- Additional material is published online only. To view please visit the journal online (http://dx.doi.org/10.1136/ bjsports-2018-099099).
${ }^{1}$ Faculty of Education, Arts and Sports, Western Norway University of Applied Sciences, Sogndal, Norway
${ }^{2}$ Department of Sport Medicine, Norwegian School of Sport Sciences, Oslo, Norway


## Correspondence to

Solveig Nordengen, Faculty of Education, Arts and Sports, Western Norway University of Applied Sciences, Norway; solveig.nordengen@hvl.no

Accepted 17 July 2018

- http://dx.doi.org/10.1136/ bjsports-2018-099778


## (D) Check for updates

© Author(s) (or their employer(s)) 2019. No commercial re-use. See rights and permissions. Published by BMJ.

To cite: Nordengen S, Andersen LB,
Solbraa AK, et al.
Br J Sports Med Epub ahead of print: [please include Day Month Year]. doi:10.1136/
bjsports-2018-099099

# Cycling is associated with a lower incidence of cardiovascular diseases and death: Part 1 - systematic review of cohort studies with meta-analysis 

Solveig Nordengen, ${ }^{\text {1,2 }}$ Lars Bo Andersen, ${ }^{1,2}$ Ane K Solbraa, ${ }^{1}$ Amund Riiser ${ }^{1}$


#### Abstract

Objectives Physical inactivity is a risk factor for cardiovascular disease (CVD). Cycling as a physical activity holds great potential to prevent CVD. We aimed to determine whether cycling reduces the risk of CVD and CVD risk factors and to investigate potential doseresponse relationships. Design Systematic review and meta-analysis of quantitative studies. Eligibility criteria for selecting studies We searched four databases (Web of Science, MEDLINE, SPORTDiscus and Scopus). All quantitative studies, published until August 2017, were included when a general population was investigated, cycling was assessed either in total or as a transportation mode, and CVD incidence, mortality or risk factors were reported. Studies were excluded when they reported continuous outcomes or when cycling and walking were combined in them. We pooled adjusted relative risks (RR) and OR. Heterogeneity was investigated using I. Results The search yielded 5174 studies; 21 studies which included 1,069,034 individuals. We found a significantly lower association in combined CVD incidence, mortality and physiological risk factors with total effect estimate 0.78 ( $95 \% \mathrm{CI}(\mathrm{Cl}): 0.74-0.82$; $\left.\mathrm{P}<0.001 ; \mathrm{I}^{2}=58 \%\right)$. Separate analyses for CVD incidence, mortality and risk factors showed estimates of RR 0.84 (CI, 0.80 to $0.88 ; \mathrm{P}<0.001 ; \mathrm{I}^{2}=29 \%$ ), RR 0.83 (Cl, 0.76 to $\left.0.90 ; \mathrm{P}<0.001 ; \mathrm{I}^{2}=0 \%\right)$, and OR 0.75 ( $\mathrm{Cl}, 0.69$ to $\left.0.82 ; P<0.001 ;\left.\right|^{2}=66 \%\right)$, respectively. We found no dose-response relationship or sex-specific difference. Conclusions Any form of cycling seems to be associated with lower CVD risk, and thus, we recommend cycling as a health-enhancing physical activity. Systematic review registration Prospero CRD42016052421.


## INTRODUCTION

The rise in non-communicable diseases (NCDs) is a growing challenge worldwide. ${ }^{12}$ In 2016, cardiovascular disease (CVD) was one of the five leading causes of years of life lost. ${ }^{3}$ Physical inactivity is associated with CVD and CVD risk factors, ${ }^{4}{ }^{5}$ and the WHO has declared physical inactivity the fourth leading risk factor for global mortality. ${ }^{6}$ Approximately a quarter of the world's adults are physically inactive. ${ }^{7}$ Globally, the level of physical activity has decreased over previous decades ${ }^{8}$ and is still decreasing. ${ }^{7}$ Multi-sectorial and multidisciplinary public health actions are needed to tackle the problem of physical inactivity. ${ }^{9}$

## What is already known?

- The rise of non-communicable diseases is a growing challenge worldwide.
- Physical inactivity is associated with CVD as well as its risk factors.
- Thus, it is necessary to increase physical activity levels by means of multi-sectorial and multidisciplinary public health actions.
- Active transport may be a promising approach to increase levels of physical activity and reduce CVD risk.


## What are the new findings?

- Cycling was associated with $22 \%$ lower risk of combined CVD risk than using passiv transport.
- There was no sex-difference or dose-response relationship of cycling and risk of CV
- Politicians, stakeholders and city planners may promote cycling as public health action.

Changes in the built environment are likely to increase the activity level among children and adults. ${ }^{10}$ Walking and cycling separately, adjusted for other physical activity, may reduce the all-cause mortality at a population level. ${ }^{11}$ Active transportation may also reduce the incidence of NCDs, including CVD. ${ }^{8}$ Therefore, active transportation may be a promising approach to increase physical activity levels and reduce CVD risk. In addition, cycling as transportation may appeal to many people who are not interested in participating in sport as a means of being physically active.

One limitation of research studies investigating active transportation is that they often combine walking and cycling. ${ }^{12}$ This is a problem since cycling often is performed at a higher exercise intensity than walking, ${ }^{13}$ and higher exercise intensity is associated with a further reduction in risk of coronary heart disease. ${ }^{14}$ Therefore, cycling may be more effective than walking in preventing CVD. ${ }^{12}$ To our knowledge, there has not been a meta-analysis examining prevention of CVD and cycling. Nevertheless, there are two meta-analyses examining CVD and active transport ${ }^{1516}$ and one literature review of cycling. ${ }^{12}$ Therefore, this systematic review with meta-analysis of cycling and CVD adds increased power to investigate the association, as data are pooled, and accounts better for
the observed heterogeneity than when walking and cycling are combined.

We aimed to assess the strength of association between cycling and (1) CVD and (2) CVD risk factors. We hypothesised there would be similar associations for men and women, and a dose-response relationship between cycling and health.

## METHODS

We conducted a systematic review with meta-analysis. The protocol was registered with the PROSPERO database on 6 December 2016 (PROSPERO ID: CRD42016052421) (http:// www.crd.york.ac.uk/PROSPERO/display_record.php?ID = CRD42016052421) and complied with Preferred Reporting Items for Systematic Reviews and Meta-Analyses 2009 guidelines. ${ }^{17}$

## Literature search

We searched for published quantitative studies (prospective, retrospective, cohort, longitudinal design and cross-sectional studies or randomised controlled trials) that examined the association of cycling with CVD or CVD risk factors to 8 August 2017. The first author (SN), in cooperation with a librarian, performed the search. Published and peer-reviewed articles in English were identified from four electronic databases: Web of Science, MEDLINE, SPORTDiscus and Scopus. The search strategy consisted of the terms 'cycling' OR 'bicycling' OR 'biking' OR 'commuter cycling' AND ‘CVD' OR
'CVD risk factors' OR 'CVD risk factor' OR 'cardiovascular disease risk factors' OR 'cardiovascular disease' OR 'cardiovascular diseases' OR 'cardiovascular disease*." In total, 5174 records were identified: Web of Science (3525), MEDLINE (via EBSCO) (522), SPORTDiscus (41)and Scopus (1086). After elimination of duplicates, 4785 records remained (figure 1). ${ }^{17}$ See online supplementary table 1, for example, of full search strategy run in MEDLINE via EBSCO. We searched the reference lists of included studies and contacted experts in the field to identify any studies that may have been missed in our electronic database search.

## Inclusion criteria and selection process

Studies were excluded if they measured domains other than cycling, such as stationary cycling, or if cycling was a part of a rehabilitation programme/intervention or investigated an unhealthy population. We had no criteria for sample size.

We included studies that (1) employed a quantitative design and studied a general population; (2) assessed cycling exposure either as a mode of transportation, or as a recreational activity; (3) measured CVD, CVD mortality or physiological CVD risk factors as an outcome and (4) reported dichotomous outcome measures.

Two reviewers (SN and AR) independently assessed the studies for eligibility with subsequent consensus by discussion.


Figure 1 Flow chart of included studies as proposed by Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement 2009. ${ }^{17}$

Table 1 Quality assessments of included studies based on the Quality Assessment Tool of Quantitative Studies ${ }^{18}$

| Study | Selection bias | Study design | Confounding factors | Blinding | Data collection | Withdraws and drop-outs | Global rating* |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Hoevenaar-Blom et al ${ }^{19}$ | Weak | Moderate | Strong | NA | Moderate | Strong | Moderate |
| Koolhaas et al ${ }^{20}$ | Weak | Moderate | Strong | NA | Moderate | Moderate | Moderate |
| Armstrong et al ${ }^{21}$ | Moderate | Moderate | Moderate | NA | Strong | Weak | Moderate |
| Blond et al ${ }^{22}$ | Weak | Moderate | Strong | NA | Moderate | Moderate | Moderate |
| Andersen et al ${ }^{23}$ | Weak | Moderate | Strong | NA | Moderate | Strong | Moderate |
| Celis-Morales et al ${ }^{24}$ | Weak | Moderate | Strong | NA | Moderate | Strong | Moderate |
| Matthews et al ${ }^{25}$ | Strong | Moderate | Strong | NA | Moderate | Strong | Strong |
| Besson et al ${ }^{26}$ | Weak | Moderate | Moderate | NA | Moderate | Weak | Weak |
| Oja et al ${ }^{27}$ | Moderate | Moderate | Strong | NA | Moderate | Strong | Strong |
| Sahlquist et al ${ }^{28}$ | Moderate | Moderate | Strong | NA | Moderate | Moderate | Strong |
| Grøntved et a ${ }^{29}$ | Moderate | Moderate | Strong | NA | Moderate | Moderate | Strong |
| Laverty et a $\beta^{30}$ | Weak | Weak | Strong | NA | Moderate | NA | Weak |
| Wen et $a^{11}$ | Moderate | Weak | Strong | NA | Moderate | NA | Moderate |
| Østergaard et al ${ }^{32}$ | Moderate | Weak | Moderate | NA | Weak | NA | Weak |
| Bere et $\mathrm{a}^{33}$ | Weak | Moderate | Moderate | NA | Moderate | Weak | Weak |
| Sahlqvist et al ${ }^{34}$ | Weak | Weak | Strong | NA | Moderate | NA | Weak |
| Millett et a ${ }^{35}$ | Moderate | Weak | Strong | NA | Moderate | NA | Moderate |
| Berger ${ }^{36}$ | Weak | Weak | Moderate | NA | Weak | NA | Weak |
| Evenson et al ${ }^{37}$ | Moderate | Weak | Strong | NA | Moderate | NA | Moderate |
| Hu et al ${ }^{38}$ | Strong | Weak | Moderate | NA | Moderate | NA | Moderate |
| Ramirez-Velez et al ${ }^{39}$ | Strong | Weak | Moderate | NA | Moderate | NA | Moderate |

*Weak, moderate and strong indicated poor, moderate and high study quality, respectively. NA, not applicable.

## Risk of bias assessment

The included studies were assessed according to the Quality Assessment Tool of Quantitative Studies. ${ }^{18}$ SN and AR independently assessed each study. Any case of disagreement was resolved by discussion. The tool consists of six components: representativeness of the target group, study design, confounding factors, blinding of both assessors and participants, reliability and validity of measures and number of withdrawals and dropouts. Each component was rated 'weak', 'moderate' or 'strong' following a standardised rating system, where 'weak' and 'strong' indicates poor and high quality, respectively. Studies with no weak components were rated as 'strong', studies with one weak component were rated as 'moderate' and studies with more than one weak component were rated as 'weak'. For detailed information of distribution of study quality, se table 1. ${ }^{19-39}$

## Contact with authors

We (SN or LBA) contacted the corresponding author when there was a lack of clarity or when additional information was needed. ${ }^{39}$

## Data extraction and main analysis

Data extraction was conducted by SN based on the main estimate exposure, which was defined in accordance with the protocol as any cycling. Main outcomes were defined a priori as CVD mortality, CVD incidence and CVD risk factors. CVD and coronary heart disease were treated as CVD for both CVD mortality and CVD incidence. In studies where relative risk (RR) was presented with more than one model of adjustment, the most conservative estimate was included. If both CVD mortality and CVD incidence were reported, ${ }^{24}$ CVD incidence was included due to higher numbers of cases.

For single risk factors, each risk factor was included in the main estimate, but not when both 'overweight or obese' and
'obesity' were reported in a single study. In this case, only 'overweight or obese' was included due to higher numbers of cases. If studies only reported high and low dose or reported men and women separately or reported more than one level of dose, we meta-analysed each study and included the combined estimate (online supplementary table 2).

Among those 10 studies reporting either CVD mortality or CVD incidence only, the following was analysed: (1) CVD incidence and total cycling, ${ }^{24}$ (2) CVD incidence and estimated total cycling, ${ }^{20-22}$ (3) CVD mortality and estimated total cycling, ${ }^{28}$ (4) CVD mortality and estimated commuter cycling, ${ }^{256}$ (5) CVD mortality and total cycling ${ }^{23} 27$ and (6) CVD incidence and estimated commuter cycling. ${ }^{19}$ We included only the estimate of highest statistical power from each study. This was important to ensure that individuals were included in the meta-analysis only once.

## Data extraction subgroup analysis

Due to a wide range in reporting of exposure and outcomes, we classified exposure as total cycling or commuter cycling. Outcomes were classified by subgroups for CVD mortality, CVD incidence, grouped CVD risk factors, and single CVD risk factors. CVD risk factors were only analysed when reported by $\geq 2$ studies (online supplementary table 4 ). This resulted in subgroup analyses of (1) overweight or obese, (2) obesity, (3) hypertension, (4) HDL-cholesterol level and (5) triglyceride level. See table 2 for details of classifications of risk factors. We analysed hypertensive versus not hypertensive. All subgroups were analysed for men, women and men and women combined.

## Dose-response

Each study was individually recoded into low-dose and highdose cycling when possible. Low dose was defined as the lowest amount of cycling reported, and high dose was defined as
Table 2 Characteristics of included studies.

| Study | Design/cohort /countries | Type of cycling | Population | Dates/years of follow up | Total N | Incidence /death | Outcome | Prevalence of cycling (\%) Total/low/high | RR/OR (95\% CI) | Dose |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  |  |  |  | Low | High |
| Hoevenaar-Blom et al ${ }^{19}$ | Prospective cohort/MORGAN/ The Netherlands | Commuter | Men, women; Aged 20-65 y at baseline | 1993-2006/9.8 | 16442 | 923/NA | Incidence | 75\%/19\%/5\% | 0.82 (0.73 to 0.92) | Regular cycling | >2.5hour/wk |
| Koolhaas et al ${ }^{20}$ | Prospective cohort/Rotterdam study/ <br> The Netherlands | General | Men, women; aged $>55 \mathrm{y}$ at baseline | 1997-2012/10.3 | 5901 | 642/NA | Incidence | 58\%/32\%/26\% | 0.78 (0.67 to 0.91) | $13 \mathrm{~min} / \mathrm{day}$ | 51 min/day |
| Armstrong et al ${ }^{21}$ | Prospective cohort/ Million Women Studyl United Kingdom | Total | Women; Aged 55.9 (SD 4.8) $y$ at baseline | 1998/9 | 497857 | 6815/NA | Incidence | Not reported* | 0.84 (0.80 to 0.88) | >0-2 hour/wk | >2 hour/wk |
| Blond et al ${ }^{22}$ | Prospective cohort/ Diet, Cancer and Health/ Denmark | Overall, commuter | Men, women; Aged 50-65 y at baseline | 1993-2013/20 | 53723 | 2892/NA | Incidence | Not reported | 0.87 (0.82 to 0.93) | >0-2.5 hour/wk $\dagger$ | >2.5hour/wkt |
| Andersen ZJ et al ${ }^{23}$ | Prospective cohort/ Diet, Cancer, and Health/ Denmark | Commuter, leisure time | Men, women; Aged 50-65 y at baseline | 1993-2010/13 | 52061 | NA/1285 | Mortality | 68\%/NA/NA | 0.78 (0.69 to 0.88) | No dose reported. $3.2 \pm 3.4$ hour/wk |  |
| Celis-Morales et al ${ }^{24}$ | Prospective cohort/ UK Biobank/ United Kingdom | Commuter | Men, women; $40-69$ y at baseline | 2007-2014/5 | 263540 | 1110/496 | Incidence, mortality | $3 \% / N A / N A$ | 0.54 (0.33 to 0.88) | Short $\ddagger$ | Long $\ddagger$ |
| Matthews et al ${ }^{25}$ | Prospective cohort/ Shanghai Women's Health Studyl <br> China | Commuter | Women; <br> Aged $40-70 \mathrm{y}$ at baseline | 1997-2004/5.7 | 67143 | NA/251 | Mortality | NA/19\%/5\% | 0.72 (0.42 to 1.23) | 0-1-3.4 METh/day | >3.5 METh/day |
| Besson et al ${ }^{26}$ | Prospective cohort/ EPIC-Norfolk/ United Kingdom | Commuter | Men, women; Aged 45-79 y at baseline | 1993-2006/7 | 14903 | NA/370 | Mortality | NA/NA/NA | 0.77 (0.51 to 1.15) | <30 min/wk | > $30 \mathrm{~min} / \mathrm{wk}$ |
| Oja et al ${ }^{27}$ | Prospective cohort/ HSE \& SHeS/ England, Scotland | Any | Men, women; Aged 30-98 y at baseline | 1991-2008/9.2 | 75014 | NA/1909 | Mortality | 10\%/5\%/5\% | 0.93 (0.76 to 1.16) | min/wk low § | $\mathrm{min} / \mathrm{wk}$ high § |
| Sahlquist et al ${ }^{28}$ | Prospective cohort/ EPIC-Norfolk/ United Kingdom | Commuter, Total | Men, women; Aged 40-79 y at baseline | 1993-2011/15.3 | $\begin{aligned} & 22450 \\ & \text { Commuter; } \\ & 13346 \end{aligned}$ | NA/1639 | Mortality | Total: 30\%/NA/NA Commuter: NA/4\%/2\% | 0.86 (0.74 to 1.00) | 1-59 min/wk | >60 min/wk |
| Grøntved et al ${ }^{29}$ | Prospective cohort/ Västerbottens Health Survey/Sweden | Commuter | Men, women; Aged 43.5 y at baseline | 1990-2011/10 | 23732 |  | Risk factors の**†† $\ddagger \ddagger$ | 24\%/NA/NA | $\begin{aligned} & 1 ; 0.85(0.73 \text { to } 0.99) \\ & 2 ; 0.87(0.79 \text { to } 0.95) \\ & 3 ; 0.85(0.76 \text { to } 0.94) \end{aligned}$ |  |  |
| Laverty et a ${ }^{30}$ | Cross sectional/ Understanding society/ United Kingdom | Commuter | Men, women; Aged 16-65 y | NA | 20458 |  | Risk factors $\\|^{* *}$ | 3\%/NA/NA | $\begin{aligned} & 1 ; 0.63(0.53 \text { to } 0.75) \\ & 2 ; 0.76(0.56 \text { to } 1.01) \end{aligned}$ |  |  |
| Wen et $a^{31}$ | Cross-sectional/ <br> New south Wales Adult Health Surveyl Australia | Commuter | Men, women; Aged $\geq 16$ y | NA | 6832 |  | Risk factors 19 | 3\%-10\%/NA/NA | 1; 0.34 (0.13 to 0.89 ) |  |  |
| Østergaard et a ${ }^{32}$ | Cross sectional/ <br> NA/ <br> Denmark | Commuter | Men, women; Aged 12-16 y |  | 3847 |  | Risk factors 19 | 62\%/NA/NA | 1; 0.55 (0.42 to 0.72) |  |  |
| Bere et al ${ }^{33}$ | Longitudinal/ ENDORSE and Youth in Balance/ The Netherlands, Norway | Commuter | Men, women; Aged 13.2 y at baseline | 2005-2008/2 | 890 |  | Risk factors 919 | 48\%/NA/NA | 1; 0.44 (0.21 to 0.88) |  |  |

Table 2 Continued

| Study | Design／cohort ／countries | Type of cycling | Population | Dates／years of follow up | Total N | Incidence ／death | Outcome | Prevalence of cycling （\％）Total／low／high | RR／OR（95\％CI） | Dose |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  |  |  |  | Low | High |
| Sahlqvist et al ${ }^{34}$ | Cross－sectional／ Bicycle Victoria／ Australia | Commuter | Men，women； Aged $\geq 18 \mathrm{y}$ | NA | 1813 |  | Risk factors ${ }^{\text {d }}$ | 100\％／NA／NA | 1； 0.67 （0．50 to 0．90） |  |  |
| Millett et al ${ }^{35}$ | Cross－sectional／ Indian Migration Study／ India | Commuter | Men，women； Aged $\geq 18 \mathrm{y}$ | NA | 3902 |  | Risk factors $\\|^{* *}$ | 45\％－68\％／NA／NA | $\begin{aligned} & 1 ; 0.66(0.55 \text { to } 0.77) \\ & 2 ; 0.51(0.36 \text { to } 0.71) \end{aligned}$ |  |  |
| Berger et $\mathrm{l}^{\beta 6}$ | Cross－sectional／ TCCS／ United States | Commuter | Men，women； Aged 20－64 y | NA | 1450 |  | Risk factorsๆ＊＊††まキ | 100\％／NA／NA | $\begin{aligned} & 1 ; 0.69(0.58 \text { to } 0.82) \\ & 2 ; 0.67(0.50 \text { to } 0.90) \\ & 3 ; 0.72(0.59 \text { to } 0.88) \\ & 4 ; 0.85(0.67 \text { to } 1.07) \end{aligned}$ |  |  |
| Evenson et al ${ }^{37}$ | Cross－sectional／YRBS／ United States | Commuter | Men，women； Youth in 6th－12th grades | NA | 4448 |  | Risk factors ${ }^{\text {d }}$ | 13\％／NA／NA | 1； 0.71 （0．52 to 0．98） |  |  |
| Huet al ${ }^{38}$ | Cross－sectional／ NA／China | Commuter | Men，women； Aged 20－49 y | NA | 3708 |  | Risk factorst† $\ddagger$ | 11\％－19\％／NA／NA | $\begin{aligned} & 3 ; 0.71(0.52 \text { to } 0.98) \\ & 4 ; 0.90(0.66 \text { to } 1.23) \end{aligned}$ |  |  |
| Ramirez－Velez et a ${ }^{39}$ | Cross－sectional／ FUPRECOL／Colombia | Commuter | Men，women； Aged 9－17．9 y | NA | 1568 |  | Risk factorsttま\＃ | 23\％／NA／NA | $\begin{aligned} & 3 ; 1.06(0.81 \text { to } 1.37) \\ & 4 ; 1.03(0.83 \text { to } 1.23) \end{aligned}$ |  |  |

[^0]the highest dose reported (table 2, characteristics of included studies). For the study by Blond et al, ${ }^{22}$ low dose was generated after meta-analysis of low ( $>0-1 \mathrm{~h} /$ week) and moderately low (1-2.5 h/week) cycling. The dose-response relationship was analysed for total cycling and commuter cycling. When both CVD incidence and CVD mortality were reported, ${ }^{24}$ CVD incidence was included in the dose-response analysis.

We reanalysed the dose-response relationship in post-hoc analysis by redefining the criteria for low and high dose. First, we redefined the cut-off for high dose as $>1 \mathrm{~h} /$ week, then as $>2 \mathrm{~h} /$ week and finally we analysed at three dosage levels. ${ }^{21}$

## Statistics

In all analyses, we ensured that individuals were not analysed more than once for the same outcome, that is, 'overweight or obese' and 'obesity.' Due to this, studies were only included once for CVD incidence and CVD mortality but may have been included in different subgroup analyses or for equivalent CVD risk factors. For analyses of CVD incidence or CVD mortality, we calculated pooled RR or pooled HR. For analyses of each CVD risk factor, we calculated adjusted OR.

All analyses were performed in Stata v.12.1 (StataCorp LP, USA), using user-written commands described by Egger et al. ${ }^{40}$ The estimates are presented as multivariate adjusted RR (CVD incidence and CVD mortality) or OR (CVD risk factors) with 95\% CIs.

We used random effect models. ${ }^{40}$ Dose-response relationships and differences between sexes were analysed using meta-regression and presented as $\beta$-coefficients and $P$ values. Heterogeneity
was assessed using the $\mathrm{I}^{2}$ statistic, Q (Cochran's heterogeneity test) and $P$ value. The $I^{2}$ statistic was calculated using Stata based on Q and df .

$$
\mathrm{I}^{2}=100 \% \times(\mathrm{Q}-\mathrm{df}) / \mathrm{Q}
$$

As proposed by Higgins et al, ${ }^{41} \mathrm{I}^{2}$ describes the percentage of total variance across studies, with values between $0 \%$ and $100 \%$, where $0 \%$ indicates no heterogeneity. Negative values were set equal to zero. ${ }^{41}$ Heterogeneity was tested in all analyses, but should be interpreted with caution when few studies were analysed due to the possibility of false homogeneity. ${ }^{41}$

Following the rule of thumb described by Sterne et al, ${ }^{42}$ the test for funnel plot asymmetry was only used when there were more than nine studies in the meta-analysis (figure 2). Sensitivity analyses, tests for heterogeneity and regression analyses are presented in online supplementary table $5 \mathrm{a}-12 \mathrm{~b}$.

## Small-study effect

The small-study effect was investigated for the total estimate CVD using the 'metabias' and 'metainf' commands as described by Egger et al. ${ }^{40}$ We also performed subgroup analyses for study quality and for CVD incidence compared with CVD mortality.

## Role of the funding source

There was no funding source for this systematic review.

## RESULTS

In total, 38 studies fulfilled the primary inclusion criteria. As the present meta-analysis comprises dichotomous outcomes only, 17


Figure 2 Forest plot of the main analysis of cycling on CVD incidence (risk ratio), CVD mortality (risk ratio), and CVD risk factors (OR). *The combined random effect estimate was 0.783 (CI: 0.744 to 0.824 ) for CVD incidence, CVD mortality and CVD risk factors combined, indicated by the diamond in the bottom of the diagram. The combined estimate was statistically significant, but were moderately heterogeneous ( $l^{2}=58 \%$ ). From the top, the first ten studies are either CVD incidence or CVD mortality estimates, and the latter studies are CVD risk factors. See table 2 details of included studies.


Figure 3 Forest plot of sensitivity analysis of main analysis on CVD incidence and CVD mortality. Total cycling is indicated by blue colour, and commuter cycling is indicated by red colour. *The combined random RR was 0.840 ( $\mathrm{CI}: 0.812$ to $0.868, \mathrm{I}^{2}=0 \%$ ) for CVD Incidence and CVD mortality, indicated by the diamond in the bottom of the diagram. For CVD incidence the combined RR was 0.837 ( $0.797-0.880, I^{2}=30 \%$ ), and for mortality the combined RR was $0.827\left(0.761-0.899, I^{2}=0 \%\right)$. The inconsistent result of homogeneity is most likely due to few studies in the separate analysis.
studies with outcomes presented only as continuous variables were excluded. Thus, the present meta-analysis included 21 studies (figure 1). Data were reanalysed of high-density lipoprotein (HDL)-cholesterol and triglyceride levels from the study of Ramírez-Vélez et al ${ }^{39}$ due to lack of clarity.

In total, 1,069,034 individuals from eight different cohorts and four different countries were included in the analysis of CVD incidence and CVD mortality. The estimates were based on 12,382 incidents and 5950 deaths during a follow-up time of $9.8 \pm 4.9$ years. Further, 72,648 individuals from 10 countries were analysed for one or more CVD risk factors. figure 1 presents detailed information regarding the review process and exclusions. table 2 summarises the characteristics of the 21 included studies. ${ }^{19-39}$

## Main analysis of outcome

For the overall analysis of CVD incidence, CVD mortality and CVD risk factors, there was a significant total effect estimate of 0.78 ( $95 \% \mathrm{CI}: 0.74$ to $0.82, \mathrm{P}<0.001 ; \mathrm{I}^{2}=58 \%$, Q $\mathrm{P}<0.001$ ) (figure 2). The RR for CVD incidence was 0.84 ( $0.80-0.88$, $\mathrm{P}<0.001 ; \mathrm{I}^{2}=30 \%, \mathrm{Q} \mathrm{P}=0.22$ ). The RR for CVD mortality was 0.83 ( $0.76-0.90 ; \mathrm{P}<0.001 ; \mathrm{I}^{2}<0 \%, \mathrm{Q} \mathrm{P}=0.58$ ). The OR for CVD risk factors was 0.75 ( $0.68-0.82 ; \mathrm{P}<0.001 ; \mathrm{I}^{2}=64 \%$, Q $\mathrm{P}<0.001$ ).

## Sensitivity analysis of total cycling and commuter cycling in the main analysis

For total cycling, there was a RR of 0.80 ( $0.71-0.90, \mathrm{P}<0.001$; $\mathrm{I}^{2}=45 \%, \mathrm{Q} P=0.16$ ) for CVD incidence and a RR of 0.84 ( $0.71-0.99, \mathrm{P}=0.037 ; \mathrm{I}^{2}=53 \%, \mathrm{Q} \mathrm{P}=0.14$ ) for CVD mortality (figure 3). For commuter cycling, there was a RR of 0.86
(0.85-0.91, $\mathrm{P}<0.001 ; \mathrm{I}^{2}<0 \%, \mathrm{Q} P=0.33$ ) for CVD incidence, a RR of 0.84 ( $0.74-0.97, \mathrm{P}=0.014 ; \mathrm{I}^{2}<0 \%, \mathrm{Q} P=0.73$ ) for CVD mortality and an OR of 0.75 (0.69-0.82, $\mathrm{P}<0.001 ; \mathrm{I}^{2}=66 \%$, Q $\mathrm{P}<0.001$ ) for CVD risk factors (figure 3).

## Subgroup analysis of total cycling

CVD incidence and CVD mortality
When performing subgroup analysis of total cycling, we found a RR of 0.806 ( $0.741-0.877, \mathrm{P}<0.001 ; \mathrm{I}^{2}=41 \%, \mathrm{Q} \mathrm{P}=0.132$ ) for combined CVD incidence and CVD mortality. Subgroup analysis showed similar results when CVD incidence was analysed separately, with a RR of $0.800\left(0.712-0.899, \mathrm{P}<0.001 ; \mathrm{I}^{2}=45 \%\right.$, $\mathrm{Q} \mathrm{P}=0.162$ ). Matthews et al ${ }^{24}$ analysed women only, and no studies analysed men separately. No studies reported results for combined or single risk factors of total cycling, and thus, all analyses of risk factors were derived from commuter cycling; see online supplementary table 10a-12b for sex differences.

CVD risk factors only
No study reported total cycling and CVD risk factors.

## Subgroup analysis of commuter cycling CVD incidence, CVD mortality and CVD risk factors

A total of 46 different estimates were reported for commuter cycling. When CVD incidence, CVD mortality and CVD risk factors were combined, there was a RR of 0.77 ( $0.73-0.82$, $\mathrm{P}<0.001 ; \mathrm{I}^{2}=53 \%, \mathrm{Q} \mathrm{P}<0.001$ ). Subgroup analysis including only CVD incidence gave a RR of 0.859 (0.814-0.907, $\mathrm{P}<0.001$; $\mathrm{I}^{2}<0 \%, \mathrm{Q} P=0.465$ ); see online supplementary table $12 \mathrm{a}-\mathrm{b}$.


Figure 4 Forest plot of sensitivity analysis of CVD risk factors for commuter cycling. *Combined OR was 0.749 ( $0.689-0.815, I^{2}=54 \%$ ) indicated by the diamond in the bottom. Red boxes indicates overweight or obese, blue box indicates hypertension, green box indicates triglycerides and yellow box indicates HDL. All risk factors independently beside HDL were significant. For detailed information of each outcome see table 6a-b in online supplementary tables.

## CVD risk factors only

CVD risk factors were reported for commuter cycling. Overweight and obesity were the most commonly reported risk factors (figure 4), and were classified according to WHO. ${ }^{43}$ In total, 'overweight or obese' or 'obesity' were reported 14 times. When analysing 'overweight or obese' and 'obesity,' there was an OR of 0.633 (0.574-0.669, $\left.\mathrm{P}<0.001 ; \mathrm{I}^{2}<0 \%, \mathrm{Q} \mathrm{P}=0.814\right)$ and OR 0.722 ( $0.631-0.826, \mathrm{P}<0.001 ; \mathrm{I}^{2}=29 \%, \mathrm{Q} \mathrm{P}=0.204$ ), respectively. There was an OR of 0.714 ( $0.566-0.900, \mathrm{P}=0.004$; $\mathrm{I}^{2}=72 \%, \mathrm{Q} \mathrm{P}=0.014$ ) for hypertension, 0.827 ( $0.712-0.961$, $\mathrm{P}=0.013 ; \mathrm{I}^{2}=52 \%, \mathrm{Q} \mathrm{P}=0.098$ ) for triglyceride level and 0.983 ( $0.822-1.176, \mathrm{P}=0.855 ; \mathrm{I}^{2}<0 \%, \mathrm{Q} \mathrm{P}=0.502$ ) for HDL-cholesterol level. Triglyceride level remained significant only when analysing men and women combined. HDL-cholesterol was the only risk factor not significant for men, women, or combined.

There was no dose-response relationship for total cycling, commuter cycling or combined total and commuter cycling (online supplementary table 7a-9b). All post-hoc analyses remained nonsignificant (coefficient $=-0.010-0.002$, $\mathrm{P}=0.648-0.909$ ).

## Small study effects

There was a significant small study effect, indicating possible publication bias (online supplementary figure 1-2).

## DISCUSSION

Cycling was associated with a $16 \%$ lower risk of CVD incidence, $17 \%$ lower risk of CVD mortality and a $25 \%$ lower risk of CVD risk factors. When CVD incidence and mortality were combined, cycling was associated with a $22 \%$ lower risk. However, the main analysis was heterogeneous ( $\mathrm{I}^{2}=58 \%$ ), possibly because
we included cross-sectional and prospective studies of populations of children and adults. To assess CVD incidence and CVD mortality, we analysed prospective cohort studies of adult populations.

Our results support those of a previous study of approximately 173,000 adults - that active transportation, especially cycling, reduces CVD risk. ${ }^{15}$ We analysed an almost 10 -fold larger population and included only cycling as an activity. Our results were slightly more consistent, and we found a stronger association for cycling compared with studies combining walking and cycling. Our results should be of interest for policy-makers and politicians, since they provide evidence of the protective effect of cycling on CVD.

## CVD risk factors

In our systematic review, the most commonly reported and most frequently reduced risk factor was overweight or obesity. In a scoping review, Brown et al ${ }^{16}$ found a small but significant reduction in body mass index with active transportation, but concluded that the effect might be smaller than indicated in the literature. However, in contrast, we found a $36 \%$ lower risk in cyclists for both overweight and obesity (OR 0.64 , CI: 0.58 to $0.70, \mathrm{I}^{2}=0 \%$ ) combined, and a $27 \%$ lower risk for obesity (OR 0.73 , CI: 0.57 to $\left.0.94, \mathrm{I}^{2}=66 \%\right)$. The relatively low heterogeneity could be erroneous, due to a smaller number of studies. ${ }^{41}$ Therefore, it is possible that our results overestimate the risk reduction associated with cycling. However, our main analysis is supported by our subgroup analysis of commuter cycling and CVD risk factors (online supplementary table 12a-b), adding strength to our conclusions.

Hypertension was the second most reduced risk factor (OR 0.71 , CI: 0.57 to 0.90 ). Two studies ${ }^{30} 36$ defined hypertension based on a self-reported diagnosis by a physician, while Grøntved et al ${ }^{29}$ used systolic and diastolic blood pressure of $>140$ and $>90 \mathrm{~mm} \mathrm{Hg}$, respectively, or usage of antihypertensive medications. Further, risk of high triglyceride level was reduced by $18 \%$ for commuter cycling compared with that of passive commuters. Finally, HDL-cholesterol level was the only non-significant, homogeneous risk factor. Cycling therefore seems to be associated with an enhanced CVD profile and thus cycling may be able to prevent CVD incidence or CVD mortality.

## Sex differences

In contrast to a previous meta-analysis, ${ }^{15}$ we found no evidence that women experienced a greater effect from cycling compared with that of men. In our systematic review, CVD incidence and CVD mortality results were mainly presented in both sexes combined, whereas CVD risk factor results more often included a sex-specific analysis. There was a tendency for women to have greater risk reduction for both high triglyceride and HDL-cholesterol levels compared with men (online supplementary table 10a-12b).

## Dose-response relationship

In contrast to previous suggestions, ${ }^{112}$ we found no difference between low-dose and high-dose cycling. Increased cycling dose was associated with lower CVD risk, especially for commuter cycling and CVD mortality. This is in accordance with the finding of Kelly et al, ${ }^{11}$ where the steepest risk reduction for all-cause mortality was for $0-101$ min per week of cycling, but with further reduction in risk among those cycling $>101 \mathrm{~min}$ per week.

When analysing the dose-response relationship, there were several challenges. First, we divided each study individually into either high or low doses based on the amount of cycling reported in each study. This resulted in heterogeneity of the definition of low and high dose: high dose in some studies ${ }^{26} 28$ was similar to low dose in other studies (See table 2 for details). Second, there were few individuals in high-dose groups compared with those in low-dose groups; this was due to the low prevalence of cycling in general and a lower prevalence of high-dose cycling. Therefore, the results regarding the dose-response relationship should be interpreted with caution. We encourage researchers to be more consistent when creating categories for cycling doses and to report data, including that of low prevalence, in each category.

## Strength and limitations

One of the greatest challenges of analysing cycling behaviour is that cycling is not a singular behaviour - often individuals engage in multiple physical activities. This means that people engaged in other forms of activities may be more likely to choose active transport as well. Even though 15 of 21 included studies adjusted for other physical activities, there may be residual confounding from leisure-time physical activity. In addition, in included studies with a low prevalence of cycling, cyclists may be a select group of individuals with superior health (and lower CVD risk profile). However, the majority of included studies adjusted for smoking status, alcohol consumption and level of education (see online supplementary Table 13 for details of adjustments).

Cycling and walking have different benefits such as an increased amount of vigorous activity ${ }^{12}$; therefore, cycling
might be more protective than walking. Forty five studies were excluded due to merged groups of walking and cycling. This might be because few of the included studies were designed to evaluate the effect of cycling but rather aimed to register activity levels in large populations. If studies were not primarily designed to investigate the independent association of cycling and CVD, this may explain the publication bias we found in our funnel plot.

All studies used self-reported measurements of cycling and aimed to register physical activity levels. Self-report measurements may have recall bias, and social desirability bias by over-reporting of activity and underestimation of body weight. There was evidence for a small-study effect, and studies of negative results were less likely to be published. ${ }^{40}$ This may have influenced our results by increasing the possibility that we overestimated the true association between cycling and CVD. On the other hand, the main analysis was primarily based on high-quality studies that consistently reported positive associations between cycling and reduction in CVD incidence and mortality. However, the results were less certain for the association between cycling and CVD risk factors since the studies included in those analyses were of moderate and low quality.

## CONCLUSION

Cyclists had lower risk of CVD incidence, CVD mortality and some CVD risk factors. Similar lower risk of CVD were observed for men and women. Health professionals, city planners and stakeholders can recommend cycling to prevent CVD and should aim to increase the amount of any cycling.

Acknowledgements We thank senior librarian Anita Svedal as well as Lise VikHaugen for considerable support during the literature search.
Contributors All authors contributed to the design of the study and reviewed the report. SN and LBA generated the hypotheses. $S N$ and AR performed the literature search. SN, AR and LBA analysed the data. SN wrote the first draft of the manuscript. LBA, AKS and AR revised the manuscript critically for important intellectual content. All authors, external and internal, had full access to all of the data (including statistical reports and tables) in the study and can take responsibility for the integrity of the data and the accuracy of the data analysis. LBA is the study guarantor.
Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.
Competing interests None declared.
Provenance and peer review Not commissioned; externally peer reviewed.

## REFERENCES

1 Lozano R, Naghavi M, Foreman K, et al. Global and regional mortality from 235 causes of death for 20 age groups in 1990 and 2010: a systematic analysis for the Global Burden of Disease Study 2010. Lancet 2012;380:2095-128.
2 Andersen LB, Mota J, Di Pietro L. Update on the global pandemic of physical inactivity. Lancet 2016;388:1255-6.
3 GBD 2016 Causes of Death Collaborators. Global, regional, and national age-sex specific mortality for 264 causes of death, 1980-2016: a systematic analysis for the Global Burden of Disease Study 2016. Lancet 2017;390:1151-210.
4 World Health Organization. Global recommendations on physical activity for health, 2010.

5 Warburton DE, Charlesworth S, Ivey A, et al. A systematic review of the evidence for Canada's Physical Activity Guidelines for Adults. Int J Behav Nutr Phys Act 2010;7:39.
6 World Health Organization. Global health risks: mortality and burden of disease attributable to selected major risks, 2009.
7 Sallis JF, Bull F, Guthold R, et al. Progress in physical activity over the Olympic quadrennium. Lancet 2016;388:1325-36.
8 Hallal PC, Andersen LB, Bull FC, et al. Global physical activity levels: surveillance progress, pitfalls, and prospects. Lancet 2012;380:247-57.
9 Reis RS, Salvo D, Ogilvie D, et al. Scaling up physical activity interventions worldwide: stepping up to larger and smarter approaches to get people moving. Lancet 2016;388:1337-48.

10 Smith M, Hosking J, Woodward A, et al. Systematic literature review of built environment effects on physical activity and active transport - an update and new findings on health equity. Int J Behav Nutr Phys Act 2017;14:158.
11 Kelly P, Kahlmeier S, Götschi T, et al. Systematic review and meta-analysis of reduction in all-cause mortality from walking and cycling and shape of dose response relationship. Int J Behav Nutr Phys Act 2014;11:132.
12 Oja P, Titze S, Bauman A, et al. Health benefits of cycling: a systematic review. Scand J Med Sci Sports 2011;21:496-509.
13 Oja P, Mänttäri A, Heinonen A, et al. Physiological effects of walking and cycling to work. Scand J Med Sci Sports 1991;1:151-7.
14 Tanasescu M, Leitzmann MF, Rimm EB, et al. Exercise type and intensity in relation to coronary heart disease in men. JAMA 2002;288:1994-2000.
15 Hamer M, Chida Y. Active commuting and cardiovascular risk: a meta-analytic review. Prev Med 2008;46:9-13.
16 Brown V, Moodie M, Mantilla Herrera AM, et al. Active transport and obesity prevention - A transportation sector obesity impact scoping review and assessment for Melbourne, Australia. Prev Med 2017;96:49-66.
17 Moher D, Liberati A, Tetzlaff J, et al. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. PLoS Med 2009;6:e1000097.
18 Thomas BH, Ciliska D, Dobbins M, et al. A process for systematically reviewing the literature: providing the research evidence for public health nursing interventions. Worldviews Evid Based Nurs 2004;1:176-84.
19 Hoevenaar-Blom MP, Wendel-Vos GC, Spijkerman AM, et al. Cycling and sports, but not walking, are associated with 10-year cardiovascular disease incidence: the MORGEN Study. Eur J Cardiovasc Prev Rehabil 2011;18:41-7.
20 Koolhaas CM, Dhana K, Golubic R, et al. Physical Activity Types and Coronary Heart Disease Risk in Middle-Aged and Elderly Persons: The Rotterdam Study. Am J Epidemiol 2016;183:729-38.
21 Armstrong ME, Green J, Reeves GK, et al. Frequent physical activity may not reduce vascular disease risk as much as moderate activity: large prospective study of women in the United Kingdom. Circulation 2015;131:721-9.
22 Blond K, Jensen MK, Rasmussen MG, et al. Prospective Study of Bicycling and Risk of Coronary Heart Disease in Danish Men and Women. Circulation 2016;134:1409-11.
23 Andersen ZJ, de Nazelle A, Mendez MA, et al. A study of the combined effects of physical activity and air pollution on mortality in elderly urban residents: the Danish Diet, Cancer, and Health Cohort. Environ Health Perspect 2015;123:557-63.
24 Celis-Morales CA, Lyall DM, Welsh P, et al. Association between active commuting and incident cardiovascular disease, cancer, and mortality: prospective cohort study. BMJ 2017;357:j1456.
25 Matthews CE, Jurj AL, Shu XO, et al. Influence of exercise, walking, cycling, and overall nonexercise physical activity on mortality in Chinese women. Am J Epidemiol 2007;165:1343-50.
26 Besson H, Ekelund U, Brage S, et al. Relationship between subdomains of total physical activity and mortality. Med Sci Sports Exerc 2008;40:1909-15.

27 Oja P, Kelly P, Pedisic Z, et al. Associations of specific types of sports and exercise with all-cause and cardiovascular-disease mortality: a cohort study of 80306 British adults. Br J Sports Med 2017;51:812-7.
28 Sahlqvist S, Goodman A, Simmons RK, et al. The association of cycling with all-cause, cardiovascular and cancer mortality: findings from the population-based EPIC-Norfolk cohort. BMJ Open 2013;3:e003797.
29 Grontved A, Koivula RW, Johansson I, et al. Bicycling to Work and Primordial Prevention of Cardiovascular Risk: A Cohort Study Among Swedish Men and Women. J Am Heart Assoc 2016;5:e004413.
30 Laverty AA, Mindell JS, Webb EA, et al. Active travel to work and cardiovascular risk factors in the United Kingdom. Am J Prev Med 2013;45:282-8.
31 Wen LM, Rissel C. Inverse associations between cycling to work, public transport, and overweight and obesity: findings from a population based study in Australia. Prev Med 2008;46:29-32.
32 Østergaard L, Grøntved A, Børrestad LA, et al. Cycling to school is associated with lower BMI and lower odds of being overweight or obese in a large population-based study of Danish adolescents. J Phys Act Health 2012;9:617-25.
33 Bere E, Oenema A, Prins RG, et al. Longitudinal associations between cycling to school and weight status. Int J Pediatr Obes 2011;6(3-4):182-7.
34 Sahlqvist SL, Heesch KC. Characteristics of utility cyclists in Queensland, Australia: an examination of the associations between individual, social, and environmental factors and utility cycling. J Phys Act Health 2012;9:818-28.
35 Millett C, Agrawal S, Sullivan R, et al. Associations between active travel to work and overweight, hypertension, and diabetes in India: a cross-sectional study. PLoS Med 2013;10:e1001459.
36 Berger AT, Qian XL, Pereira MA. Associations Between Bicycling for Transportation and Cardiometabolic Risk Factors Among Minneapolis-Saint Paul Area Commuters: A Cross-Sectional Study in Working-Age Adults. Am J Health Promot 2018;32:890117117710735.
37 Evenson KR, Huston SL, McMillen BJ, et al. Statewide prevalence and correlates of walking and bicycling to school. Arch Pediatr Adolesc Med 2003;157:887-92.
38 Hu G, Pekkarinen H, Hänninen O, et al. Relation between commuting, leisure time physical activity and serum lipids in a Chinese urban population. Ann Hum Biol 2001;28:412-21.
39 Ramírez-Vélez R, García-Hermoso A, Agostinis-Sobrinho C, et al. Cycling to School and Body Composition, Physical Fitness, and Metabolic Syndrome in Children and Adolescents. J Pediatr 2017;188:57-63.
40 Egger M, Davey-Smith G, Altman D, eds. Systematic reviews in health care: metaanalysis in context. 2nd ed: John Wiley \& Sons, 2008.
41 Higgins JP, Thompson SG, Deeks JJ, et al. Measuring inconsistency in meta-analyses. BMJ 2003;327:557-60.
42 Sterne JA, Sutton AJ, loannidis JP, et al. Recommendations for examining and interpreting funnel plot asymmetry in meta-analyses of randomised controlled trials. BMJ 2011;343:d4002.
43 World Health Organization. Obesity: preventing and managing the global epidemic: WorldHealth Organization, 2000.


[^0]:    Risk factors．
    ＊States that cycling is infrequent in this cohort．
    †Commuter cycling：Low dose $=0-1.5$ hour／week；High dose $>1.5$ hour／week
    $\ddagger$ Split into groups according to distance．
    §Groups defined by using the sex－specific medians．
    §Groups defined by using the sex－specific medians．
    II）
    ${ }_{* *}$ Hypertension（self－reported or doctor－diagnosed ${ }^{30} 36$ or systolic blood pressure or diastolic blood pressure $>140 \mathrm{and}>90 \mathrm{~mm} \mathrm{Hg}$ ，and／or use of antihypertensive medications．${ }^{29}$
    $\dagger \dagger$ ）Hypertriglyceridemia（ $>1.7 \mathrm{mmol} / \mathrm{L}^{29}$ self－reported or doctor diagnosed，${ }^{36}$ adverse $\log$ transformed scale，${ }^{38}$ or＇high triglycerides＇not defined．${ }^{39}$
    $\ddagger \ddagger)$ Low high－density lipoprotein level（self－reported or doctor－diagnosed，${ }^{36}$ adverse
    METh，metabolic equivalent hours．；NA，not applicable；RR，relative risk；wk，week．

