# Obesity but not overweight is associated with increased mortality risk 

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#### Abstract

The association between body mass index (BMI) and survival has been described in various populations. However, the results remain controversial and information from low-prevalence Western countries is sparse. Our aim was to examine this association and its public health impact in Switzerland, a country with internationally low mortality rate and obesity prevalence. We included 9,853 men and women aged 25-74 years who participated in the Swiss MONICA (MONItoring of trends and determinants in CArdiovscular disease) study (1983-1992) and could be followed up for survival until 2008 by using anonymous record linkage. Cox regression models were used to calculate mortality hazard ratios (HRs) and to estimate excess deaths. Independent variables were age, sex, survey wave, diet, physical activity, smoking, educational class. After adjustment for age and sex the association between BMI and all-cause mortality was $J$ shaped (non-smokers) or $U$ shaped (smokers). Compared to BMI 18.5-24.9, among those with BMI $\geq 30$ (obesity) HR for all-cause mortality was 1.41 ( $95 \%$ confidence interval: 1.23-1.62), for cardiovascular disease (CVD) 2.05 (1.60-2.62), for cancer 1.29 (1.04-1.60). Further adjustment attenuated the obesity-mortality relationship but the associations remained statistically significant. No significant increase was found for overweight (BMI 25-29.9). Between 4 and $6.5 \%$ of all deaths, $8.8-13.7 \%$ of CVD deaths and $2.4-3.9 \%$ of cancer deaths could be attributed to


[^0]obesity. Obesity, but not overweight was associated with excess mortality, mainly because of an increased risk of death from CVD and cancer. Public health interventions should focus on preventing normal- and overweight persons from becoming obese.

Keywords Overweight • Obesity • Mortality • Relative risk • Population attributable risk

## Abbreviations

| CVD | Cardiovascular disease |
| :--- | :--- |
| COPD | Chronic obstructive pulmonary disease |
| HR | Hazard ratio |
| ICD | International classification of diseases |
| ISCED | International standard classification of education |
| SNC | Swiss national cohort |
| WHO | World health organization |

## Introduction

Worldwide, the prevalence of overweight and obesity increased over the past decades and has become a major burden for societies. Excess weight is associated with an increased risk of disease and death, particularly from cardiovascular disease (CVD) and cancer [1-3]. The association between BMI and mortality substantially varies between populations and causes of death [1,3-6] and can change over time [2,5]. A part of this variation could be due to shortcomings originating from data assessment or induced in data analyses or interpretation. For example, in many studies, data is self-reported or does not stem from general populations but from selected groups such as physicians, [7, 8] nurses [9] or alumni [10]. Also, information from low-prevalence Western countries is sparse.

The calculation and interpretation of relative risks could become critical in countries with high prevalence because the reference population (normal weight persons) becomes an increasingly selected minority [2]. This may be an explanation for the "obesity paradox", where individuals with excess weight were found to have longer survival and fewer CVD events [6, 11]. In many studies, the $U$ shape of the association between BMI and mortality risk was more pronounced among smokers than among non-smokers [5, 12, 13]. In order to estimate the independent contribution of excess weight, one should also consider that obese persons may differ in socioeconomic status and other lifestyle behaviours, e.g., regarding diet and physical activity. It is also unclear to what extent the risk associated with excess weight measured by BMI is accounted for by intermediate CVD risk factors such as high blood pressure and cholesterol level.

We aimed at determining the risk and burden of death associated with BMI on a population level. For this purpose, our study population can be regarded as exceptional. Switzerland has lower mortality (particularly for CVD) and a lower prevalence of obesity than most other countries [14]. Obesity prevalence in Switzerland is about half of that of the European average, a third of that of the UK and a quarter of that of the US [14, 15]. BMI was based on measured weight and height, and over $90 \%$ of participants could be followed-up for up to 25 years. The database includes a large set of social, behavioural and clinical parameters thus, offering the possibility to consider potential confounders or effect modifier in the analysis and to evaluate the independent effect of excess weight.

## Methods

Study population

Included individuals (25-74 years) were participants of the Swiss MONICA (MONItoring of trends and determinants in CArdiovscular disease) study. MONICA is an international multicentre project initiated and coordinated by the World Health Organization (WHO) [16]. In Switzerland, the study has been conducted in three waves between 1983 and 1992 [17, 18]. Sampled persons were invited to attend a health examination in their community of residence and to complete a self-administered questionnaire. The participation rate varied between 54 and $78 \%$ [19]. As in virtually all MONICA centres, no provision was made for a mortality follow-up. Recently, in Switzerland, this shortcoming could be overcome by an anonymous record linkage with the Swiss National Cohort (SNC) [20]. Details of the study population are given in Table 1. For the calculation of excess deaths attributable to obesity we used
prevalence from two studies conducted in 2003 (measurement) and 2007 (self-report) [21, 22].

Record linkage procedure
In order to determine survival, data from the SNC including information on cause of death was linked to MONICA participant records. The SNC encompasses all residents of Switzerland enumerated in the national 1990 or 2000 censuses ( 6.8 and 7.3 million, respectively). Deterministic and probabilistic methods were used to link anonymised census, death and emigration records [23]. Also, record linkage of MONICA and the SNC based exclusively on anonymous records [20]. $97.8 \%$ of the eligible 10,160 MONICA participants could be linked to a census (1990: 9,737; 2000: 8,749), mortality (1,526, 1984-2008) and/or emigration record (320, 1990-2008). 83 participants of the last wave of MONICA could only be linked to the preceding 1990 census but not to a subsequent census, mortality or emigration record, thus, leaving 9,853 individuals for survival analysis. Linkage procedures and linkage success were described in detail [20].

## Exposure variables

Education, lifestyle and clinical risk factors stem from MONICA. Measurements and blood sampling procedures have been described [17-19]. BMI was calculated from measured height ( cm ) and weight ( kg ) by dividing weight by height squared ( $\mathrm{kg} / \mathrm{m}^{2}$ ). Underweight (BMI $<18.5 \mathrm{~kg} /$ $\mathrm{m}^{2}$ ), normal weight (BMI 18.5-24.9 $\mathrm{kg} / \mathrm{m}^{2}$ ), overweight (BMI $25-29.9 \mathrm{~kg} / \mathrm{m}^{2}$ ) and obesity ( $\geq 30 \mathrm{~kg} / \mathrm{m}^{2}$ ) were defined according to the WHO criteria [24].

The following educational classes were used: (i) "Mandatory": compulsory schooling (corresponding to completed 8th US grade) or less (International Standard Classification of Education, ISCED 1 and 2); (ii) "Secondary": vocational training or high school (completed 12th US grade; ISCED 3); (iii) "Tertiary": technical college, upper vocational or university education (ISCED 5) [25, 26].

In order to look for the risk factor variables providing the most robust results after adjustment, we performed sensitivity analyses with smoking status (number of cigarettes smoked daily; never, former and current smokers; regular smokers, occasional smokers, non-smokers), blood cholesterol (total cholesterol, HDL-cholesterol, ratio of total cholesterol/HDL-cholesterol), blood pressure (diastolic and systolic, derived four blood pressure categories, known hypertension, hypertension treatment). We finally selected current regular and occasional smokers and nonsmokers (including former smokers). Construction of diet and physical activity scores is described in the Web Annex.

Table 1 Characteristics (counts, means and proportions) of the study population, by BMI category, 9,853 participants of the Swiss MONICA study, 1983-92, 25-74 years at baseline

|  | Total | Body mass index category ( $\mathrm{kg} / \mathrm{m}^{2}$ )* |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | <18.5 | 18.5-24.9 | 25-29.9 | $\geq 30$ |
| Participants ( $n$ ) | 9,853 | 167 | 4,590 | 3,789 | 1,307 |
| Prevalence (\%) | 100 | 1.7 | 46.7 | 38.5 | 13.1 |
| Women (\%) | 49.6 | 86.2 | 59.6 | 37.1 | 46.0 |
| Mean age (years) | 47.2 | 44.7 (42.6-46.8) | 44.4 (44.1-44.7) | 48.9 (48.6-49.3) | 52.0 (51.4-52.6) |
| Mean follow-up time (years) | 18.6 | 18.5 (17.7-19.3) | 19.0 (18.8-19.1) | 18.4 (18.3-18.6) | 17.7 (17.5-18.0) |
| Education |  |  |  |  |  |
| Tertiary (\%) | 19.3 | 28.9 (21.9-35.9) | 23.9 (22.7-25.1) | 16.2 (15.0-17.1) | 10.9 (9.2-12.6) |
| Upper secondary (\%) | 48.7 | 51.2 (43.5-58.9) | 51.5 (50.0-52.9) | 47.7 (46.2-49.4) | 41.5 (38.8-44.2) |
| Mandatory and secondary (\%) | 32.0 | 19.9 (13.7-26.0) | 24.6 (23.4-25.9) | 36.1 (34.5-37.6) | 47.6 (44.9-50.3) |
| Smoking |  |  |  |  |  |
| Current regular (\%) | 28.6 | 31.7 (24.6-38.9) | 32.3 (31.0-33.7) | 26.6 (25.2-28.0) | 21.2 (18.9-23.4) |
| Current occasional (\%) | 4.5 | 3.6 (0.7-6.4) | 4.8 (4.2-5.4) | 4.7 (4.0-5.4) | 2.8 (1.9-3.7) |
| Non-smokers (former and never, \%) | 66.9 | 64.7 (57.3-72.0) | 62.9 (61.5-64.3) | 68.7 (67.2-70.1) | 76.0 (73.7-78.4) |
| Mean physical activity score | 3.52 | 3.47 (3.17-3.77) | 3.79 (3.74-3.84) | 3.44 (3.39-3.49) | 2.85 (2.76-2.94) |
| Mean diet score | 3.51 | 3.51 (3.35-3.67) | 3.54 (3.51-3.57) | 3.51 (3.48-3.54) | 3.39 (3.34-3.44) |
| Blood pressure (systolic/diastolic) |  |  |  |  |  |
| Mean systolic (mmHg) | 129.3 | 117.4 (115.0-119.9) | 124.5 (124.0-125.0) | 132.2 (131.6-132.8) | 138.9 (137.8-139.9) |
| Mean diastolic ( mmHg ) | 78.9 | 71.1 (69.7-72.6) | 76.1 (75.8-76.4) | 80.8 (80.4-81.1) | 84.2 (83.7-84.8) |
| $\geq 140$ or $\geq 90$ (\%) | 28.8 | 11.4 (6.5-16.2) | 19.0 (17.9-20.2) | 34.6 (33.1-36.1) | 48.6 (45.9-51.4) |
| Cholesterol (C) |  |  |  |  |  |
| Total C: HDL C (ratio) | 5.06 | 3.79 (3.62-3.97) | 4.42 (4.38-4.47) | 5.52 (5.46-5.58) | 6.15 (6.03-6.27) |
| Ratio $\geq 5$ (\%) | 44.4 | 13.0 (7.8-18.3) | 28.7 (27.4-30.0) | 56.8 (55.2-58.4) | 67.5 (64.9-70.1) |
| Deaths |  |  |  |  |  |
| All causes ( $n$ ) | 1,526 | 24 | 520 | 634 | 348 |
| Cardiovascular disease ( $n$ ) | 448 | 8 | 126 | 187 | 127 |
| Cancer ( $n$ ) | 636 | 2 | 232 | 268 | 134 |

* Based on height and weight measured at baseline

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For separate analyses we used cholesterol ratio and the four blood pressure categories according to the American Heart Association [27].

## Outcome variables

Causes of death were classified according to ICD (International Classification of Diseases) revisions 8 (ICD-8) and 10 (ICD-10). In Switzerland ICD-8 was used until 1994 followed by ICD-10 thereafter. Due to the relatively small number of deaths, causes of death had to be grouped into CVD (ICD-8: 410-438; ICD-10: I00-I99), cancer (ICD-8:

140-239; ICD-10: C00-C99; D00-D48), and non-cancer-non-CVD (remainder).

Statistical analyses
In a first step, we performed all analyses separately by sex. We found no significant sex differences, neither in obesity prevalence, nor in obesity related hazard ratio (HRs). In order to obtain more robust estimates, we decided to pool sexes. Kaplan-Meier curves were calculated for all-cause mortality as well as separately for CVD, cancer and non-cancer-non-CVD, using appropriate methods to account for competing risks. For the estimation of HRs, we fitted a Cox
regression model including relevant independent variables (age, sex, educational class, risk factors) and adjusting for study wave. The association between BMI and death was increasingly adjusted for additional variables using three models: (1) age, sex and survey wave; (2) + diet/physical activity scores and smoking; (3) + educational class. The proportional hazards assumption was tested and checked by visual inspection and seemed to be widely fulfilled. The methods for the calculation of population attributable fractions is described in the Web Annex. General descriptive analyses and survival estimations were performed with Stata 11 (Stata Corp, Texas, USA, 2009), Kaplan-Meier curves and attributable deaths were obtained with R 2.10.1 (The R Foundation for Statistical Computing, 2009).

## Results

## Descriptive crude analyses

Characteristics of participants are summarized in Table 1. The number of underweight persons was small and mainly consisted of women. In contrast, among obese, there were about as many women as men. The prevalence of obesity was higher than that based on a survey with self-reports but in line with that based on a study with measurement of height and weight in 2003 [21, 22]. Obese persons were older, particularly when compared to normal weight participants. Between BMI categories there were also differences regarding educational class and smoking status. Variations in blood pressure and cholesterol were substantially larger than in physical activity and diet.

Kaplan-Meier curves decrease when a death occurs (Fig. 1). The curves of overweight and obese persons decrease more rapidly than the curve of normal weight persons. However, one should consider that persons in the different BMI categories also differed by sex and age: normal weight persons were younger and more often women. The curve of obese persons differs more strongly from the curve of overweight persons in CVD than in cancer deaths.

## Adjusted analyses

Figure 2 shows HRs for all-cause mortality by BMI category in smokers (regular and occasional) and non-smokers (never and former). The curves have a $J$ shape (nonsmokers) or a $U$ shape (smokers), with increased HRs at one or both extremes. Under- and normal- weight smokers had in average a higher risk than obese non-smokers. In both groups, overweight was not associated with higher risk than normal weight. The increase in HRs at

BMI $<20 \mathrm{~kg} / \mathrm{m}^{2}$ was only due to an increase in men-in women, HRs remained fairly unchanged. However, the number of deaths was small and the confidence intervals were overlapping between sexes. In contrast, increase in HRs for BMI $\geq 30 \mathrm{~kg} / \mathrm{m}^{2}$ was almost identical in men and women (sex differences not shown). The curve of the entire population (smokers and non-smokers combined) is shown in Web Annex Fig. 1.

Table 2 shows all-cause and cause-specific mortality risk by BMI category. The results are adjusted with an increasing number of variables leading to three different models. For none of the cause of death groups and in none of the models, overweight was associated with increased mortality. Underweight tended to be associated with higher CVD but with lower cancer mortality. The number of cases was however, small in that group. After full adjustment, mortality from all-causes among obese persons was $36 \%$ higher than among normal weight persons. CVD mortality was more strongly associated with obesity than cancer mortality. Non-cancer-non-CVD deaths were comparably more frequent among underweight than among normal weight individuals. In underweight persons, HRs for CVD and non-cancer-non-CVD deaths were higher in men than in women but the differences did not reach statistical significance (not shown). Generally, there were no fundamental differences between age groups (Web annex Table 1). In separate analyses, adjustment for cholesterol ratio and blood pressure substantially attenuated the association of obesity with CVD mortality by about $50 \%$. Nevertheless, this association remained statistically significant. The relationship of CVD mortality with overweight was virtually not affected by the adjustment.

Figure 3 shows deaths attributable to obesity as percentage of all deaths related to all death in the corresponding group (all-cause, CVD, cancer, non-cancer-non-CVD). Prevalence rates from studies with either self-reported (empty diamonds) or measured height and weight (filled circles) used for the calculation of attributable deaths are given in the legend. Due to the higher obesity prevalence, the proportion of attributable deaths was higher when it was based on measured height and weight: $4 \%$ (self-report) and $6.5 \%$ (measurement) of all deaths were attributable to obesity. The proportions were higher for CVD (8.8 and $13.7 \%$ of all CVD deaths) than for cancer ( 2.4 and $3.9 \%$ of all cancer deaths).

## Discussion

Main results

In this general Swiss population, obesity but not overweight was associated with increased risk of dying from


Fig. 1 Kaplan-Meier curves of normal weight, overweight and obese individuals for mortality from all causes (A), CVD (B), cancer (C) and non-cancer-non-CVD (D), 9,853 participants of the Swiss MONICA study, 1983-92, 25-74 years at baseline. MONICA


MONItoring of trends and determinants in CArdiovscular disease. CVD Cardiovascular Disease. Body Mass index categories are based on height and weight measured at baseline

Fig. 2 All-cause mortality (hazard ratios with $95 \%$ confidence interval) by smoking status and BMI category, adjusted for age, sex and study wave, 9,853 participants of the Swiss MONICA study, 1983-92, 25-74 years at baseline. MONICA MONItoring of trends and determinants in CArdiovscular disease. Reference category are nonsmokers with BMI $20.0-22.4 \mathrm{~kg} / \mathrm{m}^{2}$. Asterisk represents based on height and weight measured at baseline


CVD and, to a smaller extent, from cancer. The relationship between BMI and all-cause mortality followed a $J$ shaped pattern in non-smokers and a $U$ shape pattern in smokers (Fig. 2). Among underweight persons, the tendency to an increased risk was driven by CVD (men) and non-cancer-non-CVD (men and women). However, this relationship was confounded by smoking status. The relative risk of death associated with obesity remained
significantly increased after adjustment for lifestyle factors and educational level (Table 2).

Overweight and obesity

A J- or U-shaped relationship between BMI and all-cause mortality has been documented in most but not all cohort studies [1, 3, 13, 15, 28-32]. A more pronounced $U$ shape

Table 2 Adjusted hazard ratios for all-cause and cause specific mortality, by BMI category, 9,853 participants of the MONICA study, 1983-92, 25-74 years at baseline

|  | Body mass index | y (kg |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $<18.5$ |  | 18.5-24.9 | 25-29.9 |  | $\geq 30$ |  |
|  | HR (95\% CI) | $P$ | HR | HR (95\% CI) | P | HR (95\% CI) | P |
| All cause |  |  |  |  |  |  |  |
| Model 1 | 1.32 (0.87-1.99) | 0.187 | 1.00 | 0.96 (0.85-1.08) | 0.455 | 1.41 (1.23-1.62) | $<0.001$ |
| Model 2 | 1.18 (0.78-1.80) | 0.431 | 1.00 | 0.95 (0.85-1.07) | 0.438 | 1.37 (1.19-1.58) | $<0.001$ |
| Model 3 | 1.21 (0.80-1.85) | 0.367 | 1.00 | 0.94 (0.84-1.06) | 0.329 | 1.36 (1.18-1.56) | $<0.001$ |
| Cardiovas | isease (CVD) |  |  |  |  |  |  |
| Model 1 | 1.62 (0.79-3.33) | 0.189 | 1.00 | 1.10 (088-1.38) | 0.407 | 2.05 (1.60-2.63 | $<0.001$ |
| Model 2 | 1.52 (0.74-3.13) | 0.259 | 1.00 | 1.08 (0.86-1.35) | 0.535 | 1.95 (1.52-2.51) | $<0.001$ |
| Model 3 | 1.60 (0.77-3.29) | 0.206 | 1.00 | 1.07 (0.85-1.35) | 0.551 | 1.92 (1.49-2.47) | $<0.001$ |
| Cancer |  |  |  |  |  |  |  |
| Model 1 | 0.26 (0.06-1.04) | 0.056 | 1.00 | 0.96 (0.80-1.15) | 0.647 | 1.29 (1.04-1.60) | 0.021 |
| Model 2 | 0.24 (0.06-0.98) | 0.047 | 1.00 | 0.97 (0.81-1.16) | 0.711 | 1.28 (1.03-1.60) | 0.027 |
| Model 3 | 0.25 (0.06-1.01) | 0.051 | 1.00 | 0.94 (0.79-1.13) | 0.519 | 1.25 (1.00-1.56) | 0.050 |
| Non-cance | CVD |  |  |  |  |  |  |
| Model 1 | 2.54 (1.46-4.42) | 0.001 | 1.00 | 0.86 (0.69-1.06) | 0.158 | 1.10 (0.84-1.44) | 0.488 |
| Model 2 | 2.18 (1.23-3.88) | 0.008 | 1.00 | 0.86 (0.69-1.07) | 0.174 | 1.08 (0.82-1.42) | 0.576 |
| Model 3 | 2.22 (1.25-4.0) | 0.006 | 1.00 | 0.86 (0.69-1.06) | 0.162 | 1.08 (0.82-1.42) | 0.585 |

* Based on height and weight measured at baseline

Model 1 (basic) adjusted for age, sex and survey wave
Model 2 (lifestyle) additionally adjusted for diet, physical activity and smoking
Model 3 (socio-economic status) additionally adjusted for education
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Fig. 3 Estimated excess deaths in Switzerland attributable to obesity, in percent of all causes and cause-specific deaths, 9,853 participants of the Swiss MONICA study, 1983-92, 25-74 years at baseline. MONICA MONItoring of trends and determinants in CArdiovscular disease. Analyses were adjusted for age and sex

comparable to ours was found in a MONICA-population from Poland and in several other studies $[1,5,12,13,29$, 30]. The fact that we found no increased risk in overweight persons was at variance with reports from Korea, Europe and North America and China (in persons ages $<65$ years) [1, 13, 29, 31], but in line with other European studies, reports from India, the US and with results from older populations $[6,12,15,29,32-34]$. In contrast to
overweight, obesity was associated with increased risk of death in almost all available studies.

As shown by most others and by our study, the increased mortality risk was mainly due to CVD $[1,3,13,15,29,31$, 35]: the HRs of obese versus normal weight persons were very similar to ours $[1,13,15,29,31,35]$. In accordance with our results, no or only a marginal increase in relative CVD mortality risk in overweight persons was found in

Poland and the US $[29,35]$ while a small but significant increase was reported in Europe and North America, China and Korea [1, 13, 29, 31]. Our relative risks for cancer mortality associated with obesity were somewhat higher than those found in a large Asian-Pacific sample [36]. This and two other samples showed a small but significant relative cancer risk in overweight persons, which was at odds with our finding [3, 31, 36]. In a study including participants from Europe and North America, the increased risk associated with higher BMI was substantially lower for cancer than for CVD (HR per $5 \mathrm{~kg} / \mathrm{m}^{2}$ of increase in BMI: 1.10 vs. 1.41 ) [1, 3]. In contrast, in a large Korean population cancer and CVD mortality were similarly associated with obesity [31].

In our study, lifestyle factors and educational level only moderately attenuated excess risk of obese individuals. Thus, there is little evidence for uncontrolled confounding. However, in line with most literature a part of the CVD risk associated with obesity was apparently mediated by high blood pressure and cholesterol ratio [29, 31, 35]. In a metaanalysis of 21 cohort studies, almost half of the increased risk of fatal and nonfatal CHD events in overweight and obese persons was explained by higher blood pressure and cholesterol levels [37].

Others showed that fatal and non-fatal CVD events were similarly associated with overweight and obesity [35]. Analogously, cancer incidence and morbidity were associated with excess weight in a similar manner [4]. It is therefore possible that, in Switzerland, corresponding to mortality figures, there is also no increased relative risk for CVD and cancer morbidity among overweight persons.

## Underweight

Persons with underweight only contributed a small number of deaths and results should therefore be interpreted cautiously. In accordance with others, the relative risk of death of underweight persons was higher in smokers than in nonsmokers (Fig. 2) [1, 3, 13, 31]. The tendency to higher mortality risk of underweight persons was driven by death from non-cancer to non-CVD. Our data did not offer sufficient statistical power to assess which specific causes were responsible for excess mortality in smokers and nonsmokers. Other studies showed that respiratory mortality (e.g., COPD) substantially contributed [1, 13, 31, 38]. Most but not all studies showed an increased CVD mortality at BMI below 18.5 [1, 3, 13, 29-31].

The association between low BMI and death could reflect insufficient consideration of early or chronic disease leading to both thinness and death. Our data, however, provide only little evidence for this: first, our population was comparably young and age did not appear to fundamentally affect the relationship between BMI and mortality
risk; second, adjustment for smoking only marginally attenuated estimates; third, the number of cancer deaths was negligible among underweight persons and the pattern (non-cancer-non-CVD $>\mathrm{CVD}>$ cancer) was the same as in a large study considering only healthy subjects who never smoked [3]. Furthermore, only 3.6 and $11.9 \%$ of all non-injury deaths occurred during the first 2 and 5 years after study entry. Others have shown that an increased risk of death in underweight persons was still observed among healthy participants and after exclusion of the initial 5 years of follow-up [3, 29, 39].

Excess deaths attributable to obesity
In Switzerland, nationally representative data on obesity prevalence is only available from self-reports which substantially underestimate real obesity prevalence [40]. For valid figures, excess deaths should be estimated with obesity prevalence based on measured height and weight. Since more recent data was not available, we used measured BMI data from 2003 for the calculation of excess deaths. However, this should not substantially underestimate current excess deaths because in Switzerland obesity prevalence stagnated or increased only marginally since 2003 [41].

In the US, where obesity prevalence is about twice as high as in Switzerland, approximately $29 \%$ of CVD deaths and $8 \%$ of cancer deaths were attributable to BMI $\geq 25 \mathrm{~kg} / \mathrm{m}^{2}[1,11]$. For the UK, the corresponding proportions were 23 and $6 \%$, respectively [1, 42]. In accordance with our figures, a calculation based on European data estimated that in Switzerland around 700 cancer cases/year could be attributed to overweight or obesity [43]. Probably, our estimates would be higher, had we used other markers than BMI for the definition of obesity (e.g., body fat percentage) [44]. However, a large European study proved BMI to remain significantly associated with the risk of death also in models that included waist circumference or waist-to-hip ratio [32].

Public health implications

Our findings do not support the concept that persons with overweight should decrease their BMI in order to reduce their risk of premature death. For this category (BMI 25-29.9) efforts aimed at avoiding weight gain and improving health behaviour may be more appropriate. Our study also implies that by controlling and treating risk factors in obese persons, excess mortality risk can be decreased but not eliminated. Because long term weight loss is unrealistic in obese persons, the only way to decrease excess deaths on a population level is to prevent persons from becoming obese. Our results also show that
the benefit of lower BMI does not counterweight the increased risk associated with smoking. Thus, under- and normal- weight smokers should be as consequently screened and motivated for smoking cessation as obese smokers.

## Limitations

The MONICA participants included in our study had a lower mortality and were thus, presumably more healthy than the general Swiss population [20]. However, in a large study conducted in an immediately neighbouring Austrian region, the difference between expected and observed mortality was even larger [13], but the observed patterns were generally the same as in our study. Obese persons are less likely to participate in health surveys than normal weight persons and the "healthy participant effect" could have distorted relative risks [45]. We also had only one measurement of height and weight at study entry and could not consider change in BMI during follow-up. Our obesity marker was restricted to BMI and did not include waist circumference, visceral fat or body fat percentage. To the extent that BMI imperfectly reflects adiposity, our results would tend to underestimate the deleterious effects of obesity [44]. Our information on diet and physical activity was based on a coarse assessment. However, the derived scores were significantly associated with mortality (not shown). Our information on pre-existing disease was restricted to known hypertension. Inclusion of this variable in the model only minimally affected estimates. Also for other reasons discussed above, we have little evidence that severe disease existing before study entry played a major role in our population. We also found no evidence that age affected the obesity-mortality relationship.

## Conclusion

In this study from Switzerland, obesity but not overweight was associated with an increased risk of death. Excess mortality was driven by CVD and, to a smaller extent, by cancer. After adjustment for lifestyle risk factors and for educational class, the independent effect of obesity decreased but remained significant. Our results question the targeting of overweight individuals for weight loss programs in order to decrease burden of disease. In contrast, people should be prevented from becoming obese-irrespective of their body weight. In smokers, underweight was significantly associated with premature death, mainly due to non-cancer-non-CVD causes.

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## References

1. Whitlock G, Lewington S, Sherliker P, et al. Body-mass index and cause-specific mortality in 900,000 adults: collaborative analyses of 57 prospective studies. Lancet. 2009;373:1083-96.
2. Flegal KM, Graubard BI, Williamson DF, et al. Excess deaths associated with underweight, overweight, and obesity. Jama. 2005;293:1861-7.
3. Berrington de Gonzalez A, Hartge P, Cerhan JR, et al. Body-mass index and mortality among 1.46 million white adults. N Engl J Med. 2010;363:2211-9.
4. Reeves GK, Pirie K, Beral V, et al. Cancer incidence and mortality in relation to body mass index in the million women study: cohort study. BMJ. 2007;335:1134.
5. Manson JE, Bassuk SS, Hu FB, et al. Estimating the number of deaths due to obesity: can the divergent findings be reconciled? J Womens Health (Larchmt). 2007;16:168-76.
6. Romero-Corral A, Montori VM, Somers VK, et al. Association of bodyweight with total mortality and with cardiovascular events in coronary artery disease: a systematic review of cohort studies. Lancet. 2006;368:666-78.
7. Ajani UA, Lotufo PA, Gaziano JM, et al. Body mass index and mortality among US male physicians. Ann Epidemiol. 2004;14: 731-9.
8. Gelber RP, Kurth T, Manson JE, et al. Body mass index and mortality in men: evaluating the shape of the association. Int J Obes (Lond). 2007;31:1240-7.
9. Manson JE, Willett WC, Stampfer MJ, et al. Body weight and mortality among women. N Engl J Med. 1995;333:677-85.
10. Lee IM, Manson JE, Hennekens CH, et al. Body weight and mortality. A 27 year follow-up of middle-aged men. JAMA. 1993;270:2823-8.
11. Ogden CL, Carroll MD, Curtin LR, et al. Prevalence of overweight and obesity in the United States, 1999-2004. JAMA. 2006;295:1549-55.
12. Engeland A, Bjorge T, Selmer RM, et al. Height and body mass index in relation to total mortality. Epidemiology. 2003;14: 293-9.
13. Klenk J, Nagel G, Ulmer H, et al. Body mass index and mortality: results of a cohort of 184, 697 adults in Austria. Eur J Epidemiol. 2009;24:83-91.
14. Bjorntorp P, De Jounge K, Sjostrom L, et al. The effect of physical training on insulin production in obesity. Metabolism. 1970;19:631-8.
15. Flegal KM, Graubard BI, Williamson DF, et al. Cause-specific excess deaths associated with underweight, overweight, and obesity. JAMA. 2007;298:2028-37.
16. Bothig S. WHO MONICA project: objectives and design. Int J Epidemiol. 1989;18:S29-37.
17. Wietlisbach V. Théorie et pratique de l'échantillonnage: L'exemple de l'enquête MONICA. Soz Praeventivmed. 1987;32:52-62.
18. Wietlisbach V, Paccaud F, Rickenbach M, et al. Trends in cardiovascular risk factors (1984-1993) in a Swiss region: results of 3 population surveys. Prev Med. 1997;26:523-33.
19. Wolf HK, Kuulasmaa K, Tolonen H, et al. Participation rates, quality of sampling frames and sampling fractions in the MONICA surveys. 1998; http://www.ktl.fi/publications/monica/nonres/ nonres.htm [last access 12 Jan 2010].
20. Bopp M, Braun J, Faeh D, et al. Establishing a follow-up of the Swiss MONICA participants (1984-1993): record linkage with census and mortality data. BMC Public Health. 2010;10:562.
21. Swiss Federal Statistical Office. Die Schweizerische Gesundheitsbefragung 2007 in Kürze [WWW document] URL http:// www.bfs.admin.ch/bfs/portal/de/index/infothek/erhebungen__ quellen/blank/blank/ess/03.Document.122331.pdf (accessed April 2010). 2008
22. Firmann M, Mayor V, Vidal PM, et al. The CoLaus study: a population-based study to investigate the epidemiology and genetic determinants of cardiovascular risk factors and metabolic syndrome. BMC Cardiovasc Disord. 2008;8:6.
23. Bopp M, Spoerri A, Zwahlen M, et al. Cohort profile: the Swiss national cohort: a longitudinal study of 6.8 million people. Int J Epidemiol. 2009;38:379-84.
24. Obesity: preventing and managing the global epidemic. Report of a WHO consultation. World Health Organ Tech Rep Ser. 2000; 894. http://whqlibdoc.who.int/trs/WHO_TRS_894.pdf.
25. Bopp M, Minder CE. Mortality by education in German speaking Switzerland, 1990-1997: results from the Swiss national cohort. Int J Epidemiol. 2003;32:346-54.
26. Lynch J, Smith GD, Harper S, et al. Is income inequality a determinant of population health? Part 1. A systematic review. Milbank Q. 2004;82:5-99.
27. Chobanian AV, Bakris GL, Black HR, et al. Seventh report of the joint national committee on prevention, detection, evaluation, and treatment of high blood pressure. Hypertension. 2003;42:1206-52.
28. Baldinger B, Schwarz C, Jaggy C. Cardiovascular risk factors, BMI and mortality in a cohort of Swiss males (1976-2001) with high-sum-assured life insurance cover. J Insur Med. 2006;38: 44-53.
29. Gu D, He J, Duan X, et al. Body weight and mortality among men and women in China. JAMA. 2006;295:776-83.
30. Pajak A, Topor-Madry R, Waskiewicz A, et al. Body mass index and risk of death in middle-aged men and women in Poland. Results of POL-MONICA cohort study. Kardiol Pol. 2005;62: 95-105. discussion 06-7.
31. Jee SH, Sull JW, Park J, et al. Body-mass index and mortality in Korean men and women. N Engl J Med. 2006;355:779-87.
32. Pischon T, Boeing H, Hoffmann K, et al. General and abdominal adiposity and risk of death in Europe. N Engl J Med. 2008;359:2105-20.
33. Tamakoshi A, Yatsuya H, Lin Y, et al. BMI and all-cause mortality among Japanese older adults: findings from the Japan collaborative cohort study. Obesity (Silver Spring). 2010;18:362-9.
34. Flicker L, McCaul KA, Hankey GJ, et al. Body mass index and survival in men and women aged 70-75. J Am Geriatr Soc. 2010;58:234-41.
35. Wilson PW, D'Agostino RB, Sullivan L, et al. Overweight and obesity as determinants of cardiovascular risk: the Framingham experience. Arch Intern Med. 2002;162:1867-72.
36. Parr CL, Batty GD, Lam TH, et al. Body-mass index and cancer mortality in the Asia-Pacific Cohort Studies Collaboration: pooled analyses of 424,519 participants. Lancet Oncol. 2010;11:741-52.
37. Bogers RP, Bemelmans WJ, Hoogenveen RT, et al. Association of overweight with increased risk of coronary heart disease partly independent of blood pressure and cholesterol levels: a metaanalysis of 21 cohort studies including more than 300,000 persons. Arch Intern Med. 2007;167:1720-8.
38. Sauvaget C, Ramadas K, Thomas G, et al. Body mass index, weight change and mortality risk in a prospective study in India. Int J Epidemiol. 2008;37:990-1004.
39. He J, Gu D, Wu X, et al. Major causes of death among men and women in China. N Engl J Med. 2005;353:1124-34.
40. Faeh D, Braun J, Bopp M. Underestimation of obesity prevalence in Switzerland: comparison of two methods for correction of selfreport. Swiss Med Wkly. 2009;139:752-6.
41. Faeh D, Bopp M. Excess weight in the canton of Zurich, 1992-2009: harbinger of a trend reversal in Switzerland? Swiss Med Wkly. 2010;140:w13090.
42. Health Survey for England. 2004. http://www.dh.gov.uk/en/ Publicationsandstatistics/PublishedSurvey/HealthSurveyFor England/index.htm (accessed 10 Sep 2010). 2004.
43. Ceschi M, Gutzwiller F, Moch H, et al. Epidemiology and pathophysiology of obesity as cause of cancer. Swiss Med Wkly. 2007;137:50-6.
44. Marques-Vidal P, Bochud M, Mooser V, et al. Obesity markers and estimated 10-year fatal cardiovascular risk in Switzerland. Nutr Metab Cardiovasc Dis. 2009;19:462-8.
45. Sonne-Holm S, Sorensen TI. Prospective study of attainment of social class of severely obese subjects in relation to parental social class, intelligence, and education. Br Med J (Clin Res Ed). 1986;292:586-9.

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