Review
Can Weight Loss Improve the Cardiovascular Outcomes of Patients with Obesity and Obstructive Sleep Apnea?

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Abstract: Cardiovascular events are the primary cause of mortality in patients with obstructive sleep apnea and obesity. The rising prevalence of obstructive sleep apnea in recent decades has been linked to increasing rates of obesity. Obstructive sleep apnea has also been linked with many different cardiovascular diseases including coronary artery disease, stroke, heart failure, hypertension, and atrial fibrillation. Obesity is an increasing health concern globally, in part because obesity complications such as hypertension, diabetes, and obstructive sleep apnea increase the risk of cardiovascular diseases. More than 10% weight loss may be required to prevent or reverse obesity complications. Treatment approaches to obesity include nutritional therapy, exercise therapy, pharmacotherapy, and surgical therapies. This review intends to identify the effects of weight loss on cardiovascular outcomes in patients with obesity and obstructive sleep apnea. Despite the strong association between cardiovascular diseases and obstructive sleep apnea, randomized trials have failed to demonstrate that treatment of obstructive sleep apnea reduces cardiovascular events, even in patients with established cardiovascular diseases. Weight loss in patients with obstructive sleep apnea improves HbA1c, systolic blood pressure, HDL cholesterol, and triglycerides, but thus far no changes in cardiovascular events have been shown. The combination of weight loss with continuous positive airway pressure (CPAP) appears more beneficial than either treatment in isolation. Large well-controlled trials in patients with obstructive sleep apnea to assess the effects of different weight reduction programs on cardiovascular disease are still needed.

Keywords: obstructive sleep apnea; obesity; cardiovascular diseases; weight loss

1. Introduction
Obstructive sleep apnea is characterized by repetitive upper airway collapse during sleep resulting in a temporary cessation of breathing (apnea), shallow breathing (hypopnea), or respiratory-related arousals. The apnea–hypopnea index measures the number of apnea and hypopnea events per hour of sleep. The apnea–hypopnea index is used to further classify obstructive sleep apnea as mild (5 ≤ apnea–hypopnea index < 15), moderate (15 ≤ apnea–hypopnea index < 30), or severe (apnea–hypopnea index ≥ 30) [1]. The pathophysiology of obstructive sleep apnea is commonly believed to be primarily due to anatomical anomalies of the upper airway. While a defect in the upper airway anatomy is common, it is not the only pathogenic process at work. Anatomical and non-anatomical elements have been included in a novel model of obstructive sleep apnea pathogenesis. These non-anatomical aspects include (1) impairments in the upper airway dilator muscle function during sleep, (2) respiratory control instability, and (3) low respiratory arousal threshold [2].

Obstructive sleep apnea has a global prevalence of 13–33% in middle-aged men and 6–19% in middle-aged women [3]. The rising prevalence of obstructive sleep apnea in the
recent decades has been linked to increasing rates of obesity [4]. Obstructive sleep apnea has been linked with many different cardiovascular diseases including hypertension, stroke, heart failure, coronary artery disease, and atrial fibrillation [5,6]. Adults with obstructive sleep apnea, in addition to increased risk of developing cardiovascular disease, also have worse cardiovascular outcomes [7]. We reviewed the associations between obstructive sleep apnea and cardiovascular disease and whether weight loss may improve the cardiovascular outcomes of patients with obstructive sleep apnea and obesity.

2. Obstructive Sleep Apnea Increases the Risk Factors of Cardiovascular Disease

Among the mechanisms which explain the association between obstructive sleep apnea and myocardial infarction (MI), common risk factors include male sex, age, hypertension, obesity, and smoking [8]. However, other direct effects of obstructive sleep apnea merit consideration. The combination of repetitive apnea–hypopnea, hypoxia, and arousal from sleep increases sympathetic activity, which is maintained during wakefulness, thus increasing myocardial oxygen demand [9]. The mechanistic understanding which connects obstructive sleep apnea and cardiovascular disease is poorly understood due to the diverse and complicated elements of obstructive sleep apnea and the multiple other comorbid conditions (especially obesity) impacting cardiovascular health. When obstructive apnea or hypopnea occurs, the upper airway collapses throughout sleep, affecting a complete or partial interruption of airflow even with sustained respiratory struggle. The sympathetic tone is stimulated, and respiratory work increases as opposed to the closed upper airway, increasing negative intrathoracic pressure. Stimulation of the sympathetic tone across the parasympathetic system affects heart rate and blood pressure [10]. Awakening from sleep terminates the asphyxia event, with re-establishing airflow and re-oxygenation but further increased sympathetic tone. Obstructive sleep apnea seems to be correlated with increased levels of inflammatory cytokines. Furthermore, metabolic dysregulation is observed in obstructive sleep apnea patients (with abnormalities in both fat and glucose metabolism. This contributes to atherosclerosis and endothelial damage with enhanced arterial stiffness. However, it remains unclear whether metabolic abnormalities and inflammation are exacerbated by obstructive sleep apnea, whether these are epiphenomena are due to obesity [10].

2.1. Hypertension

Obstructive sleep apnea is common in middle-aged and older people, although hypertension is also prevalent among middle-aged and older people. This increases the chance of significant complications between hypertension and obstructive sleep apnea. The level of the complications of hypertension and obstructive sleep apnea is significantly greater than expected. Hypertension is prevalent in patients with obstructive sleep apnea and contributes to vascular injury and cardiovascular events. Several pathophysiologic mechanisms contribute to the increased risk of hypertension in individuals with obstructive sleep apnea, including upregulation of neurohormonal pathways, endothelial dysfunction, and inflammation [6]. Although patients with obstructive sleep apnea have a higher incidence of hypertension [11], the inverse is also true because patients with hypertension are more likely to experience sleep-disordered breathing, especially those who have failed to respond to traditional treatment. Up to 84% of this subset of patients may have undetected obstructive sleep apnea [12].

Animal experiments have provided direct proof that obstructive sleep apnea causes hypertension. When obstructive sleep apnea is induced, it results in acute transient rises in nighttime blood pressure and ultimately culminates in persistent daytime hypertension [11]. When blood pressure is strictly regulated in rats, it also decreases sleep apneas [13].

In humans, a causal link between obstructive sleep apnea and hypertension has not been established because variables such as age and obesity confound the association. However, epidemiological data suggest that hypertension was found in approximately 50% of patients with obstructive sleep apnea (7).
2.2. Dyslipidemia

Obstructive sleep apnea is commonly associated with elevated plasma triglycerides, low-density lipoprotein cholesterol (LDL-c), and total cholesterol. Moreover, the reduction in high-density lipoprotein (HDL) may, in part, be due to deleterious oxidative processes commonly found in patients with obstructive sleep apnea [14–16]. The effect of treating obstructive sleep apnea in children with adenotonsillectomy is variable as regards the impact on lipid profiles [17] because chronic intermittent hypoxia may affect both lipid biosynthesis and lipid peroxidation [18].

2.3. Type 2 Diabetes

Inflammation in patients with type 2 diabetes is characterized by elevated levels of proinflammatory cytokines or a rising number of white blood cells in the blood or tissue. Stimulation of the inflammatory process often indicates abnormalities such as tissue injury and organ dysfunction. Obesity might cause chronic low-grade inflammation and hat is involved in type 2 diabetes. In addition, adipose-specific cytokines (leptin adiponectin, leptin, and interleukin 6 (IL-6)) are secreted by visceral adipocytes and inflammatory cytokines (tumor necrosis factor α (TNF-α)). An elevated amount of fatty tissue draining into the chemokines, portal vein, and IL-6 production can induce liver and systemic insulin resistance [19,20]. Although obesity is a serious risk factor for type 2 diabetes mellitus, coexistent severe obstructive sleep apnea may independently add to the risk. The relationship between intermittent hypoxia and insulin resistance [21] has been assessed by evaluating the effect of hypoxia–reoxygenation cycles on insulin target tissues. Rodent models suggest that chronic exposure to intermittent hypoxia induces insulin resistance [22].

2.4. Atrial Fibrillation

Obstructive sleep apnea is a common risk factor for atrial fibrillation [23]. Recurrent episodes of obstructive sleep apnea may lead to cardiac structural and electric remodeling. Repetitive episodes of obstructive sleep apnea in an animal model can cause atrial fibrosis and important changes in connexin-43 distribution and expression, thus leading to slow atrial conduction. This increases the vulnerability to arrhythmias, including atrial fibrillation [24]. Furthermore, untreated obstructive sleep apnea doubles the risk of recurrence of atrial fibrillation in patients after electrical cardioversion. Treatment of obstructive sleep apnea with CPAP attenuates the risk of atrial fibrillation [25]. Obstructive sleep apnea shares many common risk factors with atrial fibrillation. The prevalence of both atrial fibrillation and obstructive sleep apnea is rising due to increases in obesity and cardiovascular disease. The close association between obstructive sleep apnea and cardiovascular disease, and atrial fibrillation and cardiovascular disease may obscure a directly causal relationship between atrial fibrillation and obstructive sleep apnea. The interplay of the pathophysiology of these chronic diseases is complex and likely bidirectional. Obstructive sleep apnea may contribute to atrial fibrillation, and, in turn, atrial fibrillation promotes the development of obstructive sleep apnea. Nonetheless, these entities are associated with one another, independently of other cardiovascular diseases [26].

2.5. Heart Failure

Sleep apnea is predictable in patients with heart failure, with a prevalence of between 50% and 70% [27]. Mainly, central sleep apnea accounts for two-thirds of the sleep apnea cases in this population, while obstructive sleep apnea is less frequent. Central sleep apnea is a frequent concomitant finding in patients with severely impaired cardiac function [27]. Another study aimed to assess the prevalence of sleep-disordered breathing and its associated risk factors in French patients with heart failure showed that 30% of syndromes were classified as central and 70% as obstructive [28]. Coexisting sleep apnea in patients with heart failure has been associated with an increased risk of adverse outcomes, including mortality [29]. Several pathophysiological processes resulting from apneic events may explain this association. These involve stimulation of the sympathetic nervous system [30]
and increased preload and afterload resulting from perturbation of intrathoracic pressure while struggling to inspire against blocked airways [31]. Worsening hypertension, increased risk of arrhythmias including sudden cardiac death [32], and myocardial infarctions [33] are other mechanisms by which sleep apnea may worsen outcomes in patients with heart failure [34].

The relationship between obstructive sleep apnea and cardiovascular events remains unclear. A systematic review and meta-analysis conducted by Loke, Yoon K., et al. suggested that obstructive sleep apnea may be an independent risk factor for cardiovascular, stroke, and overall mortality. Due to imprecision and inconsistencies in the data, the strength of potential association between obstructive sleep apnea and ischemic heart disease remains unclear [35]. A cohort study evaluating the relationship between obstructive sleep apnea-related variables and the risk of CV events revealed that several obstructive sleep apnea-related factors other than the apnea–hypopnea index were important predictors of a composite CV outcome [36]. Hence, the need for a randomized controlled trial is crucial.

3. Continuous Positive Airway Pressure (CPAP) Reduces Risk Factors for Myocardial Infarction, Atrial Fibrillation, and Heart Failure

Randomized controlled trials demonstrated that CPAP lowers blood pressure by 2–3 mm Hg [37]. Among patients with obstructive sleep apnea and resistant hypertension, CPAP treatment for 12 weeks decreased the 24 h mean and diastolic blood pressure and increased the nocturnal blood pressure [38]. CPAP also improved dyslipidemia (decrease in total cholesterol and LDL and increase in HDL). This may contribute to a potential reduction in cardiovascular and cerebrovascular events [39]. In patients with moderate-to-severe obstructive sleep apnea, compliant CPAP usage may improve insulin secretion and insulin resistance. The latter was associated with an improvement in leptin even after short-term CPAP therapy [40,41].

Both epidemiological studies and cohort studies suggest that CPAP reduces atrial fibrillation recurrence risk after cardioversion and ablation [42,43]. In 426 individuals undergoing pulmonary vein isolation, 62 patients with verified obstructive sleep apnea who used CPAP had a higher rate of atrial fibrillation-free survival than those who did not use CPAP (72% vs. 37%). The atrial fibrillation-free survival rate among CPAP users was comparable to that of individuals without obstructive sleep apnea. However, randomized controlled trials are still awaited [44]. Not surprisingly, given the association between obstructive sleep apnea and atrial fibrillation, there is also an association between obstructive sleep apnea and stroke [45,46]. CPAP may attenuate this risk [46], but this has not been studied prospectively.

4. CPAP Does Not Reduce Myocardial Infarction

The potential associations of CPAP to reduce composite cardiovascular events, all-cause, and cardiovascular death in patients with concomitant cardiovascular disease and obstructive sleep apnea rely on data from observational studies [47]. There is currently no level 1 evidence to suggest that CPAP can prevent future cardiovascular events in patients, including obstructive sleep apnea and coronary artery disease [48]. A randomized controlled trial that studied the impact of CPAP on cardiovascular outcomes showed no reduction in long-term adverse cardiovascular outcomes in the intention-to-treat population [49]. A clinical trial comparing usual care with usual care plus CPAP therapy found that the addition of CPAP did not prevent cardiovascular events in patients with moderate-to-severe obstructive sleep apnea and preexisting cardiovascular disease [50]. However, no significant beneficial effects of CPAP were shown in patients with obstructive sleep apnea in the trials evaluating CPAP therapy on major adverse cardiovascular events [49–51]. CPAP did not reduce the rate of complex cardiovascular events at a median follow-up of 3.7 years in the SAVE (Sleep Apnea Cardiovascular Endpoints) trial that randomized 2717 participants with cerebrovascular disease with moderate-to-severe obstructive sleep
apnea or [50]. Furthermore, no effect of CPAP therapy on major adverse cardiovascular events in obstructive sleep apnea with or without cardiovascular morbidities was shown in several meta-analyses of randomized trials [52–54]. However, the study populations of the included studies are diverse, from the general population to patients with severe coronary artery disease (such as myocardial infarction), thus precluding definitive conclusions.

5. Obesity

Obesity is a chronic multifactorial disease. Obesity is caused by inherited biological and ecological factors, diet, physical activity, and exercise choices. It is a medical problem that raises the threat of other diseases and health problems, such as high blood pressure, certain cancers, diabetes, and heart disease [55].

Various risk factors, involving obesity, age, sex, heritable and race/ethnicity factors, are well-known in the pathogenesis of sleep apnea. However, obesity has usually been known to be one of the most significant sleep apnea risk factors [56]. Some cross-sectional experiments have always found a connection between the obstructed sleep apnea risk and body mass index. The registered prevalence of sleep apnea ranges from 40% to 90% in individuals with a body mass index > 40 kg/m² (severe obesity) [57,58]. Significant sleep apnea appears in over 70% of sleep apnea patients with obesity and 40% of people with obesity [59].

Obesity is the only major obstructive sleep apnea risk factor that is variable. Weight decrease in the short term of one to two years steer to better metabolic control in patients with obstructive sleep apnea [60]. A prospective study of Wisconsin citizens indicated that a 10% weight loss anticipated a 26% reduction in the sleep apnea severity apnea–hypopnea index [58]. Moreover, another cohort research observed decreased apnea occurrence after weight loss [61]. Even though the mechanisms by which weight loss decrease obstructive sleep apnea symptoms are not entirely understood, factors such as decreased visceral fat deposition play an important role [62]. Airway structures and altered neurophysiologic control of respiration during weight loss are likely also important [63]. Moreover, obesity may control the chemoreflex function through neurohormonal mediators such as leptin, which reduces when sleep apnea patients lose weight [60]. Therefore, these potential methods likely work in concert to act as a vicious cycle in the body weight gain pathogenesis and obstructive sleep apnea (Figure 1).

Figure 1. Proposed mechanisms acting as a vicious cycle in the pathogenesis of obstructive sleep apnea and obesity and the potential influences of shared and non-shared genes.
Obesity complications such as hypertension, diabetes, insulin resistance, and obstructive sleep apnea increase the risk of cardiovascular diseases [64]. Obesity may also independently contribute to atherosclerosis and coronary artery disease [47,65–67]. To prevent or reverse obesity complications, more than 10% weight loss may be required [68], and the previously suggested 5% to 10% weight loss may not be sufficient [69,70].

In the case of obesity, evaluation before the intervention should include a comprehensive weight history, a complete individual and family medical history, blood studies, a physical examination looking for signs of the complications of obesity, and a behavioral history [71]. This approach may identify primary causes for weight gain, such as an endocrine disorder, medications linked with weight gain, an underlying eating disorder, or differences in the condition that have led to decreased activity and an energy imbalance. Body mass index and waist circumference should also be included in this initial evaluation. Blood tests should consist of a measurement of thyroid function, insulin sensitivity, liver function, and lipid profile [71]. Once a diagnosis of obesity has been made, the management of obesity should be stratified. Treatment approaches for obesity include nutritional therapy, pharmacotherapy, and surgical therapies, as shown in Table 1 [72].

<table>
<thead>
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<th>Table 1. Treatment options of obesity.</th>
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<tr>
<td><strong>Lifestyle therapy</strong></td>
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<td>Nutritional therapy</td>
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<td>Exercise therapy</td>
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<td>Cognitive behavioral therapy</td>
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<td><strong>Pharmacotherapy</strong></td>
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<td>FDA-approved medication</td>
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<td>Orlistat</td>
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<td>Phentermine/topiramate (Qsymia)</td>
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<td>Naltrexone/bupropion (Contrave)</td>
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<td>Liraglutide (Saxenda)</td>
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<td>Setmelanotide (IMCIVREE)</td>
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<td>Phentermine</td>
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<td><strong>Bariatric surgery</strong></td>
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<td>Sleeve gastrectomy</td>
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<td>Gastric bypass</td>
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5.1. Weight Loss as a Treatment to Reduce Cardiovascular Events

Currently, no randomized control trials have shown that intentional weight loss reduces the cardiovascular death risk [73–75]. Non-randomized long-term follow-up data from a prospective cohort Swedish Obese Subjects study showed that bariatric surgery resulted in reduced cardiovascular mortality and occurrence of first-time (fatal and nonfatal) cardiovascular events [76]. The Look AHEAD clinical trial was designed to assess the long-term effects of nutritional therapy and exercise therapy delivered over ten years in patients with obesity and type 2 diabetes. The primary outcome was time to the incidence of a major cardiovascular disease event. The study revealed that weight loss produced improvements in HbA1c, systolic blood pressure, HDL triglycerides, and cholesterol at years 1 and 4 (all \( p \leq 0.02 \)) [77], but no changes in cardiovascular events over ten years [77]. However, the subgroup of patients who lost more than 10% of their weight did have a reduction in mortality [78]. Another two-year follow-up cohort study also demonstrated that 5–10% weight loss improves risk factors but not cardiovascular events [79].

5.2. Weight Loss as a Treatment for Obstructive Sleep Apnea

Although weight loss can facilitate obstructive sleep apnea treatment, it can rarely cure it and still requires a combination with continuous positive airway pressure [80]. A systematic review and meta-analysis conducted to assess the effect of lifestyle therapy on the oxygen desaturation index, apnea–hypopnea index, and excessive daytime sleepiness among adults revealed significant reductions in all these components after weight loss. However, most patients still had diagnosable obstructive sleep apnea [81]. The SCALE sleep apnea study compared the effects of 3.0 mg liraglutide to placebo on obstructive sleep apnea [82].
apnea severity and body weight loss in patients with obstructive sleep apnea and obesity. This study showed that 3.0 mg liraglutide produced greater reductions in ischemic heart disease, body weight, systolic blood pressure, and HbA\textsubscript{1c} in patients with obesity with moderate/severe obstructive sleep apnea, but most patients still had diagnosable obstructive sleep apnea at the end of the study [82]. A prospective multicenter study investigated the effects of a laparoscopic Roux–en–Y gastric bypass. One year after surgery, the prevalence of obstructive sleep apnea decreased, but more than half of those with obstructive sleep apnea at baseline still had it after surgery. Obstructive sleep apnea was cured in 45% and improved in another 33% of the patients, but moderate or severe obstructive sleep apnea persisted in 20% of the patients after the operation [83]. A recent prospective cohort study showed that patients with moderate-to-severe obstructive sleep apnea may lose less weight with bariatric surgery than those with milder disease [84]. A systematic review included 19 surgical (n = 525), and 20 nonsurgical (n = 825) studies assessing the body mass index and the apnea–hypopnea index before and after intervention [85]. The results showed that bariatric surgery and non-surgical weight loss both had beneficial effects on obstructive sleep apnea through body mass index and apnea–hypopnea index reduction. However, bariatric surgery may result in a significantly greater improvement in the body mass index and the apnea–hypopnea index than nonsurgical alternatives [85]. The exact relationship between bariatric surgery and nonsurgical weight loss interventions in OSA resolution remains a challenge due to the need in randomized controlled trials [85].

5.3. Weight Loss as a Treatment to Reduce Cardiovascular Events in Patients with Obstructive Sleep Apnea

Cardiovascular events are the primary cause of mortality in patients with obstructive sleep apnea and obesity [86]. CPAP combined with weight loss may improve obstructive sleep apnea pathophysiology and the apnea–hypopnea index and reduce cardiovascular risk [87,88]. A randomized 24-week trial on 181 patients with moderate to severe obstructive sleep apnea and obesity comparing the effects of CPAP, weight loss, or combined CPAP and weight loss was conducted. Combining CPAP and weight loss did not reduce CRP levels more than either intervention alone. Weight loss provided an incremental reduction in serum triglycerides and the insulin resistance level when combined with CPAP. Weight loss and CPAP did incrementally reduce blood pressure compared with either intervention alone [89]. Another randomized control trial of 42 patients with obesity, severe obstructive sleep apnea (apnea–hypopnea index > 30 events/h) and treated with CPAP for a minimum of 6 months before the study allocated patients to an intensive weight loss program or standard lifestyle recommendations over 12 months. The effect of weight loss was assessed on obstructive sleep apnea severity and metabolic variables. The intensive weight loss program effectively reduces weight and obstructive sleep apnea severity while also improving lipid profiles, glycemic control, and inflammatory markers [90]. The authors concluded that weight loss intervention is an effective strategy to improve cardiovascular risk in patients with obesity and obstructive sleep apnea [66], but whether this will reduce cardiovascular events remains determined. Table 2 summarizes the available data assessing the effect of weight loss as a treatment for cardiovascular events risk reduction, obstructive sleep apnea, and to reduce cardiovascular events in patients with obstructive sleep apnea. This potential relationship is plausible because of prevailing pathophysiological mechanisms, but definitive evidence is still lacking. There is evidence that treatment with CPAP decreases blood pressure, but the impact of weight loss in patients with resistant hypertension is more profound than in those with acute obstructive sleep apnea. Current data supporting the impact of CPAP on cardiovascular events come from nonrandomized studies, and higher-quality evidence is needed to change clinical practice. We also require using the concept of individualized medicine for this matter and focusing on genetic variations and why a few patients with OSA develop cardiovascular effects while others do not. Such research will assist in reporting on the clinical trials that are required to be performed. Weight loss in patients with obstructive sleep apnea produced improvements in HbA\textsubscript{1c}. 
systolic blood pressure, HDL cholesterol, and triglycerides, but thus far, no changes in cardiovascular events have been shown. This will require cooperation among specialists in cardiology and sleep apnea.

Table 2. Summary of weight loss as a treatment for cardiovascular events risk reduction, obstructive sleep apnea and to reduce cardiovascular events in patients with obstructive sleep apnea.

<table>
<thead>
<tr>
<th>Weight loss as a treatment to reduce cardiovascular events</th>
<th>Nonrandomized long-term follow-up data from the prospective cohort Swedish Obese Subjects study showed that bariatric surgery resulted in reduced cardiovascular mortality and occurrence of first-time (fatal and nonfatal) cardiovascular events. The Look AHEAD clinical trial showed no changes in cardiovascular events over ten years.</th>
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6. Conclusions

Despite the association between cardiovascular disease and obstructive sleep apnea, randomized trials have failed to demonstrate that obstructive sleep apnea treatment improves the outcomes of cardiovascular events, even in patients with established cardiovascular disease [50]. The combination of weight loss with continuous positive airway pressure (CPAP) appears to be more helpful than either treatment in isolation. Large well-controlled trials in patients with obstructive sleep apnea to evaluate the impact of different weight reduction programs on cardiovascular disease are still required.

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