2° outcome was higher for pts in the very low-normal SBP group AHR: 1.31 (95% CI: 1.13–1.52), in the low-normal
	normal SBP levels with risk of recurrent stroke.	<u>Categories</u> : Based on mean SBP level was very low-normal (<120 mm Hg), low-normal (120≤130 mm Hg), high-normal (130≤140 mm Hg), high	from vascular causes. <u>Key findings</u> : Recurrent stroke rates were 8.0% (95% CI: 6.8%–9.2%) for the very low- normal SBP level group, 7.2% (95% CI: 6.4%–	SBP group AHR: 1.16 (95% CI: 1.03–1.31), in the high SBP group AHR: 1.24 (95% CI: 1.11–1.39), and in the very high SBP group AHR: 1.94 (95% CI: 1.74–2.16).
	hoc analysis of a multicenter trial involving 20,330 pts (age ≥50 y) with recent noncardioembolic ischemic stroke followed up for 2.5 y	 (140≤150 mm Hg), and very high (≥150 mm Hg). 1° outcome was recurrent stroke and the 2° outcome was a composite of recurrent stroke, MI, and death due to vascular causes 	8.0%) for the low-normal SBP group, 6.8% (95% CI: 6.1%–7.4%) for the high-normal SBP group, 8.7% (95% CI: 7.9%–9.5%) for the high SBP group, and 14.1% (95% CI: 13.0%–15.2%) for the very high SBP group. Compared with pts in the high-normal SBP group, the risk of 1° outcome was higher for pts in the very low- normal SBP group AHR: 1.29 (95% CI: 1.07–	<u>Summary</u> : Among pts with recent noncardioembolic ischemic stroke, SBP levels during follow-up in the very low-normal (<120 mm Hg), high (140–≤150 mm Hg), or very high (≥150 mm Hg) range were associated with increased risk of recurrent stroke.
	Study Size: 20,330		1.56), in the high SBP group AHR: 1.23 (95% CI: 1.07–1.41), and in the very high SBP group AHR: 2.08 (95% CI: 1.83–2.37).	
Ovbiagele B, et al., 2013 (221) 22244715	<u>Aim</u> : To assess association of maintaining low- normal vs. high- normal SBP levels with risk of recurrent stroke. <u>Study type</u> : Post hoc analysis of a multicenter trial involving 3,680 pts with recent noncardioembolic ischemic stroke followed up for 2 y	Inclusion criteria: Pts with an ischemic stroke <120 d before randomization Categories: • Based on mean in-trial SBP value was low-normal (<120 mm Hg), high-normal (120 to <140 mm Hg), or high (>140 mm Hg). • 1° outcome was stroke	<u>1° outcome</u> : First recurrence of stroke of any type <u>Key findings</u> : Rate of recurrent stroke was 9.1% in the low- normal group, 6.7% in the high-normal group, and 10% in the high group. Difference in recurrent stroke rate between low-normal and high-normal groups was more prominent within the first 6 mo (low-normal, 4.5%; high-normal, 2.5%; high, 3.4%) vs. after 6 mo (low-normal, 4.6%; high-normal, 4.2%; high, 6.6%). Over study period, compared with the high-normal group, risk of the 1° outcome trended higher in the low-normal group AHR: 1.47 (95% CI: 0.94–2.29; p=0.09) and was higher in the high group AHR: 1.39 (95% CI: 1.08–1.79; p=0.01).	Summary: Results support a possible pattern of increased risk of recurrent stroke in pts with low- normal SBP levels, especially within the first 6 mo after first stroke. However, this study likely was not sufficiently powered to detect more than a strong statistical trend underlying this relationship.

Lin MP, et al., 2015 (222) <u>25765723</u>	Aim: To assess link between SBP and mortality after stroke. Study type: Analyses of nationally representative survey data (NHANES) Study Size: 455 pts	Inclusion criteria: Adults ≥20 y with self-reported stroke. Categories: Baseline SBP was as low to normal (<120 mm Hg), normal (120–140 mm Hg), and high (≥140 mm Hg).	<u>1° outcomes</u> : All-cause and vascular mortality <u>Key findings</u> : 2 y after assessment, the low to normal SBP group tended to have the highest cumulative all-cause mortality (11.5%), compared with mortality rates of 8.5% and 7.5% in the normal and high SBP groups, respectively. Similar patterns were seen with vascular mortality. After adjusting for covariates, compared with the high SBP group, the low to normal group had higher all-cause mortality AHR: 1.96 (95% CI: 1.13–3.39; p=0.017) and trended toward higher vascular mortality AHR: 2.08 (95% CI: 0.93–4.6; p=0.075). Compared with the normal BP group, the risk of all-cause and vascular mortality trended higher in low to normal BP group but did not achieve statistical significance.	Summary: After stroke, compared with SBP in the high range, low to normal SBP may be associated with poorer mortality outcomes. Study limited by self-reported nature and retrospective design.
Kim J, et al., 2014 (223) <u>24509123</u>	Aim: To investigate the association between BP and vascular events up to 10 y after stroke. Study type: Analysis of population based study (North East Melbourne Stroke Incidence Study (NEMESIS)	Inclusion criteria: 5-y survivors of stroke Categories: Stratification by quartiles of SBP Follow-up: Annually by telephone at 6, 8, and 9 y and face-to-face interview at 7 and 10 y after stroke.	 <u>1° outcomes</u>: Composite of all-cause death or nonfatal vascular event (stroke or AMI); and all-cause death alone. <u>Key findings</u>: In 5-y survivors of stroke, compared to a SBP of 131–141 mm Hg, SBP of 120 mm Hg or less was associated with a 61% greater risk of stroke, acute MI and death (HR: 1.61; 95% CI: 1.08–2.41; p=0.019). Compared to the reference category of SBP 131–141 mm Hg, there were no differences in outcome in the pts with SBP 121–130 mm Hg (p=0.491) or 142–210 mm Hg (p=0.313). Findings were not modified after adjusting for antihypertensive drug prescriptions. 	<u>Summary</u> : There appears to be a greater risk of poor outcome in long-term survivors of stroke with low SBP. This is further evidence that low SBP may result in poor prognosis.
Wang WT, et al., 2016 (224) <u>27082571</u>	<u>Aim</u> : To investigate the relative effects of BP-lowering therapies [ACEI, ARB, BB, CCBs, diuretics, and	Inclusion criteria: • RCTs comparing the effects of any of the 6 most commonly used BP-lowering drug classes [ACEI, ARB, alpha-blocker, BB, diuretics, and CCB] vs. placebo	<u>1° outcome</u> : Recurrent stroke <u>2° outcome</u> : CHD, and MACCE <u>Key findings</u> : • Compared with placebo, ACEI plus diuretic	 Virtually all BP-lowering medication classes reduced vascular events including recurrent stroke. The higher the average BP reduction between the treatment vs. control groups the larger the risk reduction in recurrent stroke events and MACCE.

	combinations of 3 drugs] in pts with a prior stroke history <u>Study size</u> : 15 RCTs composed of 39,329 participants previous stroke	or comparing 1 type of antihypertensive agent with another type on pts who have suffered from stroke or TIA s • RCTs reporting outcomes of interest with a follow-up of more than a month.	 reduced recurrent stroke (OR: 0.54; 95% CI: 0.33–0.90). ACEI plus diuretic had a higher probability of being at the best ranking position (31%). Compared with regimens not including diuretics, diuretics-based treatments resulted in a significantly larger reduction in BP (12.0mm Hg; 95% CI: 7.0–16.9), Treatment regimens including diuretics had a RR of 0.619 (95% CI: 0.515–0.743) for recurrent stroke, which was significantly lower than treatments that did not include diuretics (RR=0.882; 95% CI: 0.800–0.973) with a p value for interaction of 0.0008. None of the between-drug comparisons showed significant differences in effect on outcomes 	 Diuretic-based treatments lowered the risk of recurrent stroke more than treatments that did not include diuretics. There were no significant differences in effect on 2° stroke reduction between the various individual antihypertensive medication classes.
Katsanos AH, et al., 2017 (225) <u>27802419</u>	Aim: To assess the association of BP reduction with recurrent stroke and CV events using available RCT data on 2° stroke prevention <u>Study size</u> : 14 studies with 42,736 pts	Inclusion criteria: RCTs of antihypertensives for 2° stroke prevention pts that reported achieved BP values during the follow-up period. Exclusion criteria: Observational studies, case series, case reports, RCTs in non-IS/TIA population, and studies not reporting data on finally achieved BP values	 <u>1° outcome</u>: Recurrent stroke <u>2° outcome</u>: MI, death from any cause, and risk of CV death <u>Key findings</u>: SBP reduction linearly associated with lower risk of recurrent stroke (regression slope, 0.02; 95% CI: 0.01–0.04; p=0.049), MI (regression slope, 0.022; 95% CI: 0.002–0.041; p=0.024), death from any cause (regression slope, 0.02; 95% CI: 0.01–0.03; p=0.001), and CV death (regression slope, 0.05; 95% CI: 0.03–0.07; p<0.001). No relation was observed between the degree of SBP reduction and the risk of disabling or fatal stroke (regression slope, 0.001; 95% CI: –0.024–0.022; p=0.944). Relation of SBP reduction with ischemic or hemorrhagic stroke was not assessed due to the small number of studies with available data (<10). 	Summary: BP reduction is linearly associated with the magnitude of risk reduction in recurrent cerebrovascular and CV events, but optimal BP target not evaluated.

Data Supplement 45. RCTs and Meta-analysis Comparing PAD (Section 9.5)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
HOPE Östergren J, et al., 2004 (226) <u>14683738</u>	Aim: To assess the impact of ramipril compared to placebo on the prevention of major CV events in PAD pts in the HOPE study. Study type: Multicenter, double-blind RCT Size: 9,541 randomized in HOPE (1,725 randomized who had baseline PAD, defined by ABI with pulse detection by either Doppler or palpation)	Inclusion criteria: • ≥55 y • Existing CVD (CAD, stroke, PAD) or DM with an additional CVD risk factor (smoking, HTN, hypercholesterolemia, low HDL, microalbuminuria) Exclusion criteria: • Received ACEI or vitamin E or had uncontrolled HTN • HF or LV dysfunction *All eligible pts had 7- to 10-d run-in period, received 2.5 mg ramipril daily; those who tolerated were then assigned placebo for 10–14 d and then were randomized to1 of intervention arms or control	Intervention: Ramipril (10 mg/d): 4,645 randomized Intervention: Placebo: 4,652 randomized	 <u>1º endpoint</u>: Combined CV death, nonfatal MI, nonfatal stroke In pts with history of symptomatic PAD, comparing ramipril to placebo: RR: 0.75; 95% CI: 0.61–0.92 In pts with no history of symptomatic PAD, but severe subclinical disease defined as ABI <0.6, comparing ramipril to placebo: RR: 0.77; 95% CI: 0.55–1.09 In pts with no history of symptomatic PAD, but moderate subclinical disease defined as ABI 0.6–0.9, comparing ramipril to placebo: RR: 0.72; 95% CI: 0.56–0.92 <u>1º Safety endpoint</u>: N/A <u>Summary:</u> Ramipril prevented clinical evidence of PAD as well as in those without PAD. The relative benefit was similar in pts classified by levels of ABI, even though event rates were higher in pts with subclinical and clinical ABI. 	 Relevant 2° endpoint: Individual components of composite endpoint, all-cause mortality, hospitalizations for HF, DM complications In pts with history of symptomatic PAD, comparing ramipril to placebo: for MI, RR: 0.75 (95% CI: 0.58–0.98); for stroke, RR: 0.72 (95% CI: 0.50–1.05); for CVD mortality, RR: 0.75 (95% CI: 0.56–0.99); for total mortality, RR: 0.85 (95% CI: 0.68–1.07); for DM complications, RR: 0.87 (95% CI: 0.74–1.09); for HF, RR: 0.81 (95% CI: 0.53–1.24) In pts with no history of symptomatic PAD, but severe subclinical disease defined as ABI <0.6, comparing ramipril to placebo: for MI, RR: 0.73 (95% CI: 0.52–1.89); for CVD mortality, RR: 0.76 (95% CI: 0.46–1.25); for total mortality, RR: 0.81 (95% CI: 0.55–1.19); for DM, RR: 0.83 (95% CI: 0.55–1.19); for DM, RR: 0.83 (95% CI: 0.50–1.39); for HF, RR: 0.66 (95% CI: 0.34–1.28) In pts with no history of symptomatic PAD, but moderate subclinical disease defined as ABI 0.6–0.9, comparing ramipril to placebo: for MI, RR: 0.81 (95% CI: 0.26–0.77); for CVD mortality, RR: 0.44 (95% CI: 0.26–0.77); for CVD mortality, RR: 0.62 (95% CI: 0.42–0.90); for total mortality, RR: 0.58 (95% CI: 0.42–0.79); for diabetic complications, RR: 0.80 (95% CI: 0.53–1.21); for HF, RR: 0.69 (95% CI: 0.38–1.23)

Overlack A, et al., 1994 (227) <u>8059778</u>	Aim: To determine the effect of perindopril compared to placebo on various clinical outcomes in pt subgroups. Study type: Multicenter, double-blinded RCT (3 wk placebo run-in period, 6 wk double-blind phase) Size: 490 (54 with PAD)	Inclusion criteria: • Mild newly diagnosed essential HTN in addition to 1 concomitant diseases or therapies: hyperlipidemia, DM-2, IHD, cardiac arrhythmias, PAD, nephropathy with proteinuria, COPD, or degenerative join disease with NSAIDs • 40–75 y *Antihypertensive treatment was stopped 1 wk prior to randomization, required DBP 95–104 mm Hg Exclusion criteria: N/A	Intervention: Perindopril (4 mg/d): 253 randomized Comparator: Placebo: 237 randomized	 <u>1° endpoint</u>: ABI measured by Doppler In pts with baseline PAD, there was no difference in post-treatment Doppler Index between perindopril (0.75) vs. placebo (0.75); p>0.05 <u>1° Safety endpoint</u>: Spontaneously reported side effects: 5.5% of pts in perindopril, 3.8% of pts in placebo <u>Summary</u>: In pts with PAD, Doppler index at baseline was not different between the 2 groups and remained unchanged during treatment. Pain-free and maximal walking distances increased from baseline but there were no significant between group differences. 	 <u>Study limitations and adverse events:</u> ABI not measured by Doppler gold standard <u>Relevant 2° endpoint:</u> Pain-free walking distance (m), maximal walking distance In pts with baseline PAD, there was no difference in change in pain-free walking distance (m) between perindopril (+11 m) vs. placebo (+11 m); p>0.05 In pts with baseline PAD, there was no difference in change in maximal walking distance between perindopril (pre-trial: 318 m (SD: 45), post-trial: 323 m (SD: 43) vs. placebo (pre-trial: 333 m (SD: 43), post-trial: 369 m (SD: 46) <u>Study limitations and adverse events:</u> Short follow-up, unable to assess hard clinical outcomes
Schweizer J, et al., 1998 (228) <u>9581724</u>	Aim: To determine whether treatment with high dose verapamil prevents restenosis in pts with PAD at high risk for reoccurrence after successful PTCA. Study type: Double- blind RCT (6 mo duration)	Inclusion criteria: • PAD (based on arterial angiography and color- coded duplex ultrasound) present for >6 mo • Primary success of PTCA treatment (≥30% reduction of initial lumen constriction) • Stable angina pectoris, mild HTN and at least1	Intervention: Verapamil (240 mg/twice/d): 49 randomized Comparator: Placebo: 49 randomized	 <u>1° endpoint</u>: Percentage of diameter stenosis At 6 wk, mean % diameter stenosis in verapamil group was 46.8 (SD: 14.1) vs. placebo was 55.5 (SD: 10.0) At 6 mo, mean % diameter stenosis in verapamil group was 48.0 (SD: 11.5) vs. 	 <u>Relevant 2° endpoint:</u> Intima/media thickness was 1.2 mm (SD: 0.31) in verapamil vs. 1.9 mm (SD: 0.47), p<0.001 Septal thickness was 10.2 mm (SD: 1.1) in verapamil vs. 11.9 mm (SD: 2.3), p<0.001 Crurobrachial ratio dorsalis pedis was 0.76 (SD: 0.10) in verapamil vs. placebo was 0.72 (SD: 0.08)

	<u>Size</u> : 98 pts	additional risk factor: DM, hyperlipoproteinemia, total or subtotal vascular occlusion of dilated segmented, eccentric stenosis, residual stenosis of at least 30%, or stenosis localized in the distal superficial femoral artery <u>Exclusion criteria:</u> • History of pelvic stenosis • Previous adjuvant therapy with calcium antagonists or beta- adrenergic blocking agents • Age >75 y • Prior revascularization of same area • 1st, 2nd, or 3rd AV block, sinoatrial block, diseases of supporting or connective tissues, moderate arterial HTN with SBP >170 mm Hg and DBP >95 mm Hg		placebo was 69.6 (SD: 12.2), p<0.01 <u>1° Safety endpoint</u> : N/A <u>Summary:</u> In pts with PAD at increased risk for restenosis, the administration of high dose verapamil prevented recurrent stenosis for 6 mo after successful peripheral angioplasty and was well tolerated.	 Crurobrachial ratio tibial artery was 0.76 (SD: 0.09) in verapamil vs. placebo was 0.70 (SD: 0.10) Arterial pressure was 134/87 mm Hg (SD: 5.2/4.2) in verapamil vs. placebo was 165/97 mm Hg (6.5/4.4), p<0.001 Total vessel diameter was 8.3 mm (SD: 0.3) in verapamil vs. 7.5 mm (SD: 0.3), p<0.001 <u>Study limitations and adverse events:</u> Short follow-up, unable to assess hard clinical outcomes
NORMA Espinola-Klein C, et al., 2011 (229) 21646599	<u>Aim</u> : Evaluate the effects of treatment with the endothelium-dependent vasodilating beta 1- selective blocker nebivolol, as compared with the nonvasodilating beta 1-selective blocker metoprolol, on clinical parameters of PAD and endothelial function, and to compare the	HTN with SBP >170 mm	Intervention arms: • Nebivolol (5 mg/d): 65 randomized • Metoprolol (95 mg/d): 63 randomized	<u>1° endpoint</u> : • Change in ABI measured by Doppler • In nebivolol: initial ABI 0.62 (SD: 0.16), post-treatment ABI 0.68 (SD: 0.20), p-value for change: 0.002 • In metoprolol: initial ABI 0.63 (SD: 0.17), post- treatment ABI 0.67 (SD:	Relevant 2° endpoint:• Change in absolute claudication distance were 32.7 m in nebivolol (p- value 0.03) vs. 39.7 m in metoprolol (p- value 0.01), but no difference between 2 groups (p-value 0.54)• Changes in SBP were -5.2 mm Hg in nebivolol (p=0.001) and -3.9 mm Hg in metoprolol (p=0.01), no difference between groups

	tolerability of both drugs in pts with PAD <u>Study type</u> : Double- blinded RCT (48 wk) <u>Size</u> : 128	 DBP at time of enrollment <100 mm Hg Exclusion criteria: Premenopausal women Critical limb ischemia with rest pain, leg ulcer, gangrene, severe angina pectoris that limits exercise capacity, severe HF that 		 0.21), p-value for change: 0.04 Comparing ABI change in nebivolol to metoprolol: 0.02 (p=0.69). <u>1st safety endpoint</u>: N/A <u>Summary:</u> BB therapy was well tolerated in pts with 	 No change in flow-mediated dilatation in either group (p=0.16) <u>Study limitations and adverse events:</u> Absence of placebo group 21 total adverse events, 10 in nebivolol, 11 in metoprolol (adverse events: bradycardia, tachycardia, blurred vision, worsening HTN, edema, worsening claudication, blurred vision, erectile
		limits exercise capacity, hyperthyroidism, poorly controlled DM (HbA1c>10%) • Contraindications for BBs • Acute MI within 6 mo before screening • Previous treatment with nebivolol or carvedilol		intermittent claudication and HTN during a treatment period of 1 y. In the direct comparison, there was no significant difference between nebivolol and metoprolol.	dysfunction, edema, vertigo, temporary dysesthesia of the hands, dyspnea, skin irritation, headache, moderate diarrhea)
		*Concomitant treatment with calcium antagonists, ACEIs, angiotensin II type 1 receptor antagonists, aspirin, clopidogrel, statins, estrogens was permitted if no change in dosage had been made in the previous 3 mo before screening			
INVEST Bavry AA, et al., 2010 (230) <u>19996066</u>	<u>Aim</u> : To examine the effect of average treated BP on adverse outcomes in PAD pts with CAD and to compare 2 antihypertensive medications <u>Study type</u> : Post hoc analysis of international	Inclusion criteria: • ≥50 y • HTN, clinically stable CAD • Pt reported PAD Exclusion criteria: Contraindications to the treatment groups	Interventions: • Calcium antagonist- based strategy: verapamil with or without trandolapril • BB-based strategy: atenolol with or without hydrochlorothiazide *2° medications only given to achieve BP of	 <u>1° endpoint</u>: Composite outcome: all- cause death, nonfatal MI, nonfatal stroke No statistically significant difference in composite 1° outcome OR: 0.90 (95% CI: 0.76, 1.07) comparing calcium antagonist based group to BB based group in fully adjusted model 	Relevant 2° endpoint: N/A• This trial also notes the J-shaped relationship between BP achieved and clinical outcomes• Risk of 1° outcome was reduced most when SBP was treated to 130–140 mm Hg and DBP 60–90, as opposed to <130/80 as 2005 guidelines suggest in PAD ptsStudy limitations and adverse events:

	randomized, blinded- endpoint trial (48 wk) <u>Size</u> : 22,576 in total trial (2,699 with PAD in this analysis)		<140/90 mm Hg in all participants except for those with renal impairment or DM, BP<130/85 mm Hg	 Kaplan–Meier curve for 1° outcome shows slightly lower cumulative incidence in calcium antagonist group (log rank p=0.26) <u>1st safety endpoint</u>: N/A <u>Summary:</u> Among PAD pts, the incidence of the 1° outcome was not significantly different between treatment groups. 	 PAD was not uniformly measured or adjudicated (only based on pt report) Asymptomatic PAD was not captured
VALUE Zanchetti A, et al., 2006 (231) <u>17053536</u>	Aim: To examine the effect of valsartan vs. amlodipine on cardiac morbidity and mortality in hypertensive pts at high CV risk Study type: Prespecified additional analyses of international randomized, double-blind, parallel- group trial Size: 15,245 in total trial (2,114 with PAD)	Inclusion criteria: ≥ 50 y • HTN (untreated: 160– 210/<115 mm Hg, treated: <210/<115 mm Hg) • High risk for cardiac events (male sex, verified DM, current smoking, high cholesterol, LV hypertrophy by ECG, proteinuria on dipstick, serum creatinine 150–265 micromol/L, coronary disease diagnosis, cerebrovascular disease diagnosis, or PAD diagnosis) Exclusion criteria: • Renal artery stenosis • Pregnancy • AMI, coronary angioplasty or CABG in last 3 mo • Severe hepatic disease • Severe chronic renal failure	Interventions: • Valsartan: 7,649 total • Amlodipine: 7,596 total *No PAD-specific numbers available	 <u>1° endpoint</u>: Composite of sudden cardiac death, fatal MI, death during/after percutaneous coronary intervention or CABG, HF requiring hospitalization, nonfatal MI, or emergency procedure to prevent MI There was no significant difference in the 1° outcome by treatment group among all pts and by PAD status. Among pts with PAD, the 1° outcome occurred in 13.4% of valsartan vs. 13.6% of amlodipine pts. Among pts without PAD, the corresponding % were 10.1% and 9.9%. <u>1st safety endpoint</u>: <u>Summary</u>: The effects of treatments on occurrence of the 1° outcome did not different by PAD status. 	Relevant 2° endpoint: N/A Study limitations and adverse events: • Limited subgroup analyses, only 1° outcome reported • High-risk population limits generalizability

Piller LB, et al., 2014 (232) 25002161	Aim: To compare, by randomized treatment groups (amlodipine, lisinopril, chlorthalidone) hospitalized or revascularized PAD rates and subsequent morbidity and mortality. Study type: Post-hoc analysis of prospective, randomized, double- blinded active-control trial (ALLHAT study— amlodipine, lisinopril compared to chlorthalidone control arm) Size: 33,357 pts Aim: To evaluate the	 Congestive HF requiring ACEI therapy Pts on monotherapy with 3 blockers for both CAD and HTN Inclusion criteria: BP of 140–180/90–110 for untreated, 160/100 for treated pts Age ≥55 y Have at least1 CV risk factor (risk factors: old myocardial injury or stroke, history of coronary revascularization procedure, other documented atherosclerotic CVD PAD, history of intermittent claudication, peripheral artery revascularization or peripheral artery angioplasty, DM-2, current cigarette smoking, HDL <0.90 mmol/L, LVH, major ST depression, T-wave inversion) Exclusion criteria: Canadian pts for whom outcome measures could not be assessed (n=533) 	Intervention arms: • Amlodipine: 8,898 randomized • Lisinopril: 8,904 randomized Comparator: Chlorthalidone: 15,002 randomized *Goal BP was <140/90 in each randomized group (achieved using study drug but adding open-label agents at physician discretion when necessary) Interventions: Any	 <u>1° endpoint</u>: PAD requiring hospitalization or outpatient revascularization procedure 830 cases of PAD over 8.8 y follow-up; no significant difference between treatment groups after adjustment HR comparing amlodipine to chlorthalidone: 0.86 (95% CI: 0.72, 1.03) after full adjustment, p-value: 0.099 HR comparing lisinopril to chlorthalidone: 0.98 (95% CI: 0.83, 1.17) after full adjustment, p-value: 0.847 Kaplan Meier: Y-to-PAD was longer amlodipine vs. chlorthalidone (no difference between lisinopril and chlorthalidone) <u>1° Safety endpoint</u>: N/A 	Relevant 2° endpoint:• Post-PAD morbidity and mortality• Comparing amlodipine tochlorthalidone, no difference in post-PADmorbidity or mortality: MI, HR: 0.82 (95%CI: 0.48, 1.40); Stroke, HR: 0.86 (95% CI:0.41, 1.79); Cardiac Revascularization,HR: 1.39 (95% CI: 0.81, 2.39); HF, HR1.32 (95% CI: 0.79, 2.18); Total Mortality,HR: 0.92 (95% CI: 0.74, 1.15)• Comparing lisinopril to chlorthalidone,no difference in post-PAD morbidity ormortality: MI, HR: 0.74 (95% CI: 0.44,1.25); Stroke, HR: 0.94 (95% CI: 0.48,1.86); Cardiac Revascularization, HR:1.25 (95% CI: 0.73, 2.13); HF, HR: 1.08(95% CI: 0.65, 1.80); Total Mortality, HR:0.95 (95% CI: 0.77, 1.18)Study limitations and adverse events:• PAD not specifically collected atbaseline, thus cannot detect actualincidence (however, randomizationpresumably resulted in equal number ofbaseline PAD cases in each group)• Asymptomatic PAD likely missed(definition used in this study based onhospitalization, likely only capturing verysevere cases)Study limitations and adverse events:
al., 2011 (113) <u>21364140</u>	effect of antihypertensive treatment on 2° prevention of CVD events and all-cause mortality among pts	of antihypertensive treatment among pts with BP <140/90 mm Hg for the prevention of CVD events.	antihypertensive agent compared with placebo or no treatment.	controls, pts receiving antihypertensive medications had a pooled RR of 0.77 (95% CI: 0.61, 0.77) for stroke: 0.80 (95% CI: 0.69,	 PAD not specifically collected at baseline, thus cannot detect actual incidence (however, randomization presumably resulted in equal number of baseline PAD cases in each group)

without clinically defined	Exclusion criteria: CVD	0.93) for MI: 0.71 (95% CI:	Asymptomatic PAD likely missed
HTN.	events were not reported	0.65, 0.77) for CHF: 0.85	(definition used in this study based on
	by HTN status that	(95% CI: 0.80, 0.90) for	hospitalization, likely only capturing very
Study type: Meta-	included participants with	composite CVD events: 0.83	severe cases)
analysis including 25	and without HTN; study	(95% CI: 0.69, 0.99) for CVD	
RCTs	population did not include	mortality and 0.87 (95% CI:	
	persons with BP in the	0.80, 0.95) for all-cause	
Size: 64,162 pts without	normal or prehypertensive	mortality from random effect	
HTN.	ranges; study population	models. Results did not differ	
	did not include persons	according to trial	
	with preexisting CVD or	characteristics or subgroups	
	CVD equivalents, such as	defined by clinical history,	
	DM; antihypertensive	although no specific PAD	
	medication was not a part	subgroup was defined.	
	of the intervention;	5	
	treatment allocation was	Summary: Among pts with	
	not random; measure of	clinical history of CVD,	
	variance not reported;	including PAD, but without	
	participants were <18 y;	HTN, antihypertensive	
	there were differences	treatment was associated	
	between intervention and	with reduced risk of stroke,	
	control groups other than	CHF, composite CVD events	
	antihypertensive treatment.	and all-cause mortality.	
	Preexisting CVD included	<i>y</i>	
	PAD.		

Data Supplement 46. RCTs and Meta-analyses Comparing BP Targets in DM (Section 9.6)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# pts) / Study Comparator (# pts)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
ADVANCE Kaplan NM, et al., 2007 (233) <u>17765962</u>	<u>Aim:</u> To assess the effects of an ACEI perindopril and a diuretic indapamide combination on serious vascular events in pts with	DM-2 pts 30–55 y. <u>Inclusion criteria</u> : At least 1 of the following: history of major CVD, (stroke, MI, admission for TIA, UA, coronary	• Fixed combination of perindopril and indapamide compared with perindopril and placebo.	<u>1° endpoints</u> : Composite of CV death, nonfatal MI, nonfatal stroke, new or worsening nephropathy, or retinopathy. <u>Results</u> : After 4.3 y follow-up, pts assigned to active therapy had a reduction of SBP of 5.6 mm Hg. RR of major macro- or micro-	Summary: • This large RCT provides evidence that routine administration of fixed combination ACEI and thiazide-type diuretic therapy reduces risk of major CV events in those with at least 1 risk factor.

	DM irrespective of initial BP levels or the use of other BP- lowering drugs. <u>Study type</u> : RCT <u>Size:</u> 11,140 pts, 4.3 y follow-up	revascularization, or amputation for PVD) or at least 1 other risk factor (history of microvascular disease, microalbuminuria, proliferative diabetic retinopathy, retinal photocoagulation therapy, macular edema, blindness, cigarette smoking, high cholesterol, low HDL cholesterol, low HDL cholesterol, diagnosis of DM at least 10 y before enrollment or ≥65 y at entry <u>Exclusion criteria</u> : HbA1c target ≤6.5% or indication for insulin.		vascular events decreased by 9% (HR: 0.91; (95% CI: 0.83, 1.00), p<0.04). Death from CVD decreased by 18%; RR: 0.82 (95% CI: 0.68, 0.98) and death from any cause decreased by 14%; RR: 0.86 (95% CI: 0.75, 0.98). The effects of study treatment did not differ by initial BP or concomitant use of other treatments at baseline. The pts had at least 1 CV risk factor.	• The ADVANCE trial included DM pts both with and without HTN. In this RCT, pts were randomized to active treatment or placebo rather than to a different BP goal, so that it is impossible to determine whether the benefit was due to the treatment of HTN <i>per se</i> .
ACCORD Cushman WC, et al., 2010 (234) 20228401	Aim: To assess whether therapy targeting normal SBP (<120 mm Hg) reduces major CV events in DM-2 at high risk for CV events. Study type: RCT Size: 4,733 pts, 4.7 y follow-up	Inclusion criteria:DM-2 with HgbA1c ≥7.5%;≥40 y with CVD or ≥55y with anatomicalevidence ofatherosclerosis,albuminuria, LVH, or≥2 additional riskfactors for CVD.Exclusion criteria:BMI ≥45, serumcreatinine >1.5, andother serious illness.	• Pts were randomly assigned to intensive therapy SBP <120 mm Hg or standard therapy SBP <140 mm Hg.	<u>1° outcomes:</u> Nonfatal MI, nonfatal stroke, or CV death. <u>Results:</u> Mean SBP in the intensive therapy group was 119.3 mm Hg and in the standard therapy group was 133.5 mm Hg. The annual 1° outcome 1.87% in the intensive therapy group and 2.09% in the standard therapy group HR: 0.88; 95% CI: 0.073–1.06; p=0.20. The annual rates of death from any cause were 1.28% and 1.19% in the 2 groups, respectively (HR: 0.59; 95% CI: 0.39–0.89; p=0.01). Serious adverse events attributed to antihypertensive treatment occurred in 3.3% of the intensive therapy group and 1.3% of the standard therapy group (p<0.001).	Limitations: This trial had an open label design. The rate of adverse events in the standard therapy group was less than expected. Pts younger than 40 y or older than 79 y were not included. Summary: In pts with DM-2 and high risk for CV events, targeting SBP of <120 as compared with <140 mm Hg did not reduce the rate of composite outcome of fatal and nonfatal major CV events and was associated with greater risk for adverse events.

Margolis KL et al., 2014 (235) <u>24595629</u>	<u>Aim:</u> To compare effects of combinations of standard and intensive treatment of glycemia and BP in the ACCORD trial. <u>Study type</u> : RCT <u>Size</u> : 4,733 pts, 4.7 y follow-up	Inclusion criteria: Type 2 DM with HgbA1c ≥7.5%; ≥40 y with CVD or ≥55 y with anatomical evidence of atherosclerosis, albuminuria, LVH, or at least 2 additional risk factors for CVD. Exclusion criteria: BMI ≥45, serum creatinine >1.5, and other serious illness.	• Pts were randomly assigned to intensive therapy SBP<120 mm Hg or standard therapy SBP<140 mm Hg.	<u>1° outcomes:</u> Nonfatal MI, nonfatal stroke, or CV death. <u>Results</u> : In the BP trial, risk of the 1° outcome was lower in the groups intensively treated for glycemia HR: 0.67 (95% CI: 0.50, 0.91), BP HR: 0.74 (95% CI: 0.55, 1.00), or both HR: 0.71 (95% CI: 0.52, 0.96) compared with combined standard BP and glycemia treatment. For 2° outcomes, MI was significantly reduced by intensive glycemia treatment and stroke by intensive BP treatment; most other HRs were neutral or favored intensive treatment groups.	Limitations: 2° analysis; results analyzed across individual cells of a factorial design with shorter follow- up than originally intended reducing power to detect meaningful differences and interactions; results may not apply to younger, healthier diabetics. Conclusions: Either intensive BP or glycemia control reduced major CVD compared with combined standard treatment, but the combination was no better than the individual intensive interventions.
Soliman EZ et al., 2015 (236) <u>26459421</u>	Aim: To compare effects of combinations of standard and intensive control of BP on the risk of LVH in the ACCORD trial. Study type: RCT Size: 4,331 pts, 4.7 y follow-up	Inclusion criteria: DM- 2 with HgbA1c ≥7.5%; ≥40 y with CVD or ≥55 y with anatomical evidence of atherosclerosis, albuminuria, LVH, or at least 2 additional risk factors for CVD. Exclusion criteria: BMI ≥45, serum creatinine >1.5, and other serious illness.	• Pts were randomly assigned to intensive therapy SBP<120 mm Hg or standard therapy SBP<140 mm Hg.	<u>1° outcomes:</u> Nonfatal MI, nonfatal stroke, or CV death. <u>Results</u> : The outcome measures were electrocardiographic LVH defined by Cornell voltage (binary variable) and mean Cornell index (continuous variable). The baseline prevalence of LVH (5.3% vs. 5.4% ; p=0.91) and the mean Cornell index ($1,456$ vs. $1,470$ μ V; p=0.45) were similar in the intensive (n=2,154) and standard (n=2,177) BP- lowering arms, respectively. However, after median follow-up of 4.4 y, intensive, compared with standard, BP lowering was associated with a 39% lower risk of LVH (OR: 0.61; 95% CI: 0.43–0.88; p=0.008) and a significantly lower adjusted mean Cornell index ($1,352$ vs. $1,447$ μ V; p<0.001). The lower risk of LVH associated with intensive BP lowering during follow-up was because of more regression of baseline LVH and lower rate of developing new LVH, compared with standard BP lowering. No interactions by age, sex, or race were observed.	Limitations: 2° analysis; open-label design; LVH defined by EKG and not by echo or cardiac MRI; results may not apply to younger, healthier diabetics. Conclusions: Targeting a SBP of <120 mm Hg when compared with <140 mm Hg in pts with HTN and DM produces a greater reduction in LVH

Xie X, et al.,	Aim: To assess the	Inclusion criteria:	• 5 RCTs (6,960	1° outcomes: Major CV events, defined as	Study limitations: Only 6,960 pts
2015 (21)	efficacy and safety	RCTs with different BP	pts) enrolled only	MI, stroke, HF or CV death, separately and	with DM were included in the total
<u>26559744</u>	of intensive BP	targets or different BP	pts with DM and 6	combined; nonvascular and all-cause	study size of 44,989 pts.
	lowering strategies.	changes between more	trials (2,809 pts)	mortality; ESKD; and adverse events; new	
		vs. less intense therapy	specifically recruited	onset microalbuminuria/macroalbuminuria or	Conclusions: The absolute CV
	Study type:	with at least 6 mo	pts with CKD.	change from micro- to macroalbuminuria and	benefits were greatest in trials in
	Systematic review	follow-up.		retinopathy in pts with DM.	which all enrolled pts had vascular
	and meta-analysis				disease, renal disease or DM.
		Exclusion criteria:		Results: Pts in the more intensive BP-	However, only 6,960 of the 44,989
	Size: 19 trials with	Trials that did not		lowering treatment group had mean BP	pts had DM and no sub-analysis for
	44,989 pts; 3.8 y of	assess a different		133/76 mm Hg compared with 140/81 mm	DM was provided; however, the
	follow-up.	target or relevant		Hg in the less intensive group. Intensive BP-	outcome benefits were qualitatively
		outcome.		lowering treatment achieved RR reductions	most striking for pts with DM, CKD
				for major CV events: 14% (95% CI: 4–22),	and/or vascular disease.
				MI: 13% (95% CI: 0–24), stroke: 22% (95%	
				Cl: 10–32), albuminuria: 10% (95% Cl: 3–	
				16), and retinopathy progression: 19% (95%	
				CI: 0–34). However, more intensive	
				treatment had no clear effects on HF: RR:	
				15% (95% CI: -11–34), CV death: 9% (-11–	
				26), total mortality: 9% (95% CI: -3–19), or	
				ESKD: 10% (95% CI: -6–23). The reduction in major CV events was consistent across pt	
				groups, and additional BP lowering had a	
				clear benefit even in pts with SBP <140 mm	
				Hg. The absolute benefits were greatest in	
				trials in which all enrolled pts had vascular	
				disease, renal disease, or DM. Serious	
				adverse events associated with BP lowering	
				were only reported by 6 trials and had an	
				event rate of 1%–2% per y in intensive BP	
				lowering group pts, compared with 0.9% in	
				the less intensive treatment group (RR: 1.35;	
				95% CI: 0.93–1.97). Severe hypotension was	
				more frequent in the more intensive	
				treatment regimen (RR: 2.68; 95% CI: 1.21-	
				5.89; p=0.015), but the absolute excess was	
				small (0.3% vs. 0.1% per pt-y for the duration	
				of follow-up).	

ACCOMPLISH Weber MA, et al., 2010 (237) 20620720	Aim: To determine which combination therapy in pts with HTN and DM most effectively decreases CV events. Study type: RCT Size: 2,842 pts with DM from the ACCOMPLISH study of 6,946 pts; 30 mo follow-up	Inclusion criteria: HTN and DM with high risk for CV events. Exclusion criteria: BMI >45; serum Cr >1.5; other serious illness	Pts were randomly assigned to benazepril plus amlodipine or benazepril plus hydrochlorothiazide. BPs were 145/79 at baseline.	<u>1° outcomes</u> : Composite of death from CV causes, nonfatal MI, nonfatal stroke, hospitalization for angina, resuscitation after sudden cardiac arrest, and coronary revascularization. <u>Results</u> : The mean achieved BP was 131.5/72.6 and 132.7/73.7 in the B + A and B + H groups, respectively, during the 30 mo of follow-up. There were 8.8% and 11% 1° events, respectively (HR: 0.79; 95% CI: 0.684–0.92; p=0.003). In the pts with DM there were clear coronary benefits with B + A, including both acute clinical events (p=0.013 and revascularizations (p=0.024). There were no unexpected adverse events.	Summary: In pts with DM and HTN, combining an ACEI with a CCB, compared with hydrochlorothiazide, was superior in reducing CV events.
ASCOT Ostergren J, et al., 2008 (238) <u>18854748</u>	Aim: To compare the effects of an amlodipine-based regimen vs. and atenolol-based regimen on CV outcomes in pts with DM <u>Study type:</u> RCT (BP lowering arm of ASCOT) <u>Size:</u> 5,137 pts with DM, minimum 4 y follow-up	Inclusion criteria: Pts 40–65 y with HTN (>160/100 mm Hg) or treated HTN and DM plus 2 additional CV risk factors: PAD, previous stroke or TIA, male sex, ≥55 y, microalbuminuria, smoking, total cholesterol to HDL ratio ≥6, or family history of CHD.	• Pts were randomly assigned to an amlodipine- based regimen with addition of perindopril as required or an atenolol-based regimen with addition of a thiazide as required and therapy titrated as required to achieve target BP of 130/80 mm Hg.	<u>1° outcomes:</u> Fatal CHD and nonfatal MI. <u>Results</u> : BPs were 136/75 (amlodipine and 137/76 (atenolol) at the end of study. There was a 3/1.9 mm Hg lower BP in pts on amlodipine. The amlodipine-based regimen reduced CV events and procedures compared to the atenolol-based regimen (HR 0.86; 0.76-0.98; p=0.026). Fatal and nonfatal strokes were reduced by 25% (p=0.017), PAD by 48% (p=0.004) and noncoronary vascularization procedures by 57% (p=0.001).	Summary: In the large DM subgroup of the BP-lowering arm of ASCOT, the benefits of an amlodipine-based treatment compared with an atenolol-based treatment on the incidence of total CV events and procedures was significant.
SHEP Kostis JB, et al., 2005 (239) <u>15619390</u>	Aim: To assess the long-term mortality rate of pts with DM pts in the SHEP trial randomly assigned to stepped care with chlorthalidone or placebo.	Inclusion criteria: Isolated systolic HTN (SBP 160–219 mm Hg) with DBP <90 mm Hg. Exclusion criteria: Pts with insulin–dependent DM and those who	• Pts were randomly assigned to chlorthalidone or placebo. If BP remained above goal, atenolol or placebo was added.	<u>1° outcomes:</u> CV mortality rate <u>Results</u> : BP was 11.1/3.4 mm Hg lower in the active treatment group at the end of the study. Diuretic treatment in pts with DM was strongly associated with long-term CV mortality rate (AHR: 0.668 (95% CI: 0.526,	Summary: Chlorthalidone-based treatment improved long-term outcomes in pts with DM.

		required diuretic		0.848) and total mortality rate: 0.805 (95%	
	Study type: RCT	therapy.		Cl: 0.680, 0.952).	
	<u>Size:</u> 4,732 pts; follow-up 14.3 y				
ROADMAP Menne J, et al., 2012 (240) 22418908	<u>Aim:</u> To assess whether olmesartan compared to placebo delays the onset of albuminuria in pts with DM and HTN. <u>Study type:</u> RCT <u>Size:</u> 4,020 pts; follow-up 3.2 y	Inclusion criteria: Pts with HTN defined as BP ≥130/80 mm Hg and at least 1 CV risk factor.	• Pts were randomly assigned to olmesartan or placebo. Additional antihypertensive therapy except for ACEs and ARBs to lower BP.	<u>1° outcome:</u> Time to onset of microalbuminuria. <u>Results</u> : Average BP was 126.3/74.7 and 129.5/76.6, respectively (significant not stated). Olmesartan delayed the onset of microalbuminuria by 25% (0.75; 95% CI: 0.61–0.92; p=0.007). CV events were comparable in the 2 groups.	<u>Summary:</u> Pts with better BP reduction are less likely to develop microalbuminuria. Treatment with an ARB delayed the onset of microalbuminuria independently of baseline BP and degree of BP reduction.
ABCD Estacio RO, et al., 1998 (241) <u>9486993</u>	Aim: To compare the effects of "intensive" compared with "moderate" BP treatment on 24-h creatinine clearance (GFR) in pts with DM and HTN. Study type: RCT – open label Size: 472 pts; follow- up 5 y	Inclusion criteria: Pts with HTN defined as DBP ≥90 mm Hg and DM-2	Pts were randomly assigned to "intensive" treatment (DBP<75 mm Hg and "moderate" treatment (DBP 80– 89 mm Hg) with a combination of nisoldipine and enalapril as the initial antihypertensive medication.	 <u>1° outcome</u>: Change in 24-h creatinine clearance. <u>Results</u>: The mean BP achieved was 132/78 in the intensive group and 138/86 in the moderate control group. During the 5-y follow-up period, there was no difference in GFR between the groups. After the first y of antihypertensive treatment, GFR stabilized in both the intensive and moderate groups with normal albumin excretion or microalbuminuria. In contrast, pts with overt albuminuria demonstrated steady decline in GFR whether on intensive or moderate therapy. Neither was there a significant difference in the progression from normal to micro- or micro-to overt albuminuria. Intensive therapy demonstrated a lower overall incidence of deaths, 5.5% vs. 10.7%; p=0.037 (2° endpoint). 	Limitations: Open-label design; the definition of DM was 2 fasting blood glucose measurements >140 mg/dL as opposed to >126 today; serious side effects were not reported. Risk of bias due to a greater proportion of pts with established CVD at baseline assigned to the standard BP target. Summary: BP control of 138/86 or 132/78 with either nisoldipine or enalapril as the initial antihypertensive agent appeared to stabilize renal function in HTN pts with type 2 DM without overt albuminuria over a 5-y period. For the ABCD trials, only ABDC (H) included strictly pts with HTN and DM. The quality of evidence is low due to imprecision and risk of bias.
Hypertension Optimal Treatment (HOT trial)	Aim: To assess the optimum target DBP	Inclusion criteria: Pts with HTN defined as	 Pts were randomly assigned to 1 of 3 DBP target 	<u>1° outcomes:</u> Major CV events, MI, stroke, CV mortality and total mortality.	Limitations: Open-label design; the definition of DM-2 fasting blood glucose measurements >140 mg/dL

Hansson L, et al., 1998 (242) <u>9635947</u>	in the treatment of HTN. <u>Study type</u> : RCT <u>Size:</u> 1,501 pts in the DM subgroup; follow-up 3.8 y	DBP 100–115 mm Hg and DM.	groups: ≤90, ≤85, or ≤80 mm Hg.	<u>Results:</u> In the group randomized to <80 mm Hg, the risk of major CV events was halved in comparison to the target <90. CV mortality was lower in the <80 group compared to the other groups.	as opposed to >126 today; serious side effects were not reported; potential bias due to subgroup analysis. <u>Summary</u> : In pts with DM and HTN, intensive lowering of BP was associated with a low rate of CV events. The quality of evidence is low to very low due to imprecision and risk of bias.
UKPDS 1998 (243) 9732337	Aim: To determine whether tight control of BP prevents macrovascular and microvascular complications in pts with DM-2. Study type: RCT Size: 1,148 hypertensive pts with type 2 DM Follow-up: 8.4 y	Inclusion criteria: Fasting plasma glucose concentration >6 mmol/l in 2 mornings. Exclusion criteria: Ketonuria >3 mmol/l; history of MI in the previous y; current angina or HF; >1 major vascular episode; serum creatinine concentration >175 µmol/l; retinopathy requiring laser treatment; malignant HTN; an uncorrected endocrine abnormality; an occupation that would preclude insulin treatment; a severe concurrent illness; inadequate understanding or unwillingness to enter the study.	• Pts were randomized to tight BP control (target BP<150/85 mm Hg) or less tight BP control (target <180/105 mm Hg),	 <u>1° outcomes:</u> 1) First clinical endpoint related to DM (sudden death, death from hyperglycemia or hypoglycemia, fatal or nonfatal MI, angina, HF, stroke, renal failure, amputation, vitreous hemorrhage, retinal photocoagulation, blindness in 1 eye or cataract extraction). 2) Death related to DM. 3) Death from all causes. <u>Results</u>: BP in the tight BP control group was 144/82 compared with the group assigned less tight control (154/87), p<0.0001. Reductions in risk in the group assigned tight BP control compared with those of the less tight control group were 24% (95% CI: 8%–38%; p=0.0046) in DM related endpoints; 32% in deaths related to DM (95% CI: 6%–51%; p<0.019; 44% in strokes (95% CI: 11%–36%; p<0.0092 in microvascular endpoints, predominantly due to risk of retinal photocoagulation. 	Limitations: DBP targets were high (85 mm Hg in the tight control group and 105 mm Hg in the less tight control group) and similar to the cutoffs for the no treatment groups in trials comparing treatment with no treatment. UKPDS evaluated lowering both SBP and DBP so it is impossible to separate the outcomes effects of DBP. Therefore, the evidence is of low quality. Summary: Tight BP control in pts with HTN and DM-2 achieved a clinically important reduction in the risk of death related to DM, progression of DM retinopathy and deterioration of visual acuity, but the quality of evidence is low.
Arguedas JA, et al., 2013 (244) <u>24170669</u>	Aim: To determine if "lower" BP targets (any target <130/85	Inclusion criteria: RCTs in which individuals were	• Pts with HTN and DM were randomly assigned to the	<u>1° outcomes:</u> Total mortality, total serious adverse events, MI, stroke, CHF, and ESRD.	Conclusions: Evidence from RCTs does not support BP targets lower

mm Hg) are	randomized to a "lower"	intensive or	Results: Only 1 trial (ACCORD) compared	than standard targets in pts with
associated with	compared with a	standard BP control	outcomes associated with 'lower' (<120 mm	HTN and DM.
reduction in mortality	"standard" BP target.	group.	Hg) or 'standard' (<140 mm Hg) SBP targets	
and morbidity	June 1991	3 • • F	in 4734 pts. Despite achieving a significantly	
compared to	Exclusion criteria:		lower BP (119.3/64.4 mm Hg vs. 133.5/70.5	
"standard" BP	Studies that did not		mm Hg, p<0.0001), and using more	
targets (<140–	meet the inclusion		antihypertensive medications, the only	
160/90–100 mm Hg)	criteria. Excluded		significant benefit in the group assigned to	
in pts with DM.	studies were UKPDS		'lower' SBP was a reduction in the incidence	
	1998, HTN in Diabetes		of stroke: RR: 0.58; (95% CI: 0.39–0.88;	
Study type: Meta-	Study IV 1996, SANDS		p=0.009), absolute risk reduction 1.1%. The	
analysis of RCTs.	2008, Lewis 1999 and		effect of SBP targets on mortality was	
analysis of ICCTS.	the Steno-2 study.		compatible with both a reduction and	
Size: 5 RCTs	the Sterio-2 study.		increase in risk: RR: 1.05 (95% CI: 0.84,	
recruiting a total of			1.30), low quality evidence. Trying to achieve	
7,314 ps.			the 'lower' SBP target was associated with a	
7,514 ps.			significant increase in the number of other	
Mean follow-up: 4.5			serious adverse events: RR: 2.58, (95% CI:	
			•	
У			1.70-3.91; p<0.00001, absolute risk increase	
			2.0%. 4 trials (ABCD-H, ABCD-N, ABCD-2V,	
			and a subgroup of HTN Optimal Treatment)	
			specifically compared clinical outcomes	
			associated with 'lower' vs. 'standard' targets	
			for DBP in pts with DM. The total number of	
			pts included in the DBP target analysis was	
			2580. Pts assigned to 'lower' DBP had a	
			significantly lower achieved BP: 128/76 mm	
			Hg vs. 135/83 mm Hg; p<0.0001. There was	
			a trend towards reduction in total mortality in	
			the group assigned to the 'lower' DBP target:	
			RR: 0.73 (95% CI: 0.53–1.01), mainly due to	
			a trend to lower non-CV mortality. There was	
			no difference in stroke: RR: 0.67, (95% CI:	
			0.42–1.05), in MI: RR: 0.95 (95% CI: 0.64–	
			1.40) or in CHF: RR: 1.06 (95% CI: 0.58–	
			1.92), low-quality evidence. End-stage renal	
			failure and total serious adverse events were	
			not reported in any of the trials. A sensitivity	
			analysis of trials comparing DBP targets <80	
			mm Hg (as suggested in clinical guidelines)	

				vs. <90 mm Hg showed similar results. There was a high risk of selection bias for every outcome analyzed in favor of the 'lower' target in the trials included for the analysis of DBP targets.	
Palmer SC, et al., 2015 (245) <u>26009228</u>	Aim: To investigate the benefits and harms of BP- lowering drugs in adults with DM Study type: Network meta-analysis of RCTs. Size: 157 studies in 43,256 pts mostly with DM and CKD. Mean follow-up: 4.5 y	Inclusion criteria: Pts ≥18 y with DM and CKD and were treated in clinical trials that compared any orally administered antihypertensive agent alone or in combination with a 2nd antihypertensive agent or combination, placebo, or control. Exclusion criteria: Pts who underwent kidney transplantation or dialysis.	N/A	<u>1° outcomes</u> : All-cause mortality and ESKD (need for dialysis or transplantation). <u>Results:</u> No drug regimen was more effective than placebo for reducing all-cause mortality. However, compared with placebo, ESRD was significantly less likely after dual treatment with an ARB and an ACEI: OR: 0.62 (95% CI: 0.43–0.90) and after ARB monotherapy: OR: 0.77 (95% CI: 0.65–0.92). No regimen significantly increased hyperkalemia or acute kidney injury, although combined ACEI and ARB treatment had the lowest rank among all interventions because of borderline increases in estimated risks of these harms; OR: 2.69 (95% CI: 0.97–7.47) for hyperkalemia; OR: 2.69 (95% CI: 0.98– 7.38) for acute kidney injury.	Limitations: Effects of BP treatment on CV events and related mortality were uncertain. Data for the outcome of ESKD were restricted largely to pts with macroalbuminuria. Acute kidney injury was poorly defined with low quality of evidence. Conclusions: No BP-lowering strategy prolonged survival in adults with DM and CKD. ACEIs and ARBs, alone or in combination, were the most effective strategies against ESKD. Any benefits of combined ACEI and ARB treatment need to be balanced against potential harms of hyperkalemia and acute kidney injury.
Turnbull F, et al., 2005 (246) <u>15983291</u>	Aim: To determine the benefits associated with different treatment regimens in pts with and without DM and whether there are important differences in the effects of different BP-lowering regimens in these 2 pt groups. <u>Study type:</u> Meta- analysis of RCTs.	Inclusion criteria: Randomization of pts between a BP-lowering agent and a control (placebo or less intensive BP-lowering regimen) or randomization of pts between regimens based on different classes of BP-lowering drugs. Exclusion criteria: Studies not meeting the above criteria.	N/A	<u>1° outcomes</u> : Nonfatal stroke or death from cerebrovascular disease; nonfatal MI or death from CAD; HF causing death or requiring hospitalization; total CV events; total CV deaths; and total mortality. <u>Results</u> : Total major CV events were reduced to a comparable extent in individuals with and without DM by regimens based on ACEIs, calcium antagonists, ARBs and diuretics/ BBs (p<0.19 for all). There was limited evidence that lower BP goals produced larger reductions in total major CV events in pts with vs. without DM (p<0.03).	Limitations: No analysis of renal outcomes, risk of new DM or progression of existing DM; combined comparison of persons taking diuretics and BB s; some studies selected pts on the basis of the presence or absence of DM. Summary: Effects of BP-lowering agents on major CV events were broadly comparable for pts with and without DM.

ALLHAT Whelton PK, et al., 2005 (247) 15983290	Size: 27 RCTs including 158,709 pts (33,395 with DM and 125,314 without DM). Follow-up: Minimum 1,000 pt-y Aim: To determine the optimal first step antihypertensive drug therapy in DM- 2 or impaired fasting blood glucose levels and specifically whether treatment with a CCB or ACEI decreases clinical complications compared to treatment with a thiazide type diuretic. Study type: RCT Size: 31,512 pts stratified into type 2 DM (13,101), IFG (1,399) and normoglycemia	Inclusion criteria: Pts ≥55 y with HTN and at least 1 other risk factor for CHD. Exclusion criteria: No history of DM or no fasting glucose measurement or nonfasting glucose level ≥110 mg/dL.	• Pts were randomly assigned to double-blind first- step treatment with chlorthalidone 12.525 mg/d, amlodipine 2.5–10 mg/d or Lisinopril 10–40 mg/d.	<u>1° outcomes:</u> Fatal CHD and nonfatal MI <u>Results</u> : There was no significant difference in RR (RR) for the 1° outcome in DM or NG pts assigned to amlodipine or lisinopril vs. chlorthalidone or in IFG pts assigned to lisinopril vs. chlorthalidone RR: 1.73 (95% CI: 1.10, 2.72). A significantly higher RR was noted for the 1° outcome in IFG pts assigned to amlodipine vs. chlorthalidone. Stroke was more common in NG pts assigned to lisinopril vs. chlorthalidone RR: 1.31 (95% CI: 1.10, 1.57). HF was more common in DM and NG pts assigned to amlodipine RR: 1.39 (95% CI: 1.22, 1.59) and 1.30 (95% CI: 1.12, 1.51), respectively or lisinopril: 1.15 (95% CI: 1.00–1.32) and 1.19 (95% CI: 1.02, 1.39), respectively vs. chlorthalidone.	Limitations: Microalbuminuria was not measured. Summary: Our results provide no evidence of superiority for treatment with CCBs or ACEIs compared with a thiazide-type diuretic during first- step antihypertensive therapy in DM, IFG, or NG.
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Data Supplement 47. Nonrandomized Trials, Observational Studies, and/or Registries in DM (Section 9.6)

Study Acronym;Study Type/Design;Patient PopulationPrimary Endpoint and ResultsSummary/ConclAuthor;Study Size (N)(include P value; OR or RR; & 95% Cl)Comment(s)Year PublishedYear PublishedStudy Size (N)Study Size (N)	clusion (s)
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ADVANCE Hata J, et al., 2013 (248) 23926207	<u>Aim</u> : To assess the effects of visit-to-visit SBP variability and maximum SBP on the risks of macrovascular or microvascular outcomes by using data from the ADVANCE trial. <u>Study type:</u> Observational analysis <u>Size:</u> 8,811 pts	Inclusion criteria: Pts had not experienced major macro- or microvascular events during first 2 y of the ADVANCE trial Exclusion criteria: None	<u>1° endpoint</u> : Composite of CV death, nonfatal MI, nonfatal stroke, new or worsening nephropathy, or retinopathy. <u>Results:</u> Major macro- and micro-vascular events were associated with SBP variability even after adjustment for mean SBP and other confounding factors. For the highest 10% variability, HR: 1.54 (95% CI: 0.99, 2.39) for macrovascular events; for microvascular events, HR: 1.84 (95% CI: 1.19, 2.84).	<u>Summary:</u> Visit-to-visit SBP variability and maximum SBP are independent risk factors for macro- and micro-vascular events.
ADVANCE-ON Zoungas S, et al., 2014 (249) 25234206	Aim: To determine whether the mortality benefit that had been observed among pts originally assigned to BP- lowering therapy were still evident at the end of 6-y follow-up Study type: Observational analysis Size: 8,494 pts	Inclusion criteria: Pts with DM who participated in post- trial follow-up for 6 y Exclusion criteria: See above	<u>1° endpoint</u> : Death from any cause and major macrovascular complications (a composite of nonfatal MI, nonfatal stroke, or death from any CV cause. <u>Results:</u> The reductions in the risk of death from any cause and of death from CV causes that had been observed in the group receiving active BP-lowering treatment during the ADVANCE trial were attenuated but significant at the end of the post-trial follow- up. HRs were 0.95 (95% CI: 0.84–0.99; p=0.03) and 0.88 (95% CI: 0.77–0.99; p=0.04), respectively.	Summary: Benefits were attenuated but still present at the end of 6 y.
ROADMAP Mene J, et al., 2014 (250) 24772521	Aim: To determine whether the ROADMAP olmesartan medoxomil treatment resulted in a potential long-term micro- and macro-vascular benefit. <u>Study type</u> : Observational analysis <u>Size</u> : 1,758 pts; 3.3 y follow- up	Inclusion criteria: See above Exclusion criteria: See above	<u>1° endpoint:</u> See above <u>Results:</u> The original ROADMAP study showed a 23% reduction in microalbuminuria despite good and comparable BP control in both groups. Pts who developed microalbuminuria had a higher incidence of cardio- and cerebrovascular events: OR: 1.77 (95% Cl: 1.03–3.03; p=0.039) compared to those in whom this was not the case. DM retinopathy and HF requiring hospitalization also were reduced.	Summary: renal artery stenosis blockade might cause a sustained reduction in micro- and macro-vascular events.
Edmin C, et al., 2015 (251)	Aim: Determine associations between BP-lowering	Inclusion criteria: All RCTs of BP-lowering treatment in	BP-lowering drug vs. placebo: 26 RCTs	Limitations: Reliability of this meta-analysis is limited by the scarcity of large trials with

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<u>25668264</u>	treatment and presence of	which entire trial population	More intensive vs. less intensive BP	achieved SBP levels in the 120–130 mm Hg
	vascular disease in DM-2	had DM-2 or in which the	lowering: 7 RCTs	range. The relatively short follow-up of
		results of a DM subgroup	 BP-lowering vs. another drug: 17 RCTs 	included trails may have prevented
	Study type: Large meta-	were obtained. Studies were		associations of BP-lowering treatment with
	analysis of	included regardless of the	Results: Baseline BP: A 10-mm Hg SBP	vascular outcomes from being observed,
	40 high quality RCTs (1/1966–	presence or absence of	reduction was associated with a significantly	particularly for outcomes such as HF and
	10/2014) judged low risk of	defined HTN.	lower risk of all-cause mortality RR: 0.87	renal failure, which are often a consequence
	bias		(95% CI: 0.78–0.96), CVD events RR: 0.89	of MI or albuminuria, respectively.
		Exclusion criteria: Trials	(95% CI: 0.80–0.98), and stroke events RR:	
	Size: 100,354 pts with DM; all	conducted predominantly in	0.73 (95% CI: 0.64–0.83). The associations	Summary:
	trials >1,000 pt-y of follow-up	pts with type 1 DM were	for HF and renal failure were not significant.	 This large meta-analysis of 40 RCTs
	BP-lowering drug vs. placebo:	excluded.	For microvascular events, a 10-mm reduction	provides evidence that BP lowering is
	26 RCTs		in SBP was associated with a lower risk of	associated with lower risks of outcomes in pts
			retinopathy RR: 0.87 (95% CI: 0.76–0.99) and	with initial mean SBP \geq 140 mm Hg compared
	More intensive vs. less		albuminuria RR: 0.83 (95% CI: 0.79–0.87).	with those <140 mm Hg with the exception of
	intensive BP lowering: 7 RCTs		· · · · · · · · · · · · · · · · · · ·	stroke, albuminuria and retinopathy. When
	BP-lowering vs. another		Stratified by initial SBP:	trials were stratified by achieved SBP
	drug: 17 RCTs		Trials stratified by SBP >140 to <140 mm Hg	treatment was associated with lower risks only
	3		showed significant interactions for all-cause	in the <130 mm Hg stratum for stroke and
			mortality RR: 0.73 (95% CI: 0.64–0.84) vs.	albuminuria.
			1.07 (95% CI: 0.92–1.26), CVD RR: 0.74	• This meta-analysis shows that although BP
			(95% CI: 0.65–0.85) vs. RR: 0.96 (95% CI:	lowering was not associated with a lower risk
			0.88–1.05), CHD RR: 0.73 (95% CI: 0.61–	of CVD or CHD events at a baseline SBP
			0.87) vs. RR: 0.97 (95% CI: 0.86–1.10), HF	<140 mm Hg, it does observe lower risks of
			RR: 0.75 (95% CI: 0.59–0.94) vs. RR: 0.97	stroke, retinopathy and progression of
			(95% CI: 0.79–1.19) and albuminuria RR:	albuminuria.
			0.71 (95% CI: 0.63–0.79) vs. RR: 0.86 (95%	This study provides evidence that for
			Cl: 0.81–0.99).	individuals at high risk for these outcomes
				(history of cerebrovascular disease or mild
			Stratified by achieved SBP:	nonproliferative retinopathy), commencement
			Trials stratified by SBP achieved in the	of therapy below an initial SBP of 140 mm Hg
			treatment group \geq 130 or <130 mm Hg and	and treatment to SBP <130 may be indicated.
			the associations of a 10-mm Hg SBP	and treatment to SDF < 150 may be indicated.
			reduction compared between the strata	
			showed significant interactions for all-cause	
			mortality RR: 0.75 (95% CI: 0.65–0.86) vs.	
			RR: 1.06 (95% CI: 0.90–1.265), CVD RR:	
			0.74 (95% CI: 0.64–0.85) vs. RR: 0.96 (95%	
			CI: 0.88–1.05), CHD RR: 0.70 (95% CI: 0.58–	
			0.83) vs. RR: 0.97 (95% CI: 0.85–1.10), HF	

			RR: 0.75 (95% CI: 0.59–0.95) vs. RR: 1.00 (95% CI: 0.81–1.23) and albuminuria RR: 0.71 (95% CI: 0.64–0.79) vs. RR: 0.86 (95% CI: 0.81–0.90) with higher risk in the ≥130 mm Hg group. Stratified by class of medications: Few differences were observed in the association between BP-lowering treatment and outcomes for regimens based on different classes of medications, except HF, in which diuretics were associated with lower RR: 0.83 (95% CI: 0.72–0.95) than all other classes. This was driven largely by the results of ALLHAT.	
Cheng J, et al., 2014 (252) <u>24687000</u>	Aim: To separately evaluate the effects of ACEIs and ARBs on all-cause mortality, CV deaths, and major CV events in pts with DM Study type: Meta-analysis of 35 high quality RCTs (1966– 2012) Size: 56,444 pts with DM; all trials had follow-up of at least 12 mo	Inclusion criteria: RCTs including post hoc analyses and subgroups for DM with median follow-up of at least 12 mo. Comparisons with placebo, no treatment or other antihypertensive drugs, including ACEIs and ARBs. Exclusion criteria: Cross- over trials	• ACEIs significantly reduced the risk of all- cause mortality by 13% (RR: 0.87; 95% CI: 0.78–0.98), CV deaths by 17% (RR: 0.83; 95% CI: 0.70–0.99), and major CV events by 14% (RR: 0.86; 95% CI: 0.77–0.95), including MI by 21% (RR: 0.79; 95% CI: 0.65–0.95) and HF by 19% (RR: 0.81; 95% CI: 0.71–0.93). Treatment with ARBs did not significantly affect all-cause mortality (RR: 0.94 (95% CI: 0.82–1.08), CV death rate (RR: 1.21 (95% CI: 0.81–1.80) and major CV events (RR: 0.94; 95% CI: 0.85–1.01) with the exception of HF (RR: 0.70; 95% CI: 0.59–0.82).	Summary: • RCTs comparing ACEs vs. active drugs/placebo/no treatment: 26 RCTs (12 active drugs, 11 placebo) • RCTs comparing ARBs vs. active drugs/placebo/no treatment: 13 RCTs (3 active drugs, 10 placebo) • This meta-analysis provides evidence that ACEIs reduce all-cause mortality, CV mortality, and major CV events in pts with DM, whereas ARBs had no benefits on these outcomes.
Arguedas JA, et al., 2013 (244) <u>24170669</u>	Aim: To determine if "lower" BP targets (any target <130/85 mm Hg) are associated with reduction in mortality and morbidity compared to "standard" BP targets (<140–160/90–100 mm Hg) in pts with DM. <u>Study type</u> : Meta-analysis of RCTs.	Inclusion criteria: RCTs in which individuals were randomized to a "lower" compared with a "standard" BP target. Exclusion criteria: Studies that did not meet the inclusion criteria. Excluded studies were UKPDS 1998, HTN in Diabetes Study IV	<u>1° outcomes:</u> Total mortality, total serious adverse events, MI, stroke, CHF, and ESRD. <u>Results</u> : Only 1 trial (ACCORD) compared outcomes associated with 'lower' (<120 mm Hg) or 'standard' (<140 mm Hg) SBP targets in 4734 pts. Despite achieving a significantly lower BP (119.3/64.4 mm Hg vs. 133.5/70.5 mm Hg, p<0.0001), and using more antihypertensive medications, the only significant benefit in the group assigned to 'lower' SBP was a reduction in the incidence	<u>Conclusions:</u> Evidence from RCTs does not support BP targets lower than standard targets in pts with HTN and DM.

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	Size: 5 RCTs recruiting a total	1996, SANDS 2008, Lewis	of stroke: RR: 0.58 (95% CI: 0.39–0.88;	
	of 7,314 ps.	1999 and the Steno-2 study.	p=0.009), absolute risk reduction 1.1%. The	
			effect of SBP targets on mortality was	
	Mean follow-up: 4.5 y		compatible with both a reduction and increase	
			in risk: RR: 1.05 (95% CI: 0.84–1.30), low-	
			quality evidence. Trying to achieve the 'lower'	
			SBP target was associated with a significant	
			increase in the number of other serious	
			adverse events: RR: 2.58 (95% CI: 1.70–3.91;	
			p<0.00001), absolute risk increase 2.0%. 4	
			trials (ABCD-H, ABCD-N, ABCD-2V, and a	
			subgroup of HOT) specifically compared	
			clinical outcomes associated with 'lower' vs.	
			'standard' targets for DBP in pts with DM. The	
			total number of pts included in the DBP target	
			analysis was 2580. Pts assigned to 'lower'	
			DBP had a significantly lower achieved BP:	
			128/76 mm Hg vs. 135/83 mm Hg,	
			p<0.0001. There was a trend towards	
			reduction in total mortality in the group	
			assigned to the 'lower' DBP target: RR: 0.73	
			(95% CI: 0.53–1.01), mainly due to a trend to	
			lower non- CV mortality. There was no	
			difference in stroke: RR: 0.67 (95% CI: 0.42–	
			1.05), in MI: RR: 0.95 (95% CI: 0.64–1.40) or	
			in CHF: RR: 1.06 (95% CI: 0.58–1.92), low	
			quality evidence. End-stage renal failure and	
			total serious adverse events were not	
			reported in any of the trials. A sensitivity	
			analysis of trials comparing DBP targets <80	
			mm Hg (as suggested in clinical guidelines)	
			vs. <90 mm Hg showed similar results. There	
			was a high risk of selection bias for every	
			outcome analyzed in favor of the 'lower' target	
			in the trials included for the analysis of DBP	
			targets.	
Cushman WC, et	Aim: To assess whether	Inclusion criteria: Type 2	Pts were randomly assigned to intensive	Limitations: This trial had an open label
al., 2010 (234)	therapy targeting normal SBP	DM with HgbA1c \geq 7.5%; \geq 40	therapy SBP<120 mm Hg or standard therapy	design. The rate of adverse events in the
20228401	(<120 mm Hg) reduces major	y with CVD or \geq 55 y with	SBP<140 mm Hg.	standard therapy group was less than
		anatomical evidence of	_	
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	CV events in type 2 DM at high risk for CV events.	atherosclerosis, albuminuria, LVH, or at least 2 additional risk factors for CVD.	<u>1° outcomes</u> : Nonfatal MI, nonfatal stroke, or CV death.	expected. Pts younger than 40 y or older than 79 y were not included.
	<u>Study type</u> : RCT <u>Size</u> : 4,733 pts, 4.7 y follow- up	risk factors for CVD. <u>Exclusion criteria</u> : BMI ≥45, serum creatinine >1.5, and other serious illness.	<u>Results:</u> Mean SBP in the intensive therapy group was 119.3 mm Hg and in the standard therapy group was 133.5 mm Hg. The annual 1° outcome 1.87% in the intensive therapy group and 2.09% in the standard therapy group HR: 0.88 (95% CI: 0.073–1.06; p=0.20). The annual rates of death from any cause were 1.28% and 1.19% in the 2 groups, respectively (HR: 0.59; 95% CI: 0.39–0.89; p=0.01). Serious adverse events attributed to antihypertensive treatment occurred in 3.3% of the intensive therapy group and 1.3% of the standard therapy group (p<0.001).	<u>Summary</u> : In pts with type 2 DM and high risk for CV events, targeting SBP of <120 as compared with <140 mm Hg did not reduce the rate of composite outcome of fatal and nonfatal major CV events and was associated with greater risk for adverse events.
Hartley L, et al., 2014 (253) <u>25436436</u>	Aim: To determine the effectiveness of transcendental meditation for the 1° prevention of CVD Study type: Literature review of RCTs Size: 4 trials with a total of 430 pts	Inclusion criteria: ≥3 mo duration, healthy adults or adults at high risk of CVD, comparison of no or minimal intervention. Exclusion criteria: Multi- factorial interviews	1° outcomes: Clinical CVD events and major CVD risk factors <u>Results:</u> No conclusions of the effectiveness of transcendental meditation for the 1° prevention of CVD	<u>Limitations</u> : Limited evidence <u>Summary</u> : No conclusions as to the effectiveness of transcendental meditation for the 1° prevention of CVD. There was considerable heterogeneity between trials and the included studies were small, short-term, and at overall serious risk of bias.
Schmieder RE, et al., 2007 (254) 17416265	Study type: Topic review	Inclusion criteria: N/A Exclusion criteria: N/A	<u>1° outcomes</u> : N/A Results: N/A	Limitations: N/A Summary: N/A
Lv, et al., 2013 (127) <u>23798459</u>	Aim: To assess the renal and CV effects of intensive BP lowering in people with CKD Study type: Systematic review Size: 9,287 pts with CKD and 1,264 kidney failure events	 <u>Inclusion criteria</u>: N/A <u>Inclusion criteria</u>: Randomized trials of pts with CKD assigned to different target BP that reported kidney failure and CV events. 11 trials on 9,287 pts with CKD and 1,264 kidney failure events (doubling of serum creatinine, 50% decline in GFR or ESKD) Included AASK, REIN-2, 	<u>Results</u> . N/A <u>Results</u> . Compared with standard regimens, more intensive BP lowering reduced risk of composite endpoint HR: 0.82 (95% CI: 0.68– 0.98) and ESKD HR: 0.79 (95% CI: 0.67– 0.93). Effect was modified by proteinuria (p=0.006) and markers of trial quality. Intensive BP lowering reduced the risk of kidney failure HR: 0.73 (95% CI: 0.62–0.86) but not in pts without proteinuria at baseline HR: 1.12 (95% CI: 0.67–1.87). No clear effect on CV events or death.	Limitations: All trials used open label, in 2 pts were blinded, substantial variability in design quality. There was substantial variability in BP targets by MAP, SBP and DBP or only DBP. Most trials did not include pts with diabetic kidney disease Summary: • Renal outcomes: 7 trials (N=5308) recorded a total of 1,264 kidney failure events. A -7.7 mm Hg difference in SBP and a -4.9 mm Hg

MDRD, Wuhl (children), Toto, Schrier plus 5 trials with CKD subgroups, also included the late nonrandomized follow-up studies for AASK and MDRD • BP targets varied substantially between trials. 2 trials targeted mean BP <92 mm Hg for the intensive treatment arm, and 107 mm Hg in the standard treatment arm. 1 trial aimed for BP <130/80 mm Hg vs. a DBP of 90 mm Hg, 1 study targeted <120/80 mm Hg vs. 135– 140/85–90 mm Hg, and 4 studies had DBP <75–80 mm Hg vs. from 80–90 mm Hg. A trial involving pediatric pts targeted a 24-h mean BP <the 50th="" percentile,<br="">compared with the 50th to 95th percentiles in the control</the>	 difference in DBP seen between treatment arms. Overall, a more intensive regimen reduced risk of composite kidney failure events by 17% (HR: 0.82; 95% CI: 0.68, 0.98), reduced the risk of ESKD alone by 18% (pooled HR for composite outcomes: 0.79; 95% CI: 0.67, 0.93). Intensive BP lowering had no effect on kidney failure in pts who did not have proteinuria (3 trials involving 1,218 pts (HR: 1.12; 95% CI: 0.67–1.87), but it did reduce the risk of progressive kidney failure by 27% (5 trials involving 1,703 pts (HR: 0.73; 95% CI: 0.62–0.86) in pts who did have proteinuria at baseline. CV outcomes: major CV events reported in 5 trials (472 CV events in 5,308 pts with CKD). Intensive BP lowering did not reduce risk of CV events in pts with CKD, but the CIs remained wide (RR: 1.09 (95% CI: 0.83, 1.42). 6 trials reported stroke outcomes (197 events in 5,411 pts), 5 trials reported MI (138 events in 4.212 pts) and 5 trials reported UE
trial involving pediatric pts targeted a 24-h mean BP <the 50th="" percentile,<="" td=""><td>Intensive BP lowering did not reduce risk of CV events in pts with CKD, but the CIs remained wide (RR: 1.09 (95% CI: 0.83, 1.42). 6 trials reported stroke outcomes (197</td></the>	Intensive BP lowering did not reduce risk of CV events in pts with CKD, but the CIs remained wide (RR: 1.09 (95% CI: 0.83, 1.42). 6 trials reported stroke outcomes (197

Data Supplement 48. Atrial Fibrillation (Section 9.8)

Study Acronym; Author; Year Published	Aim of Study	Study Type	Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoints	P Value; OR, HR, or RR; & 95% Cl	Study Limitations & Adverse Events
Jibrini, et al., 2008 (255) <u>18223352</u>	<u>Aim</u> : To assess the effectiveness of ACEIs and ARBs in the prevention of AF, and to identify those clinical entities in which RAAS inhibition would most likely benefit the pts.	<u>Study type</u> : Meta-analysis	• 11 published studies; 55, 989 pts (26,973 pts in intervention, 29,016pts in comparator)	Inclusion criteria: Studies of RAAS blockade in CHF, MI, electrical cardioversion, and HTN) with incidence of AF noted during follow-up. Exclusion criteria: Studies without the measurement of AF or use of RAAS blockade.	Intervention: RAAS blockade Intervention: Placebo, amlodipine, BB or thiazide diuretic	<u>1° endpoint</u> (efficacy) and results: AF occurrence or reoccurrence.	Treatment with RAAS blockers reduced RR of AF in pts with HTN by 23% (p<0.001), by 11% in pts after MI (p<0.05), by 51% after electrical cardioversion (p<0.001), by 32% in pts with HF (p<0.001) and by 19% overall (p<0.001).	Not a comprehensive analysis of all antihypertensive. Adverse events not catalogued in meta-analysis.
Zhao et al., 2015 (256) <u>26668582</u>	<u>Aim</u> : To investigate the effectiveness and safety of ACEIs or angiotensin II receptor blockers (ARBs) on preventing AF in essential hypertensive pts. <u>Study type</u> : Meta-analysis	Intervention: RAAS blockade, n=20,491 <u>Comparator</u> : BB/calcium antagonist, n=22,401	Inclusion criteria: RCTs on the effects of ACEI/ ARBs on essential hypertensive pts. Exclusion criteria: Non- RCTs, subjects who were not treated with ACEI or ARB, and trials not	<u>1° endpoint</u> : AF occurrence or reoccurrence.	• ACEI/ARBs reduced the incidence of AF recurrence compared to calcium antagonists (RR: 0.48; 95% CI: 0.40–0.58; p<0.00001) or b-blockers (RR: 0.39; 95% CI: 0.20– 0.74; p=0.005). ACEI/ARBs may reduce the incidence of AF recurrence and CHF, with fewer serious adverse effects, but did not prevent new onset of AF.	N/A	• Doxazosin was associated with a higher incidence (2%) of AF/AFL prior to having the drug discontinued by the trial. Excluding doxazosin, there was no relationship between treatment drug and AF/AFL incidence.	• 2° analysis of RCT.

Size: 10 studies,	mentioning of AF prevention.		
n=42,892			

Data Supplement 49. Valvular Heart Disease (Section 9.9)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Study Intervention (# patients) / Study Comparator (# patients)	Patient Population	Endpoints	P Value; OR, HR, or RR; & 95% Cl	Study Limitations & Adverse Events
Healey et al., 2005 (257) <u>15936615</u>	<u>Aim</u> : Systematic review of all RCT evaluating the benefit of trials of ACEI and ARBs in prevention of AF <u>Study type</u> : Meta- analysis <u>Size</u> : 11 studies included with 56,308 pts	Intervention: n=27,089 RAAS blockade <u>Comparators</u> : n=29,220 placebo or active control antihypertensive	Inclusion criteria: Studies of RAAS blockade in CHF, MI, electrical cardioversion, and HTN) with incidence of AF noted during follow-up Exclusion criteria: Studies without the measurement of AF or use of RAAS blockade.	<u>1° endpoint</u> : AF occurrence or reoccurrence	• ACEIs and ARBs reduced RR of AF by 28% (p=0.0002), greatest in pts with HF [RR reduction: 44%; p=0.007). No significant reduction in AF in pts with HTN (RR reduction: 12%; p=0.4), but 1 trial found a significant 29% reduction in pts with LVH. Following cardioversion there was a large effect (48% RR reduction; 95% CI: 21%– 65%).	ACEIs and ARBs appear to be effective in prevention of AF probably limited to pts with systolic LV dysfunction or HTN LVH
Jibrini et al., 2008 (255) <u>18223352</u>	Aim: To assess the effectiveness of ACEIs and ARBs in the prevention of AF, and to identify those clinical entities in which RAAS inhibition would most likely benefit the pts. Study type: Meta- analysis	Intervention: n=26,973 RAAS blockade <u>Comparators</u> : n=29,016 placebo, amlodipine, BB or thiazide diuretic	Inclusion criteria: Studies of RAAS blockade in CHF, MI, electrical cardioversion, and HTN) with incidence of AF noted during follow-up Exclusion criteria: Studies without the measurement of AF or use of RAAS blockade.	<u>1° endpoint</u> : AF occurrence or reoccurrence.	• Treatment with RAAS blockers reduced RR of AF in pts with HTN by 23% (p<0.001), by 11% in pts after MI (p<0.05), by 51% after electrical cardioversion (p<0.001), by 32% in pts with HF (p<0.001) and by 19% overall (p<0.001).	N/A

	Size: 11 studies, 55,989 pts					
Zhao et al., 2015 (256) <u>26668582</u>	Aim: To investigate the effectiveness and safety of ACEIs or angiotensin II receptor blockers (ARBs) on preventing AF in essential hypertensive pts. <u>Study type</u> : Meta- analysis <u>Size</u> : 10 studies, n=42,892	Intervention: RAAS blockade, n=20,491 Comparator: BB/calcium antagonist, n=22,401	Inclusion criteria: RCTs on the effects of ACEI/ ARBs on essential hypertensive pts. Exclusion criteria: Non- RCTs, subjects who were not treated with ACEI or ARB, and trials not mentioning of AF prevention.	<u>1° endpoint</u> : AF occurrence or reoccurrence.	• ACEI/ARBs reduced the incidence of AF recurrence compared to calcium antagonists (RR: 0.48; 95% CI: 0.40–0.58; p<0.00001) or b-blockers (RR: 0.39; 95% CI: 0.20–0.74; p=0.005). ACEI/ARBs may reduce the incidence of AF recurrence and CHF, with fewer serious adverse effects, but did not prevent new onset of AF.	N/A
Hansson et al., 1999 (258) <u>10030325</u>	Aim: CAPP Trial was designed to compare the effects of ACE inhibition and conventional therapy on CV morbidity and mortality in pts with HTN. Study type: RCT Size: 10,985	Intervention: Captopril, n=5,592 Comparator: 5,493 pts were allocated to diuretics or BBs	Inclusion criteria: Pts aged 25–66 y with a measured DBP of ≥100 mm Hg on 2 occasions were included. Exclusion criteria: 2° HTN, serum creatinine concentration of more than 150 micromol/L, and disorders that required treatment with BB.	 <u>1° endpoint</u>: Fatal and nonfatal MI and stroke, and other CV deaths. <u>2° endpoint</u>: New or deteriorated IHD and CHF, AF, DM, TIA s, and death from all causes. 	• Captopril and conventional treatment did not differ in rates of all cardiac events— fatal and nonfatal MI, other CV deaths and sudden deaths, IHD, CHF, or AF (0.94; p=0.30).	N/A
Hansson et al., 1999 (259) <u>10577635</u>	Aim: STOPH-2 aimed to compare the effects of conventional and newer antihypertensive drugs on CV mortality and morbidity in elderly pts.	Intervention: n=2205 pts treated with ACEI <u>Comparator</u> : n=2,213 pts treated with BB or diuretic combination or n=2,196 pts treated with CCB	Inclusion criteria: HTN with BP ≥ 180 mm Hg systolic, aged 70– 84 y Exclusion criteria: Outside of the age range (n=14)	<u>1° endpoint</u> : CV death <u>2° endpoint</u> : CV events, DM and AF	• Old and new antihypertensive drugs were similar in prevention of CV mortality or major events. Decrease in BP was of major importance for the prevention of CV events. No difference in AF frequency was found (5.3% with ACEI, 4.1% with CCB and 5.2% with older drugs).	N/A

	Study type: RCT					
	<u>Size</u> : 6,614					
Wachtell et al., 2005 (260) <u>15734615</u>	Aim: LIFE trial aimed to determine whether angiotensin II receptor blockade is better than beta- blockade in	Intervention: n=4,298 treated with losartan Comparator: n=4,182 treated	Inclusion criteria: Hypertensive pts with LVH by echo Exclusion criteria: Prior AF history in 342 pts	<u>1° endpoint</u> : new onset of AF <u>2° endpoint</u> : None	• New-onset AF occurred in 150 pts randomized to losartan vs. 221 to atenolol (6.8 vs.10.1 per 1,000 person-y; RR: 0.67; 95% CI: 0.55–0.83; p<0.001) despite	N/A
	preventing new- onset AF. <u>Study type</u> : RCT	with atenolol			similar BP reduction. Pts receiving losartan tended to stay in sinus rhythm longer (p=0.057) than those receiving atenolol.	
	<u>Size</u> : 9,193					
Haywood et al., 2009 (261)	<u>Aim</u> : To investigate incidence of development of	Intervention: n=42,418 on diuretics	Inclusion criteria: Essential HTN with BP >140/90 without	<u>1° endpoint</u> : ECG evidence of AF/AFL on follow-up of HTN and	• AF/AFL occurred in 641 pts on follow-up. Incidence did not differ by class of	• Doxazosin group was limited by higher cardiac event rates and
<u>19926008</u>	AF/AFL in pts enrolled in this comparative trial of antihypertensives (ALLHAT). <u>Study type</u> : RCT	Comparator: n=39,056	medications, >180 systolic if on medications <u>Exclusion criteria</u> : Not meeting inclusion criteria	dyslipidemia	antihypertensive, other than increased frequency in the doxazosin group by 33% vs. chlorthalidone group (p=0.05 after risk adjustment).	early termination of this portion of the trial.
	<u>Size</u> : 81,474					
Julius et al., 2004 Julius, 2004 610}	<u>Aim</u> : The Valsartan Antihypertensive Long-term Use Evaluation (VALUE) trial: does valsartan	Intervention: n=7,649 on valsartan Comparator:	Inclusion criteria: Hypertensive pts, ≥50 y with DM, current smoking, high total cholesterol, LVH by ECG,	<u>1° endpoint</u> : Cardiac mortality, morbidity, HF, stroke, all-cause death, new onset DM	• AF occurred in 2.4% with valsartan and 2.0% with amlodipine; p=0.1197.	N/A
<u>15207952</u>	reduce cardiac morbidity and mortality more than	n=7,596 on amlodipine	proteinuria on dipstick and CKD (not end-stage)	Safety endpoint: Hypotension, syncope		
	amlodipine for the same degree of BP reduction in in hypertensive pts at high CV risk.		Exclusion criteria: ESRD, renal artery stenosis, pregnancy, AMI, PTCA or CABG within the past 3 mo, clinically	2° endpoint: AF		

Study type: RCT	relevant valvular disease, cerebrovascular accident in the past 3 mo, severe	
<u>Size</u> : 15,245	hepatic disease, severe chronic renal failure, CHF requiring ACEI therapy and	
	pts on monotherapy with blockers for both CAD and HTN.	

Data Supplement 50. RCTs and Meta-analysis Comparing Valvular Heart Disease (Section 9.9)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Study Intervention (# patients)/Study Comparator (# patients)	Patient Population	Endpoints	P Value; OR, HR, or RR; & 95% Cl	Study Limitations & Adverse Events
SCOPE-AS Chockalingam A, et al., 2004 (262) <u>15077102</u>	Aim: To determine the clinical tolerance and efficacy of the ACEI enalapril in the setting of symptomatic severe AS. Study type: RCT Size: 56 pts	Intervention: Enalapril 2.5 mg BID increasing to 10 mg BID (37 pts) <u>Comparator</u> : Placebo (19 pts)	Inclusion criteria: Severe aortic stenosis (aortic valve area <0.75 cm ² , mean aortic gradient >50 mm Hg, or aortic valve Doppler jet >4.5 m/s) and symptomatic NYHA class III or IV dyspnea or angina Exclusion criteria: Persistent hypotension (SBP <90 or mean BP <60), severe mitral stenosis (mitral valve orifice <1.0 cm ²), known intolerance for ACEI, and renal dysfunction (serum creatinine >2.5 mg/dL).	 <u>1° endpoint</u>: Improvements in Borg dyspnea index and 6-min walk distance at 1 mo <u>Safety endpoint</u>: Development of hypotension <u>2° endpoint</u>: Minor ACEI intolerance, cough, presyncope, improvement in NYHA class, and echo parameters 	• Pts who tolerated enalapril (n=34) had significant improvement in NYHA class, Borg index (5.4 \pm 1.2 vs. 5.6 \pm 1.7; p=0.03), and 6- min walk distance (402 \pm 150 vs. 376 \pm 174; p=0.003) compared with control pts.	• Treatment with enalapril resulted in hypotension in 3 of 5 pts with LV dysfunction and congestive HF had hypotension.

SEAS Rieck ÅE Hypertension, 2012 (263) 22647889	Aim: To determine the impact of HTN on LV structure and outcome during progression of aortic valve stenosis Study type: RCT observational substudy of SEAS trial Size: 1616 pts	Intervention: 1,340 pts with HTN Comparator: 276 pts without HTN	Inclusion criteria: Pts 45- 85 y who had asymptomatic, mild-to- moderate aortic valve stenosis, as assessed on echo, with a peak aortic- jet velocity of 2.5–4 m per second, were eligible for the study.	<u>1° endpoint</u> : Echo LV mass; MACE; mortality	 HTN predicted 51% higher incidence of abnormal LV geometry at final study visit independent of other confounders (p<0.01). HTN was associated with a 56% higher rate of ischemic CV events and a 2-fold increased mortality (both p<0.01). 	No specific randomized intervention for HTN.
Eleid MF, et al., 2013 (264) <u>23956211</u>	Aim: To evaluate the hemodynamic effects of vasodilator therapy in pts with LGSAS Study type: Nitroprusside infusion Size: 24	Intervention: Infusion of IV sodium nitroprusside to reduce BP and arterial afterload (18 pts with hypertensive LGSAS) <u>Comparator</u> : Baseline hemodynamics (6 pts with low EF LGSAS)	Inclusion criteria: Symptomatic pts with HTN (aortic SBP >140 mm Hg) and low-gradient (mean gradient <40 mm Hg) severe aortic stenosis (aortic valve area <1 cm (2)) with preserved EF (EF >50%). Exclusion criteria: Moderate or severe concomitant valvular heart disease (e.g., aortic, mitral or tricuspid regurgitation), reduced left ventricular EF (>50%), age <18 y, and complex CHD.	1° endpoint:Nitroprusside reducedmean PA pressure(25±10 mm Hg) and LVend-DBP (11±5 mm Hg;p<0.001 for both	• Treatment of HTN with vasodilator therapy results in a lowering of the total LV afterload, with a decrease in LV filling pressures and PA pressures.	No translation to clinical or ambulatory vasodilator use.
RIAS Trial Bull S, et al., 2015 (265) 25796267	<u>Aim</u> : To determine if ACEIs improve outcomes in AS. <u>Study type</u> : RCT <u>Size</u> : 100	Intervention: Ramipril ramped up from 2.5 to 5 to 10 mg for 1 y (50 pts) Comparator: Placebo (50 pts)	Inclusion criteria: Pts >18 y with moderate or severe aortic stenosis (valve area <1.5 cm ² , or peak velocity >3.0 m/s [peak valve gradient >36 mm Hg]), 2 who were asymptomatic as judged by pt-reported symptoms,	 <u>1° endpoint</u>: Adverse events; laboratory abnormalities; change in LVM from baseline to 12 mo measured by CMR. <u>2° endpoint</u>: Change in LV EF and function by CMR and echo, change in 	• Reduction in LVM in the ramipril group vs. placebo group (mean change -3.9 vs. +4.5 g, respectively; p=0.0057); preserved tissue Doppler systolic velocity compared with placebo (+0.0 vs0.5 cm/s;	• A larger clinical outcome trial to confirm these findings and explore their clinical relevance is required.

			and who did not have indications for valve replacement surgery. <u>Exclusion criteria</u> : Any other significant (>mild) VHD, excess hypo- or HTN (BP <100/40 or >200/110 mm Hg). Intolerance of ACEIs or ARBs or their prescription over the previous 3 mo	BNP); and change in distance walked on exercise tolerance testing.	p=0.04); trend to less progression of the aortic stenosis (valve area 0.0 cm ² vs0.2 cm ² in the placebo arm; p=0.067).	
Scognamiglio R, et al., 1994 (266) <u>8058074</u>	<u>Aim</u> : To assess whether vasodilator therapy reduces or delays the need for valve replacement <u>Study type</u> : RCT <u>Size</u> : 143	Intervention: Nifedipine 20 mg Q12 H (69 pts) Comparator: Digoxin 0.25 mg daily (74 pts)	Inclusion criteria: Asymptomatic pts with isolated, chronic, severe aortic regurgitation and normal LV systolic function Exclusion criteria: Worsening aortic regurgitation within 6 mo, DBP above 90 mm Hg, CAD, aortic valve gradient ≥ 20 mm Hg, other valvular or CHD, poor quality echo or an LV EF <50%.	<u>1° endpoint</u> : Frequency of valve replacement	• At 6 y, a 34% of the digoxin group had undergone valve replacement, but only 15% of the nifedipine group (p<0.001)	• No placebo group, and digoxin is a poor comparator due to toxicity which is now recognized.
Evangelista A, et al., 2005 (267) <u>16192479</u>	Aim: To identify the possible beneficial effects of vasodilator therapy on LV function and the need for aortic-valve replacement. Study type: RCT Size: 95 pts	Intervention: Nifedipine 20 mg Q12 H or enalpril 20 mg daily (32 pts nifedipine, 32 pts enalapril) Comparator: Placebo (31 pts)	Inclusion criteria: Consecutive pts with asymptomatic, chronic, severe aortic regurgitation and normal LV function Exclusion criteria: LVEF <50%, AF, CAD or other nonaortic VHD	<u>1° endpoint</u> : Frequency of valve replacement	• Rate of aortic-valve replacement was similar among the groups: 39% in the control group, 50% in the enalapril group, and 41% in the nifedipine group (p=0.62).	N/A

Scognamiglio R, et al., 1994 (266) <u>8058074</u>	Aim: To assess whether vasodilator therapy delays need for valve replacement in pts with asymptomatic severe AR. Study type: RCT Size: 143 pts	Intervention: 69 pts received nifedipine Comparator: 74 pts received digoxin	Inclusion criteria: Severe aortic regurgitation without symptoms Exclusion criteria: DBP >90, recent worsening of aortic regurgitation, mixed aortic stenosis / aortic regurgitation or any additional valve disease, LVEF <50.	<u>1° endpoint</u> : Worsening symptoms, LVEF decline to <50% or both, requiring valve replacement surgery	• 15% met criteria for valve replacement with nifedipine, but 34% did with digoxin (p<0.001)	No placebo control.
Evangelista A, et al., 2005 (14) <u>16192479</u>	Aim: To assess whether vasodilator therapy delays need for valve replacement in pts with asymptomatic severe AR. Study type: RCT Size: 95 pts	Intervention: 32 pts received enalapril; 32 pts received nifedipine <u>Comparators</u> : 31 pts received placebo	Inclusion criteria: Severe aortic regurgitation without symptoms Exclusion criteria: Not listed.	<u>1° endpoint</u> : Worsening symptoms, LVEF decline to <50% or both, requiring valve replacement surgery	• 41% met criteria for valve replacement with nifedipine, 50% did with enalapril, and 39% in the control group (p=0.62)	• BP of 145/75 average between the 3 groups, indicate lack of severity. Post-Rx BP is not reported.

Data Supplement 51. RCTs Comparing Race/Ethnicity (Section 10.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Leenen F, et al., 2006 (268) <u>16864749</u>	Study type: RCT comparison of an alpha blocker, ACEI, or CCB, each compared to a thiazide-type diuretic. This is post hoc comparison between	 >50 y Lisinopril (n=9,054); Amlodipine (9,048) African American 15,085 (35.5%) White 11,580 (47.0%) 	 Amlodipine vs. Lisinopril 	 No significant difference in 1° outcome (nonfatal MI and fatal CHD) or other prespecified outcomes: CHD, 1° outcome plus revascularization and hospitalized 	• In African Americans, Lisinopril less effective than amlodipine for BP reduction (mean follow-up BP 2.7/1.6 mm Hg higher with Lisinopril) and in reducing strokes (RR:1.51; 95% CI: 1.22–1.86) and

	CCB vs. ACEI incl in race subgroup.			angina, composite CVD, HF, ESRD, except strokes	combined CVD (RR: 1.13; 95% CI:1.02–1.24; p=0.025)
Wright JT et al. 2008 (269) <u>18227370</u>	<u>Study type</u> : Race subgroup comparison of RCT comparison of an ACEI or CCB compared to a thiazide-type diuretic on nonfatal or fatal CHD in pts with metabolic syndrome	 >50 y African American n=12,818 Non-African American n=24,473 	Chlorthalidone vs. Amlodipine, or Lisinopril	 No difference in 1° outcome (nonfatal MI and fatal CHD). Other prespecified outcomes: CHD, 1° outcome plus revascularization and hospitalized angina, composite CVD, stroke, HF, ESRD 	 In African Americans with metabolic/cardiometabolic syndrome: Amlodipine similar for chlorthalidone for all outcomes but inferior for HF (HR: 1.50; 95% Cl: 1.18–1.90) and combined CVD (HR: 1.14; 95% Cl: 1.00–1.29). Lisinopril less effective for SBP reduction by 4 mm Hg; combined CHD (HR: 1.19 (95% Cl: 1.01, 1.40); combined CVD (HR: 1.24; 95% Cl: 1.09–1.40); stroke (HR: 1.37; 95% Cl: 1.07–1.76); HF (HR: 1.49; 95% Cl: 1.17–1.90); and ESRD (HR: 1.70; 95% Cl: 1.13–2.55)
Wright JT, et al., 2009 (270) <u>19433694</u>	Study type: Race subgroup comparison of RCT comparison of an alpha blocker vs. a thiazide-type diuretic Size: 9,061	 >50 y (35.5% African American) 	Chlorthalidone vs. Doxazosin	No difference in 1° outcome (nonfatal MI and fatal CHD). Other prespecified outcomes: CHD, 1° outcome plus revascularization and hospitalized angina, composite CVD, stroke, HF, ESRD	• In African Americans: combined CVD (HR: 1.28; 95% CI: 1.16–1.42); HF (HR: 1.84; 95% CI: 1.51–2.24); stroke HR (CI): 1.10–1.73)
SPRINT Wright JT Jr, et al., 2015 (114) 26551272	Aim: To test the effectiveness of a goal SBP<120 mm Hg vs. a goal SBP<140 mm Hg for the prevention of CVD in pts with SBP≥130 mm Hg at baseline. Study type: RCT	Inclusion criteria: SBP≥130 mm Hg, with upper limit varying as number of pre-trial BP- lowering meds increased. age ≥50 y Presence of at least 1 of the following: • Clinical or subclinical CVD • CKD stage 3 or greater • Age≥75 y	Intervention: Intensive BP- lowering treatment to goal SBP<120 mm Hg Comparison: • Standard BP-lowering treatment to goal SBP<140 mm Hg • Net treatment difference ~3 drugs (2.8) on average vs. 2 drugs (1.8) on average	1° endpoint: CVD (MI, ACS, stroke, HF, CVD death) HR: 0.75 (0.64–0.89) Other endpoints: • Total deaths: • Total deaths: 0.73 (0.60–0.90) • 1° or death: 0.78 (0.67–0.90) • Components of 1° composite mostly consistent in direction other than ACS – no difference. CKD outcomes:	Summary: • More intensive SBP lowering to a goal of <120 mm Hg with achieved mean of ~121 mm Hg resulted in less CVD and lower total mortality over 3.26 y in comparison with a goal SBP<140 mm Hg and achieved SBP of ~135 mm Hg. • There were small increases in some expected SAEs. Perhaps unexpected, a sizable

	Size: 9361 participants followed median of 3.26 y	 Framingham General CVD risk≥15% in 10 y <u>Exclusion criteria:</u> Major ones included DM, history of stroke, ESRD (eGFR <20) 	• During the trial, mean SBP was 121.5 vs. 134.6.	 1° in CKD pts: reduction in GFR of ≥50% or ESRD 0.89 (0.42–1.87) Incident albuminuria: 0.72 (0.48–1.07) In pts without CKD: reduction in GFR ≥30% and to <60 3.49 (2.44–5.10) Incident albuminuria: 0.81 (0.63–1.04) Adverse events: SAEs: 1.04; p=0.25 Significant absolute increases seen in intensive group for hypotension (1%), syncope (0.6%), electrolyte abnormality (0.8%), acute kidney injury/acute 	increase in reduced eGFR in the non-CKD group and AKI/ARF overall was observed in the intensive group. While of uncertain etiology and significance, there is speculation this could be an acute hemodynamic effect, especially given the findings regarding albuminuria. <u>Limitations:</u> Few participants were untreated at baseline ~9%, so SPRINT provides little if any insight at present regarding BP-lowering medication initiation for untreated people with SBP
				 renal failure (1.6%) over the study period. 1.7% fewer pts had orthostatic hypotension in intensive group; p=0.01. 	130–139.
VA Coop 1967 (262) <u>4862069</u>	Study type: RCT to examine effect of treatment of severe HTN Size: 143	 54% African American DBP 115–129 mm Hg 	• HCTZ, Reserpine, Hydralazine vs. placebo	• CVD or stroke events, Grade 3 or 4 retinopathy, doubling of creatinine or BUN. Study terminated early for 27 events vs. 2 events (placebo vs. active)	N/A
VA Coop 1970 (271) <u>4914579</u>	Study type: RCT to examine effect of treatment of mild to moderately severe HTN	 42% African American DBP 90–115 mm Hg 	• HCTZ, Reserpine, Hydralazine vs. placebo	• CVD or stroke events, Grade 3 or 4 retinopathy, doubling of creatinine or BUN	
	<u>Size</u> : 380				

HTN Detection and Follow-up Program (HDFP) 1979 <u>6480895</u> (272)	Study type: RCT; comparison of stepped care at academic centers vs. usual care provided by community Size: 10,950 pts	 44% African American 30–69 y 	• Chlorthalidone, Reserpine, Hydralazine, Guanethidine vs. referral to community care	• 23% decrease in mortality in African Americans on Stepped Care	N/A
LIFE Dahlof B, et al. 2002 <u>11937178</u> (14)	Study type: RCT comparison of an ARB compared to a BB on CVD	 55-80 y (mean 66.9 y) African American 533 (6) White 8,503 (92) Asian 43 (0.5) Hispanic 100 (1) Other 14 (0.2) 	Losartan vs. Atenolol	• Interaction of race and treatment on CVD events (p=0.005) CVD increased 55% in African Americans in the Losartan group	N/A
VALUE Julius S, et al. 2006 (265) <u>16864741</u> (273)	Study type: RCT comparison of an ARB vs. a CCB on CVD	 >50 y (mean 67.3 y) African American 658 (4.3) White 13,643 (89.1) Asian 535 (3.5) Other 474 (3.1) 	 Valsartan vs. Amlodipine 	• CVD increased ~20% (NS) in African Americans in Valsartan group	N/A
AASK Norris K, et al. 2006 <u>17059993</u> (174)	Study type: RCT comparison of 2 BP targets and 3 drug regimens on renal outcomes Size: 1,094 pts	 18–70 y; African Americans; eGFR: 25–65 mL/min/1.73 m² 	• MAP of <92 mm Hg compared to MAP 102–107 mm Hg and an ACEI or CCB each compared to a BB	• No difference between BP targets. ACEI > BB > CCB	N/A
ALLHAT 2002 (274) <u>12479763</u>	Study type: RCT comparison of an alpha blocker, ACEI, or CCB, each compared to a thiazide-type diuretic Size: 42,418	 >50 y African American 15,085 (35.5) White 19,977 (47.0) Hispanics 5,299 (12.5) 	Chlorthalidone vs. Doxazosin, Amlodipine, or Lisinopril	 No difference in 1° outcome (nonfatal MI and fatal CHD) 	• Chlorthalidone (and amlodipine was superior in reducing BP by 4/1 mm Hg and CVD events (stroke and CVD) vs. lisinopril in African Americans
INVEST Pepine CJ, et al., 2003 (275)	Study type: RCT comparison of CCB plus an ACEI	 ≥ 50 y with HTN and CHD 36% Hispanic 	Verapamil/trandolapril vs. Atenolol/ HCTZ	•No difference in 1° outcome (nonfatal MI, nonfatal stroke, all- cause mortality). Mean SBP	N/A

<u>14657064</u>	compared to a BB plus a thiazide diuretic <u>Size</u> : 22,576	13% African American49% White		reduction Hispanics vs. non- Hispanic pts (-21.3 vs17.4 mm Hg; p<0.001)	
Wright JT, et al., 2005 (276) <u>15811979</u>	Study type: Race subgroup comparison of RCT comparison of an alpha blocker, ACEI, or CCB compared to a thiazide-type diuretic	 >50 y African American, n=11,792 Non-African American, n=21,565 	Chlorthalidone vs. Amlodipine, or Lisinopril	No difference in 1° outcome (nonfatal MI and fatal CHD). Other prespecified outcomes: CHD, 1° outcome plus revascularization and hospitalized angina, composite CVD, stroke, HF, ESRD	• In African Americans: Amlodipine similar to chlorthalidone for all outcomes but inferior for HF (HR: 1.37; 95% Cl: 1.24–1.51). Lisinopril less effective for SBP reduction by 4 mm Hg, stroke (HR: 1.40; 95% Cl: 1.17–1.68), combined CVD (HR: 1.19; 95% Cl: 1.09– 1.30), HF (HR: 1.30; 95% Cl: 1.10–1.54).

Data Supplement 52. RCTs Comparing Women With Hypertension (Section 10.2.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Turnbull F, et al., 2008 (277) <u>18852183</u>	<u>Aim:</u> Assess sex differences in response to BP treatment <u>Study type:</u> Meta- analysis of 31 RCTs <u>Size:</u> 103,268 men, 87,349 women	Mean ages: • Women: 63.0 y • Men: 61.7 y	Intervention: N/A Comparator: N/A	<u>1° endpoint</u> : Nonfatal stroke or death from cerebrovascular disease (ICD 430–438); (ii) nonfatal MI or deaths from CHD, excluding SCD (ICD 410–414); (iii) HF causing death or requiring hospitalization (ICD 428); (iv) total major CV events (stroke, CHD events, HF, other CV death); (v) total CV deaths (ICD 396–459); and (vi) total mortality Safety endpoint : N/A	<u>Summary</u> : Achieved BP reductions were comparable for men and women in every comparison made. For the 1° outcome of total major CV events there was no evidence that men and women obtained different levels of protection from BP- lowering or that regimens based on ACEIs, calcium antagonists, ARBs, or diuretics/BBs were more effective in1 sex than the other (all p-homogeneity >0.08).
Wing L, et al., 2003 (278) <u>12584366</u>	Aim: Comparison of ACE vs. Diuretic on incident CVD	Inclusion criteria: Pts 65–84 y	Intervention: ACE Comparator: Diuretic	Endpoint: All CV events or death from any cause Safety endpoint: N/A	<u>Summary</u> : Among male subjects, HR: 0.83 (95% CI: 0.71–0.97; p=0.02); among female subjects, HR: 1.00 (95% CI: 0.83–1.21; p=0.98); the p value for

	<u>Study type:</u> Practice- based RCT open label treatment, blinded event <u>Size:</u> 6,083 pts	Exclusion criteria: Life- threatening illness, contraindication to an ACEI or diuretic, a plasma creatinine concentration of more than 2.5 mg per deciliter (221 micromol per liter), malignant hypertension, or dementia	Note: Clinicians chose which ACE or diuretic		the interaction between sex and treatment-group assignment was 0.15.
Fletcher A, et al., 1988 (279) <u>2907053</u>	Aim: Monitoring event rates in pts assigned to treatment by clinicians <u>Study type:</u> Observational <u>Size:</u> 2,607	Inclusion criteria: Age >18 y Exclusion criteria: N/A	Intervention: N/A	<u>1° endpoint</u> : Total mortality incident "IHD" <u>Safety endpoint</u> : N/A	Summary: BBs reduced mortality in men but not women (p<0.01)
Forette F, et al., 2002 (280) <u>12374512</u>	Aim: Legacy follow-up for dementia prevention Study type: RCT with legacy follow-up Size: 2,902 in the legacy follow-up	Inclusion criteria: Age ≥60 y Exclusion criteria: HTN 2° to a disorder that needed specific medical or surgical treatment; congestive HF; dissecting aortic aneurysm; serum creatinine concentration at presentation of 180 micromol/I or more; stroke or MI in the y before the study; dementia; substance abuse; any disorder prohibiting a sitting or standing position; any severe concomitant or non-CVD	Intervention: Nitrendipine + HCTZ Comparator: Placebo	<u>1° endpoint</u> : Incidence of dementia <u>2° endpoint</u> : Cognitive decline measured by MMSE <u>Safety endpoint</u> : N/A • Cases Active: 21 • Cases Placebo; 43 • Rate 3.3 vs. 7.4 cases/1,000 pt y 0.38 (95% CI: 0.23–0.64; p<0.001) • MMSE: No impact	 Study discontinued early for CVD benefit so a legacy follow-up with both groups (off protocol) yielded a follow-up of 3.7 y SBP was 149 mm Hg in treatment vs. 156 mm Hg in control arm Summary dementia: Compared with the controls, long-term antihypertensive therapy reduced the risk of dementia by 55%, from 7.4–3.3 cases per 1,000 pt-y (43 vs. 21 cases; p<0.001). After adjustment for sex, age, education, and entry BP, the relative HR associated with the use of nitrendipine was 0.38 (95% CI: 0.23, 0.64), p<0.001. Lack of impact on MMSE not surprising given low sensitivity to change and large sample size

Study Acronym (if applicable) Author Year	Study Type/Design*; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Pucci M, et al., 2015 (281) <u>25612630</u>	Study type: Review of published reports of fetotoxicity of ACE/ARB antihypertensives in the first trimester of pregnancy. Usually case/control design. Size: N/A	Inclusion criteria: Pregnant women receiving ACE/ARB in the 1 st trimester of pregnancy only and comparable controls Exclusion criteria: Use of ACE/ARB later in pregnancy	<u>1° endpoint</u> : Adverse outcomes of pregnancy <u>Results</u> : Adverse events are higher in pregnancies of women who receive ACE/ARB in the first trimester of pregnancy but results are not independent of known confounders	 Fetotoxicity in the first trimester of pregnancy cannot be definitely attributed to ACE/ARB treatment; data are inconclusive. Other known causes of fetotoxicity may be responsible for increased risk in the first trimester (HTN, obesity, undiagnosed DM, other anti-hypertensives)
Moretti ME, et al., 2012 <u>22203847</u> (282)	Study type: Case control comparing pts exposed to ACE/ARB in the first trimester to healthy controls and those on other anti-hypertensives Size: 388 total pts (equally divided)	Inclusion criteria: Mothers calling into the Mother Risk Program re: medication toxicity during pregnancy Exclusion criteria: Non- English speaking	<u>1° endpoint</u> : Malformations and adverse fetal outcomes <u>Results</u> : No difference among groups but study under-powered	Supportive of above review
Ferrer RL, et al., 2000 (283) 11094241	Study type: Meta-analysis Size: 46 observational studies and randomized control trials	Inclusion criteria: Pre- specified quality entrance criteria Exclusion criteria: N/A	 <u>1° endpoint</u>: Adverse pregnancy outcomes <u>Results</u>: Maternal HTN increases risk for 1) perinatal mortality (OR: 3.4:1) and 2) placental abruption 	 HTN by itself is associated with adverse perinatal outcomes ACEIs independently are responsible for some outcomes
			(2.1:1)ACEIs are associated with fetopathy (fetal renal failure)	

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*Quality assessment analysis may need to be applied on a case-by-case basis for controversial studies (by ERC chairs).

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
SPRINT Senior Williamson JD, et al., 2016 (190) <u>27195814</u>	Aim: Intensive SBP goal <120 mm Hg) vs. standard (SBP goal <140) Study type: RCT Size: 2,636; 30% met criteria for being classified as ambulatory frail Mean follow-up:3.1 y	Inclusion criteria: Men and women age 75+; mean age 79.8 y; 38% women; 17% black, 74% Caucasian Exclusion criteria: Nursing home residents; prevalent DM, stroke, Class III/IV HF, dementia	Intervention: Medications and dietary advice to achieve SBP of <120 mm Hg <u>Comparator:</u> Medications and dietary advice to achieve SBP of <140 mm Hg • Achieved SBP: Intensive=123.4 mm Hg Standard=134.8 mm Hg	 <u>1° endpoint</u>: Composite CVD outcome (AMI, non-MI ACS, stroke, HF, CVD death. <u>Results</u>: 102 events in the intensive treatment group vs. 148 events in the standard treatment group; HR: 0.66; 95% CI: 0.51–0.85 and all-cause mortality (73 deaths vs. 107 deaths, respectively; HR: 0.67; 95% CI: 0.49–0.91. No difference in falls, orthostatic hypotension, or overall SAEs. NNT for 1° outcome=27 and NNT for all-cause mortality=41 	Limitations: Does not apply to nursing home pts or those with dementia or advance Conclusions: Intensive SBP is safe and effective for lowering CVD events and total mortality in adults ≥75 y

Data Supplement 54. RCT for Older Persons (Section 10.3.1)

Data Supplement 55. RCTs Comparing Hypertensive Crises and Emergencies (Section 11.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
CLUE Peacock WF, et al., 2011 (284) 21707983	<u>Aim:</u> Compare safety and efficacy of IV nicardipine vs. labetalol in the management of acute HTN. <u>Study type:</u> RCT	Inclusion criteria: SBP ≥180 mm Hg on 2 consecutive occasions 10 min apart in the ED.	• 110 pts randomized to nicardipine; 116 to labetalol. End-organ damage preceded randomization in 63%% with no difference between the groups. The target BP range (TR; at the discretion of the	Results: Within 39 min, nicardipine pts reached TR than labetalol pts (91.7 vs. 82.5%; p=0.039). Of 6 BP measurements taken 5 min apart, nicardipine pts had a higher rate of 5 and 6 SBP measures in the TR than labetalol pts (47.3 vs. 32.8%;	Limitations: Study unblinded; large number of pts without end-organ damage (which usually defines a hypertensive emergency); physicians ordered fewer dose titrations of labetalol than nicardipine; thus, lack of BP decline might have been due to insufficient dosing by physicians hesitant to administer successively increasing doses of labetalol as recommended by the FDA.

	<u>Size:</u> 226 pts		treating physician) was defined as SBP ± 20 mm Hg. • Dosing titrations were	p=0.026). Rescue medications did not differ between the nicardipine and labetalol groups. Nicardipine	<u>Conclusions</u> : Pts treated with nicardipine are more likely to reach the physician-specified TR than those treated with labetalol. In this study (2014), initial SBP was not a predictor of the
			those recommended by the FDA.	pts were more likely in the TR than labetalol pts (OR: 2.73; 95% CI: 1.1–6.7; p=0.028).	ability to achieve the pre-specified TR in 30 min. Subgroup analysis demonstrated the similar results for sub-populations with end-organ damage (n=141) and renal dysfunction (n=104).
Liu-DeRyke X, et al., 2013 (285) <u>23760911</u>	Aim: Compare ability of IV nicardipine and labetalol to lower BP in acute hemorrhagic or ischemic stroke. Study type: RCT (pseudo- randomization) Size: 54 pts	Inclusion criteria: Pts with acute hemorrhagic or ischemic stroke who were at or exceeded AHA guidelines BP recommendations. Exclusion criteria: Traumatic brain injury; intracranial neoplasm, received antihypertensive medication within previous 24 h, brain stem herniation, immediate brain death, acute MI, or bradycardia <50 bpm.	• 28 pts randomized to labetalol and 26 to nicardipine. Goal BP defined using the latest consensus recommendations.	Results: All pts receiving nicardipine achieved BP goal Compared with 61% in the labetalol group (p<0.001). 89% of the nicardipine group achieved goal within 60 min vs. 25% in the labetalol group (p<0.001). The nicardipine group had better maintenance of BP, greater percent of time spent within goal and less BP variability compared with the labetalol group (p<0.001). Less rescue medication had to be given to the nicardipine than the labetalol group (p<0.001).	<u>Limitations</u> : Very small; pseudo-randomization. <u>Conclusions</u> : In acutely hypertensive stroke pts, a superior BP-lowering response was achieved with nicardipine over labetalol. Despite this, there was no significant difference in clinical outcomes.
CATIS He J, et al., 2014 (202) <u>24240777</u>	Aim: Evaluate whether immediate BP reduction in pts with acute ischemic stroke would reduce death and major disability in 14 d or hospital discharge. Study type: RCT Size: 4,071 pts	Inclusion criteria: Pts had nonthrombolysed ischemic stroke within 48 h of onset and elevated SBP. Baseline SBP was 166.7 mm Hg in the antihypertensive treatment group and 165.6 mm Hg in the control group.	• This was a Chinese multicenter, single- blinded, blinded endpoints RCT conducted in 26 hospitals in China. 2,038 pts were randomized to receive antihypertensive treatment and 2,033 were randomized to the control group. The trial was designed to test a BP	Results: In the antihypertensive treatment group, SBP was reduced from 166.7 to 144.7 mm Hg (-12.7%) within 24 h and in the control group from 165.6 to 152.9 mm Hg (-7.2%) (absolute difference -9.1 mm Hg; 95% CI: -10.2– -8.1; p<0.001). Mean SBP was 137.3 mm Hg in the antihypertensive treatment	Limitations: Study excluded pts with BP ≥220/120 mm Hg, so the results do not apply to such pts. Pts treated acutely with thrombolytic therapy were excluded. Trial performed exclusively in Chinese pts. <u>Conclusions</u> : Among pts with acute ischemic stroke, BP reduction with antihypertensive medications, compared to absence of antihypertensive medications, did not reduce the likelihood of death and major disability at 14 d or hospital discharge.

INTERAC-2 Anderson CS, et al., 2013 (191) 23713578	Study type: RCT Size: 2,839 pts	•To compare the management strategy of targeting SBP<140 mm Hg within 1 h with the current guideline strategy of targeting SBP to <180 mm Hg with the use of agents of the physicians' choosing.	 reduction strategy rather than the efficacy of specific antihypertensive drugs. Pts in the control group discontinued their home BP medications. <u>1° outcome</u>: Combination of death and major disability at 14 d or hospital discharge. This was an international, multicenter, prospective randomized open-treatment, blinded endpoint trial. The pts had onset of spontaneous ICH within 6 h of enrollment. <u>1° outcome</u>: Death or major disability, defined as a score of 3-6 on the modified Rankin scale, at 90 d. 	group and 146.5 mm Hg in the control group at the 7 th d of randomization (absolute difference -9.3 mm Hg; 95% Cl: -10.1– -8.4; p<0.001). The 1° outcome did not differ between treatment groups (OR: 1.00; 95% Cl: 0.88–1.14) at 14 d or hospital discharge. The 2° outcome of death and major disability at 3 mo post- treatment follow-up did not differ between the groups. <u>Results:</u> 719 of 1,382 pts receiving intensive treatment as compared to 785 of 1,412 pts receiving guideline- recommended treatment had a 1° outcome event [OR with intensive treatment: 0.87; 95% Cl: 0.75–1.01; p=0.06). Ordinal analysis showed significantly lower modified Rankin scores with intensive treatment (OR for greater disability: 0.87; 95% Cl: 0.77–1.00; p=0.04). Mortality was 11.9% in the group receiving guideline- recommended treatment and 12.0% in the group receiving guideline- recommended treatment. Nonfatal serious events were not significantly different between the groups.	Limitations: No major limitations. Conclusions: In pts with ICH, intensive lowering of BP resulted in a borderline significant reduction in the rate of death or severe disability at 90 d. An ordinal analysis of modified Rankin scores indicated improved functional outcomes with intensive lowering of BP. Intensive BP reduction was shown to be safe and to result in significantly better health-related quality of life.
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PRONTO Peacock WF, et al., 2014 (286) <u>24655702</u>	<u>Study type:</u> RCT <u>Size:</u> 104 pts	• To determine the efficacy and safety of clevidipine vs. standard-of-care (SOC) iv antihypertensive therapy in hypertensive acute HF.	 This was a randomized, open-label, active control study of clevidipine vs. standard-of-care in ED pts with acute HF with SBP ≥160 mm Hg. <u>1° outcome</u>: Co-1° endpoints were median time to and % attaining a SBP within a prespecified TR at 30 min. 	Results: More clevidipine pts reached target BP reduction (71%) than did those receiving standard-of- care (37%) and clevidipine was faster to target (p=0.0006). Serious adverse events were similar between clevidipine and standard-of- care.	<u>Limitations</u> : Small study, open-label design. <u>Conclusions</u> : In hypertensive acute HF, clevidipine safely and rapidly reduced BP and improved dyspnea more effectively that standard- of-care.
Farias S, et al., 2014 <u>13849948</u> (287)	Aim: To determine if achievement of target BP is less likely in pts with higher initial BP using a post hoc analysis in a pt subset from CLUE <u>Study type</u> : RCT Post-hoc Analysis <u>Size</u> : 223 pts	Inclusion criteria: SBP ≥180 mm Hg on 2 consecutive occasions 10 min apart in the ED. Exclusion criteria: Contraindication to giving either a BB or CCB or clinical scenarios in which a compelling agent was indicated.	 This was a post hoc analysis of CLUE, an RCT, in which pts were dichotomized using the median presenting SBP as the partition point. Individuals above and below the median were evaluated as to the proportion achieving the 1° outcome. <u>1° outcome</u>: Achievement of target SBP range within 30 min. 	Results: Early achievement of target SBP was independent of presenting SBP.	Limitations: 2° analysis of the 1° CLUE study; SBP control only evaluated for the first 30 min posttreatment; no inclusion of critically ill pts; 80% of enrolled subjects were African-American. Conclusions: Presenting SBP does not appear to affect the ultimate ability to reduce BP for pts with marked, acute HTN in the ED when treated with either IV nicardipine or IV labetalol.

Data Supplement 56. RCTs Assessing Impact of Hypertension Therapy on Dementia Incidence (Section 11.3)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
SHEP Applegate WB, et al., 1994 (288) <u>7944835</u>	<u>Aim:</u> Compare loss of instrumental activities of daily living by SBP	Inclusion criteria: 60–80 y (mean 71.6 y)	Intervention: Chlorthalidone + Atenolol or reserpine	<u>1° endpoint</u> : Loss of dementia- related functions (instrumental activities of daily living)	Relevant 2° endpoint: Incidence of surrogate markers for dementia

	traatmaantuka	Evolucion oritorio, Lister:		Casas	Cummon u Noncignificant 1/0/ laura
	treatment vs.	Exclusion criteria: History	0	Cases	Summary: Nonsignificant 16% lower
	placebo	and/or signs of major CVDs	Comparator:	Active: 37	incidence of incident instrumental
	0	(e.g., previous MI, coronary	Placebo	Placebo: 44	activity of daily living disability.
	Study type: RCT	artery surgery, major		• p=0.84 (0.54,1.31)	However, assignment to the placebo
		arrhythmias, conduction	<u>SBP</u>	 No cognitive function instrument 	group and the resulting occurrence of
	<u>Size</u> : 4,736	defect, recent stroke, carotid	Treatment/Placebo	included in trial	CV events independently predicted
		artery disease, ≥2 TIAs and	difference:		missed assessments. However, when
	Duration: 5 y	signs or symptoms in a single	-12 mm Hg		20%–30% and 40%–80% of the
		neurological distribution); other	Achieved mean		subjects who missed the assessment
		major diseases (e.g., cancer,	SPB: 143 mm Hg in		were assumed to be
		alcoholic liver disease,	treatment group vs.		cognitively/functionally impaired,
		established renal dysfunction)	155 mm Hg in		assignment to active treatment reduced
		with competing risk factors for	placebo group		the risk of these outcomes. Thus, in the
		the 1° endpoint; stroke;			SHEP study, the cognitive and
		presence of medical			functional evaluations were biased
		management problems (e.g.,			toward the null effect by differential
		insulin dependent DM, history			dropout. This might have obscured the
		of dementia, evidence of			appraisal of a protective effect of
		alcohol abuse); bradycardia;			treatment on the cognitive and
		people maintained on BBs,			functional decline of older hypertensive
		diuretics, other			adults
		antihypertensive drugs,			
		anticoagulants.			
Syst-Eur	Aim: Incident	Inclusion criteria: ≥60 y	Intervention:	Endpoint: Dementia (defined by	Summary: Trial stopped early for
Forette F, et al.,	dementia		Nitrendipine ±	MMSE)	positive effect on CVD outcomes.
1998 (289)		Exclusion criteria: HTN 2° to	enalapril ± HCTZ	,	1
9802273	Study type: RCT	a disorder that needed specific	1	Cases:	
		medical or surgical treatment;	Comparator:	• Active: 11	
	Size: 2,418 pts	congestive HF; dissecting	Placebo	Placebo: 21	
		aortic aneurysm; serum		• (3.8 vs. 7.7 per 1,000 pt-y)	
	Duration: 2 y	creatinine concentration at	SBP	• p=0.05	
		presentation of 180	treatment/placebo	- p=0.03	
		micromoles/I or more; stroke or	difference:		
		MI in the y before the study;	-8.3 mm Hg		
		dementia; substance abuse;	Achieved SBP in		
		any disorder prohibiting a	152 mm Hg		
		sitting or standing position; any	treatment arm; 160		
		CVD	arm		
		severe concomitant or non- CVD	mm Hg placebo arm		

Syst-Eur (legacy follow-up) Forette F, et al., 2002 (280) <u>12374512</u>	<u>Aim:</u> Legacy follow- up for dementia prevention <u>Study type</u> : RCT with legacy follow- up <u>Size</u> : 2,902 pts <u>Duration:</u> 3.7 y	Inclusion criteria: ≥60 y Exclusion criteria: HTN 2°ary to a disorder that needed specific medical or surgical treatment; congestive HF; dissecting aortic aneurysm; serum creatinine concentration at presentation of 180 micromoles/l or more; stroke or MI in the y before the study; dementia; substance abuse; any disorder prohibiting a sitting or standing position; any severe concomitant or non- CVD	Intervention: Open label follow-up of Syst-Eur pts originally assigned to Nitrendipine ± enalapril ± HCTZ vs. placebo SBP Treatment/Placebo difference: -7.0 mm Hg Achieved SBP in 149 mm Hg treatment arm 156 mm Hg placebo arm	<u>1° endpoint</u> : Incidence of dementia <u>Endpoint 2:</u> Cognitive decline measured by MMSE <u>Safety endpoint</u> : N/A • Cases active: 21 • Cases placebo; 43 • Rate 3.3 vs. 7.4 cases/1,000 pt-y • 0.38 (95% CI: 0.23–0.64; p<0.001)	 This legacy follow-up with both groups (off protocol) yielded a follow-up of 3.7 y SBP was 149 mm Hg in treatment vs. 156 mm Hg in control arm Summary dementia: Compared with the controls, long-term antihypertensive therapy reduced the risk of dementia by 55%, from 7.4–3.3 cases per 1,000 pt-y (43 vs. 21 cases; p<0.001). After adjustment for sex, age, education, and entry BP, the RH rate associated with the use of nitrendipine was 0.38; 95% CI: 0.23–0.64; p<0.001. Lack of impact on MMSE not surprising given low sensitivity to change and large sample size
SCOPE Lithell H, et al., 2003 (290) <u>12714861</u>	<u>Aim:</u> Incident dementia (cognitive decline as 2° outcome) <u>Study type:</u> RCT <u>Size:</u> 4,964 <u>Duration:</u> 3.7 y	Inclusion criteria: 70–89 y (mean 76 y) Exclusion criteria: Prevalent dementia; 2° HTN, SBP >180 mm Hg, orthostatic hypotension, need for antihypertensive treatment other than hydrochlorothiazide during run-in; stroke or MI within 6 mo; decompensated HF; serum creatinine>180 micromole/I (men) or>140 micromole/I (women);	Intervention: Candesartan ± HCTZ Comparator: Placebo ± Rx for community based SPB standard SPB standard SBP Treatment/Placebo difference: -3.2 mm Hg	Endpoint: Incident dementia Also decline in MMSE Dementia Cases: Active: 62 Placebo: 57 p=1.08 (0.75–1.56) Cognitive decline slower in treatment group	 Summary: Mean follow-up 3.7 y. Treatment group SBP=144 mm Hg and placebo 147 mm Hg; thus, relatively minimal differences in achieved SBP between arms There were no significant differences between the treatment groups in either dementia or cognitive decline.
PROGRESS Tzourio C, et al., 2003 (291) <u>12742805</u>	AIM: Dementia with or without recurrent stroke Study type: RCT Size: 6,105 pts	Inclusion criteria: Prior stroke or TIA, any adult age	Intervention: Perindopril ± indapamide Comparator: Placebo	Endpoint: Dementia alone or with recurrent stroke Dementia cases: Only stroke- related dementia reduction of 34% (95% CI: 3–55), p=0.03.	Summary: Dementia alone was not affected in this trial. Only dementia associated with incident cerebrovascular accident

	Duration: 3.9 y		SBP Treatment/Placebo difference: -9.4 mm Hg • Achieved SBP in 138 mm Hg treatment arm 147 mm Hg placebo arm		
Hypertension in the Very Elderly Trial	Aim: Incident dementia 2º aim	Inclusion criteria: ≥80 y	Intervention: Indapamide ±	1° endpoint: Incident dementia	Summary: Stopped early due to benefit in 1° outcome.
cognitive function		Exclusion criteria: Prevalent	Perindopril	Events:	
assessment	Study type: RCT	dementia		Treatment=126	
(HYVET-Cog)			Comparator:	Placebo=137	
Peters R, et al., 2008 (292)	<u>Size</u> : 3,336		Placebo	 14% reduction not significant HR: 0.86 (95% CI: 0.67–1.09) 	
18614402	Duration: 2.2 y		SBP	TIK. 0.80 (95% CI. 0.07–1.09)	
			treatment/placebo		
			difference:		
			- 15 mm Hg		
			Target SBP 150		
			mm Hg		
			 Achieved SPB in treatment arm=146 		
			mm Hg		

Data Supplement 57. RCTs for Patients Undergoing Surgical Procedures (Section 11.5)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
POISE Study Group, et al., 2008 (293) <u>16875901</u>	<u>Aim</u> : Definitively establish the effects of BB therapy in pts undergoing noncardiac surgery	Inclusion criteria: Pts undergoing noncardiac surgery with, or at risk for ASVD	Intervention: extended release metoprolol succinate Comparator: Placebo	<u>1° endpoint</u> : Composite of CV death, NF MI, NF cardiac arrest <u>Results</u> : Fewer pts taking metoprolol than placebo reached the 1° endpoint, HR: 0.84; 95% CI 0.70–0.99; p=0.0399.	<u>Limitations</u> : No data for pts <45 y, no data for pts undergoing cardiac surgery <u>Conclusions</u> : This study highlights combined benefits and

Study type: RCT	However more in metoprolol group had	risk of BB regimen in noncardiac
	death HR: 1.33; 1.03–1.74; p=0.0317 and	surgery and importance of pt
<u>Size:</u> 8,351	more had stroke HR: 2.17; 1.26–3.74;	physician discussion in deciding
	p=0.0053.	upon its use.

Data Supplement 58. Observational and Nonrandomized Studies for Patients Undergoing Surgical Procedures (Section 11.5)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Howell SJ, et al., 2004 (294) <u>15013960</u>	Study type: A systematic review and meta-analysis Size: 30 observational studies	Inclusion criteria: Available crude OR for association between HTN and periop CV complications along with variance Exclusion criteria: N/A. Studies defining HTN solely on admission BP	<u>1° endpoint</u> : Periop CV complications <u>Results</u> : Pts with SBP >180 or DBP >110 mm Hg more prone to periop ischemia, arrhythmias, and CV lability OR: 1.35 (1.17–1.56).	 Pts with SBP >180 or DBP >110 mm Hg more prone to periop ischemia, arrhythmias, and CV lability OR: 1.35 (1.17–1.56). But there was no evidence that deferring surgery in such pts reduces periop risk Conclude that planned surgery should not be deferred on basis of single admission BP. History of target organ damage more important than preop BP in predicting complications
Hart GR and Anderson RJ, 1981 (295) <u>6114720</u>	<u>Study type</u> : Literature review <u>Size</u> : 72 pts BB s, 148 pts Clonidine	Inclusion criteria: Symptoms on cessation of BBs or clonidine Exclusion criteria: CP Bypass, carotid endarterectomy	<u>1° endpoint</u> : CV symptoms or events after abrupt cessation of BBs or clonidine <u>Results</u> : Symptoms of anxiety, chest pain with tachycardia, HTN, myocardial ischemia; less frequently MI may occur on abrupt withdrawal of BB or Clonidine	• Summary of case reports. CV events such as tachycardia, HTN, angina, myocardial ischemia or infarction can occur after abrupt withdrawal of BB or Clonidine. No information on incidence.
Shammash JB, et al., 2001 (296) <u>11136500</u>	Study type: Prospective observational study Size: 140 pts	Inclusion criteria: Review of 140 pts undergoing vascular surgery at university hospitals Exclusion criteria: N/A	<u>1° endpoint</u> : In-hospital mortality <u>Results</u> : 50% mortality in 8 pts with BB discontinued vs. 1.5% mortality in pts with BB continued. OR: 65.0; p=0.001	• Discontinuing BB immediately after vascular surgery may increase the risk of postoperative CV morbidity and mortality

Lindenauer PK, et al.,	Study type: Retrospective	Inclusion criteria:	1° endpoint: In-hospital mortality	Periop BB therapy is associated with a reduced
2005 (297)	cohort	Age >18 y, major		risk of in-hospital death among high-risk, but not
<u>16049209</u>	Size: 122,338 pts	noncardiac surgery	Results: On BB therapy, mortality in low risk (RCRI =0) OR: 1.43 (1.29–1.58) to	low-risk pts undergoing major noncardiac surgery.
	<u>5126</u> . 122,550 pts	Exclusion criteria:	high risk (RCRI) OR 4 or higher OR 0.57	
		contraindication to	(0.42–0.76)	
		BB therapy	· · ·	
Wallace AW, et al.,	Study type: Retrospective	Inclusion criteria:	1° endpoint: 30-d and 1-y mortality	Periop BB therapy based upon periop Cardiac
2010 (298) 20864832	study	All surgical pts at SF VAMC	Deculto: Addition of PD thorapy	Risk Reduction protocol is associated with a reduction in 30-d and 1-y mortality. Periop
20004032	Size: 38,779 operations	VAIVIC	<u>Results</u> : Addition of BB therapy associated with reduction in 30-d OR: 0.52	withdrawal of BB is associated with increased
		Exclusion criteria:	(0.33–83; p=0.006) and 1-y OR: 0.64	mortality
		N/A	(0.51–0.79; p<0.0001) mortality	,
Andersson C, et al	Study type: Retrospective	Inclusion criteria:	1° endpoint: 30-d risk of MACE and all-	Among pts with IHD undergoing noncardiac
2014 (299) 24247428	cohort study	Pts with IHD undergoing	cause mortality	surgery, use of BB associated with lower risk of 30
24247420	Size: 28,263 pts	noncardiac surgery	Results: Among pts with HF BB Rx HR:	d MACE and mortality only among those with HF or recent MI
	<u>0120</u> , 20,200 pts	nonedralae surgery	0.78 (0.67–90) for MACE and all-cause	
		Exclusion criteria:	mortality 0.80 (0.70-0.92) all-cause	
		N/A	mortality; and with recent Hx MI HR: 0.60	
			(0.42–0.86) MACE, 0.80 (0.53–1.21) all- cause mortality	
Hoeks SE, et al.,	Study type: Prospective	Inclusion criteria:	1° endpoint: 1-y mortality	Periop BB use was independently associated
2007 (300)	survey	Pts 18 y and older	<u> </u>	with lower risk of 1-y mortality while periop
<u>16935011</u>		undergoing	Results: 1 y BB use had lower mortality	withdrawal was associated with higher risk of 1 y
	<u>Size</u> : 771 pts	peripheral vascular	c/w non-BB users (HR: 0.4; 95% CI: 0.2-	mortality
		surgery	0.7); BB withdrawal had increased	
		Exclusion criteria:	mortality c/w nonusers (HR: 2.7; 95% CI: 1.2–5.9)	
		N/A	1.2 0.7)	
Barrett TW, et al	Study type: Retrospective	Inclusion criteria:	1° endpoint: Long-term mortality, median	The use of propensity-adjusted BB c/w use
2007 (301)	cohort study	Pts undergoing	follow-up 2.7 y	reduced long-term mortality by 16%
<u>17702038</u>	<u>Size</u> : 3,062 pts	vascular surgery	Results: Use of BB over study period c/w	
	<u>0120</u> . 0,002 pt3	Exclusion criteria:	no BB reduced mortality (HR: 0.84; 95%	
		N/A	Cl: 0.73–0.96; p=0.0106)	
London MJ, et al.	Study type: Retrospective	Inclusion criteria:	1° endpoint: All-cause 30-d mortality and	BB therapy was associated with lower rates of
2013 (302)	cohort analysis	Pts undergoing major	cardiac morbidity (cardiac arrest, or non-Q	30-d all-cause mortality in pts with ≥ 2 Revised
<u>23613075</u>	Size: 136,745 pts	noncardiac surgery	wave MI	Cardiac Index Factors
	<u>Jize</u> . 130,743 pts			

		Exclusion criteria: N/A	Results: BB exposure lower 30-d mortality in pts with 2 or more RCIF (RR: 0.63; 95% CI: 0.50–0.80; p<.001)	
Turan A, et al. 2012 (303) <u>22253266</u>	Study type: Matched observational study Size: 79,228 pts	Inclusion criteria: Pts with noncardiac surgery	<u>1° endpoint</u> : Intraoperative and post- operative upper airway complications, in- hospital complications, and 30-d mortality	• No association found between use of ACEIs and intraoperative or postoperative upper airway complications, in-hospital complications, or 30-d mortality
		Exclusion criteria: N/A	<u>Results</u> : ACEI usage was not associated with either 30-d mortality (OR: 0.93; 95% CI: 0.73–1.19; p=0.22	
Rosenman DJ, et al 2008 (304) <u>18698608</u>	Study type: Review of observational and randomized studies	Inclusion criteria: Adult pts, most >18 y, nonemergent surgery, using ACEI	<u>1° endpoint</u> : Hypotension requiring vasopressors at or shortly after induction of anesthesia	• Pts receiving immediate preoperative ACEI or ARA were more likely to develop hypotension requiring vasopressors at or shortly after induction of anesthesia. Sufficient data were not present to
	<u>Size</u> : 434 pts	or ARA chronically Exclusion criteria: N/A	<u>Results</u> : Pts receiving preoperative ACEI or ARA more likely to develop hypotension requiring vasopressors. RR: 1.51; 95% CI: 1.14–2.01	assess other outcomes.
Roshanov P.S., et al. 2017 (305) <u>27775997</u>	Study type: International prospective cohort	Inclusion criteria: Pts at least 44 y undergoing	<u>1° endpoint</u> : 30-d all-cause death, stroke, or myocardial injury	• Withholding ACEI/ARB before major noncardiac surgery was associated with a lower risk of death and postoperative vascular events.
	Size: 14,687 pts	noncardiac surgery requiring overnight hospital admission <u>Exclusion criteria</u> : N/A	<u>Results</u> : ACEI/ARB users who withheld ACEI/ARB in the 24 H before surgery were less likely to suffer death, MI or stroke 0.82; 95% CI: 0.70–0.96; p=0.01	

Data Supplement 59. RCTs of Adherence and Compliance with Fixed Dose Combinations Regimens (Section 12.1.1)

Study Acronym;	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if any);
Author;	Study Type;		(# patients) /	(Absolute Event Rates,	Study Limitations;
Year Published	Study Size (N)		Study Comparator	P value; OR or RR; & 95%	Adverse Events
			(# patients)	CI)	

COMFORT Matsumura K, et al., 2012 (306) <u>22447014</u>	<u>Aim</u> : Evaluate whether a combination pill of antihypertensive drugs improves medication adherence in hypertensive pts vs. use of single agents. <u>Study type</u> : Multicenter, open, RCT at 29 sites in Japan. Adherence assessed by pill count. <u>Size</u> : 207 pts	Inclusion criteria: • ≥20 y agent with HTN • Could be treated with an ARB and diuretic Exclusion criteria: • Extremely high BP (≥200 mm Hg SBP or ≥120 mm Hg DBP) • Serious renal or liver dysfunction • Taking >4 tablets, excluding study drugs	Intervention: Combination tablet of (Losartan 50 mg/HCTZ 12.5 mg; n=103) <u>Comparator</u> : ARB and a thiazide diuretic as separate agents (n=104)	<u>1° endpoint</u> : Adherence rates as assessed by pill count 98% in both groups (p=0.89) over entire study period (0–6 mo). <u>Safety endpoint</u> : No differences in serious adverse events (1% vs. 1%; p=0.99) or mild adverse events (6% vs. 10%; p-0.31)	 <u>2° endpoint</u>: No significant difference in mean SBP and DBP (0.3 and 0.1 mm Hg respectively; p=0.84/0.96). <u>Study limitations</u>: Adherence rate very high for both groups and likely does not represent real-world rates. Short duration (6 mo) and thus does not provide much information on medication persistence (continuation of drug therapy long-term) Possible selection bias with 2 run-in phases Different healthcare system (Japan) with medications provided through public medical insurance
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Data Supplement 60. Nonrandomized Trials, Observational Studies, and/or Registries of Antihypertensive Medication Adherence Strategies (Section 12.1.1)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% Cl)	Summary/Conclusion Comment(s)
Schroeder K, et al., 2004 (307) <u>15078641</u>	Study type: Systematic review of RCTs. Size: 38 studies testing 58 different interventions containing data on 15,519 pts; 9 studies assessed simplification of dosing regimen	Inclusion criteria: • Database search for all RCTs, all languages, in Cochrane Controlled Trials Register, MEDLINE, EMBASE, and CINAHL (all y through 2002) • Population of interest were pts with essential HTN in primary care, outpatient, or community setting • Interventions aimed to increase adherence to BP-lowering medication • Reported outcome was adherence	 <u>1° endpoints</u>: Adherence as assessed by pill counts, self-report, or electronic monitoring system <u>Results</u>: 9 studies assessed simplification of dosing regimen, 7 of which compared adherence associated with frequency of administration (twice daily vs. once daily [n=6] or 3 times daily vs. twice daily [n=1]). All studies examining effect of dosing frequency demonstrated improved adherence (range: 8%, 19.6% improvement; p<0.01 for all). 	• Adherence to antihypertensive medication was significantly improved with once daily vs. multiple daily dosing regimens. Most studies used an electronic monitoring system. Limitations in the systematic review include heterogeneity in pts, interventions, and outcomes, and the majority of studies were of low quality. In addition, different definitions of adherence in the RCTs make it difficult to examine the precise relationship of adherence to BP control.

		• RCT where pt care in intervention group(s) compared to either no intervention or usual care	• Only 1 of the 7 studies demonstrated improved BP control (change in SBP 6 mm Hg; p<0.01). However, different medications used for comparison (once daily amlodipine 5 mg vs. diltiazem SR 60 mg twice daily).	
Iskedjian M, et al., 2002 (308) <u>11911560</u>	<u>Study type</u> : Meta-analysis <u>Size</u> : 8 studies involving a total of 11,485 observations (1,830 for once daily dosing, 4,405 for twice daily dosing, 4,147 for >twice daily dosing, 9,655 for maximum daily dose).	 Inclusion criteria: Database search of MEDLINE, EMBASE, and International Pharmaceutical Abstracts (1980– 1998) 1° studies that compared adherence rates between different dosing regimens Prospective trials (e.g., RCTs, cohort studies), retrospective studies, database analyses Any published study using an instrument to measure adherence, but must have used some measurement tool in each comparison group. Adherence rates to solid, oral dosage form for treatment of HTN of at least 10 wk duration 	 1° endpoints: Medication adherence rates compared between once daily and maximum daily dose, once daily and twice daily, twice daily and >twice daily Results: Average adherence rates with once daily dosing were greater compared to maximum daily dose regimens (91.4% [SD=2.2%] vs. 83.2% [SD=3.5%]; z=4.46; p<0001.) Average adherence rates with once daily dosing were greater compared to twice daily dosing regimens (92.7% [SD=2.3%] vs. 87.1% [SD=2.9%]; z=2.22; p=0.026.) There was no difference in adherence rates between regimens dosed twice daily or greater than twice daily (90.8% [SD=4.7%] vs. 86.3% [SD=6.7%]; z=1.82; p=0.069). 	• Antihypertensive regimens dosed once daily were associated with significantly improved adherence compared to twice daily or maximum daily dose regimens.
Claxton AJ, et al., 2001 (309) <u>11558866</u>	Study type: Systematic review Size: 76 studies	Inclusion criteria: • Database search of MEDLINE, Psychinfo, HealthStar, Health & Psychological Instruments, and Cochrane library 1986–2000 • Compliance rates assessed using electronic monitoring device • Data pooled to calculate mean compliance with once daily, twice daily, 3 times daily, and 4 times daily dosing regimens	 <u>1° endpoints</u>: Mean compliance rates by prescribed dose regimen <u>Results:</u> 26 studies evaluated CVD; 17 HTN only. For all studies, mean dose-taking compliance defined as number of appropriate doses taken during each d was 79% for once daily, 69% for twice daily, 65% for 3 times daily and 51% for 4 times daily dosing (p≤0.001 for once daily vs. 3 times daily, once daily vs. 4 times daily, and twice daily vs. 4 times daily; no statistically significant between once daily vs. twice daily or twice daily vs. 3 times daily dosing). 	• Medication compliance as measured by electronic monitoring devices were improved with less frequent dosing. Once-daily dosing was associated with the greatest rate of compliance. Limitations of this analysis include heterogeneity of studies and disease states studied.

Sherrill B, et al., 2011 (310) 22142349	Study type: Meta-analysis to compare health resource use cost, adherence, and	Inclusion criteria: • Database search of PubMed, EMBASE, The Cochrane Library,	 For 14 studies that assessed ability to take doses within prescribed time frame, once daily regimens were associated with better dose-time compliance (74% ± 31%) compared to twice daily (58% ± 23%) or 3 times daily (46% ± 8%); formal statistical analysis not conducted due to too few studies. <u>1° endpoints</u>: Health care costs, adherence, persistence 	Medication adherence and persistence was significantly greater with SPC than free-equivalent
22142347	persistence between groups of pts taking antihypertensives as SPCs vs. free-equivalent components. <u>Size</u> : 15 retrospective database studies in HTN	 ENBASE, The Countrie Library, and EconLit (no limit on publication dates) English-language publications Clinical trial or observational study (e.g., database or registry) that compared SPC with free-equivalent components Data on compliance, adherence, persistence, and/or health care costs and/or resource use (unadjusted cost analyses) 	 <u>Results:</u> All-cause total costs were estimated to be lower with SPC vs. free-equivalent components free-equivalent components by \$2,039 (95% CI: \$1030, \$3047) in 2009 dollars and HTN/CV- related costs were lower by \$709 (95% CI: \$117, \$1,032), 2009 dollars. Adherence as measured by MPR was greater for SPC vs. free-equivalent components (total inverse variance 13.31; 95% CI: 8.26–18.35). Persistence to therapy was greater with SPC than free-equivalent components (risk ratio: 2.13; 95% CI: 1.11–4.09) 	with SPC than nee-equivalent components. Costs were also significantly lower with SPC than with free-equivalent components. However, cost data should be interpreted with caution considering unadjusted costs were used in this meta-analysis. In addition, heterogeneity was present in analyses of each outcome. This meta-analysis did not include the observational study by Yang et al. as that study used an adjusted analysis methodology.
Yang W, et al., 2010 (311) <u>20629600</u>	Study type: Observational analysis using multivariate regression-adjusted analysis to compare compliance/persistence, health care resources, and cost associated with SPC or FC antihypertensives over 6 mo study period both nationally and at the state level. Size: 579,581 pts (382,476 SPC and 197,375 FC) identified in MarketScan Database (2006–2008)	Inclusion criteria: • Pts in MarketScan Database • Diagnosis of HTN based on ICD- 9 codes 401.xx and 405.xx • Pts initiated on any of the following SPC treatments or the same FC: ARB + CCB, ARB + HCTZ, ACEI + HCTZ • For SPC cohort, at least 1 prescription filled in observational window • For FC cohort, pts filled individual components separately within 15 d of each other and with 15 d overlap of supply • ≥18 y	 Endpoints: 1° outcome: Compliance and persistence with the index therapy (SPC or FC) measured by MPR within 6 mo of index date 2° outcomes: Healthcare resource utilization (number of all-cause hospitalizations, number ER visits, number CV hospitalizations, and CV-related ER visits) and health care costs (all cause medical costs, all-prescription drug costs, CV-related medical service costs, and HTN prescription-related drug costs) Results: Compliance nationally as assessed by MPR was improved in pts taking SPC vs. FC antihypertensives (difference=11.6%; 95% CI: 11.4%–11.7%). 	• This large observational study found that medication compliance/persistence to antihypertensives was improved with SPC compared to FC using an adjusted multivariate regression model. All-cause medical costs were also decreased with the used of SPC antihypertensives, although prescription costs were greater.

		 Continuous eligibility in database for 6 mo after index date Valid 3-digit zip code in database 	 Treatment discontinuation rates were lower with SPC vs. FC antihypertensives (40.7% vs. 59.3%; 95% CI: 0.46–0.48). There were fewer all-cause hospitalizations and ER visits in SPC vs. FC pts IRR: 0.77 (95% CI: 0.75–0.79) and IRR: 0.87 (95% CI: 0.86, 0.89), respectively. All-cause medical costs were reduced with SPC vs. FC (-\$208; 95% CI: -\$302– -\$114), but antihypertensive prescription costs were greater (\$53; 95% CI: \$51–\$55). 	
Gupta, et al., 2010 (312) <u>20026768</u>	Study type: Meta-analysis to assess compliance, adherence, persistence, BP control, and safety with FDC antihypertensives compared to their free components Size: 15 studies (n=32,331) with ≥1 evaluated outcome; 3 cohort studies and 2 trials of compliance (n=17,999); 3 cohort studies on persistence (n=12,653); 5 trials of adverse drug effects of FDCs (n=1,775); 9 trials of BP change (n=1,671)	 Inclusion criteria: Database search of PubMed (1966–February 2008), Web of Science (1970 to April 2008), and the Cochrane Controlled Trial (1800–April 2008). Clinical trials or cohort studies included if published in English and compared an FDC of hypertensive agents with free-drug combination of its components. Extractable data reported including compliance (or adherence), persistence, BP- lowering effects, adverse effects 	 <u>1° endpoint</u>: Compliance (or adherence) and persistence to therapy BP-lowering efficacy Adverse effects <u>Results:</u> Use of FDC therapy was associated with a 21% increase in compliance, both in the cohort studies (n=5) and clinical trials (OR: 1.21; 95% Cl: 1.00–1.47) and (OR: 1.21; 95% Cl: 1.03–1.43). There was a 50% increase in persistence with therapy, but this was not statistically significant (OR: 1.54: 95% Cl: 0.95–2.49). Analysis of all 6 retrospective cohort studies indicated that FDC therapy was associated with a 29% increase in compliance and persistence to therapy (OR: 1.29; 95% Cl: 1.11–1.50). No sign of heterogeneity of publication bias. FDC therapy was associated with a nonsignificant reduction in SBP (-4.1 mm Hg; 95% Cl: -9.8–1.5 mm Hg; p=0.15) and DBP (-3.1 mm Hg; 95% Cl: -7.1–0.9 mm Hg; p=0.13) compared to free-drug combinations. Strong evidence of heterogeneity but no evidence of publication bias. FDC therapy was associated with a 20% nonsignificant decrease in adverse effects (OR: 	• Use of FDC therapy is associated with significant improvements in compliance and persistence to antihypertensive therapy and possible improvement in BP control and decreased risk of adverse effects.

Bangalore S, et al., 2007 (313) <u>17679131</u>	Study type: Meta-analysis to assess if compliance is improved with FDC therapy compared to free-drug regimens in chronic diseases including HTN, HIV, tuberculosis, and DM	Inclusion criteria: • Database search of MEDLINE (1966–2005) • Studies included if published in English and compared an FDC with free-drug combination of its components and reported medication compliance (adherence)	 0.80; 95% CI: 0.58, 1.11) compared to freedrug combinations. <u>1° endpoint</u>: Compliance, considered as either adherence or persistence to medication therapy <u>Results:</u> Use of FDC therapy was associated with a 26% decreased risk of noncompliance vs. freedrug combinations (pooled RR: 0.74 (95% CI: 0.69, 0.80), p<0.0001) in all diseases states. 	• Use of FDC combination therapy in hypertensive pts was associated with a 24% decreased risk of noncompliance compared to use of free-drug regimens.
	Size: 9 studies total (n=20,242), 4 of which were in hypertensive populations (n=17,175)	or persistence	 There was no evidence of heterogeneity. In hypertensive pts, FDC was associated with 24% decreased risk of noncompliance (pooled RR: 0.76 (95% CI: 0.71, 0.81), p<0.0001) compared to free-drug regimen. There was no evidence of publication bias. Marked heterogeneity in how compliance was measured among studies 	
Kumagai N, et al., 2013 (314) <u>23072348</u>	Study type: Prospective, multicenter, observational study of pts converted from free-drug combinations of an ARB and amlodipine to the same product as a FDC. Size: 196 pts	Inclusion criteria: • Outpatients with essential HTN • Self-measured home BP • Prescribed FDC of an ARB (8 mg candesartan, 80 mg valsartan, or 40 mg telmisartan) and 5 mg) and 5 mg amlodipine • Pts divided into 2 groups: Group 1 received an ARB and amlodipine in the morning as free drug combinations and Group 2 took ARB in the morning and amlodipine in the evening. After 1 mo, both groups converted to once daily FDC product. Exclusion criteria: • Severe renal of liver dysfunction • Severe HF • Prescription of time-specific packs	 Endpoints: Adherence to antihypertensive therapy as measured by self-reporting Self-monitored BP measurements and clinical BP measurements before and after switch to FDC antihypertensive therapy. Drug costs Results: Self-monitoring BP measurements taken during early morning was lower with FDC compared to free-drug combinations (-5 mm Hg SBP, -2 mm Hg DBP; p<0.01 for both) Average clinic BP was lower with FDC compared to free-drug combination (-5 mm Hg SBP, -2 mm Hg SBP; p<-0.01). Self-reported adherence was improved with FDC vs. free-combination agents (~99% vs. 95% p<0.01). SBP was significantly lower in the group with improved adherence (~7.5 mm Hg) 	• Use of FDC with an ARB and amlodipine was associated with improved adherence, lower BP, and decreased health care costs compared to free-drug combination therapy. Limitations to this study include the observational design, low numbers of pts, use of self-reported adherence, short follow-up period, non-U.S. country with a different health care system (Japan), and very high baseline rate of adherence (~95%) as well post-switch to FDC (~99%), which is not what is seen in usual practice.

			 compared to the group without improved drug adherence (~4 mm Hg; p<0.05). Healthcare costs were decreased by 31% per pt from 17,075 yen (\$216.93 USD; Aug. 2012) to 11,815 yen (\$150.10 USD; Aug. 2012) over the 3 mo period. 	
Mazzaglia G, et al., 2009 (315) <u>19805653</u>	Study type: Retrospective cohort Size: 18,046 pts	Inclusion criteria: Newly diagnosed and treated hypertensive pts ≥35 y initially free of CVD identified from Italian general pt registry. Exclusion criteria: CHD, cerebrovascular disorders, congestive HF who had been hospitalized for CABG or coronary angioplasty, those recovered in a cardiology ward before index diagnosis, incident CV event in the 180 d after index diagnosis, pts receiving nitrates	1° endpoint: Describe adherence to antihypertensive therapy and its associate with concurrent drug use, comorbidities, and CV risk factors. Adherence was estimated by calculating the proportion of days which pt had pills available during the follow-up. <u>Results:</u> At baseline (6 mo after index diagnosis), adherence rates were high (≥80% proportion of days covered) in 8.1% of pts, intermediate (40-79% proportion of d covered) in 4.5%, and low (≤40% proportion of d covered) in 51%. Multiple drug treatment (1.62; 95% CI: 1.43–1.83), dyslipidemia (1.52; 95% CI: 1.24–1.87), DM (1.40; 95% CI: 1.15–1.71), obesity (1.50; 95% CI: 1.26–1.78) and antihypertensive combination therapy (1.29; 95% CI: 1.15–1.45) were associated with high adherence to treatment (p<0.001).	• High adherence was associated with a 38% decreased risk of CV events compared with low adherence. Combination therapy associated with 29% improved adherence compared to monotherapy.
Jackson KC, et al., 2008 (316) <u>18803997</u>	<u>Study type</u> : Retrospective cohort study <u>Size</u> : 908 pts	Inclusion criteria: • ≥18 y and diagnosis of HTN • Benefit-eligible for pharmacy claims • Antihypertensive naive (no prescription fill for antihypertensive drug ≥110 d prior to index date) • Received 1 of 3 regimens: 1.) 2 pill regimen with valsartan + amlodipine, 2.) 2-pill regimen with valsartan/HCTZ in FDC + amlodipine, 3.) 3-pill regimen with valsartan + HCTZ + amlodipine as free-drug components	<u>1° endpoint</u> : Adherence as measured by MPR <u>Results</u> : 224 pts received valsartan + amlodipine, 619 received valsartan/HCTZ + amlodipine, and 65 received valsartan + HCTZ + amlodipine. MPR ratios were 75.4% with valsartan + amlodipine, 73.1% with valsartan/HCTZ + amlodipine, and 60.5% with valsartan + HCTZ + amlodipine (p=0.005). Older age was associated with improved MPR (75.2% for those \geq 64 y. vs. 69.6% for 18 to <36 y; p=0.023).	• An inverse relationship existed between the number of pills and adjusted MPR, with lower adherence noted in 3-pill regimens vs. 2-pill regimens.

		Exclusion criteria: Pts who received <2 prescription fills, did		
		not continuously have prescriptions refilled for each medication, or switched from1 medication to		
		another without a time overlap		
Dickson M, et al., 2008 (317) <u>18303937</u>	Study type: Retrospective cohort study Size: 5,704 pts	Inclusion criteria: • 65–100 y on index date • Received at least 2 prescriptions for study drugs (amlodipine/benazepril FDC n=2336] or DHP-CCB and ACEI as separate agents [n=3368] between 1997–2001 • Continuously eligible for Medicaid for 12 mo following index date Exclusion criteria: • >180 d of hospitalization • <30 d of study drug supply • Any nursing home claims during the12 mo follow-up period	 <u>1° endpoint</u>: Determine rates of compliance (MPR) and total costs of care (defined as sum of payments for Medicaid claims for ambulatory care, hospital claims, prescription drug claims, and Medicare ross claims) in pts treated with FDC amlodipine/benazepril vs. a DHP-CCB and ACEI prescribed as free-combination agents. <u>Results</u>: MPR was significantly higher for pts receiving FDC compared with free-combination therapy (63.5% vs. 49%; p<0.05). Average total cost of care (2002 value) was \$3,179 with FDC compared to \$5,236 with free-combination agents (p<0.0001). Multivariate regression analysis indicated an increase of 0.5% for each 1-unit increase in MPR, and for each comorbidity there was a 10.4% increase. Total cost of care for FDC group was 12.5% lower than free-combination group (p<0.003) 	• FDC combination therapy with amlodipine/benazepril was associated with better compliance than a DHP-CCB and ACEI as free- combination agents. FDC was also associated with lower total costs of care.

Data Supplement 61. RCTs and Meta-analysis on Strategies to Promote Lifestyle Modification (Section 12.1.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% Cl)	Relevant 2° Endpoint; Study Limitations; Adverse Events Summary
Artinian NT, et al., 2010 (318) 20625115	<u>Aim</u> : To provide evidence-based recommendations on implementing PA and dietary interventions among adults,	Inclusion criteria: Included studies were limited to adult pts ≥18 y; English language; randomized controlled or quasi-experimental designs	Cognitive-behavioral strategies for promoting behavior change including Goal Setting, Self- Monitoring, Frequent and Prolonged Contact, Feedback and Reinforcement, Self-Efficacy Enhancement, Incentives, Modeling, Problem Solving, Relapse Prevention, Motivational	Variable, too numerous to summarize here.	• Variable, too numerous to summarize here.

Eckel RH, et al.,	including adults of racial/ethnic minority and/or socioeconomically disadvantaged populations. Study type: Literature review, evidence synthesis and recommendations using ACC/AHA evidence grading. Size: 70 studies, including 65 RCTs published from 1997–2007.	or meta-analyses; focused on the effects of diet or PA interventions on weight, BP, PA level, aerobic and resistance exercise, fitness, or consumption of calories, fruits, vegetables, fiber, total fat, saturated fat, cholesterol or salt <u>Exclusion criteria</u> : Feeding trials, observational studies of specific nutrients, and observational studies of aerobic capacity were excluded. Given the varying goals and outcomes of the different identified intervention studies, when possible we used a common measure of effect size to quantify and compare the success of each intervention. Inclusion criteria: N/A	Interviewing; also Intervention Processes or Delivery Strategies, including Targeting Single Behaviors Versus Multiple Behaviors, Print- or Media-Only Delivery Strategies, Group, Individual, Technology, and Multicomponent- Based Delivery Strategies, Group-Based Interventions, Individual-Focused Interventions, Computer/Technology-Based Interventions, and Multicomponent Intervention Delivery Strategies; also, Special Considerations for Interventions With Minority and Socioeconomically Disadvantaged Populations, including Setting in Which Healthcare Is Delivered, Peer/Lay Led Versus Professionally Led, Cultural Sensitivity, Literacy Level Sensitivity, Barriers to Behavior Change, and Acculturation. In addition, Fostering Initiation and Maintenance of Behavior Change. <u>Comparator</u> : Usual care or other comparison group	N/A	N/A
2013 (319) 24239922	Guideline	Exclusion criteria: N/A	group		

Data Supplement 62. RCTs, Meta-analyses, and Systematic Reviews on the Effect of Structured, Team-based Care Interventions for Hypertension Control (Section 12.2)

Study Acronym;	Aim of Study;	Patient Population	Study Intervention (#	Endpoint Results	Relevant 2° Endpoint (if any);
Author;	Study Type;		patients) /	(Absolute Event Rates, P	Study Limitations;
Year Published	Study Size (N)		Study Comparator (#	value; OR or RR; & 95% Cl)	Adverse Events
			patients)		Summary

Brownstein JN, et al., 2007 (320) <u>17478270</u>	Aim: Examine the effectiveness of community health workers in supporting the care of pts with HTN Study type: Systematic review Size: 14 studies, including 8 RCTs	Inclusion criteria: Studies examining the effects of an intervention involving community health workers on the care of pts with HTN Exclusion criteria: Studies that focused exclusively on outcomes among community health workers and those involving peers who merely led support groups	Intervention: Community health workers as HTN care team members. Community health workers were broadly defined as health workers who were trained as part of an intervention, had no formal paraprofessional designation, and had relationship with the community being served. The community health workers, predominantly women, were recruited from the community, and resembled the pts in race/ethnicity and socioeconomic background. Roles included: (1) providing health education and information to pts and families; (2) ensuring that pts received services necessary for BP control; (3) providing direct services, including measuring and monitoring BP; (4) providing social support to the pts and their family members; and (5) serving as mediators between pts and the healthcare and social service systems.	<u>1º endpoint</u> : Differences between groups in BP control groups favored community health worker groups over control and ranged from 4%– 46% over 6–24 mo, across 7 RCTs; though 1 RCT showed no difference between groups. <u>Safety endpoint</u> : N/A	 <u>2° endpoints:</u> Appointment keeping: significant improvements ranging from 19%–39% (relative changes) over 12–24 mo in community health worker intervention Adherence to medications: Range of findings included significant improvement in community health worker intervention group compared with control, between-group differences ranged from 8%–14%; 26% greater compliance among pts receiving intense community health worker interventions; and 17% significant improvement in adherence to medication with counseling by community health workers. <u>Limitations</u>: High level of heterogeneity of the populations, settings, interventions, and outcomes <u>Summary</u>: Including community health workers as part of the HTN care team resulted in significant improvements BP control, appointment keeping, and adherence to antihypertensive medications, primarily among low income, urban African Americans.
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Carter BL, et al., 2009 (321) <u>19858431</u>	Aim: Determine potency of interventions for BP involving nurses and pharmacists Study type: Meta- analysis Size: 37 RCTs of team- based HTN care involving nurse or pharmacist intervention	Inclusion criteria: RCT of team-based HTN care involving nurse or pharmacist intervention Exclusion criteria: Absence of above	Intervention: Team-based HTN care involving nurse or pharmacist intervention in nearly all studies involving nurses or pharmacists in clinics, consistent and dedicated case management activities were provided that were distinct from traditional nursing or pharmacist duties. However, pharmacists in community pharmacies usually had to incorporate the intervention with traditional medication dispensing functions. Comparator: Usual care	<u>1° endpoint</u> : OR (95% Cl) for controlled BP were nurses: 1.69 (1.48, 1.93); pharmacists within primary care clinics: 2.17 (1.75, 2.68); and community pharmacists: 2.89 (1.83, 4.55). Mean (SD) reductions in SBP were: nurse intervention, 5.84 (8.05) mm Hg; pharmacists in clinics, 7.76 (7.81) mm Hg; and community pharmacists, 9.31 (5.00) mm Hg. There were no significant differences between nurse and pharmacist effects (p \geq 0.19). <u>1° Safety endpoint</u> : N/A	 Stepwise regression was used to compare studies that included a given intervention strategy with studies that did not. Several individual components of the interventions were associated with significant reductions in mean SBP including pharmacist recommended medication to physician (-27.21 mm Hg; p=0.002), counseling about lifestyle modification (-12.63 mm Hg; p=0.03), pharmacist performed the intervention (-11.70 mm Hg; p=0.03), use of a treatment algorithm (-8.46 mm Hg; p<0.001), completion of a drug profile and/or medication history (-8.28 mm Hg; p=0.001), and the overall intervention potency score assigned by the study reviewers (p<0.001). The factors associated with a reduction in DBP were: referral was made to a specialist (-19.61 mm Hg; p=0.04), providing pt education about BP medications (-17.60 mm Hg; p=0.003), completion of a drug profile and/or medication history (-7.27 mm Hg; p=0.006), pharmacist performed the intervention (-3.94 mm Hg; p=0.04).
					Summary: Interventions involving pharmacists or nurses were associated with significantly improved BP control.
Clark CE, et al., 2010 (322) <u>20732968</u>	<u>Aim</u> : Review trials of nurse led interventions for HTN in primary care to clarify the evidence base, establish whether nurse prescribing is an important intervention	Inclusion criteria: RCT of nursing intervention for HTN Exclusion criteria: Absence of above	Intervention: Interventions were categorized as nurse support delivered by either telephone, community monitoring or nurse led clinics. These were held in either primary care or 2° care. 1 study used alternate	<u>1° endpoint</u> : • Compared with usual care, Interventions that included a stepped treatment algorithm showed greater reductions in SBP (weighted MD -8.2 mm Hg (95% CI: -11.5– -4.9);	<u>Summary</u> : Nurse led interventions that included a stepped treatment algorithm or nurse led prescribing showed significantly greater reductions of SBP and DBP than usual care. Telephone monitoring was associated with higher achievement of study targets for BP. Community monitoring showed lower

	Study type: Meta- analysis Size: 32 RCTs of nursing intervention for HTN		sessions with nurses at home and in general practice. 14 studies included a stepped treatment algorithm and 9 included nurse prescribing in the protocol. <u>Comparator</u> : Usual care	 Nurse prescribing showed greater reductions SBP, -8.9 mm Hg, (95% CI: -12.5 5.3), and DBP, -4.0 mm Hg, (95% CI: -5.32.7); Telephone monitoring showed higher achievement of BP targets (RR: 1.24; 95% CI: 1.08–1.43); Community monitoring showed greater reductions in (weighted MD) SBP, -4.8 mm Hg, (95% CI: -7.02.7), and DBP, -3.5 mm Hg, (95% CI: -4.52.5). Safety endpoint: N/A 	outcome SBP, greater reductions in SBP and DBP, and, although pooling of data was not possible, greater achievement of study BP targets.
Proia KK, et al., 2014 (323) <u>24933494</u>	Aim: Examine current evidence on the effectiveness of team- based care in improving BP outcomes (update of prior systematic review) <u>Study type</u> : Systematic review <u>Size</u> : 52 studies of team-based primary care for pts with 1° HTN	Inclusion criteria: Study of team-based care; conducted in a high-income economy; reported at least 1 BP outcome of interest; included a comparison group or had an interrupted time-series design with at least 2 measurements before and after the intervention; targeted populations with 1° HTN or populations with comorbid conditions such as DM as long as the 1° focus of the intervention; and did not	Intervention: Team-based care was defined as adding new staff or changing the roles of existing staff to work with a PCP for HTN care. Team members who collaborated with pts and PCPs were predominantly nurses (28 studies); pharmacists (15 studies); both nurses and pharmacists (5 studies); or community health workers, integrated care managers, or behavioral interventionists (4 studies). Key roles included HTN medication management, active pt follow-up, and adherence and self- management support. Interventions were usually	 <u>1° endpoint</u>: Proportion with controlled BP: Absolute percentage point (pct pt) change in pts with controlled BP from 33 studies comparing team- based care to usual care: median effect estimate was 12 pct pts (IQI=3.2–20.8 pct pts). Most individual effect estimates in the favorable direction were significant (p<0.05). Reduction in SBP (44 studies): The median reduction in SBP was 5.4 mm Hg (IQI=2.0–7.2 mm Hg). Most individual effect estimates were significant (p<0.05). Reduction in DBP: The overall median reduction in 	 <u>2° endpoints</u>: Compared with pts in usual care, the proportion of pts receiving team-based care with "high" medication adherence (defined as taking medications as prescribed >80% of the time) increased by a median of 16.3 pct pts (9 studies). <u>Stratified analyses for BP outcomes</u>: Team member role in medication management: Larger improvements in BP outcomes than overall estimates were demonstrated when team members could make changes to medications independent of the PCP or team members could provide medication recommendations and make changes with the PCP's approval as compared to team members providing only adherence support and information on medication and HTN.

		Exclusion criteria: Inclusion of populations with 2° HTN (e.g., pregnancy) or with a history of CVD (e.g., MI)	implemented across multiple settings in the healthcare system and in the community, where they were implemented in pharmacies and through home outreach visits. Comparator: Usual care	DBP was 1.8 mm Hg (IQI=0.7–3.2 mm Hg) from 38 studies. Safety endpoint: No harm to pts was identified from team- based care interventions in the included studies or the broader literature.	 Number of team members added: Adding ≥2 members demonstrated larger improvements in the proportion of pts with controlled BP and reduction in DBP compared to adding only 1; median reductions in SBP were similar regardless of team size. Improvement in the proportion of pts with controlled BP was similar for studies from both healthcare and community settings. <u>Limitations</u>: Included studies reported significant differences in pt demographics between intervention and comparison groups at baseline, possible contamination within intervention and implemented interventions. <u>Summary</u>: There is strong evidence that team- based care is effective in improving BP outcomes, especially when pharmacists and nurses are part of the team.
Santschi V, et al., 2014 (324) <u>24721801</u>	Aim: Assess effect of pharmacists interventions on BP and determine potential determinants of heterogeneity <u>Study type</u> : Meta- analysis <u>Size</u> : 39 RCTs were included with 14,224 pts	Inclusion criteria: RCT of pharmacist intervention delivered by a pharmacist alone or in collaboration with other healthcare professionals Exclusion criteria: Absence of above	Intervention: Pharmacist intervention delivered by a pharmacist alone or in collaboration with other healthcare professionals. Pharmacist interventions mainly included pt education, feedback to physician, and medication management. Comparator: Usual care	<u>1° endpoint</u> : Pharmacist interventions were associated with a large reduction in systolic and DBP of -7.6 mm Hg (95% CI: -9.0– -6.3 mm Hg) and -3.9 mm Hg (95% CI: -5– -2.8 mm Hg), respectively <u>Safety endpoint</u> : N/A	<u>Summary</u> : Pharmacist interventions, alone or in collaboration with other healthcare professionals, improved BP management

Shaw RJ, et al., 2014 (325) <u>25023250</u>	Aim: Determine whether nurse-managed protocols are effective for outpatient management of pts with DM, HTN, and hyperlipidemia (HTN RCT outcomes only included here) Study type: Meta- analysis Size: 12 RCTs, with 10,362 pts, of nurse- managed protocols for	Inclusion criteria: RCT of nurse- managed protocols for outpatient management of HTN Exclusion criteria: Absence of above	Intervention: Involvement of a registered nurse or a licensed practical nurse functioning beyond the usual scope of practice, such as adjusting medications and conducting interventions based on a written protocol. All studies used a nurse who titrated medications by following a protocol. Comparator: Usual care	 <u>1° endpoint</u>: SBP and DBP decreased by 3.68 mm Hg (95% CI: 1.05–6.31 mm Hg) and 1.56 mm Hg (95% CI: 0.36–2.76 mm Hg), respectively, with high variability (l²>70%) Nurse-managed protocols were more likely to achieve target BP than control protocols (OR: 1.41; 95% CI: 0.98–2.02), though difference was not significant and treatment effects were highly variable (Q 35.20; l²=74%). 	 Included studies of low/good quality as well as moderate/fair, and high quality Descriptions of interventions and protocols were limited <u>Summary</u>: Nurse-managed protocols for HTN care were associated with a mean decrease in SBP and DBP but not increase in HTN control.
Carter BL, et al.,	outpatient management of HTN <u>Aim</u> : Evaluate if a	Inclusion criteria:	Intervention:	Safety endpoint: N/A 1° endpoint: BP control at 9	2° endpoints:
2015 (326) 25805647	 <u>him</u>: Evaluate if a physician/pharmacist collaborative model would be implemented as determined by improved BP control and whether long-term BP control could be sustained <u>Study type</u>: Cluster RCT <u>Size</u>: 32 primary care offices from 15 states enrolled 625 pts with uncontrolled HTN; 54% from racial/ethnic minority groups and 50% with DM or CKD 	Offices were required to have an onsite clinical pharmacist must have practiced in the office. Pts were eligible if they were English or Spanish speaking, ≥18 y with uncontrolled BP as measured by the SC on the baseline visit. Exclusion criteria: Absence of above	Pharmacist conducted medical record review and a structured interview with the subject, including 1) a medication history; 2) an assessment of knowledge of BP medications, dosages and timing, and potential side effects; and 3) other barriers to BP control (e.g., side effects and nonadherence). The model recommended a telephone call at 2 wk, structured face- to-face visits at baseline, 1, 2, 4, 6, and 8 mo and additional visits if BP remained uncontrolled. The pharmacist created a care plan with recommendations for the physician to adjust	<u>a reinpoint</u> . BP control at 9 mo was 43% in intervention offices compared with 34% in control group (adjusted OR: 1.57 (95% CI: 0.99, 2.50), p=0.059). <u>Safety endpoint</u> : N/A	 The adjusted difference in mean SBP/DBP between the intervention and control groups for all pts at 9 mo was -6.1/-2.9 mm Hg (p=0.002 / p=0.005, respectively), and it was -6.4/-2.9 mm Hg (p=0.009 / p=0.044, respectively) in pts from racial or ethnic minorities. BP control and mean BP were significantly improved in pts from racial minorities in intervention offices at 18 and 24 mo (p=0.048 and p<0.001) compared with the control group. <u>Summary</u>: Although the results of the 1° outcome (BP control) were negative, the key 2° endpoint (mean BP) was significantly improved in the intervention group. Thus, the findings for 2° endpoints suggest that team-based care using clinical pharmacists significantly

therapy based on the JNC-	reduced BP in subjects from racial
7, and the BP goals were	minority groups.
<140/90 mm Hg for	
uncomplicated HTN or	
<130/80 mm Hg for pts with	
DM or CKD. The	
pharmacists did not follow	
algorithms or protocols other	
than JNC-7. Physicians	
were free to accept or to	
reject any recommendation	
or to modify the plan.	
Recommendations	
to pts focused on	
medication education,	
improving adherence, and	
strategies to implement	
lifestyle modifications.	
Comparator: Pharmacists	
in control offices were	
instructed to avoid	
intervention for study pts	
with HTN, but they could	
provide usual care curbside	
consultations if physicians	
specifically asked questions.	

Data Supplement 63. Electronic Health Records and Patient Registries (Section 12.3.1)

Study Acronym Author Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events Summary
Bardach NS, et al., 2013 (327) <u>24026600</u>	<u>Aim:</u> To assess the effect of P4P incentives on quality in EHR-enabled small practices in the	• Participating clinics (n=42 for each group) had similar baseline characteristics, with	• A city program provided all participating clinics with the same EHR software with decision support	• Intervention clinics had greater adjusted absolute improvement in rates of appropriate antithrombotic prescription (12.0% vs.	• Although the effect of the intervention was lower than the 10% improvement that we estimated a priori, the absolute risk reduction for BP control among pts with DM was 7.8% (NNT, 13). This

	context of an established QI initiative. <u>Study type and size:</u> A cluster-randomized trial of small (<10 clinicians) primary care clinics in New York City from April 2009 through March 2010.	a mean of 4,592 (median, 2,500) pts at the intervention group clinics and 3,042 (median, 2,000) at the control group clinics.	and pt registry functionalities and QI specialists offering technical assistance. • Incentivized clinics were paid for each pt whose care met the performance criteria, but they received higher payments for pts with comorbidities, who had Medicaid insurance, or who were uninsured (maximum payments: \$200/pt; \$100,000/clinic). Quality reports were given quarterly to both the intervention and control groups.	6.1%, difference: 6.0% (95% CI: 2.2%, 9.7%), p=0.001 for interaction term), BP control (no comorbidities: 9.7% vs. 4.3%, difference: 5.5% (95% CI: 1.6%, 9.3%), p=0.01 for interaction term; with DM: 9.0% vs. 1.2%, difference: 7.8% (95% CI: 3.2%, 12.4%), p=0.007 for interaction term; with DM or ischemic vascular disease: 9.5% vs. 1.7%, difference: 7.8% (95% CI: 3.0%, 12.6%), p=0.01 for interaction term), and in smoking cessation interventions (12.4% vs. 7.7%, difference: 4.7% (95% CI: -0.3%, 9.6%), p=0.02 for interaction term). Intervention clinics performed better on all measures for Medicaid and uninsured pts except cholesterol control, but no differences were statistically significant.	suggests that, for every 13 pts seeing incentivized clinicians, 1 more pt would achieve BP control. The 7.8% absolute change in BP control for pts with DM represents a 46% relative increase in BP control among intervention pts compared with the baseline of 16.8%. Further research is needed to determine whether this effect of the P4P intervention on BP control increases or decreases over time. However, this NNT to achieve BP control through incentives, taken together with the large relative increase in percentage of pts with BP control and the potential effect of BP control on risk of ischemic vascular events, suggests a reasonable opportunity to reduce morbidity and mortality through P4P as structured in this study. <u>Limitations:</u> Some clinics exited the program after randomization, with more control clinics leaving than intervention clinics. Additionally, this intervention occurred in the setting of a voluntary QI program. This may reflect a high level of intrinsic motivation to improve among practices in the study, as demonstrated by engagement with the QI specialists
Banerjee D, et al., 2012 (328) <u>22031453</u>	Study type: 3-y, cross-sectional sample using pt EHRs.	 251,590 pts ≥18 y. Underlying HTN was defined as 2 or more abnormal BP readings ≥140/90 mm Hg and/or pharmaceutical treatment. Appropriate HTN diagnosis was defined by the reporting of ICD-9 codes (401.0– 	• To identify prevalent and incident HTN cases in a large outpatient healthcare system, examine the diagnosis rates of prevalent and incident HTN, and identify clinical and demographic factors	• The prevalence of HTN was 28.7%, and the diagnosis rate was 62.9%. The incidence of HTN was 13.3%, with a diagnosis rate of 19.9%. Predictors of diagnosis for prevalent HTN included older age, Asian, African American, higher BMI, and increased number	• Outpatient EHR diagnosis rates are suboptimal, yet EHR diagnosis of HTN is strongly associated with treatment. Targeted efforts to improve diagnosis should be a priority.

		401.9). Factors associated with HTN diagnosis were assessed through multivariate analyses of pt clinical and demographic characteristics.	associated with appropriate HTN diagnosis.	of ABP readings. Predictors for incident HTN diagnosis were similar. In pts with 2 or more abnormal BP readings, HTN diagnosis was associated with significantly higher medication treatment rates (92.6% vs. 15.8%; p<0.0001).	
Jaffe MG, et al., 2013 (329) 23989679	Aim: Study the effect of a multipronged, system-based, OI approach on HTN control. Study type: Observational Size: All pts with HTN in the KPNC system were included	Inclusion criteria: 350,000 pts in the KPNC system with HTN in 2001, increasing to 650,000 in 2009 Eligibility: • ≥2 HTN diagnoses coded in primary care visits in the prior 2 y • ≥1 primary care HTN diagnoses and 1 or more hospitalizations with a 1° or 2° HTN diagnosis in the prior 2 y • ≥1 primary care HTN diagnoses and 1 or more filled prescriptions for HTN medication within the prior 6 mo, or • ≥1 primary care HTN diagnoses and 1 or more stroke-related hospitalizations or a history of coronary disease, HF, or DM	Intervention: KPNC HTN Program includes: HTN registry, HTN control monitoring and feedback system, evidence-based practice guidelines, medical assistant BP recheck program, and promotion of single polypill formulation (lisinopril- hydrochlorothiazide) Comparator: Insured pts in California from 2006–2009 who were included in the HEDIS commercial measurement by California health insurance plans participating in the NCQA quality measure reporting process. A 2° comparison group was included to obtain the reported national mean NCQA HEDIS commercial rates of	 <u>1° endpoint</u>: HTN control rates in KPNC pts with HTN improved from 43.6% (95% CI: 39.4%, 48.6%) in 2001 to 80.4% (95% CI: 75.6%, 84.4%) by the end of the study period (p<0.001 for trend). By comparison, national mean NCQA HEDIS commercial measurement HTN control increased from 55.4%–64.1%. California mean NCQA HEDIS commercial rates of HTN control were similar to those reported nationally from 2006–2009 (63.4%– 69.4%). <u>1° Safety endpoint</u>: N/A 	• A system-based approach to HTN control that includes performance measurement and QI strategies led to a significant improvement in HTN control (80%, compared to 44% baseline control) in a large population of pts in a managed care health plan.

Rakotz MK, et al., 2014 (330)	<u>Aim</u> : The goal of this study was to develop	• Of the 139,666 active adult primary care pts in	HTN control from 2001–2009 from health plans that participated in the NCQA HEDIS quality measure reporting process. • In phase 1, we reviewed EHRs using	• Of the 1,033 at-risk pts who remained active during	• Although we used multiple algorithms to identify pts with elevated BP
25024244	a technology-based strategy to identify pts with undiagnosed HTN in 23 primary care practices and integrate this innovation into a continuous QI initiative in a large, integrated health system.	these 23 practices, 47,822 already had a diagnosis of HTN, white- coat HTN, pre-HTN, or elevated BP. The 3 screening algorithms for undiagnosed HTN were applied to the remaining pts' EHRs. There were 1,586 pts who met the criteria of 1 or more of the algorithms and were therefore considered at risk for undiagnosed HTN.	algorithms designed to identify pts at risk for undiagnosed HTN. We then invited each at-risk pt to complete an automated office BP protocol. In phase 2, we instituted a QI process that included regular physician feedback and office- based computer alerts to evaluate at-risk pts not screened in phase 1. Study pts were observed for 24 additional mo to determine rates of diagnostic resolution. After phase 1, we established a continuous QI initiative to further evaluate pts who remained at risk for undiagnosed HTN. In this 24-mo follow-up phase (phase 2), all primary care physicians received monthly lists of their pts who continued to be at risk for undiagnosed HTN.	phase 2, 740 (72%) were classified by the end of the follow-up period: 361 had HTN diagnosed, 290 had either white coat HTN, pre- HTN, or elevated BP diagnosed, and 89 had normal BP. By the end of the follow-up period, 293 pts (28%) had not been classified and remained at risk for undiagnosed HTN.	readings, it is unlikely that we identified all pts with undiagnosed HTN.

Borden WB, et al.,	Aim: The purpose of	Using the National CV	These pts were contacted by staff via telephone or letter to arrange a follow-up appointment. These pts remained on the physicians' lists until an automated office BP evaluation was completed or an ICD-9 diagnosis was entered into the chart that indicated the pt's at-risk status had been resolved. In addition, when an at-risk pt arrived for an office visit for any reason, a best practice advisory was prominently displayed on that pt's EHR screen to notify the medical assistant and physician that an automated office BP measurement was needed. N/A	• Of 1,185,253 pts in the	Among U.S. ambulatory cardiology
Borden WB, et al., 2014 (331) <u>25447261</u>	<u>Aim</u> : The purpose of this study was to examine the effect of the 2014 expert panel BP management recommendations on pts managed in U.S. ambulatory CV practices.	• Using the National CV Data Registry PINNACLE Registry, we assessed the proportion of 1,185,253 pts who met the 2003 and 2014 panel recommendations, highlighting the populations of pts for whom the BP goals changed.	N/A	• Of 1,185,253 pts in the study cohort, 706,859 (59.6%) achieved the 2003 JNC-7 goals. Using the 2014 recommendations, 880,378 (74.3%) pts were at goal. Among the 173,519 (14.6%) for whom goal achievement changed, 40,323 (23.2%) had a prior stroke or TIA, and 112,174 (64.6%) had CAD. In addition, the average Framingham risk score in	• Among U.S. ambulatory cardiology pts with HTN, nearly 1 in 7 who did not meet JNC-7 recommendations would now meet the 2014 treatment goals.

	а	this group was $8.5 \pm 3.2\%$, and the 10-y atherosclerotic	
	C	CVD risk score was 28.0 \pm	
	1	19.5%.	

Data Supplement 64. RCTs, Meta-analyses, and Systematic Reviews on the Effect of Telehealth Interventions to Improve Hypertension Control (Section 12.3.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events Summary
Burke LE, et al., 2015 (332) <u>26271892</u>	<u>Aim</u> : Review of the Scientific Literature on mHealth Tools Related to CVD Prevention <u>Study type</u> : Systematic review <u>Size</u> : 69 studies of the use of mobile technologies to reduce CVD risk behaviors	Inclusion criteria Studies of electronic and mobile technology tools in CV prevention; published from 2004–2014 in English language; enrolling adults except for smoking cessation, for which adolescents were also included; conducted in the U.S. and in developed countries. Exclusion criteria: Absence of above.	Intervention: Mobile technologies to reduce CVD risk behaviors- varied across studies <u>Comparator</u> : Varied across studies.	<u>1º endpoint</u> : Varied across studies. <u>1º Safety endpoint</u> : N/A	Summary: mHealth or mobile technologies have the potential to transform the delivery of health- related messages and ongoing interventions targeting behavior change. Moreover, the use of monitoring devices (e.g., Bluetooth- enabled BP monitors and blood glucose monitors) permits the sharing of important pt self-management parameters with healthcare providers in real time and the delivery of feedback and guidance to pts when they need it. Furthermore, using mHealth tools for monitoring provides the clinician data that far exceed what can be measured in the brief clinical encounter and reflect the status of physiological or behavioral measures in the person's natural setting.
Liu S, et al., 2013 (333) <u>23618507</u>	<u>Aim</u> : Assess the efficacy of e- counselling in reducing BP	Inclusion criteria: 1) Trials that investigated the effect of Internet-based lifestyle interventions on SBP and DBP, 2) trials that included	Intervention: Internet- based intervention as preventive e-counselling or advice using Web sites or e-mails to modify exercise or diet as a	<u>1° endpoint</u> : MD in BP reduction (Internet- based – usual care): SBP: -3.8 mm Hg (95% CI: - 5.63– -2.06), I ² =61	• Behavior change techniques that were used in more than 50% of the successful internet-based interventions included the following: providing information on consequences of behavior in general

			<u> </u>		
	Study type:	supplemental components	means of improving BP	DBP: -2.1 mm Hg (95% CI: -	(86%), incorporating feedback on
	Systematic review,	such as mobile text	control. These Internet-	3.51– -0.65), l ² =57	performance (86%), prompting self-
	meta-analysis	messages, telephone, or	based interventions were		monitoring of behaviors (71%), and
		in-person support, 3)	primarily self-guided,	Influence of intervention	giving instructions on how to perform
	Size: 13 RCTs or case-	intervention duration of at	and access was gained	attributes:	the targeted behavior change (71%).
	control studies	least 8 wk, and 4) SBP	via desktop computer,	Intervention duration:	
		and DBP reported as 1° or	laptop, tablet, or smart	Long-term (≥6 mo)	Summary: Internet-based
		2° outcome, measured at	phone. The duration of	intervention: SBP -5.8 mm Hg	interventions reduced SBP and DBP
		a clinic or office.	each intervention had to	(95% CI: -4.3– -4.1)	significantly compared to usual care.
			be at least 8 wk in order	Short-term (<6 mo)	Internet-based interventions had
		Exclusion criteria:	to achieve clinically	intervention: SBP -3.47 mm	greater effect on BP lowering if they
		Absence of above.	meaningful outcomes,	Hg (95% CI: -5.2– -1.7)	were 1) long-term (≥ 6 mo) in
			including the pt's ability	DBP mean reduction: results	duration, and 2) used >5 behavior
			to learn and adhere to	not reported, not statistically	change techniques.
			complex new behaviors,	significant.	
			and to allow for sufficient	# of behavior change	
			time to demonstrate a	techniques:	
			stable reduction in BP.	≥5 behavior change	
			The majority (9/13) of	techniques: SBP -5.92 mm	
			interventions had	Hg (95% CI: -7.43– -4.42) /	
			supplemental	DBP -2.45 mm Hg (95% CI: -	
			components that were	3.50– -1.41)	
			not internet-based, such	<5 behavior change	
			as text messages, in-	techniques: SBP -2.69 mm	
			person visits, and live	Hq (95% CI: -4.61– -0.78) /	
			support and 10/13	DBP -0.02 mm Hg (95% CI: -	
				1.20–1.17)	
			targeted both exercise and diet behaviors.	1.20-1.17)	
				10 Cofety and sint N/A	
			Componenter, House core	1° Safety endpoint: N/A	
			Comparator: Usual care		
			with no internet-based		
On hand Carlad	Also Declaradata (laster ten seltente	strategy.		L factoria da com
Omboni S, et al.,	Aim: Review data from	Inclusion criteria:	Intervention: HBPT had	1° endpoint: Compared to	Limitations:
2013 (334)	RCTs on the	English language	to be based on the use	usual care, HBPT improved:	HBPT intervention features
<u>23299557</u>	effectiveness of HBPT	 Published up to Feb. 	of an electronic	Office SBP by 4.71 mm Hg	(telemonitoring systems and self-
	vs. usual care with	2012	automated BP monitor	(95% CI: 6.18–3.24;	monitoring programs) as well as
	respect to improvement	 RCT testing HBPT vs. 	storing values obtained	p<0.001); I ² =52.2%; p=0.003	inclusion criteria and demographic
	of BP control,	usual care.	at the pt's home and	 Office DBP by 2.45 mm Hg 	and clinical characteristics of the
	healthcare resources		transferring them to a	(95% CI: 3.33–1.57;	comparative groups varied across
	utilization and costs,		remote computer	p<0.001); I ² =40.4%; p=0.048	

ana <u>Siz</u> witi not on	udy type: Meta- halysis ze: 23 unique RCTs th 7037 pts (though of all studies reported all outcomes of terest)	(wired or wireless), a modem or an Internet connection. At least 1 self BP measurement had to be available for each pt in the intervention group. Comparator: Usual care	mm Hg nondiabetic pts and <130/80 mm Hg diabetic pts): RR: 1.16 (95% CI: 1.04–1.29; $p<0.001$); $I^2=69\%$; $p<0.001$ 2° endpoint : Compared to usual care, HBPT improved: • Greater prescription of antihypertensive medications: weighted MD 0.40 (95% CI: 0.17–0.62; $p<0.001$); $I^2=84.2\%$; $p<0.001$ • Lower number of office visits: weighted MD -0.18 (95% CI: -0.37–0.00); $I^2=32.7\%$; $p=0.146$ • Quality of life physical component of SF-12 or SF-36 questionnaire: weighted MD 2.78 (95% CI: 1.15–4.41); $I^2=0.0\%$; $p=0.853$ • There was no difference between HBPT and usual care in: • Therapeutic adherence [92% HBPT vs. 90% usual care; between-group difference +1.30% (95% CI: - 2.31–4.90; $p=0.481$), $I^2=0.00\%$; $p=0.888$) • Quality of life mental component of SF-12 or SF-36 questionnaire: weighted MD -	 Most studies were powered to test differences in BP lowering, not 2° outcomes <u>Summary</u>: HBPT yielded greater SBP and DBP reductions and a larger proportion of pts achieving BP control than usual care. HBPT vs. usual care resulted in greater prescription of antihypertensive medications and fewer office visits but no difference in therapeutic adherence. Healthcare costs were higher with HBPT than usual care, but when HBPT-related costs were excluded, medical costs were similar between groups. Use of HBPT vs. usual care improved quality of life physical component but not mental. Authors note that the amount of office BP reduction attributable to HBPT was in line with that observed in RCTs of antihypertensive drugs compared with placebo. The estimate was also larger than that usually related to HBP self-monitoring, which speaks in favor of a possible added value of the teletransmission approach.
			I ² =0.00%; p=0.888) • Quality of life mental	approacn.

				HBPT group vs. usual care: weighted MD 662.92 (95% CI: 540.81–785.04) euros per pt; I ² =99.6%; p<0.001, but costs were similar when only medical costs (excluding HBPT-related costs) were considered (-12.4; 95% CI: - 930.52–906.23) euros; p=0.767. <u>Safety endpoint</u> : No difference was observed in the risk of adverse events (RR: 1.22; 95% CI: 0.86– 1.71; p=0.111)	
Verberk W, et al., 2011 (335) 21527847	Aim: Examine the usefulness of telecare for HTN management Study type: Meta- analysis Size: 9 RCTs with 2,501 pts	Inclusion criteria: 1) Published in the English language, 2) pts were diagnosed as hypertensive and performed BP self- measurement at home, 3) RCTs that compared telecare of BP with usual care, 4) data were transmitted to healthcare providers by telephone, modem, Internet, or mail, and 5) either change in BP or the number of pts that reached their target BP was an outcome and was provided in the study. Date restrictions not reported. Exclusion criteria: Absence of above	Intervention: Telecare for HTN management (treatment and/or coaching). Telecare involved a data transmission process to collect data on a pt's health status to allow remote HTN management. Procedures varied in length and frequency of contact and method of delivery (i.e., often telephone or cell phone with or without internet/computer; with or without behavioral counseling by nurse or pharmacist), often as an adjunct to "usual care" clinical visits.	<u>1° endpoint</u> : Difference in BP Reduction (Telecare-Usual care): • SBP 5.2 ± 1.5 mm Hg (95% Cl: 2.31–8.07) • DBP 2.1 ± 0.8 mm Hg (95% Cl: 0.52–3.69) <u>Safety endpoint</u> : N/A	Limitations: Telecare intervention methods varied greatly across studies Summary: Telecare led to a greater decrease in SBP and DBP compared with usual care. Telecare seems a valuable tool to support HTN management.

Agarwal R, et al., 2011 (27) 21115879	Aim: Quantify both the magnitude and mechanisms of benefit (including effect on therapeutic inertia) of home BP monitoring on BP reduction. Therapeutic inertia was defined as no change in medications combined with uncontrolled BP. Study type: Systematic review and meta-analysis Size: 37 RCTs with 9,446 pts. Trial settings included community (n=5), dialysis unit (n=2), general practices (n=18), hospitals and general practice (n=1), and hospital-based outpatient units (n=11).	Inclusion criteria: Studies that randomized pts to control or home BP monitoring group Exclusion criteria: Absence of above	Intervention: Home BP monitoring as an adjunct to usual care for HTN Comparator: Usual care with BP monitoring in clinic	<u>1° endpoint</u> : Compared with usual care alone, home- based BP monitoring: •Reduced SBP: -2.63 mm Hg (95% CI: -4.24 – -1.02) and • Reduced DBP: -1.68 mm Hg (95% CI: -2.58– -0.79) • Greater reduction in SBP by HBPM interventions was seen with added telemonitoring (effect size -3.20; 95% CI: - 4.66– -1.73) vs. home BP monitoring (effect size -1.26; 95% CI: -2.20– -0.31; p=0.029). This finding is relevant to telemonitoring	 <u>2° endpoints:</u> More frequent reductions in antihypertensive medication (presumably due to identification of white coat HTN): RR: 2.02 (95% CI: 1.32–3.11) Lowered therapeutic inertia (i.e., unchanged medication despite elevated BP: RR for unchanged medication 0.82 (95% CI: 0.68–0.99) <u>Limitations</u>: Different inclusion and exclusion criteria, different BP measurement techniques, drug titration protocols, pt populations, and duration of follow-up across studies likely introduced significant heterogeneity in effect size. <u>Summary</u>: Home BP monitoring leads to a small but significant reduction in SBP and DBP. Greater reduction in SBP is seen when HBPM is accompanied by specific programs to titrate antihypertensive drugs. 1 such strategy is telemonitoring, in which BP readings obtained at home are relayed to the provider who can then take appropriate action, thus reducing therapeutic inertia.
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Data Supplement 65. RCTs and Observational Studies that Report on the Effect of Performance Measures and on Hypertension Control (Section 12.4.1)

Study Acronym;	Aim of Study;	Patient Population	Study Intervention (# patients)	Endpoint Results	Relevant 2° Endpoint (if any);
Author;	Study Type;		1	(Absolute Event Rates, P	Study Limitations;
Year Published	Study Size (N)		Study Comparator (# patients)	value; OR or RR; & 95%	Adverse Events
				CI)	

Svetkey LP, et al., 2009 (336) <u>19920081</u>	Aim: Study the effect of physician intervention and/or pt intervention vs. usual care, to assess the impact of education, monitoring, and feedback protocol to help improve HTN control Study type: Nested 2×2 RCT Size: 8 primary care practices, 32 physicians, 574 pts	Inclusion criteria: Practices: matched pairs (intervention vs. usual care) by specialty (internal medicine vs. family physician) and by pt socioeconomic mix. All physicians were invited to participate. <u>Pt eligibility:</u> ≥25 y, hypertensive by billing code. <u>Pt exclusion:</u> Self- reported CKD, CVD event within past 6 mo, pregnant, breastfeeding, or planning a pregnancy.	Physician Intervention: 18 mo of online training, self- monitoring, quarterly feedback reports.Pt Intervention: 20 weekly group sessions for 6 mo, followed by 12 monthly telephone counseling contacts, focused on weight loss, DASH dietary patter, exercise, and reduce sodium intake.Comparator:Usual care	<u>1° endpoint</u> : Pt intervention + physician intervention group had greatest BP lowering at 6 mo (-9.7 mm Hg ± 12.7), but at 18 mo there was no significant difference between groups. <u>1° Safety endpoint</u> : N/A	• This trial suggests that pt level monitoring and feedback, in combination with physician level monitoring and feedback, provides additional 6 mo BP control above and beyond usual care. The impact of the intervention diminished after the weekly pt group sessions ended and monthly telephone calls began instead.
Jaffe MG, et al., 2013 (329) <u>23989679</u>	Aim: Study the effect of a multipronged, system- based, QI approach on HTN control. Study type: Observational Size: All pts with HTN in the KPNC system were included	Inclusion criteria: 350,000 pts in the KPNC system with HTN in 2001, increasing to 650,000 in 2009 Eligibility: • ≥2 HTN diagnoses coded in primary care visits in the prior 2 y • ≥1 primary care HTN diagnoses and 1 or more hospitalizations with a 1° or 2° HTN diagnosis in the prior 2 y • ≥1 primary care HTN diagnoses and 1 or more filled prescriptions for HTN medication within the prior 6 mo, or	Intervention: KPNC HTN Program includes: HTN registry, HTN control monitoring and feedback system, evidence- based practice guidelines, medical assistant BP recheck program, and promotion of single polypill formulation (lisinopril-hydrochlorothiazide) Comparator: Insured pts in California from 2006–2009 who were included in the HEDIS commercial measurement by California health insurance plans participating in the NCQA quality measure reporting process. A 2° comparison group was included to obtain the reported national mean NCQA HEDIS commercial rates of HTN control from 2001–	 <u>1° endpoint</u>: HTN control rates in KPNC pts with HTN improved from 43.6% (95% CI: 39.4%–48.6%) in 2001 to 80.4% (95% CI: 75.6%–84.4%) by the end of the study period (p<0.001 for trend). By comparison, national mean NCQA HEDIS commercial measurement HTN control increased from 55.4%–64.1%. California mean NCQA HEDIS commercial rates of HTN control were similar to those reported nationally from 2006–2009 (63.4%–69.4%). 	• A system-based approach to HTN control that includes performance measurement and QI strategies led to a significant improvement in HTN control (80%, compared to 44% baseline control) in a large population of pts in a managed care health plan.

		• ≥1 primary care HTN diagnoses and 1 or more stroke-related hospitalizations or a history of coronary disease, HF, or DM	2009 from health plans that participated in the NCQA HEDIS quality measure reporting process.	<u>1° Safety endpoint</u> : N/A	
al., 2013 (337) 23536132	Aim: Study the effect of an audit-based education intervention to guidelines/prompts, vs. usual care, to help improve BP control in pts with CKD Study type: Cluster RCT Size: 93 general practices (30 audit-based education intervention, 32 Guidelines/prompts, and 31 usual care)	Inclusion criteria: All pts with CKD in the participating practices	Intervention: Audit-based education vs. guidelines/prompts Comparator: Usual care	<u>1° endpoint</u> : SBP was significantly lower in the audit-based education group (-2.41 mm Hg; 95% CI: 0.59–4.29). There was no significant change in BP in the other 2 groups. <u>1° Safety endpoint</u> : No reports of harm.	• This trial suggests that an intervention that includes specific performance and feedback reports improves BP control in pts with CKD, compared to usual care. To the contrary, the use of practice guidelines and prompts did not improve BP control compared to usual care.

Data Supplement 66. RCTs, Meta-analyses, and Systematic Reviews on Quality Improvement Strategies on Hypertension Treatment Outcomes (Section 12.4.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events Summary
Walsh JM, et al., 2006 (338) <u>16799359</u>	<u>Aim</u> : Assess the effectiveness of QI strategies in lowering BP <u>Study type</u> : Systematic review	Inclusion criteria: Trials, controlled before–after studies, and interrupted time series evaluating QI interventions targeting HTN control and reporting BP outcomes.	Intervention: QI interventions targeting some component of provider behavior or organizational change to improve HTN control <u>Comparator</u> : Contemporaneous	 The majority of articles described interventions consisting of more than 1 strategy with the median number of QI strategies per comparison =3. Results are organized below by type of QI strategy. Variety of strategies used 	Limitations: Studies varied by design, population, sample size, setting, and methodological quality. Definition of each QI strategy varied across studies. Few studies assessed a single QI strategy; because most studies included more than 1 QI strategy, it could not be discerned which individual QI strategies had the

Size: 44 al	ticlos Evolucion oritorio.	observation of cohorts	SBP/DBP, median reduction:	greatest effects or whether certain
reporting 5		differing primarily with	4.5 mm Hg (IQR: 1.5–11.0)/	combinations of individual QI
compariso		respect to exposure to	2.1 mm Hg (IQR: -0.2–5.0)	strategies were more "potent" than
	subpopulations (e.g., HTN	the QI intervention	SBP/DBP control: 16% (IQR:	others.
	in pts with alcoholism)		10.3–32.2)/ 6% (IQR: 1.5–	
			17.5)	Summary: QI strategies are
			Provider reminders	associated with improved HTN
			SBP/DBP, median reduction:	control. QI strategies improved SBP
			1.2 mm Hg (IQR: 1.0–1.9)/ 0.3	and the proportion of pts achieving
			mm Hg (IQR: -0.2–1.7)	SBP control and had a more modest
			DBP control: 5% (IQR: 2.0–	effect on DBP and the proportion of
				pts achieving DBP control. Team
			7.0)	change (i.e., a focus on HTN by
			 Facilitated relay of clinical 	
			data	someone in addition to the pt's
			SBP/DBP, median reduction:	physician) had the largest effect on
			8.0 mm Hg (IQR: 2.5–12.3)/	both SBP and DBP. All of the
			1.8 mm Hg (IQR: -0.1–4.5)	strategies assessed may be beneficial
			SBP/DBP control: 25% (IQR:	in terms of clinically meaningful
			17.0-34.2)/ 2% (IQR: 1.6-5.0)	reductions in BP under some
			Audit and feedback	circumstances and in varying
			SBP/DBP, median reduction:	combinations.
			1.5 mm Hg (IQR: 1.2–1.7)/ 0.6	
			mm Hg (IQR: 0.4–1.0)	
			SBP/DBP control: -3.5% (IQR:	
			-5.7–1.4)/ 2.0% (IQR: 1.7–4.3)	
			 Provider education 	
			SBP/DBP, median reduction:	
			3.3 mm Hg (IQR: 1.2–5.4)/ 0.6	
			mm Hg (IQR: -0.7v3.4)	
			SBP/DBP control: 11% (IQR:	
			1.4–13.1)/ 4% (IQR: 1.7–11.3)	
			Pt education	
			SBP/DBP, median reduction:	
			8.1 mm Hg (IQR: 3.3–11.8)/	
			3.8 mm Hg (IQR: 0.6–6.7)	
			SBP/DBP control: 19% (IQR:	
			11.4–33.2)/ 17% (IQR: 11.4–	
			24.5)	
			 Promotion of self- 	
			management	

				SBP/DBP, median reduction: 3.3 mm Hg (IQR: 2.6–10.1)/ 2.8 mm Hg (IQR: 0.4–6.7) SBP/DBP control: 13%/ 9% (IQR: 5.3–11.4) • <u>Pt reminders</u> SBP/DBP, median reduction: 3.3 mm Hg (IQR: 2.3–4.5)/ 0.4 mm Hg (IQR: -2.4–5.0) DBP control: 2% (IQR: 1.1– 9.4) • <u>Team change</u> SBP/DBP, median reduction: 9.7 mm Hg (IQR: 4.2–14.0) (p<0.05)/ 4.2 mm Hg (IQR: 0.2–6.8) (p<0.05) SBP/DBP control: 22% (IQR: 9.0–33.8)/ 17% (IQR: 5.7– 24.5) • <u>Financial incentives</u> SBP/DBP, median reduction: - 13.3 mm Hg/ 0.0 mm Hg (IQR: -2.0–2.5) DBP control: 4% (IQR: -1.1– 9.4) Safety endpoint: N/A	
Carter BL, et al., 2009 (321) <u>19858431</u>	<u>Aim</u> : Determine potency of interventions for BP involving nurses and pharmacists <u>Study type</u> : Meta- analysis <u>Size</u> : 37 RCTs of team-based HTN care involving nurse or	Inclusion criteria: RCT of team-based HTN care involving nurse or pharmacist intervention Exclusion criteria: Absence of above	Intervention: Team- based HTN care involving nurse or pharmacist intervention In nearly all studies involving nurses or pharmacists in clinics, consistent and dedicated case management activities were provided that were distinct from traditional nursing or	 <u>1° endpoint:</u> OR (95% CI) for controlled BP were: nurses: 1.69 (1.48, 1.93); pharmacists within primary care clinics: 2.17 (1.75, 2.68); and community pharmacists: 2.89 (1.83, 4.55). Mean (SD) reductions in SBP were: nurse intervention: 5.84 (8.05) mm Hg; pharmacists in clinics: 7.76(7.81) mm Hg; and 	• Stepwise regression was used to compare studies that included a given intervention strategy with studies that did not. Several individual components of the interventions were associated with significant reductions in mean SBP including pharmacist recommended medication to physician (-27.21 mm Hg; p=0.002), counseling about lifestyle modification (-12.63 mm Hg; p=0.03), pharmacist performed the intervention (-11.70 mm Hg; p=0.03), use of a treatment

	pharmacist intervention		pharmacist duties. However, pharmacists in community pharmacies usually had to incorporate the intervention with traditional medication dispensing functions. <u>Comparator</u> : Usual care	community pharmacists: 9.31 (5.00) mm Hg. • There were no significant differences between nurse and pharmacist effects (p≥0.19). <u>Safety endpoint</u> : N/A	algorithm (-8.46 mm Hg; p<0.001), completion of a drug profile and/or medication history (-8.28 mm Hg; p=0.001), and the overall intervention potency score assigned by the study reviewers (p<0.001). The factors associated with a reduction in DBP were: referral was made to a specialist (-19.61 mm Hg; p=0.04), providing pt education about BP medications (- 17.60 mm Hg; p=0.003), completion of a drug profile and/or medication history (-7.27 mm Hg; p=0.006), pharmacist performed the intervention (-4.03 mm Hg; p=0.04), or nurse performed the intervention (-3.94 mm Hg; p=0.04). <u>Summary</u> : Interventions involving pharmacists or nurses were associated with significantly improved BP control.
Agarwal R, et al., 2011 (27) <u>21115879</u>	Aim: Quantify both the magnitude and mechanisms of benefit (including effect on therapeutic inertia) of home BP monitoring on BP reduction. Therapeutic inertia was defined as no change in medications combined with uncontrolled BP. Systematic Review and Meta-analysis	Inclusion criteria: Studies that randomized pts to control or home BP monitoring group Exclusion criteria: Absence of above	Intervention: Home BP monitoring as an adjunct to usual care for HTN <u>Comparator</u> : Usual care with BP monitoring in clinic	<u>1° endpoint</u> : Compared with usual care alone, home-based BP monitoring: • Reduced SBP: -2.63 mm Hg (95% CI: -4.24 – -1.02) and • Reduced DBP: -1.68 mm Hg (95% CI: -2.58 – -0.79) • Greater reduction in SBP by home BP monitoring interventions was seen with added telemonitoring effect size: -3.20 (95% CI: -4.66 – 1.73) vs. home BP monitoring effect size: -1.26; 95% CI: - 2.20 – -0.31; p=0.029. <u>Safety endpoint</u> : N/A	 <u>2° endpoints</u>: More frequent reductions in antihypertensive medication (presumably due to identification of white coat HTN): RR: 2.02; 95% CI: 1.32–3.11 Lowered therapeutic inertia (i.e., unchanged medication despite elevated BP: RR for unchanged medication 0.82 (95% CI: 0.68–0.99) <u>Limitations</u>: Different inclusion and exclusion criteria, different BP measurement techniques, drug titration protocols, pt populations, and duration of follow-up across studies likely introduced significant heterogeneity in effect size.

Anchala R, et al., 2012 (339) 23071713	Size: 37 RCTs with 9446 pts. Trial settings included community (n=5), dialysis unit (n=2), general practices (n=18), hospitals and general practice (n=1), and hospital-based outpatient units (n=11). <u>Aim</u> : Evaluate the role of decision support systems in prevention of CVD among pts <u>Study type:</u> Systematic review and meta-analysis <u>Size</u> : 10 studies with 5 studies reporting effect on BP (BP results only reported here)	Inclusion criteria: 1) Cross-sectional, case control, cohort, and RCTs, 2) Studies conducted among adult pts ≥18, 3) studies on prevention of CV disorders (MI, stroke, CHD, peripheral vascular disorders and HF) and management of HTN, 4) studies on interventions including: decision support systems, clinical decision supports systems, computerized decision support systems, clinical decision making tools and medical decision making Exclusion criteria: Absence of above	Intervention: Decision support systems, clinical decision supports systems, computerized decision support systems, clinical decision making tools and medical decision making in the management of HTN <u>Comparator</u> : Usual care	1° endpoint: • Reduction in SBP (5 studies): 2.32 mm Hg (95% CI: -3.960.69) • Reduction in DBP (2 studies): 0.42 mm Hg (95% CI: -2.30-1.47) Safety endpoint: N/A	Summary: • Home BP monitoring leads to small but significant reduction in SBP and DBP. Greater reduction in SBP is seen accompanied by specific programs to titrate antihypertensive drugs. One such strategy is telemonitoring, in which BP readings obtained at home are relayed to the provider who can then take appropriate action. Limitations: • Small number of studies of varied quality. • Interventions varied across studies. Summary: Clinical decision support resulted in modest reduction of SBP and no significant reduction of DBP.
Proia KK, et al., 2014 (323) <u>24933494</u>	<u>Aim</u> : Examine current evidence on the effectiveness of team- based care in improving BP outcomes (update of	Inclusion criteria: Study of team-based care; conducted in a high- income economy; reported at least 1 BP outcome of interest; included a comparison group or had	Intervention: Team- based care was defined as adding new staff or changing the roles of existing staff to work with a PCP for HTN care. Team members who	 <u>1° endpoint</u>: Proportion with controlled BP: Absolute percentage point (pct pt) change in pts with controlled BP from 33 studies comparing team-based care to usual care: median effect 	<u>2° endpoints</u> : Compared with pts in usual care, the proportion of pts receiving team-based care with "high" medication adherence (defined as taking medications as prescribed >80% of the time) increased by a median of 16.3 pct pts (9 studies).

		improving when pha of the tea	rmacists and nurses are part
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Data Supplement 67. Nonrandomized Trials, Observational Studies, and/or Registries of Effect of Quality Improvement Strategies on Hypertension Treatment Outcomes (Section 12.4.2)

Study Acronym; Author; Year Published	Study Type/Design; Study Size (N)	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Thomas KL, et al., 2014 (340) <u>25351480</u>	Study type: Community- based HTN QI program [multifaceted BP control program using a web-based health portal (Heart360), community health coaches, and PA guidance] to improve HTN control in a diverse community setting Design: Pre-post study without a concurrent control Size: 1756 pts with HTN from 8 clinics: • Median age, 60 y • Female, 65.6% • African American, 76.1%	Inclusion criteria: Individuals from pt sites >18 y with a previous billing diagnosis of HTN (ICD-9 code 401.X) or a previous clinical diagnosis of HTN in the medical record. Exclusion criteria: Did not reside in Durham County or had a neurocognitive disorder that prevented enrollment	<u>1° endpoint</u> : 1) Difference in SBP and DBP from enrollment (BP obtained in the clinic at enrollment) to the last BP as measured in clinic within 6 mo after enrollment, 2) proportion of pts that achieved BP <140/90 mm Hg by last clinic visit within 6 mo, and 3) proportion of pts with BP <140/90 mm Hg or drop in SBP ≥10 mm Hg by last visit relative to their enrollment BP. <u>Results:</u> • Mean change in BP: -4.7 mm Hg (SD ± 21.4) / -2.8 mm Hg (SD ± 11.8) after 6 mo • BP control (<140/90 mm Hg) rate: Increased from 51% at baseline to 63% at 6 mo • Proportion with BP<140/90 or ≥10 mm Hg decrease in SBP at 6 mo was 69% • Among those who were in tiers 1 (BP=140/90– 159/99 mm Hg) and 2 (BP≥159/99 mm Hg) at enrollment (n=889), BP change was -8.8 mm Hg (SD ± 15.8) / -5.0 mm Hg (SD ± 10.0) and -23.7 mm Hg (SD ± 26.5) / -10.1 mm Hg (SD ± 14.1), respectively.	<u>Summary:</u> A multicomponent- tiered HTN program that included team-based care with PAs and community health coaches was associated with improved BP control in a diverse community- based population. Though the web-based approach presented technical challenges for some pts, there was a direct association between higher use of Heart360 and larger recorded BP declines as entered into Heart360. This provides some indirect evidence that those pts who were more engaged with their BP self-monitoring achieved better BP control.
Jaffe MG, et al., 2013 (329) <u>23989679</u>	Study type: Quasi- experimental evaluation of multi-faceted QI program that included 1) Health system- wide HTN registry, 2) HTN control rates (with provider audit and feedback), 3)	Inclusion criteria: Pts identified with HTN within an integrated health care delivery system (KPNC) from 2001–2009	<u>1° endpoint</u> : BP control using NCQA HEDIS measures <u>Results:</u> BP control increased from 44%–80% from 2001–2009 with the KPNC QI program compared to 55.4% to 64.1% for the national mean and 63.4% to	Summary: Implementation of a large-scale HTN program was associated with a significant increase in HTN control compared with state and national control rates.

	Exclusion criteria: None stated	69.4% for the Ca mean from 2006 to 2009 NCQA HEDIS commercial measurement comparison groups.	
Size: Kaiser HTN registry increased from 349,937 pts in 2001 to 652,763 in 2009.			

Data Supplement 68. RCTs Comparing Financial Incentives (Section 12.5)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events Summary
Peterson LA, et al., 2013 (341) <u>24026599</u> Hysong, SJ, et al., 2012 (342) <u>23145846</u>	Aim: To test the effect of explicit financial incentives to reward guideline recommended HTN care. Study type: Cluster randomized trial of 12 VA Outpatient clinics with 5 performance periods and a 12-mo washout Size: 83 PCPs and 42 nonphysician	• Study population was providers, not pts: a minimum of 5 fulltime PCPs from 12 hospital- based primary care clinics in 5 A Networks. Then, the clinics were randomized to 1 of 4 study groups, 1) physician level (individual) incentives, 2) practice- level incentives, 3) physician-level plus practice-level (combined) incentives, and 4) no incentives (control).	Interventions: Education, Financial Incentives, Audit and Feedback; Intervention group pts received up to 5 incentive payments in their paychecks ~every 4 mo and were notified each time a payment was posted. Comparator: 4 different groups,1 paid incentives at the practice level,1 paid incentives at the physician level, 1 paid	<u>1° endpoint</u> : In unadjusted analyses, the percentage of pts either with controlled HTN or receiving an appropriate response increased for each incentive group between baseline and final performance period, 75% to 84% in the individual group, 80% to 85% in the practice group, and 79% to88% in the combined group. Performance did not change in control group, 86%. The adjusted estimated ab-solute change over the study of the pts meeting the combined BP or	Summary: • Mean (SD) total payments over the study were \$4,270 (\$459), \$2672 (\$153), and \$1,648 (\$248) for the combined, individual, and practice- level interventions, respectively. Change in BP control or appropriate response to uncontrolled BP compared with the control group was significantly greater only in the individual incentives group. Change in guideline-recommended medication use was not significant compared with the control group. The effect of the incentive was not sustained after a washout.

	personnel (e.g., nurses, pharmacists). <u>Main Outcomes and</u> <u>Measures:</u> Among a random sample, number of pts achieving guideline- recommended BP thresholds or receiving an appropriate response to uncontrolled BP, number of pts prescribed guideline- recommended medications, and number who developed hypotension.		for both levels and the 4 th paid no incentives. (19–20 physicians in each group)	appropriate response measure was 8.84% (95% CI: 4.20%– 11.80%) for the individual group, 3.70% (95% CI: 0.24%, 7.68%) for the practice group, 5.54% (95% CI: 1.92%–9.52%) for the combined group, and 0.47% (95% CI: -3.12%–4.04%) for the control group. The adjusted estimated absolute difference over the study in the change between the proportion of the physician's pts achieving BP control or receiving an appropriate response for the individual incentive group and the controls was 8.36% (95% CI: 2.40%–13.00%; p=0.005).	• Financial incentives may constitute an insufficiently strong intervention to influence goal commitment when providers attribute performance to external forces beyond their control.
Karunaratne K, et al., 2013 (343) <u>23658247</u>	Aim: The aim of this study was to evaluate the effectiveness of renal indicators outlined in P4P on the management of HTN in primary care. To estimate the cost implications of the resulting changes in prescribing patterns of antihypertensive medication following introduction of such indicators. Study type: Prospective cohort study using a large primary care database.	Inclusion criteria: A total of 10,040 pts had confirmed stage 3–5 CKD in the 2 y pre-QOF and formed the study cohort. Exclusion criteria: None	Intervention: The implementation of national estimated GFR reporting and the inclusion of renal- specific indicators in a primary care P4P system since April 2006 has promoted identification and better management of risk factors related to CKD. In the UK, the P4P framework is known as the QOF. Comparator: N/A	 Mean age of the cohort at the start of the study period was 64.8 y, 55% were female. In those pts with stage 3–5 CKD 83.9% were hypertensive, defined by a pre-P4P BP of >140/85 or currently taking antihypertensive medication. The proportion of pts with CKD 3–5 attaining the BP target of 145/80 increased from 41.5% in the pre-QOF period to 50.0% in the post-QOF period. This increase was even more marked for those with HTN in the pre-QOF period (28.8%–45.1%). In the hypertensive pts, mean BP fell from 146/79 mm Hg to 140/76 in the first 2 y post-P4P [p<0.01, analysis of variance]. 	Summary: Population BP control has improved since the introduction of P4P renal indicators, and this improvement has been sustained. This was associated with a significant increase in the use of antihypertensive medication, resulting in increased prescription cost. Longer-term follow-up will establish whether or not this translates to improved outcomes in terms of progression of CKD, CVD and pt mortality.

	This cohort was taken from a database collated as part of a clinical decision support system used to assist the management of CKD in primary care. <u>Size</u> : 90,250 pts on general practitioner registers with a valid serum creatinine estimation in the 6-y study period. A total of 10 040 pts had confirmed stage 3–5 CKD in the 2 y pre- QOF and formed the study cohort.			BP reduction was sustained in the last 2 y of the study, 139/75 (p<0.01, analysis of variance). The proportion of hypertensive pts taking ACEIs or angiotensin blockers increased, this was also sustained in the third time period. An increase in the prescribing of diuretics, CCBs and BBs was also observed. The additional cost of increased prescribing was calculated to be euro 25.00 per hypertensive pt based on GP prescription data.	
Serumaga B, et al., 2011 (344) <u>21266440</u>	Aim: The aim of this study was to evaluate the effectiveness of renal indicators outlined in P4P on the management of HTN in primary care. To estimate the cost implications of the resulting changes in prescribing patterns of antihypertensive medication following introduction of such indicators. Study type: Interrupted time series study	Inclusion criteria: Pts with HTN diagnosed between Jan. 2000–Aug. 2007. Exclusion criteria: None	Intervention: The UK P4P incentive (the Quality and Outcomes Framework), which was implemented in April 2004 and included specific targets for general practitioners to show high quality care for pts with HTN (and other diseases). Comparator: None	• After accounting for secular trends, no changes in BP monitoring: level change: 0.85 (95% CI: -3.04-4.74), p=0.669 and trend change: -0.01, (95% CI: -0.24-0.21), p=0.615, control: -1.19 (95% CI: -2.06- 1.09), p=0.109 and -0.01 (95% CI: -0.06-0.03), p=0.569, or treatment intensity; 0.67: (95% CI: -1.27-2.81), p=0.412 and 0.02 (95% CI: -0.23-0.19, p=0.706 were attributable to P4P. P4P had no effect on the cumulative incidence of stroke, MI, renal failure, HF, or all- cause mortality in both treatments experienced and newly treated subgroups.	Summary: Good quality of care for HTN was stable or improving before P4P was introduced. P4P had no discernible effects on processes of care or on HTN related clinical outcomes. Generous financial incentives, as designed in the UK P4P policy, may not be sufficient to improve quality of care and outcomes for HTN and other common chronic conditions.

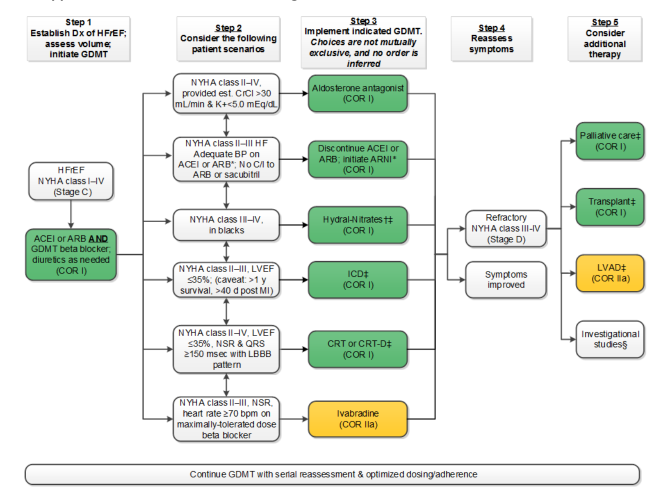
Bardach NS, et	Size: 470,725 pts with HTN diagnosed between Jan 2000– Aug 2007. al., Aim: To assess the	Participating clinics	A city program	 Intervention clinics had greater 	Summary: In our study, although the
2013 (327) 24026600	 and the intervention of the inter	(n=42 for each group) had similar baseline characteristics, with a mean of 4,592 (median, 2,500) pts at the intervention group clinics and 3,042 (median, 2,000) at the control group clinics.	 A city program provided all participating clinics with the same EHR software with decision support and pt registry functionalities and QI specialists offering technical assistance. Incentivized clinics were paid for each pt whose care met the performance criteria, but they received higher payments for pts with comorbidities, who had Medicaid insurance, or who were uninsured (maximum payments: \$200/pt; 100,000/clinic). Quality reports were given quarterly to both the intervention and control groups. 	adjusted absolute improvement in rates of appropriate antithrombotic prescription 12.0% vs. 6.1%, difference: 6.0% (95% CI: 2.2%–9.7%; p=0.001 for interaction term), BP control (no comorbidities): 9.7% vs. 4.3%, difference: 5.5% (95% CI: 1.6%–9.3%; p=0.01 for interaction term); with DM: 9.0% vs. 1.2%, difference: 7.8% (95% CI: 3.2%–12.4%; p=0.007 for interaction term); with DM or ischemic vascular disease: 9.5% vs. 1.7%, difference: 7.8% (95% CI: 3.0%–2.6%; p=0.01 for interaction term), and in smoking cessation interventions (12.4% vs. 7.7%), difference: 4.7% (95% CI: -0.3%–9.6%; p=0.02 for interaction term). Intervention clinics performed better on all measures for Medicaid and uninsured pts except cholesterol control, but no differences were statistically significant.	effect of the intervention was lower than the 10% improvement that we estimated a priori, the absolute risk reduction for BP control among pts with DM was 7.8% (NNT, 13). This suggests that, for every 13 pts seeing incentivized clinicians, 1 more pt would achieve BP control. The 7.8% absolute change in BP control for pts with DM represents a 46% relative increase in BP control among intervention pts compared with the baseline of 16.8%. Further research is needed to determine whether this effect of the P4P intervention on BP control increases or decreases over time. However, this NNT to achieve BP control through incentives, taken together with the large relative increase in percentage of pts with BP control and the potential effect of BP control on risk of ischemic vascular events, suggests a reasonable opportunity to reduce morbidity and mortality through P4P as structured in this study.

					motivation to improve among practices in the study, as demonstrated by engagement with the QI specialists
Maimaris W, et al., 2013 (345) <u>23935461</u>	Aim: To assess strategies for influencing HTN care including procurement of essential medications, the existence of simple national guidelines for HTN management, introduction of financial incentives for health care practitioners to diagnose or treat HTN, and enhanced health insurance coverage. <u>Study type</u> : Systematic review examining the effect of national or regional health system arrangements on HTN care and control	Study selection criteria based on: 1) HTN awareness. Defined as pts with clinically measured hypertensives who have been diagnosed by a health care professional as hypertensive. 2) HTN treatment. Defined as the use of at least 1 antihypertensive medication in a pt with known HTN. 3) Antihypertensive medication adherence. Defined as consistently taking the antihypertensive medication regimen as prescribed by the health care provider. 4) HTN control: defined as the achievement of BP<140/90 mm Hg (or other explicitly defined threshold) in individuals being treated for HTN, or, alternatively, measured by the mean BP amongst individuals with HTN.	• The screening process is described using an adapted PRISMA flowchart. 5,514 articles were screened by title and abstract for inclusion. The full text of 122 of the 5,514 articles was obtained and assessed for eligibility. 53 studies met eligibility criteria for this review. 51 of the included studies were quantitative and 2 were qualitative. Of the 51 quantitative studies, 1 was an RCT; 12 were cohort studies, 2 of which were retrospective; 3 were case-control studies; 32 were cross-sectional studies; and 3 were ecological studies. 42 of the 53 studies (79%) were carried out in countries classified by the World Bank as high- income countries, 36 of which were in the U.S. 6 studies were carried out in upper middle-income countries, 3 in lower middle-income	 Health insurance status: 15 cross-sectional studies reported comparisons of HTN outcomes in insured and uninsured pts. 8 of these 15 studies reported that insurance was associated with improved HTN treatment, control or medication adherence. The 7 other cross- sectional studies that compared HTN outcomes in insured pts and uninsured pts, reported no significant negative or positive associations between insurance status and HTN outcome. Medication costs or medication co-payments: All 6 of these studies reported significant associations between reduced co-payments or costs and improved HTN control or medication adherence. Co-payments for medical care: 14 quantitative studies measured the association of medication co-payments or costs with HTN control or treatment adherence, 9 of which were set in the U.S., and 1 in each of Cameroon, China, Finland, Israel, and Brazil. 2 of the 14 studies had a low risk of bias. 7 of the 14 studies were cohort studies, 1 was a case- control study, and 6 were cross- sectional studies. All 7 cohort 	• Although lacking longitudinal studies, we found a large positive association between having a routine physician or place of care for HTN management and treatment, awareness, control, and adherence to antihypertensive treatment, again in the U.S. publication and reporting bias noted by authors.

countries, and 1 in a	studios reported associations
	studies reported associations
low-income country.	between increased medication
	costs or co-payments and
	reductions in HTN control or
	reduced adherence to
	antihypertensive medication,
	although for 1 of these 7 cohort
	studies, the association between
	increased copayments and
	reduced medication adherence
	was only found for low
	medication co-payments, and at
	high co-payment levels
	medication adherence was
	actually found to increase (OR
	for medication adherence vs.
	baseline of 1 for \$0 co-
	payments was 0.72 for \$1-\$9
	co-payments (p=0.05), 1.02 for
	\$10-\$29 co-payments (p=0.05),
	and 1.32 for co-payments . \$30
	(p=0.05)
	Physician remuneration
	models: 2 studies evaluated the
	association of physician
	remuneration models with HTN
	control or treatment adherence,
	1 an ecological study set in
	Canada, and 1 a U.S. cross-
	sectional study. Neither study
	had a low risk of bias. The U.S.
	study reported improved rates of
	HTN control amongst pts treated
	under a capitation model
	compared to fee-for service pts
	(adjusted OR for HTN control:
	1.82 (95% CI: 1.02–3.27) for
	capitation vs. fee-for-service
	pts). The Canadian study
	reported highest rates of HTN

	treatment and control among practices using a capitation model, compared to fee-for- service and salary model. HTN awareness levels were highest	
	in practices with a fixed salary remuneration model.	

Additional Data Supplement Tables and Figures



Data Supplement A. Treatment of HFrEF Stages C and D

Colors correspond to COR in Table 1. For all medical therapies dosing should be optimized and serial assessment exercised.

*See text for important treatment directions.

[†]Hydral-Nitrates Green Box- The combination of ISDN/HYD with ARNI has not been robustly tested. BP response should be carefully followed.

‡See 2013 HF guideline.

§Participation in investigational studies is also appropriate for stage C, NYHA class II and III HF.

ACEI indicates angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; ARNI, angiotensin receptor-neprilysin inhibitor; BP, blood pressure; bpm, beats per minute; C/I, contraindication; CRT-D, cardiac resynchronization therapy-device; COR, class of recommendation; Dx, diagnosis; GDMT, guideline-directed management and therapy; HF*r*EF, heart failure with reduced ejection fraction; ICD, implantable cardioverter-defibrillator; ISDN/HYD, isosorbide dinitrate hydral-nitrates; LBBB, left bundle-branch block; LVEF, left ventricular

ejection fraction; LVAD, left ventricular assist device; NSR, normal sinus rhythm; and NYHA, New York Heart Association.

How often do you:	Response:
 Forget to take your high BP medicine? Decide NOT to take your high BP medicine? Eat salty foods Shake salt on your food before you eat it? Eat fast food? 	1. All of the Time 2. Most of the Time 3. Some of the Time 4. None of the Time
 Make the next appointment before you leave the doctor's office? Miss scheduled appointments? Forget to get prescriptions filled? Run out of high BP pills? Skip your high BP medicine before you go to the doctor? Miss taking your high BP pills when you feel better? Miss taking your high BP pills when you feel sick? Take someone else's high BP pills? Miss taking your high BP pills when you are careless? 	Medication taking subscale: Items 1, 8,9,10,11,12,13,14. Reducing sodium intake subscale: Items 3,4,5. Appointment keeping subscale: Item 6,7.

Data Supplement B. Medication Adherence Assessment Scales

BP indicates blood pressure.

Data Supplement C. Categories Defining Normal BP, Elevated BP, and Stages 1, 2, and 3 Hypertension

		SBP (mm Hg) →					
		<120	120–129	130–139	140–159	160+	
(mm Hg)	<80	Normal	Elevated	Stage 1	Stage 2	Stage 2	
DBP (n	80–89	Stage 1	Stage 1	Stage 1	Stage 2	Stage 2	
\downarrow	90–99	Stage 2	Stage 2	Stage 2	Stage 2	Stage 2	
	100+	Stage 2	Stage 2	Stage 2	Stage 2	Stage 2	

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Stages 1, 2, and 3 refer to the stage of hypertension.

BP indicates blood pressure; DBP, diastolic blood pressure; and SBP, systolic blood pressure.

Class	Drug	Dosage Strengths (mg/mg)	Daily Frequency*
2-drug combinations			
ACE Inhibitors + Thiazide	Benazepril/Hydrochlorothiazide	10/12.5, 20/12.5, 20/25	1
	Captopril/Hydrochlorothiazide	25/15, 50/15, 25/25, 50/25	2
	Enalapril/Hydrochlorothiazide	5/12.5, 10/25	1 or 2
	Fosinopril/Hydrochlorothiazide	10/12.5, 20/12.5	1
	Lisinopril/Hydrochlorothiazide	10/12.5, 20/12.5, 20/25	1
	Moexipril/Hydrochlorothiazide	7.5/12.5, 15/12.5, 15/25	1 or 2
	Quinapril/Hydrochlorothiazide	10/12.5, 20/12.5, 20/25	1 or 2
ARBs + Thiazide	Azilsartan/Chlorthalidone	40/12.5, 40/25	1
	Candesartan/Hydrochlorothiazide	16/12.5, 32/12.5, 32/25	1
	Eprosartan/Hydrochlorothiazide	600/12.5, 600/25	1
	Irbesartan/Hydrochlorothiazide	150/12.5, 300/12.5, 300/25	1
	Losartan/Hydrochlorothiazide	50/12.5, 100/12.5, 100/25	1 or 2
	Olmesartan/Hydrochlorothiazide	20/12.5, 40/12.5, 40/25	1
	Telmisartan/Hydrochlorothiazide	40/12.5, 80/12.5, 80/25	1
	Valsartan/Hydrochlorothiazide	80/12.5, 160/12.5, 320/12.5,	1
		160/25, 320/25	_
CCB – dihydropyridine + ACEIs	Amlodipine/Benazepril	2.5/10, 5/10, 5/20, 10/20, 5/40, 10/40	1
	Enalapril/Felodipine	5/5	1
	Perindopril/Amlodipine	3.5/2.5, 7/5, 14/10	1
CCB – dihydropyridine + ARB	Amlodipine/Olmesartan	5/20, 10/20, 4/40	1
	Amlodipine/Valsartan	5/160, 10/160, 5/320, 10/320	1
	Telmisartan/Amlodipine	40/5, 80/5, 40/10, 80/10	1
CCB – nondihydropyridine + ACEIs	Trandolapril/Verapamil	2/180, 1/250, 2/240, 4/240	1
Beta blocker + Thiazide	Atenolol/Chlorthalidone	50/25, 100/25	1
	Bisoprolol/Hydrochlorothiazide	2.5/6.25, 5/6.25, 10/6.25	1
	Metoprolol succinate/Hydrochlorothiazide	25/12.5, 50/12.5, 100/12.5	1
	Metoprolol tartrate/ Hydrochlorothiazide	50/25, 100/25, 100/50	1 or 2
	Nadolol/Bendroflumethiazide	40/5, 80/5	1
	Propranolol/Hydrochlorothiazide	40/25, 80/25	1 or 2
Direct renin inhibitor + CCB – dihydropyridine	Aliskiren/amlodipine	150/5, 150/10, 300/5, 300/10	1
Direct renin inhibitor + Thiazide	Aliskiren/ Hydrochlorothiazide	150/12.5, 150/25, 300/12.5, 300/25	1
Direct renin inhibitor + CCB – dihydropyridine	Aliskiren/Amlodipine	150/12.3, 150/23, 300/12.3, 300/23	1
Direct renin inhibitor + Thiazide	Aliskiren/Hydrochlorothiazide	150/12.5, 150/25, 300/12.5, 300/25	1
Central acting agent + Thiazide	Clonidine/Chlorthalidone		1 1 or 2
Central acting agent + miazide	Methyldopa/Hydrochlorothiazide	0.1/15, 0.2/15, 0.3/15 250/15, 250/25	2
Diuratic potassium sparing	Amiloride/Hydrochlorothiazide		1
Diuretic- potassium sparing + Thiazide	Triamterene/Hydrochlorothlazide	5/50	1
Diuretic- aldosterone antagonist +		37.5/25, 75/50	
Thiazide	Spironolactone/ Hydrochlorothiazide	25/25	1 or 2
3-drug combinations			1
ARB + CCB – dihydropyridine + Thiazide	Amlodipine/Valsartan/ Hydrochlorothiazide	5/160/12.5, 10/160/12.5, 5/160/25, 10/160/25, 10/320/25	1
	Olmesartan/Amlodipine/ Hydrochlorothiazide	20/5/12.5, 40/5/12.5, 40/5/25, 40/10/12.5, 40/10/25	1
Direct renin inhibitor + CCB – dihydropyridine + Thiazide	Aliskiren/Amlodipine/Hydrochlorothiazide	150/5/12.5, 300/5/12.5, 300/5/25, 300/10/12.5, 300/10/25	1
	in the EDA approved labeling http://dailumed	nlm nih gov/dailumad/indox.cfm)	

Data Supplement D. Fixed-Dose Combination Antihypertensive Drugs

*Dosages may vary from those listed in the FDA approved labeling http://dailymed.nlm.nih.gov/dailymed/index.cfm).

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ACE indicates angiotensin-converting enzyme; ARB, angiotensin-receptor blocker; and CCB, calcium channel blocker.

From Chobanian et al. JNC 7. (347)

Data Supplement E. Examples of Hypertension Quality Improvement Strategies

Quality Improvement Strategy	Examples
Audit and feedback on performance	 Feedback of performance to individual providers Benchmarking – provision of outcomes data from top performers for comparison with provider's own data Performance measures, quality indicators and reports Use of registries to track BP control status at system and provider levels
Provider education	 In person, online, or other education to improve BP measurement and management skills Training to improve communication, cultural competency, and ability to inspire and support lifestyle modification
Patient education	 Intensive education strategies promoting hypertension self- management Cultural and linguistic tailoring of materials to increase acceptability
Promotion of self-management	Reduce barriers for patients to receive and adhere to medications and to implement lifestyle modification
Patient reminder systems	Postcards, calls, texts, or emails to patients
(for follow-up appointments, BP checks, and self-management)	Telehealth-delivered reminders
System change	 Standardization of BP measurement using an automated device and standardized protocol Screening to identify all patients eligible for hypertension management Systematic follow-up of patients for the initiation and intensification of antihypertensive therapy Decision support to providers to guide protocol-based treatment decisions Physician or other clinical champion designated to lead hypertension care improvement initiatives Hypertension specialist available for consult Partner with community resources to support BP management

BP indicates blood pressure.

Adapted with permission from Walsh et al. (348).

Data Supplement F. Barriers and Improvement Strategies in Antihypertensive Medication Adherence (349-353)

Barriers	Improvement Strategies
Patient Level	
 Multiple comorbid conditions requiring complex medication regimens Convenience factors (e.g., dosing frequency) Health beliefs Behavioral factors Lack of involvement in the treatment decision-making process Issues with treatment of asymptomatic diseases (e.g., treatment side effects) Resource constraints Suboptimal health literacy 	 Educate patients about hypertension, consequences of hypertension, and possible adverse effects of medications Collaborate with patient to establish goals of therapy and plan of care Maintain contact with patients; consider telehealth approaches (Section 12.3.2). Integrate pill-taking into daily routine activities of daily living with adherence support tools such as reminders, pillboxes, packaging, or other aids Use motivation interventions to support medication adherence and lifestyle modification efforts Use medication adherence scales to facilitate identification of barriers and facilitators to and behaviors associated with adequate adherence Address health literacy Teach-back method Empower patients to ask questions Use visual, interactive education Health literacy universal precautions tool kit Provide medication list/pictorial medication schedule
Provider and Health System Levels	6 Trovide medication had pictorial medication schedule
 Prescription of complex drug regimens Inadequate communication with patient about regimen, adverse effects, treatment goals Inadequate communication among multiple providers Office visit time limitations Limited access to care, pharmacies, prescription refills 	 Assess for nonadherence and explore barriers to medication adherence Use a multifactorial approach to optimize adherence Participate in training to enhance communication skills and increase cultural competence Use a multifactorial approach to optimize adherence Reduce complexity of medication regimen Utilize agents that are dosed once daily over those which require multiple daily doses Utilize fixed-dose combination agents when available and simplify drug regimens Consider overall side effect profile and preferentially use agents that are well tolerated Use low-cost and generic antihypertensives from drug classes where RCTs have demonstrated a reduction in cardiovascular events when appropriate (354) Use team-based care approaches (Section 12.2) Use health information technology-based approaches (Section 12.3)

RCTs indicate randomized controlled trials.

Data Supplement G. Examples of Strategies to Promote Lifestyle Modification Interventions in Patients With Hypertension (318, 319, 355-361)

	Lifestyle Modification Intervention	References
Tobacco Cessation	Ask all adults about tobacco use	(361, 362)
	 Advise them to stop using tobacco 	
	Provide behavioral interventions	
	Consider pharmacotherapy for tobacco cessation	
Weight Loss	• Offer or refer obese adults to intensive cognitive and behavioral interventions aimed at to improve weight status and other risk factors for important health outcomes.	(355, 356)
Sodium Reduction	 Offer or refer to behavioral counselling aimed at reduced intake of dietary sodium Encourage use of food labels to choose lower sodium products 	
Alcohol	 Screen adults ≥18 y of age for alcohol misuse and provide persons engaged in risky or hazardous drinking with behavioral counseling interventions to reduce alcohol misuse. 	(357, 358)
Physical Activity and Diet	 Use medium- to high-intensity behavioral counseling interventions to improve intermediate health outcomes; addressing barriers, such as lack of access to affordable healthier foods, transportation barriers and poor local safety. 	(359, 360)

Data Supplement H. Responsibilities and Roles of the Hypertension Team

Hypertension Team Responsibilities

- Communication and care coordination among various team members, the patient and family members or other support persons.
- Effective use of evidence-based diagnosis and management guidelines
- Regular, structured follow-up mechanisms and reminder systems to monitor patient progress
- Engage patients in their care by shared decision making
- Medication adherence support and appropriate education about hypertension medication
- Medication addition and titration using evidence-based treatment algorithms
- Use of evidence-based tools and resources designed to maximize self-management (including health behavior change, lifestyle modification, etc.)
 Follow a single personalized plan of area based when patient characteristics and peeds

 Follow a single, personalized plan of 	f care based upon patient characteristics and needs
Individual Hypertension Team Members	Roles (examples)
Primary Care Physician, Physician	Routine and complex hypertension care, managing primary care
Assistant, Advanced Practice Nurse	issues.
Cardiologist	Routine and complex hypertension care, especially for patient with
	cardiac disease or high risk for major cardiovascular events.
Nephrologist, Endocrinologist,	Management of complex hypertension care, especially due to
Hypertension Specialist	secondary causes, and/or resistant hypertension.
Nurse (including in-office, home care,	Accurate assessment of BP, medication reconciliation, patient
internal and external population health	education, self-management, lifestyle modification and adherence.
personnel)	
Clinical Pharmacist	Comprehensive medication management, which involves identification
	and documentation of medication-related problems, initiating,
	modifying, and discontinuing medication to address identified
	problems, and educating patients on their medication regimen.
Dietician	Ongoing patient-centered counseling to assess dietary habits and
	preferences, set and monitor goals for healthy lifestyle
Social Worker	Assess for psychosocial, cultural and financial barriers, find solutions
	to overcome these barriers.
Community Health Providers	Assess for psychosocial, cultural and financial barriers, identify and
	promote acceptable community-based resources to overcome these
	barriers.

BP indicates blood pressure.

Data Supplement I. Examples of Telehealth Strategies and Technologies to Promote Effective Hypertension Management

Telehealth strat	egies
Automated	BP data capture and transmission of the patient's self-measured BP
Self-manage	ement support including education, reminders, and feedback that is automated or delivered by a
healthcare	professional
Medication	titration and follow-up monitoring protocols/algorithm
Prescription	n refill reminders
Medication	adherence assessments
Self-monito	ring of lifestyle behaviors
Integration	of behavior change techniques, including in person or e-counseling
Case/care/p	population health management
Commonly used	I telehealth technologies
Wired "land	l line" telephone
Wireless sm	nart phone applications
• Internet-ba	sed website via computers and handheld devices
Text message	ging
E-mail mess	saging
Social netw	orking and social media websites/applications
Wireless BP	measurement devices
Electronic p	ill dispensers/counters

BP indicates blood pressure.

Data Supplement J. Publicly Available Performance Measures Used to Assess Hypertension Care Quality Services (363-367)

Quality Measure	Source	Description	Additional information
Controlling High BP PQRS Measure #236; NQF #0018	NCQA	Percentage of patients 18–85 y of age who had a diagnosis of hypertension and whose BP was adequately controlled (<140/90 mm Hg during the measurement period)	Used in the CMS, PQRS, MSSP, Medicare Advantage "Stars" ratings; component of Commercial Health Plan HEDIS quality measure set
Comprehensive Diabetes Care: BP Control (<140/90 mm Hg) NQF #0061	NCQA	The percentage of patients 18–75 y of age with DM (type 1 and type 2) whose most recent BP level taken during the measurement y is <140/90 mm Hg	 Used for: Accreditation Decision-making by businesses about health plan purchasing Decision-making by consumers about health plan/provider choice External oversight/Medicaid External oversight/Medicare External oversight/State government program Internal quality improvement Public reporting
Adult Kidney Disease: BP Management PQRS #122	PCPI, RPA	Percentage of patient visits for those patients ≥18 y of age with a diagnosis of CKD (stage 3, 4, or 5, not receiving renal replacement therapy) with a BP<140/90 mm Hg OR ≥140/90 mm Hg with a documented plan of care	Used in PQRS
Percentage of patients ≥18 y of age with BP documented in the medical record (every 2 y if <120/80 mm Hg, every y if 120–139/80–89 mm Hg)	ICSI	This measure is used to assess the percentage of patients age 18 y of age and older with BP documented in the medical record (every 2 y if <120/80 mm Hg, every y if 120– 139/80–89 mm Hg)	Used for internal quality improvement
Controlling High BP for People with Serious Mental Illness NQF #2602	NCQA	The percentage of patients 18–85 y of age with serious mental illness who had a diagnosis of hypertension and whose BP was adequately controlled during the measurement	 Current Use: Accreditation Decision-making by businesses about health plan purchasing Decision-making by consumers about health plan/provider choice External oversight/Medicaid External oversight/state government program \internal quality improvement
Diabetes Care for People with Serious Mental Illness: BP Control (<140/90 mm Hg) NQF #2606	NCQA	The percentage of patients 18–75 y of age with a serious mental illness and DM (type 1 and type 2) whose most recent BP reading during the measurement year is <140/90 mm Hg	 Current Use: Accreditation Decision-making by businesses about health plan purchasing Decision-making by consumers about health plan/provider choice External oversight/Medicaid

Quality Measure	Source	Description	Additional information
			 External oversight/state government program Internal quality improvement
Hypertension diagnosis and treatment: percentage of adult patients ≥18 y of age diagnosed with hypertension who are not at goal for hypertension and have received counseling on diet and physical activity in the past 12 mo	ICSI	Used to assess the percentage adult patients ≥ 18 y of age diagnosed with hypertension who are not at goal for hypertension and have received counseling on diet and physical activity in the past 12 mo	Used for Internal Quality Improvement
Ambulatory care sensitive conditions: age-standardized acute care hospitalization rate for conditions where appropriate ambulatory care prevents or reduces the need for admission to the hospital per 100,000 population <75 y of age	СІНІ	Used to assess the age-standardized acute care hospitalization rate for conditions where appropriate ambulatory care prevents or reduces the need for admission to the hospital per 100,000 population <75 y of age	Used for: Monitoring health state(s) National health policymaking National reporting State/Provincial health policymaking
Hypertension: the relative resource use by members with hypertension during the measurement y	NCQA	Used to assess the relative resource use by members with hypertension by reporting total standard cost and service frequency for all services for which the organization has paid or expects to pay during the measurement y	 Used for: Accreditation External oversight/Medicaid External oversight/Medicare External oversight/State government program Monitoring and planning Public reporting

BP indicates blood pressure; CIHI, Canadian Institute for Health Information; CKD, chronic kidney disease; CMS, Centers for Medicare and Medicaid Services; DM, diabetes mellitus; HEDIS, healthcare Effectiveness Data and Information Set; ICSI, Institute for Clinical Systems Improvement; MSSP, Medicare Shared Savings Program; NCQA, National Committee for Quality Assurance; NQF, National Quality Forum; OR, odds ratio; PCPI, Physician Consortium for Performance Improvement; and PQRS, Physician Quality Reporting System; and RPA, Renal Physicians Association.

Data Supplement K. Online Quality Improvement Resources for Treatment and Control of Hypertension

<u>American College of Cardiology/American Heart Association/Centers for Disease Control</u> Science Advisory for the Effective Approach to High Blood Pressure Controlⁱ

http://content.onlinejacc.org/article.aspx?articleid=1778408

<u>American Medical Association</u> Measure, Act and Partner (M.A.P.) to help patients control blood pressure and ultimately prevent heart disease

http://www.ama-assn.org/ama/pub/about-ama/strategic-focus/improving-health-outcomes/improving-blood-pressure-control.page

<u>United States Health and Human Services (HHS)/Centers for Disease Control (CDC)</u> Million Hearts Campaign Evidence-based Treatment Protocols for Improving Blood Pressure Control

http://millionhearts.hhs.gov/resources/protocols.html

Department of Defense/Veterans' Affairs

http://www.healthquality.va.gov/guidelines/CD/htn/

Kaiser Permanente Hypertension Management programs to improve blood pressure control

http://kpcmi.org/how-we-work/hypertension-control/

Institute for Clinical Systems Improvement (ICSI) Hypertension Diagnosis and Treatment Guidelines

https://www.icsi.org/guidelines more/catalog guidelines and more/catalog guidelines/catalog cardiovascular g uidelines/hypertension/

New York Health and Hospitals Corporation (HHC) Hypertension Collaborative Care Pathway

http://millionhearts.hhs.gov/Docs/NYC HHC Hypertension Protocol.pdf

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Author Relationships With Industry and Other Entities (Comprehensive)—2017 ACC/AHA/AAPA/ABC/ACPM/AGS/APhA/ASH/ASPC/NMA/PCNA Guideline for the Prevention, Detection, Evaluation, and Management of High Blood Pressure in Adults (October 2017)

Committee Member	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness	Salary
Paul K. Whelton (Chair)	Tulane University School of Hygiene and Tropical Medicine—Show Chwan Professor of Global Public Health	None	None	None	• NIH–SPRINT trial† (PI)	None	None	None
Robert M. Carey (Vice Chair)	University of Virginia—Dean Emeritus and University Professor, Department of Medicine	None	None	None	• NIH†	None	None	None
Wilbert S. Aronow	Westchester Medical Center and New York Medical College—Professor of Medicine	None	None	None	None	None	None	None
Donald E. Casey, Jr	Thomas Jefferson College of Population Health—Adjunct Faculty; Alvarez & Marsal Ipo4health— Principal and Founder	None	None	None	None	None	None	None
Karen J. Collins	Collins Collaboration— President	None	None	None	None	North Carolina A&T State University Alumni Association‡	None	None

Cheryl Dennison Himmelfarb	John Hopkins University— Professor of Nursing and Medicine, Institute for Clinical and Translational Research	MedThink Communicatio ns	None	None	 Helene Fuld Health Trust† NIH† 	• Preventive Cardiovascular Nurses Association‡	None	None
Sondra M. DePalma	PinnacleHealth CardioVascular Institute—Physician Assistant; American Academy of PAs— Director, Regulatory and Professional Practice	American Society of Hypertension	None	None	None	• Accreditation Council for Clinical Lipidology‡	None	None
Samuel Gidding	Alfred I. Dupont Hospital for Children—Chief, Division of Pediatric Cardiology, Nemours Cardiac Center	 Familial Hypercholester olemia Foundation‡ Regenxbio 	None	None	 Familial Hypercholesterolemia Foundation‡ NIH† 	• Cardiology Division Head‡	None	None
David C. Goff, Jr*	Colorado School of Public Health— Professor and Dean, Department of Epidemiology	None	None	None	None	None	None	None
Kenneth A. Jamerson	University of Michigan Health System—Professor of Internal Medicine and Frederick G.L. Huetwell Collegiate Professor of Cardiovascular Medicine	• American Society of Hypertension	None	None	• NIH/NIDDK/NHLBI†	 American Society of Hypertension‡ International Society of Hypertension In Blacks‡ Bayer Healthcare Pharmaceuticals 	None	None

Daniel W. Jones	University of Mississippi Medical Center— Professor of Medicine and Physiology; Metabolic Diseases and Nutrition— University Sanderson Chair in Obesity Mississippi Center for Obesity Research— Director, Clinical and Population	None	None	None	None	None	None	None
Eric J. MacLaughlin	Science Texas Tech University Health Sciences Center— Professor and Chair, Department of Pharmacy Practice, School of Pharmacy	• American Society of Hypertension	None	None	None	 AHA‡ American College of Clinical Pharmacy‡ American Pharmacists Association‡ Texas Tech University Health Sciences Center† NIH 	None	None
Paul Muntner	University of Alabama at Birmingham— Professor, Department of Epidemiology	 Amgen Inc. National Center for Health Statistics[†] 	None	None	 AHA† Amgen Inc.† NIH† 	None	None	None
Bruce Ovbiagele	Medical University of South Carolina— Pihl Professor and Chairman of Neurology	None	None	None	None	None	None	None
Sidney C. Smith, Jr	University of North Carolina at Chapel Hill—Professor of Medicine; Center for Cardiovascular Science and Medicine—Director	None	None	None	None	None	None	None

Crystal C. Spencer	Spencer Law, PA— Attorney at Law	None	None	None	None	 AHA‡ Dermatologic Surgery Associates† Hospital Corporation of America† 	None	None
Randall S. Stafford	Stanford Prevention Research Center— Professor of Medicine; Program on Prevention Outcomes— Director	None	None	None	None	None	None	None
Sandra J. Taler	Mayo Clinic— Professor of Medicine, College of Medicine	None	None	None	None	None	None	 American Society of Hypertension Clinical Specialist Program† American Society of Nephrology†
Randal J. Thomas	Mayo Clinic— Medical Director, Cardiac Rehabilitation Program	None	None	None	None	None	None	None
Kim A. Williams, Sr	Rush University Medical Center— James B. Herrick Professor; Division of Cardiology— Chief	None	None	None	None	None	None	None
Jeff D. Williamson	Wake Forest Baptist Medical Center— Professor of Internal Medicine; Section on Gerontology and Geriatric Medicine—Chief	None	None	None	None	None	None	None

Jackson T. Wright, Jr	Case Western	None	None	None	None	Northeast Ohio	None	None
	Reserve					Neighborhood		
	University—					Health Centers‡		
	Professor of							
	Medicine; William							
	T. Dahms MD							
	Clinical Research							
	Unit—Program							
	Director; University							
	Hospitals Case							
	Medical Center-							
	Director, Clinical							
	Hypertension							
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†Significant relationship. ‡No financial benefit.

AAPA indicates American Academy of Physician Assistants; ABC, Association of Black Cardiologists; ACC, American College of Cardiology; ACPM, American College of Preventive Medicine; AGS, American Geriatrics Society; AHA, American Heart Association; APhA, American Pharmacists Association; ASH, American Society of Hypertension; ASPC, American Society for Preventive Cardiology; NIDDK, National Institute of Diabetes and Digestive and Kidney Diseases; NHLBI, National Heart, Lung, and Blood Institute; NIH, National Institutes of Health; NMA, National Medical Association; PCNA, Preventive Cardiovascular Nurses Association; and PI, principal investigator.