	ICMJE DISCLOSURE FORM				
Da	Date: 4/12/2023				
Yo	Your Name: Bettina Mittendorfer				
Ma	anuscript Title:	Beta-cell function after Roux-en-Y gastric bypass surgery or reduintake alone in people with obesity	Beta-cell function after Roux-en-Y gastric bypass surgery or reduced energy intake alone in people with obesity		
Ma	anuscript Number (if kı	nown):170307-INS-CMED-2	170307-INS-CMED-2		
cor aff ind The epi tha	In the interest of transparency, we ask you to disclose all relationships/activities/interests listed below that are related to the content of your manuscript. "Related" means any relation with for-profit or not-for-profit third parties whose interests may be affected by the content of the manuscript. Disclosure represents a commitment to transparency and does not necessarily indicate a bias. If you are in doubt about whether to list a relationship/activity/interest, it is preferable that you do so. The author's relationships/activities/interests should be defined broadly. For example, if your manuscript pertains to the epidemiology of hypertension, you should declare all relationships with manufacturers of antihypertensive medication, even if that medication is not mentioned in the manuscript. In item #1 below, report all support for the work reported in this manuscript without time limit. For all other items, the time frame for disclosure is the past 36 months.				
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		Time frame: Since the initial planning of the work			
1	All support for the	□ None			

NIH grants R01 DK037948, R01 DK101578, manuscript (e.g., The payments were made to the institution funding, provision P30 DK056341, P30 DK020579, and UL1 and the funding sources had no role in the TR002345, and grants from the American design and conduct of the study; collection, of study materials, Diabetes Association (1-18-ICTS-119), and management, analysis, and interpretation of medical writing, the Longer Life Foundation (2019-011). the data; preparation, review, or approval of article processing the manuscript; and decision to submit the charges, etc.) manuscript for publication. No time limit for this item. Click the tab key to add additional rows Time frame: past 36 months 2 Grants or None contracts from any entity (if not indicated in item #1 above). 3 Royalties or \boxtimes None licenses

		Name all entities with whom you have this relationship or indicate none (add rows as needed)	Specifications/Comments (e.g., if payments were made to you or to your institution)
4	Consulting fees	None None	
5	Payment or honoraria for lectures, presentations, speakers bureaus, manuscript writing or educational events	None	
6	Payment for expert testimony	None	
7	Support for attending meetings and/or travel	None	
8	Patents planned, issued or pending	None	
9	Participation on a Data Safety Monitoring Board or Advisory Board	None	
10	Leadership or fiduciary role in other board, society, committee or advocacy group, paid or unpaid	None	

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11	Stock or stock options	[⊠] None	
12	Receipt of equipment, materials, drugs, medical writing, gifts or other services	[⊠] None	
13	Other financial or non-financial interests	⊠ None	
Plea	Please place an "X" next to the following statement to indicate your agreement: I certify that I have answered every question and have not altered the wording of any of the questions on this form.		

ICMJE DISCLOSURE FORM			
Date:		4/12/2023	
Your Name:		Bruce W. Patterson	
Manuscript Title:		Beta-cell function after Roux-en-Y gastric bypass surgery or reduced energy intake alone in people with obesity	
Manuscript Number (if	known):	170307-INS-CMED-2	
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	#1 above).		
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ICMJE DISCLOSURE FORM

ICIVIJE DISCLOSORE FORIVI		
Date:	4/12/2023	
Your Name:	Faidon Magkos	
Manuscript Title:	Beta-cell function after Roux-en-Y gastric bypass surgery or reduced energy intake alone in people with obesity	
Manuscript Number (if known):	170307-INS-CMED-2	
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ICMJE DISCLOSURE FORM			
Date:		4/12/2023	
Your Name:		Mihoko Yoshino	
Manuscript Title:		Beta-cell function after Roux-en-Y gastric bypass surgery or reduced energy intake alone in people with obesity	
Manuscript Number (if I	known):	170307-INS-CMED-2	
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ICMJE DISCLOSURE FORM

Date:	_4/12/2023		
Your Name:	David P. Bradley Beta-cell function after Roux-en-Y gastric bypass surgery or reduced energy intake alone in people with obesity		
Manuscript Title:			
Manuscript Number (if known):	Manuscript Number (if known): 170307-INS-CMED-2		
In the interest of transparency, we ask you to disclose all relationships/activities/interests listed below that are related to the content of your manuscript. "Related" means any relation with for-profit or not-for-profit third parties whose interests may be affected by the content of the manuscript. Disclosure represents a commitment to transparency and does not necessarily indicate a bias. If you are in doubt about whether to list a relationship/activity/interest, it is preferable that you do so.			
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ICMJE DISCLOSURE FORM

Date:	4/12/2023		
Your Name:	J. Christopher Eagon		
Manuscript Title:	Beta-cell function after Roux-en-Y gastric bypass surgery or reduced energy intake alone in people with obesity		
Manuscript Number (if known): 170307-INS-CMED-2			
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ICMJE DISCLOSURE FORM			
Date:		4/12/2023	
Your Name:		Samuel Klein	
Manuscript Title:		Beta-cell function after Roux-en-Y gas intake alone in people with obesity	stric bypass surgery or reduced energy
Manuscript Number (if known): _ 170307-INS-CMED-2			
content of your manuscr affected by the content indicate a bias. If you ar The author's relationship epidemiology of hyperte that medication is not m In item #1 below, report	In the interest of transparency, we ask you to disclose all relationships/activities/interests listed below that are related to the content of your manuscript. "Related" means any relation with for-profit or not-for-profit third parties whose interests may be affected by the content of the manuscript. Disclosure represents a commitment to transparency and does not necessarily indicate a bias. If you are in doubt about whether to list a relationship/activity/interest, it is preferable that you do so. The author's relationships/activities/interests should be defined broadly. For example, if your manuscript pertains to the epidemiology of hypertension, you should declare all relationships with manufacturers of antihypertensive medication, even if that medication is not mentioned in the manuscript. In item #1 below, report all support for the work reported in this manuscript without time limit. For all other items, the time frame for disclosure is the past 36 months.		
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STROBE Statement—checklist of items that should be included in reports of observational studies

	Item No	Recommendation	Page number
Title and abstract	1	(a) Indicate the study's design with a commonly used term in the title or the abstract	1 and 2
		(b) Provide in the abstract an informative and balanced summary of what was	2
		done and what was found	
Introduction			
Background/rationale	2	Explain the scientific background and rationale for the investigation being reported	4-6
Objectives	3	State specific objectives, including any prespecified hypotheses	5-6
Methods			
Study design	4	Present key elements of study design early in the paper	6-8
Setting	5	Describe the setting, locations, and relevant dates, including periods of	6-8
S		recruitment, exposure, follow-up, and data collection	
Participants	6	(a) Cohort study—Give the eligibility criteria, and the sources and methods of	6-7
1		selection of participants. Describe methods of follow-up	
		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	
		Cross-sectional study—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	
		Case-control study—For matched studies, give matching criteria and the	
		number of controls per case	
Variables	7	Clearly define all outcomes, exposures, predictors, potential confounders, and	7-8
Variables	,	effect modifiers. Give diagnostic criteria, if applicable	, 0
Data sources/	8*	For each variable of interest, give sources of data and details of methods of	7-8
measurement	O	assessment (measurement). Describe comparability of assessment methods if	7-0
measarement		there is more than one group	
Bias	9	Describe any efforts to address potential sources of bias	N/A
Study size	10	Explain how the study size was arrived at	N/A
Quantitative variables	11	Explain how due study size was arrived at Explain how quantitative variables were handled in the analyses. If applicable,	7-8
Quantitative variables	11	describe which groupings were chosen and why	7-0
Statistical methods	12	(a) Describe all statistical methods, including those used to control for	8
Statistical methods	12	confounding	O
		(b) Describe any methods used to examine subgroups and interactions	N/A
		(c) Explain how missing data were addressed	N/A
		(d) Cohort study—If applicable, explain how loss to follow-up was addressed	N/A
		Case-control study—If applicable, explain how matching of cases and controls	IN/A
		was addressed	
		Cross-sectional study—If applicable, describe analytical methods taking account of sampling strategy	
			N T / A
ontinued on nevt page		(\underline{e}) 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, completing follow-up, and analysed	Tables 1 and 2
		(b) Give reasons for non-participation at each stage	
		(c) Consider use of a flow diagram	
Descriptive	14*	(a) Give characteristics of study participants (eg demographic, clinical, social) and	Tables
data		information on exposures and potential confounders	1 and
			2
		(b) Indicate number of participants with missing data for each variable of interest	
		(c) Cohort study—Summarise follow-up time (eg, average and total amount)	
Outcome data	15*	Cohort study—Report numbers of outcome events or summary measures over time	N/A
		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 (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	yes
		(c) If relevant, consider translating estimates of relative risk into absolute risk for a	
		meaningful time period	
Other analyses	17	Report other analyses done—eg analyses of subgroups and interactions, and sensitivity	N/A
		analyses	
Discussion			
Key results	18	Summarise key results with reference to study objectives	6-9
Limitations	19	Discuss limitations of the study, taking into account sources of potential bias or	9-12
		imprecision. Discuss both direction and magnitude of any potential bias	
Interpretation	20	Give a cautious overall interpretation of results considering objectives, limitations,	9-12
		multiplicity of analyses, results from similar studies, and other relevant evidence	
Generalisability	21	Discuss the generalisability (external validity) of the study results	9-12
Other information	on		
Funding	22	Give the source of funding and the role of the funders for the present study and, if	3
		applicable, for the 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.