Supplemental Table S2. Checklist. STROBE Statement.

	Item No	Recommendation	Page Number
Title and abstract	1	(a) Indicate the study's design with a commonly used	2; Abstract,
		term in the title or the abstract	paragraph 2
		(b) Provide in the abstract an informative and balanced	2; Abstract,
		summary of what was done and what was found	paragraph 2
Introduction			
Background/rationale	2	Explain the scientific background and rationale for the	6; Introduction,
		investigation being reported	paragraph 6
Objectives	3	State specific objectives, including any prespecified	6; Introduction,
		hypotheses	paragraph 6
Methods			
Study design	4	Present key elements of study design early in the paper	6; Methods,
			paragraphs 1 - 3
Setting	5	Describe the setting, locations, and relevant dates,	6; Methods,
		including periods of recruitment, exposure, follow-up,	paragraphs 1, 2,
		and data collection	6
Participants	6	(a) Give the eligibility criteria, and the sources and	6; Methods,
		methods of selection of participants. Describe methods	paragraph 2
		of follow-up	
		(b) For matched studies, give matching criteria and	N/A
		number of exposed and unexposed	
Variables	7	Clearly define all outcomes, exposures, predictors,	6; Methods,
		potential confounders, and effect modifiers. Give	paragraph 2
		diagnostic criteria, if applicable	
Data sources/	8*	For each variable of interest, give sources of data and	6; Methods,
measurement		details of methods of assessment (measurement).	paragraphs 2, 4,
		Describe comparability of assessment methods if there	5
		is more than one group	
Bias	9	Describe any efforts to address potential sources of	14; Discussion,
		bias	paragraph 8
Study size	10	Explain how the study size was arrived at	8; Results,
			paragraph 1
Quantitative variables	11	Explain how quantitative variables were handled in the	7; Methods,
		analyses. If applicable, describe which groupings were	paragraph 6
		chosen and why	
Statistical methods	12	(a) Describe all statistical methods, including those	7; Methods,
		used to control for confounding	paragraph 6
		(b) Describe any methods used to examine subgroups	7; Methods,
		and interactions	paragraph 6
		(c) Explain how missing data were addressed	8; Results,
			paragraph 1
		(d) If applicable, explain how loss to follow-up was addressed	N/A
		(<u>e</u>) Describe any sensitivity analyses	N/A
Results		,	

Participants	13*	(a) Report numbers of individuals at each stage of study—eg numbers potentially eligible, examined for eligibility, confirmed eligible, included in the study,	8; Results, paragraph 1
		completing follow-up, and analysed	21/2
		(b) Give reasons for non-participation at each stage	N/A
		(c) Consider use of a flow diagram	N/A
Descriptive data	14*	(a) Give characteristics of study participants (eg	8; Results,
		demographic, clinical, social) and information on	paragraph 2
		exposures and potential confounders	
		(b) Indicate number of participants with missing data	8; Results,
		for each variable of interest	paragraph 1
		(c) Summarise follow-up time (eg, average and total amount)	N/A
Outcome data	15*	Report numbers of outcome events or summary	8; Results,
		measures over time	paragraph 3
Main results	16	(a) Give unadjusted estimates and, if applicable,	8; Results,
		confounder-adjusted estimates and their precision (eg,	paragraph 3, 4,
		95% confidence interval). Make clear which	5, 6
		confounders were adjusted for and why they were	
		included	
		(b) Report category boundaries when continuous	N/A
		variables were categorized	
		(c) If relevant, consider translating estimates of relative	N/A
		risk into absolute risk for a meaningful time period	
Other analyses	17	Report other analyses done—eg analyses of subgroups	9; Results,
		and interactions, and sensitivity analyses	paragraph 5, 6
Discussion			
Key results	18	Summarise key results with reference to study	11; Discussion,
		objectives	paragraph 1
Limitations	19	Discuss limitations of the study, taking into account	14; Discussion,
		sources of potential bias or imprecision. Discuss both	paragraph 8
		direction and magnitude of any potential bias	
Interpretation	20	Give a cautious overall interpretation of results	14; Conclusion,
		considering objectives, limitations, multiplicity of	paragraph 1
		analyses, results from similar studies, and other	
		relevant evidence	
Generalisability	21	Discuss the generalisability (external validity) of the	13; Discussion,
		study results	paragraph 6
Other information		· · · · · · · · · · · · · · · · · · ·	
Funding	22	Give the source of funding and the role of the funders	N/A
		for the present study and, if applicable, for the original	
		study on which the present article is based	

^{*}Give information separately for exposed and unexposed groups.