# UC Irvine UC Irvine Previously Published Works

**Title** Obesity Paradox in Patients on Maintenance Dialysis

**Permalink** https://escholarship.org/uc/item/30g9c2mf

Authors Kalantar-Zadeh, Kamyar Kopple, Joel D

Publication Date

2006

DOI

10.1159/000095319

### **Copyright Information**

This work is made available under the terms of a Creative Commons Attribution License, available at <u>https://creativecommons.org/licenses/by/4.0/</u>

Peer reviewed

## **Obesity Paradox in Patients on Maintenance Dialysis**

#### Kamyar Kalantar-Zadeh, Joel D. Kopple

Division of Nephrology and Hypertension, Los Angeles Biomedical Institute at Harbor-UCLA Medical Center, Torrance, Calif.; David Geffen School of Medicine at UCLA and UCLA School of Public Health, Los Angeles, Calif., USA

#### Abstract

Overweight (body mass index  $[BMI] = 25-30 \text{ kg/m}^2$ ) and obesity  $(BMI > 30 \text{ kg/m}^2)$  have become mass phenomena with a pronounced upward trend in prevalence in most countries throughout the world and are associated with increased cardiovascular risk and poor survival. In patients with chronic kidney disease (CKD) undergoing maintenance hemodialysis an 'obesity paradox' has been consistently reported, i.e., a high BMI is incrementally associated with better survival. While this 'reverse epidemiology' of obesity is relatively consistent in maintenance hemodialysis patients, studies in peritoneal dialysis patients have yielded mixed results. A similar obesity paradox has been described in patients with chronic heart failure as well as in 20 million members of other distinct medically 'at risk' populations in the USA. Possible causes of the reverse epidemiology of obesity include: (1) time-discrepancies between the competing risks for the adverse events that are associated with overnutrition and undernutrition; (2) sequestration of uremic toxins in adipose tissue; (3) selection of a gene pool favorable to longer survival in dialysis patients during the course of CKD progression, which eliminates over 95% of the CKD population before they commence maintenance dialysis therapy; (4) a more stable hemodynamic status; (5) alterations in circulating cytokines; (6) unique neurohormonal constellations; (7) endotoxin–lipoprotein interactions; and (8) reverse causation. Examining the causes and consequences of the obesity paradox in dialysis patients can improve our understanding of similar paradoxes observed both for other conventional risk factors in chronic dialysis patients, such as blood pressure and serum cholesterol, and in other populations, such as patients with heart failure, cancer or AIDS or geriatric populations.

Copyright © 2006 S. Karger AG, Basel

*Founding Source*: Supported by a Young Investigator Award from the National Kidney Foundation, a research grant from DaVita, Inc., and the National Institute of Diabetes, Digestive and Kidney Disease grant # DK61162 (for KKZ).

#### Introduction

Individuals with chronic kidney disease (CKD) stage 5 who undergo maintenance dialysis treatment have a high mortality rate, currently 20% per year in the USA and 10–15% in Europe [1]. This high mortality has not changed substantially in recent years despite many advances in dialysis techniques and patient care [2]. Cardiovascular disease is the main cause of death in dialysis patients [3]. The currently estimated chronic dialysis population of 350,000 patients in the USA grows constantly and fast, and is projected to reach over one-half million by 2010 and over one million by 2020 [2].

It was once believed that the traditional cardiovascular risk factors and/or conditions related to dialysis treatment and technique are the main causes of poor clinical outcome; however, recent randomized controlled trials including the 4D Trial [4] and the HEMO and ADEMEX studies [5, 6] failed to show an improvement of mortality by lowering serum cholesterol levels or by increasing dialysis dose, respectively. Hence, it is not unlikely that conditions other than the traditional risk factors are related to the enormous cardiovascular epidemic and high death rate in this population.

An increasing number of epidemiologic studies, based on analyses of large samples of dialysis patients and national databases, have indicated paradoxical and inverse associations between classical cardiovascular risk factors and mortality in dialysis patients [7, 8]. Indeed, a worse survival among dialysis patients has been observed with a low, rather than a high, body mass index (BMI) [9] or weight-for-height [10], blood pressure [11], and serum concentrations of cholesterol [12], homocysteine [13] and creatinine [14]. Even more ironically are findings indicating that high values of these risk factors are paradoxically protective and associated with improved survival. This phenomenon has been referred to as 'reverse epidemiology' [7] or 'altered risk factor pattern' [8]. These epidemiologic findings have contributed to the growing confusion and have left physicians with the ongoing dilemma as to whether or not to treat obesity, hypercholesterolemia, hypertension, or hyperhomocysteinemia in chronic dialysis patients [8]. Among the above-mentioned cardiovascular risk factors that are inversely associated with mortality, the so-called 'obesity paradox' has been the most consistent and most extensively studied [15, 16]. In this manuscript, the paradoxical predictability of mortality of measures of body size and several hypotheses that have been advanced to explain these paradoxes are reviewed critically.

#### **Obesity in the General Population**

In recent years, overweight  $(BMI = 25-30 \text{ kg/m}^2)$  and obesity  $(BMI > 30 \text{ kg/m}^2)$  have become mass phenomena with a pronounced upward trend in

Kalantar-Zadeh/Kopple

prevalence in virtually all developed and developing countries [17, 18]. The prevalence of obesity has reached epidemic proportions in the USA: It ranged between 13 and 15% between 1960 and 1980 [19-21], but doubled to 23 and 31% during 1988–1994 and 1999–2000, respectively [17, 22]. Obesity is a strong risk factor for the development of diabetes mellitus, atherosclerotic cardiovascular disease, cancer and even CKD [23-26]. However, despite detrimental effects of being overweight, 'obese nations live paradoxically longer than ever' [18]. The increasing prevalence of obesity may be understood in the light of evolution, because energy metabolism is asymmetric with energy accumulation being the necessary condition of survival during 'hard times' [18]. According to this theory, this genetic characteristic, the so-called thrifty gene(s), was necessary for survival of humanity, because during the course of history there was never a long period of uninterrupted food abundance, whereas famines and other hardships occurred frequently. Therefore, fat accumulation, when food was available, meant survival at times of hardship. In contrast, the potential detrimental effects of overnutrition and overweight generally only became manifest at an older age to which most people did not live and therefore were not very relevant to survival [18]. The foregoing model may explain why in chronic disease states obesity confers survival advantages [27].

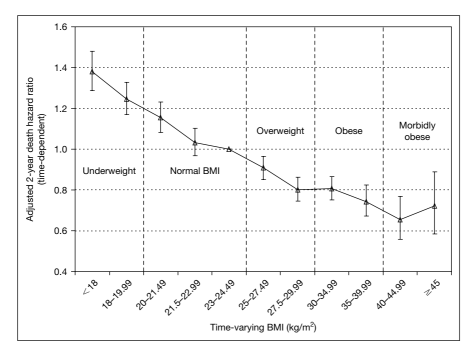
#### **Obesity and Survival in Hemodialysis Patients**

Obesity has recently been reported to be a risk factor of the CKD progression to stage 5 [26], although this epidemiologic observation may be severely confounded by a strong survival bias [28], especially because many CKD patients die before they reach stage 5 [29]. Maintenance hemodialysis (MHD) patients appear to have a lower BMI as compared with age- and sex-matched controls from the general population [30–32].

Most epidemiologic studies have shown an inverse association between larger body size and lower death risk in MHD patients, independent of other markers of nutritional status [15]. The Diaphane Collaborative Study [33] appears to be the first to report the association between low BMI and high death rate in a cohort (1972–1978) of 1,453 mostly nondiabetic French MHD patients, which was confirmed 15 years later by Leavey et al. [34] (3,607 MHD patients) using the United States Renal Data System (USRDS) database. Fleischmann et al. [35] identified for the first time the so-called 'obesity paradox', i.e., a significantly higher survival rate in overweight and obese MHD patients (BMI  $\geq$  27.5 kg/m<sup>2</sup>) compared to those with a normal weight (BMI = 20–27.5 kg/m<sup>2</sup>) and underweight (BMI < 20 kg/m<sup>2</sup>) [35]. Wolf et al. [36] (9,165 MHD patients) and Port et al. [37] (45,967 incident MHD patients) reported similar paradoxical associations in the USRDS database. The *Dialysis Outcomes and Practice Patterns Study* (DOPPS) [38] (9,714 MHD patients in the USA and Western Europe from 1996 to 2000) confirmed an inverse BMI–mortality relationship in MHD subpopulations defined by continent, race, gender, tertiles of severity of illness and comorbid conditions. Glanton et al. [39] (151,027 incident dialysis patients from the USRDS) found that the obesity paradox was not uniform across different gender and race/ethnicity subgroups and was stronger in African-Americans. Johansen et al. [40] (418,055 maintenance dialysis patients in the USRDS data) found that even morbid obesity was associated with increased survival, except for in Asian-Americans. High BMI and Benn's index were also associated with a reduced risk of hospitalization. Survival rates based on estimates of adiposity and fat mass yielded similar results, and adjustments of body weights for differences in lean body mass did not substantially alter the paradoxical associations [40].

Some investigators have studied body size surrogates other than BMI. Kopple et al. [10] examined weight adjusted for height percentiles in 12,965 MHD patients from the Fresenius database and found that those patients with greater weight-for-height had lower mortality rates. Lowrie et al. [41] (43,334 MHD patients from the Fresenius database) examined body surface area and weight divided by height (wt/ht) in addition to the BMI. The log of risk decreased in rough linear fashion for weight, weight-for-height, and body surface area [41]. Beddhu et al. [42] (70,028 incident MHD patients in the USRDS database) used 24-hour urine creatinine excretion as a indicator of muscle mass and concluded that higher muscle mass was a stronger predictor of survival than was higher total body weight in heavy MHD patients. However, their data showed that obesity was associated with better survival within each fat/muscle category [42]. The inherent associations of urine creatinine with renal function, muscle mass and meat intake may restrict the generalizability of the foregoing conclusions [43].

Kalantar-Zadeh et al. [9] recently examined the effects of both absolute magnitude of BMI (using the time-dependent Cox model) and changes in BMI over time on all-cause and cardiovascular mortality in a 2-year nonconcurrent cohort of 54,535 MHD patients in the national database of the second largest dialysis care provider in the USA (DaVita, Inc.). They found that obesity, including morbid obesity (BMI >  $35 \text{ kg/m}^2$ ), was associated with survival advantages in virtually all subgroups of age, gender, race, dialysis vintage, serum albumin, and Kt/V (fig. 1). Moreover, they showed for the first time that independently of almost any BMI level, weight loss is associated with increased mortality, whereas weight gain confers survival advantages [9] (fig. 2). Finally, in another recent study, Kalantar-Zadeh et al. [44] measured total body fat directly in 535 MHD patients over 3.5 years and found that not only a lower

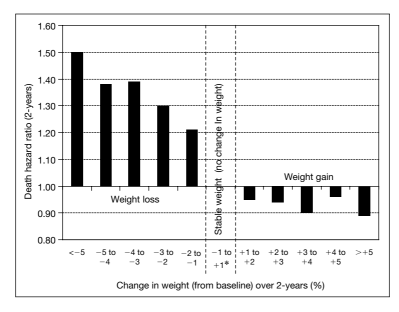


*Fig. 1.* Adjusted death hazard ratio in 54,535 MHD patients (7/2001–6/2003); recreated bases on data from reference [9].

total body fat was associated with higher mortality, but loss of body fat over time was associated with increased death risk.

#### **Obesity and Survival in Peritoneal Dialysis Patients**

Some [45–50], but not all [51–53], studies in chronic peritoneal dialysis (CPD) patients have reported inverse weight-mortality relationships [15]. In the CANUSA study, a 1% difference in the percent lean body mass was associated with a 3% change in the relative risk of death [45, 46]. McCusker et al. [47], Chung et al. [49], and Johnson et al. [48] found a significantly decreased survival rate in CPD patients with a lower body weight. The largest epidemiologic study of body weight and survival in CPD patients included nearly 46,000 CPD patients in 1990s [50], and showed that overweight and obese CPD patients had longer survival than those with normal BMI. These findings could not be adequately explained by lower rates of renal transplantation or lower technique survival rates. Abbott et al. [52] compared 1,675 MHD and 1,662 CPD patients and found that



*Fig. 2.* Relative risk of death for changes in weight over time; recreated bases on data from reference [9].

5-year survival based on BMI cutoff of  $30 \text{ kg/m}^2$  in CPD patients was not different than in MHD.

Several studies in CPD patients either have not found any survival advantage for obesity or have indicated a higher risk of death in obese CPD patients [53]. McDonald et al. [51] examined 9,679 CPD patients in Australia or New Zealand over an 11-year interval and found that obesity was independently associated with death and technique failure except among patients of New Zealand Maori/Pacific Islander origin. Stack et al. [54] examined CPD–MHD differences in a cohort of 134,728 new ESRD patients from the USRDS and concluded that the selection of HD over PD was associated with a survival advantage in patients with large body habits [54]. Beddhu et al. [42] hypothesized that the survival advantages of obesity is due to muscle mass both in MHD and CPD patients [55] using urinary creatinine as a indicator of muscle mass [56].

#### Other Populations with a Reverse Epidemiology

Patients with chronic heart failure (CHF) [57], geriatric populations [58], and patients with malignancy [59], AIDS [60], chronic obstructive pulmonary disease [61], or rheumatoid arthritis [62] also exhibit a risk factor reversal.

Kalantar-Zadeh/Kopple

*Table 1.* Potential mechanisms that may result in the observed paradoxical associations between obesity and better survival in dialysis patients

Kidney Disease Wasting (Malnutrition–inflammation complex syndrome) Time discrepancy between competitive risk factors: overnutrition vs. undernutrition Unusual genetic constellation due to survival selection during CKD progression Sequestration/storage of uremic toxins in fat tissue Anti-inflammatory cytokines related to body mass, including adiponectins Tumor necrosis factor alpha receptors Endotoxin-lipoprotein hypothesis Stability of hemodynamic status in obese patients Neurohormonal alterations in obesity Alteration of conventional risk factors in uremic milieu ('beyond Framingham') Reverse causation Survival bias

There appear to be at least 20 million Americans who may have a reverse epidemiology pattern [63]. There are striking similarities in the reported paradoxes between the patients with CHF, currently almost 5 million individuals in the USA, and MHD patients [64, 65]. Furthermore, millions of the fast-growing population of octogenarians and nonagenarians in the industrial countries appear to display a reverse epidemiology [63]. Studying the causal factors that engender the obesity paradox in chronic dialysis patients may confer better insight into understanding the pathophysiology and public health consequences of these phenomena in other populations.

### Pathophysiology of the Obesity Paradox

Several hypotheses have been advanced to explain the obesity paradox in dialysis patients (table 1) [8, 15, 16]. Obesity and weight gain may be associated with a more stable short-term hemodynamic constellation and improved hemodynamic tolerance to afterload-reducing agents, especially because overweight and obese patients with heart failure tend to have higher systemic blood pressure values [66]. Thus, obese patients might better tolerate removal of large volumes of fluid during the hemodialysis procedure with less likelihood of hypotension. Obesity may mitigate stress responses and the heightened sympathetic and renin–angiotensin activity; the latter are associated with a poor prognosis in heart failure and fluid overload states such as in dialysis patients [67]. Hence, better outcome is expected if angiotensin axis can be blocked [68]. Altered cytokine and neuroendocrine profiles of obese patients may also play

a role in conferring survival advantages to obese patients. Adipose tissue produces adiponectins [69], as well as soluble tumor necrosis factor alpha receptors which may neutralize the adverse biologic effects of tumor necrosis factor alpha [70]. It is also possible, although not yet proven, that the uremic milieu or volume overload modifies the cardiovascular constellations so that factors 'beyond Framingham paradigms' are more relevant for survival [71].

It has been postulated that higher concentrations of total cholesterol (lipoproteins) are beneficial for dialysis and CHF patients, since a richer pool of lipoproteins can actively bind to and remove circulating endotoxins; hence, the increase pool of these lipoproteins may attenuate the propensity of endotoxins which would otherwise cause inflammation and subsequent atherosclerosis if unbound [72]. This so-called 'endotoxin–lipoprotein' hypothesis was originally advanced to explain the hypercholesterolemia paradox in CHF patients [73].

It is also possible that uremic toxins are more effectively sequestered when abundant adipose tissue is present. It has been shown that weight loss and reduction in adipose tissue is associated with the imminent release of and significant increase in circulating lipophilic hexachlorobenzene and other chlorinated hydrocarbons [74]. Weight loss may also be associated with reduced skeletal muscle oxidative metabolism, leading to a mitigated anti-oxidant defense [75]. These findings may provide one explanation for why body fat loss has recently been found to be associated with increased death risk in dialysis patients [44].

It is, of course, possible that BMI is not a cause but a consequence of conditions that lead to poor outcome in dialysis patients or in similar populations with a paradoxical risk factor profile. 'Reverse causation' is a known possible source of bias in epidemiologic studies that examine associations without the direction of the causal pathway [76]. Comoribid states may lead to kidney disease wasting or cardiac cachexia and also to higher rate of mortality. However, even if the reverse causation is a cause of the reverse epidemiology, it does not explain why obesity including morbid obesity is associated with better outcome than the traditional normal or healthy weights in dialysis patients.

Of the currently estimated 20 million individuals with CKD in the USA [77], it is projected that over 90% will die before advancing to end-stage renal disease [29]. Hence, only less than 5% of the large CKD pool will be the 'unlucky lucky' individuals to reach the dialysis facility chair [7]. This may lead to a significant 'survival selection' [64] resulting in genetic constellations in dialysis patients that may be significantly different than their early CKD predecessors [78]. According to this theory, those few CKD patients who reach ESRD may have either a more accelerated rate of progression of CKD or special genes that protected them against the fatal ravages of cardiovascular disease which is inherent to CKD. Whether this is called 'survival bias' or 'survival

selection' (similar to evolutionary natural selection), maintenance dialysis patients must be genetically or phenotypically dissimilar to their CKD predecessors who do not survive and may not have the survival characteristics and epidemiological features of their progenitors.

Survival advantages that exist in obese dialysis patients may, in the *short-term*, outweigh the harmful effects of these risk factors in causing cardiovascular disease and death in the *long-term* [7]. In other words, dialysis patients may not live long enough to die of the adverse effects of overnutrition, because they are more likely to die much faster of the consequences of undernutrition [64]. This so-called time-discrepancy between the two sets of competing risk factors, i.e., short-term killers (malnutrition–inflammation complex) vs. long-term killers (obesity and overnutrition), can explain why obesity treatment may be irrelevant or even harmful in many (but not all) dialysis patients if the issue at hand is the short-term survival. Currently 2/3 of all dialysis patients in the USA die within 5 years of commencing dialysis, a 5-year survival worse than many cancer patients [64]. Hence, treatment of malnutrition–inflammation complex, also known as kidney disease wasting, should be the target of efforts to improve survival in maintenance dialysis patients.

#### **Conclusions and Future Steps**

Studying the obesity paradox and other similar phenomena in dialysis patients leads to additional questions: Is the reverse epidemiology a true entity with clinical and public health implications in millions of patients with CKD, CHF, advanced age, malignancy, AIDS, etc., or is it a statistical fallacy that needs to be 'controlled away'? [79]. At which CKD stage does the reverse epidemiology start and in whom does it develop? Which groups of dialysis patients have a stronger, weaker or no obesity paradox? Can the so-called 'reversal of the reverse epidemiology' (or 'back-to normal) phenomenon upon successful renal transplantation of dialysis patients or with frequent (daily and/or nocturnal) dialysis treatment be confirmed? [64]. If this reversal of the altered risk factor relationships in these patients are real, what are the mechanisms for this phenomenon? What should be our therapeutic targets for body weight-forheight in our CKD and maintenance dialysis patients? Should we revise the current guidelines that recommend that obese dialysis patients on transplant waiting lists should lose weight as a prerequisite for renal transplantation? [40]. For that matter, what should be our therapeutic targets in maintenance dialysis patients for other clinical characteristics for which the usual risk factor relationships to mortality are altered, such as blood pressure or serum cholesterol or phosphorus, to mention a few? Is the evidence for these altered risk factor relationships sufficiently established to justify proposing to research granting agencies, the funding of randomized, prospective interventional trials to examine the appropriate therapeutic targets for BMI or some of these other clinical targets in chronic dialysis patients?

The field of altered risk factor relationships is in its infancy and appears to be evolving quickly. It is possible that, in the long run, overweight patients, if they survive sufficiently long, will suffer from more cardiovascular consequences. Until more information is available, it is prudent to avoid causal inferences for such observational data.

#### References

- 1 Goodkin DA, Mapes DL, Held PJ: The dialysis outcomes and practice patterns study (DOPPS): how can we improve the care of hemodialysis patients? Semin Dial 2001;14:157–159.
- 2 United States Renal Data System: Excerpts from the USRDS 2004 Annual Data Report. Am J Kid Dis 2005;45(suppl 1):S1–S280.
- 3 Foley RN, Parfrey PS, Sarnak MJ: Epidemiology of cardiovascular disease in chronic renal disease. J Am Soc Nephrol 1998;9:S16–S23.
- 4 Wanner C, Krane V, Marz W, Olschewski M, Mann JF, Ruf G, Ritz E: Atorvastatin in patients with type 2 diabetes mellitus undergoing hemodialysis. N Engl J Med 2005;353:238–248.
- 5 Eknoyan G, Beck GJ, Cheung AK, Daugirdas JT, Greene T, Kusek JW, Allon M, Bailey J, Delmez JA, Depner TA, Dwyer JT, Levey AS, Levin NW, Milford E, Ornt DB, Rocco MV, Schulman G, Schwab SJ, Teehan BP, Toto R: Effect of dialysis dose and membrane flux in maintenance hemodialysis. N Engl J Med 2002;347:2010–2019.
- 6 Paniagua R, Amato D, Vonesh E, Correa-Rotter R, Ramos A, Moran J, Mujais S: Effects of increased peritoneal clearances on mortality rates in peritoneal dialysis: ADEMEX, a prospective, randomized, controlled trial. J Am Soc Nephrol 2002;13:1307–1320.
- 7 Kalantar-Zadeh K, Block G, Humphreys MH, Kopple JD: Reverse epidemiology of cardiovascular risk factors in maintenance dialysis patients. Kidney Int 2003;63:793–808.
- 8 Kopple JD: The phenomenon of altered risk factor patterns or reverse epidemiology in persons with advanced chronic kidney failure. Am J Clin Nutr 2005;81:1257–1266.
- 9 Kalantar-Zadeh K, Kopple JD, Kilpatrick RD, McAllister CJ, Shinaberger CS, Gjertson DW, Greenland S: Association of morbid obesity and weight change over time with cardiovascular survival in hemodialysis population. Am J Kidney Dis 2005;46:489–500.
- 10 Kopple JD, Zhu X, Lew NL, Lowrie EG: Body weight-for-height relationships predict mortality in maintenance hemodialysis patients. Kidney Int 1999;56:1136–1148.
- 11 Kalantar-Zadeh K, Kilpatrick RD, McAllister CJ, Greenland S, Kopple JD: Reverse epidemiology of hypertension and cardiovascular death in the hemodialysis population: the 58th annual fall conference and scientific sessions. Hypertension 2005;45:811–817.
- 12 Nishizawa Y, Shoji T, Ishimura E, Inaba M, Morii H: Paradox of risk factors for cardiovascular mortality in uremia: is a higher cholesterol level better for atherosclerosis in uremia? Am J Kidney Dis 2001;38:S4–S7.
- 13 Kalantar-Zadeh K, Block G, Humphreys MH, McAllister CJ, Kopple JD: A low, rather than a high, total plasma homocysteine is an indicator of poor outcome in hemodialysis patients. J Am Soc Nephrol 2004;15:442–453.
- 14 Lowrie EG, Lew NL: Death risk in hemodialysis patients: the predictive value of commonly measured variables and an evaluation of death rate differences between facilities. Am J Kidney Dis 1990;15:458–482.
- 15 Kalantar-Zadeh K, Abbott KC, Salahudeen AK, Kilpatrick RD, Horwich TB: Survival advantages of obesity in dialysis patients. Am J Clin Nutr 2005;81:543–554.

Kalantar-Zadeh/Kopple

- 16 Kalantar-Zadeh K: Causes and consequences of the reverse epidemiology of body mass index in dialysis patients. J Ren Nutr 2005;15:142–147.
- 17 Kuczmarski RJ, Flegal KM, Campbell SM, Johnson CL: Increasing prevalence of overweight among US adults. The National Health and Nutrition Examination Surveys, 1960 to 1991. JAMA 1994;272:205–211.
- 18 Lev-Ran A: Human obesity: an evolutionary approach to understanding our bulging waistline. Diabetes Metab Res Rev 2001;17:347–362.
- 19 Birkner R: Plan and Initial Program of the Health Examination Survey. Vital Health Stat 1 1965;125:1–43.
- 20 Miller HW: Plan and operation of the health and nutrition examination survey. United States 1971–1973. Vital Health Stat 1 1973;1:1–46.
- 21 McDowell A, Engel A, Massey JT, Maurer K: Plan and operation of the Second National Health and Nutrition Examination Survey, 1976–1980. Vital Health Stat 1 1981;1:1–144.
- 22 Flegal KM, Carroll MD, Ogden CL, Johnson CL: Prevalence and trends in obesity among US adults, 1999–2000. JAMA 2002;288:1723–1727.
- 23 Byers T: Body weight and mortality. N Engl J Med 1995;333:723–724.
- 24 Manson JE, Willett WC, Stampfer MJ, Colditz GA, Hunter DJ, Hankinson SE, Hennekens CH, Speizer FE: Body weight and mortality among women. N Engl J Med 1995;333:677–685.
- 25 Lew EA, Garfinkel L: Variations in mortality by weight among 750,000 men and women. J Chronic Dis 1979;32:563–576.
- 26 Hsu C-y, McCulloch CE, Iribarren C, Darbinian J, Go AS: Body mass index and risk for end-stage renal disease. Ann Intern Med 2006;144:21–28.
- 27 Kalantar-Zadeh K, Abbott KC, Salahudeen AK, Kilpatrick RD, Horwich TB: Survival advantages of obesity in dialysis patients. Am J Clin Nutr 2005;81:543–554.
- 28 Kalantar-Zadeh K, Kopple JD: Body mass index and risk for end-stage renal disease. Ann Intern Med, 2006;144:21–28.
- 29 Keith DS, Nichols GA, Gullion CM, Brown JB, Smith DH: Longitudinal follow-up and outcomes among a population with chronic kidney disease in a large managed care organization. Arch Intern Med 2004;164:659–663.
- 30 United States Renal Data System: US Department of Public Health and Human Services. Public Health Service. Bethesda, National Institutes of Health, 2003.
- 31 Kopple JD: Nutritional status as a predictor of morbidity and mortality in maintenance dialysis patients. ASAIO J 1997;43:246–250.
- 32 Kalantar-Zadeh K, Kilpatrick RD, Kopple JD, Stringer WW: A matched comparison of serum lipids between hemodialysis patients and nondialysis morbid controls. Hemodial Int 2005;9: 314–324.
- 33 Degoulet P, Legrain M, Reach I, Aime F, Devries C, Rojas P, Jacobs C: Mortality risk factors in patients treated by chronic hemodialysis. Report of the Diaphane Collaborative Study. Nephron 1982;31:103–110.
- 34 Leavey SF, Strawderman RL, Jones CA, Port FK, Held PJ: Simple nutritional indicators as independent predictors of mortality in hemodialysis patients. Am J Kidney Dis 1998;31:997–1006.
- 35 Fleischmann E, Teal N, Dudley J, May W, Bower JD, Salahudeen AK: Influence of excess weight on mortality and hospital stay in 1346 hemodialysis patients. Kidney Int 1999;55:1560–1567.
- 36 Wolfe RA, Ashby VB, Daugirdas JT, Agodoa LY, Jones CA, Port FK: Body size, dose of hemodialysis, and mortality. Am J Kidney Dis 2000;35:80–88.
- 37 Port FK, Ashby VB, Dhingra RK, Roys EC, Wolfe RA: Dialysis dose and body mass index are strongly associated with survival in hemodialysis patients. J Am Soc Nephrol 2002;13: 1061–1066.
- 38 Leavey SF, McCullough K, Hecking E, Goodkin D, Port FK, Young EW: Body mass index and mortality in 'healthier' as compared with 'sicker' haemodialysis patients: results from the Dialysis Outcomes and Practice Patterns Study (DOPPS). Nephrol Dial Transplant 2001;16:2386–2394.
- 39 Glanton CW, Hypolite IO, Hshieh PB, Agodoa LY, Yuan CM, Abbott KC: Factors associated with improved short term survival in obese end stage renal disease patients. Ann Epidemiol 2003;13: 136–143.
- 40 Johansen KL, Young B, Kaysen GA, Chertow GM: Association of body size with outcomes among patients beginning dialysis. Am J Clin Nutr 2004;80:324–332.

Obesity Paradox

- 41 Lowrie EG, Li Z, Ofsthun N, Lazarus JM: Body size, dialysis dose and death risk relationships among hemodialysis patients. Kidney Int 2002;62:1891–1897.
- 42 Beddhu S, Pappas LM, Ramkumar N, Samore M: Effects of body size and body composition on survival in hemodialysis patients. J Am Soc Nephrol 2003;14:2366–2372.
- 43 Kalantar-Zadeh K, Abbott KC, Salahudeen AK: Reverse epidemiology of obesity in dialysis patients: fat or muscle? Am J Clin Nutr 2005;82:910–911.
- 44 Kalantar-Zadeh K, Kuwae N, Wu DY, Shantouf RS, Fouque D, Anker SD, Block G, Kopple JD: Associations of body fat and its changes over time with quality of life and prospective mortality in hemodialysis patients. Am J Clin Nutr 2006;83:202–210.
- 45 Canada-USA Peritoneal Dialysis Study Group: Adequacy of dialysis and nutrition in continuous peritoneal dialysis: association with clinical outcomes. J Am Soc Nephrol 1996;7:198–207.
- 46 Hakim RM, Lowrie E: Obesity and mortality in ESRD: is it good to be fat? Kidney Int 1999;55: 1580–1581.
- 47 McCusker FX, Teehan BP, Thorpe KE, Keshaviah PR, Churchill DN, for the Canada-USA (CANUSA) Peritoneal Dialysis Study Group: How much peritoneal dialysis is necessary for maintaining a good nutritional status? Kidney Int 1996;56:(Suppl)S56–S61.
- 48 Johnson DW, Herzig KA, Purdie DM, Chang W, Brown AM, Rigby RJ, Campbell SB, Nicol DL, Hawley CM: Is obesity a favorable prognostic factor in peritoneal dialysis patients? Perit Dial Int 2000;20:715–721.
- 49 Chung SH, Lindholm B, Lee HB: Influence of initial nutritional status on continuous ambulatory peritoneal dialysis patient survival. Perit Dial Int 2000;20:19–26.
- 50 Snyder JJ, Foley RN, Gilbertson DT, Vonesh EF, Collins AJ: Body size and outcomes on peritoneal dialysis in the United States. Kidney Int 2003;64:1838–1844.
- 51 McDonald SP, Collins JF, Johnson DW: Obesity is associated with worse peritoneal dialysis outcomes in the Australia and New Zealand patient populations. J Am Soc Nephrol 2003;14: 2894–2901.
- 52 Abbott KC, Glanton CW, Trespalacios FC, Oliver DK, Ortiz MI, Agodoa LY, Cruess DF, Kimmel PL: Body mass index, dialysis modality, and survival: analysis of the United States Renal Data System Dialysis Morbidity and Mortality Wave II Study. Kidney Int 2004;65:597–605.
- 53 Aslam N, Bernardini J, Fried L, Piraino B: Large body mass index does not predict short-term survival in peritoneal dialysis patients. Perit Dial Int 2002;22:191–196.
- 54 Stack AG, Murthy BV, Molony DA: Survival differences between peritoneal dialysis and hemodialysis among 'large' ESRD patients in the United States. Kidney Int 2004;65:2398–2408.
- 55 Ramkumar N, Pappas LM, Beddhu S: Effect of body size and body composition on survival in peritoneal dialysis patients. Perit Dial Int 2005;25:461–469.
- 56 Foley RN: Body mass index and survival in peritoneal dialysis patients. Perit Dial Int 2005;25: 435–437.
- 57 Curtis JP, Selter JG, Wang Y, Rathore SS, Jovin IS, Jadbabaie F, Kosiborod M, Portnay EL, Sokol SI, Bader F, Krumholz HM: The obesity paradox: body mass index and outcomes in patients with heart failure. Arch Intern Med 2005;165:55–61.
- 58 Stevens J, Cai J, Pamuk ER, Williamson DF, Thun MJ, Wood JL: The effect of age on the association between body-mass index and mortality. N Engl J Med 1998;338:1–7.
- 59 Yeh S, Wu SY, Levine DM, Parker TS, Olson JS, Stevens MR, Schuster MW: Quality of life and stimulation of weight gain after treatment with megestrol acetate: correlation between cytokine levels and nutritional status, appetite in geriatric patients with wasting syndrome. J Nutr Health Aging 2000;4:246–251.
- 60 Chlebowski RT, Grosvenor M, Lillington L, Sayre J, Beall G: Dietary intake and counseling, weight maintenance, and the course of HIV infection. J Am Diet Assoc 1995;95:428–432; quiz 433–425.
- 61 Wilson DO, Rogers RM, Wright EC, Anthonisen NR: Body weight in chronic obstructive pulmonary disease. The National Institutes of Health Intermittent Positive-Pressure Breathing Trial. Am Rev Respir Dis 1989;139:1435–1438.
- 62 Escalante A, Haas RW, del Rincon I: Paradoxical effect of body mass index on survival in rheumatoid arthritis: role of comorbidity and systemic inflammation. Arch Intern Med 2005;165: 1624–1629.

- 63 Kalantar-Zadeh K, Kilpatrick RD, Kuwae N, Wu DY: Reverse epidemiology: a spurious hypothesis or a hardcore reality? Blood Purif 2005;23:57–63.
- 64 Kalantar-Zadeh K, Abbott KC, Kronenberg F, Anker SD, Horwich TB, Fonarow GC: Epidemiology of dialysis patients and heart failure patients; special review article for the 25th anniversary of the Seminars in Nephrology. Semin Nephrol 2006;26:118–133.
- 65 Kalantar-Zadeh K, Block G, Horwich T, Fonarow GC: Reverse epidemiology of conventional cardiovascular risk factors in patients with chronic heart failure. J Am Coll Cardiol 2004;43: 1439–1444.
- 66 Horwich TB, Fonarow GC, Hamilton MA, MacLellan WR, Woo MA, Tillisch JH: The relationship between obesity and mortality in patients with heart failure. J Am Coll Cardiol 2001;38:789–795.
- 67 Schrier RW, Abraham WT: Hormones and hemodynamics in heart failure. N Engl J Med 1999;341:577–585.
- 68 Yusuf S, Sleight P, Pogue J, Bosch J, Davies R, Dagenais G: Effects of an angiotensin-convertingenzyme inhibitor, ramipril, on cardiovascular events in high-risk patients. The Heart Outcomes Prevention Evaluation Study Investigators. N Engl J Med 2000;342:145–153.
- 69 Stenvinkel P, Marchlewska A, Pecoits-Filho R, Heimburger O, Zhang Z, Hoff C, Holmes C, Axelsson J, Arvidsson S, Schalling M, Barany P, Lindholm B, Nordfors L: Adiponectin in renal disease: relationship to phenotype and genetic variation in the gene encoding adiponectin. Kidney Int 2004;65:274–281.
- 70 Mohamed-Ali V, Goodrick S, Bulmer K, Holly JM, Yudkin JS, Coppack SW: Production of soluble tumor necrosis factor receptors by human subcutaneous adipose tissue in vivo. Am J Physiol 1999;277:E971–E975.
- 71 McClellan WM, Chertow GM: Beyond framingham: cardiovascular risk profiling in ESRD. J Am Soc Nephrol 2005;16:1539–1541.
- 72 Niebauer J, Volk HD, Kemp M, Dominguez M, Schumann RR, Rauchhaus M, Poole-Wilson PA, Coats AJ, Anker SD: Endotoxin and immune activation in chronic heart failure: a prospective cohort study. Lancet 1999;353:1838–1842.
- 73 Rauchhaus M, Coats AJ, Anker SD: The endotoxin-lipoprotein hypothesis. Lancet 2000;356: 930–933.
- 74 Jandacek RJ, Anderson N, Liu M, Zheng S, Yang Q, Tso P: Effects of yo-yo diet, caloric restriction, and olestra on tissue distribution of hexachlorobenzene. Am J Physiol Gastrointest Liver Physiol 2005;288:G292–G299.
- 75 Imbeault P, Tremblay A, Simoneau JA, Joanisse DR: Weight loss-induced rise in plasma pollutant is associated with reduced skeletal muscle oxidative capacity. Am J Physiol Endocrinol Metab 2002;282:E574–E579.
- 76 Macleod J, Davey Smith G: Psychosocial factors and public health: a suitable case for treatment? J Epidemiol Community Health 2003;57:565–570.
- 77 Jones CA, McQuillan GM, Kusek JW, Eberhardt MS, Herman WH, Coresh J, Salive M, Jones CP, Agodoa LY: Serum creatinine levels in the US population: third National Health and Nutrition Examination Survey. Am J Kidney Dis 1998;32:992–999.
- 78 Kalantar-Zadeh K, Balakrishnan VS: The kidney disease wasting: inflammation, oxidative stress and diet-gene interaction. Hemodial Int 2006, in press.
- 79 Liu Y, Coresh J, Eustace JA, Longenecker JC, Jaar B, Fink NE, Tracy RP, Powe NR, Klag MJ: Association between cholesterol level and mortality in dialysis patients: role of inflammation and malnutrition. JAMA 2004;291:451–459.

Kamyar Kalantar-Zadeh, MD, PhD, MPH Associate Professor of Medicine and Pediatrics Division of Nephrology and Hypertension Los Angeles Biomedical Research Institute at Harbor-UCLA Medical Center 1124 West Carson Street Torrance, CA 90502 (USA) Tel. +1 310 222 3891, Fax +1 310 782 1837, E-mail kamkal@ucla.edu