Stability and change of body mass index as a predictor of disability pension

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Abstract
Aims: To investigate whether stability or change in body mass index (BMI) predict disability pension (DP) due to
musculoskeletal diagnosis (MSD) when controlling for familial confounding. Methods: Our study cohort consisted of
were included and DPs were collected from the national pension registers until the end of 2004. Cox proportional hazards
regression models with Hazard Ratios (HR) and 95% Confidence Intervals (CI) were used for statistical analyses. Results:
General DP was granted to 2853 individuals and DP due to MSD to 1143 individuals during the 23-year follow-up. A one-
unit increase in BMI in both 1975 (HR 1.08, 95% CI 1.05, 1.10) and 1981 (HR 1.06, 95% CI 1.04, 1.07), as well as the
stability of and change in BMI from 1975 to 1981 were all associated with an increased risk of DP. These associations held
in the analyses controlling for multiple covariates (age, sex, socioeconomic status, education, marital status, leisure-time
physical activity, and musculoskeletal pain), and mainly also familial confounding, that is, genetics and shared environment.
HR for stable obesity was 2.28 (95% CI 1.69, 3.08) for DP due to MSD, and 1.91 (95% CI 1.56, 2.34) for general DP in
the fully adjusted models. Conclusions: BMI is an early predictor of general DP and also of DP due to MSD. Owing
to the independency of various covariates and potentially also familial confounding, BMI may possibly have a
direct effect on the risk of DP.

Key Words: Sickness absence, disability pension, musculoskeletal disorders, body mass index, prospective design

Background
Permanent incapacity to work carries a burden at both the societal and individual level. In many countries, people with permanent incapacity to work are granted a disability pension (DP) to compensate for decreased income. Many diseases and accidents may give rise to permanently decreased work capacity, for which reason a medical certificate is commonly required before a DP is granted. However, societal, work, and individual characteristics may enhance or diminish the impact of the disease or accident, and these are often considered when making the decision to grant a DP. One such characteristic is relative weight, which at its extreme is manifested as obesity. Obesity, which may even increase [1], is furthermore associated with the occurrence and severity of many chronic diseases [2] that are behind the granting of DPs and, therefore, cause both direct and indirect costs to society [3]. Even in the absence of concomitant diseases, severe obesity can impede work capacity. Obesity by itself can result in stigmatization at workplaces, thus contributing to a diminished work capacity.

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Epidemiological studies of DP, either as diagnosis-specific or in general, have suggested that relative weight, commonly assessed as body mass index (weight per height squared, BMI) plays a role in the process of becoming incapable of working [4–10]. However, most of these studies, even a study based on partially the same data from the Finnish Twin Cohort, have investigated BMI at one time point only, i.e. ignoring the potential change or stability of relative weight [6,7,9–11]. Furthermore, if the change in or stability of BMI has been investigated, the evaluations have been limited to comparisons between decrease and increase vs. stability. Examples of this include a study of Swedish twins with two time points, 25 years apart, preceding six years follow-up and another study of the Finnish working-age population with two time points, 5 years apart [5,12], a study with focus on temporary work incapacity in terms of sickness absence spells [12], or a study focused only on obesity, such as a very large study of over one million Swedish military conscripts [13]. Hence, there is a need to take a closer look at the relationship between BMI and work incapacity, and BMI’s contribution to the risk of DP.

A fact that needs to be taken into account is genetics, which is known to influence both BMI [14] and DP [15–17]. Therefore, it is possible that observed associations between BMI and DP are confounded by genetic or shared environmental (likely related to family background and childhood) influences, as suggested by an earlier study partially based on the same dataset [11], and a study of Swedish twins [5]. A strong method for controlling these influences are within-family comparisons, especially the co-twin control method that compares twin pairs who are discordant in the factors of interest (in this case BMI and DP). The co-twin control method also extends the case-control design, as the same-sex twins are matched in genetic background (totally in the case of monozygotic (MZ) pairs, and partially in the cases of dizygotic (DZ) pairs), parental characteristics, most childhood experiences and exposures, and have perfect matching age. Therefore, the co-twin control design is particularly useful, as it provides strong evidence of possible causal associations, compared with observational epidemiological studies of unrelated individuals. While studying discordant twin pairs, no association is expected between BMI and DP if genetics and shared environmental influences (i.e. familial factors) are of importance. Instead, if results of discordant twin pairs are similar to the results within the whole cohort (i.e. between all individuals) then factors specific to each individual are more important. The latter finding would support a causal association.

The aim of this study was to investigate whether stability or change in BMI predict DP due to musculoskeletal diagnosis (MSD), to evaluate if an association exists with DP in general, and to control whether familial confounding has an influence on these associations.

Methods

The baseline questionnaire of the Finnish Twin Cohort includes data from a comprehensive set of questions on sociodemographic, health, and lifestyle factors mailed in 1975 to all same-sex Finnish twins born between 1880 and 1957, and to both co-twins who were still alive (response rate 89%) [10,18]. In 1981, the same twins were mailed a follow-up questionnaire irrespective of whether they had responded to the baseline questionnaire (response rate 84%). A total of 24,043 twin individuals born between 1910 and 1957 were available for this study. For the analyses, only the twin individuals who responded to both questionnaires with information on BMI, had not retired from work before the date of the 1981 questionnaire, and were resident in Finland in 1981 were included. Hence, the final study sample comprised 17,169 twin individuals (52% women), including 2259 complete MZ pairs, 4545 complete DZ pairs, and 3561 twin individuals whose co-twin did not fulfill the selection criteria.

Data regarding DP was obtained from the official Finnish pension registers [15]. In the Finnish pension system, a medically confirmed illness, disease, or injury which essentially restricts or prevents working is a requirement for a disease-based early retirement pension (DP or individual early retirement pension for employees aged 58–64) to be granted. However, as part of the evaluation of work capacity, functional capacity, occupational skills, education, work tasks, and work history are always assessed. The insurance institutions make the final decision regarding a person’s work incapacity. For this study, DP in general, and DP due to MSD was selected on the basis of the International Classification of Diseases (ICD) ICD-10 codes M00–M99, and the corresponding codes in ICD-8 and ICD-9, as encoded by the Finnish insurance institutions.

Information on mortality and migration were used for the assessment of censoring the data derived from the Population Register Centre of Finland. Unique personal identification codes, which are assigned to all Finnish residents, were used for the record linkages. The follow-up time was from the date of the 1981 questionnaire to the date when DP was awarded, until the person began to receive an old age pension, to the date of death/emigration, or to 31 December 2004, whichever occurred first.
BMI (kg/m²) was computed from self-reported weight and height; the validity of self-reported BMI values in this cohort was found to be high [19]. First, BMI was analyzed as a continuous trait in 1975 and 1981 in the regression models. Then the change between 1975 and 1981 was assessed by including BMI in 1975 and BMI in 1981 in the same model to reflect on BMI level and change. For descriptive purposes, and to reflect the change between 1975 and 1981, the WHO classification was used. Based on this, BMI was categorized into < 18.5 (underweight), 18.5–25 (normal weight), 25–30 (overweight), and > 30 (obesity). The stability of and change in BMI was assessed using combinations of five categories: stable normal weight (reference); stable overweight; stable obesity; decreased BMI (any change from 1975 to a lesser BMI category in 1981); and increased BMI (any change from 1975 to a higher BMI category in 1981), including only those with full BMI data at both time points and who were eligible for follow-up. The number of individuals with stable underweight (underweight both in 1975 and 1981) was so low (< 6%) that they were omitted from the analyses.

In this study, the covariates included from the 1975 questionnaire were age, sex, and socioeconomic status based on occupation. From the 1981 questionnaire, we included: education (nine categories by years of education, converted into years of education); marital status (dichotomized to those living with someone vs. single); estimates of leisure-time physical activity measured by monthly frequency, mean duration, and mean intensity, and computed to metabolic equivalent (MET) values [20]; and musculoskeletal pain assessed by the incidence of pain in the lower back, neck or shoulder area that had affected work capacity in recent years (yes vs. no). The responses to these three musculoskeletal pain items were used to calculate a summary pain score (0–3 locations) [11].

Statistical analysis
All the analyses were performed using version 12.1 of the Stata statistical software. DP and DP due to MSD as the outcome variables were used to calculate Cox proportional HRs with 95% CIs using the follow-up time in days. Due to dependent observations in the sample – i.e. sampling twin pairs instead of unrelated individuals – pair identity was used to cluster all the analyses in order to adjust the standard errors for a lack of statistical independence within pairs [21]. In all models with the whole cohort, age was included as a continuous variable, and sex was stratified to provide men and women with their own baseline hazards to control for the effects of sex. The proportional hazard assumptions were tested by Schoenfeld residuals for BMI in 1975 and BMI in 1981 in separate models accounting for age and sex. No violations were detected.

First, we conducted basic models accounting for age and sex. In order to illustrate the linear trend in BMI in 1975 and 1981, we calculated the HRs with 95% CIs for both DP and DP due to MSD, adjusting for age and stratifying with sex for each value within each respective year’s BMI, and presented the information in a graph. Due to the low numbers of individuals scoring the highest and lowest values of BMI, values below 18 kg/m² were collapsed into a single value of 18 kg/m², and values ranging from 32 kg/m² to 47 kg/m² were collapsed into 32 kg/m². The reference group for the linear trend evaluation was set to the mean value, this being a value of 22 kg/m² for BMI in 1975, and a value of 23 kg/m² for BMI in 1981.

Next, we calculated separate models for DP and DP due to MSD, with all the covariates included in the models. The selection of covariates was due to their significance in the prediction of DP, some of which were based on earlier studies of partially the same cohort [9,11], but also due to their potential association with BMI.

The third set of analyses included twin pairs discordant for DP or DP due to MSD, applying conditional Cox proportional hazards models [22]. In these conditional models, the follow-up time to DP was analyzed in relation to the follow-up time of the co-twin. This means that one twin had a DP during the follow-up but that his/her co-twin had not been granted DP. The effects of potentially confounding familial factors can be controlled in the models when twin pairs are stratified, hence, allowing each twin pair to have their own baseline hazard. The number of discordant pairs for DP was 398 MZ and 1075 DZ pairs, whereas for DP due to MSD, it was 164 MZ and 516 DZ pairs. By using BMI as a continuous trait, we maximized the information from the available discordant pairs.

Finally, we tested the interaction between BMI and age separately for men and women, but also the interaction between BMI and sex while accounting for age. The interaction term was added to the models and the statistical significance of the interaction term was tested by the log-likelihood ratio test.

Results
Table I presents the baseline descriptive information regarding those granted DP or DP due to MSD, and those without DP during the follow-up. The mean BMI was slightly higher among men than women,
but also among those granted DP or DP due to MSD than among those with no DP in both 1975 and 1981 (Table I). The proportion of those maintaining stable normal weight (from 1975 to 1981) was 60–68% among those without DP, and 40–57% among those with DP or DP due to MSD.

Each one-unit increase in BMI in both 1975 and 1981 was significantly associated with an increased risk of DP and DP due to MSD (Table II). The linear trends for the HR with 95% CI for each value of BMI in 1975 and 1981 are shown in Figure 1 for DP, and in Figure 2 for DP due to MSD. These associations were also evident when marital status, education, socioeconomic status, leisure-time physical activity, and musculoskeletal pain locations were accounted for in the analyses, but also in the analyses that mutually adjusted the BMI in 1975 and 1981 (models 2–3, Table II). The analyses of the effect of a one-unit increase in BMI among discordant twin pairs (models 4–6, Table II) indicated independence from familial confounding, although the slightly attenuated point estimates in MZ twins suggest that the possibility of familial confounding cannot be fully ruled out.

The stability of and change in BMI from 1975 to 1981 indicated an elevated risk of both DP and DP due to MSD for all BMI categories when compared to those with stable normal weight (Table III). The association remained in the analyses when covariates were accounted for, but in the models accounting for familial confounding, the point estimates attenuated slightly in MZ twins.

Interaction between BMI and sex was not significant for DP or DP due to MSD when age was adjusted for. BMI and age showed no significant interaction for DP, but among women, the interaction was significant for DP due to MSD ($p = .009$), whereas among men it was not. The effect of BMI on DP risk was lower among younger than older women.

Table I. Frequencies and percentages of background factors and BMI categories for DP due to MSD, general DP (also including MSD), and no DP with complete BMI data from 1975 to 1981 and eligible for follow-up (1981–2004).

<table>
<thead>
<tr>
<th></th>
<th>DP due to MSD (n = 1143)</th>
<th>DP in general (n = 2853)</th>
<th>No DP (n = 14,316)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Men</td>
<td>Women</td>
<td>Men</td>
</tr>
<tr>
<td>Age at baseline (1975)</td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
</tr>
<tr>
<td>Education years (1975)</td>
<td>37.1</td>
<td>8.8</td>
<td>37.8</td>
</tr>
<tr>
<td>Leisure-time physical activity (MET)</td>
<td>6.4</td>
<td>1.6</td>
<td>6.7</td>
</tr>
<tr>
<td>BMI 1975</td>
<td>4.5</td>
<td>4.4</td>
<td>4.1</td>
</tr>
<tr>
<td>BMI 1981</td>
<td>25.0</td>
<td>2.9</td>
<td>24.0</td>
</tr>
<tr>
<td>Socioeconomic status</td>
<td>%</td>
<td>%</td>
<td>%</td>
</tr>
<tr>
<td>Upper white collar</td>
<td>3</td>
<td>2</td>
<td>7</td>
</tr>
<tr>
<td>Lower white collar</td>
<td>20</td>
<td>24</td>
<td>22</td>
</tr>
<tr>
<td>Skilled worker</td>
<td>54</td>
<td>39</td>
<td>50</td>
</tr>
<tr>
<td>Unskilled worker</td>
<td>13</td>
<td>18</td>
<td>11</td>
</tr>
<tr>
<td>Farmer</td>
<td>10</td>
<td>14</td>
<td>8</td>
</tr>
<tr>
<td>Other</td>
<td>0</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Musculoskeletal pain</td>
<td>44</td>
<td>62</td>
<td>56</td>
</tr>
<tr>
<td>Married</td>
<td>37.1</td>
<td>35.3</td>
<td>34.2</td>
</tr>
</tbody>
</table>

but also among those granted DP or DP due to MSD than among those with no DP in both 1975 and 1981 (Table I). The proportion of those maintaining stable normal weight (from 1975 to 1981) was 60–68% among those without DP, and 40–57% among those with DP or DP due to MSD.

Each one-unit increase in BMI in both 1975 and 1981 was significantly associated with an increased risk of DP and DP due to MSD (Table II). The linear trends for the HR with 95% CI for each value of BMI in 1975 and 1981 are shown in Figure 1 for DP, and in Figure 2 for DP due to MSD. These associations were also evident when marital status, education, socioeconomic status, leisure-time physical activity, and musculoskeletal pain locations were accounted for in the analyses, but also in the analyses that mutually adjusted the BMI in 1975 and 1981 (models 2–3, Table II). The analyses of the effect of a one-unit increase in BMI among discordant twin pairs (models 4–6, Table II) indicated independence from familial confounding, although the slightly attenuated point estimates in MZ twins suggest that the possibility of familial confounding cannot be fully ruled out.

The stability of and change in BMI from 1975 to 1981 indicated an elevated risk of both DP and DP due to MSD for all BMI categories when compared to those with stable normal weight (Table III). The association remained in the analyses when covariates were accounted for, but in the models accounting for familial confounding, the point estimates attenuated slightly in MZ twins.

Interaction between BMI and sex was not significant for DP or DP due to MSD when age was adjusted for. BMI and age showed no significant interaction for DP, but among women, the interaction was significant for DP due to MSD ($p = .009$), whereas among men it was not. The effect of BMI on DP risk was lower among younger than older women.
Discussion

Earlier studies have shown that BMI at one time point plays a role in the process of a person becoming incapable of working [4–11]. In addition, the few existing studies that have evaluated the change or stability of BMI over several time points have indicated a risk of temporary work incapacity [5,12]. This prospective cohort study may have been among the first to shed further light on the role of relative weight over two time points separated by six years in the risk of DP among over 17,000 twin individuals. In this study, each unit increase in BMI, and stable overweight and obesity, were both strong predictors of DP, but particularly for DP due to MSD over the 23-year follow-up. Since this prospective dataset was collected from twins, we had the unique possibility to control, not only for the measured covariates that potentially influence both BMI and DP using a standard epidemiological analysis, but also for familial confounding through the use of co-twin control design. Thus, we took into account covariates such as marital status, education, socioeconomic status, leisure-time physical activity, and musculoskeletal pain locations, but also unknown or unmeasured factors shared by co-twins, including genetic background. The co-twin control design applies analyses of discordant twin pairs, and is a powerful tool, as individuals who differed in DP and BMI within a pair were also optimally matched cases and controls, as they were
same-sexed twin pairs. The results suggest independence of the observed association from familial influences. Nevertheless, we cannot rule out the influence of familial influences due to attenuated point estimates among MZ twins. However, the results mainly remained unchanged in both analyses for each unit of BMI and analyses for BMI categories over time, reflecting stability and change between 1975 and 1981. As the analyses may have lacked power, they should be interpreted with caution.

Figure 2. Age- and sex-adjusted hazard ratios for each value of BMI in 1975 and 1981 (reference values the means, respectively) predicting the risk for disability pension due to musculoskeletal diagnoses. Error bars indicate 95% CIs.

Table III. Cox proportional hazard ratios with 95% CIs of stability and change of BMI categories between 1975 and 1981 for those granted DP or DP due to MSD during the 23-year follow-up (1981–2004).

<table>
<thead>
<tr>
<th>DP due to MSDa</th>
<th>Stable normal weight (n = 514)</th>
<th>Stable overweight (n = 297)</th>
<th>Stable obesity (n = 57)</th>
<th>Decreased BMI (n = 76)</th>
<th>Increased BMI (n = 195)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model 1: All individuals, age and sex adjusted</td>
<td>HR</td>
<td>95% CI</td>
<td>HR</td>
<td>95% CI</td>
<td>HR</td>
</tr>
<tr>
<td>Model 2: All individuals, full covariate adjustment</td>
<td>1.00</td>
<td>–</td>
<td>1.62</td>
<td>1.39, 1.89</td>
<td>2.59</td>
</tr>
<tr>
<td>Model 3: All pairs</td>
<td>1.00</td>
<td>–</td>
<td>1.44</td>
<td>1.22, 1.68</td>
<td>2.28</td>
</tr>
<tr>
<td>Model 4: DZ pairs</td>
<td>1.00</td>
<td>–</td>
<td>1.65</td>
<td>1.14, 2.39</td>
<td>2.14</td>
</tr>
<tr>
<td>Model 5: MZ pairs</td>
<td>1.00</td>
<td>–</td>
<td>1.35</td>
<td>0.61, 2.95</td>
<td>2.40</td>
</tr>
<tr>
<td><strong>DPa</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Model 1: All individuals, age and sex adjusted</td>
<td>1.00</td>
<td>–</td>
<td>1.45</td>
<td>1.32, 1.60</td>
<td>2.19</td>
</tr>
<tr>
<td>Model 2: All individuals, full covariate adjustment</td>
<td>1.00</td>
<td>–</td>
<td>1.32</td>
<td>1.19, 1.46</td>
<td>1.91</td>
</tr>
<tr>
<td>Model 3: All pairs</td>
<td>1.00</td>
<td>–</td>
<td>1.23</td>
<td>1.00, 1.52</td>
<td>1.54</td>
</tr>
<tr>
<td>Model 4: DZ pairs</td>
<td>1.00</td>
<td>–</td>
<td>1.29</td>
<td>1.02, 1.63</td>
<td>1.55</td>
</tr>
<tr>
<td>Model 5: MZ pairs</td>
<td>1.00</td>
<td>–</td>
<td>1.02</td>
<td>0.64, 1.64</td>
<td>1.43</td>
</tr>
</tbody>
</table>

aThe number of individuals (n) is given for the analyses of the whole cohort (models 1 and 2), whereas in the discordant pair analyses, the analyses were for the pairs (not shown in the table, models 3–5) and included 398 MZ and 1075 DZ pairs for DP, and 164 MZ and 516 DZ pairs for DP due to MSD.
between BMI and age may play a minor role in predicting DP, or that this interrelationship should merit further studies with an even larger sample.

Since BMI is known to have a strong genetic component [14,24] and DP is also moderately heritable [15,17,25], it was considered possible that familial influences would play a role in the associations between BMI and future DP. However, familial factors had no effect on the associations between BMI and DP. This may suggest that unique environmental factors correlate between BMI and DP. Further, the results suggest a causal association between BMI and DP. However, more studies with even larger datasets are needed to confirm these findings, as we may have lacked power, particularly in the separate analyses of MZ and DZ twins. Alternatively, the putative causal association can be tested through Mendelian randomization analyses, which also typically require very large sample sizes due to the modest effect sizes associated with known genetic variants.

Our long follow-up, 23 years for DP, indicates that BMI during the early working career may be an early predictor of subsequent work incapacity. From a public health perspective, BMI is easy to measure and is usually assessed routinely in health check-ups. One potential pathway in the association between BMI and DP might be the mediating role of societal, work, and individual characteristics that may influence the risk of disease or accident, and the consequences of these, such as DP. Mediating factors may play a role in the decision to grant DP, i.e. severe obesity may complicate working in some occupations, consequently impairing the work capacity, or being underweight may prevent participation in social activities due to stigmatization and hence, lead to increased stress and work loading. Furthermore, the curvilinear association of BMI – i.e. both the extremes (underweight and severe obesity) – are linked with diseases, but also with work capacity [4], and, therefore, should be investigated and treated for any potential underlying medical condition to clarify the role in the process of the development of work incapacity. Hence, the results emphasize the societal costs of obesity as well as the costs for the obese individuals. High BMI or BMI that shows an increasing trend could trigger a more detailed evaluation of the individual’s life situation, health behavior, and work capacity in an attempt to prevent work incapacity that could lead to DP. Across individual, organizational or societal levels, BMI would be a potential target for health promotion through tailored interventions due to the global pandemic of overweight and obesity [1].

This study has several strengths, including the use of longitudinal design with prospective follow-up from 1981 to 2004, and nationally representative high quality registry data of DP with detailed diagnoses and dates. Furthermore, we had comprehensive survey data for a large cohort of Finnish twins, which permitted us to apply co-twin control design to control for familial confounding as well as a variety of other covariates. In addition, we were able to investigate BMI data from two time points separated by six years, enabling us to evaluate stability or change in BMI with the risk of permanent work incapacity. This has rarely been done before. However, we also need to address some limitations. BMI was based on self-reported weight and height, with known downward bias [26,27], although the validity of BMI values in this cohort was high [19]. This may have diluted the observed effect compared to objectively assessed BMI data, which may be hard to obtain in large samples with tens of thousands of individuals, as in this study. Despite the relatively large dataset with over 17,000 individuals, the power was not optimal for running the analyses for MZ and DZ twin pairs separately. Another potential limitation may lay in the specific welfare system and society of Finland. Therefore, the results may be applicable to other Nordic countries that have similar welfare models and societies, but somewhat less so to other countries. Furthermore, a limitation is that our follow-up was limited to the years between 1981 and 2004. It is obvious that the work life and occupational health care have changed during these 23 years. Therefore, this prospective design should be replicated with follow-up until more recent years to establish if the association between BMI and DP would be generalized to the current working life.

Conclusions

Relative weight in terms of BMI is an early and independent predictor for DP and DP due to MSD. BMI may possibly have a causal effect on the risk of DP, although the influence of genetic confounding merits further investigation.

Conflict of interest

None declared.

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