Long Term Prognostic Impact of Sex-specific Longitudinal Changes in Blood Pressure. The EPIC-Norfolk Prospective Population Cohort Study

Tiberiu A. Pana (1) 1,2, Robert N. Luben (1) 3, Mamas A. Mamas (1) 4, John F. Potter (1) 5,6, Nicholas J Wareham⁷, Kay-Tee Khaw (1) 3, and Phyo K. Myint (1) 1,2,5,6*

¹Aberdeen Diabetes and Cardiovascular Centre, School of Medicine, Medical Sciences & Nutrition, University of Aberdeen, Room 4:013, Polwarth Building, Foresterhill, Aberdeen AB25 2ZD, UK; ²Ageing Clinical and Experimental Research (ACER) Team, University of Aberdeen, Aberdeen, UK; ³Clinical Gerontology Unit, Department of Public Health & Primary Care, University of Cambridge, Worts Causeway, Cambridge CB1 8RN, UK; ⁴Keele Cardiovascular Research Group, Keele University, Stoke-on-Trent, University Road, Staffordshire ST5 5BG, UK; ⁵Department of Clinical Neuroscience, Norfolk and Norwich University Hospital, Norwich, UK; ⁶Norwich Cardiovascular Research Group, Norwich Medical School, University of East Anglia, Norwich Research Park, Centrum, Colney Lane, Norwich NR4 7UG, UK; and ⁷MRC Epidemiology Unit, Institute of Metabolic Science, University of Cambridge School of Clinical Medicine, Cambridge CB2 0SL, UK

Received 16 January 2021; revised 26 March 2021; editorial decision 25 May 2021; accepted 28 May 2021

Aims

We aimed to determine the sex differences in longitudinal systolic and diastolic blood pressure (SBP and DBP) trajectories in mid-life and delineate the associations between these and mortality (all-cause, cardiovascular, and non-cardiovascular) and incident cardiovascular disease (CVD) in old age.

Methods and results

Participants were selected from the European Prospective Investigation into Cancer, Norfolk (EPIC-Norfolk) cohort study. Sex-specific trajectories were determined using group-based trajectory models using three clinic BP measurements acquired between 1993 and 2012 (mean exposure ~12.9 years). Multivariable Cox regressions determined the associations between trajectories and incident outcomes over the follow-up (median follow-up 9.4 years). A total of 2897 men (M) and 3819 women (F) were included. At baseline, women were younger (F-55.5, M-57.1), had a worse cardiometabolic profile and were less likely to receive primary CVD prevention including antihypertensive treatment (F-36.0%, M-42.0%). Over the exposure period, women had lower SBP trajectories while men exhibited more pronounced SBP decreases over this period. Over the follow-up period, women had lower mortality (F-11.9%, M-20.5%) and CVD incidence (F-19.8%, M-29.6%). Compared to optimal SBP (≤120 mmHg) and DBP (≤70 mmHg) trajectories, hypertensive trajectories were associated with increased mortality and incident CVD in both men and women during follow-up at univariable level. These associations were nevertheless not maintained upon extensive confounder adjustment including antihypertensive therapies.

Conclusion

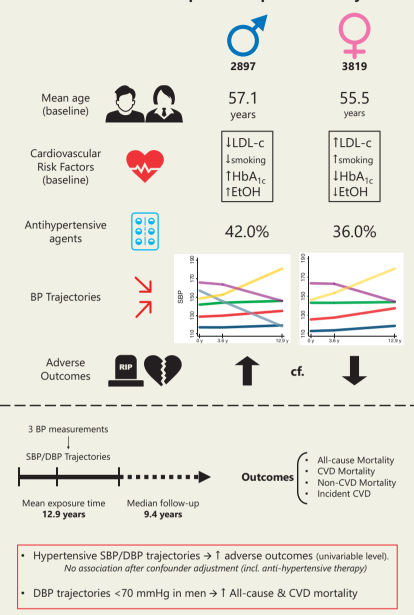
We report sex disparities in CVD prevention which may relate to worse cardiometabolic profiles and less pronounced longitudinal SBP decreases in women. Effective anti-hypertensivetherapy may offset the adverse outcomes associated with prolonged exposure to high blood pressure.

^{*} Corresponding author. Tel: +44 (0) 1224 437841, Email: phyo.myint@abdn.ac.uk

[©] The Author(s) 2021. Published by Oxford University Press on behalf of the European Society of Cardiology.

Graphical Abstract

EPIC-Norfolk Prospective Population Study



Keywords Blood pressure • Longitudinal • Trajectory • Sex-specific • Mortality • Cardiovascular disease

Introduction

While the magnitude of blood pressure (BP) elevation predicts the strength of association between hypertension and adverse outcomes, $^{1-3}$ long-term exposure to elevated BP values is also important. Long-term BP trajectories allow additional factors including antihypertensive treatment and ageing-related changes in arterial stiffness and BP 4 to be considered. Group-based trajectory modelling (GBTM) is a

data-driven approach which allows the derivation of clusters of individuals exhibiting statistically similar longitudinal trajectories of a given parameter, such as BP.⁵ Unlike approaches which define longitudinal trajectories *a priori*, GBTM does not rely on assumptions based on subjective and inflexible ex *ante* assignment rules.⁶ GBTM therefore enables the identification of new and previously unrecognized longitudinal trajectories.⁶ In this data driven approach, the clusters thus identified do not represent distinct fixed entities but rather convenient groupings

of individuals following similar trajectories. Individuals are assigned to each trajectory group based on a probability of group membership and therefore the interpretation of such trajectories depends on these considerations. ^{5,6}

Hypertensive trajectories are associated with an increased risk of incident stroke,⁷ renal disease,⁸ cardiovascular disease (CVD),^{9–11} and mortality.^{12–14} Nevertheless, significant sex differences have been demonstrated in BP trajectories.¹⁵ These may also mediate sex differences in CVD epidemiology, given recent findings of such differences in the relationship between longitudinal BP trajectories and incident atrial fibrillation (AF).¹¹ Nevertheless, no studies have previously evaluated the associations between sex-specific longitudinal BP trajectories and mortality and overall incident CVD. Furthermore, current hypertension guidelines lack sex-specific recommendations.^{1–3}

In this study, we aimed to determine the sex differences in longitudinal systolic BP (SBP) and diastolic BP (DBP) trajectories in mid-life and delineate the associations between these and mortality (all-cause, cardiovascular, and non-cardiovascular) and incident CVD in older age using data from the European Prospective Investigation into Cancer—Norfolk Cohort (EPIC-Norfolk).

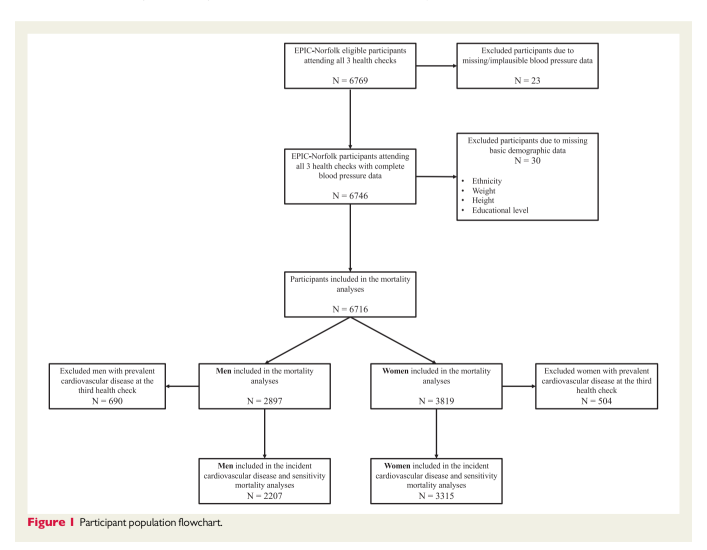
Methods

Ethical considerations

This study was conducted in accordance with the Declaration of Helsinki (1975) and later amendments. Ethical approval was obtained from the Norwich Ethics Committee. All participants gave informed signed consent for the examination of medical records and use of the data. The data that support the findings of this study are available from the corresponding author, upon reasonable request.

Data source

Participants were selected from the European Prospective Investigation into Cancer, Norfolk (EPIC-Norfolk) prospective cohort study. Study recruitment methods have been previously described. ¹⁶ In brief, men and women aged 40–79 (at study baseline) from 35 General Practices in Norfolk, England were invited to participate. Three health checks (HCs) occurred between 1993–1998 (study baseline), 1998–2000, and 2004–2012. At each HC, data on age, demographic characteristics, behavioural parameters, SBP and DBP measurements, and medication were collected. Self-reported comorbidities were ascertained during the first and second HCs. A follow-up questionnaire obtained between 2000 and 2006 ascertained self-reported comorbidities before the third HC. ¹⁷



Inclusion and exclusion criteria

Figure 1 illustrates the participant flowchart. Out of 6769 participants who attended the first three HCs of the EPIC-Norfolk study, 6716 (2897 men and 3819 women) were included in the mortality analyses, after the exclusion of 53 participants with missing or implausible blood pressure data. A total of 690 men and 504 women with prevalent CVD at the third HC (2004–2012) were further excluded from the incident CVD analyses, including 2207 men and 3315 women.

Definition of exposure, confounders, and outcomes

Outcomes

All-cause mortality was ascertained using death certificate data from the Office of National Statistics. 18 Cardiovascular mortality was ascertained using death certificate data and International Classification of Disease 10 (ICD-10) codes (I10-79) and ICD-9 codes (401-448) obtained through record linkage with the National Health Service (NHS) hospital information system and ENCORE (East Norfolk Commission Record) to allow notification of any hospital admission. Incident CVD was defined as the first date of any hospital admission/primary care diagnosis with a diagnosis comprised within the ICD-10 codes of I11-I79 and ICD-9 codes of 402-448, excluding diagnoses of essential hypertension (ICD-10: I10 and ICD-9: 401). Previously published validation studies of random samples from EPIC-Norfolk assessing the diagnoses of stroke¹⁸ and heart failure¹⁹ have shown that these parameters were ascertained with high accuracy. Furthermore, the United Kingdom National Health Service (NHS) captures almost all incident events and the EPIC-Norfolk study participants were registered with a General Practitioner and assigned an NHS number, allowing extremely robust record linkage. Therefore, the outcomes employed in our study were ascertained with high accuracy. Participants were followed up until the end of March 2018.

Exposures

Blood pressure measurements were acquired by a trained nurse after participants had been seated for 3 minutes in a quiet room. Two readings were taken using an aneroid Accutorr Sphygmomanometer (Datascope, UK) using an appropriately sized cuff with the participants' arm in the horizontal position in line with the mid-sternum.²⁰ The mean of the two readings was then recorded.

Longitudinal SBP and DBP trajectories across the first three HCs (mean exposure 12.9 years) were determined separately for men and women using Group-Based Trajectory Models (GBTM) and the *traj* Stata plugin. Trajectories were modelled using the censored normal distribution. The selection of the GBTM model with an optimal number of quadratic groups was informed by the Bayesian Information Criterion (BIC). The model with the least negative BIC which contained at most six distinct groups was chosen, ensuring that no group contained less than 1% of the considered population.

Confounders

Potential confounders considered were measured at the third HC (age, sex, ethnicity, body mass index (BMI), physical activity levels, low-density lipoprotein cholesterol, smoking status, and units of alcohol drunk) or ascertained from a follow-up questionnaire obtained before (2000–2006) the third HC (self-reported cancer, asthma, and chronic obstructive pulmonary disease). Covariates were chosen based on clinical judgement and previous literature.^{7,11,12,22–24} The estimated glomerular filtration

rate was computed using the creatinine values measured at the third HC using the Modification of Diet in Renal Disease (MDRD) formula. Comorbid CVD was defined as a self-reported diagnosis of angina, myocardial infarction, cerebrovascular disease or peripheral vascular disease on/before the follow-up questionnaire or incident CVD (ICD-10: I11-I79, ICD-9: 402-448) reported during the exposure period. Co-morbid diabetes mellitus was defined as a self-reported diagnosis of diabetes reported on/before the follow-up questionnaire, glycated haemoglobin levels >6.5% (47.5 mmol/mol) or self-reported anti-diabetic medication at the third HC.

Statistical analysis

Data were analysed using Stata 15.1 SE (StataCorp 2017, Stata Statistical Software: Release 15, College Station, TX, USA: StataCorp LLC). A 5% threshold of statistical significance was used (P < 0.05).

Missing data

Six variables collected at the third HC contained missing data. Table 1 details the proportion of missing data for these variables. Supplementary material online, Table S1 summarizes third HC characteristics stratified by whether data on any of these variables were missing. A total of 2364 (35.2%) participants had missing data on at least one variable. Those were significantly more likely to be younger and have higher incidence of adverse events (all-cause mortality, cardiovascular mortality, and incident CVD). Data missingness was likely dependent only on observed but not unobserved data, and subsequently missing-at-random.²⁶ A multiple imputation by chained equation algorithm with 20 iterations was implemented. Variables were imputed using predictive mean matching drawing from five nearest neighbours. Age, sex, ethnicity and third HC data: weight, height, educational level, SBP, DBP, heart rate, pre-existing comorbidities, medication, and two Nelson-Aalen cumulative hazard functions (incident all-cause and cardiovascular mortality) were used as predictors.

Descriptive statistics

Participant characteristics at the third HC were compared between men and women using the χ^2 test, student's *t*-test or Mann–Whitney U test as appropriate.

Cox regression analyses

For the mortality analyses, participants were followed-up from the third HC until either death or end of follow-up. Participants in the incident CVD analyses were followed-up from the third HC until either the incidence of a cardiovascular event, death or end of follow-up. Given that for this analysis we determined cause-specific hazard ratios, ²⁷ death was considered a censoring event. The median follow-up time was calculated separately for men and women using the reverse Kaplan–Meier method.

Sex-specific Cox regression models were computed for all outcomes of interest. A further sensitivity Cox regression analysis was undertaken assessing the relationship between BP trajectories and all-cause mortality only amongst participants without prevalent CVD at the third HC. The satisfaction of the proportional hazards assumption for the exposures was verified using log-negative-log plots. The BP trajectory group containing the lowest measurements was assigned as reference for all analyses. Sequentially adjusted models were constructed: Model A—Univariable; Model B—Multivariable adjustment for age and ethnicity; Model C—Model B + BMI, physical activity level, smoking and alcohol consumption;

Continued

Table I Third health check characteristics and incident outcomes of included participants from the European Prospective Investigation in Cancer (EPIC)-Norfolk (unless otherwise stated), stratified by sex

	Men 2897	Women 3819	P-value
Age, mean (SD)			
1st health check (1993–1998)	57.13 (8.02)	55.54 (7.81)	<0.001
2nd health check (1998–2000)	60.75 (8.07)	59.15 (7.87)	<0.001
3rd health check (2004–2012)	70.07 (8.25)	68.47 (8.05)	<0.001
Ethnicity, N (%)			0.359
White	2883 (99.52)	3810 (99.764)	
Black	2 (0.07)	2 (0.05)	
South Asian	3 (0.10)	2 (0.05)	
Other	9 (0.31)	5 (0.13)	
Weight (kg), mean (SD)	81.19 (12.04)	68.08 (12.44)	<0.001
Height (cm), mean (SD)	173.50 (6.58)	160.54 (6.18)	<0.001
Body mass index (kg/m²), mean (SD)	26.95 (3.58)	26.41 (4.56)	<0.001
Systolic blood pressure (mmHg), mean (SD)	,	,	
1st health check	134.70 (16.28)	129.42 (16.98)	<0.001
2nd health check	134.66 (16.57)	129.66 (17.35)	<0.001
3rd health check	136.62 (15.39)	135.94 (17.15)	0.096
Diastolic blood pressure (mmHg), mean (SD)	,	,	
1st health check	83.44 (10.57)	79.12 (10.38)	<0.001
2nd health check	83.35 (10.70)	78.92 (10.55)	<0.001
3rd health check	79.40 (9.59)	77.16 (8.98)	<0.001
Estimated glomerular filtration rate ^a (mL/min/1.73 m ²), mean (SD)	73.17 (17.40)	72.44 (20.93)	0.206
Creatinine (mmol/L), mean (SD)	93.24 (20.86)	73.29 (16.56)	<0.001
Missing, N (%)	872 (30.10)	1184 (31.00)	0.427
HbA _{1c} (%), mean (SD)	5.84 (0.67)	5.80 (0.56)	0.010
Missing, N (%)	206 (7.11)	331 (8.67)	0.020
LDL cholesterol (mmol/L), mean (SD)	2.91 (0.97)	3.36 (0.97)	<0.001
Missing, N (%)	217 (7.49)	335 (8.77)	0.058
Units of alcohol drunk, median (IQR)	6.00 (1.00–12.00)	2.00 (0.00–6.00)	<0.001
Missing, N (%)	97 (3.35)	153 (4.01)	0.158
Educational Level, N (%)	77 (3.33)	155 (1.01)	<0.001
None	635 (21.92)	1115 (29.20)	\0.00
O-level	279 (9.63)	522 (13.67)	
A-level	1384 (47.77)	1581 (41.40)	
University degree	599 (20.68)	601 (15.734)	
Physical activity level, N (%)	377 (20.00)	001 (13.734)	<0.001
Inactive	1086 (37.49)	1381 (36.16)	\0.00 1
Moderately inactive	731 (25.23)	` ,	
Moderately active	519 (17.92)	1232 (32.26)	
•	, ,	644 (16.87)	
Active	520 (17.95)	511 (13.38)	
Missing	41 (1.42)	51 (1.34)	0.024
Smoking status, N (%)	00 (2.44)	450 (444)	0.024
Yes	90 (3.11)	159 (4.16)	
No Missing	2762 (95.34)	3604 (94.37)	
Missing	45 (1.55)	56 (1.47)	
Pre-existing co-morbidities	004 (20.44)	047 (24 20)	.0.00
Cardiovascular disease, N (%)	881 (30.41)	817 (21.39)	<0.00
Diabetes mellitus, N (%)	315 (10.87)	265 (6.94)	<0.001
Cancer, N (%)	178 (6.14)	369 (9.66)	<0.001
Asthma, N (%)	247 (8.53)	390 (10.21)	0.020
Chronic obstructive pulmonary disease, N (%)	215 (7.42)	363 (9.51)	0.003

Table I Continued

	Men 2897	Women 3819	P-value
Drug therapy			
Aspirin, N (%)	758 (26.16)	500 (13.09)	<0.001
Lipid-lowering agents, N (%)	814 (28.1)	730 (19.11)	<0.001
Non-steroidal anti-inflammatory drugs, N (%)	908 (31.34)	794 (20.79)	<0.001
Anti-diabetic drugs, N (%)	168 (5.80)	103 (2.70)	<0.001
Antihypertensive agents, N (%)	1218 (42.04)	1375 (36.00)	<0.001
ACE inhibitors, N (%)	616 (21.26)	486 (12.73)	<0.001
Beta-blockers, N (%)	433 (14.95)	415 (10.87)	<0.001
Loop diuretics, N (%)	119 (4.11)	170 (4.45)	0.492
Other diuretics, N (%)	326 (11.25)	563 (14.74)	<0.001
Angiotensin receptor blockers, N (%)	174 (6.01)	264 (6.91)	0.136
Calcium channel blockers, N (%)	407 (14.05)	428 (11.21)	<0.001
Incident outcomes ^b			
Mortality, N (%)			
All-cause	595 (20.54)	453 (11.86)	<0.001
Cardiovascular	160 (5.52)	133 (3.48)	<0.001
Non-cardiovascular	435 (15.02)	320 (8.38)	<0.001
Incident cardiovascular disease, c N (%)	653 (29.59)	656 (19.79)	<0.001

Statistically significant results (P < 0.05) are highlighted in bold.

ACE, angiotensin-converting enzyme; HbA_{1c}, glycated haemoglobin; IQR, interquartile range; LDL, low-density lipoprotein; SD, standard deviation.

Model D—Model C + pre-existing comorbidities (CVD, diabetes mellitus, cancer, asthma, chronic obstructive pulmonary disease) and serum low-density lipoprotein cholesterol; Model E—Model D + antihypertensive treatment.

Results

Descriptive statistics

Table 1 summarizes the participant sex-specific characteristics at the third HC. A total of 2897 (43.1%) men and 3819 (56.9%) women were included. Compared to men [mean age (standard deviation) = 70.07 (8.25) years], women were younger [68.47 (8.05) years]. SBP and DBP measurements were higher amongst men than women. Compared to men, women had lower rates of self-reported comorbid CVD, diabetes mellitus, cancer, asthma, and chronic obstructive pulmonary disease on or before the third HC. Women also had lower rates of self-reported therapy with aspirin, lipid-lowering agents, anti-diabetic, and anti-hypertensive agents. Over the follow-up period, men had higher rates of incident mortality (all-cause, cardiovascular, and non-cardiovascular) as well as incident CVD.

Longitudinal blood pressure trajectories

There were six distinct SBP trajectories amongst men included in the mortality analyses (*Figure 2* and Supplementary material online, *Table S2*), which were characterized according to the 2018 ESC guidelines. A total of 14.7% belonged to trajectory 1 (stable optimal SBP), 45.4% to trajectory 2 (stable normal/high normal SBP), 29.9% to trajectory

3 (stable grade 1 hypertension), 2.2% to trajectory 4 (well-controlled grade 1 hypertension), 1.2% to trajectory 5 (grade 1 \rightarrow grade 2 hypertension) and 6.6% to trajectory 6 (grade 2 \rightarrow grade 1 hypertension). Amongst women included in the mortality analyses, five SBP trajectories were revealed: 23.6% belonged to trajectory 1 (stable optimal SBP), 44.2% to trajectory 2 (rising normal/high normal SBP), 26.8% to trajectory 3 (stable grade 1 hypertension), 3.0% to trajectory 4 (grade 2 \rightarrow grade 1 hypertension), and 2.4% to trajectory 5 (grade 1 \rightarrow grade 2 hypertension), respectively.

Amongst men included in the incident CVD analyses, 5 SBP trajectories were revealed (*Figure 3* and Supplementary material online, *Table S3*): 19.4% belonged to trajectory 1 (borderline stable optimal SBP), 50.2% to trajectory 2 (rising normal/high normal SBP), 21.3% to trajectory 3 (rising grade 1 hypertension), 4.1% to trajectory 4 (grade 1 hypertension \rightarrow high-normal SBP), and 5.0% to trajectory 5 (grade 2 \rightarrow grade 1 hypertension). Amongst women included in the incident CVD analyses, there were five similar SBP trajectories: 12.6% belonged to trajectory 1 (stable optimal SBP), 37.8% to trajectory 2 (rising normal/high-normal SBP), 37.2% to trajectory 3 (high-normal SBP \rightarrow grade 1 hypertension), 2.4% to trajectory 4 (grade 1 \rightarrow grade 2 hypertension), and 9.9% to trajectory 5 (decreasing grade 1 hypertension) and

All analysed groups revealed five distinct DBP trajectories, which were similar in all groups: trajectory 1 (low optimal DBP), trajectory 2 (high optimal DBP), trajectory 3 (normal/high-normal DBP), trajectory 4 (grade 1 hypertension \rightarrow normal DBP), and trajectory 5 (stable grade 1/grade 2 hypertension).

^aEstimated glomerular filtration rate was calculated using the *Modification of Diet in Renal Disease* formula.²⁵

blncident outcomes measured during the follow-up period from the third health check (2004–2012) until the end of March 2018, resulting in a median follow-up of 9.44 years.

funcident cardiovascular disorders reported only amongst patients without pre-existing cardiovascular disease at the third health check (N = 2207 men; 3315 women).

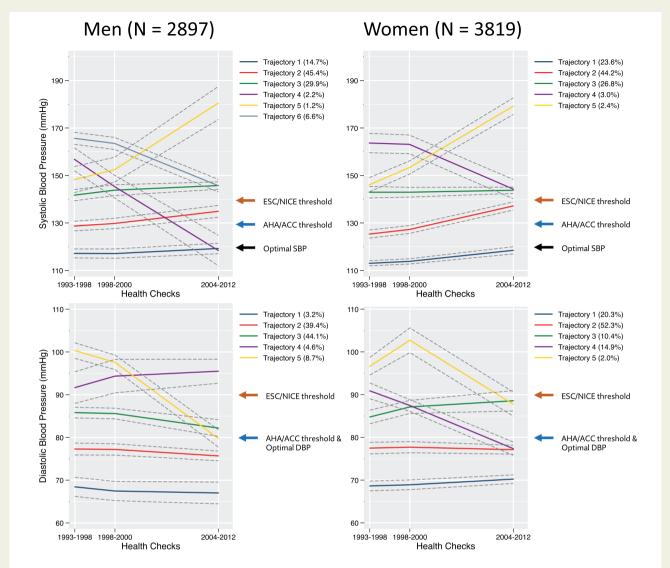


Figure 2 Blood pressure trajectories amongst men (n = 2897) and women (n = 3819) included in the mortality analyses. SBP trajectories in men: Trajectory 1—stable optimal SBP, Trajectory 2—stable normal/high normal SBP, Trajectory 3—stable grade 1 hypertension, Trajectory 4—well-controlled grade 1 hypertension, Trajectory 5—grade 1 → grade 2 hypertension, Trajectory 6—grade 2 → grade 1 hypertension. SBP trajectories in women: Trajectory 1—stable optimal SBP, Trajectory 2—rising normal/high normal SBP, Trajectory 3—stable grade 1 hypertension, Trajectory 4—grade 2 → grade 1 hypertension, Trajectory 5—grade 1 → grade 2 hypertension. DBP trajectories: Trajectory 1—low optimal DBP, Trajectory 2—high optimal DBP, Trajectory 3—normal/high-normal DBP, Trajectory 4—grade 1 hypertension → normal DBP, Trajectory 5—stable grade 1/grade 2 hypertension. 95% confidence intervals are represented as dashed grey lines.

Cox regression analyses

Median follow-up periods (interquartile range) were 9.4 (8.0–10.9) and 9.4 (7.8–11.0) years for men and women, respectively. Univariable analyses (Model A) showed that compared to SBP trajectory 1 participants (stable optimal SBP), the other SBP trajectories were associated with increased risk of incident all-cause, cardiovascular, and non-cardiovascular mortality amongst both sexes (*Figure 4* and Supplementary material online, *Tables S4*–S6). In men, trajectories 3–6 were associated with two-fold increases in the risk of all-cause mortality, while trajectories 2–6 were associated with up to three-fold increases for the same outcome (*Figure 4*). Nevertheless,

these associations with any of the mortality outcomes were not revealed upon confounder adjustment (Models B–E). In men, similar patterns were revealed between DBP trajectories and incident mortality. In univariable analyses (Model A), trajectories 2–4 were associated with 35–50% reductions in the risk of incident all-cause mortality compared to trajectory 1 (low optimal DBP), which were also not revealed upon confounder adjustment (Models B–E). Nevertheless, there were a few significant associations between DBP trajectories and cardiovascular mortality amongst men. Compared to trajectory 1 (low optimal DBP), trajectories 2 (high optimal DBP), and 3 (normal/high-normal DBP) were associated with significantly

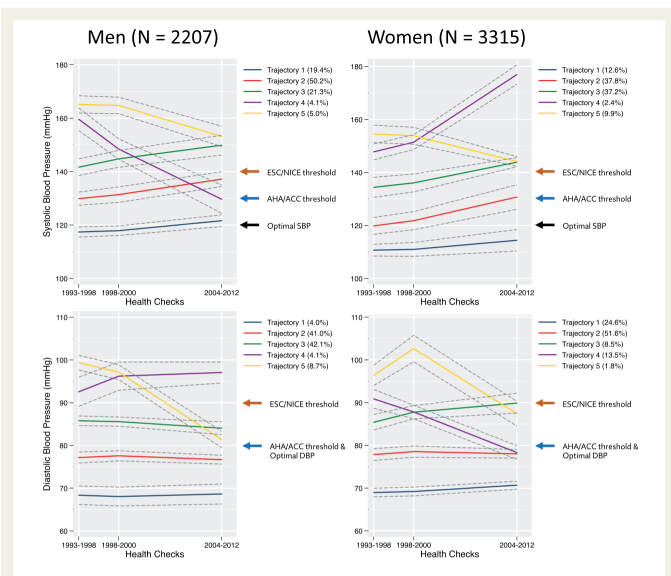


Figure 3 Long-term blood pressure patterns amongst men (n = 2207) and women (n = 3315) participants included in the incident cardiovascular disease analyses. SBP trajectories in men: Trajectory 1—borderline stable optimal SBP, Trajectory 2—rising normal/high normal SBP, Trajectory 3—rising grade 1 hypertension, Trajectory 4—grade 1 hypertension \rightarrow high-normal SBP, Trajectory 5—grade 2 \rightarrow grade 1 hypertension. SBP trajectories in women: Trajectory 1—stable optimal SBP, Trajectory 2—rising normal/high normal SBP, Trajectory 3—high-normal SBP \rightarrow grade 1 hypertension, Trajectory 4—decreasing grade 1 hypertension, Trajectory 5—grade 1 \rightarrow grade 2 hypertension. DBP trajectories: Trajectory 1—low optimal DBP, Trajectory 2—high optimal DBP, Trajectory 3—normal/high-normal DBP, Trajectory 4—grade 1 hypertension \rightarrow normal DBP, Trajectory 5—stable grade 1/grade 2 hypertension. 95% confidence intervals are represented as dashed grey lines.

lower cardiovascular mortality in all models considered (Models A–E).

The sensitivity all-cause mortality analysis considering only participants without prevalent CVD at the third health checked revealed results consistent with the main analyses (Supplementary material online, *Table S7*).

Amongst men, compared to SBP trajectory 1 (stable optimal SBP), trajectories 2–5 were associated with significant increases between 30% and three-fold in incident CVD in the univariable model (Model A). The associations between trajectories 2–4 and incident CVD were not revealed after confounder adjustment (Models B–E).

Nevertheless, the association between trajectory 5 (grade $2 \rightarrow$ grade 1 hypertension) was revealed upon adjustment for age, ethnicity, BMI, physical activity level, smoking, and alcohol consumption and pre-existing comorbidities (Models B–D), but not after further adjustment for antihypertensive treatment (Model E) (*Figure 5*, Supplementary material online, *Table S8*). There were no statistically significant relationships between DBP trajectories 2-5 (compared to trajectory 1) and incident CVD amongst men across all fully adjusted models considered.

Amongst women, compared to SBP trajectory 1 (stable optimal SBP), trajectories 3–5 were associated with significant two- to three-

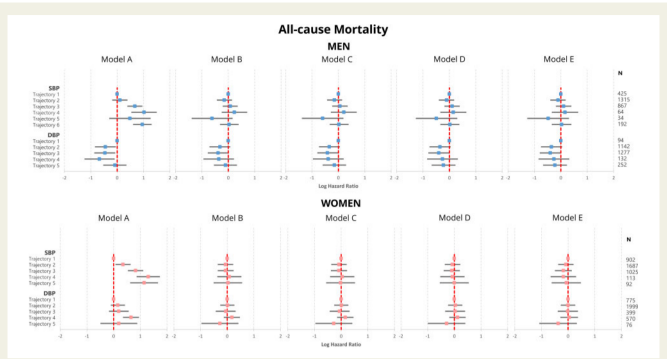


Figure 4 Results of Cox regressions assessing the association between blood pressure phenotypes and all-cause mortality. Model A—Univariable; Model B—Multivariable adjustment for age and ethnicity; Model C—Model B + body mass index, physical activity level, smoking and alcohol consumption; Model D—Model C + pre-existing co-morbidities (cardiovascular disease, diabetes mellitus, cancer, asthma, chronic obstructive pulmonary disease) and serum low-density lipoprotein cholesterol; Model E—Model D + antihypertensive treatment. SBP trajectories in men: Trajectory 1—borderline stable optimal SBP, Trajectory 2—rising normal/high normal SBP, Trajectory 3—rising grade 1 hypertension, Trajectory 4—grade 1 hypertension \rightarrow high-normal SBP, Trajectory 5—grade 2 \rightarrow grade 1 hypertension. SBP trajectories in women: Trajectory 1—stable optimal SBP, Trajectory 2—rising normal/high normal SBP, Trajectory 3—high-normal SBP \rightarrow grade 1 hypertension, Trajectory 4—decreasing grade 1 hypertension, Trajectory 5—grade 1 \rightarrow grade 2 hypertension. DBP trajectories: Trajectory 1—low optimal DBP, Trajectory 2—high optimal DBP, Trajectory 3—normal/high-normal DBP, Trajectory 4—grade 1 hypertension \rightarrow normal DBP, Trajectory 5—stable grade 1/grade 2 hypertension. Median (interquartile range) follow-up was 9.4 (8.0–10.9) years amongst both men and women, respectively. DBP, diastolic blood pressure; SBP, systolic blood pressure.

fold increases in the risk of incident CVD at univariable level (Model A). The associations between trajectories 2-4 and incident CVD were not revealed upon confounder adjustment (Models B-E). Nevertheless, the association between trajectory 5 (decreasing grade 1 hypertension) and incident CVD was revealed upon adjustment for age and ethnicity (Model B), but not after further confounder adjustment (Models C-E). Compared to DBP trajectory 1 (low optimal DBP), trajectories 2-5 were associated with significant 30% to twofold increases in the risk of incident CVD amongst women. The associations between trajectories 2, 3, and 5 and incident CVD were not revealed upon confounder adjustment (Models B-E), while the association between trajectory 4 (grade 1 hypertension → normal DBP) and incident CVD was revealed upon adjustment for age, ethnicity, BMI, physical activity and smoking, alcohol consumption (Models B-C) but not after further adjustment for pre-existing comorbidities and antihypertensive treatment (Models D-E).

Discussion

In this prospective cohort study with long-term historical BP data spanning \sim 13 years, we determined the sex differences in

longitudinal BP trajectories in mid-life and have delineated their associations with mortality (all-cause, cardiovascular, and noncardiovascular) and incident CVD in later life. We found that women were significantly younger, but demonstrated a worse cardiometabolic profile at baseline with higher baseline LDL-c levels, lower levels of physical activity, and higher prevalence of smoking. Women were also less likely to receive primary CVD prevention, antihypertensive, and anti-diabetic treatment. They nevertheless tended to have SBP trajectories characterized by lower mean measurements, while men tended to exhibit more pronounced decreases in SBP over the duration of the exposure period. Despite overall lower mortality and incident CVD rates amongst women, the excess risk of these adverse outcomes associated with hypertensive trajectories was higher in women than in men at univariable level. While the univariable associations between most longitudinal BP trajectories and mortality were rendered non-significant upon only age and ethnicity adjustment amongst both sexes, the association between hypertensive BP trajectories and incident mortality amongst men was maintained after comprehensive adjustment for age, ethnicity, lifestyle factors, comorbidities, and baseline LDL-c levels. Further adjustment for antihypertensive treatment rendered this association non-significant,

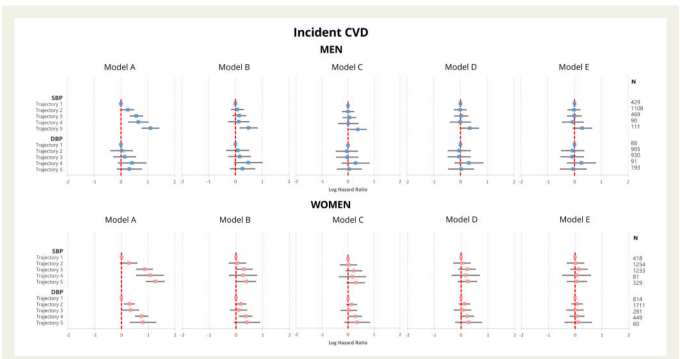


Figure 5 Results of Cox regressions assessing the association between blood pressure phenotypes and incident cardiovascular disease amongst participants without prevalent cardiovascular disease at the third health check of the EPIC-Norfolk study. Model A—Univariable; Model B—Multivariable adjustment for age and ethnicity; Model C—Model B + body mass index, physical activity level, smoking and alcohol consumption; Model D—Model C + pre-existing co-morbidities (cardiovascular disease, diabetes mellitus, cancer, asthma, chronic obstructive pulmonary disease) and serum low-density lipoprotein cholesterol; Model E—Model D + antihypertensive treatment. SBP trajectories in men: Trajectory 1—borderline stable optimal SBP, Trajectory 2—rising normal/high normal SBP, Trajectory 3—rising grade 1 hypertension, Trajectory 4—grade 1 hypertension \rightarrow high-normal SBP, Trajectory 5—grade 2 \rightarrow grade 1 hypertension. SBP trajectories in women: Trajectory 1—stable optimal SBP, Trajectory 2—rising normal/high normal SBP, Trajectory 3—high-normal SBP \rightarrow grade 1 hypertension, Trajectory 4—decreasing grade 1 hypertension, Trajectory 5—grade 1 \rightarrow grade 2 hypertension. DBP trajectories: Trajectory 1—low optimal DBP, Trajectory 2—high optimal DBP, Trajectory 3—normal/high-normal DBP, Trajectory 4—grade 1 hypertension \rightarrow normal DBP, Trajectory 5—stable grade 1/grade 2 hypertension. Median (interquartile range) follow-up was 9.4 (8.0–10.9) and 9.4 (7.8–11.0) years amongst men and women, respectively. DBP, diastolic blood pressure; SBP, systolic blood pressure.

suggesting that the long-term adverse effects of hypertension may be offset by appropriate and timely antihypertensive treatment.

Several previous investigations assessed the relationship between long-term BP burden and CVD. A dose-response relationship between cumulative exposure to elevated BP and incident CVD and cardiovascular mortality has been previously reported. 13,28-31 Furthermore, several investigations have reported that prolonged exposure to elevated BP trajectories is associated with increased atherosclerotic burden,²² incident stroke,⁷ heart failure,⁹ AF,¹¹ overall CVD, ⁹ all-cause mortality, ²⁴ and cardiovascular mortality. ¹² Recent findings indicate that long-term BP trajectories may differ by sex. 15 Indeed, sex differences have also been recently reported for incident AF. 11 Amongst 7670 men and 8376 women from a Norwegian prospective cohort with a mean age at the beginning of the exposure period of \sim 40 years (mean age of our cohort at the beginning of the exposure period was 56.2 years), stronger associations between elevated/hypertensive BP trajectories and incident AF were documented amongst women than in men.¹¹ However, no previous investigations have assessed the relationship between the different phenotypes of longitudinal BP changes and adverse outcomes separately amongst men and women. Our study is the first to report these relationships.

Our results suggest that men exhibit higher SBP trajectories than women, with men in the reference SBP trajectory having a mean SBP of \sim 120 mmHg throughout the exposure period while women in the reference group ~110 mmHg. Nevertheless, more pronounced SBP decreases were recorded in men. This may be attributed to a tendency of undertreating hypertension in women, illustrated by lower utilization of antihypertensive agents in women than in men at the end of the exposure period (36% vs. 42%, respectively). Our results are largely consistent with previous findings suggesting a larger unused potential for cardiovascular prevention by BP reduction strategies in women than in men.³² Furthermore, we also found that compared to low optimal DBP (<70 mmHg), DBP trajectories characterized by high normal values (80-90 mmHg) were associated with lower risk of all-cause and cardiovascular mortality in men, even after full multivariable adjustment. Such associations were nevertheless not revealed in women (who had lower CVD prevalence at baseline) or in participants included in the incident CVD analyses (who were free of prevalent CVD at baseline), suggesting that long-term exposure DBP <70 mmHg may be deleterious in patients with preexisting CVD, which has been previously reported in larger studies with shorter follow-up. 33,34

The overall lack of association between the other BP trajectories and outcomes may be related to the characteristics of the included participant sample: mean BMI \sim 26.5–26.9 kg/m², estimated glomerular filtration rate ~73 mL/min/1.73 m², glycated haemoglobin levels \sim 5.8% (39.9 mmol/mol), LDL cholesterol levels \sim 3.0 mmol/L, \sim 95% non-smokers. Furthermore, a high proportion of participants were undergoing CVD primary prevention with aspirin, lipid-lowering, and antihypertensive agents at the end of the exposure period. The included sample thus comprised of relatively healthy participants undergoing appropriate primary prevention in whom incident CVD would be less likely to result in a fatal event. This may also be explained by survivorship bias, in which the inclusion of a healthier sub-population surviving over a ~13-year-old period spanning the first three HCs of the EPIC-Norfolk study may have led to the underestimation of the mortality and incident CVD risk. Further studies replicating our analytic strategy are therefore warranted to determine the same associations in other cohorts with differing distributions of ethnicity, cardiovascular risk profile and comorbidities to ensure the external validity of our findings.

Our results may inform primary CVD prevention practice by highlighting the importance of sex differences in the natural course and management of hypertension from mid-life onwards. While women exhibited BP trajectories characterized by initial lower values, men received more aggressive antihypertensive therapy, resulting in more pronounced longitudinal BP decreases. Suboptimal primary CVD prevention amongst women is also reflected in poorer cardiometabolic health at baseline and higher relative risk increases associated with exposure to non-optimal BP values in women at univariable level. These results therefore reflect important between-sex disparities in primary prevention which need to be addressed in order to ensure appropriate and fair provision of care. Therefore, a programme of systematic population screening for hypertension with a special emphasis on early diagnosis in women should be implemented in order to address the sex disparities highlighted by our study. Patient education and regular follow-up are also be warranted to ensure appropriate compliance with antihypertensive therapy. Furthermore, the results of our study in addition to previous literature^{33,34} highlight that lowering DBP to values <70 mmHg may be deleterious in men as well as patients with pre-existing CVD and such stringent BP treatment targets should be avoided in these populations.

Our study benefited from several strengths. We used data from EPIC-Norfolk, a large, prospective cohort with robust ascertainment of exposures, confounders and outcomes. Furthermore, EPIC-Norfolk benefits from long-term follow-up, allowing the determination of long-term BP trajectories over 12 years and the adjudication of outcomes over a median follow-up of 9.4 years after the exposure period. We were able to control for wide range of important demographic, lifestyle, social and medical factors. Furthermore, we employed robust statistical methods to adjudicate longitudinal BP trajectory group membership.

Naturally, there were also limitations. We included >99% White Caucasian participants, and thus ethnicity-stratified analyses could

not be performed. Further studies are required to determine these associations amongst other ethnicities. BP trajectories were adjudicated based on measurements from three HCs. Nevertheless, no other BP measurements were acquired and therefore BP variations occurring between these HCs were not considered, introducing potential uncertainty regarding the development of BP trajectories between the HCs. Furthermore, we used self-reported comorbidities, which may lead to inaccuracies. We nevertheless employed a combined comorbidity definition also employing medication data, biomarkers measured at the HCs and incident diagnoses reported during the exposure period in order to minimize potential inaccuracies in the ascertainment of self-reported comorbidities. Furthermore, having only included a subgroup of the EPIC-Norfolk prospective study which attended the first three HCs of the study spanning \sim 13 years, our analyses may be prone to survivorship bias. Nevertheless, this is an inherent limitation of any study analysing longitudinal changes in BP. Despite adjusting for a wide range of participant characteristics at the beginning of the follow-up, data describing the evolution of these co-variates over the follow-up were not available, which did not allow longitudinal co-variate changes to be considered. As an observational study, we were not able to account for residual confounders.

In conclusion, using data from a large-scale prospective cohort study, we determined the sex differences in longitudinal BP trajectories in midlife and delineated the associations between these and mortality and incident CVD in older age. While men exhibited higher BP longitudinal trajectories than women, these were characterized by more pronounced BP decreases throughout the exposure period of the study, which may be attributed to the relative undertreatment of women. This suggests important between-sex disparities in primary CVD prevention. Exposure to hypertensive SBP and DBP trajectories was associated with higher mortality and incident CVD risk amongst both sexes at univariable level. Nevertheless, these associations were not maintained upon extensive confounder adjustment including antihypertensive therapy, suggesting that effective therapy may offset the adverse outcomes associated with prolonged exposure to high BP. Our results also highlight that longitudinal exposure to low DBP values <70 mmHg may be independently associated with higher risk of all-cause and cardiovascular mortality amongst men.

Supplementary material

Supplementary material is available at European Journal of Preventive Cardiology online.

Acknowledgements

We are grateful to all the participants who have been part of the project and to the many members of the study teams at the University of Cambridge who have enabled this research. We would also like to acknowledge the principal investigators and staff of the EPIC-Norfolk study.

Funding

The EPIC-Norfolk study (doi:10.22025/2019.10.105.00004) has received funding from the Medical Research Council (MR/N003284/1 and MC-UU_12015/1) and Cancer Research UK (C864/A14136).

Conflict of interest: none declared.

References

- Williams B, Mancia G, Spiering W, Agabiti Rosei E, Azizi M, Burnier M, Clement DL, Coca A, de Simone G, Dominiczak A, Kahan T, Mahfoud F, Redon J, Ruilope L, Zanchetti A, Kerins M, Kjeldsen SE, Kreutz R, Laurent S, Lip GYH, McManus R, Narkiewicz K, Ruschitzka F, Schmieder RE, Shlyakhto E, Tsiouffis C, Aboyans V, Desormais I; ESC Scientific Document Group. 2018 ESC/ESH Guidelines for the management of arterial hypertension. Eur Heart J 2018:39:3021–3104.
- National Institute for Health and Care Excellence. Overview | Hypertension in adults: diagnosis and management | Guidance | NICE, NICE; 2019.
- 3. Whelton PK, Carey RM, Aronow WS, Casey DEJ, Collins KJ, Dennison HC, DePalma SM, Gidding S, Jamerson KA, Jones DW, MacLaughlin EJ, Muntner P, Ovbiagele B, Smith SCJ, Spencer CC, Stafford RS, Taler SJ, Thomas RJ, Williams KAS, Williamson JD, Wright JTJ. 2017 ACC/AHA/AAPA/ABC/ACPM/AGS/APA/ASH/ASPC/NMA/PCNA Guideline for the prevention, detection, evaluation, and management of high blood pressure in adults: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice guidelines. Hypertension 2018;71:e13—e115.
- 4. Sun Z. Aging, arterial stiffness, and hypertension. *Hypertension* 2015;**65**:252–256.
- Shearer DM, Thomson WM, Broadbent JM, McLean R, Poulton R, Mann J. Highrisk glycated hemoglobin trajectories established by mid-20s: findings from a birth cohort study. BMJ Open Diabetes Res Care 2016;4:e000243.
- Nagin DS, Odgers CL. Group-based trajectory modeling in clinical research. Annu Rev Clin Psychol 2010;6:109–138.
- Portegies MLP, Mirza SS, Verlinden VJA, Hofman A, Koudstaal PJ, Swanson SA, Ikram MA. Mid- to late-life trajectories of blood pressure and the risk of stroke: the Rotterdam Study. *Hypertension* 2016;67:1126–1132.
- Zheng W, Mu J, Chu C, Hu J, Yan Y, Ma Q, Lv Y, Xu X, Wang K, Wang Y, Deng Y, Yan B, Yang R, Yang J, Ren Y, Yuan Z. Association of blood pressure trajectories in early life with subclinical renal damage in middle age. J Am Soc Nephrol 2018:29:2835–2846.
- Smitson CC, Scherzer R, Shlipak MG, Psaty BM, Newman AB, Sarnak MJ, Odden MC, Peralta CA. Association of blood pressure trajectory with mortality, incident cardiovascular disease, and heart failure in the cardiovascular health study. Am J Hypertens 2017;30:587–593.
- Lee C-L, Wang J-S. Systolic blood pressure trajectory and cardiovascular outcomes: an analysis using data in the Systolic Blood Pressure Intervention Trial. Int J Clin Pract 2020;74:e13450.
- Sharashova E, Wilsgaard T, Ball J, Morseth B, Gerdts E, Hopstock LA, Mathiesen EB, Schirmer H, Løchen M-L. Long-term blood pressure trajectories and incident atrial fibrillation in women and men: the Tromsø Study. Eur Heart J 2020;41: 1554–1562.
- Tielemans SMAJ, Geleijnse JM, Laughlin GA, Boshuizen HC, Barrett-Connor E, Kromhout D. Blood pressure trajectories in relation to cardiovascular mortality: The Rancho Bernardo Study. J Hum Hypertens 2017;31:515–519.
- Zhang Y, Vittinghoff E, Pletcher MJ, Allen NB, Zeki Al Hazzouri A, Yaffe K, Balte PP, Alonso A, Newman AB, Ives DG, Rana JS, Lloyd-Jones D, Vasan RS, Bibbins-Domingo K, Gooding HC, de Ferranti SD, Oelsner EC, Moran AE. Associations of blood pressure and cholesterol levels during young adulthood with later cardiovascular events. J Am Coll Cardiol 2019;74:330–341.
- Ravindrarajah R, Hazra NC, Hamada S, Charlton J, Jackson SHD, Dregan A, Gulliford MC. Systolic blood pressure trajectory, frailty, and all-cause mortality >80 years of age: cohort study using electronic health records. *Circulation* 2017; 135:2357–2368.
- Ji H, Kim A, Ebinger JE, Niiranen TJ, Claggett BL, Bairey Merz CN, Cheng S. Sex differences in blood pressure trajectories over the life course. JAMA Cardiol 2020; 5:255–226.
- Day N, Oakes S, Luben R, Khaw KT, Bingham S, Welch A, Wareham N. EPIC-Norfolk: study design and characteristics of the cohort. European Prospective Investigation of Cancer. Br J Cancer England 1999;80(Suppl 1):95–103.

 EPIC-Norfolk. About EPIC-Norfolk: Timeline. https://www.epic-norfolk.org.uk/ about-epic-norfolk/timeline/. (07 June 2021).

- Sinha S, Myint PK, Luben RN, Khaw K-T. Accuracy of death certification and hospital record linkage for identification of incident stroke. BMC Med Res Methodol 2008:8:74.
- Pfister R, Michels G, Wilfred J, Luben R, Wareham NJ, Khaw K-T. Does ICD-10 hospital discharge code I50 identify people with heart failure? A validation study within the EPIC-Norfolk study. Int. J. Cardiol. Netherlands 2013;4413– 4414
- Bain LKM, Myint PK, Jennings A, Lentjes MAH, Luben RN, Khaw K-T, Wareham NJ, Welch AA. The relationship between dietary magnesium intake, stroke and its major risk factors, blood pressure and cholesterol, in the EPIC-Norfolk cohort. Int J Cardiol 2015;196:108–114.
- Jones BL, Nagin DS. A note on a stata plugin for estimating group-based trajectory models. Sociol Methods Res 2013;42:608–613.
- Allen NB, Siddique J, Wilkins JT, Shay C, Lewis CE, Goff DC, Jacobs DRJ, Liu K, Lloyd-Jones D. Blood pressure trajectories in early adulthood and subclinical atherosclerosis in middle age. JAMA 2014;311:490–497.
- Tielemans SMAJ, Geleijnse JM, Menotti A, Boshuizen HC, Soedamah-Muthu SS, Jacobs DRJ, Blackburn H, Kromhout D. Ten-year blood pressure trajectories, cardiovascular mortality, and life years lost in 2 extinction cohorts: the Minnesota Business and Professional Men Study and the Zutphen Study. J Am Heart Assoc 2015;4:e001378.
- Wang A, Zhang Y, Li J, Zhao Q, Cao Y, Li J, Zhang R, Chen S, Gao J, Wu S. High SBP trajectories are associated with risk of all-cause death in general Chinese population. J Hypertens 2018;36:1299–1305.
- Levey AS, Bosch JP, Lewis JB, Greene T, Rogers N, Roth D. A more accurate method to estimate glomerular filtration rate from serum creatinine: a new prediction equation. Modification of Diet in Renal Disease Study Group. Ann Intern Med 1999;130:461–470.
- 26. White IR, Royston P, Wood AM. Multiple imputation using chained equations: issues and guidance for practice. *Stat Med* 2011;**30**:377–399.
- Noordzij M, Leffondre K, van Stralen KJ, Zoccali C, Dekker FW, Jager KJ. When do we need competing risks methods for survival analysis in nephrology? Nephrol Dial Transplant 2013;28:2670–2677.
- Allen N, Berry JD, Ning H, Van Horn L, Dyer A, Lloyd-Jones DM. Impact of blood pressure and blood pressure change during middle age on the remaining lifetime risk for cardiovascular disease: the cardiovascular lifetime risk pooling project. *Circulation* 2012;**125**:37–44.
- Pool LR, Ning H, Wilkins J, Lloyd-Jones DM, Allen NB. Use of long-term cumulative blood pressure in cardiovascular risk prediction models. JAMA Cardiol 2018; 3:1096–1100
- 30. Satoh M, Ohkubo T, Asayama K, Murakami Y, Sugiyama D, Yamada M, Saitoh S, Sakata K, Irie F, Sairenchi T, Ishikawa S, Kiyama M, Ohnishi H, Miura K, Imai Y, Ueshima H, Okamura T; EPOCH-JAPAN Research Group. Lifetime risk of stroke and coronary heart disease deaths according to blood pressure level: EPOCH-JAPAN (Evidence for Cardiovascular Prevention From Observational Cohorts in Japan). Hypertension 2019;73:52–59.
- Teramoto K, Nadruz Junior W, Matsushita K, Claggett B, John JE, Skali H, Solomon S, Cheng S, Shah AM. Mid- to late-life time-averaged cumulative blood pressure and late-life cardiac structure, function, and heart failure. *Hypertension* 2020;**76**:808–818.
- 32. Boggia J, Thijs L, Hansen TW, Li Y, Kikuya M, Björklund-Bodegård K, Richart T, Ohkubo T, Jeppesen J, Torp-Pedersen C, Dolan E, Kuznetsova T, Olszanecka A, Tikhonoff V, Malyutina S, Casiglia E, Nikitin Y, Lind L, Maestre G, Sandoya E, Kawecka-Jaszcz K, Imai Y, Wang J, Ibsen H, O'Brien E, Staessen JA. Ambulatory blood pressure monitoring in 9357 subjects from 11 populations highlights missed opportunities for cardiovascular prevention in women. *Hypertension* 2011;**57**:397–405.
- 33. Vidal-Petiot E, Ford I, Greenlaw N, Ferrari R, Fox KM, Tardif J-C, Tendera M, Tavazzi L, Bhatt DL, Steg PG. Cardiovascular event rates and mortality according to achieved systolic and diastolic blood pressure in patients with stable coronary artery disease: an international cohort study. Lancet 2016;388: 2142–2152.
- 34. Böhm M, Schumacher H, Teo KK, Lonn EM, Mahfoud F, Mann JFE, Mancia G, Redon J, Schmieder RE, Sliwa K, Weber MA, Williams B, Yusuf S. Achieved blood pressure and cardiovascular outcomes in high-risk patients: results from ONTARGET and TRANSCEND trials. *Lancet* 2017;389:2226–2237.