

Supplementary Material. Systematic review: associations of testosterone with men's health.

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29 **2. Additional details on systematic searches and screening**

30
31 We used online search tools to identify available published (MEDLINE, EMBASE) and grey
32 literature (OpenGrey, Mednar) *items* (journal articles, reports, theses, webpage articles)
33 reporting on suitable prospective cohort *studies* (the underlying unique sources of data). We
34 used OpenGrey and Mednar because both were free search tools that we considered likely to
35 identify additional grey literature items and studies in an expanded search beyond the
36 mainstream publications. Mednar is a medically-focussed search engine of public and deep
37 web resources, excluding subscription services.[1] OpenGrey is a searchable database
38 containing citations for items including technical or research reports, theses, conference
39 papers, and other types of grey literature.[2] Literature searches were conducted on 18-22
40 July 2019, with no date restrictions set.

41
42 Where possible (as functionality varied among the different tools), we placed the following
43 restrictions on the search: items reporting on the results of a research study, longitudinal or
44 prospective cohort studies, not of hormone therapy or deprivation treatments. Due to study
45 timeframe and language translation limitations, we opted to search for only those items that
46 were reported in the English language. The terms and full criteria used for the MEDLINE
47 search are provided in Supplementary table 1, and the PRISMA checklist as Supplementary
48 table 5.

49
50 Selection criteria were set as applicable to the planned sets of IPD meta-analyses
51 (Supplementary table 2).[3] Only items reporting on prospective population-based cohort
52 studies, adults of combined sexes or of men alone, with individuals free of the disease at
53 baseline, were sought. Items reporting a different design for the analysis of longitudinal data,
54 such as nested case-control or case-cohort design, were also considered acceptable. A

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55 minimum of five years follow-up was selected, to ensure a sufficient number of incident
56 events for statistical modelling. We excluded items that did not measure testosterone using
57 mass spectrometry, which is regarded to be the 'gold standard' method,[4] although
58 testosterone was not required to be mentioned in the title or abstract, nor modelled as the
59 primary exposure variable. Selected items were to be studies of humans, reported in English,
60 and reporting on analyses of at least one of the AIMS outcomes.

61

62 Two reviewers (RJM, JH) independently screened the de-duplicated items against these pre-
63 specified criteria. To optimise efficiency, the selection of items proceeded in two steps. Title
64 and abstract screenings (Step 1) were followed by full text screening of items selected in Step
65 1 (Step 2). If an item was selected for exclusion, then the main reason for that decision was
66 recorded. If there was uncertainty in the decision to exclude, in Step 1 the reviewer selected
67 "include" (in Step 1) or "maybe" (in Step 2). At the end of each step, the two reviewers
68 sought to achieve consensus, through discussion, for each item that did not achieve
69 agreement. Exclusion reasons were used to inform discussions for achieving consensus. Items
70 with a consensus decision of "maybe" were further investigated by Reviewer 1 (RJM) using
71 information external to the systematic searches and screenings (reading further details of
72 methods used in cited articles, and from correspondence with authors or other researchers
73 currently working on the research study).

74

75 This screening procedure was adjusted to accommodate the different types of items reviewed
76 (published articles, theses, webpage articles, unpublished reports; Supplementary table 3). A
77 pilot set of title-only screenings for 30 randomly chosen articles suggested that sufficient

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78 information was contained within the titles alone for the purpose of Step 1 screenings.^a
79 Therefore, in cases when an abstract was not available, only the titles were screened. Website
80 items identified by the Mednar search tool were the type of item that most often did not have
81 abstract or summary text, and in these cases the webpage text was reviewed in place of an
82 abstract (Supplementary table 3).
83
84 Endnote X8[5] was used for collating and storing the citations returned from literature
85 searches, and for de-duplicating and storing the selected references. The full citations,
86 including abstracts, were exported from Endnote for uploading into Rayyan[6], which is a
87 free web tool that was used for screening, recording exclusion decisions, and downloading
88 selection results.
89
90 The literature search identified 2,177 items (1,738 published and 439 from grey literature),
91 with 1,994 items remaining after duplicates had been removed, and after excluding two
92 Mednar items that had insufficient information available to review (Fig. 1). Supplementary
93 table 4 shows the frequencies of returned items by search terms present in the titles and
94 abstracts. Most (72.7%) had the word “cancer”, and 1,107 (55.5%) of these had the word
95 “prostate cancer”, in the title or abstract. This, combined with frequent mentions of
96 “androgen deprivation” (29.2%), “radiotherapy” (18.6%), and “brachytherapy” (8.3%), show
97 that items reporting aspects of testosterone deprivation or suppression for treating prostate
98 cancer were a predominant feature of the returned items. Different types of returned items
99 included 1,764 published articles, 111 webpage articles, 81 theses, and 38 unpublished
100 reports/other documents, and the percentages without abstract or webpage text screened in

^a 30 titles were initially screened at random. 18 were flagged as not suitable, leaving 12 as potentially suitable. Subsequent Step 1 screening of titles with abstracts selected 25 of these articles for exclusion, with 5 retained for Step 2 (full text screening). All 5 were flagged as being potentially suitable in the pilot set of title-only screenings.

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101 Step 1 were 2.6%, 1.8%, 24.7%, 65.8%, respectively (i.e., 4.7% overall).

102

103 One thousand nine hundred sixty-eight items were excluded, five items were classified as
104 “Maybe”, and one item could not be screened because the full text version was not available,
105 leaving n = 20 suitable items selected (Fig. 1). Most (92.1%) of the exclusions were made
106 from reviewing titles and abstracts at Step 1, with a further 6.6% excluded from screening of
107 the 157 full text items in Step 2. Inter-reader agreement was a Cohen's Kappa $\kappa = 0.69$
108 (or 96.0 percent agreement) for Step 1 and $\kappa = 0.82$ (or 98.1 percent agreement) for Step 2.
109 Percentages of items with search terms (AIMS outcomes) in the title or abstract increased
110 after Step 1 in most cases except for “cancer” and “prostate cancer” (Supplementary table 4).
111 This reflects many exclusions in Step 1 that were of items reporting research on testosterone
112 deprivation or suppression treatments for prostate cancer.

113

114 The systematic approach to literature searching and screening is widely held to be beneficial
115 to identifying studies that otherwise may not have been considered for inclusion, and thus to
116 minimise the prospect for reviewer biases affecting study selections and summary results.[7]
117 This process is not perfect though, and in our case it did not identify two prospective cohort
118 studies that were known to be suitable, prior to commencing this review (FHS, MAILES).[3]
119 In the case of MAILES, this was one of the more recently commenced of the selected studies,
120 with its cohort profile article published in 2014,[8] and accordingly has had a comparatively
121 short timeframe within which to analyse and publish suitable findings. In the case of FHS,
122 associations of endogenous testosterone with male health outcomes had previously been
123 investigated and published, but not using mass spectrometry for measuring testosterone.[9,
124 10] Those articles were identified in the literature search but had been excluded on account of
125 assay method. Only relatively recently have testosterone measures been re-assayed for FHS

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126 participants using mass spectrometry methods.[11] One article by Pencina et al[12] was
127 possibly within scope but not identified because it had not been entered into the MEDLINE
128 database prior to the literature search (article entry date = 14 May 2020). Furthermore, an
129 article that presented suitable estimates from one of the selected studies by Yeap et al[13]
130 was not identified from the literature search because it did not have “prospective”, “follow-
131 up”, “cohort study” or “longitudinal study” terms in its title or abstract, nor any of the
132 corresponding MeSH terms listed (refer to Supplementary table 1 for search terms used).

133

134 In expanding our literature search to unpublished grey literature, it successfully located one
135 suitable item, which was a link to a Web MD webpage article, with further details published
136 in a conference abstract by Sueoka et al[14] that would otherwise have not been returned
137 from searching only the MEDLINE and EMBASE databases.

138

*Supplementary Material. Systematic review: associations of testosterone with men's health.*139 **3. Tables**140 Supplementary table 1. Full electronic search strategy used for MEDLINE database.

141

142 The following is the search that was conducted on 18 July 2019 using MEDLINE.

143

144 1. Testosterone/ or Androgens/

145 2. (testosterone or androgen* or sex hormone* or sex steroid*).ti.

146 3. (testosterone or androgen*).ab.

147 4. cardiovascular diseases/ or heart diseases/ or heart failure/ or vascular diseases/ or stroke/

148 or myocardial infarction/ or coronary disease/ or cerebrovascular disorders/

149 5. (cardiovascular or stroke or myocardial infarction or heart failure).ti.

150 6. neoplasms/ or colorectal neoplasms/ or lung neoplasms/ or prostatic neoplasms/

151 7. cancer.ti.

152 8. mortality/ or mortality.ti.

153 9. dementia/ or cognition/ or dementia.ti. or cognit*.ti.

154 10. Aging/psychology or Neuropsychological Tests/

155 11. 1 or 2 or 3

156 12. 4 or 5 or 6 or 7 or 8 or 9 or 10

157 13. 11 and 12

158 14. longitudinal studies/ or prospective studies/ or follow-up studies/ or cohort studies/

159 15. (prospective or follow-up or cohort study or longitudinal study).ti,ab.

160 16. 14 or 15

161 17. 13 and 16

162 18. (exogenous or replacement or therapy or hormone treatment).ti.

163 19. Hormone Replacement Therapy/

164 20. 18 or 19

165 21. 17 not 20

166 22. limit 21 to humans

167 23. limit 22 to english language

168 24. limit 23 to (adaptive clinical trial or address or autobiography or bibliography or

169 biography or case reports or clinical trial, all or clinical trial, phase i or clinical trial, phase ii

170 or clinical trial, phase iii or clinical trial, phase iv or clinical trial, veterinary or clinical trials,

171 veterinary as topic or clinical trial protocol or clinical trial or comment or controlled clinical

172 trial or dictionary or editorial or lecture or legislation or meta analysis or practice guideline or

173 pragmatic clinical trial or published erratum or randomized controlled trial or retracted

174 publication or "retraction of publication" or "review" or "scientific integrity review" or

175 "systematic review")

176 25. Retrospective Studies/ or Case-Control Studies/ or (retrospective analysis or case-

177 control).ti.

178 26. 24 or 25

179 27. 23 not 26

180

181 Notes:

182

183 Terms with a trailing "/" are MeSH terms and those with a trailing "*" are truncated search

184 strings. Beforehand, a search of PROSPERO was conducted for another suitable strategy but

185 none were found. However, the above strategy is based upon one that has been used for a

186 similar study.[15] This search strategy is also published in the protocol article for the

187 Androgens In Men Study.[3]

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188 **Supplementary table 2: Selection criteria for screening items returned from the literature search.** If neither Include nor Exclude could be selected for Step
189 1, then reviewer selected "Include".

	Exclude	Include	Rationale	Used in Step 1		Used in Step 2
				Title & Abstract		Full-text
				Title only (no abstract)	Title & Abstract	
Article type:	Reviews, comments/opinion pieces, systematic reviews, dictionary, fact sheet, website information about diseases, fact sheets, etc.	Research study article / report, or an article that specifically refers to the results of one (e.g., a webpage referring to unpublished data).	These searches were of both published and unpublished scientific literature for the purpose of identifying prospective cohort studies that are likely to have the relevant data for planned IPD meta-analyses	Yes	Yes	Yes
Study type:	Retrospective or cross-sectional designs, case studies, case-control, surveys, RCTs or other trials, experiments, evaluation of androgen / testosterone therapy / deprivation / HRT or the effectiveness of any other type of intervention / surgery / treatment, genetics, etc.	Prospective cohort study.	A prospective cohort study design is of incident health outcomes for investigating etiology or disease risk for a cohort free of disease at baseline, and ideally should be representative of the local population, but may or may not be some demographic subset: e.g., age range, sex, ethnicity type.	Yes	Yes	Yes
Population (at baseline/date of recruitment to study)	Studies of juveniles only Studies of females only Individuals with some specific health condition/characteristic or following surgery / other medical treatment for specific illness	Adults (18 yr or older) Not females only Community-dwelling men	The study is of community-dwelling men.	Yes	Yes	Yes
Exposure (at baseline)	Do not exclude studies that do not model testosterone as the exposure: although it should be shown that it was measured for participants. If not mentioned in Step 2 then Exclude.	Endogenous testosterone	This will be the focal exposure for all IPD meta-analyses. However, as we are focussing on the identification of only those studies who have suitable androgen measurements available in IPD data, then testosterone does not necessarily need to be modelled as the focal exposure in included items. It is likely that details on the methods will be available only from full-text review.	Only if available	Only if available	Yes
	Testosterone not measured using mass spectrometry	Testosterone assay of serum or plasma sample using mass spectrometry (lc-ms or gc-ms)		Only if available	Only if available	Yes
Outcome (at follow-up)	Incident outcome not one of those type of events specified for inclusion.	Diagnosis/event of: cardiovascular disease (any); cancer (any); dementia. Deaths (any cause); deaths due to any type of cardiovascular disease; deaths due to any type of cancer. Cognition change / outcome	These are the outcomes for the planned IPD meta-analyses so it is important to seek IPD datasets from those studies who have already modelled these outcomes. We refer to these as the "AIMS outcomes".	Yes	Yes	Yes
	Less than 5 years of follow-up data	Five or more years of follow-up data, with outcomes identified using systematic follow-up or data linkage.	As consistent across all included studies for IPD meta-analyses and set <i>a priori</i> . Likely that this will be available only from full-text review so not included Step 1.	No	No	Yes
Language	Title and/or abstract of article not in English	Title and/or abstract of article in English	As limited by the timeframe of this study and the native language of reviewers (a practicality).	Yes	Yes	Yes
Species	Studies not of humans	Studies of humans	We are studying humans.	Yes	Yes	Yes

190

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191 Supplementary table 3: Adaptation of screening rules for different types of published and
 192 unpublished items.
 193

Item Type	Step 1	Step 2
Published article	Screen title (and abstract ^a)	Screen full text article
Thesis	Screen title (and abstract ^a)	Screen full thesis
Unpublished report / other document	Screen title (and abstract ^{a,b})	Screen full document
Webpage	Screen title and webpage ^c	Screen full text article/document as identified from the webpage, or from a google search of information provided about the article, from the webpage.

194

195 ^a = when an abstract was available, otherwise title-only decisions were made (see
 196 Supplementary table 2).

197 ^b = or, if not an abstract, other suitable document summary, as returned by the search tool.

198 ^c = for webpage articles, the webpage text served as the proxy for an abstract, with the
 199 proviso that the reviewer did not navigate to additional webpages during Step 1.

200

*Supplementary Material. Systematic review: associations of testosterone with men's health.*201 Supplementary table 4. Words mentioned in the titles or abstracts of reviewed items.^a

Word(s)	Step 1 items (n=1,994)	Step 2 items (n=158)	Selected items (n=20)
<i>Search terms (AIMS outcomes)</i>			
cancer	1,449 (72.7)	72 (45.6)	6 (30.0)
colorectal cancer	9 (0.5)	4 (2.5)	2 (10.0)
lung cancer	10 (0.5)	6 (3.8)	2 (10.0)
prostate cancer	1,107 (55.5)	40 (25.3)	2 (10.0)
cardiovascular	219 (11.0)	49 (31)	15 (75.0)
heart failure	29 (1.5)	2 (1.3)	1 (5.0)
stroke	31 (1.6)	12 (7.6)	4 (20.0)
myocardial infarction	33 (1.7)	7 (4.4)	1 (5.0)
mortality	232 (11.6)	45 (28.5)	9 (45.0)
dementia	22 (1.1)	8 (5.1)	2 (10.0)
cognit*	87 (4.4)	20 (12.7)	4 (20.0)
<i>Other frequently observed (not search terms)</i>			
androgen deprivation	583 (29.2)	2 (1.3)	0 (0.0)
androgen receptor	235 (11.8)	10 (6.3)	0 (0.0)
brachytherapy	165 (8.3)	0 (0.0)	0 (0.0)
breast cancer	153 (7.7)	9 (5.7)	0 (0.0)
radiotherapy	371 (18.6)	0 (0.0)	0 (0.0)

202 ^a = Items summarised as numbers (percentages); *= wildcard character designating truncation

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Supplementary table 5. PRISMA Checklist.

Section/topic	#	Checklist item	Reported on page #
TITLE			
Title	1	Identify the report as a systematic review, meta-analysis, or both.	1
ABSTRACT			
Structured summary	2	Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.	2-3
INTRODUCTION			
Rationale	3	Describe the rationale for the review in the context of what is already known.	4-5
Objectives	4	Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).	5. For this type of review it is PEO instead.
METHODS			
Protocol and registration	5	Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.	5
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.	5, Supplementary table 2
Information sources	7	Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.	5-6
Search	8	Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.	Supplementary table 1
Study selection	9	State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).	5-6, Supplementary tables 2-3.

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Section/topic	#	Checklist item	Reported on page #
Data collection process	10	Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.	6, Suppl. Data.
Data items	11	List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.	6, Suppl. Data.
Risk of bias in individual studies	12	Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.	6, Supplementary table 8.
Summary measures	13	State the principal summary measures (e.g., risk ratio, difference in means).	7
Synthesis of results	14	Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., I^2) for each meta-analysis.	7
Risk of bias across studies	15	Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).	7
Additional analyses	16	Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.	7
RESULTS			
Study selection	17	Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram.	8, Fig. 1
Study characteristics	18	For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations.	Supplementary tables 6-7, 9-10
Risk of bias within studies	19	Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).	9, Supplementary table 8
Results of individual studies	20	For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.	10-12, Figs 3-4; Supplementary figure 3

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Section/topic	#	Checklist item	Reported on page #
Synthesis of results	21	Present results of each meta-analysis done, including confidence intervals and measures of consistency.	10-12, Figs 3-4; Supplementary figure 3
Risk of bias across studies	22	Present results of any assessment of risk of bias across studies (see Item 15).	10-11, Supplementary figure 2
Additional analysis	23	Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).	10-12, Fig 2; Supplementary figures 1-3
DISCUSSION			
Summary of evidence	24	Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).	12-16
Limitations	25	Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias).	14-15
Conclusions	26	Provide a general interpretation of the results in the context of other evidence, and implications for future research.	13-16
FUNDING			
Funding	27	Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.	17

From: Moher D, Liberati A, Tetzlaff J, Altman DG, The PRISMA Group (2009). Preferred Reporting Items for Systematic Reviews and Meta-Analyses: The PRISMA Statement. PLoS Med 6(7): e1000097. doi:10.1371/journal.pmed1000097

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Supplementary table 6. Attributes of selected items.

Item	Article	Country	Study name [§]	Baseline**			Follow-up (relevant outcomes)		
				No. adult males	Baseline period	Age (yr) Mean (sd)	T (nmol/L) Mean (sd)	Length of follow-up (yr) (person-years) [¶]	AIMS Longitudinal Outcomes (no. of events analysed)
<i>Selected from systematic review</i>									
1	Srinath, 2015[16]	USA	ARIC	1,558	1996-98	63.1 (5.6)	13.9 (5.7)	Md=12.8 (CHD); Md=13.1 (HF) (25,374; HF)	Coronary Heart Disease (CHD; 287) Heart Failure (HF; 104) CHD deaths (29) All-cause deaths (347)
2	Srinath, 2016[17]	USA	ARIC	1,558	1996-98	63.1 (5.6)	13.9 (5.7)	Md=14.1 (27,311)	Ischemic Stroke (79)
3	Chan, 2016[18]	Australia	BHS	1,804	1994-95	50.3 (16.8)	13.6 (4.9)	Mn=14.9 (31,930)	CVD events (234; 399)*** CVD deaths (71; 141)*** All-cause deaths (191; 319)***
4	Chasland, 2017[19]	Australia	BHS	1,649	1994-95	49.8 (15.3)	13.7 (4.9)	Tot=20	CVD events (415) CVD deaths (127)
5	Chan, 2018[20]	Australia	BHS	1,574	1994-95	51.1 (14.7)	13.5 (4.8)	Tot=20	Prostate cancer (116) Lung cancer (22) Colorectal cancer (48) Cancer (any; 289)
6	Hsu, 2015[21]	Australia	CHAMP	853	2005-07	76.9 (5.5)	14.6 (6.2)	Tot=5	Cognitive decline (95)
7	Hsu, 2016[22]	Australia	CHAMP	1,705	2005-07	76.9 (5.5)	14.9 (6.6)	Md=6.9; Tot=10 (11,764)	Cancer deaths (151) CVD deaths (185) Other deaths (174) All-cause deaths (510)
8	Hsu, 2018[23]	Australia	CHAMP	1,651	2005-07	76.9 (5.5)	14.7 (6.4)	Tot=5	All-cause deaths (382) CVD deaths (cases not reported) Cancer deaths (cases not reported) Other deaths (cases not reported) Change in: MMSE, SF-12 (Mental).

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Item	Article	Country	Study name [§]	Baseline**				Follow-up (relevant outcomes)	
				No. adult males	Baseline period	Age (yr) Mean (sd)	T (nmol/L) Mean (sd)	Length of follow-up (yr) (person-years) [¶]	AIMS Longitudinal Outcomes (no. of events analysed)
9	Rosenberg, 2018[24]	USA	CHS	1,019	1994	76.3 (4.9)	13.2 (6.2)	Md=9.5 (10,716)	Atrial Fibrillation (304)
10	Shores, 2014a[25]	USA	CHS	1,032	1994	76.5 (5.2)	13.5 (6.1)	Md=10; Tot=16 (19,220)	Ischemic stroke (114)
11	Shores, 2014b[26]	USA	CHS	NR	1994	NR	NR	Md=8.9 (CVD events) Md=10.8 yr (All-cause deaths). (9,184; CVD events)	CVD events (436) CVD deaths (157) All-cause deaths (777)
12	Lee, 2013[27]	Europe ^{§§}	EMAS	2,736	2003-05	59.2 (10.7)	16.5 (6)	Md=4.3; Tot=5 (14,486)	Cancer (any) Myocardial Infarction (MI) Heart Failure, Other heart conditions Stroke Cognitive function All-cause deaths (193)
13	Pye, 2014[28]	Europe ^{§§}	EMAS	2,599	NR	60 (11)	NR	Md=4.3; Tot=5 (11,140)	Cancer deaths (60) CVD deaths (56) All-cause deaths (147)
14	Chan, 2017[29]	Australia	HIMS	3,690	2001-04	77 (3.6)	13.1 (4.9)	Md=9.1, 9.2; Tot=11 (38,665)	Prostate cancer (348) Lung cancer (107) Colorectal cancer (137)
15	Ford, 2018[30]	Australia	HIMS	4,069	2001-04	NR	NR	Md=10.5; Tot=12 (44,404)	Dementia (499)
16	Yeap, 2014[31]	Australia	HIMS	3,690	2001-04	NR	NR	Mn=6.6 (2.3 sd) (28,036)	MI (344) Stroke (300)

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Item	Article	Country	Study name [§]	Baseline**			Follow-up (relevant outcomes)	
				No. adult males	Baseline period	Age (yr) Mean (sd)	T (nmol/L) Mean (sd)	Length of follow-up (yr) (person-years) [¶]
17	Ohlsson, 2010[32]	Sweden	MrOS	2,644	2001-04	75.5 (3.2)	15.6 (6.5)	Mn=4.5† (11,880) CVD deaths (123) Cancer deaths (127) All-cause deaths (328)
18	Ohlsson, 2011[33]	Sweden	MrOS	2,416	2001-04	75.4 (3.2)	15.7 (6)	Md=5.1 (11,605) CVD events (485) Chronic Heart Disease events (302) Cerebrovascular events (225)
19	Tivesten, 2014[34]	Sweden	MrOS	2,416	2001-04	75.4 (3.2)	15.7 (6)	Md=5.2 (12,070; CHD) Cerebrovascular Disease (225; CBD) (12,137; CBD)
20	Kische, 2017[35]	Germany	SHIP	1,962	1997-01	49.5 (16.3)	15.6 (6.1)	Tot=10 Change in cognitive status
Decision = "Maybe". Item selected based on additional information								
21	LeBlanc, 2010[36]	USA	MrOS	1,602	NR	NR	NR	Mn=4.5†† (26,977) Cognitive function (and change in) Cognitive decline
22*	Sueoka, 2010[14]	USA	MrOS	697	2000-05	72 (5.5)	14.5 (5.1)	Av=3.9†† (6,247) Coronary Heart Disease events (100)
Other. Additional studies selected based on information external to the systematic review								
	No articles were selected.	USA	FHS	3,352[12]	1998-05	59.6 (9.1)[12] 49.4 (13.8)[11]	20.7 (8.0)[12]	Tot=10 (for Atrial Fibrillation)[37] Cardiovascular outcomes[37, 38] Deaths[37] Cause-specific deaths[38] Cancer[39]
	No articles were selected.	Australia	MAILES	1,632[40]	2002-06[8]	54.1 (11.4)[40]	17.3 (5.7)[40]	Md=4.95; IQR=4.35-5.00[40] (12,686) CVD events Deaths (99)[8] Cause-specific deaths[8]

§ Study name abbreviations: 'ARIC'= Atherosclerosis Risk in Communities; 'BHS'=Busselton Health Study; 'CHAMP'=The Concord Health and Ageing in Men Project; 'CHS'= Cardiovascular Health Study; 'FHS'= the Framingham Heart Study; 'HIMS'=The Health In Men Study; 'EMAS'=European Male Ageing Study; 'MAILES'= The Men Androgen Inflammation Lifestyle Environment and Stress study; 'MrOS Sweden'=The MrOS Osteoporotic Fractures in Men study in Sweden; 'MrOS USA' = The MrOS Osteoporotic Fractures in Men study USA; 'SHIP'=Study of Health in Pomerania SHIP.

§§ = UK, Italy, Belgium, Poland, Sweden, Spain, Hungary, Estonia

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¶ 'Md'=median; 'Mn'=mean; 'Av'=average; 'Tot'=total follow-up for the cohort (i.e., maximum, rounded down to nearest whole year); 'IQR'=interquartile range. Unless provided in text, person-years was calculated by multiplying the median, mean, or average length of follow-up by the total number of adult male participants.

* = Note that this is a published conference abstract so is not technically a "Full Text" item.

** = Baseline statistics reported for whole cohort; 'NR' = statistics not reported for whole cohort; Means and standard deviations calculated by firstly transforming into standard units (for T: nmol/L) and then, where required, transforming from quartile statistics using the Box-Cox method of McGrath et al.[41]

*** = First number is for individuals without CVD at baseline.

† = Total follow-up exceeded 5 years, from baseline visit (2001-04) to end of mortality data collection (March 1, 2008).

†† = Note that since there was no published follow-up estimate exceeding 5 years (a requirement for selection) and it was not clear, based on the article information alone, whether the total follow-up was at least 5 years, these items were initially classified as "Maybe". The length of follow-up for collection of AIMS outcome data was determined to be satisfactory from subsequent correspondence with MrOS USA researchers.

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Supplementary table 7. Exposure levels, outcome assessment, covariates.*

Study	Article	Longitudinal measure of association	Exposure (testosterone)	Outcome ascertainment	Covariates
ARIC	Srinath, 2015[16]	HR	T quartiles	CVD events and deaths identified by annual questionnaires and continuous surveillance, independent from hospital admissions data (ICD codes). Cause of death from death certificates.	Age, race/centre, BMI, waist circumference, smoking status, diabetes mellitus, hypertension, LDL, HDL.
	Srinath, 2016[17]	HR	T tertiles	Definite or probable stroke events identified from hospital admissions, annual phone calls, study examinations adjudicated by a physician, with secondary physician adjudication if it disagreed with a computer algorithm.	Age, race, centre, BMI, waist circumference, smoking status, diabetes mellitus, hypertension, LDL, HDL.
BHS	Chan, 2016[18]	HR	T quartiles (results not shown), Continuous T.	Linked hospital admissions and deaths records (ICD codes)	Age, smoking, vigorous exercise, alcohol, BMI, diabetes, CVD, COPD, non-skin cancer, systolic blood pressure, hypertension, lipid lowering therapy, cholesterol, HDL, triglycerides, C-reactive protein, creatinine
	Chasland, 2017[19]	HR	Categories: Low (L) v High (H) T, physical activity(PA) LT+LPA, LT+HPA, HT+LPA, HT+HPA	Linked hospital admissions and deaths records (ICD codes)	Age, prevalent CVD, smoking, waist circumference, cholesterol, HDL, lipids medication, diabetes, systolic blood pressure, hypertension medication
	Chan, 2018[20]	HR	T quartiles, Continuous T.	Linked cancer and death registry records (ICD codes)	Age, marital status, occupation, smoking, alcohol consumption, leisure time physical activity, BMI, diabetes
CHAMP	Hsu, 2015[21]	Slope estimate (change in MMSE on baseline hormone level or longitudinal change in hormone level)	Continuous T, cFT	Clinic assessment: MMSE, Informant Questionnaire on Cognitive Decline as initial screen, followed by clinical assessment to diagnosis categories: normal cognition, MCI, dementia. During follow-up: A decline in MMSE by ≥ 3 points	Age, BMI, smoking status, years of education, depression score (GDS)
	Hsu, 2016[22]	RR	Continuous T, cFT	Deaths identified from 4-monthly phone calls or deaths registry. Cause of death identified on death certificates independently by 2 medical practitioners.	Age, BMI, smoking status, comorbidity score
	Hsu, 2018[23]	HR, RR (Death outcomes); Slope estimates (MMSE, SF-12 Mental)	Categories: Low (<20 th centile) v Normal T combinations with Low (<20 th centile) v Normal cFT	Cause of death identified on death certificates independently by 2 medical practitioners.	Age, BMI, smoking status, comorbidity score
CHS	Rosenberg, 2018[24]	HR	Continuous T and cFT, T and cFT quintiles	Independently verified from ECGs taken annually for participants and from hospital discharge diagnoses	Age (stratified), race, education, income, clinic, smoking status, diabetes mellitus, BMI, loop diuretics, height, hypertension, depressed left ventricular ejection fraction, kidney function, systolic blood pressure, SHBG

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Study	Article	Longitudinal measure of association	Exposure (testosterone)	Outcome ascertainment	Covariates
	Shores, 2014a[25]	HR	Continuous T, cFT (linear & non-linear), T categories	Medicare data, hospital records, imaging studies, autopsy results, death certificates, physician interviews data used for adjudications by committee, which included a neurologist.	Age, systolic blood pressure, anti-hypertensive medications, atrial fibrillation, diabetes, smoking, lipid-lowering drugs, HDL, cholesterol, creatinine, fasting glucose, diabetes medications.
	Shores, 2014b[26]	HR	Continuous T, cFT (linear or non-linear) categories: Q1, Q2-4	Medicare data, hospital records, imaging studies, autopsy results, death certificates, physician interviews data used for adjudications by committee, which included a neurologist.	Age, race, site, smoking status, alcohol consumption, hypertensive use, HDL, BMI, waist circumference, diabetes, SHBG.
EMAS	Lee, 2013[27]	N/A	No modelling of longitudinal outcomes reported	MI, heart failure, other heart conditions, cancers, stroke identified from postal questionnaire, MMSE for participants ≥65 yr old from clinic assessments Variable methods for data capture + validation among centres.	No modelling of longitudinal outcomes reported
	Pye, 2014[28]	HR	T, cFT categories: quintiles, low v eugonadal T, LOH status.	Deaths identified from follow-up postal questionnaire or enquiry if no reply received, with 89% of deaths verified from death certificates, death registers, or medical/hospital records.	Age, site, BMI, smoking status, general health.
HIMS	Chan, 2017[29]	SHR	Continuous T, cFT.	Linked hospital admissions, death and cancer registry records (ICD, ICD-O-3 codes).	Age, BMI, smoking status, physical activity, alcohol consumption, diabetes mellitus, HDL, triglycerides, prior cancer diagnosis.
	Ford, 2018[30]	HR	Continuous T, cFT Quartile categories of T, cFT	Linked data (ICD codes) from inpatient and outpatient mental health services, hospital admissions, community aged care services, cancer and death registries.	Age, baseline cognitive function, depression, BMI, hypertension, CVD, plasma homocysteine.
	Yeap, 2014[31]	HR	T, cFT as quartile categories	Linked hospital admissions, death and cancer registry records (ICD codes).	Age, education, smoking status, BMI, waist to hip ratio, hypertension, dyslipidemia, diabetes, creatinine, prior cancer or existing CVD. Also SHBG for models with T.
MrOS Europe	Ohlsson, 2010[32]	HR (in relation to DHEA, DHEA-S)	No: T modelled as a covariate only	Linked data (death and hospital discharge registries), death certificates.	Age, site, BMI, C-reactive protein, ApoB/A1, smoking status, diabetes, hypertension, prior CVD, prior cancer, low testosterone (in lowest quartile), low estradiol
	Ohlsson, 2011[33]	HR	T, cFT as quartile categories, T as binary categories.	Linked data (death and hospital discharge registries), death certificates.	Age, morning sample, site, BMI, ApoB/A1, physical activity, smoking status, diabetes, hypertension
	Tivesten, 2014[34]	HR (in relation to DHEA, DHEA-S)	No: T modelled as a covariate only	Linked data (death and hospital discharge registries), death certificates.	Age, morning sample, site, BMI, ApoB/A1, C-reactive protein, estradiol, testosterone (i.e., continuous T), SHBG, eGFR, smoking status, diabetes, hypertension.
SHIP	Kische, 2017[35]	Slope estimate (change in MMSE on baseline hormone)	T, cFT as continuous and as 10-year age group quartile categories.	MMSE score.	Age, BMI, smoking status, alcohol consumption, physical activity, hypertension, occupational status, education level, civil status, baseline MMSE.

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Study	Article	Longitudinal measure of association	Exposure (testosterone)	Outcome ascertainment	Covariates
MrOS USA	LeBlanc, 2010[36]	Change in mean score RR of clinically important decline	cFT quartiles and continuous cFT and T (data not shown)	Cognitive tests at the baseline and follow-up visit from Part B of the Trail Making Test (Trails B) and the Modified Mental State Examination (3MS). Calculated from pre-defined drop in scores.	Age group, education level, race, general health, alcohol consumption, clinic, physical and mental health, physical activity, medications used at baseline, other sex steroids, SHBG.
	Sueoka, 2010[14]	HR	T quartiles	CHD events identified from 3-monthly contacts with participants. Incident events were reviewed and adjudicated by cardiologist using clinical records.	Age, clinic, BMI, blood pressure, lipid levels, smoking, hypertension, diabetes, use of lipid-lowering agents
FHS	N/A – no items were selected.	N/A	N/A	AF measured and adjudicated by cardiologists. Mortality data from death certificates, hospital or institutional records, obituaries, or direct notification[37] Medical records of CVD events reviewed by panel of experienced investigators. A heart study neurologist examined most participants with suspected stroke[38] Medical records of cancer diagnoses reviewed by two independent reviewers, with majority confirmed by pathology reports.[39]	N/A
MAILES	N/A – no items selected.	N/A	N/A	Self-reported and clinical follow-up data, death registry (linked data)[8]	N/A

* ApoB/A1 = apolipoprotein-B to apolipoprotein-A1 ratio; BMI = body mass index; cFT = calculated free testosterone; CHD = coronary heart disease; COPD = chronic obstructive pulmonary disease; CVD = cardiovascular disease; DHEA = dehydroepiandrosterone; DHEA-S = DHEA sulfate; ECG = electrocardiogram; eGFR = estimated glomerular filtration rate; GDS = Geriatric Depression Scale; HDL = high-density lipoprotein; HR = hazard ratio; ICD = International Classification of Diseases; ICD-O-3 = International Classification of Diseases for Oncology; LDL = low-density lipoprotein; LOH = late-onset hypogonadism; MCI = mild cognitive impairment; MMSE = mini-mental state examination; N/A = not applicable; Q1=quartile 1; Q2-4=quartiles 2 to 4 combined; RR = relative risk; SF-12 = The Short Form (12) Health Survey; SHBG = sex hormone binding globulin; SHR = subhazard ratio, as estimated from competing-risks regression; T = total endogenous testosterone.

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Supplementary table 8. Newcastle-Ottawa Quality Assessment Scale for Cohort Studies: selected articles.

Article	Study	Selection (4 stars)	Comparability (2 stars)	Outcome (3 stars)	Notes on Selection	Notes on Outcome
Srinath 2015[16]	ARIC	****	**	**		Losses to f/u not mentioned; linked data ^b
Srinath 2016[17]	ARIC	****	**	**		Losses to f/u not mentioned; linked data ^b
Chan 2016[18]	BHS	****	**	**		Losses to f/u not mentioned; linked data ^b
Chasland 2017[19]	BHS	***	**	**	Prevalent cases not excluded ^a	Losses to f/u not mentioned; linked data ^b
Chan 2018[20]	BHS	****	**	**		Losses to f/u not mentioned; linked data ^b
Hsu 2015[21]	CHAMP	****	**	***		
Hsu 2016[22]	CHAMP	***	**	***	Prevalent cases not excluded ^c	
Hsu 2018[23]	CHAMP	***	**	***	Prevalent cases not excluded ^c	
Rosenberg 2018[24]	CHS	****	**	**		Losses to f/u not mentioned; linked data ^d
Shores 2014a[25]	CHS	****	**	**		Losses to f/u not mentioned; linked data ^d
Shores 2014b[26]	CHS	****	**	**		Losses to f/u not mentioned; linked data ^d
Lee 2013[27]	EMAS	NA	NA	NA	No modelling of longitudinal outcomes reported	
Pye 2014[28]	EMAS	***	**	***	Prevalent cases not excluded ^a	
Chan 2017[29]	HIMS	****	**	**		Losses to f/u not mentioned; linked data ^b
Ford 2018[30]	HIMS	****	**	**		Losses to f/u not mentioned; linked data ^b
Yeap 2014[31]	HIMS	***	**	**	Prevalent cases not excluded ^a	Losses to f/u not mentioned; linked data ^b
Ohlsson 2010[32]	MrOS Sw.	NA	NA	NA	Testosterone was not the exposure variable in this article	
Ohlsson 2011[33]	MrOS Sw.	****	**	***		
Tivesten 2014[34]	MrOS Sw.	NA	NA	NA	Testosterone was not the exposure variable in this article	
Kische 2017[35]	SHIP	***	**	***	Prevalent cases not excluded ^c	
LeBlanc 2010[36]	MrOS USA	****	**	*		Bias from loss to f/u; F/u OK: additional steps ^f
Sueoka 2010[14]	MrOS USA	***	**	*	Prevalent cases not excluded ^a	F/u OK: additional steps ^f
Additional item not selected but included in DR-MA:						
Yeap 2014b[13]	HIMS	***	**	**	Prevalent cases not excluded ^a	Losses to f/u not mentioned; linked data ^b

'NA' = Not applicable (see Notes); 'f/u' = follow-up (of incident events); 'DR-MA' = dose-response meta-analyses of published estimates.

^a = The influence of prevalent cases was statistically adjusted by including prevalent status as a model predictor.

^b = Follow-up of cases was assumed to be almost complete because analyses were of linked administrative data.

^c = The influence of prevalent cases was statistically adjusted by incorporating into a comorbidity status model predictor.

^d = Follow-up of cases was assumed to be almost complete because analyses were of linked administrative data (with expert adjudications).

^e = Outcome was change in cognition score, with baseline score (prevalent status) included as a model predictor.

^f = Total length of follow-up period was not reported but determined to be satisfactory from correspondence with MrOS USA researchers.

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Supplementary table 9. Extracted hazard ratio data for dose-response meta-analyses (DR-MAs).*

Article	Study	Outcome	Testosterone	Units	HR	95% CI	Notes
Chan 2016[18]	BHS	All-cause mortality	<10.20	nmol/L	ref.		
Chan 2016	BHS	All-cause mortality	10.20 - <13.04	nmol/L	0.84	(0.62-1.14)	
Chan 2016	BHS	All-cause mortality	13.04 - <16.58	nmol/L	0.86	(0.62-1.19)	
Chan 2016	BHS	All-cause mortality	≥16.58	nmol/L	0.9	(0.62-1.3)	
Pye 2014[28]	EMAS	All-cause mortality	<11.65	nmol/L	1.1	(0.6-1.8)	
Pye 2014	EMAS	All-cause mortality	11.65-14.61	nmol/L	0.7	(0.4-1.3)	
Pye 2014	EMAS	All-cause mortality	14.61-17.28	nmol/L	0.7	(0.4-1.3)	
Pye 2014	EMAS	All-cause mortality	17.28-21.20	nmol/L	1.2	(0.7-2)	
Pye 2014	EMAS	All-cause mortality	>21.20	nmol/L	ref.		
Srinath 2015[16]	ARIC	All-cause mortality	≤288.4	ng/dL	0.96	(0.7-1.34)	
Srinath 2015	ARIC	All-cause mortality	288.5-377.6	ng/dL	0.99	(0.72-1.35)	
Srinath 2015	ARIC	All-cause mortality	377.7-480.1	ng/dL	1	(0.74-1.35)	
Srinath 2015	ARIC	All-cause mortality	≥480.2	ng/dL	ref.		
Shores 2014b[26]	CHS	All-cause mortality	<278	ng/dL	1.06	(0.88-1.29)	
Shores 2014b	CHS	All-cause mortality	≥278	ng/dL	ref.		
Yeap 2014b[13]	HIMS	All-cause mortality	0.25-9.82	nmol/L	ref.		Fully-adjusted model + SHBG
Yeap 2014b	HIMS	All-cause mortality	9.82-12.53	nmol/L	0.81	(0.68-0.98)	
Yeap 2014b	HIMS	All-cause mortality	12.56-15.75	nmol/L	0.75	(0.61-0.92)	
Yeap 2014b	HIMS	All-cause mortality	15.79-46.50	nmol/L	0.77	(0.61-0.97)	
Yeap 2014b	HIMS	All-cause mortality	0.25-9.82	nmol/L	ref.		Fully-adjusted model + LH
Yeap 2014b	HIMS	All-cause mortality	9.82-12.53	nmol/L	0.84	(0.7-1.01)	
Yeap 2014b	HIMS	All-cause mortality	12.56-15.75	nmol/L	0.81	(0.67-0.97)	
Yeap 2014b	HIMS	All-cause mortality	15.79-46.50	nmol/L	0.89	(0.73-1.07)	
Hsu 2016[22]	CHAMP	All-cause mortality		ng/mL	1.17	(1.03-1.32)	Per SD decrease in T. RR estimate used.
Chan 2016	BHS	CVD mortality	<10.20	nmol/L	ref.		

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Article	Study	Outcome	Testosterone	Units	HR	95% CI	Notes
Chan 2016	BHS	CVD mortality	10.20 - <13.04	nmol/L	1.12	(0.7-1.78)	
Chan 2016	BHS	CVD mortality	13.04 - <16.58	nmol/L	1.39	(0.86-2.25)	
Chan 2016	BHS	CVD mortality	≥16.58	nmol/L	1.25	(0.69-2.25)	
Chasland 2017[19]	BHS	CVD mortality	<13.1	nmol/L	ref.		Total PA, "Low" PA, NS PA x T: these estimates were used
	BHS	CVD mortality	≥13.1	nmol/L	1.25	(0.77-2.03)	
Chasland 2017	BHS	CVD mortality	<13.1	nmol/L	0.69	(0.4-1.2)	Total PA, "High" PA, NS PA x T
Chasland 2017	BHS	CVD mortality	≥13.1	nmol/L	0.8	(0.48-1.35)	
Pye 2014	EMAS	CVD mortality	<11.65	nmol/L	1	(0.4-2.2)	
Pye 2014	EMAS	CVD mortality	11.65-14.61	nmol/L	0.5	(0.2-1.4)	
Pye 2014	EMAS	CVD mortality	14.61-17.28	nmol/L	0.4	(0.2-1.2)	
Pye 2014	EMAS	CVD mortality	17.28-21.20	nmol/L	1.1	(0.5-2.4)	
Pye 2014	EMAS	CVD mortality	>21.20	nmol/L	ref.		
Srinath 2015	ARIC	CVD mortality	≤288.4	ng/dL	1.36	(0.45-4.08)	
Srinath 2015	ARIC	CVD mortality	288.5-377.6	ng/dL	1.26	(0.73-3.7)	
Srinath 2015	ARIC	CVD mortality	377.7-480.1	ng/dL	0.57	(0.16-1.99)	
Srinath 2015	ARIC	CVD mortality	≥480.2	ng/dL	ref.		
Shores 2014b	CHS	CVD mortality	<278	ng/dL	1.28	(0.94-1.75)	
Shores 2014b	CHS	CVD mortality	≥278	ng/dL	ref.		
Yeap 2014b	HIMS	CVD mortality	0.25-9.82	nmol/L	ref.		
Yeap 2014b	HIMS	CVD mortality	9.82-12.53	nmol/L	0.82	(0.61-1.11)	
Yeap 2014b	HIMS	CVD mortality	12.56-15.75	nmol/L	0.79	(0.58-1.09)	
Yeap 2014b	HIMS	CVD mortality	15.79-46.50	nmol/L	0.79	(0.56-1.11)	
Hsu 2016	CHAMP	CVD mortality		ng/mL	1.11	(0.93-1.32)	Per SD decrease in T. RR estimate used
Srinath 2016[17]	ARIC	Stroke / CBD	≤317.7	ng/dL	1.47	(0.83-2.61)	
	ARIC	Stroke / CBD	317.8-441.2	ng/dL	ref.		
Srinath 2016	ARIC	Stroke / CBD	≥441.3	ng/dL	1.15	(0.62-2.14)	
Shores 2014a[25]	CHS	Stroke / CBD	<200	ng/dL	1.46	(0.77-2.75)	
Shores 2014a	CHS	Stroke / CBD	200-400	ng/dL	0.9	(0.56-1.45)	

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Article	Study	Outcome	Testosterone	Units	HR	95% CI	Notes
Shores 2014a	CHS	Stroke / CBD	400-600	ng/dL	ref.		
Shores 2014a	CHS	Stroke / CBD	600-800	ng/dL	1.73	(0.88-3.39)	
Shores 2014a	CHS	Stroke / CBD	>800	ng/dL	1.69	(0.51-5.60)	
Yeap 2014[31]	HIMS	Stroke / CBD	0.25-9.82	nmol/L	ref.		
Yeap 2014	HIMS	Stroke / CBD	9.82-12.53	nmol/L	0.8	(0.59-1.09)	
Yeap 2014	HIMS	Stroke / CBD	12.56-15.75	nmol/L	0.72	(0.52-0.99)	
Yeap 2014	HIMS	Stroke / CBD	15.79-46.5	nmol/L	0.56	(0.39-0.81)	
Ohlsson 2011[33]	MrOS(Sw)	Stroke / CBD	≤340	ng/dL	ref.		
Ohlsson 2011	MrOS(Sw)	Stroke / CBD	341-438	ng/dL	ref.		
Ohlsson 2011	MrOS(Sw)	Stroke / CBD	439-549	ng/dL	ref.		Quartile 4 vs. quartiles 1 to 3 of testosterone
Ohlsson 2011	MrOS(Sw)	Stroke / CBD	≥550	ng/dL	0.76	(0.55-1.05)	
Ohlsson 2011	MrOS(Sw)	CVD	≤340	ng/dL	ref.		
Ohlsson 2011	MrOS(Sw)	CVD	341-438	ng/dL	1.02	(0.80-1.30)	
Ohlsson 2011	MrOS(Sw)	CVD	439-549	ng/dL	0.96	(0.75-1.23)	
Ohlsson 2011	MrOS(Sw)	CVD	≥550	ng/dL	0.71	(0.54-0.93)	
Chan 2016	BHS	CVD		nmol/L	1.03	(0.92-1.15)	Per SD increase in T
Chasland 2017	BHS	CVD	<13.1	nmol/L	ref.		Total PA, "Low" PA, NS PA x T: these estimates were used
Chasland 2017	BHS	CVD	≥13.1	nmol/L	1.09	(0.83-1.44)	
Chasland 2017	BHS	CVD	<13.1	nmol/L	0.93	(0.70-1.23)	Total PA, "High" PA, NS PA x T
Chasland 2017	BHS	CVD	≥13.1	nmol/L	1.04	(0.78-1.38)	
Shores 2014b	CHS	CVD	<278	ng/dL	1.11	(0.87-1.43)	
Shores 2014b	CHS	CVD	≥278	ng/dL	ref.		
Yeap 2014	HIMS	CVD: MI	0.25-9.82	nmol/L	ref.		
Yeap 2014	HIMS	CVD: MI	9.82-12.53	nmol/L	1.07	(0.79-1.44)	
Yeap 2014	HIMS	CVD: MI	12.56-15.75	nmol/L	1.03	(0.76-1.41)	
Yeap 2014	HIMS	CVD: MI	15.79-46.5	nmol/L	0.92	(0.66-1.28)	
Srinath 2015	ARIC	CVD: HF	≤288.4	ng/dL	0.77	(0.46-1.29)	
Srinath 2015	ARIC	CVD: HF	288.5-377.6	ng/dL	0.72	(0.43-1.21)	

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Article	Study	Outcome	Testosterone	Units	HR	95% CI	Notes
Srinath 2015	ARIC	CVD: HF	377.7-480.1	ng/dL	0.87	(0.53-1.42)	
Srinath 2015	ARIC	CVD: HF	≥480.2	ng/dL	ref.		
Chan 2018[20]	BHS	Cancer	<10.17	nmol/L	ref.		
Chan 2018	BHS	Cancer	10.17-<12.95	nmol/L	0.72	(0.53-0.99)	
Chan 2018	BHS	Cancer	12.95-<16.49	nmol/L	0.71	(0.51-0.98)	
Chan 2018	BHS	Cancer	≥16.49	nmol/L	0.81	(0.57-1.14)	
Chan 2018	BHS	Cancer: Prostate	<10.17	nmol/L	ref.		
Chan 2018	BHS	Cancer: Prostate	10.17-<12.95	nmol/L	0.62	(0.37-1.03)	
Chan 2018	BHS	Cancer: Prostate	12.95-<16.49	nmol/L	0.75	(0.46-1.23)	
Chan 2018	BHS	Cancer: Prostate	≥16.49	nmol/L	0.58	(0.33-1.01)	
Chan 2018	BHS	Cancer: Colorectal		nmol/L	1.04	(0.76-1.42)	Per SD increase in T
Chan 2018	BHS	Cancer: Lung		nmol/L	0.65	(0.39-1.09)	Per SD increase in T
Chan 2017[29]	HIMS	Cancer: Prostate		nmol/L	1.00	(0.90-1.12)	Per SD increase in T.
Chan 2017	HIMS	Cancer: Colorectal		nmol/L	0.96	(0.80-1.15)	Per SD increase in T
Chan 2017	HIMS	Cancer: Lung		nmol/L	1.30	(1.06-1.60)	Per SD increase in T
Hsu 2016	CHAMP	Cancer mortality		ng/mL	1.30	(1.02-1.65)	Per SD decrease in T. RR estimate used.
Ford 2018[30]	HIMS	Dementia	Not reported	nmol/L	1.39	(1.04-1.85)	
Ford 2018	HIMS	Dementia			1.31	(1.00-1.73)	
Ford 2018	HIMS	Dementia			1.23	(0.93-1.61)	
Ford 2018	HIMS	Dementia			ref.		
Ford 2018	HIMS	Dementia	SD for cohort not reported	nmol/L	1.11	(1.01-1.21)	Per SD decrease in T

* = Estimates were also reported for all-cause and CVD mortality mortality for the CHAMP study in another of the selected articles,[23] but were not used because they were reported for combinations of free testosterone and total testosterone, and so were not comparable to the above published estimates.

CBD = cerebrovascular disease; CVD = cardiovascular disease; HF = heart failure; LH = luteinising hormone; MI = myocardial infarction; NS = non-significant result at *a priori* selected threshold for test; PA = physical activity level; ref. = referent level; RR = relative risk; SD = standard deviation; SHBG = sex hormone-binding globulin; T = endogenous total testosterone concentration.

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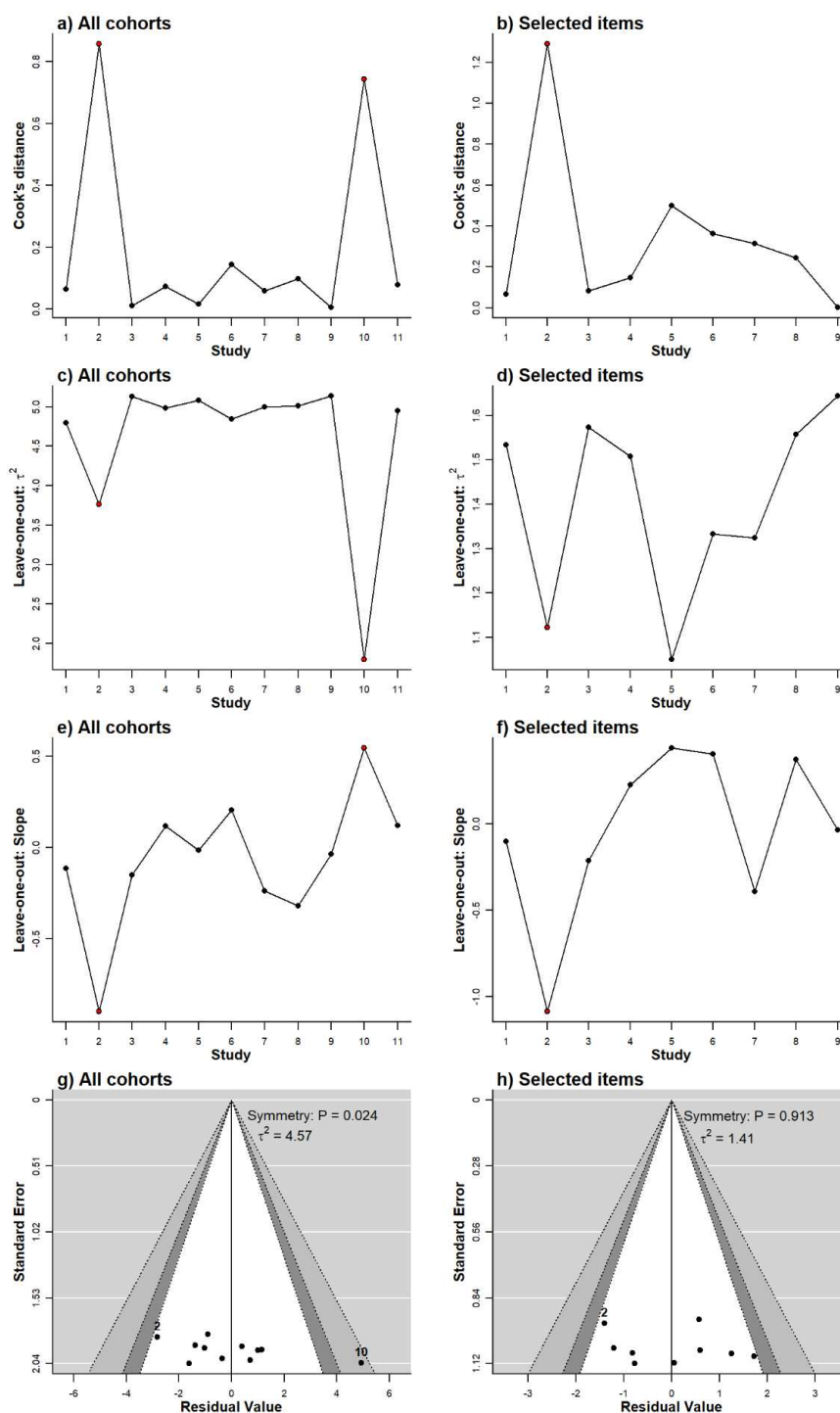
Supplementary table 10. Published estimates for selected studies investigating associations of total testosterone with cognitive status or decline.*

Article	Study	Outcome	Testosterone	Units	Effect size parameter	Estimate	95% CI	p value	Notes
Hsu 2015[21]	CHAMP	Cognitive change	Baseline	ng/mL	Slope: Per	0.012	NR	0.7	Longitudinal change in MMSE score
Hsu 2015	CHAMP	Cognitive change	Longitudinal	ng/mL	unit decline	0.067	NR	0.03	Longitudinal change in MMSE score
Hsu 2015	CHAMP	Cognitive decline	Baseline	ng/mL	Odds Ratio	NR	NR	NR	Longitudinal decline in MMSE \geq 3 points. Non-significant association (data not shown)
Hsu 2018[23]	CHAMP	Baseline cognition	Baseline NN	nmol/L	Slope	0	0	NR	MMSE** at baseline
Hsu 2018	CHAMP	Baseline cognition	Baseline NL	nmol/L	Slope	0.1	-0.8-1.1	NR	MMSE at baseline
Hsu 2018	CHAMP	Baseline cognition	Baseline LN	nmol/L	Slope	0.02	-1.0-1.02	NR	MMSE at baseline
Hsu 2018	CHAMP	Baseline cognition	Baseline LL	nmol/L	Slope	0.8	-0.5-0.4	NR	MMSE at baseline
Hsu 2018	CHAMP	Cognitive change	Baseline NN	nmol/L	Slope	0	0	NR	Longitudinal change in MMSE score
Hsu 2018	CHAMP	Cognitive change	Baseline NL	nmol/L	Slope	0.007	-0.5-0.5	NR	Longitudinal change in MMSE score
Hsu 2018	CHAMP	Cognitive change	Baseline LN	nmol/L	Slope	-0.2	-0.7-0.4	NR	Longitudinal change in MMSE score
Hsu 2018	CHAMP	Cognitive change	Baseline LL	nmol/L	Slope	-0.004	-0.3-0.3	NR	Longitudinal change in MMSE score
Kische 2017[35]	SHIP	Cognitive change	Baseline	nmol/L	Slope	0.02	-0.15-0.20	\geq 0.05	Longitudinal change in MMSE score after 5 years
Kische 2017	SHIP	Cognitive change	Baseline	nmol/L	Slope	0.01	-0.22-0.24	\geq 0.05	Longitudinal change in MMSE score after 10 years
LeBlanc 2010[36]	MrOS US	Baseline cognition	Baseline	nmol/L	-	NR	NR	\geq 0.29	3MS score (includes MMSE)
LeBlanc 2010	MrOS US	Follow-up cognition	Baseline	nmol/L	-	NR	NR	\geq 0.29	3MS score (includes MMSE)
LeBlanc 2010	MrOS US	Cognitive change	Baseline	nmol/L	-	NR	NR	\geq 0.29	3MS score (includes MMSE)
LeBlanc 2010	MrOS US	Baseline cognition	Baseline	nmol/L	-	NR	NR	\geq 0.63	Trails B: test of executive function and motor speed
LeBlanc 2010	MrOS US	Follow-up cognition	Baseline	nmol/L	-	NR	NR	\geq 0.63	Trails B: test of executive function and motor speed
LeBlanc 2010	MrOS US	Cognitive change	Baseline	nmol/L	-	NR	NR	\geq 0.63	Trails B: test of executive function and motor speed

* NR = Not reported; NN = Normal (>10.2 nmol/L) total testosterone (T), normal (>156 pmol/L) calculated free testosterone (cFT); NL = normal T, low (<156 pmol/L) cFT; LN = low (<10.2 nmol/L) T, normal cFT; LL = low T, low cFT. MMSE = mini-mental state examination; 3MS = modified mini-mental state examination.

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4. Figures.

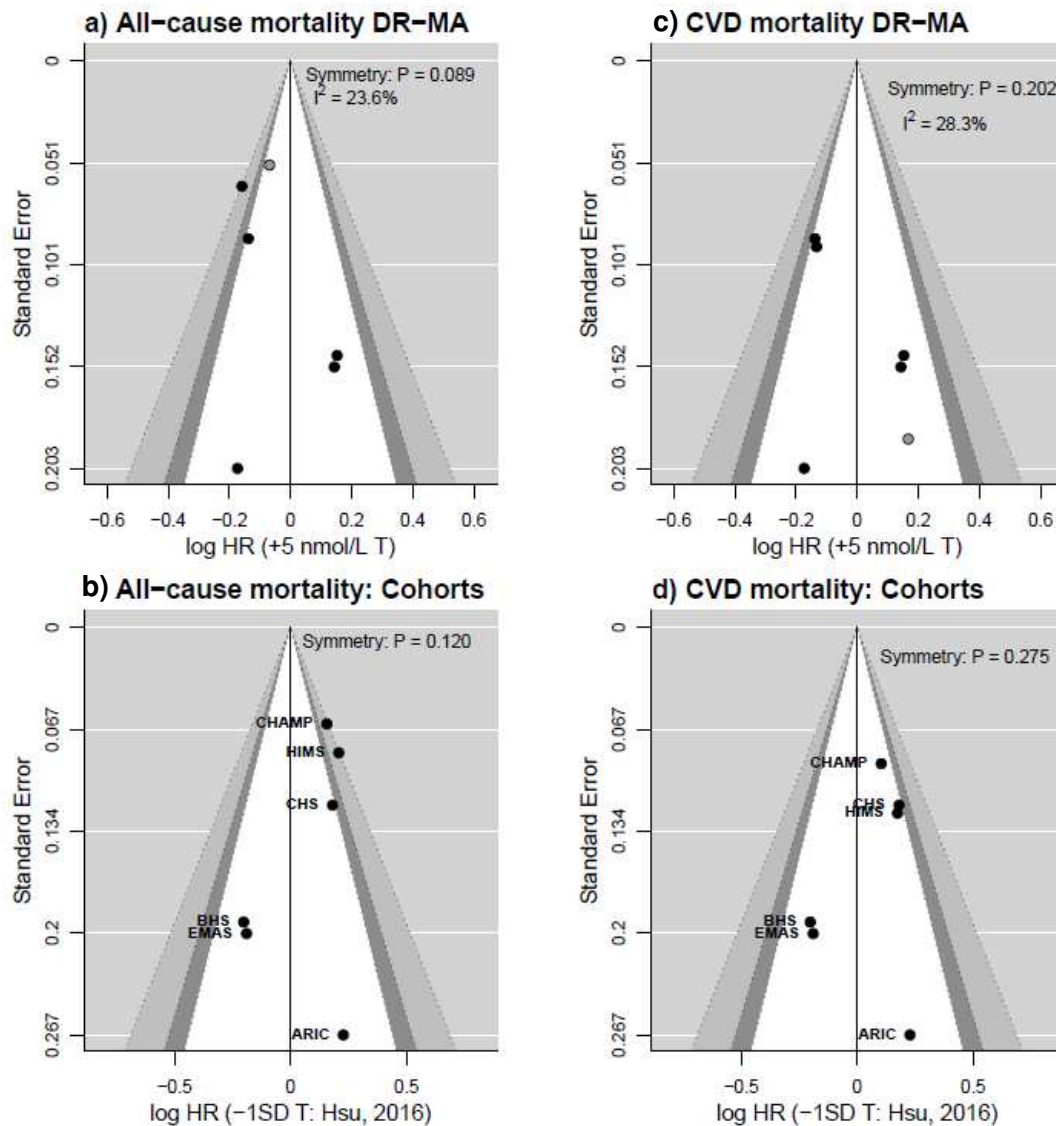


Supplementary figure 1. Meta-regression diagnostics. Meta-regression diagnostics showing the influence of studies on model fit (a,b), τ^2 (estimated amount of total heterogeneity: c,d), estimated slope (e,f), and distribution of residuals with funnel plots (g,h). Analysis repeated for all 11 cohort studies (a,c,e,g) and for 9 studies with selected articles (b,d,f,h). In cases where more than one article was available per cohort study, the article with the largest sample

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size was used. Highlighted estimates for cohort study 2 (BHS) were those from Chan et al.[18] (N=1,804) and for study 10 (FHS) were from Pencina et al.[12] (N=720). In funnel plots: light grey + dark grey + white shading = 99% pseudo confidence interval (CI); dark grey + white shading = 95% CI; white shading = 90% CI.

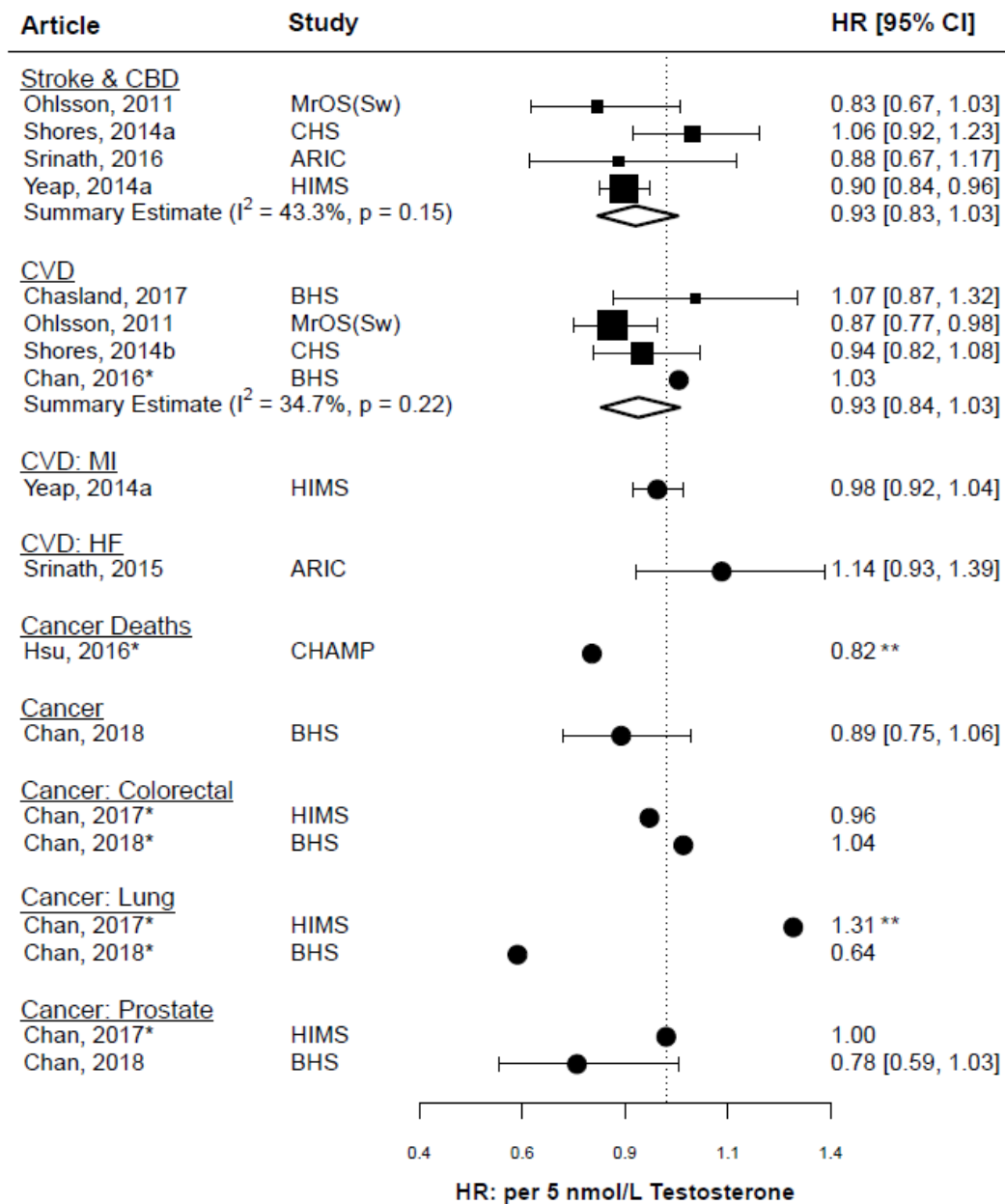
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Supplementary figure 2. Funnel plots for dose-response meta-analyses. Contour-enhanced funnel plots showing the distribution of log hazard ratio (HR) estimates for all-cause mortality (a, b) and mortality caused by cardiovascular disease (CVD) (c, d) attributed to a 5 nmol/L increase (a, c), or a 1.9 nmol/L (1SD in Hsu et al. 2016[22]) decrease (b, d), in endogenous testosterone concentration. Log HR values and standard errors were calculated using generalised least squares regression of published estimates.[42, 43] In cases where more than one article was available per cohort study, the article with the largest sample size was used. Estimates represented by black dots in (a) and (c) were analysed in respective dose-response meta-analyses (DR-MA; results presented in Figs. 3, 4). The grey dot in (a) is the estimate for Yeap et al. 2014b[13] and in (c) is the estimate for Chasland et al. (2017)[19]; these estimates were substituted for others for the HIMS and BHS studies respectively for alternative summary estimates (i.e., the grey summary estimates presented in Figs. 3, 4). Estimates presented in (b) and (d) are shown for a more complete assessment of funnel plot symmetry: estimates are plotted for all studies with estimates, including those that did not have sufficient information for including in the DR-MA. In funnel plots: light grey + dark

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grey + white shading = 99% pseudo confidence interval (CI); dark grey + white shading = 95% CI; white shading = 90% CI.



Supplementary figure 3: Forest plot of published hazard ratio (HR) estimates: association of testosterone with other AIMS outcomes. Plotted estimates for other outcomes, as listed in the AIMS protocol article,[3] have been standardised to the HR for a 5nmol/L increase in testosterone. The size of squares are scaled to the precision of estimates, as used for obtaining the corresponding summary estimate for that outcome (diamonds). Estimates presented as circles were not used to obtain a summary estimate and so the size of circles is not scaled to estimated precision. * = 95 % confidence intervals (CIs) were not calculable for these

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estimates, which were reported as the HR per standard deviation change (see Supplementary table 9).

** = for 'per SD' estimates: to show that the published HR 95% CIs did not overlap 1. 'CBD' = Cerebrovascular disease; 'CVD' = cardiovascular disease; 'MI' = myocardial infarction; 'HF' = heart failure. Study-specific estimates presented for MrOS Sweden (Ohlsson, 2011)[33]; CHS (Shores, 2014a, b)[25-26]; ARIC (Srinath, 2015; 2016)[16-17]; HIMS (Yeap, 2014a, b)[13,31]; BHS (Chan, 2016; Chasland, 2017; Chan, 2018)[18-20].

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