Dysfunctional eating behaviours and personality traits in severely obese patients undergoing bariatric surgery

The PSYMO study – a randomized controlled trial

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A dissertation for the degree of Philosophiae Doctor – February 2014
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Preface

The patients who have undergone bariatric surgery at the Morbid Obesity Centre, Vestfold Hospital Trust have given birth to the PSYMO-study. The research idea was born after a couple of years of working with post-surgical follow-up groups (2 years after surgery). In almost all groups, there were patients complaining that “their heads had not yet understood that their stomachs were much smaller” and that they still craved energy-dense food that they could not eat anymore. Despite comprehensive preparations before surgery, increasingly more patients in those groups claimed that they missed their dumping episodes that had the force to “punish” them every time they ate sugar and fatty food. Furthermore, some patients could tell that just a few weeks after surgery, they found out how much chocolate they could eat at a time without activating any dumping symptoms. My clinical experience was that too many of those patients had either re-developed the same dysfunctional eating behaviours as before the surgery, or developed new maladaptive eating patterns with grazing and emotional eating. For some patients, it was as if the body signals of dumping, which appear to work as Skinnerian punishment mechanisms, were not sufficient to change their eating behaviours permanently.

In this light, bariatric surgery is not only a topic for surgeons and internists. Therefore, this project hoped to expand knowledge and understanding of how cognitive behavioural therapy may improve dysfunctional eating behaviours and affective symptoms. Additionally, we aimed to give a deeper elaboration of important psychological features and mechanisms that underlie eating behaviours and how these may contribute in the understanding of morbid obesity.
Acknowledgements

This thesis is the result of the randomized controlled trial PSYMO carried out at the Morbid Obesity centre, Vestfold Hospital Trust, Tønsberg in the period 2011-2014. This work has been supported by an unrestricted educational grant from The Norwegian Health authorities, South-East, which funded my position as a PhD student.

My deepest gratitude goes to my former leader, Pediatrician Martin Handeland from whom I have learned a lot and who let me spend time at work to study and start writing my protocol. Thank you so much, Martin!

Morbid obesity is surely a disease that you can measure in pounds and centimetres, but nevertheless, in my clinical research and practice, I have recognized that the most important issue for patients may be the illness of being morbidly obese and their perceptions of having this disease. Therefore, I want to express my deepest thankfulness to all the PSYMO participants who have contributed with their willingness, positive attitudes and hours and hours of participation so that this research could be done.

As a PhD fellow, one does not research alone! Even though I worked as a health behaviour therapist with no external funding, professor and Head of the Morbid Obesity centre and my co-supervisor, Jøran Hjelmesæth has from the very beginning been extremely encouraging and supportive in designing a new study, and helped me out writing a research protocol so that we could apply for external funding. He deserves the greatest gratitude for the invaluable contributions he has made from the early beginning until the completion of this thesis!

In the search for an experienced and excellent main supervisor in psychology, I used Google! I found that professor Jan Rosenvinge at the The Arctic University of Norway, University of Tromsø, had done a lot of interesting work in the intersection of eating disorders, eating pathology and morbid obesity. He was (and still is!) a very busy man, but he was extremely positive and helpful from the very beginning. He also connected his colleague, my co-
supervisor professor Oddgeir Friborg, to this study. Please accept my heartfelt gratitude and thanks for all your helpfulness, support and the extremely important contributions and honest feedback in this study.

For a PhD fellow doing clinical research and especially a randomized controlled trial, one is completely dependent on a very well-organised clinic with well-planned logistic. I feel a special gratitude to Astrid Hillestad for having an excellent over-view of the patients admitted for bariatric surgery, Randi S. Lund for very interesting discussions of health related quality of life in women with obesity and Mariann Mathisen (Head of the Hospital Library) to set up systematic literature searches and alerts! Especially, I am filled with profound gratitude to my former research assistant Åshild Skulstad-Hansen for organising patients’ logistics with impressive accuracy at the one year follow-up measurements. I also extend gratitude to both former and current colleagues at the Morbid Obesity Centre for their contributions and support. Especially

In keeping up the good spirit, my special thanks go to my colleagues “The very best Line Kristin Johnson”, Jan Magnus Fredheim and Njord Nordstrand – for your crazy humour and comments! My deep gratitude also goes to my colleague Dag Hofsø (“My Martin”) who always showed exemplary organisational skills and offered his sincere willingness to help out making nice figures and tables for my oral presentation in Liverpool.

I would also like to express my deep thanks to two of my previous teachers: My former French teacher in high school, Sigurd Reppesgård, and supervisor and professor at the Norwegian School of Management, Øyvind Martinsen. You are both very gifted teachers and supervisors, and you have meant more to my career than you can imagine.

To all my good friends, but first of all the group of girlfriends who I have had since childhood (“Flyfillene”): Ellen, Inger, Liv-Grethe, Rikke, Tone and Tove Gro (alphabetically sited). You have all been extremely curious, encouraging and responsive about my work. You are
simply the best! I also want to express my gratefulness to my dear friends in “the Beaujolais Nouveau group” who have given me so much joy and laughter.

My thoughts also go to my father, who died in 2001, but who gave me so much inspiration and encouragement in education and home-work when I grew up.

I am also extremely thankful for all the attention and interest in my work from my mother, my lovely daughters-in-law Eline and Kamma, my American family: “cousoeur” Lisa, her husband Jay and “my American son” Carl. Thank you, Lisa for patiently practicing with me over Skype when I was going to have my oral presentation at the European Congress of Obesity research (ECO) in 2013!

To you, my dearest Geir, my best friend, coach, mentor and husband: You are the very best!

And finally, my everlasting gratitude goes to my two sons, Sindre and Erlend. Thank you so much for being patient, helpful, funny and encouraging and for expressing that you are proud of Mum!

(I also do hope that my family will forgive me forgetting that I invited everyone over for family-dinner and that I was still working with this thesis when the dinner-guests arrived - without any dinner in the house. A PhD-project is overwhelming! It steals all the cognitive capacity that you may have.)

February, 2015

Hege Gade

*I am now doing three things: concerts, conducting, and teaching, and they each support each other. I learn to see things from different perspectives and listen with different ears. The most important thing that you need to do is really listen.*

(Itzhak Perlman)
### Abbreviations

<table>
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<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tr>
<td>BE</td>
<td>Binge eating (symptoms)</td>
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<td>BED</td>
<td>Binge eating disorder</td>
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<td>BMI</td>
<td>Body Mass Index</td>
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<td>BS</td>
<td>Bariatric surgery</td>
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<td>CR</td>
<td>Cognitive restraint</td>
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<td>DE</td>
<td>Dysfunctional eating behaviours</td>
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<td>ED(s)</td>
<td>Eating disorders</td>
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<td>EE</td>
<td>Emotional eating</td>
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<td>EB(s)</td>
<td>Eating behaviour(s)</td>
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<td>MO</td>
<td>Morbid obesity/Morbid obese</td>
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<td>RYGB</td>
<td>Roux-en-Y Gastric Bypass</td>
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<td>PSYMO</td>
<td>Psychology in morbid obesity</td>
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<td>SG</td>
<td>Sleeve gastrectomy</td>
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<td>UE</td>
<td>Uncontrolled eating</td>
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List of papers


1 Introduction

1.1 Classification and prevalence of obesity

The World Health Organization defines obesity as a chronic disease and has determined cut-off points for overweight and obesity (1). Using the Body Mass Index (BMI) (weight in kilograms divided by the square of height in meters (kg/m\(^2\))) values between 25 and 29.9 define overweight, whilst a BMI score ≥ 30 indicates obesity. Being morbidly obese indicates a BMI ≥ 40 or a BMI ≥ 35 with at least one comorbid condition (i.e. hypertension, Type 2 diabetes, or sleep apnea).

According to the Norwegian epidemiological study HUNT (2), which includes 90,000 participants, the obesity epidemic in Norway is neither slowing down nor halting. Findings from this study indicate that there has been an increase in BMI in all weight groups, and around 4-5% of the Norwegian population are estimated to have morbid obesity (MO). Moreover, the increase in BMI has been largest in the youngest age groups. These findings suggest that there is a strong demand for efficacious treatment strategies and intervention programmes that can help people to lose weight.

1.2 Eating and eating behaviours

Obesity has become a major cause of morbidity and mortality, and the obesity epidemic seems more related to an increase in energy intake than a more sedentary life style. Data from the American Food consumption surveys from 1977-2006 (3, 4) indicated that the total energy intake has increased significantly during this period, and so does the prevalence of obesity. The consumption of more energy-dense food, larger portion sizes and more frequent “snacking” and drinking have been characterised as important eating behaviours (EB)
explaining the significant increase in the prevalence of obesity (4, 5). Hence, positive energy balance is considered the most important aetiology factor for the development of obesity (6).

1.2.1 Hunger and dieting

The regulation of appetite and body-weight is a complicated process including neural systems and hunger- and appetite hormones (7). From neurophysiological and endocrinological perspectives, eating is a response to hunger- and appetite hormones like for instance ghrelin, a gut-hormone which stimulates the appetite and the growth hormone release (8). The seminal experiment *The Biology of Human Starvation* (9) presents insight in how starvation and hunger influence affective symptoms and mood, as well as a cognitive shift consisting of an intense preoccupations with thoughts of food. The participants in this study became depressed, irritable, their libido changed. However, hunger as an internal cue has a comprehensive impact on mood, cognition and EBs.

These early findings are of special interest considering how dieting may influence EB. The main change of EB in dieting is a cognitive restraint of eating, meaning that the intake of energy is below the energy expenditure. From an evolutionary perspective, dieting may be considered as an abnormal or dysfunctional eating behaviour (DE), as it in the long run may be life-threatening to consume fewer calories than needed. As an EB, dieting, or food restriction as shown in the starvation experiment (9), may influence both cognition and emotions, and in some people, the “dieting behaviour” itself also leads to overeating (10). From dieting to overeating - one DE may trigger another. The dieting may elicit shifts in cognition and a feeling of losing control when one is not completely compliant to the diet, and these cognitions and emotional states may trigger overeating. Furthermore, in persons who have had a massive weight loss, the overeating behaviour may be considered as a life-saving strategy. Moreover, in some studies, dieting behaviours have been found to predict increased risk for weight gain (11-13). According to the restraint theory (14), the self-induced restraint
of eating and overeating co-occur, and that it is the dieting that causes overeating when a break in a diet occurs. Ruining the diet temporarily seems to have a cognitive influence on the eating behaviour. The overeating may be interpreted as a physiological process, but the overeating seems to be strongly mediated by cognition (15).

1.2.2 Eating behaviours in morbidly obese

DE can be defined as exerting rigid control, or loss of control over eating, or eating for emotional reasons rather than hunger or appetite. This can aggravate to conditions close to or satisfying DSM-5-criteria for Other Specified Feeding or Eating Disorder (OSFED) or a binge eating disorder (BED). The development of obesity may be seen as a consequence of DE as the behaviour is associated with overconsumption of energy dense food (16-20). DE is maintained partly by the reinforcing effect of alleviating negative moods or experiences of stress and activation (21-23). DE has, thus been operationalized as emotional eating (EE), uncontrolled eating (UE) and cognitive restraint of eating (CR) (24). EE serves to regulate negative mood states like feeling lonely, anxious or depressed. On the other hand, people with UE tend to lose control over their eating behaviour when feeling hungry while people with CR tend to overly restrict their food intake in order to control weight, body shape or body composition. All three patterns have all been identified among MO-patients (25). Recently, Gelinas et al (26) presented findings indicating that morbidly obese patients admitted for bariatric surgery (BS) perceive eating in response to negative affect, overeating and food desirability or eating in response to positive affect/social cues as three main reasons for their weight gain. The patients’ perceptions concur with findings indicating that EE and UE are associated with higher body weight (27). Furthermore, in patients undergoing BS, less improvements in UE and EE postoperatively are related to a higher BMI (28).
Among patients with morbid obesity (MO) the DSM-IV definition of binge eating disorder (BED) vary between 6 – 49% and binge eating (BE) symptoms (of any kind) between 6-64% (29-35). Moreover, 10-25% of obese patients considered for or completing bariatric surgery (BS) have DE (33, 36-39). A recent study (40) comparing ED symptoms in normal, overweight and obese ED-patients, indicates that the obese patients display even more severe ED symptoms (poorer body-image and feelings of lacking control over food intake) than normal weight patients. Moreover, findings from the Look AHEAD-study (41) also demonstrate that participants with DE are significantly heavier than those without.

For patients eligible for bariatric surgery (BS), DE has been reported both prior (35, 42) to and after BS (21, 32, 43, 44), and DE may also impair sustained weight loss post-surgery (22, 32, 34, 45, 46). For most of those patients experiencing weight regain postoperatively, the major regain tend to start after one year post-surgically.

Thus, DE seems to be strongly related to obesity in general and weight regain in patients undergoing BS in particular, but less is known about the underlying mechanisms of DE.

1.3 What affects dysfunctional eating behaviours in morbidly obese?

1.3.1 The impact of personality

Personality is a psychological construct of large impact as it permeates most aspects of humans’ cognitive and emotional life and behaviour (47). According to Ozer et al (48), the construct of personality refers to the stable personal characteristics that are revealed in a particular pattern of behaviour, and these characteristics are able to predict individual differences in behaviour. Additionally, personality relates to factors causing disease (49), and as such also life outcomes for the individual. The mechanisms underpinning DE are poorly understood, but certain personality traits may be good candidates. There are individual
differences in how different characteristics are distributed. (47). In a representation of the Five Factor Model of personality, five dimensions represent basic tendencies of thoughts, feelings and behaviours. Those basic tendencies are **Neuroticism** (anxiety, angry hostility, depression, self-consciousness, impulsiveness and vulnerability), **Extraversion** (warmth, gregariousness, assertiveness, activity, excitement seeking, positive emotions), **Openness for experience** (fantasy, aesthetic feelings, actions, ideas and values), **Agreeableness** (trust, straightforwardness, altruism, compliance, modesty and tender-mindedness) and **Conscientiousness** (competence, order, dutifulness, achievement striving, self-discipline and deliberation). These traits may be seen as individual characteristics accounting for particular patterns of behaviour across various contexts. People with higher scores on neuroticism have more pronounced emotional dysfunction with more negative feelings which may act as drivers in adopting dysfunctional behaviours. Particularly, higher levels of neuroticism are related to an impaired ability to cope with stress and stress-related disorders. Eating patterns may then be negatively influenced by neuroticism. In addition to the genetic factors of being predisposed to weight gain, body weight reflects our EB. DE observed among morbidly obese patients seem related with personality (29, 36, 50-52), and in particular, neuroticism (52-55). Neuroticism may imply a tendency to be sensitive towards excessive availability of food (52), and in the development of MO this trait may affect a disability to adjust EB. Recent findings indicate that the facet impulsiveness may be an important risk factor for the tendency to engage in addictive food-intake (56). Thus, neuroticism may be seen as a psychological risk factor in the development of MO. According to the findings of Canetti (57), EE mediated the relation between neuroticism and body weight, meaning that there may be a causal pathway via emotionally regulated eating explaining the mechanism between neuroticism and BMI. Personality traits may have a basic influence on how the individual adapt to the environment, and the characteristic adaptations may include personal strivings and attitudes.
As personality traits are quite stable over time (58), EB are persistently influenced by these traits. Hence, personality traits, like neuroticism, may be clinically important to address in addition to mood disorders as such traits are part of the psychological makeup of an individual that always exert a potential negative effect. In the study by Terracciano et al (59), they found that personality traits were associated with both underweight and overweight. Participants with overweight showed significantly lower scores on conscientiousness and significantly higher impulsiveness than those with normal weight. Additionally, those who scored in the higher levels of impulsiveness were in average 4 kg heavier. High levels of neuroticism imply a tendency to give in to sudden temptation to eat, and may be a strong predictor of the development of obesity and DE.

It is not known how gender, mood and affective symptoms, respectively may relate to personality traits and EB. Though, in female pre-bariatric obese patients, two broad personality subtypes have been observed (60). A more well-adjusted subtype shows a profile of high emotional stability, extraversion, conscientiousness and agreeableness, while a less well-adjusted subtype may be characterized by the opposite. The latter subtype also tends to display significantly more binge eating (BE) and DE in order to regulate emotions.

Concerning successful weigh reduction maintenance, calorie restriction has been identified as the strongest predictor of a sustained weight loss (61), and in a large Australian sample, conscientiousness was found to be inversely related to obesity (62). People who succeed in maintaining weight loss and calorie restriction tend to score significantly lower on neuroticism and hostility and higher on conscientiousness (63, 64).

1.3.2 Gender and personality traits

Women constitute the majority of morbidly obese persons including those seeking BS (65), and DE is also more prevalent among women in the general population (66). Moreover, EE seems to be more frequent among morbidly obese women (22). In the obesity literature small
to negligible gender differences have been reported in how EB and personality traits may be related (55), and inconsistent findings may be due to small sample sizes and failures to include BS-patients. Findings indicate that individual differences in personality traits and EB may account for the fact that some manage long-term calorie restriction and some do not (63). Thus, those who manage such long-term calorie restriction behaviours display lower neuroticism and low responsiveness to external cues. The fact that EE is more prominent in female patients admitted for BS, and that EE may be a “high risk” EB in both the development of obesity and weight regain after surgery, expanding knowledge about gender differences is important. The previously described associations between DE and gender in patients admitted for BS are not sufficient for the understanding of this relationship. Therefore, underlying mechanisms such as personality traits, symptoms of anxiety and depression need to be explored.

1.3.3 The role of poor mental health

Substantial comorbidity of mental disorders has been reported among MO patients (36, 39, 67-73), and this may be attenuated for MO patients admitted for BS compared with MO-patients seeking conservative obesity treatment (30). Studies indicate that in patients admitted for BS, affective and mood disorders might be more prevalent in female patients (72, 74). A recent review (72), however, shows widely disparate prevalence figures, i.e. 36-72% for lifetime, and 20-55% for a current DSM-IV symptom diagnosis of notably major depressive disorder, general anxiety disorder, BED and dysthymic disorder. This diversity of prevalence rates may to some extent be accounted for by sample characteristics and the use of different assessment methods (71).

Mood disorders, such as depression, may disrupt self-regulatory sleep and EB (75), and depression in obese patients may therefore affect the appetite and craving for energy-dense
food (76). Additionally, depression is associated with higher BMI in the population of obese (30).

As indicated in *Figure 1*, mood in general may affect self-rewarding mechanisms of food (77).

![Figure 1. Complex two-way relationship linking food intake, mood and obesity](image)

*Figure by Singh, M. in Mood, food and obesity (78)*

Recent findings show that the hunger hormone ghrelin is involved in stress and reward-oriented behaviours and regulates anxiety (79). Anxiety may in turn affect stress-related eating and the motivation to eat (80). Anxiety may further prompt worries about future consequences of non-sustainable changes and a reluctance to implement behavioural changes. As discussed by others (81, 82), the role of anxiety for eating behaviour may be different than that of depression. In the period following BS, symptoms of depression usually show a larger reduction than symptoms of anxiety (83, 84).

For MO patients it may be even more complicated to regulate eating: Neurophysiology studies (85, 86) indicate that glucocorticoids, like for instance the stress hormone cortisol, stimulate appetite and may have a major effect on food intake and fat storage (87-89). We may thus expect that the higher levels of anxiety and depression symptoms, which may imply higher cortisol-levels over time, also may have an impact on eating and the development of DE. To add, there are findings demonstrating that a stress-induced EB may have an impact on
food choices (80, 89), indicating that the nutrients that are “craved” tend to be energy-dense, often consisting of sugar and fat.

1.3.4 Environmental factors

Developmentally, EB can be viewed as patterns or habits of eating which have been acquired from the observing, imitating and internalizing of care-givers’ food choices and feeding behaviours (90). Hence, the development of DE, as well as the child’s weight status, seem closely related to parents’ restrictive feeding behaviour (91), and the family environment. The availability of food has an impact on EB, and for a variety of reasons, we are tempted to eat when we are exposed to external cues like seeing or smelling food, either in commercial publicities or in real life (92, 93).

EB may as well be associated with positive emotions and cues (26) leading for instance, to more eating in familiar social settings (94).

1.3.5 Interactive processes

Environmental cues may also in turn influence the individuals’ adaptations. One example may serve to illustrate a dynamic process: A patient with a high neuroticism – score (traits) has developed low self-esteem and a feeling of being a person who is unable to control needs or impulses (adaptations). In an obesogenic environment, the patient is constantly surrounded by highly available energy-dense food which smells and looks very tasty (environmental cues).

In an obesogenic environment, the high neuroticism-score may represent a vulnerability of developing both DE and obesity. Hence, given these circumstances, a high neuroticism-score may imply practical health consequences.

To conclude, most likely, there is a complicated interplay between homeostatic regulators, heritage, internal and external cues which interact in the development of both EB in general and DE in particular. Additionally, there are also good reasons to assume that people who are genetically more disposed for weight gain, struggle even more to maintain healthier self-
regulation in eating when they are exposed to environmental food cues. Due to the fact that females are overrepresented among individuals with MO, and among patients undergoing BS, there is a need to expand our knowledge about the interplay between gender, DE and personality.

1.4 Treatment of obesity

1.4.1 Nonsurgical treatment

From a medical point of view, a successful long-term treatment is weight reduction – to achieve a significantly lower BMI over time in order to prevent serious complications like hypertension, diabetes and sleep apnea (95). A significant meaningful weight loss of ≥ 5% may be achieved in different sorts of non-surgical interventions. In the Look AHEAD-study (96) 50% of obese participants with diabetes achieved this goal through intensive interventions to change lifestyle. Using calorie restriction, physical activity and psycho-educational guidance only, life style changes among MO patients is difficult, but weight loss of ≥ 5% may be achievable (97). Researchers are still searching for new approaches in the treatment of obesity, and the study of Wadden (98) demonstrates that one third of the patients achieved long-term clinically meaningful weight loss with enhanced counselling. A clinical meaningful weight loss may result in a significant improvement of physical health. On the other hand, a weight reduction of ≥ 5% of maximum weight may not be sufficient for many patients with MO in order to lower the risk of medical complications and psychological maladjustment. For those patients, surgical treatment may be an option.

1.4.2 Surgical treatment

BS is considered when other treatments fail. Selection criteria for BS include BMI ≥ 40, or BMI ≥ 35 with at least the presence of one co-morbid condition (i.e. diabetes, hypertension or sleep apnea) and previous failures of weight loss attempts (99-101). In Norway the Roux-en-
Y gastric bypass (RYGB) and the sleeve gastrectomy (GS) (Figure 2) are the most common BS-procedures (102). In RYGB, the effects of limiting food intake and the nutrition absorbed are combined, while GS is a restrictive surgical procedure which only limits food intake (65).

![Figure 2. Roux-en-Y gastric bypass (left), gastric sleeve (right). (Figure by K. Toverud) (103)](image)

1.4.3 The outcomes treatment

1.4.3.1 Surgical treatment

Both BS-procedures are designed to restrict food intake and nutrient malabsorption, and may result in significant long-term weight loss and reduction in DE (104-107). Other important factors explaining the significant weight reduction are also the decrease of hunger and the increase of satiety (108, 109). Nevertheless, long-term weight regain has been observed in approximately 20% of the patients (110-113) starting off rather immediately (within six months) after surgery (107). DE seems to increase the risk of relapse (22, 112-115). Findings from the SOS study (28) demonstrate that a) smaller decreases in UE during the first year and b) higher levels of UE at six months and 1 year postoperatively predict less-successful long-term weight loss after BS. In contrast, CR was unrelated to both short- and long-term weight
loss. These results confirm earlier findings (116) showing that loss of control over eating in the same post-surgical time interval predicted sub-optimal weight loss after surgery. The identification of post-surgical DE in these studies may be seen in relation to the development of different weight trajectories identified by Courcoulas et al (107) showing a variability in the amount of weight loss after the first postoperative year. These weight trajectories may be related to a redevelopment of DE postoperatively.

As many patients demonstrate DE both prior to (29, 33, 36-39) and after BS (21, 32, 39, 43, 44), the former may in fact predict a failure to maintain a successful weight loss (34, 45, 46) because such DE is characterised by overconsumption of energy dense food, overweight and obesity (17, 18, 89). Several attempts to identify possible psychological predictors in explaining sub-optimal weight loss after BS (22, 51, 115, 117-121) have, however, failed in providing consistent findings due to the diversity in measurements and terminology. One study (45) reports a development of grazing eating behaviours, but with less amount of food, while Toussi (121) identified BED as a predictor, while other studies report no predictive value (116), notably with respect to weight loss (115), and rather, that suboptimal weight loss is predicted by the level of post-operative BED-symptoms. Recently, Konttinen et al (28) published ten year follow-up data from the SOS study concerning the predictive values of eating behaviours and weight change after surgery. They concluded that lower levels of uncontrolled eating at 6 months and 1 year after surgery, and greater decrease in these EB, predicted significantly better short- and long-term weight loss (28).

1.4.3.2 Non-surgical treatments

In most of the nonsurgical treatment programmes including for instance mindfulness, cognitive behavioural therapy (CBT) and various forms of therapeutic support, weight reduction is the main target. In the literature a huge variety of behavioural interventions have been studied. A main conclusion is that patients lose weight, but most patients relapse into
substantial long-term weight-regain (122), and hence, the outcome is worse than for patients receiving BS. It has been reported that approximately 20% of overweight and obese may achieve a long term weight loss of 10% from the maximum weight (123, 124). In successful long-term weight losers, improved EBs have been identified (124). This may demonstrate the difficulty in maintaining clinically meaningful weight loss, but also a treatment ineffectivity. Psychological treatments may delay weight regain, but the effects are modest (125, 126).

In contrast, the large multicentre randomised controlled trial Look AHEAD study (96), which included a manual-based intervention with CBT and motivational interviewing (MI) indicated that at least half of the patients in the intensive lifestyle group achieved a better restriction and control of their EB. The intensive intervention lasted for four years, and thereafter the participants were offered maintenance counselling. During the first year the participants were seen weekly for the first 6 months, and thereafter, group-sessions were offered every second week. In addition, individual visits were offered monthly. By comparison, the control group received educational- and social support sessions the first year in addition to the clinical diabetes visits. These results supported that a highly intensive manual-based intervention may have a favourable effects on EB and maintenance of weight loss.

1.4.3.3 Combining surgery and CBT

Concerning pre-surgical interventions, Ashton et al (127, 128) found that a brief group-based CBT-intervention improved binge eating behaviours before surgery. Additionally, a subgroup of patients who were identified as positive responders to the treatment had a significant larger weight loss 1 year after surgery. In the study of Abiles et al (129), psychological comorbidities were alleviated after a three months CBT intervention, both in patients with or without BED. In addition, other forms of pre-surgical intervention have been conducted, such
as preoperative counselling (130) which did not result in any significant group-differences in weight loss one year after surgery.

Regarding the impact of post-surgical interventions, Sarwer et al (131) found that dietary counselling had a short-term effect on DE and weight as the significant effects waned already six months following BS and stayed the same 1 year later, as well. A number of studies have examined if CBT may alleviate BED symptoms (127, 128, 132-134). As DE might represent different variations or grades of severity of binge eating behaviour and that CBT is known to be the first-line therapy of treating BED (135), one might expect that a pre-surgical CBT intervention may alleviate DE. Additionally, CBT is among the best known empirically-supported treatment for a range of mental disorders, in particular anxiety and mood disorder (136). The studies of Ashton et al (127, 128) and Abiles et al (129) showed that pre-surgical interventions based on more psychological principles may reduce both affective symptoms and DE. These studies had a non-randomised design without control groups and do not permit causal conclusions about the effectiveness of a pre-surgical CBT intervention. Therefore, there is a lack of knowledge whether CBT before surgery may have an impact both before and 1 year after surgery.

Notably, CBT results in significant improvements in psychosocial functioning, but it does not produce weight loss (137). These findings may somehow be seen as contradictory: If the patients experience more control over EB and fewer episodes of overeating, one could logically presume that they would lose weight as well. Some of the core issues concerning weight reduction and changes in DE seem to be that weight-changes are affected by a spectrum of biological factors that strongly regulate weight, and EBs might therefore be considered as a moderator in weight regulation, and not necessarily a causal factor.
Therefore, regarding surgery for obesity, one may hypothesise that a combination of CBT and BS rather than BS alone may result in better maintenance of functional EB, and psychological health and possibly, weight loss as well.

In summary, approximately 70% of all patients undergoing BS are women, and an expanded knowledge concerning gender-related differences in DE in this population is needed. Moreover, high rates of DE are found both before and after surgery. First, based on data from the SOS study, uncontrolled eating following surgery seems to predict long-term suboptimal weight loss (28). Second, manual-based intensive interventions that include CBT techniques are related to significantly better weight loss and improvements in DE in obese patients with diabetes (96). Third, pre-surgical CBT interventions addressing DE (127, 128) are expected to facilitate weight loss. With this literature review in mind, there exist, to the best of our knowledge, no RCT that has examined whether a CBT intervention that is offered before surgery may improve DE, anxiety and depressive symptoms and weight loss 1 year after BS.
2 Aims of the thesis

The aims of this thesis are:

1. To explore gender-related differences in DE in patients admitted for BS, and the mediating role of specific personality traits, and symptoms of depression and anxiety, respectively.
2. To investigate whether CBT reduced DE in patients planned for BS.
3. To investigate one year after BS whether there were statistically significant differences between the two treatment arms with regard DE, mood and anxiety symptoms and weight, respectively.
3 Materials and methods

3.1 Study designs

All data were collected at baseline (Paper I), immediately after (Paper II), and one year after (Paper III) a 10-week CBT-intervention. Paper I used a cross-sectional design, while the remaining papers used a randomised controlled design.

3.1.1 Participants

A number of 167 patients were eligible during the inclusion period from September 2011 to April 2012. Patients were consecutively accepted for BS at the Morbid Obesity Centre (Vestfold Hospital Trust), and invited to participate in the present RCT study. This is a tertiary care centre located in Tønsberg, serving approximately one million inhabitants in Southern Norway.

3.2 Procedure

Four months prior to BS, immediately after the final CBT-intervention, and at a 1-year follow-up (December 2012-November 2013) patients were referred to a web-based form collecting demographic information as well as data about personality traits, eating behaviours, and affective and depressive symptoms. The questionnaires data based on validated and standardized instruments. The anthropometric measures were collected at the hospital. After the measurements, the patients were randomised into the treatment arm, i.e. a 10-week CBT-treatment and the usual care condition, by using a web-based programme (www.randomizer.org) to allocate blocks of patients. The participants were informed about the result of the randomization immediately after screening.
3.3 Attrition

Baseline data (Paper I) comprised the 102 patients who consented to participate, randomised to CBT (N= 50) or usual care (N=52), respectively. At the post-treatment (Paper II) four patients (two in each treatment condition) were lost early in the treatments due to trial fatigue. Despite an effort to reach out to the participants by invitation letters and offers to arrange flexible meetings 14 patients refused to participate (5 in the intervention - and 9 in the control group) in the follow-up (Paper III). The flow of the study participants is shown in Figure 3.
Figure 3. Flow of patients in the PSYMO study
3.4. Measurements/instruments

3.4.1 Anthropometric measures

Anthropometric measures were collected with patients in an upright position wearing light clothing and no shoes. Waist circumference was measured at the midpoint between the lowest rib margin and the iliac crest, whilst hip circumference was measured at the level of the major trochanter. Neck circumference was measured below the larynx and perpendicular to the long axis of the neck. Waist, hip and neck circumference were measured with a tape measure to the nearest cm. Height was measured with a wall mounted stadiometer to the nearest cm, whilst weight was measured to the nearest 0.5 kg.

3.4.2 Eating behaviours

The Norwegian version of the Three Factor Eating Questionnaire (TFEQ R-21) validated for measuring dysfunctional eating behaviours in obese individuals (24, 138) was used. It consists of 21 items comprising the subscales “emotional eating” (EE; 6 items), “uncontrolled eating” (UE; 9 items) and “cognitive restraint of eating” (CR; 6 items). The domain scores range was 0-100, and with 100 indicating maximal level of dysfunction.

3.4.3 Personality traits

The Norwegian version of the NEO Personality Inventory-Revised (NEO PI-R) (139) was used. The NEO PI-R is used world-wide and is based on the Five Factor Model (FFM) of personality (140). It consists of 240 items using a five-point Likert scale to measure five domains of personality: Neuroticism (N), Extraversion (E), Openness (O), Agreeableness (A) and Conscientiousness (C). The domain scores are each composed of six facet scores, and hence, mean raw scores were created for the five domain scores and the 30 facet scores and used in the correlation/regression analyses. Higher scores indicate that a trait is more present. Gender adjusted standardized T-scores (M=50, SD=10) were used for descriptive purposes.
3.4.4 Anxiety and depression

Anxiety and depression were measured by the Hospital Anxiety and Depression Scale (HADS) (141). The HADS is a self-report measure of non-vegetative affective symptoms (141, 142) where seven items assess depression (HADS-D) and seven items measure anxiety (HADS-A), respectively. Items are scored 0-3 yielding a range of 0-21 within each subscale. A cut-off \( \geq 8 \) is used in Norway to indicate a probable diagnosis of depression or anxiety (143).

3.5 Pre-surgical interventions

3.5.1 Pre-surgery treatment

During the four months prior to surgery all patients (both groups) received up to three consultations, either from a medical doctor, a dietician, a nurse or a physiotherapist. The patients received educational materials concerning nutritional recommendations, detailed information about the mandatory low calorie diet the last three weeks before surgery as well as guidance about recommended physical activity level and intensity.

3.5.2 The CBT intervention

The intervention group received in addition a series of 10 sessions (five at the hospital and five as scheduled telephone calls to the patients at home) based on principles from self-determination theory and motivational interviewing, focusing on an accepting communication style to lower resistance and increase patients’ perceived competence and intrinsic motivation to change (144). This was focused during the first two sessions. The remaining sessions were bases on principles from CBT. Table 1 provides an overview of the contents of the sessions.
<table>
<thead>
<tr>
<th>Sessions</th>
<th>Session content</th>
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<tbody>
<tr>
<td><strong>Session 1</strong>&lt;br&gt;<em>(At the centre)</em>&lt;br&gt;<em>Both groups</em></td>
<td>• Establishing rapport with the patient in order to facilitate a good therapeutic working alliance.&lt;br&gt;• Provide information about the interventions to all patients&lt;br&gt;• Conduct the baseline measurements and perform the randomization and informing the patients about their allocated group.</td>
</tr>
<tr>
<td><strong>Session 2</strong>&lt;br&gt;<em>(At the centre)</em></td>
<td>• Introduction to the underlying principles of the therapy (working transparently, collaboratively, time-limited and using a manual)&lt;br&gt;• Inform the patient about CBT and the treatment plans in the study&lt;br&gt;• Psycho-education focusing on the relationships between eating behaviours, cognitive- and behavioral patterns, affect-regulation and obesity, thus introducing the patients for the CBT model.&lt;br&gt;• Introducing and explaining home-work sheets for session 3 and 4</td>
</tr>
<tr>
<td><strong>Sessions 3 + 4</strong>&lt;br&gt;<em>(By telephone-calls)</em></td>
<td>• Reviewing the patient’s home-work sheets&lt;br&gt;• Recognizing and addressing dysfunctional eating behaviours&lt;br&gt;• Working with the patient’s behavioral eating patterns (what triggers eating), and the associated cognitions and emotions.&lt;br&gt;• Providing the patient’s means to assess their own perception about recognizing improvement in dysfunctional cognitions and eating behaviours.</td>
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<tr>
<td><strong>Session 5</strong>&lt;br&gt;<em>(At the centre)</em></td>
<td>• Coping with situational “triggers” that may lead to dysfunctional cognitive- and eating behavioral patterns&lt;br&gt;• Working with the patient’s cognitive- and behavioral eating patterns (“triggers”, cognition, emotion and eating behaviour)&lt;br&gt;• Introducing and explaining home-work sheets for sessions 6 &amp; 7</td>
</tr>
<tr>
<td><strong>Session 6 &amp; 7</strong>&lt;br&gt;<em>(By telephone calls)</em></td>
<td>• Reviewing the patient’s home-work sheets&lt;br&gt;• Continuing the intervention techniques&lt;br&gt;• Reinforcing positive changes in eating behaviours</td>
</tr>
<tr>
<td><strong>Session 8</strong>&lt;br&gt;<em>(At the centre)</em></td>
<td>• Continuation or refining intervention techniques (as session 5) by guiding the patient in avoiding situational “triggers” and making a plan for practicing new eating behaviours.&lt;br&gt;• Introducing and explaining home-work sheets for sessions 9 &amp; 10</td>
</tr>
<tr>
<td><strong>Session 9 &amp; 10</strong>&lt;br&gt;<em>(By telephone calls)</em></td>
<td>• Reviewing the patient’s home-work sheets&lt;br&gt;• Continuation or refining intervention techniques</td>
</tr>
<tr>
<td><strong>Session 11</strong>&lt;br&gt;<em>(At the centre)</em></td>
<td>• Relapse prevention&lt;br&gt;• Ending of treatment and helping the patient to maintain positive changes</td>
</tr>
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</table>
As for the CBT intervention, the manual (Table 1) was designed to fit this particular study sample, which may provide a better fit than using manuals developed for obese patients in general or for eating disorders (134, 145, 146). However, it was based on the fundamental principle from CBT. As most patients struggle with DE both before and after surgery, a key element of the intervention was to improve DE. However, patients also struggled with other facets of eating problem, and the intervention thus also contained interventions to address the regularity of eating or to change cognitive self-talk about affect-regulated eating and thereby establishing less emotional-triggered eating. Thus, the manual was based on the principles from eating disorder specific CBT manuals, i.e. that a behavioural change is the most powerful way of changing cognitive change (147).

However, both cognitive and behavioural techniques were used, including psycho-education, and the ABC-model to counter negative automatic thoughts, rumination and worries and how cognition and emotions influence DE. In most of the sessions, monitoring records (sheets) to depict associations between situational (triggers), cognition, emotional activation and eating were extensively used. These records were also used as home-work assignments in conjunction to the hospital sessions, but new home-work tasks were not initiated during the sessions run by telephone calls. In the initial session, functional analyses were included aiming to engage the patients in the treatment, to define which roles the DE-symptoms may have in the maintenance of DE and to “transform” the patients’ symptoms of DE to useful aims of treatment.

3.6 Statistics

The SPSS version 17 was used for all statistical analyses. In all papers data are presented as mean (SD), mean (95 % CI) or number (%). Moreover, between- group differences were examined using either analysis of covariance (ANCOVA), independent sample t-test (for continuous data), linear mixed modelling analyses or Fisher’s exact tests (categorical data).
ANCOVA compares the two post-test group mean scores after adjusting for baseline scores. Effect sizes are reported as Hedges’ g indicating the differences between the groups in standard deviations. Values between 0.20 -0.49, from 0.50-0.79 and from 0.80 and higher are considered as small, moderate and high respectively (146). Associations between the continuous variables were examined with Pearson’s bivariate correlations and hierarchical multiple stepwise regression analyses. A hierarchical approach was used to assess how much of the variation in dysfunctional eating patterns that was related to three blocks of variables: 1) demographic information (i.e. age, gender, educational level status, and BMI), 2) affective symptoms (i.e. anxious and depressed mood), and 3) personality traits (i.e. NEO PI-R).

Moreover, the multiple mediation analysis was conducted (Paper I) using an SPSS macro by Preacher & Hayes (148) as it allows covariate control. Given the number of analyses, the alpha levels needed to deem a statistical test as significant were set to < .001 in the bivariate analyses (tables 1 and 2) and < .01 in the multivariate analysis.

In paper III, linear mixed regression models were built including three factors: a within factor (examining change across time), a between factor (examining differences between the treatment arms), and a final random intercept factor (adjusting for individual differences at baseline). The within factor had three levels (repeated three times: T0, T1 and T3). All outcome variables measured at baseline (T0), post CBT-intervention/pre-surgically (T1), and one year post-surgery (T2) represented time factors. The between factor had two levels: the control (G0) and intervention (G1) group. The two-way interaction (Group × Time) thus represented a test of the hypothesized treatment effect. Conducting post-hoc tests were imperative in clarifying whether there were any remaining treatment effects at T2. The restricted maximum likelihood estimation method was used.
3.7 Ethics

The study was approved by the Regional Committee for Medical and Health Research Ethics (reference number 2010/2071a), registered in the ClinicalTrials.gov-registry (identifier NCT01403558), and conducted according to the Helsinki declaration.

3.8 Funding

The PSYMO-study has been fully funded by an unrestricted research grant from the South-Eastern Regional Health Authority, Norway.
4 Results

4.1 Paper I

Psychological correlates to dysfunctional eating patterns among morbidly obese patients accepted for bariatric surgery

Our major finding was that personality traits were more firmly related to DE than mood disorder symptoms. Anxiety and depression were weakly related to such patterns when personality traits were controlled for. Neuroticism and conscientiousness were the prime personality traits involved in EE and UE/CR, and could be ascribed a vulnerable and protective role, respectively. Moreover, EE occurred more often in female patients as previously reported (22, 55, 149). A multiple mediation analysis was conducted to examine if personality, depression or anxiety played a role as mediators for the relationship between gender and EE. Personality was the only significant mediator. The initial direct path (gender-EE, controlled for BMI) was $\beta = -.46$ ($p < .001$), but dropped to $\beta = -.28$ ($p < .001$) after adding personality as the indirect path $\beta = -.18$ (CI 95% -.29 to -.09). The indirect path explained 39% (.18/.46) of the total variance in EE (Figure 4).

Figure 4. The mediation model with standardised beta coefficients
4.2 Paper II

Efficacy of a cognitive behavioural intervention on dysfunctional eating behaviours in patients admitted for bariatric surgery. A randomised controlled trial

The patients in the CBT group had significant improvements in DE, anxiety and depression compared with the control group patients. A significant reduction in BMI was also observed.

The between-group effect sizes for the improvements varied from high (uncontrolled eating \( g = -.90, p = \leq .001 \)), cognitive restraint \( g = .92, p = \leq .001 \)), to moderate (emotional eating \( g = -.67, p = \leq .001 \)), anxiety \( g = -.62, p = \leq .001 \)), depression \( g = -.73, p = \leq .001 \)) and low (BMI \( g = -.24, p = .004 \)). Adjusted between group differences at follow-up for EE, UE and CR were -19 (95% CI, -26 to -12), -19 (95 % CI, -25 to -14) and 20 (95% CI, -28 to -13) respectively, all \( p \leq .001 \). For anxiety and depression the adjusted between group differences were -2.5 (95 % CI, -3.5 to -1.4) and -2.8 (95 % CI, -3.9 to -1.6) respectively, both \( p \leq .001 \).

Concerning BMI and body weight, the adjusted between group differences were -1.1 (95 % CI, -1.8 to -.35, \( p = .004 \)), and -3 (95 % CI, -5.1 to -.84, \( p = .004 \)).

As shown in *Figure 5* and 6, the patients in the intervention group changed their EBs, affective symptoms and mood significantly in a positive direction. (These change score-figures were given as an oral presentation at the European Conference of Obesity (ECO) in Liverpool, May 2013).
4.3 Paper III

The impact of a pre-surgical cognitive behavioral therapy (CBT) on dysfunctional eating behaviours, affective symptoms and body weight one year after bariatric surgery. A randomised controlled trial

Contrary to expectations, patients in both treatment arms were indistinguishable at the one-year follow-up. Importantly, however, there were positive changes in both groups over time,
and the two groups revealed different patterns of changes in all EBs and affective symptoms.

In contrast, the patterns of weight loss did not differ between the groups.

**Dysfunctional eating behaviours**

Pairwise post-hoc tests revealed significant improvements in EE, UE and CR in the CBT group between T0 and T1 ($M_{\text{diff}} = -22.62$, $M_{\text{diff}} = -19.49$ and $M_{\text{diff}} = 25.93$, respectively, all $p$’s < .001). Further improvements between T1 and T2 were evident for EE and UE only ($M_{\text{diff}} = -8.48$, $p = .02$; $M_{\text{diff}} = -11.90$, $p < .001$, respectively).

The improvement in the control group was only evident between T1 and T2 for EE and UE ($M_{\text{diff}} = -19.97$, $p < .001$; $M_{\text{diff}} = -24.30$, $p < .001$), and between T1 and T3 for CR ($M_{\text{diff}} = 9.50$, $p = .02$).

**Group differences:** The post-hoc tests revealed significant group differences favouring CBT only at T1 for EE, UE and CR ($M_{\text{diff}} = -14.61$, $p < .01$; $M_{\text{diff}} = -15.60$, $p < .01$; $M_{\text{diff}} = -18.96$, $p < .001$, respectively), but not at T2.

**Anxiety and depression**

There was a significant reduction in anxiety and depression symptoms in the CBT group between T0 and T1 ($M_{\text{diff}} = -1.88$ and $M_{\text{diff}} = -2.69$, respectively, both $p$’s < .001), and between T1 and T2 for depression only ($M_{\text{diff}} = -1.04$, $p = .03$). In the control group, the anxiety score did not change significantly, whereas depression scores went down significantly between T1 and T2 ($M_{\text{diff}} = -2.68$, $p < .001$).

**Group differences:** No significant group differences were observed for anxiety, whereas a significant group difference favouring CBT was evident at T1 ($M_{\text{diff}} = -1.71$, $p < .01$), but not at T2.
BMI

BMI declined significantly in both groups after surgery, T1-T2 (CBT and control group; $M_{\text{diff}} = -12.60$ and $-13.79$, both $p$’s < .001), however no significant between group differences were revealed at any time point.
5 Discussion

5.1 Main findings

The main findings from this thesis were:

- The personality traits neuroticism and conscientiousness were more strongly related to DE than symptoms of mood and anxiety. Anxiety and depression were weakly related to DE when personality traits were controlled for. An additional finding was that neuroticism partially mediated this relationship, thus indicating a putative mechanism. Moreover, EE occurred more often in female patients.

- The patients in the intervention group improved their DE significantly, thus supporting the potential benefit of using CBT intervention for this group. Patients were less inclined to eat for emotional reasons and to lose control of their eating following the intervention, as well as a better ability to regulate or stop eating when satisfied. A supplementary benefit was a reduction in the symptoms of anxiety and depression. Finally, a small, yet statistically significant reduction in BMI was found.

- One year after BS, the two groups were indistinguishable in terms of weight loss, and both groups had comparable improvements in DE and symptoms of anxiety and depression. However, the onset of improvement in the CBT-group was earlier in all DE, affective and mood symptoms.

5.2 Discussions of main findings

5.2.1 The impact of CBT in the treatment of DE in patients admitted for BS

*Before surgery*, the patients in the CBT group showed a significant reduction in DE and a moderate alleviation of anxiety and depression compared to the control group. In addition,
patients in the CBT group lost about 3 kg body weight. The follow-up one year after surgery, demonstrates that the two groups had comparable improvements in all outcomes. Importantly, however, there were positive changes in both groups over time, yet with different patterns of changes in all EBs and affective symptoms.

5.2.2 CBT and changes in DE before and after surgery

Previous studies (127-129) have found effects of a CBT-intervention for BE and psychological comorbidities. These finding are relevant considering the cognitive and behavioural similarities between BE and DE as outlined in the introduction part of this thesis. Also, CBT has proven effective to alleviate affective and mood disorders (136, 150). On the one hand, our findings (Paper II) concur with the previous sparse number of comparable studies: DE and symptoms of anxiety and depression were significantly improved before surgery. On the other, these studies had a non-randomised design with no control groups, thus it is uncertain whether the improvements were caused by the CBT intervention or by other factors. Our findings are based on a randomised controlled design which provides more reliable conclusions.

Considering the purpose of establishing an evidence base for a treatment effect of a psychological intervention additive to a BS-effect, the non-significant group differences at the one-year follow-up may be discouraging. Here, several models of explanation may be possible. A delayed effect may be concealed by the relatively brief follow-up interval. The fact that the great variability in the amount and trajectory of postoperative weight loss is firstly seen two years after BS (107), may indicate that the effects of a CBT intervention may take longer to show up. Furthermore, this may indicate that the effects of the CBT intervention may be stronger when the biological effects of BS start to wane after the first year of surgery.
Another model relates to an *incomplete delivery* of the intervention. For instance, half of the 10-week intervention was given as planned telephone calls. Even though these telephone-sessions included specific CBT elements, they may have been less effective. In the absence of face-to-face communication, the scheduled telephone appointments might have been perceived less seriously and, possibly, less committing than those given in the hospital setting. This option clearly points to the role of non-specific alliance factors as well as the genuine role of face-to-face contact to attain maximal effect of the CBT interventions. Additionally, due to the relatively brief CBT intervention, case formulations were not included. This limitation may have created an intervention less tailored to individual needs, thus possibly being experienced as less personally relevant, and further reducing personal commitments. On the other hand, functional analyses included in the intervention may have countered this potential negative effect.

Several studies (28, 151, 152) show that BS has a comprehensive impact on DE, affective symptoms and weight. In the PSYMO study, the results indicate that the CBT intervention exceeded usual care in terms of *an earlier onset* of reduction of DE, anxiety and depression, which in turn indicates more functional coping with daily stress and more control over eating. These patterns of results also indicate that the advantageous improvements before surgery might be less “weight-loss dependent”, meaning that it is the significant weight loss itself, which promotes the improvements in DE, affective symptoms and mood.

An expanded understanding of DE and the change of these behaviours, the comprehensive effect of BS should be considered. BS patients have a reduced pouch after surgery. This makes it difficult to eat large portions of food, and bypassing the duodenum has most probably a major effect on the neuroendocrine regulation of hunger and appetite (153-155), DE as well as mood. In sum, these direct effects of BS may “rule out” any kind of long-term
effects from psychological interventions. As seen in the *Starvation experiment*” (9), hunger is a very strong driver and affects both cognition, emotions and DE. From a neuro-hormonal perspective, the postoperative altering of for instance the hunger hormone ghrelin might also partly explain the significant improvements in mental health in these patients, not only the weight loss itself.

5.2.3 The interrelationship between DE and body weight before surgery

Despite the significant improvements in DE in the CBT group before BS, the mean weight loss was only 3 kg. The CBT group reported significantly improvement in DE at the post-test, i.e. more restrained eating, less use of energy-dense food to regulate negative emotions, and fewer episodes of UE and impulse-driven snacking. Although the aim of the intervention was *not* weight reduction, with these improvements, one could expect that the intervention group on average would lose more than 3 kg during the 10 week intervention in terms of a significant lower intake of calories. Even in studies where weight reduction is the principal target, psychological methods *alone* might not have a significant impact on weight loss maintenance (145), meaning that weight reduction may in fact be rather resistant to psychological interventions. Studies of CBT for BE show the same tendencies: The patients report fewer episodes of DE after the therapy, but they do not lose weight (135). As discussed in part 1.3, the drivers for food intake are complex (156, 157) being controlled by signals from monitoring organs (158). This aspect may lend further support to the notion that perceiving significant improvements in DE may not automatically imply catabolic states and significant weight loss in all patients with MO.
5.2.4 Changes in body weight one year after BS

The postoperative weight outcomes may, to a certain extent, be comparable with a recent study (159) showing that including both pre- and post-surgical psychological support failed to provide significantly differences in weight loss one year after surgery. In the PSYMO study, patients in both groups reduced their weight on average approximately 40 kg. Most of the patients experience maximum weight change during the first year after BS. One may assume that the body may be resistant to further weight loss the first year after surgery (104, 160). In this regard, a pre-surgical CBT intervention will not have any additional impact beyond the surgery.

In retrospect, it might have been too optimistic to expect that the CBT intervention could have any additional effect beyond the surgery. However, additive effects may be conceived in a different way. Thus, the effect of an earlier onset of behavioural and cognitive change may be clinically important as the experience that change is possible even before the expectant change due to BS may promote self-efficacy and coping.

5.2.5 Common and specific factors

Both common and specific factors may affect the treatment outcome. The common factors include contextual factors, notably the therapist-patient alliance, which consists of the interplay between therapist and patient, including a mutual understanding of the goals of treatment and how to achieve them. It is known that satisfaction with treatment might be linked to how treatment is delivered (161, 162), and one important element in explaining therapeutic change is the quality of the therapeutic interaction between patient and therapist (163, 164).

Patients admitted for BS do definitely have common characteristics, but this is also a heterogeneous group with for instance different patterns of DE, and the way the CBT
treatment was delivered must be considered to be more flexible than strictly monitored by the manual. This implies that there may have been differences in the “doses” of the CBT treatment.

The failure not to include measurements of therapeutic alliance and manual-adherence makes it difficult to sort out common and specific effects of the CBT intervention. This opens the possibility of potent change factors unrelated to the CBT-specific treatment.

5.2.6 Gender-related differences in DE and the impact of mediators

General population studies (2, 165) show a higher probability of MO among women, and that around 70% of the patients admitted for BS are female (105). Moreover, in MO patients admitted for BS, higher rates of psychopathology are associated with female gender (72). One of the main findings (Paper I) was that female patients admitted for BS displayed significantly more DE. However, the exact nature of the relationship between gender and DE is not known. Gender differences in EBs in general may expand the understanding of gender and DE (166, 167). EBs may be explained both by genetic- and environmental factors concerning the regulation of energy balance, hunger, appetite and reward-seeking behaviour (168, 169).

One possible explanation of “the female morbid obesity-DE –axis”, is that emotion-regulated eating may be a determinant of food intake (157). EE is associated with higher consumption of sweet and fatty food, and this association is stronger in women (170, 171). A Finnish study including over 5000 adolescents showed that stress-related eating was more common in female than in male participants (172). One possible explanation is that a female vulnerability to emotion-regulated eating affects BMI, but one might also suspect the other way around – that being a women with a high BMI affects EE. The fact that morbidly obese women report having significantly lower health-related quality of life in obesity-specific emotional measurements than men (173), may also indicate that females with MO suffer significantly
more because of their weight. The negative emotional states of for instance shame, guilt and distress may in some women trigger EE.

A score $\geq 60$ on the TFRQ R-21 may be considered as dysfunctional for EE, yet this is not well established. Additional analyses (not shown in this thesis) indicate that 64% of the female patients and 15% of the males had a score of $\geq 60$ in EE at baseline. While gender-related differences were not the main focus of Paper III, additional sub-analyses showed that even one year after surgery, female patients display significantly more EE than males. In fact, there were significant correlations between postoperative EE and BMI in women but not in men.

DE observed in MO patients seems associated with personality traits (29, 36, 55) and particular with the trait neuroticism (53-55). The second major finding in Paper I demonstrates how personality traits and gender may account for individual differences in EE. The fact that patients admitted for BS have significantly higher levels of neuroticism than obese patients not admitted for surgery (174), indicates that the surgery group of MO are even further psychologically vulnerable than MO in general. The novelty of the findings indicates that neuroticism is a mechanism that partly explains the relationship between gender and EE. The mediating role of neuroticism may indicate that this trait is a key factor to modify in female bariatric surgery patients’ DE. This stands in contrast to Elfhag and Morey (55) who found negligible gender differences in the correlations between DE and personality traits, like neuroticism. A highly likely explanation of these contradictive findings is that Elfhag and Morey (55) used standardised T-scores adjusting for gender differences whereas we used unadjusted raw-scores that do not mask naturally occurring gender differences in personality traits. Affective symptoms and mood, which have been identified in other studies as possible mechanisms (175) were in our study weakly related to EE when personality traits were
controlled for. One possible interpretation is that personality permeates most aspects of humans’ cognitive and emotional life and behaviour, and it covers broader aspects of vulnerability than just depression and anxiety.

5.3 Methodological considerations

5.3.1. External validity

The target population in this study was patients with MO who were accepted for BS. The participants were consecutively admitted to surgery in a large tertiary care centre. Regarding the representativeness of the participants who were randomised, the baseline measurements concur with previous reports including BMI, gender, EB, anxiety and depression (28, 65, 151, 152, 176), personality (55, 60) and socio-economic status (177, 178). On the other hand, patients admitted for BS may be a heterogeneous population, and our findings may only be generalised to the population admitted for BS in public hospitals. Furthermore, while 167 patients were invited to participate, 53 patients declined for unknown reasons not further probed due to restrictions laid down in the ethical approval document. Another issue that may affect the external validity is “program contamination”. The patients in the CBT-group were explicitly advised not to share homework sheets, or any other materials connected to the treatment, with the patients in the control group. Despite this solicitation, we have no guarantee whether any contamination of treatment effect may have occurred, meaning that participants potentially have “shared treatment” with the control group. Being aware of the fact that many patients admitted for BS prefer connecting to a variety of “BS-groups” on Facebook, a treatment contamination may have occurred.
5.3.2 Internal validity

The recording of anthropometric measures was conducted according to standards of such measurements, and they were performed by the same researcher, which may imply consistency in these assessments.

The measurements of personality traits, EBs, affective symptoms and mood are based on self-report assessments. The use of a web-based data collection method in a structured hospital setting have strengthened the reliability by reducing the loss of variance and the need for imputation due to scattered missing data. Being requested to complete instruments displayed by a computer may in principle be unfamiliar to some patients, yet the fact that only one participant had very limited computer skills, and that assistance was offered during the completion does not represent a limitation to this study. Furthermore, the instruments are commonly used in this patient population (151, 152, 176, 179, 180) and may imply a reduction of error variance.

Another concern relevant for validity is the impact of “impression management”. In some countries like in the USA, pre-surgical evaluations may be decisive for being eligible for surgery. In these evaluations, underreporting of anxiety, depression and eating pathology has been identified (181). In the PSMO study, impression management measurements were not included, because at the time of the baseline measurements, all participants were already accepted for surgery. Therefore, we do not suspect any significant and systematic underreporting of symptoms.

In conclusion, using self-administered, well-established and validated assessment methods in a controlled hospital setting are methodological strengths.

**Paper I** had a cross-sectional design, which does not allow inferring causality. An attempt to identify predictors of DE was made, but the results of the mediation analyses need to be
replicated to generalise the findings. In considering the large number of significant test that were conducted implying an uncertainty of whether $n$ was sufficient, the alpha levels were lowered to 0.001 in the bivariate analyses and 0.01 in the multivariate regression analyses.

In Paper II, analysis of covariance (ANCOVA) was used to examine the treatment effects. By having a randomised controlled design with non-significant differences between the two groups would minimise this source of error. Moreover, not losing statistical power and to reduce the within-group error variance were the main reasons for using ANCOVA.

The calculation of sample size was based on the primary outcomes in Paper II, and the sample-size in Paper III may have been under-powered. In evaluating the treatment effects, we conducted a multilevel model using linear mixed models to study the time development in the two groups. Mixed models analyses were well-suited in the setting where repeated measurements were made on the same population, as these analyses are better suited to handle drop-out of patients over time.

In principle, the randomised controlled design makes it possible to draw causal conclusions (182). However, in the present study, the failure to include measurements of therapeutic alliance and manual adherence makes it difficult to sort out common and specific effects of the CBT intervention. This opens the possibility of potent change factors unrelated to the CBT-specific treatment.

On the one hand, the PSYMO study may be considered as a pragmatic study in the sense that it was conducted within a routine clinical context for a specified patient group (183) (i.e. MO-patients seeking BS in a specialist care centre). Moreover, a manual-based intervention was used, the attrition rates were low to moderate, and there were no statistical significant differences in baseline measurements between participants and drop-outs. Of note, there was only one researcher doing the CBT intervention, and without an additional placebo group or a
group receiving another therapy intervention, there may be certain doubts whether this is a pragmatic study.

5.4 Implications for treatment

About 4-5% of MO people in Norway qualify for BS (2). The number of patients undergoing BS is high, but most probably the amount of BS has been stable in Scandinavia during the last two years (102, 184). We consider the data from the PSYMO study to be representative for public hospital treatment-seeking morbidly obese patients admitted for bariatric surgery. As our data show, the prevalence of general psychopathology and DE is very high, and our findings of gender-related differences may imply more individualised and tailored psychological treatments for this patient population. A pre-surgical treatment based on psychological principles may improve patients’ control over food intake, reduce emotion-regulated eating as well as symptoms of anxiety and depression. Such improvements may result in better compliance with dietary recommendations.

5.5 Implications for further research

The use of RCT is sparse within this field, and should become a standard procedure in the search for treatment improvements. Also, future studies should search for clusters of personality traits which may predict relapse of DE and suboptimal weight loss or weight regain after BS.

Furthermore, the present findings highlight the need to focus on the nature and content of a CBT intervention as well as the optimal time of delivering it. Considering our one year follow-up results, it might be useful to test our CBT intervention post-surgically, as previous findings indicate that DE also may occur later than one year postoperatively and result in
impaired outcomes of BS. Finally, future studies should include a measure of therapeutic alliance to control for nonspecific factors initiating change.
6 Conclusions

This thesis may provide two main conclusions:

First, the thesis provides provisional new knowledge about how personality traits and gender may account for individual differences in EE among patients admitted for BS and the clinically relevant interplay between gender, psychological health and personality constructs.

Second, the thesis indicates an immediate impact of a pre-surgical CBT intervention on DE, affective symptoms, mood and body weight, but lends no support for a superior effect of CBT (plus BS) compared with a usual care (plus BS) at follow-up one year later.
7 References


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8 Appendix

- NEO PI-R
- Three Factor Eating Questionnaire (TFEQ R-21)
- Hospital Anxiety and Depression Scale (HADS)
- Paper I
- Paper II
- Paper III
NEO PI-R
Revidert NEO Personlighetsinventorium (NEO PI-R)
Spørsmålshefte
Paul T. Costa Jr., Ph. D. og Robert R. McCrae, Ph. D.

Instruksjoner for utfylling


På det vedlagte svarskjemaet kan du skrive navnet ditt eller en kode på den øverste linjen dersom du har blitt bedt om det. Skriv også ned datoen for utfylling, alder, kjønn, utdanning og yrke nederst på arket.

Spørreskjemaet inneholder 240 spørsmål. Les hvert spørsmål nøye, og sett en ring rundt det svaralternativet som passer best i forhold til om du er enig eller uenig i utsagnet.

Sett en ring rundt «SU» dersom utsagnet er helt uriktig eller dersom du er svært uenig.

Sett en ring rundt «U» dersom utsagnet er nokså uriktig eller dersom du er uenig.

Sett en ring rundt «N» dersom utsagnet er omtrent like riktig som uriktig, hvis du ikke kan bestemme deg, eller hvis du er nøytral i forhold til utsagnet.

Sett en ring rundt «E» dersom utsagnet stort sett er riktig eller hvis du er enig.

Sett en ring rundt «SE» dersom utsagnet er helt riktig eller dersom du er svært enig.


Besvar alle spørsmålene/utsagnene. Merk deg at alle svarene er markert nedover i kolonnene på svararket. Vær vennlig å kontrollere at svaret for hvert spørsmål blir markert på korrekt sted på svararket.

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© Norsk oversettelse ved Øyvind L. Martinsen, Hilmar Nordvik og Laila Eriksen Østbo
ISBN 82-05-32573-1
1. Jeg er ikke en person som pleier å bekymre seg.
2. Jeg liker virkelig de fleste mennesker jeg møter.
3. Jeg har en svært livlig fantasi.
5. Jeg er kjent for min Klokskap og sunne fornuft.
7. Jeg skyr menneskemengder.
8. Det som angår kunst og estetikk, er ikke spesielt viktig for meg.
10. Jeg vil heller holde mine valgmuligheter åpne enn å planlegge alt på forhånd.
11. Jeg føler meg sjelden ensom eller nedfor.
12. Jeg er dominerende, sterk og selvhevende.
13. Uten sterke følelser ville livet være uinteressant for meg.
15. Jeg forsøker å utføre alle oppgaver jeg blir pålagt på en samvittighetsfull måte.
16. I omgang med andre mennesker er jeg alltid redd for å dumme meg ut.
17. Jeg har en bedagelig stil i både arbeid og fritid.
18. Jeg har ganske fåste vaner.
20. Jeg engasjører meg lite og tar ting som de kommer.
23. Jeg liker ofte å leke med teorier eller abstrakte ideer.
24. Jeg har ingenting imot å skryte av meg selv og det jeg har utrettet.
25. Jeg er ganske flink til å tilpasse tempoet slik at jeg får gjort ting i tide.
26. Jeg føler meg ofte hjelpeløs og ønsker at andre skal løse problemene mine.
27. Jeg har aldri bokstavelig talt hoppet av glede.
28. Jeg tror at studenter bare blir forvirret og villedet av å høre kontroversielle talere.
29. Politiske ledere må være mer oppmerksomme på menneskelige sider av politikken sin.
30. I årenes løp har jeg gjort noen temmelig dumme ting.
31. Jeg blir lett skremt.
32. Jeg har ikke særlig glede av å småprate med folk.
33. Jeg forsøker å styre tankene mine inn i realistiske baner og unngå fantasiflukt.
34. Jeg tror at de fleste mennesker i bunn og grunn er velmenende.
35. Jeg tar ikke borgerpikter som f.eks. å stemme ved valg særlig alvorlig.
36. Jeg er en avbalansert person.
37. Jeg liker å ha mange mennesker rundt meg.
38. Jeg er noen ganger fullstendig oppslukt av den musikken jeg lytter til.
39. Om nødvendig er jeg villig til å manipulere folk for å få det som jeg vil.
40. Jeg holder mine åndedeler ordentlige og rene.
41. Noen ganger føler jeg meg fullstendig verdiås.
42. Noen ganger klarer jeg ikke å hevde meg så mye som jeg burde.
43. Jeg har sjelden sterke følelser.
44. Jeg forsøker å være høflig mot alle jeg møter.
45. Noen ganger er jeg ikke så pålitelig eller til å stole på som jeg burde være.
46. Jeg føler meg sjelden sjent når jeg er blant folk.
47. Når jeg gjør noe, gjør jeg det med stor iver og energi.
48. Jeg synes det er interessant å lære og utvikle nye hobbyer.
49. Jeg kan være sarkastisk og skarp når jeg må.
50. Jeg har klare mål og arbeider systematisk for å nå dem.
51. Jeg har problemer med å motstå mine lyster.
52. Jeg ville ikke like å feriere på et sted med mye natteliv, spill og underholdning.
53. Jeg synes filosofiske diskusjoner er kjedelige.
54. Jeg vil helst ikke snakke om meg selv og det jeg har utrettet.
55. Jeg kaster bort mye tid før jeg kommer i gang med arbeidet.
56. Jeg føler at jeg er i stand til å mestre de fleste av mine problemer.
57. Jeg har noen ganger opplevd intens glede eller ekstase.
58. Jeg tror at lover og velferdspolitikk burde endres for å avspeile behovene i en verden i endring.
59. Jeg er nøktern og usentimental i mine holdninger.
60. Jeg tenker gjennom ting før jeg tar en beslutning.
61. Jeg føler meg sjelden redd eller engstelig.
62. Jeg er kjent som en varm og vennlig person.
63. Jeg har et rikt fantasiliv.
64. Jeg tror de fleste mennesker vil utnytte deg hvis du lar dem gjøre det.
65. Jeg holder meg informert og tar vanligvis fornuftige avgjørelser.
66. Jeg er kjent for å være en hissig og oppførende person.
67. Jeg foretrekker vanligvis å gjøre ting alene.
68. Jeg synes det er kjedelig å se på ballett eller moderne dans.
69. Jeg kunne ikke lure noen selv om jeg ønsket det.
70. Jeg er ikke en utpreget systematisk person.
71. Jeg er sjelden trist eller deprimeret.
72. Jeg har ofte vært leder i grupper jeg har tilhørt.
73. Det er viktig for meg hva jeg føler om ting.
74. Noen mennesker ser på meg som kald og beregnende.
75. Jeg betaler min gjeld punktlig og i sin helhet.
76. Til tider har jeg vært så skamfull at jeg bare har ønsket å gjemme meg.
77. Jeg har en tendens til å arbeide langsamt, men jevnt og trutt.
78. Når jeg først finner den riktige måten å gjøre noe på, holder jeg meg til den.
79. Jeg nøler med å uttrykke mitt sinne selv når det er berettiget.
80. Når jeg starter et selvforbedringsopplegg, lar jeg det vanligvis skli ut etter noen få dager.
81. Jeg har ikke særlig problemer med å motstå fristelser.
82. Jeg har av og til gjort ting bare for «kicket» eller spenningens skyld.
83. Jeg liker å lose problemer eller «tankenøtter».
84. Jeg er bedre enn de fleste mennesker, og jeg vet det.
85. Jeg er en produktiv person som alltid får arbeidet unna.
86. Når jeg er svært stresset, føles det av og til som jeg går i stykker.
87. Jeg er ikke en munter optimist.
88. Jeg synes vi skal holde oss til våre religiøse autoriteter når det gjelder avgjørelser i moralske spørsmål.
89. Vi kan aldri gjøre for mye for de fattige og de eldre.
90. Av og til handler jeg først og tenker etterpå.
91. Jeg føler meg ofte anspent og nervøs.
92. Mange ser på meg som noe kald og fjern.
93. Jeg liker ikke å kaste bort tiden på å dagdrømme.
94. Jeg tror at de fleste mennesker jeg har med å gjøre, er ærlige og til å stole på.
95. Jeg kommer ofte opp i situasjoner uten å være fullt forberedt.
96. Jeg blir ikke betraktet som en nærtakende eller temperamentsfull person.
97. Jeg føler virkelig behov for andre mennesker når jeg er lenge for meg selv.
98. Jeg er fascinert av de mønstrene jeg finner i kunst og natur.
100. Jeg liker å ha alt på sin plass slik at jeg vet akkurat hvor det er.
101. Jeg har av og til opplevd en dyp følelse av skyld eller synd.
102. I møter lar jeg vanligvis andre stå for snakkningen.
103. Jeg legger sjelden vekt på de følelser jeg har i øyeblikket.
104. Jeg forsøker som regel å være omtenksom og hensynsfull.
105. Av og til bruker jeg når jeg legger kabal.
106. Jeg blir ikke særlig forlegen om folk gjør nar av meg og erter meg.
107. Jeg føler det ofte som om jeg strutter av energi.
108. Jeg prøver ofte ny og fremmed mat.
109. Hvis jeg ikke liker folk, lar jeg dem få vite det.
110. Jeg arbeider hardt for å nå mine mål.
111. Når jeg får livrettene mine, har jeg en tendens til å spise for mye.
112. Jeg unngår helst filmer som er sjokkerende eller skremmende.
113. Noen ganger mister jeg interessen når folk snakker om svært abstrakte, teoretiske ting.
114. Jeg forsøker å være ydmyk.
115. Jeg har problemer med å få meg selv til å gjøre det jeg burde gjøre.
116. Jeg holder hodet kaldt i krisesituasjoner.
117. Noen ganger bøbler jeg av lykke.
118. Jeg tror at de forskjellige oppfatninger av rett og galt som folk i andre samfunn har, kan være riktige for dem.
119. Jeg har ingen sympati for tiggere.
120. Jeg vurderer alltid konsekvensene før jeg handler.
121. Jeg er sjelden bekymret for fremtiden.
122. Jeg liker veldig godt å snakke med folk.
123. Jeg liker å konsentrere meg om en fantasi eller dagdrøm, utforske dens muligheter og la den vokse og utvikle seg.
125. Jeg er stolt av min gode vurderingsvevne.
126. Jeg føler ofte avsky for folk jeg er nødt til å ha med å gjøre.
127. Jeg foretrekker jobber der jeg kan arbeide alene uten å bli forstyrret av andre mennesker.
128. Dikt og poesi har lite eller ingen virkning på meg.
129. Jeg ville ha tre å bli ansett som en hykker.
130. Det virker som om jeg aldri greier å organisere meg selv.
131. Jeg har en tendens til bebreide meg selv når noe går galt.
132. Andre holder seg ofte til meg når avgjørelser skal tas.
133. Jeg opplever et bredt spekter av stemninger og følelser.
134. Jeg er ikke kjent for min gavmildhet eller sjenerøsitet.
135. Når jeg forplikter meg til noe, kan en alltid stole på at jeg følger opp.
136. Jeg føler meg ofte underlegen i forhold til andre.
137. Jeg er ikke så kvikk og livlig som andre mennesker.
139. Når jeg er blitt fornærmet, forsøker jeg bare å tilgi og glemme.
140. Det føles ikke som om noe driver meg fremover.
141. Jeg gir sjelden etter for mine impulser eller innfald.
142. Jeg liker å være der det skjer noe.
143. Jeg liker å arbeide med «tankenøtter» hvor jeg må vri hjernen.
144. Jeg har svært høye tanker om meg selv.
145. Når jeg starter på et prosjekt, fullfører jeg det nesten alltid.
146. Jeg har ofte vanskelig for å bestemme meg.
147. Jeg betrakter meg ikke som spesielt lett til sinns.
148. Jeg tror det er viktigere å være lojal mot sine idealer og prinsipper enn å ha et «åpent sinn».
149. Menneskelige behov burde alltid prioriteres fremfor økonomiske hensyn.
150. Jeg tar ofte ting på «på sparket».
151. Jeg bekymrer meg ofte for ting som kan gå galt.
152. Jeg synes det er lett å være blind og utadvendt overfor fremmede.
153. Hvis jeg føler at jeg holder på å drømme meg bort, sørger jeg som regel for å konsentrere meg om arbeid eller annen aktivitet i stedet.
154. Min umiddelbare reaksjon er å stole på folk.
155. Det ser ut som om jeg ikke lykkes helt med noe som helst.
156. Det skal mye til for å få meg sint.
157. Jeg vil heller feriere på en populær strand enn i en isolert hytte i skogen.
158. Visse typer musikk slutter aldri å fascinere meg.
159. Noen ganger lurer jeg folk til å gjøre det jeg vil de skal gjøre.
161. Jeg har ikke høye tanker om meg selv.
162. Jeg vil heller gå mine egne veier enn å være en leder for andre.
163. Jeg registrerer sjelden stemninger eller følelser som ulike omgivelser kan skape.
164. De fleste jeg kjenner, liker meg.
165. Jeg holder meg strengt til mine etiske prinsipper.
166. Jeg føler meg vel i nærver av mine sjefer eller andre autoriteter.
167. Det virker som om jeg vanligvis har det travelt.
168. Av og til gjør jeg forandringer i huset bare for å prøve noe annet.
169. Hvis noen starter en krangel, er jeg parat til å ta igjen.
171. Noen ganger spiser jeg så mye at jeg blir kvalm.
172. Jeg elsker spenningen på berg og dalbaner.
173. Jeg er lite interessert i å spekulere over universets natur eller menneskets vilkår.
174. Jeg føler ikke at jeg er bedre enn andre, uansett hvilken tilstand de er i.
175. Når et prosjekt blir for vanskelig, er jeg tilbøyelig til å starte på et nytt.
176. Jeg greier meg ganske bra i en krise.
177. Jeg er en munter, livlig person.
178. Jeg anser meg selv for å være visdyst og tolerant overfor andre menneskers livsstil.
179. Jeg mener at alle mennesker fortjener respekt.
180. Jeg tar sjelden forhastede avgjørelser.
181. Jeg har færre redsler enn folk flest.
182. Jeg er sterkt følelsesmessig knyttet til vennene mine.
183. Som barn likte jeg sjelden fantasilek.
184. Jeg pleier å tro det beste om folk.
185. Jeg er en svær kompetent person.
186. Til tider har jeg følt meg harm og forbiritet.
187. Sosiale sammenkomster er vanligvis kjedelige for meg.
188. Noen ganger når jeg leser dikt eller ser et kunstverk, føler jeg en gysning eller en bølge av begeistring.
189. Til tider herser eller snisker jeg med folk for å få dem til å gjøre det jeg vil.
190. Jeg er ikke tvangsmessig opptatt av rengjøring.
192. I samtal er jeg en tendens til å stå for det meste av snakkingen.
194. Jeg ser på meg selv som en nestekjærlig person.
195. Jeg forsøker å gjøre ting nøyaktig slik at de ikke må gjøres om igjen.
196. Hvis jeg har sagt eller gjort noe galt mot noen, har jeg fryktelig vanskelig for å se dem i øynene etterpå.
197. Livet mitt er hektisk.
198. Når jeg skal feriere, foretrekker jeg å dra tilbake til et kjent og kjært sted.
199. Jeg er sta og egenrådig.
200. Jeg streber etter å gjøre det utmerket i alt jeg gjør.
201. Noen ganger gjør jeg ting på impuls som jeg senere angår på.
203. Jeg har mye intellektuell nysgjerrighet.
204. Jeg vil heller rose andre enn selv bli rost.
205. Det er så mange småjobber som må gjøres at jeg av og til ganske enkelt ignorerer dem.
206. Når alt ser ut til å gå galt, kan jeg likevel ta gode avgjørelser.
207. Jeg benytter sjelden ord som «fantastisk» eller «sensasjonelt» for å beskrive mine opplevelser.
208. Jeg mener at hvis folk ikke vet hva de tror på når de er 25, er det noe galt med dem.
209. Jeg har sympati for dem som har vært mindre heldige enn meg.
211. Av og til får jeg skremmende tanker.
212. Jeg er personlig interessert i de menneskene jeg arbeider sammen med.
213. Jeg ville ha vansker med bare å la tankene vandre uten kontroll eller styring.
214. Jeg har stor tiltro til menneskets natur.
215. Jeg er effektiv og produktiv i mitt arbeid.
216. Selv mindre ergelser kan være frustrerende for meg.
217. Jeg liker selskaper med mange mennesker.
218. Jeg liker å lese dikt som legger vekt på følelser og forestillinger fremfor rene fortellinger.
219. Jeg er stolt av min skarpsindighet i å håndtere mennesker.
220. Jeg bruker mye tid på å lete etter ting jeg har forlagt.
221. Når noe går galt, blir jeg altså ofte motløs og får lyst til å gi opp.
222. Det er ikke lett for meg å ta føringen i en situasjon.
223. Merkelige ting – som visse dufter eller navn på fjerne steder – kan vekke sterke sinnsstemma
n i meg.
224. Jeg strekker meg langt for å hjelpe andre mennesker hvis jeg kan.
225. Jeg må virkelig være syk for å bli borte fra jobben en dag.
226. Når folk jeg kjenner gjør noe dumt, blir jeg flau på deres vegne.
227. Jeg er en svært aktiv person.
228. Jeg følger alltid den samme ruten når jeg skal et sted.
229. Jeg kommer ofte opp i argumentering/uenighet med min familie og mine kolleger.
230. Jeg er litt av en arbeidsnarkoman.
231. Jeg er alltid i stand til å holde mine følelser under kontroll.
232. Jeg liker å være i tilskuermengden ved sportsarrangører.
233. Jeg har et vidt spekter av intellektuelle interesser.
234. Jeg er en suveren person.
235. Jeg har stor selvdisciplin.
236. Jeg er ganske følelsessmessig stabil.
237. Jeg har lett for å le.
238. Jeg mener at den «nye moral», som tillater det meste, ikke er noen moral i det hele tatt.
239. Jeg vil heller være kjent som «barmhjertig» enn som «rettfærdig».
240. Jeg tenker meg om to ganger før jeg svarer på et spørsmål.

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<tr>
<th>Avsnitt</th>
<th>Påstand</th>
<th>Svarer 1</th>
<th>Svarer 2</th>
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<th>Svarer 4</th>
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<tr>
<td>1.</td>
<td>Jeg tar med hensikt små porsjoner for å holde kroppsvekten nede.</td>
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<td>2.</td>
<td>Når jeg føler meg urolig, oppdager jeg ofte at jeg spiser.</td>
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<td>3.</td>
<td>Av og til når jeg begynner å spise, er det akkurat som om jeg ikke klarer å slutte.</td>
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<td>4.</td>
<td>Når jeg føler meg nedstemt, spiser jeg ofte for mye.</td>
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<td>5.</td>
<td>Jeg unngår visse typer mat fordi de er fetende for meg.</td>
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<td>Stemmer ikke i det hele tatt</td>
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<td>7.</td>
<td>Når jeg er ansprent eller ”oppgiret”, føler jeg ofte trang til å spise.</td>
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<td>Stemmer helt</td>
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<td>Stemmer ikke i det hele tatt</td>
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<tr>
<td>8.</td>
<td>Jeg får ofte så lyst på mat at magen føles som et stort hull som ikke kan fylles.</td>
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<tr>
<td>9.</td>
<td>Jeg har alltid lyst på mat, så det er vanskelig for meg å slutte å spise før jeg har spist opp alt på tallerkenen.</td>
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<tr>
<td>10.</td>
<td>Når jeg føler meg ensom, trøster jeg meg selv med å spise.</td>
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<td></td>
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<td>Stemmer ikke i det hele tatt</td>
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<tr>
<td>11.</td>
<td>Jeg holder bevisst igjen ved måltidene for å ikke gå opp i vekt.</td>
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<td>Stemmer ikke særlig bra</td>
<td>Stemmer ikke i det hele tatt</td>
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</tr>
<tr>
<td>12.</td>
<td>Når jeg kjenner lukten av en biff som steokes eller ser en saftig kjøttbit, er det veldig vanskelig å la være å spise selv om jeg akkurat har avsluttet måltidet.</td>
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<tr>
<td></td>
<td>Stemmer helt</td>
<td>Stemmer ganske bra</td>
<td>Stemmer ikke særlig bra</td>
<td>Stemmer ikke i det hele tatt</td>
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</tr>
</tbody>
</table>

Sett ett kryss i avkrysningsboksen til venstre for det svaret som passer best.

   ☐ Stemmer helt
   ☐ Stemmer ganske bra
   ☐ Stemmer ikke særlig bra
   ☐ Stemmer ikke i det hele tatt

14. Hvis jeg kjenner meg ille til mote, forsøker jeg å dempe ubehaget med å spise.
   ☐ Stemmer helt
   ☐ Stemmer ganske bra
   ☐ Stemmer ikke særlig bra
   ☐ Stemmer ikke i det hele tatt

15. Når jeg ser noe som ser veldig godt ut, får jeg ofte så lyst på det at jeg må det spise med en gang.
   ☐ Stemmer helt
   ☐ Stemmer ganske bra
   ☐ Stemmer ikke særlig bra
   ☐ Stemmer ikke i det hele tatt

   ☐ Stemmer helt
   ☐ Stemmer ganske bra
   ☐ Stemmer ikke særlig bra
   ☐ Stemmer ikke i det hele tatt

17. Hvor ofte unngår du å ha fristende mat tilgjengelig?
   ☐ Nesten aldri
   ☐ Sjelden
   ☐ Ofte
   ☐ Nesten alltid

18. Hvor sannsynlig er det at du bevisst spiser mindre enn det du vil ha?
   ☐ Usannsynlig
   ☐ Ikke særlig sannsynlig
   ☐ Ganske sannsynlig
   ☐ Veldig sannsynlig

19. Fortsetter du å spise selv om du ikke er sulten længer?
   ☐ Aldri
   ☐ Sjelden
   ☐ Iblant
   ☐ Minst en gang i uken

20. Hvor ofte har du lyst på mat?
   ☐ Bare til måltidene
   ☐ Iblant mellom måltidene
   ☐ Ofte mellom måltidene
   ☐ Nesten alltid

21. På en skala fra 1 til 8, der 1 står for ingen begrensning (spiser hva jeg vil, når jeg vil) og 8 står for streng begrensning (begrenser alltid matinntaket, gir aldri etter), hvor på skalaen befinner du deg?
   Sett en ring rundt det tallet som passer best for deg.

   1  2  3  4  5  6  7  8

   Spiser hva jeg vil, når jeg vil

   Begrenser alltid matinntaket, gir aldri etter

### HAD

**Hospital Anxiety & Depression Scale (januar 1999)**

<table>
<thead>
<tr>
<th>Navn: ___________________________</th>
<th>Fødselsdato: ________________</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dato for utfylling: _______________</td>
<td>Pasient nr.: ________________</td>
</tr>
<tr>
<td>Behandler: _________________________</td>
<td></td>
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</tbody>
</table>

#### Rettledning

Legen er klar over at følelser spiller en stor rolle ved de fleste sykdommer. Hvis legen vet mer om følelser, vil han/hun bli bedre i stand til å hjelpe deg.

Her kommer noen spørsmål om hvorledes du føler deg. For hvert spørsmål setter du kryss for ett av de fire svarene som best beskriver dine følelser den siste uken. Ikke tenk for lenge på svaret – de spontane svarene er best.

1. **Jeg føler meg nervøs og urolig**
   - 3 Mesteparten av tiden
   - 2 Mye av tiden
   - 1 Fra tid til annen
   - 0 Ikke i det hele tatt

2. **Jeg gleder meg fortsatt over tingene slik jeg pleide før**
   - 0 Avgjort like mye
   - 1 Ikke fullt så mye
   - 2 Bare lite grann
   - 3 Ikke i det hele tatt

3. **Jeg har en urofølelse som om noe forferdelig vil skje**
   - 3 Ja, og noe svært ille
   - 2 Ja, ikke så veldig ille
   - 1 Litt, bekymrer meg lite
   - 0 Ikke i det hele tatt

4. **Jeg kan le og se det morsomme i situasjoner**
   - 0 Like mye nå som før
   - 1 Ikke like mye nå som før
   - 2 Avgjort ikke som før
   - 3 Ikke i det hele tatt

5. **Jeg har hodet fullt av bekymringer**
   - 3 Veldig ofte
   - 2 Ganske ofte
   - 1 Av og til
   - 0 En gang i blant

6. **Jeg er i godt humør**
   - 3 Aldri
   - 2 Noen ganger
   - 1 Ganske ofte
   - 0 For det meste
7. Jeg kan sitte i fred og ro og kjenne meg avslappet

- 0 Ja, helt klart
- 1 Vanligvis
- 2 Ikke så ofte
- 3 Ikke i det hele tatt

8. Jeg føler meg som om alt går langsommere

- 3 Nesten hele tiden
- 2 Svært ofte
- 1 Fra tid til annen
- 0 Ikke i det hele tatt

9. Jeg føler meg urolig som om jeg har sommerfugler i magen

- 0 Ikke i det hele tatt
- 1 Fra tid til annen
- 2 Ganske ofte
- 3 Svært ofte

10. Jeg bryr meg ikke lenger om hvordan jeg ser ut

- 3 Ja, jeg har sluttet å bry meg
- 2 Ikke som jeg burde
- 1 Kan hende ikke nok
- 0 Bryr meg som før

11. Jeg er rastløs som om jeg stadig må være aktiv

- 3 Uten tvil svært mye
- 2 Ganske mye
- 1 Ikke så veldig mye
- 0 Ikke i det hele tatt

12. Jeg ser med glede frem til hendelser og ting

- 0 Like mye som før
- 1 Heller mindre enn før
- 2 Avgjort mindre enn før
- 3 Nesten ikke i det hele tatt

13. Jeg kan plutselig få en følelse av panikk

- 3 Uten tvil svært ofte
- 2 Ganske ofte
- 1 Ikke så veldig ofte
- 0 Ikke i det hele tatt

14. Jeg kan glede meg over gode bøker, radio og TV

- 0 Ofte
- 1 Fra tid til annen
- 2 Ikke så ofte
- 3 Svært sjelden

_Takk for utfyllingen!

Sum A: 

\[1+3+5+7+9+11+13= \text{__________}

Sum D: 

\[2+4+6+8+10+12+14= \text{__________}

Sum A + D: 

\[\text{__________}\]
Selvutfylling på sju angst- og depresjonsspørsmål.

**Sum A eller Sum D:**
En skår på 11 eller mer regnes for å være et tilfelle av angst eller depresjon som vil trenge nærmere utredning (med SPIFA for eksempel) og eventuelt behandling. En skår på 8-10 anses som et mulig tilfelle, og lavere skår uttrykker en viss symptombelastning, som kan ha betydning samlet sett, men som i seg selv ikke krever spesifikk behandling av angst eller depresjon.

**Sum A + Sum D:**

Dersom inntil to spørsmål på HAD er ubesvart, vil det være mulig å beregne totalskår. Sumskåren deles med antallet besvarte spørsmål og svaret ganges med 14. Dette gir estimert totalskår.

Referanser: