BMI, Weight Stability and Mortality among Adults without Clinical Co-Morbidities: A 22-Year Mortality Follow-Up in the Finnish Twin Cohort

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Key Words
Body weight · Cause of death · Mortality · Weight changes · Discordant twins

Summary
Aim and Method: Cause-specific mortality was studied in relation to body mass index (BMI) and weight stability (defined as less than 1 BMI unit change during a 6-year period) in 15,424 initially healthy twin subjects from the Finnish Twin Cohort, first examined in 1975, re-examined in 1981, and then followed over 22 years (1982–2003). Additionally, death discordant twin pairs were studied to assess whether body weight differences are associated with mortality independent of childhood factors and genetic background. Deaths and cause of death were ascertained from national registries. Associations with mortality were estimated by Cox proportional hazards model for all individuals and conditional logistic regression analysis for pairwise analyses. Results: Mortality increased with increasing BMI for all causes and coronary heart disease (CHD) in men, and there were no associations for all natural causes, cerebrovascular disease, and violent deaths. After adjustment for multiple co-variates and changes in co-variates between 1975 and 1981, BMI was associated with CHD mortality in all men (hazard ratio (HR) = 1.22, 95% CI 1.06–1.41) and in men with stable weight between 1975 and 1981 (HR = 1.26, 95% CI 1.03–1.55). Overall risk of death and cause-specific mortality was not associated with BMI in women. Conclusion: Among clinically healthy subjects at low risk of death, BMI appears to be associated with CHD risk in men.

Introduction
Several epidemiological studies have demonstrated either linear or J- or U-shaped relationship between mortality and body weight [1–7]. Recent large population-based studies have suggested that mortality is lower among normal-weight subjects than among underweight, overweight or obese subjects [8, 9]. However, it has been debated where on the scale between overweight and obesity the risk for mortality actually increases as some recent studies have shown that mortality is lower among overweight than normal-weight subjects [10].

Most studies on weight and mortality are based on body mass index (BMI), which is dependent on the size of the lean and the fat body mass [11]. The risks for co-morbidities like cardiovascular diseases and type 2 diabetes tend to increase linearly with increasing BMI [12–16].

Studies of weight change and mortality are beset with methodological problems like confounding by smoking, intentional weight loss and pre-existing morbidities, which make interpretations and comparisons of results from various studies difficult. On one hand, modest weight loss leads to metabolic benefits [17–23], but on the other hand weight loss has been associated with excess mortality in cohort studies [14, 15, 24]. Cohort studies [25] as well as prospective longitudinal studies have concluded that stable weight may promote health over time [16, 26, 27]. In our recent study from the Finnish Twin Cohort on healthy overweight and obese subjects, intentional weight loss over 6 years predicted an almost two-fold increase in mortality compared to stable weight in initially healthy overweight and obese subjects [28], with weight gain also predicting increased risk of death. Since any single measurement
or report of concurrent body weight will provide a mixture of subjects with stable body weight and changed body weight, it is possible that the BMI-mortality association when based on a single point correspondingly reflects mixed effects of stable and changed BMI on mortality.

It is also possible that the hitherto observed associations between weight change and mortality are confounded by genetic or early environmental influences. Applying the co-twin control method, based on comparison of twin pairs discordant for the phenotype under study, it may be possible to disentangle effects of the individual environment, causing weight differences, and effects attributable to genes and early shared environment [29]. A limited number of studies with co-twin control design have been published, a few of them focused elsewhere [28]. Further exclusions due to missing data in co-variates are described below.

Our previous study focused on the intentional weight loss and mortality in overweight and obese subjects over 6 years. The aim of the present study was to further examine how the whole range of BMI assessed at two time points 6 years apart, and stable body weight over time, are related to all-cause and cause-specific mortality during a 22-year follow-up among subjects who were not working in 1981 (i.e. being on early or disability pension, unemployed or otherwise not employed) as described in detail elsewhere [28]. Further exclusions due to missing data in co-variates are described below.

BMI was calculated from the reported height in the 1975 survey and weight in both surveys. Self-reported current height and weight generally have high accuracy [38, 39] and the accuracy of the reports used in this study sample was supported by an analysis of another sub-study of the Finnish Twin Cohort [40].

The subjects were categorized as underweight (BMI under 18.5 kg/m²), normal weight (18.5 to <25 kg/m²), overweight (25 to <30 kg/m²) or obese (≥30 kg/m²) according to a BMI-based classification of the International Obesity Task Force (IOTF) [13]. Three classes of normal-weight subjects were used (18.5 to <22 kg/m², 22 to <23.5 kg/m², 23 to <25 kg/m²) Nor-
mal-weight subjects with BMI 18.5 to <22 kg/m² were used as a reference group in mortality analyses. Overweight subjects were split as over and under BMI of 27 kg/m² while the number of obese subjects did not permit subgroup analyses. Given gender differences in both body composition and mortality, all analyses were done separately for men and women.

Mortality analyses were also conducted using BMI z-scores, i.e. units of 1 standard deviation (SD) for BMI in 1975 or 1981, computed separately by sex in order to investigate if there is a change in the association between these two time points. Weight-stable subjects were defined as those whose BMI changed by less than 1 BMI unit between 1975 and 1981, while weight-unstable subjects were those who had changed by either loss or gain by 1 BMI unit or more.

The lifestyle factors (smoking, binge drinking, leisure physical activity, life dissatisfaction) were asked at both surveys and recorded in simpler categories in order to avoid overloading the multivariate models. Each of these risk factors were then categorized as being present both in 1975 and 1981, only in 1975, only in 1981, or absent in both surveys, which was coded by three dummy variables [32]. Schooling (eight categories of years of schooling), social class (six categories) and income (coded as eight levels) in 1975 were also included as potential confounders. A history of hypertension was based on both questionnaire information (self-report of physician-diagnosed hypertension, regular use of antihypertensive medications), and register information on the right to full reimbursement of physician-diagnosed hypertension, regular use of antihypertensive medications was included as a dichotomous measure. Physical activity at leisure and hypertension were included in the model but were regarded as mediators influenced by obesity, rather than confounders. Some of the co-variate information was missing, which led to the final number of subjects.

The participants were followed up until death, migration or the end of 2003, and a total of 1,163 subjects (787 men and 376 women) died. Underlying causes of death were available from Statistics Finland [41] for all but 15 subjects. The ascertainment of causes of death was based on forensic autopsy in 40% of cases. Natural and violent causes were analyzed separately. Among natural causes, coronary heart disease (CHD; ICD 8 and 9 410-414, ICD-10 I 20-I29) and cerebrovascular disease (CVD; ICD 8 and 9 430-438, ICD-10 I60-I69) were analyzed separately and as a joint outcome. The group of violent deaths included any accidental deaths and suicides. The standardized mortality ratio of the subjects without clinically overt disease was 40% of the mortality of all the twins, who had replied to both questionnaires [28].

<table>
<thead>
<tr>
<th>BMI class</th>
<th>1975</th>
<th>1981</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of subjects</td>
<td>Number of deaths</td>
<td>Number of subjects</td>
</tr>
<tr>
<td>Underweight &lt;18.5 kg/m²</td>
<td>700</td>
<td>20</td>
</tr>
<tr>
<td>Normal weight 18.5 to &lt;22 kg/m²</td>
<td>3,926</td>
<td>140</td>
</tr>
<tr>
<td>22 to &lt;23.5 kg/m²</td>
<td>1,209</td>
<td>58</td>
</tr>
<tr>
<td>23.5 to &lt;25 kg/m²</td>
<td>859</td>
<td>63</td>
</tr>
<tr>
<td>Overweight 25 to &lt;27 kg/m²</td>
<td>569</td>
<td>47</td>
</tr>
<tr>
<td>27 to &lt;30 kg/m²</td>
<td>368</td>
<td>34</td>
</tr>
<tr>
<td>Obese ≥30 kg/m²</td>
<td>136</td>
<td>14</td>
</tr>
<tr>
<td>Total</td>
<td>7,767</td>
<td>376</td>
</tr>
</tbody>
</table>

| Table 1. Number of subjects at baseline and number of deaths to the end of 2003 among clinically healthy subjects from the Finnish Twin Cohort in classes of BMI in 1975.

Statistical Analyses

Total mortality from 1982 through 2003 was analyzed by the Cox proportional hazards regression model with time (the number of days scaled as years) since response to the 1982 survey using Stata software [28, 40]. The estimated regression coefficients and standard errors were converted to hazard ratios (HR) with 95% confidence intervals (95% CI). The analyses were done using an adjusted model including the statistically significant co-variates. Age in 1981 was used as a built-in stratification variable (age intervals 24–29 years, 30–34 years, 35–44 years, > 45 years) allowing age to be modelled as a continuous variable within the age intervals mentioned. The proportionality of hazards were checked by examining ‘log-log’ plot, i.e. –ln{–ln(survival)} curves for categories of the variables versus ln(analysis time) and using Schoenfeld residuals. As the twin cohort was ascertained as twin pairs, observations (i.e. twin individuals) in twin pairs may be correlated. Therefore we adjusted for this in the estimation of standard errors of the HR and the p values by using a robust variance estimate that adjusts for within-pair correlation [45]. Two-tailed p values below 0.05 were considered statistically significant.

Results

Out of 15,494 study subjects, 1,163 (of which 787 were men) died during the 22-year follow-up (table 1). Overweight subjects made up 18% of the sample in 1975 while only 2% were obese. In 1981, 24% were overweight and 3% obese, men being more often overweight and obese than women at both time periods.
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The mean BMI (± SD) in 1981 was 24.3 ± 2.85 kg/m² in men and 22.6 ± 3.3 kg/m² in women. The mean change in BMI between 1975 and 1981 was 0.76 ± 1.61 kg/m². The mean age of women and men was 36.0 and 36.1 years in 1981, respectively.

**Direct BMI Measurement and Subsequent Mortality**

The total mortality increased approximately linearly with baseline BMI for men, but did not show any such relation in women (table 2) when adjusted for age and taking into account the significant co-morbidities of BMI by using them as exclusion criteria. The increase in risk for men was 15% (95% CI 7–24%) per BMI z-score unit in 1975 and 11% (95% CI 3–20%) per BMI z-score unit in 1981. Including a quadratic or higher-order terms for BMI did not improve model fit.

After including further co-variates and changes in them into the Cox regression models (age in 1981, smoking, binge drinking, physical activity at leisure, hypertension, education, social class and income) the associations among men became weaker with the increase per BMI score in 1981 being 7% (95% CI –1 to 15%) (table 3). This relationship was not modified by hypertension (p for interaction = 0.21). When hypertension was excluded from the model the increase per BMI score was 8.5% (95% CI 1–17%).

Among men, the increase in mortality during the 22-year follow-up appeared approximately linear across all categories of baseline BMI (fig. 2), with a significant difference from the reference category only among moderately overweight subjects (BMI 27–30 kg/m²).

For women no association of BMI with mortality was seen (table 2 and fig. 2). The women BMI class BMI, age-adjusted, HR (95% CI)

<table>
<thead>
<tr>
<th>BMI class</th>
<th>1975</th>
<th>1981</th>
</tr>
</thead>
<tbody>
<tr>
<td>Underweight</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;18.5 kg/m²</td>
<td>1.22 (0.76–1.97)</td>
<td>1.43 (0.82–2.51)</td>
</tr>
<tr>
<td>Normal weight</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18.5 to &lt;22 kg/m²</td>
<td>1.00 (reference)</td>
<td>1.00</td>
</tr>
<tr>
<td>22 to &lt;23.5 kg/m²</td>
<td>0.82 (0.60–1.13)</td>
<td>0.95 (0.70–1.30)</td>
</tr>
<tr>
<td>23.5 to &lt;25 kg/m²</td>
<td>1.04 (0.77–1.40)</td>
<td>0.93 (0.67–1.30)</td>
</tr>
<tr>
<td>Overweight</td>
<td></td>
<td></td>
</tr>
<tr>
<td>25 to &lt;27 kg/m²</td>
<td>0.91 (0.64–1.30)</td>
<td>1.16 (0.84–1.60)</td>
</tr>
<tr>
<td>27 to &lt;30 kg/m²</td>
<td>0.88 (0.59–1.32)</td>
<td>0.88 (0.59–1.30)</td>
</tr>
<tr>
<td>Obese</td>
<td></td>
<td></td>
</tr>
<tr>
<td>≥30 kg/m²</td>
<td>1.07 (0.61–1.88)</td>
<td>1.06 (0.65–1.75)</td>
</tr>
</tbody>
</table>

| Per SD of BMI | 0.96 (0.87–1.07) | 0.99 (0.88–1.11) |

*Subjects with BMI 18.5 to <22 kg/m² were the reference category.*

**Table 2.** Relative risk (HR and 95% CI) of death by end of 2003 by weight category and BMI class in 1975 or 1981 for men and women. Models adjusted for age both for 1975 and 1981 BMI

**Table 3.** Relative risk (HR) of mortality by cause of death by end of 2003 per SD of BMI in 1981 for men and women. Models adjusted for age and all co-variates for all subjects and for weight-stable subjects

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had a nonsignificantly increased risk of death among underweight subjects (table 2), accounted for mainly by smoking: among never smokers, the point estimate of mortality among those with BMI <18.5 kg/m² was not increased compared with normal-weight subjects (HR 1.00, 95% CI 0.49–2.50) while among current smoker women, the HR was 1.19 (95% CI 0.57–2.51).

Among men the mortality risk increased with increasing BMI in 1981 only in those with CHD (HR 1.22 per BMI SD, 95% CI 1.06–1.41) (table 3). The risks approximately doubled from the leanest normal weight (BMI 18.5 to <22 kg/m²) to those with BMI ≥27 kg/m² in an approximately linear manner by category of BMI (table 4). The combined category of CHD and CVD also showed an increased risk (table 3). Again, this relationship between BMI and CHD risk in men was not modified by hypertension (p for interaction = 0.48). For other cause-of-death categories no significant associations were observed. For women, no associations were seen (table 3) with any causes of death.

Stable BMI and Subsequent Mortality
When these analyses were restricted to those with stable weight (less than 1 BMI unit change between 1975 and 1981), the risk remained elevated for CHD among men (HR 1.26, 95% CI 1.03–1.55); and the point estimate (not significant) for combined CHD and CVD did not change for men. Overall the risk estimates were very similar for all subjects and for weight-stable subjects only. Among the women with stable weight, BMI was inversely associated with risk of CVD, but this was based on 12 deaths only (table 3).

Analysis in Twin Pairs Discordant for All-Cause and CHD Mortality
There were no statistically significant intra-pair differences in baseline BMI or BMI at 6-year follow up among twin pairs discordant for death of all causes, whether defined as one having died and the other still alive or when comparing the first death versus later death. For pairs discordant for death from CHD, there was an increased risk of death associated with BMI z-score at baseline (crude OR 2.27, 95% CI 1.29–4.01, p = 0.005; OR 2.44, 95% CI 1.20–4.95, p = 0.014, when adjusted for physical activity, smoking, drinking, life satisfaction and hypertension) and at 6-year follow-up (crude OR 1.89, 95% CI 1.12–3.18, p = 0.017; adjusted OR 2.93, 95% CI 1.33–6.49, p = 0.008) among male pairs. No association was

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**Table 4.** HRs for CHD mortality by weight category and BMI class among men in 1975 and 1981.

<table>
<thead>
<tr>
<th>BMI class</th>
<th>1975</th>
<th>1981</th>
</tr>
</thead>
<tbody>
<tr>
<td>Underweight</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;18.5 kg/m²</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>Normal weight</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18.5 to &lt;22 kg/m²</td>
<td>1.03 (0.53–2.00)</td>
<td>1.09 (0.53–2.23)</td>
</tr>
<tr>
<td>22 to &lt;23.5 kg/m²</td>
<td>1.97 (1.09–3.57)</td>
<td>1.51 (0.78–2.93)</td>
</tr>
<tr>
<td>Overweight</td>
<td></td>
<td></td>
</tr>
<tr>
<td>25 to &lt;27 kg/m²</td>
<td>1.88 (1.02–3.46)</td>
<td>1.81 (0.94–3.46)</td>
</tr>
<tr>
<td>27 to &lt;30 kg/m²</td>
<td>2.42 (1.29–4.51)</td>
<td>1.95 (1.00–3.81)</td>
</tr>
<tr>
<td>Obese</td>
<td></td>
<td></td>
</tr>
<tr>
<td>≥30 kg/m²</td>
<td>2.00 (0.82–4.86)</td>
<td>1.83 (0.77–4.31)</td>
</tr>
</tbody>
</table>

– = Not estimated.  
Note: Models adjusted for all covariates (leisure time physical activity, smoking, hypertension, binge drinking, life satisfaction, income, schooling and social class).
to weight change and the degree of fluctuation in weight in larger studies are needed to tease out the factors contributing interest was the weight-stable part of the sample. Other and so was not examined in more detail, as the primary group of heterogeneous in the causes of the actual weight change, and to examine stable weight, as a better measure of ‘true’ BMI study, the focus in separating stable and unstable weight was stable weight may predict longevity [26, 27]. In the present earlier prospective studies on women and men, suggests that J-shaped associations with certainty. However, this study, like adjusting of the study population, resulting in few suffering from co-morbidities or being at high risk of co-morbidities throughout the range of BMI so that the population may be considered representative of a cohort of initially healthy adults. The overall mortality of our sample was considerably skewed distribution) did not independently predict risk of death overall or from CHD among men or women.

Discussion

The main finding of this study is that in the present large cohort of healthy adults with long mortality follow-up mortality appears to increase with increasing BMI only in men, in particular in those with CHD. Stable BMI showed the same relation to increased risk for CHD mortality among these initially healthy men. Analyses on twin pairs discordant for death showed that BMI differences between twins in a pair, who are more similar to each other than randomly selected people in the population, increases risk for CHD mortality in men. Several studies have reported a J- or U-shaped relationship between mortality and body weight [1, 3, 6]. However, in the present study the relationship between 22-year mortality and BMI was approximately linear in men and absent in women. This may be explained by the rigorous exclusions and adjustments of the study population, resulting in few suffering from co-morbidities or being at high risk of co-morbidities from the 1970s. During the last decades smoking has decreased in men, slightly increased in women, and overweight, obesity and weight gain relate to increased mortality of all causes and cardiovascular diseases in particular [3, 16]. Thus, we were not surprised to see overweight as a risk factor for all-cause and cardiovascular mortality, particularly among men. Observing the same association between stable BMI and later mortality confirms that the associations based on BMI values assessed in single time points are not primarily driven by the subgroups who have had changing BMI before that time point. It should be noted that the present study does not address the general level of mortality among those with stable weight versus those with unstable weight.

Our results of discordant twin pairs confirm that overweight is a risk factor for cardiovascular mortality. The earlier studies, based on co-twin control design, which we have retrieved, did not focus on weight and cause-specific mortality. Swan and Carmelli [46] studied weight changes among US male veterans who quit smoking, and Eisen et al. [33] studied smoking, alcohol consumption and obesity among 1,911 male US MZ veterans. The results from these studies show that past smokers were heavier than their co-twins. The earlier studies from the Finnish Twin Cohort based on co-twin control design have shown that smoking and sedentary lifestyle were risk factors for CHD in men, and high alcohol consumption relates to increased total mortality [32, 37]. The association between CHD mortality and weight was seen in our discordant male pairs even after adjusting for other lifestyle factors. The extent to which this relationship is influenced by shared genes could not be conclusively determined in the present study due to the relatively small numbers of MZ in the discordant pair analysis.

Observational studies may not be effective to detect the causality between mortality and BMI at large [47]. The degree of confounding by co-morbidity varies between different studies and may partly be explained by reverse causation [48]. While there is evidence for underweight and obesity being harmful for health, it is possible that modest degree of overweight without significant co-morbidities influences mortality less than expected. Recent studies have suggested that overweight may not be related to excess mortality compared to normal weight [49].

The association between obesity and cardiovascular risk factors may change over time [50]. Most importantly, the association between BMI and mortality may be significantly modified by smoking, yet the confounding effect is likely to vary significantly between different cohorts [51, 52]. The results from the present sample may not be fully representative for the situation today as the present results describe lifestyle factors from the 1970s. During the last decades smoking has decreased in men, slightly increased in women, and overweight and obesity have increased in the Finnish population.

Like in numerous earlier studies, the present analyses are based on BMI. The well-known pitfall of BMI is that it is a crude measure which does not estimate different body com-

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position compartments [53]. BMI is to a considerable degree determined by genetic factors, whereas non-shared environmental effects are greater determinants of adult weight changes [40]. Other measures of body composition, such as measures of abdominal adiposity [5], may be more specific to be used in studies on weight and mortality [54]. The relationship between different body composition measures and mortality may also vary by age [55]. The complexity of BMI-mortality relationship was studied by Grominger [56] in a semiparametric analysis concluding that BMI, or any other categorized weight variable, may give us no or false information. The obtained estimates between BMI and mortality might be secondary to both mediators between obesity and mortality as well as to causes of obesity that influence mortality independently of obesity, like diet, physical activity and cardiovascular fitness. [56, 57]. In a large male US cohort, the relationship between CVD mortality and metabolic syndrome risk factors was entirely explained by cardiovascular fitness [58]. It is possible that metabolic syndrome risk factors do cluster in the same individuals, but we did not analyze this type of clustering in our paper. The increase in CHD risk associated with BMI in men clinically healthy at baseline may reflect future higher incidence of diabetes and hypertension among overweight and obese men compared to normal-weight men, or it may reflect the inflammatory processes seen already in young adults with acquired obesity of recent origin [59].

In spite of known problems with BMI, we chose to use it as a weight variable because comparable weight measures are needed in epidemiological studies. This study is based self-reported weights, which are possibly biased. However, the comparison of self-reported and measured weights has been ascertained in a sub-sample of this cohort and found to be sufficiently valid [40]. The use of other body composition measures could complement our knowledge on BMI-mortality association, but no such data on fat distribution or body composition was available [60–62].

The advantage of the present study is that it is based on a large nationwide cohort with longitudinal data and long follow-up on mortality, with exclusions of chronic diseases done carefully by using different sources of information. The Finnish registries also provide reliable data on mortality and cause of death. Besides using the cohort as a population-based cohort, the data also allows the specific methodological co-twin control approach of twin data. The twin pair analyses complement our individual-based analyses and allow a new contribution on the role of genetic and common environment effects on BMI and mortality.

The main finding of this study is the increasing mortality, particularly due to CHD, with increasing BMI in clinically healthy men. Among women without clinically overt co-morbidities, BMI was not associated with overall mortality or cause-specific mortality. Among men, persistent acquired overweight is a considerable risk factor for CHD mortality.

 Disclosure
The authors declared no conflict of interests.

References
5 Pischon T, et al: General and abdominal adiposity measures of abdominal adiposity [5], may be more specific to be used in studies on weight and mortality [54]. The relationship between different body composition measures and mortality may also vary by age [55]. The complexity of BMI-mortality relationship was studied by Grominger [56] in a semiparametric analysis concluding that BMI, or any other categorized weight variable, may give us no or false information. The obtained estimates between BMI and mortality might be secondary to both mediators between obesity and mortality as well as to causes of obesity that influence mortality independently of obesity, like diet, physical activity and cardiovascular fitness. [56, 57]. In a large male US cohort, the relationship between CVD mortality and metabolic syndrome risk factors was entirely explained by cardiovascular fitness [58]. It is possible that metabolic syndrome risk factors do cluster in the same individuals, but we did not analyze this type of clustering in our paper. The increase in CHD risk associated with BMI in men clinically healthy at baseline may reflect future higher incidence of diabetes and hypertension among overweight and obese men compared to normal-weight men, or it may reflect the inflammatory processes seen already in young adults with acquired obesity of recent origin [59].

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