Chronic kidney disease after nephrectomy: a clinically-significant entity?

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Abstract: Worldwide, the kidney is the ninth and 14th most common primary site of cancer in men and women respectively (1). Surgical management with either radical or partial nephrectomy is the mainstay of treatment. Surgical resection of functional kidney parenchyma is associated with reductions in glomerular filtration rate, and can lead to the development of chronic kidney disease (CKD); however, there is currently debate as to whether CKD secondary to surgical removal of a kidney is of clinical significance. Here, it will be argued that CKD is of clinical significance regardless of aetiology, due to the higher cardiovascular and mortality risk which is associated with low glomerular filtration rate.

Keywords: Nephrectomy; kidney cancer; renal cell carcinoma; chronic kidney disease (CKD); estimated glomerular filtration rate

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Introduction

Worldwide, the kidney is the 9th and 14th most common primary site of cancer in men and women respectively (1). Either radical (total) or partial (nephron-sparing) nephrectomy is the most common approach to management of patients with kidney cancer.

There is a large volume of literature evaluating postoperative kidney function following nephrectomy. In a recent systematic review, we identified 312 studies published after the year 2000 which considered kidney function as an outcome after nephrectomy (2). Interestingly, less than 5% of these studies followed patients for longer than an average of five years, and less than 2% considered an average follow-up time beyond 7.5 years (2). Many studies have aimed to describe the benefits of partial nephrectomy compared with radical nephrectomy for improving both postoperative kidney function and overall survival (3,4). While partial nephrectomy undoubtedly leads to nephron mass preservation and higher postoperative kidney function on average, there is still controversy as to benefits in terms of survival outcomes. Indeed, many well-powered observational studies have noted survival benefits for partial nephrectomy compared with radical nephrectomy (3), whereas others, including the only randomised clinical trial comparing the two procedures (5), have reported that this not the case. This apparent incongruity has perpetuated the argument that reductions in kidney function secondary to nephrectomy are not as clinically-significant as chronic kidney disease (CKD) due to a medical aetiology.

It will be argued here that surgical loss of kidney parenchyma should be viewed as a risk factor for worsening kidney function over time; and, that incident CKD [defined as an estimated glomerular filtration rate (eGFR) <60 mL/min per 1.73 m² persisting for a duration of three months or greater] is of clinical significance, regardless of the underlying cause. This will be achieved by evaluating studies which report survival and kidney functional outcomes in: (I) otherwise healthy adults who have donated a kidney; (II) patients with kidney tumours who have undergone partial or radical nephrectomy; and (III) patients who have undergone surgical management of kidney cancer, with or without comparisons made with the general population.
**Living kidney donors**

Perhaps the strongest evidence for the view that surgical removal of nephrons should be considered as a risk factor for future CKD comes from population studies of living kidney donors. Historically, it was understood that living kidney donors were unlikely to experience adverse consequences, and this belief was preserved through studies showing better health outcomes for donors compared with non-donors, which either failed to consider baseline health status as a potential confounding factor, or did not consider an adequate follow-up time to observe adverse events related to unilateral nephrectomy (see Table 1 for a summary of selected studies).

A well-cited example of the former is a population-based study published in 1997 by Fehrman-Ekholm and colleagues, which suggested that Swedish living kidney donors had longer survival times compared with the general population. They reported that 20-year survival was 85% for living donors (compared with 66% predicted survival, based on data from the general population) (6), a result which was most likely a consequence of not adjusting for baseline health status, as suggested by the study's authors. An example of the latter is a population-based study from Ontario, Canada, published by Garg et al. in 2012 (7). This study reported that kidney donors were less likely to experience death or a major cardiovascular event compared with matched non-donors, with a median follow-up of 6.5 years (the rate of death or major cardiovascular event per 1,000 person-years was 4.1 and 2.8 for non-donors and donors, respectively) (7). This study adequately accounted for baseline health status by matching donors to members of the general population, restricting to only the healthiest, by excluding from the analysis anyone with a medical comorbidity which may preclude donation; however, it is possible that the follow-up time was not adequate to discern the adverse effects of unilateral nephrectomy.

In more recent studies with longer follow-up periods, a higher likelihood of death or end-stage kidney disease (ESKD) has been reported following kidney donation (Table 1). In a study which evaluated 1,901 Norwegian kidney donors, Mjøen and colleagues reported higher rates of cardiovascular and all-cause mortality, and ESKD [adjusted hazard ratio (aHR): 1.52, 1.48, and 11.40, respectively] compared with matched non-donors, with a median time to ESKD of 18.7 years (8). Similarly, in the United States, using data from the Organ Procurement and Transplantation Network, Muzaale and colleagues found that the 15-year absolute risk of ESKD in donors was 30.8 per 10,000, compared with only 3.9 per 10,000 in matched non-donors (9). Using data from the same source, Wainright and colleagues recently reported that potential risk factors for ESKD in kidney donors included male sex, higher body mass index, black or Hispanic race, and older age; however, they also reported interactions between race and age (10).

These data all seem to suggest that, when considering a long-enough follow-up time and adequately accounting for baseline health status, kidney donors generally have a small but significantly higher absolute risk of developing ESKD than people who do not undergo unilateral nephrectomy. This inference is supported further by data indicating that patients who have undergone nephrectomy in childhood have a higher risk of cardiovascular disease and premature mortality (11). This is consistent with Brenner's hypothesis, that single-nephron hyperfiltration secondary to functional strain may initiate or perpetuate progressive deterioration of kidney function, and concomitant proteinuria hypertension, with adverse consequences only becoming apparent decades after unilateral nephrectomy (12,13). This therefore supports the argument that removal of functional kidney parenchyma is associated with a higher risk of clinically-significant kidney functional deterioration, even in healthy patients.

**Radical vs. partial nephrectomy**

Although a number of observational studies have reported survival benefits for patients undergoing partial compared with radical nephrectomy (3,4), a phase III randomised trial comparing radical and partial nephrectomy, which was conducted by the European Organisation for Research and Treatment of Cancer (EORTC), reported that this was not the case, and a survival benefit was actually observed for patients managed with radical nephrectomy (14). These conflicting results have generated a great deal of debate as to whether CKD due to surgical removal of nephrons has clinical significance, and because of the fact that the results of randomised controlled trials are generally thought to present higher-quality evidence compared with observational studies, a lot of emphasis has been placed on the results of the EORTC trial. Acknowledging the difficulty of conducting a methodologically-rigorous randomised trial for a surgical intervention, we argue here that because of methodological issues with the design of the trial, and potentially-flawed assumptions in the
Table 1 Summary of selected studies in living kidney donors

<table>
<thead>
<tr>
<th>First author, year</th>
<th>Population</th>
<th>Findings</th>
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<tbody>
<tr>
<td>Fehrman-Ekholm, 1997</td>
<td>• All living kidney donors (Sweden) who donated a kidney between 1964 and 1994 (n=430); • Comparison with national mortality data from the general population</td>
<td>• 41 (9.5%) donors died (range: 15 months–31 years post-donation); • 20-year survival was 85% (predicted survival 66%), indicating survival in living donors was 29% (95% CI: 20–38%) better than in the general population</td>
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<td>Garg, 2012</td>
<td>• All living kidney donors (Ontario, Canada) who donated a kidney between 1992 and 2009 (n=2,028); • Matched healthy non-donors, identified from the general population (n=20,280)</td>
<td>• 16 (0.78%) donors died and 26 (1.3%) experienced a major cardiovascular event (median follow-up: 6.8 years); • 365 (1.8%) non-donors died and 287 (1.4%) experienced a major cardiovascular event (median follow-up: 6.4 years); • Primary outcome—death or major cardiovascular event—less frequent in donors compared with non-donors (2.8 and 4.2 per 1,000 person-years, respectively; aHR: 0.66, 95% CI: 0.48–0.90); • Secondary outcome—death-censored major cardiovascular event—no significant differences between donors and non-donors (1.7 and 2.0 per 1,000 person-years, respectively; aHR: 0.85, 95% CI: 0.57–1.27)</td>
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<td>Mjøen, 2013</td>
<td>• All living kidney donors (Norway) who donated a kidney between 1963 and 2007 (excluding marginal donors; n=1901)*; • Matched healthy non-donors identified from the Health Study of Nord-Trøndelag, enrolled between 1984 and 1987 (n=32,621)</td>
<td>• 224 (12%) donors died (30% due to cardiovascular disease); 9 (0.47%) of donors developed ESKD; • For donors, the median time to ESKD was 18.7 years and the median follow-up time was 15.1 years; • Incidence of ESKD in donors was approximately 302 per million person-years (compared with 100 per million person-years in the general population); • 2,425 (7.4%) non-donors died (28.4% due to cardiovascular disease); 22 (0.07%) of non-donors developed ESKD; • For non-donors, the median follow-up time was 24.9 years; • Kidney donors had a higher rate of all-cause (aHR: 1.48, 95% CI: 1.17–1.88) and cardiovascular mortality (aHR: 1.52, 95% CI: 0.95–2.43), and ESKD (aHR: 11.40, 95% CI: 4.43–29.40)</td>
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<td>Muzaale, 2014</td>
<td>• All living kidney donors (USA) who donated a kidney between 1994 and 2011 (n=96,217), identified from the OPTN; • Matched healthy non-donors identified from the Third National Health and Nutrition Examination Survey (USA), enrolled between 1988 and 1994 (n=96,217)</td>
<td>• 99 (0.10%) donors developed ESKD and mean time to ESKD was 8.6 years; median follow-up time was 7.7 years; • 36 (0.04%) non-donors developed ESKD; mean time to ESKD was 10.7 years; median follow-up time was 15.0 years; • 15-year absolute risk of ESKD in donors and healthy non-donors was 30.8 and 3.9 per 10,000, respectively</td>
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<td>Wainright, 2018</td>
<td>• All living kidney donors (USA) who donated a kidney between 1994 and 2016 (n=123,526), identified from the OPTN</td>
<td>• 218 (0.18%) donors developed ESKD and median time to ESKD was 11.1 years • Male sex (aHR: 1.75), higher BMI (aHR per 5-unit increase: 1.34), black or Hispanic race (aHR: 2.79; 1.29), and older age (aHR: 1.26) were associated with higher risk of ESKD • 20-year absolute risk of ESKD varied between 8 and 111 per 10,000 in 20-year-old white females and 20-year-old black males, respectively</td>
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* donors aged >70 or <20 years, or with a body mass index >30 or <17 kg/m², blood pressure >140/90 mmHg or taking blood pressure medications, or a pre-donation eGFR <70 mL/min per 1.73 m². aHR, adjusted hazard ratio; CI, confidence interval; ESKD, end-stage kidney disease; OPTN, Organ Procurement and Transplantation Network.
interpretation of the results, this clinical trial does not provide evidence that CKD secondary to nephrectomy is of less clinical significance than CKD due to other causes (15,16).

The EORTC trial was initially designed to rule out a 10% difference in 5-year overall survival between radical nephrectomy and partial nephrectomy for simple kidney tumours ≤5 cm at largest diameter, with a hypothesis that radical nephrectomy was associated with worse survival. There were 310 participants recruited between 1992 and 1998 in Europe and participants were randomised 1:1 to each treatment arm. The primary outcome was subsequently changed to rule out a 3% difference in 5-year overall survival. Based on power calculations (where α=0.05 and β=0.20), 300 and 1,300 patients were designated as the minimum sample size to detect a 10% and 3% difference in 5-year survival, respectively. After the change, recruitment was opened up also to patients in the USA and Canada, between 1998 and 2003. In total, 541 patients were recruited from 40 different centres, including the 310 participants recruited between 1992 and 1998 (268 randomised to partial nephrectomy, 273 randomised to radical nephrectomy; 39 patients assigned partial nephrectomy were managed with radical nephrectomy and 16 patients assigned to radical nephrectomy were managed with partial nephrectomy) (5,17). Although this trial had sufficient power to address the initial goal of ruling-out a 10% difference in 5-year survival, it was underpowered to rule-out a 3% difference. It was also underpowered to evaluate differences in the incidence of ESKD. Due to a high proportion of crossover between treatment arms, there is a high risk that confounding by indication also impacted results.

The EORTC trial found that, with a median follow-up of 6.7 years, 85.7%, 64.7% and 1.5%; and 64.7%, 6.3% and 1.6%, of patients developed an eGFR <60, <30 or <15 mL/min per 1.73 m², after radical and partial nephrectomy, respectively (14). It was also reported that 10-year survival was 81.1% and 75.7% in patients randomised to be managed with radical and partial nephrectomy, respectively (5.4% difference in survival) (5).

A major issue with drawing the conclusion that CKD secondary to nephrectomy is not as significant as CKD of a non-surgical aetiology from this study is simply that the trial was not designed to answer this research question: this observation would be reasonable in the context of hypothesis generation, but was far from adequate to infer causality. Although kidney function was considered as a planned secondary analysis of this trial, this was only evaluated in the context of comparing postoperative kidney function (eGFR, continuous and by CKD stage) by surgery type. It was reported that kidney function was better in patients managed with partial nephrectomy (14); however, no analysis was performed evaluating survival mediated by postoperative kidney function. It was therefore unclear how many patients who died also had impaired kidney function. Interestingly, post hoc subgroup analyses of this study showed that for patients with preoperative SCr >25% of the upper limit of normal, the risk of mortality following partial nephrectomy was lower than radical nephrectomy (aHR: 0.8, 95% CI: 0.1–3.4) compared with patients whose SCr was ≤25% of the upper limit of normal, where this effect reversed (aHR: 1.6, 95% CI: 1.0–2.3) (18).

Despite a clear difference in postoperative eGFR between radical and partial nephrectomy, due to the reasonably short follow-up time of this trial (6.7 years) (14), it is also unlikely that a long-enough amount of time had passed to fully appreciate the effect of nephron reduction on risk of mortality, similar to potential limitations in some studies evaluating living kidney donors. Due to these limitations in the EORTC trial’s design, it could be argued that less significance should be placed on inferences related to the clinical significance of postoperative kidney function generated from these results.

Another potential issue with drawing these conclusions about the clinical significance of CKD from this trial is that the argument relies on an inherent assumption that, apart from the amount of functional parenchyma being removed, radical and partial nephrectomy are essentially equivalent. This is not the case, and it is very likely that differences between radical nephrectomy and partial nephrectomy, which extend beyond the degree of surgical nephron reduction between the two procedures, are more than trivial. Regarding potential adverse effects, Shuch et al. discussed this eloquently, arguing that partial nephrectomy should not be viewed as “protective” but rather “less harmful” than radical nephrectomy (19).

A recent systematic review of 21 studies evaluating postoperative complications following nephrectomy reported that partial nephrectomy was associated with a higher rate of complications compared with radical nephrectomy [relative risk (RR): 1.7, 95% CI: 1.3–2.2] (4). This demonstrates that clinically-significant differences between the two procedures exist, which could affect long-term health outcomes, but which are essentially ignored in arguments relating to the association between kidney function and mortality.
It should also be noted that the safety profile of partial nephrectomy has changed significantly since its introduction into clinical practice. As the EORTC trial was conducted over more than a decade (1992 to 2003) during periods where the safety profile of partial nephrectomy was variable, it is possible that variations in surgical technique, and subsequent sequelae of this, affected results. Miller et al. demonstrated that, in an American population-based retrospective study of patients managed with both radical nephrectomy and partial nephrectomy, between 1991 and 1999, there was no statistical difference in the risk of developing ESKD following partial nephrectomy compared with radical nephrectomy (aHR: 1.3, 95% CI: 0.9–1.8); but, between 2000 and 2002 the point estimates reversed, showing ESKD was less likely in patients managed with partial nephrectomy (aHR: 0.7, 95% CI: 0.6–0.9) (20).

No data have been presented comparing mortality rates by era from the EORTC trial, and it is unlikely that these analyses would be appropriately powered, given the trial’s small sample size. It is possible that earlier cases where patients were managed with partial nephrectomy may have disproportionately influenced survival estimates compared with more recent cases, leading to an inflated estimate of the overall mortality risk associated with partial nephrectomy. It is also possible that the ESKD cases associated with partial nephrectomy were from the earlier era, where ESKD was more common after partial nephrectomy. This is particularly relevant given the low event count associated with the development of ESKD in this trial. This remains a point of concern in the interpretation of these results.

Given that the EORTC trial was not adequately powered to assess outcomes related to kidney function and ESKD, had a limited follow-up time, and did not take into account differences between the procedures outside of the amount of functional parenchyma being resected, this trial does not provide strong evidence that CKD subsequent to surgical resection of kidney tissue is of less significance than medical causes of CKD.

**Downstream effects of “surgical” CKD**

An argument that is often made to minimise the perceived risk of CKD following nephrectomy is that there is not strong evidence of subsequent eGFR decline (progressive CKD) in patients with “surgical” CKD. Although this statement is true, this is a common feature of a large number of patients who develop CKD (eGFR <60 mL/min per 1.73 m²) regardless of underlying aetiology. Many patients with stage 3 CKD will not experience a decline in eGFR, have prolonged periods of non-progression, will demonstrate non-linear eGFR decline, or have a widely-fluctuating eGFR or relapsing-remitting disease trajectory (21-23). Notwithstanding, all patients with stage 3 CKD have a higher mortality and cardiovascular risk, compared with patients without CKD. It is therefore important to correlate postoperative eGFR with hard clinical end-points such as mortality, as the absence of progressive CKD does not equate with the absence of risk for adverse outcomes.

One of the largest databases of nephrectomy patients is maintained by the Cleveland Clinic in the USA, and in this section, we will primarily discuss studies which utilise these data. General findings of these studies demonstrate that, compared with patients who do not develop CKD after nephrectomy, patients who do develop CKD, and had CKD prior to surgery, have a higher risk of all-cause mortality (for example, aHR: 1.2, 95% CI: 1.0–1.4; and aHR: 2.0, 95% CI: 1.7–2.3, respectively, with a median follow-up of 9.4 years, n=4,299) (24). A common conclusion of these studies is that “medical”, “surgical” and “medical/surgical” CKD (referring to the broad underlying aetiology, or combination thereof, driving reductions in kidney function) are associated with different risks of clinically-significant end-points. We will argue that these distinctions are arbitrary, and that the key factor that determines risk is the postoperative eGFR, regardless of the underlying cause.

Nephrectomy in patients with preoperative CKD, and patients without preoperative CKD who go on to develop postoperative CKD, can lead to adverse outcomes and progressive eGFR decline, caused by either maladaptation to nephron mass reduction, or because of underlying damage to the kidneys. In patients with clinically-evident CKD prior to nephrectomy, this damage has already been identified. In patients without clinically-evident CKD prior to nephrectomy, this damage may not be present, or may only be mild, and may never have become symptomatic in the absence of surgical mass reduction. Undergoing nephrectomy modifies this, and patients are subsequently classed as having CKD. Although the distinction of CKD before surgery is important in terms of individual patient prognosis: that patients with CKD prior to undergoing nephrectomy are at higher risk of adverse events compared with patients who do not; extrapolating this to infer that new-onset CKD after surgery is of less significance than new-onset CKD of other causes is not appropriate, as these patient groups are not comparable in this respect.

If patients who develop CKD after nephrectomy are
considered to have had subclinical CKD before surgery, then the reason for the difference in terms of overall survival between the two groups (patients with CKD before surgery and patients with incident CKD after nephrectomy) could be attributed to lead-time bias. If the underlying pathological processes are considered to be essentially equivalent between subclinical and clinically-evident CKD, then the only difference is the initial time of onset. As patients with evidence of CKD before surgery are more likely to have had an earlier onset of the underlying pathology contributing to CKD, they are therefore more likely to die as a consequence of CKD than patients who develop clinically-evident CKD only after undergoing nephrectomy. Therefore, if causal inference is intended, it cannot be concluded that new-onset CKD after nephrectomy is a less significant disease process compared with prevalent CKD, based on the findings that patients with “medical/surgical” CKD have a higher mortality rate than patients with just “surgical” CKD. Although it is more likely that patients with prevalent CKD will experience adverse events, this finding is not unexpected. Mortality risk is inversely proportional to eGFR, regardless of the population of interest (25–28). In terms of evaluating the clinical significance of post-nephrectomy CKD, the important comparison in these studies was between patients with incident CKD after nephrectomy and patients who did not develop CKD after nephrectomy—this comparison showed that patients with incident CKD had significantly higher mortality rates.

Therefore, the next relevant question is: how does the risk of mortality for patients with new-onset CKD after surgery compare to that of patients with medical causes of CKD who do not undergo nephrectomy? This was partially addressed using the Cleveland Clinic Surgical Registry dataset by Demirjian et al., who compared all-cause mortality and non-renal cancer mortality in patients with “surgical” and “medical/surgical” CKD (n=1,097 and 1,053, respectively) and a cohort of 42,658 patients with CKD which was not secondary to nephrectomy, who were managed by nephrologists at the Cleveland Clinic (29). The authors showed that patients with “medical” and “medical/surgical” CKD had similar risk of all-cause and non-renal cancer mortality, and that the risk was higher than for patients who developed CKD only after nephrectomy. This study was limited by the fact that patients who did not develop CKD after nephrectomy were not included. The risk of bias introduced by performing a head-to-head comparison of these patients was also quite large, given that patients referred to a nephrologist will typically have at least moderately-severe CKD, which again introduces the risk of lead-time bias (30). It is unclear what the exclusion criteria were for the comparison group of patients with CKD of a medical aetiology and, unlike patients undergoing nephrectomy, they have no clear precipitating event to designate T₀, which makes the interpretation of time-to-event analyses difficult.

A different approach is to compare the RR for mortality for patients with and without new-onset CKD after nephrectomy to a well-established value in the literature, comparing patients with and without medical CKD in the general population, with a comparable follow-up period. Tonelli et al. conducted a systematic review of 39 studies which included populations of patients who were at risk of CKD, and had kidney function and mortality data recorded (31). Overall there were 1,371,990 patients included, with a median follow-up of 4.5 (0.8–16.0) years. They reported that patients with CKD (eGFR <60 mL/ min per 1.73 m²) had an increased risk of all-cause mortality (RR: 1.8, 95% CI: 1.3–2.3) compared with patients with a postoperative eGFR ≥60 mL/min per 1.73 m².

Data from another study which used the Cleveland Clinic Surgical Registry conducted by Wu et al., who evaluated mortality after nephrectomy in patients who did and did not develop CKD after surgery (n=931 and 2,202, respectively), was compared with the mortality estimates from the general population discussed above (26). We calculated the RR comparing non-kidney cancer-related mortality in patients who did and did not develop new-onset CKD after nephrectomy, using methods described previously (32). This analysis demonstrated that patients who developed new-onset CKD had an increased risk of mortality than those who did not (RR: 1.4, 95% CI: 1.1–1.8). The 10-year mortality risk was reasonably similar (RR: 1.5, 95% CI: 1.2–1.8). Although the point-estimates are slightly lower than those reported by Tonelli et al., there was substantial overlap between the 95% CIs, which tends to indicate that patients with new-onset CKD after surgery were at increased risk of death compared with patients who did not develop CKD after nephrectomy, and at comparable risk to patients with CKD of a medical aetiology compared with patients without CKD in the general population (Figure 1). As this is a relative measure, there is an underlying assumption that patients who do not develop CKD following surgery (i.e., postoperative eGFR >60 mL/min per 1.73 m²) were at similarly comparable risk of death compared with patients without CKD who do not undergo
nephrectomy. Based on the results of studies using the Cleveland Clinic Surgical Registry, it could be extrapolated that new-onset CKD (CKD, defined as a GFR <60 mL/min per 1.73 m²) after nephrectomy was associated with an increased risk of mortality compared with patients who developed CKD after oncological nephrectomy with patients who did not develop CKD. The RR for the general population was reported in the systematic review of patients with and without CKD in the general population conducted by Tonelli et al. There is substantial overlap between the 95% CI of these two estimates. RR, relative risk; CKD, chronic kidney disease; GFR, glomerular filtration rate.

Summary

There is a flaw in the argument that CKD secondary to surgical nephron reduction is of less clinical significance than CKD of other causes does not have a strong evidence base. Finally, we showed that, even in the absence of progressive CKD, developing CKD (eGFR <60 mL/min per 1.73 m²) after nephrectomy is associated with a higher risk of mortality compared with patients who do not develop CKD after nephrectomy, and that this risk is essentially equivalent with the risk of mortality for patients with CKD of any cause.

Thresholds for CKD stages in clinical guidelines are based on the fact that 60 mL/min per 1.73 m² is approximately half of the maximum physiological eGFR of the average person, is able to be distinguished accurately by estimating equations, and is associated with increased risk of adverse events (23). In clinical practice, the distinction of CKD stages is arbitrary, and patients can experience eGFR changes in a fluctuating or relapsing-remitting fashion. It is clear that not all patients with an eGFR <60 mL/min per 1.73 m² from a medical cause will experience further decline in kidney function, and many even experience improvements (21), not unlike patients undergoing nephrectomy; but the fact that not all patients who experience new-onset CKD after nephrectomy will experience a progressive decline in kidney function is cited as a reason for not considering it to have the same clinical significance as CKD of a medical cause (35). In the community, a patient with an eGFR fluctuating around 60 mL/min per 1.73 m² will not be referred to a nephrologist, unless a clear pattern of rapid decline is present, or there is another indication such as uncontrolled hypertension; but, they should undergo regular monitoring of eGFR and urinary albumin-creatinine ratio, and subsequently referred if function begins to deteriorate (30).

Conclusions

Patients who develop new-onset CKD after surgical management of kidney tumours should be considered differently to patients who do not, as, at a population level, they have increased risk of adverse events, including all-cause and cardiovascular mortality, and ESKD. Although a patient who ends up with an eGFR fluctuating around 50–60 mL/min per 1.73 m² after nephrectomy should not be rushed off to a nephrologist, unless a clear pattern of rapid decline is present, or there is another indication such as uncontrolled hypertension; but, they should undergo regular monitoring of eGFR and urinary albumin-creatinine ratio, and subsequently referred if function begins to deteriorate (30).
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Footnote

Conflicts of Interest: The author has no conflicts of interest to declare.

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