Preventive Nephrology: The Role of Obesity in Different Stages of Chronic Kidney Disease

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Abstract
Background: Obesity is increasing worldwide and has become a nontraditional risk factor in chronic kidney disease (CKD). Summary: Obesity-related nephropathy may aggravate renal complications of the metabolic syndrome and progress to advanced CKD stages, while obesity in early stages of CKD is clearly related to the development of kidney disease. A high body mass index (BMI) in advanced CKD stages and dialysis is an advantage for survival (so called “obesity paradox”). A high lean body to fat mass index indicates a beneficial state of body composition. In contrast, loss of muscle mass with increasing fat mass causes “sarcopenia obesity,” which is related to unfavorable outcomes in renal replacement therapy. Obesity (BMI > 30–35) in renal transplant recipients is associated with a higher risk of complications such as delayed graft function, increased rates of rejection, and graft loss. While conservative management of morbid obesity is failing in most cases, bariatric surgery seems to be an option in some cases to improve renal complications in the early stage of CKD or in transplant candidates. Key Message: In conclusion, obesity is increasingly prevalent among CKD patients. Adequate management with respect to the specific role of obesity in different stages of CKD should be integrated in routine renal care.

Introduction

The rising obesity pandemic has received major attention in general medicine and nephrology in the last few years [1, 2]. Globally, a high body mass index (BMI) accounted for 4.0 million deaths and more than two thirds of deaths related to a high BMI were due to cardiovascular disease [1]. The prevalence of obesity among children and adults has doubled since 1980 and has shown a continuous increase in many countries including the United States, Africa, and Asia, with highest numbers of obese children in China and India [1].

In view of chronic kidney disease (CKD) as a global public health concern, actions were taken to reduce major traditional and nontraditional risk factors, like obesity, as key determinants of poor health outcomes [3]. CKD is an
Important risk amplifier within diabetes, hypertension, and cardiovascular disease, conditions which are closely linked to obesity. The hazard ratio (HR) for end-stage kidney disease in obese adolescents is estimated to be between 7 in nondiabetic and almost 20 in diabetic people [3]. Thus, reduction of lifestyle-related risks is a cornerstone of mitigating the public health impact of diabetes, hypertension, and obesity. Primary intervention of obesity includes education, lifestyle, diet, exercise, weight management, and stress reduction.

Because of the limited effects of conservative and medical treatment of morbid obesity [4, 5] bariatric surgery has raised increasing interests in the renal community [6, 7]. Long-term results in patients after bariatric intervention are promising in ameliorating the prevalence of CKD, diabetes, and hypertension [8–11]. Kidney-related complications of bariatric surgery itself as acute kidney insufficiency or stone formation in specific procedures (Roux-en-Y gastric bypass and biliopancreatic diversion with duodenal switch), which affect oxalate and citrate secretion, may be considered [6].

The aim of this paper is to give a brief overview of the ambiguous role of obesity (so-called “obesity paradox”) in CKD, which is different in early stages with the origin of a specific obese nephropathy, in advanced CKD and in dialysis, and in the course of kidney transplantation (Table 1).

### Obesity in the Early Stage of CKD

Obesity (BMI >30) can cause or worsen CKD. Genetic (mono-, polygenetic) and epigenetic aspects contribute to the development of obesity [2, 12, 13] but the exact risk estimates of these aspects for the development of kidney disease remain unclear. Metabolic effects of different ad-

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**Table 1. Effect of overweight (BMI >30) in different stages of CKD (for details, see text)**

| Condition | CKD 1–4 | CKD 5 and CKD 5D | Kidney transplantation |
|-----------|---------|-----------------|-----------------------|
| Renal function | Origin of specific obesity-associated nephropathy | ? Modifies decline of renal residual function | ? Impaired graft function |
| Risk modification | Depending on metabolic aspects (diabetes control, hypertension, lipid profile) | Improved survival independent of co-morbidity | Increased risk of surgical complications |
| | Endocrine status (fertility, adrenal function, hypothyroidism, pituitary functioning) | ? Benefits may be different in the age groups and within dialysis modalities | Decreased graft survival |
| | Genetic factors | | ? Increased mortality risks due to metabolic complications |
| Preventive measurement | Diet modification | Adequate dialysis | Aggressive treatment of comorbidity risks (glucose control, lipids, smoking, hypertension) |
| | Exercise | Sufficient nutritional intake (calories, protein) | Diet modification |
| | Behavior changes | Regular assessment of body weight and conditions which may cause weight loss | Exercise/counseling |
| | Psychological counseling | | |
| | Social support | | |
| Therapeutic aspects | Medication | Standard nutritional recommendation according to treatment modality | ? Adoption of immunosuppressive regimen (steroids) |
| | Bariatric surgery | Nutritional intervention to avoid weight loss (<25 BMI) | Bariatric surgery |
| Further aspects | Definition of benefits and risks of complications by bariatric surgical intervention | Normalization of body weight in transplant candidates (BMI <30–35) | Special donor care to reduce overweight and prevent OAN |

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?, denotes limited evidence; OAN, obesity-associated nephropathy.
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Ipokines (leptin, resistin, visfatin) and downregulation of adiponectin may contribute to hemodynamic and structural renal lesions via insulin resistance, increased insulin blood level, RAAS activation, oxidative stress, and micro-inflammation [2, 7, 14]. Risks of obesity-associated CKD seems to be different in metabolically healthy subjects and in those with metabolic syndrome [2]. A recent 8-year study in Japanese people (BMI > 25) revealed a crude incidence proportion of CKD in metabolically healthy obese phenotype (i.e., high levels of insulin sensitivity, low prevalence of hypertension, favorable fasting glucose, lipid, and inflammation profile) of 6.7 versus 10.9% (odds ratio 1.44 [95% CI 0.8–2.77] vs. 2.80 [95% CI 1.45–5.35]) in metabolically abnormal subjects [15].

Morphology of obesity-associated nephropathy (OAN) has been derived from autopsy and biopsy studies. Increased kidney weight and hypertrophy of individual nephrons are common findings in OAN. Consistently, a 3-fold increased glomerular size compared to non-obese subjects is found in several studies while glomerular density in the cortex is reduced. The number of glomerular capillaries appears to be increased suggesting de novo formation of microvessels (review by Tsuboi et al. [16]). Onset of albuminuria is associated with focal segmental glomerulosclerosis (FSGS). The extent of FSGS seems to be different in different stages of obesity [16]. Finally, FSGS is accompanied by progressive tubule interstitial fibrosis (Fig. 1a, b) in accordance with loss of kidney function. Conditions of reduced nephron mass as low birth weight, congenital anomalies of the kidney and urinary tract, or nephrectomy contribute significantly to the progression of CKD in obese individuals.

Indication for renal biopsy may result from rapid progressive loss of kidney function, active urinary sediment findings, and extrarenal features suggesting non-OAN as found in single cases with glomerular or systemic disease. While regular biopsy technique may fail in obese individuals, transjugular access is an option [17].

In the general population, obesity (BMI > 30) increases the risk of CKD, onset of albuminuria in CKD stages 1–2 as well as progression to CKD stages 3 and higher in the general population [18]. The course of CKD may be modulated by different causes of reduced kidney mass, genetic aspects, and metabolic phenotype. Medical RAAS blockade to influence endothelial dysfunction and associated tubulointerstitial fibrosis may be an option to prevent obese nephropathy [14]. Interestingly, decrease of proteinuria has been observed in a single-center study in patients with biopsy-proven obese nephropathy after losing weight by a medical supervised standard weight loss program [19].

**Obesity in Advanced CKD and Dialysis: A Paradox**

The paradoxical finding that even morbid obesity predicts better outcome in dialysis is well-known since 1999 when first reports from large US cohort and single-center studies were published [20, 21].

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**Fig. 1.** Histological features of obesity-related nephropathy (percutaneous renal biopsy of a 48-year-old woman, courtesy Prof. Kerstin Amann, Erlangen, Germany). a Prominent global and focal segmental glomerulosclerosis (FSGS). PAS. ×10. b FSGS and severe tubulointerstitial fibrosis. PAS. ×20.
ies have shown that any gain in body weight is associated with better survival in advanced CKD and dialysis (review by Kalantar-Zadeh et al. 22) (Fig. 2). Analysis of death rates in advanced CKD stages suggest that BMI >40 was not associated with higher risks of death after excluding diabetes and hypertension from the risk model [22]. While larger body size by fluid retention is associated with poorer outcomes gaining body weight with increased fat and muscle mass (fat-free lean body mass) is favorable [2, 22]. Anthropometry such as mid-arm muscle circumference can be used to estimate lean body mass. Dual energy X-ray absorptiometry is routinely used for the assessment of fat mass in dialysis patients. Findings from different studies suggest that survival advantages from gaining fat mass is lower than from gaining lean body mass [22]. Using waist circumference as surrogate of intra-abdominal or visceral fat shows that each 10-cm increase in waist circumference was associated with higher all-cause and cardiovascular death [23]. The superiority of survival advantages of (gaining) muscle compared to fat mass was confirmed by calculating the relation of lean to fat mass index which results in more favorable outcomes in patients with advanced CKD stages with a high lean/fat tissue index [24].

New insights of the meaning of muscle mass compared to fat mass in body weight gain comes from the “sarcopenia obesity” concept [25]. Increasing BMI in dialysis patients does not exclude concurrent muscle wasting. Sarcopenia and its individual criteria (hand grip strength, slow gait speed) are associated with mortality in hemodialysis patients [26]. Otherwise, intervention by structured exercise programs, which increase muscle mass in these patients, is proven to ameliorate functional parameters, body composition, quality of life, and survival [27–29].

The impact of BMI variations may vary by different dialysis populations and age. Most studies are done in hemodialysis patients. While increase in muscle weight in dialysis seems to be a defined advantage in both modalities, peritoneal dialysis and hemodialysis, the role of gaining body weight by fat mass in peritoneal dialysis patients is less well defined [21]. Furthermore, intentional weight loss may differ from unintentional wasting in these populations. In older age groups, hemodialysis patients with stable weight had a longer survival than elderly who lose or gain weight [30]. Death rates of obese younger dialysis patients (<65 years) are 1.7-fold higher than for the older group (>65 years) after adjustment for comorbidity and treatment modality [31].

Future studies are needed that advance our understanding of the existence of the obesity paradox. Results from these studies may create more efforts to define a more elaborated therapeutic management of obese patients along with functional improvement, quality of life, and survival.

**Fig. 2.** The obesity paradox: reverse association of BMI and survival in CKD patients as compared to the general population (reprint from Kalantar-Zadeh et al. [22]).

**The Role of Obesity in Kidney Transplantation**

The impact of obesity (BMI >30) in kidney transplant recipients has been recognized for more than 30 years [2]. Cumulate evidence from numerous observational studies
and registries including more than 138,000 transplant patients revealed (a) a higher proportion of delayed graft function (odds ratio 1.68, 95% CI 1.39–2.03), (b) an increased risk of death-censored graft loss (HR 1.06, 95% CI 1.01–1.12), but (c) no significant long-term survival risk (HR 1.24, 95% CI 0.9–1.70) [32]. A second meta-analysis including more than 240,000 patients shows similar results for delayed graft function and a slightly increased risk of patient death (HR 1.19, 95% CI 1.10–1.31), increased risks for the presence of biopsy-proven acute rejection (HR 1.51, 95% CI 1.24–1.78), and allograft loss (HR 1.54, 95% CI 1.38–1.68) [33]. Thus, recently published calculators for kidney graft survival include BMI in the risk model [34].

As a consequence, loss of body weight is recommended in renal transplant candidates [34, 35]. Obviously, many patients are unable to achieve significant weight loss from conventional measurements. Bariatric surgery might become an option for this group of patients (summary in Hossain et al. [35]). While gastric banding is no longer the method of choice to reduce body weight and metabolic complications in the long term, newer techniques like sleeve gastrectomy and intestinal bypass procedures are increasingly performed [36, 37]. In view of the limited data, it seems to be premature to advocate routine bariatric surgery for obese kidney recipients. Nevertheless, supervised loss of body weight to achieve a BMI < 30–35 before transplantation is recommended to improve kidney graft survival.

The role of donor to recipient size mismatching is poorly defined today. A recently published single-center study estimates the impact of donor body weight to recipient body weight. Donor to recipient mismatching in the highest group of donor body weight showed no increased risk for 1 and 5 years of allograft survival [38].

Summary and Conclusion

Obesity has emerged as a new risk factor in the development of CKD. Obesity can cause a specific renal nephropathy or contribute to renal complications in metabolic syndrome. Weight reduction by conservative management or metabolic (bariatric) surgery ameliorates renal complications. In advanced CKD as well as in end-stage kidney disease, a high BMI is associated with better survival. Losing weight in the stage of dialysis treatment must be avoided. Maintenance of body weight by adequate nutrition and physical activity is mandatory to avoid weight loss or development of sarcopenia. The phenomenon of “sarcopenia obesity” must be recognized by assessing body composition together with functional testing (handgrip strength, gait speed). Structured exercise programs – mainly at the time of dialysis treatment – are proven to increase muscle mass and functional outcomes. Because high BMI (> 30–35) is associated with higher risks of transplant complications, reduction of body weight should be achieved in transplant candidates. Results of bariatric surgery in this group are limited but promising. Further research needs to focus on strategies to reduce the burden of obesity in early CKD stages, maintaining adequate body weight and body composition in advanced stages of CKD and dialysis, and the development of strategies to reduce transplantation risks in obese recipients. Thus, nephrologists have become players in a multidisciplinary team approach to fight against obesity as a pandemic health problem. Distinct knowledge of the different role of obesity in the trajectory of CKD is crucial for the adequate management of obese patients.

Disclosure Statement

The author has no conflicts of interest to disclose.

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