STROBE Statement-checklist of items that should be included in reports of observational studies

	Item No.	Recommendation	Page No.	Relevant text from manuscript
Title and abstract	1	(a) Indicate the study's design with a commonly used term in the title or the abstract		1
		(b) Provide in the abstract an informative and balanced summary of what was done and what was found		1
Introduction				
Background/rationale	2	Explain the scientific background and rationale for the investigation being reported		3
Objectives	3	State specific objectives, including any prespecified hypotheses		4
Methods				
Study design	4	Present key elements of study design early in the paper		1,4
Setting	5	Describe the setting, locations, and relevant dates, including periods of recruitment, exposure, follow-up, and data collection		4, 5; Supplementary file page 2
Participants	6	( <i>a</i> ) <i>Cohort study</i> —Give the eligibility criteria, and the sources and methods of selection of participants. Describe methods of follow-up		4, 5; Supplementary file page 2
		Case-control study-Give the eligibility criteria, and the sources and methods of case		
		ascertainment and control selection. Give the rationale for the choice of cases and controls		
		<i>Cross-sectional study</i> —Give the eligibility criteria, and the sources and methods of selection of participants		
		(b) Cohort study—For matched studies, give matching criteria and number of exposed and unexposed		Table 1, Table S1
		<i>Case-control study</i> —For matched studies, give matching criteria and the number of controls per case		
Variables	7	Clearly define all outcomes, exposures, predictors, potential confounders, and effect modifiers. Give diagnostic criteria, if applicable		4, 5; Supplementary file page 2
Data sources/	8*	For each variable of interest, give sources of data and details of methods of assessment		4-6; Supplementary file page 2,
measurement		(measurement). Describe comparability of assessment methods if there is more than one group		3
Bias	9	Describe any efforts to address potential sources of bias		6, Supplementary file page 4
Study size	10	Explain how the study size was arrived at		5

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Quantitative	11	Explain how quantitative variables were handled in the analyses. If applicable, describe which	7
variables		groupings were chosen and why	
Statistical	12	(a) Describe all statistical methods, including those used to control for confounding	7
methods		(b) Describe any methods used to examine subgroups and interactions	7
		(c) Explain how missing data were addressed	Not applicable
		(d) Cohort study—If applicable, explain how loss to follow-up was addressed	Not applicable
		Case-control study—If applicable, explain how matching of cases and controls was addressed	
		Cross-sectional study—If applicable, describe analytical methods taking account of sampling	
		strategy	
		( <u>e</u> ) Describe any sensitivity analyses	Not applicable
Results			
Participants	13*	(a) Report numbers of individuals at each stage of study-eg numbers potentially eligible, examined	Not applicable (post mortem study)
-		for eligibility, confirmed eligible, included in the study, completing follow-up, and analysed	
		(b) Give reasons for non-participation at each stage	Not applicable (post mortem study)
		(c) Consider use of a flow diagram	Not applicable (post mortem study)
Descriptive data	14*	(a) Give characteristics of study participants (eg demographic, clinical, social) and information on	Table 1, Table S1
		exposures and potential confounders	
		(b) Indicate number of participants with missing data for each variable of interest	Table 1, Table S1
		(c) Cohort study—Summarise follow-up time (eg, average and total amount)	Not applicable (post mortem study)
Outcome data	15*	Cohort study-Report numbers of outcome events or summary measures over time	Not applicable (post mortem study)
		Case-control study-Report numbers in each exposure category, or summary measures of exposure	
		Cross-sectional study—Report numbers of outcome events or summary measures	
Main results	16	(a) Give unadjusted estimates and, if applicable, confounder-adjusted estimates and their precision	7, 8,9; Table 2
		(eg, 95% confidence interval). Make clear which confounders were adjusted for and why they were	
		included	
		(b) Report category boundaries when continuous variables were categorized	Not applicable
		(c) If relevant, consider translating estimates of relative risk into absolute risk for a meaningful time	Not applicable
		period	

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Other analyses	17	Report other analyses done-eg analyses of subgroups and interactions, and sensitivity analyses	Not applicable
Discussion			
Key results	18	Summarise key results with reference to study objectives	1,11
Limitations	19	Discuss limitations of the study, taking into account sources of potential bias or imprecision. Discuss	14-15
		both direction and magnitude of any potential bias	
Interpretation	20	Give a cautious overall interpretation of results considering objectives, limitations, multiplicity of	2, 13-15; Graphic abstract
		analyses, results from similar studies, and other relevant evidence	
Generalisability	21	Discuss the generalisability (external validity) of the study results	15
Other information	on		
Funding	22	Give the source of funding and the role of the funders for the present study and, if applicable, for the	16
		original study on which the present article is based	

\*Give information separately for cases and controls in case-control studies and, if applicable, for exposed and unexposed groups in cohort and cross-sectional studies.

**Note:** An Explanation and Elaboration article discusses each checklist item and gives methodological background and published examples of transparent reporting. The STROBE checklist is best used in conjunction with this article (freely available on the Web sites of PLoS Medicine at http://www.plosmedicine.org/, Annals of Internal Medicine at http://www.annals.org/, and Epidemiology at http://www.epidem.com/). Information on the STROBE Initiative is available at www.strobe-statement.org.