Impact of acute kidney injury in elderly versus young deceased donors on post-transplant outcomes: A multicenter cohort study

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We investigated the impact of acute kidney injury (AKI) in elderly deceased-donors (DDs) vs. AKI in young DDs on post-transplant clinical outcomes. A total of 709 kidney transplant recipients (KTRs) from 602 DDs at four transplant centers were enrolled. KTRs were divided into young-DDKT and elderly-DDKT groups according to the age of DD of 60 years. Both groups were subdivided into non-AKI-KT and AKI-KT subgroups according to AKI in DDs. We investigated short-term and long-term clinical outcomes of non-AKI-DDKT and AKI-DDKT subgroups within young-DDKT and elderly-DDKT groups. The incidence of DGF in the AKI-DDKT subgroup was higher and the allograft function within 12 months after KT in the AKI-DDKT subgroup was lower than those in the non-AKI-DDKT subgroup in both young-DDKT and elderly-DDKT groups. Death-censored allograft survival rate was significantly lower in the AKI-elderly-DDKT subgroup than that in the non-AKI-elderly-DDKT subgroup, but it did not differ between AKI-young-DDKT and non-AKI-young-DDKT subgroup. In multivariable analysis, AKI-elderly-DDKT was an independent risk factor for allograft failure (hazard ratio: 2.648, 95% CI: 1.170–5.994, p = 0.019) and a significant interaction between AKI and old age in DDs on allograft failure was observed (p = 0.001). AKI in elderly DDs, but not in young DDs, can significantly affect long-term allograft outcomes of KTRs.

With a tremendous increase in the number of patients with end-stage renal disease, donor shortage in kidney transplantation (KT) has become a worldwide crucial issue to solve. In this regard, the use of kidneys from elderly deceased donors (DDs) has been proposed as an important strategy for solving this donor shortage. It has been reported that the proportion of elderly brain death patients was higher than that of younger brain death patients and that KT from elderly DDs could give survival benefit in comparison with those remaining on dialysis. In addition, in contrast to living donor KT, kidney donation from elderly DD is fully free from donor safety issue. These reasons justify the use of kidney from elderly DDs.
Acute kidney injury (AKI) is very commonly detected in individuals with brain death state for various reasons. Donor shortage also has driven the use of kidney from DDs with AKI. Previous studies including our own have shown that the long-term allograft outcome of KT from DDs with AKI is not significantly different from that of KT from DDs without AKI. However, with respect to transplant-naive individuals, long-term renal outcomes after AKI episode depend on the underlying status of the kidney. Renal outcomes after AKI episode depend on the underlying status of the kidney. Renal outcomes of AKI in elderly DDs might have adverse impact on long-term post-transplant allograft outcomes in their corresponding recipients. However, this has not been fully investigated yet.

Based on these backgrounds, the aim of this study was to investigate whether the impact of AKI in elderly DDs on post-transplant clinical outcomes might differ from that of AKI in young DDs. For this, short-term and long-term clinical outcomes of KT from DDs without AKI (non-AKI-DDKT subgroup) versus those for KT from DDs with AKI (AKI-DDKT subgroup) within young-DDKT and elderly-DDKT groups were compared. The interaction between AKI and old age in DDs on post-transplant allograft survival was also investigated.

**Results**

**Comparison of baseline characteristics between young-DDKT and elderly-DDKT groups and between non-AKI and AKI subgroups within each group.** The median follow-up period of this study was 62.5 (interquartile range: 42.3–91.7) months. The proportion of HTN was significantly higher in the elderly DD group than that in the young DD group (41.8% vs. 17.0%, p < 0.001). There was no significant difference in the proportion of AKI or the distribution of AKI severity between young DD and elderly DD groups. KDPI score was significantly higher in the elderly-DDs than that in the young-DDs. In corresponding recipients, the mean age and the proportion of DM as primary renal disease were higher in the elderly-DDKT group than those in the young-DDKT group (63.6 ± 3.0 vs. 46.7 ± 8.5 years, p < 0.001; 27.0% vs. 18.9%, p = 0.054). The proportion of re-transplants was significantly lower in the elderly-DDKT group than that in the young-DDKT group (5.4% vs. 11.9%, p = 0.046) (see Supplementary Table S1).

In the subgroup analysis, mean donor age, BMI, the proportion of cause of donor death by CVA, and KDPI score were significantly higher in the subgroup of young DDs than that in the subgroup of non-AKI DDs. However, in the group of elderly DDs, there were no significant differences in donor age, gender, BMI, HTN, DM, cause of donor death by CVA, or baseline GFR between non-AKI DDs and AKI DDs subgroups. KDPI score was higher in AKI-DDs subgroup than that in the non-AKI-DDs subgroup in both young- and elderly-DDKT group. In corresponding recipients, proportions of anti-thymocyte globulin were significantly higher in young-DDKT and elderly-DDKT groups with AKI in comparison with those without AKI (33.8% vs. 18.2%, p < 0.001 and 43.8% vs. 19.1%, p = 0.008, respectively) (Table 1).

**Impact of AKI in DDs on the development of delayed graft function between young-DDKT and elderly-DDKT groups.** The incidence of delayed graft function (DGF) was significantly higher in the young-DDKT group than that in the elderly-DDKT group (p < 0.05) (Fig. 1A). The incidence of DGF was significantly lower in the elderly-DDKT group than that in the young-DDKT group (41.8% vs. 17.0%, p < 0.001). There was no significant difference in the incidence of DGF between elderly-DDKT with AKI subgroup and elderly-DDKT without AKI subgroup (Fig. 1C). In the young-DDKT group, AKI stage 3 showed higher incidence of DGF than non-AKI, AKI stage 1, and AKI stage 2 (p < 0.05) (Fig. 1D). In the elderly-DDKT group, the incidence of DGF showed higher tendency in AKI stage 3 than non-AKI, AKI stage 1, and AKI stage 2, but it did not show statistical significance. In the multivariable analysis, donor AKI was an independent risk factor for the development of DGF, adjusted by recipient age, recipient gender, donor gender, donor age, prior KT, induction and maintenance of immunosuppressant (hazard ratio: 3.604, 95% CI: 2.279–5.698, p < 0.001). However, donor age did not show any significance for the development of DGF (Table 2).

**Impact of AKI in DDs on the change of allograft function between young-DDKT and elderly-DDKT groups.** Allograft function between 2 weeks and 12 months post-KT was significantly lower in the elderly-DDKT group than that in the young-DDKT group (Fig. 2A). In the young-DDKT group, allograft function through the first 12 months post-KT was significantly lower in the AKI-young-DDKT subgroup than that in the non-AKI-young-DDKT subgroup (p < 0.05) (Fig. 2B). In the elderly-DDKT group, allograft function between 2 days and 2 weeks post-KT was significantly lower in the AKI-elderly-DDKT subgroup than that in the non-AKI-elderly-DDKT subgroup (p < 0.05), although it did not differ between the two subgroups at 1 year from KT (Fig. 2C). According to the AKI severity, those with AKI stage 3 had lower allograft function from 1 week after KT to the first 12 months post-KT in both young-DDKT group and elderly-DDKT group compared to those in the non-AKI subgroup (p < 0.05 respectively) (Figs. 2D,E). The incidence of biopsy-proven acute rejection (BPAR) within the first year after KT did not differ significantly between the young-DDKT group and the elderly-DDKT group or between two subgroups in young-DDKT or elderly-DDKT group.

**Impact of AKI in DDs on death-censored allograft survival between young-DDKT and elderly-DDKT groups.** A total of 71 (71/709, 10.0%) cases had graft failure, including 57 (8.0%) cases in the young-DDKT group (28 (3.9%) patients in the non-AKI-young-DDKT subgroup and 29 (4.1%) patients in the AKI-young-DDKT subgroup) and 14 (2.0%) in the elderly-DDKT group (3 (0.4%) patients in the non-AKI-elderly-DDKT subgroup and 11 (1.6%) patients in the AKI-elderly-DDKT subgroup). No significant difference was detected in the distribution of causes of allograft failure between the young-DDKT group and the
elderly-DDKT group or between non-AKI-KT and AKI-KT subgroups within young-DDKT or elderly-DDKT group (Table 3). Death-censored allograft survival rate was significantly lower in the elderly-DDKT group than that in the young-DDKT group ($p < 0.05$) (Fig. 3A). In subgroup analysis, death-censored allograft survival rate was significantly lower in the AKI-elderly-DDKT subgroup than in the non-AKI-elderly-DDKT subgroup ($p < 0.05$) (Fig. 3C), although it was not significantly different between non-AKI-young-DDKT and the AKI-young-DDKT subgroups (Fig. 3B). There was no significant difference according to the AKI stage in the young-DDKT group (Fig. 3D), but AKI stage 3 showed the lowest death-censored allograft survival rate in the AKI-young-DDKT subgroups (Fig. 3B). There was no significant difference according to the AKI stage in the elderly-DDKT group in comparison with non-AKI-KT and AKI-KT subgroups within young-DDKT or elderly-DDKT group (Fig. 3E). In the multivariable analysis, AKI-elderly-DDKT was an independent risk factor for allograft failure (hazard ratio: 2.648, 95% CI: 1.706–3.994, $p = 0.012$) after adjusting by transplant years (1996–2005 vs. 2006–2010 vs. 2011–2017), transplant centers, recipient age, recipient gender, donor gender, DGF, and acute rejection. There was a significant interaction between AKI in DDs and elderly DDs for allograft failure ($p$ for interaction = 0.002) (Table 4).

### Table 1. Comparison of clinical and laboratory parameters according to acute kidney injury in young- or elderly donors. Values are expressed as means ± SDs, n (%). Abbreviations: DDKT, deceased donor kidney transplantation; AKI, acute kidney injury; CVA, cerebrovascular accident; ESRD, end-stage renal disease, HLA, human leukocyte antigen; PRA, panel reactive antibody; CKD-EPI, chronic kidney disease-epidemiology collaboration; KDPI, kidney donor profile index

| Variables                        | Young-DDKT | Elderly-DDKT | p for Trend | p for Trend |
|----------------------------------|------------|--------------|-------------|-------------|
| Non-AKI-KT                       | AKI-KT     | Non-AKI-KT   | AKI-KT      |             |
| Donors                           | n = 246    | n = 276      | n = 34      | n = 46      |
| Age at KT (years)                | 39.2 ± 14.2| 44.3 ± 11.8  | <0.001      | 63.9 ± 3.7  | 63.8 ± 3.5  | 0.916 |
| Gender (Male: Female)            | 151: 95    | 209: 67      | <0.001      | 20: 14      | 33: 13      | 0.243 |
| Body mass index (kg/m²)          | 22.4 ± 3.8 | 23.8 ± 3.6   | <0.001      | 24.1 ± 3.1  | 22.9 ± 2.7  | 0.088 |
| Hypertension, n (%)              | 42 (17.4)  | 46 (16.7)    | 0.816       | 14 (42.4)   | 19 (41.3)   | 1.000 |
| Diabetes mellitus, n (%)         | 20 (8.3)   | 22 (8.0)     | 1.000       | 2 (6.1)     | 10 (21.7)   | 0.065 |
| Cause of donor death - CVA, n (%)| 154 (62.6) | 212 (76.8)   | <0.001      | 27 (79.4)   | 31 (67.4)   | 0.125 |
| Baseline GFR (ml/min/1.73 m²) (CKD-EPI) | 86.8 ± 30.2 | 82.4 ± 21.1 | 0.100       | 76.4 ± 21.9 | 75.7 ± 13.3 | 0.898 |
| GFR at allocation (ml/min/1.73 m²) (CKD-EPI) | 102.5 ± 31.5 | 40.0 ± 23.3 | <0.001      | 81.2 ± 24.0 | 34.5 ± 20.7 | <0.001 |
| KDPI score (%)                   | 50.3 ± 22.7| 65.4 ± 19.8  | <0.001      | 91.3 ± 5.4  | 94.4 ± 4.9  | 0.011 |
| Transplant year, n (%)           | 0.060      |              |             | 0.808       |
| Gender (Male: Female)            | 156: 108   | 193: 141     | 0.802       | 30: 17      | 39: 25      | 0.844 |
| Body mass index (kg/m²)          | 22.9 ± 3.4 | 23.2 ± 4.1   | 0.261       | 23.6 ± 3.6  | 23.6 ± 3.6  | 0.974 |
| Hypertension, n (%)              | 215 (81.4) | 285 (85.3)   | 0.222       | 41 (87.2)   | 58 (90.6)   | 1.000 |
| Diabetes mellitus, n (%)         | 45 (17.0)  | 68 (20.4)    | 0.344       | 11 (23.4)   | 19 (29.7)   | 0.521 |
| Dialysis duration, years         | 8.3 ± 8.3  | 8.2 ± 10.8   | 0.924       | 6.4 ± 5.2   | 8.4 ± 14.5  | 0.354 |
| Glomerulonephritis               | 147 (55.7) | 143 (42.8)   | 0.131       | 55.1 ± 9.5  | 54.2 ± 7.6  | 0.591 |
| Diabetes mellitus                | 34 (12.9)  | 58 (17.4)    |            | 3 (6.4)     | 4 (6.3)     | 0.635 |
| Hypertension                     | 36 (13.6)  | 65 (19.5)    |            | 10 (21.3)   | 15 (23.4)   | 0.579 |
| Cause of ESRD, n (%)             | 0.016      |              |            | 0.824       |
| Cold ischemic time (min)         | 255.4 ± 123.8 | 244.7 ± 127.3 | 0.321      | 257.8 ± 110.6 | 238.2 ± 113.6 | 0.383 |
| HLA mismatch number              | 3.5 ± 1.6  | 3.7 ± 1.4    | 0.054       | 3.8 ± 1.7   | 3.9 ± 1.5   | 0.865 |
| Induction, n (%)                 | <0.001     |              |            | 0.008       |
| Main immunosuppressant           | 216 (81.8) | 221 (66.2)   | 0.38        | 38 (80.9)   | 36 (56.3)   | 0.008 |
| Tacrolimus: Cyclosporine          | 48 (18.2)  | 113 (33.8)   | 0.016       | 9 (19.1)    | 28 (43.8)   | 0.434 |
| PRA > 50%, n (%)                 | 26 (19.4)  | 33 (15.5)    | 0.380       | 3 (10.7)    | 7 (16.7)    | 0.729 |

Impact of AKI in DDs on patient survival between young-DDKT and elderly-DDKT groups. A total of 45 (45/709, 6.3%) patients died in this population. Among them, 37 (5.2%) patients were included in the young-DDKT group (19 (2.7%) patients in the non-AKI-young-DDKT subgroup and 18 (2.5%) patients
in the AKI-young-DDKT subgroup and 8 (1.1%) patients were included in the elderly-DDKT group (2 (0.3%) patients in the non-AKI-elderly-DDKT subgroup and 6 (0.8%) patients in the AKI-elderly-DDKT subgroup). There was no significant difference in the dispersion of the causes of patient death (Table 3). There

Figure 1. Comparison of the incidence of DGF (A) between young-DDKT and elderly-DDKT groups and between non-AKI-DDKT and AKI-DDKT subgroups within (B) young-DDKT and (C) elderly-DDKT groups. Comparison of the incidence of DGF according to the AKI stage in (D) young-DDKT and (E) elderly-DDKT groups. *p < 0.05 vs. young-DDKT, †p < 0.05 vs. non-AKI.

Table 2. Risk factors for development of delayed graft function in the deceased donor kidney transplantation. *Adjusted by recipient age, recipient gender, donor gender, donor age, prior KT, induction and maintenance immunosuppressant. Abbreviations: DDKT, deceased donor kidney transplantation; CVA, cerebrovascular accident; PRA, panel reactive antibody; HLA, human leukocyte antigen; CsA, cyclosporine A; ATG, antithymocyte globulin.
was also no significant difference in patient survival rate between young-DDKT and elderly-DDKT groups, or between non-AKI-young-DDKT and AKI-young-DDKT subgroups, or between non-AKI-elderly-DDKT and AKI-elderly-DDKT subgroups (see Supplementary Fig. S2).

| Variables                        | Young DDKT |          | Elderly DDKT |          |
|----------------------------------|------------|----------|--------------|----------|
|                                  | AKI-KT     | p for Trend | Non-AKI-KT   | AKI-KT   | p for Trend |
| Causes of graft failure, n (%)   |            |          |              |          |            |
| Acute rejection                  | 6 (21.4)   | 0.275    | 0            | 3 (27.2) | 1.000       |
| Chronic allograft dysfunction    | 12 (42.9)  | 19 (65.5) | 2 (66.7)     | 4 (36.4) |
| Chronic antibody mediated rejection | 1 (3.6)  | 0        | 0            | 1 (9.1)  |
| Recurrent glomerulonephritis     | 2 (7.1)    | 1 (3.4)  | 1 (33.3)     | 2 (18.2) |
| BK virus-associated nephropathy  | 5 (17.9)   | 1 (3.4)  | 0            | 1 (9.1)  |
| Unknown                          | 2 (7.1)    | 2 (7.0)  | 0            | 0        |
| Causes of death, n (%)           |            | 0.036    |              |          | 1.000       |
| Cardiovascular disease           | 3 (15.8)   | 4 (22.2) | 0            | 2 (33.3) |
| Infection                        | 5 (26.3)   | 8 (44.4) | 2 (100.0)    | 3 (50.0) |
| Malignancy                       | 4 (21.1)   | 0        | 0            | 1 (16.7) |
| Bleeding                         | 2 (10.5)   | 1 (5.6)  | 0            | 0        |
| Hepatic failure                  | 3 (15.8)   | 0        | 0            | 0        |
| Unknown                          | 2 (10.5)   | 5 (27.8) | 0            | 0        |

Table 3. Comparison of clinical outcomes according to acute kidney injury in young- or elderly donors. Values are expressed as means ± SDs, n (%). Abbreviations: DDKT, deceased donor kidney transplantation; AKI, acute kidney injury.
Discussion
In this study, we investigated the impact of AKI occurred in elderly-DDs on the clinical outcomes after KT. As a result, we found that AKI itself did not show significant influence on long-term allograft outcomes (see Supplementary Fig. S3(A)). However, KT from elderly DDs with AKI showed worse long-term allograft survival than that from elderly DDs without AKI. This finding was not shown in transplant from young-DDs. Especially, KT from elderly DDs with AKI stage 3 showed worse long-term allograft survival than that other DDKT (see Supplementary Fig. S3(B)). Our results show that strategies to prevent or minimize the occurrence of AKI in DDs, especially in elderly, might be needed to improve long-term allograft outcomes.

First, we compared the clinical characteristics between young-DDs and elderly-DDs. Incidences of HTN and KDPI score were significantly higher in elderly-DDs. Since the presence of HTN and higher KDPI score could an suggest underlying chronic tissue injury, those donors could be diagnosed with CKD17. In the subgroup analysis, age and proportion of death due to CVA and BMI were significantly higher in the AKI-DD subgroup than in the non-AKI-DD subgroup of the young-DDKT group. It was difficult to see the direct-cause-result relationship because data were cross-sectional. However, higher BMI and CVA might be related to the development of AKI in these patients18–20.

Next, we compared short-term post-transplant clinical outcomes between young-DDKT and elderly-DDKT groups and between non-AKI and AKI subgroups. In consistent with our previous study 11,12,21, the incidence of DGF was significantly higher or showed higher tendency when DDs had AKI irrespective of age of donor. In terms of the change of allograft function, allograft function at 1 year from KT showed lower value in the

| Table 4. Hazard ratios of allograft failure according to donor age (>60) or acute kidney injury. aAdjusted by transplant years (1996–2005 vs. 2006–2010 vs. 2011–2017), transplant centers, recipient age, recipient gender, donor gender, delayed graft function, acute rejection. Abbreviations: AKI, acute kidney injury; DDKT, deceased donor kidney transplantation; HR, hazard ratio; C.I., confidence interval. |
|-----------------|-----------------|-------|-----------------|-------|
| Non-AKI-young DDKT | Reference       | P     | Non-AKI-young DDKT | Reference |
| AKI-young DDKT | 0.820 (0.488–1.380) | 0.456 | 1.091 (0.615–1.935) | 0.767 |
| Non-AKI-elderly DDKT | 0.763 (0.231–2.517) | 0.657 | 0.825 (0.190–3.583) | 0.798 |
| AKI-elderly DDKT | 2.462 (1.212–5.002) | 0.013 | 2.648 (1.170–5.994) | 0.019 |
| P-value for interaction | 0.002 | | |

Figure 3. Comparison of death-censored allograft survival rate (A) between young and elderly-DDKT groups and between non-AKI-DDKT and AKI-DDKT subgroups within (B) young-DDKT and (C) elderly-DDKT groups. Death-censored allograft survival rate according to AKI stage in (D) young-DDKT and (E) elderly-DDKT groups.
elderly-DDKT group than in the young-DDKT group. It may be because baseline capacity to recover is lower in high-KDPI subgroups due to pre-existing permanent functional loss of nephron in the elderly donor at baseline as we mentioned above. In the subgroup analysis, the AKI-young-DDKT group showed lower allograft function at 1 year after KT than the non-AKI-young-DDKT group. As we mentioned above, it may be because AKI-young-DDKT showed older age and lower eGFR at baseline. Hence, their capacity to recover might be limited. In contrast, allograft function at 1 year after KT showed similar value between non-AKI-elderly-DDKT and AKI-elderly-DDKT subgroups. It might be because the baseline eGFR did not differ between the two subgroups. When we analyzed the change of allograft function until 1 year from KT according to AKI stage, although the allograft function was the lowest at 1 week after KT from DD with AKI stage 3 in both young-DDKT and elderly-DDKT groups, it showed similar values to those who took kidneys from DDs with stage 1 or 2 AKI at 1 year from KT. All these findings suggest that recently developed AKI in DDs might be a strong risk factor for DGF. For allograft function recovery at 1 year, baseline functional status could have a more significant impact.

Our major issue is whether AKI in DDs has different influence on long-term allograft survivals between young-DDKT and elderly-DDKT groups. As we expected, AKI in DDs showed significant adverse effect on allograft survivals in elderly-DDKT, especially in AKI stage 3, but not in the young-DDKT group. Interestingly, the AKI-elderly-DDKT subgroup showed significantly worse allograft survival than the other three subgroups (non-AKI-elderly-DDKT, non-AKI-young-DDKT, and AKI-young-DDKT subgroups) (see Supplementary Fig. S1). In multivariable analysis using Cox regression hazard model, co-existence of AKI and elderly-DD was a significant risk factor for allograft failure. Both AKI in DDs and elderly DDs showed a significant interaction on allograft failure. However, either AKI in DDs or elderly in DD alone did not show significance in multivariable analysis as shown in previous reports including ours.

Finally, above findings could be explained by decreased functional nephron mass at baseline in kidney from elderly DDs. In addition, development of inflammation by AKI beyond CKD state can accelerate the progression of kidney fibrosis. Hence, AKI in transplant naive patients can show more significant impact in elderly patients than in young patients in terms of a long term renal prognosis.

In contrast to our expectation, the proportion of chronic allograft dysfunction as the cause of allograft failure was not different between non-AKI-elderly-DDKT and AKI-elderly-DDKT subgroups. At first, we expected that allograft failure due to progressive allograft injury after AKI might develop as non-specific “chronic allograft nephropathy” instead of immunologic process such as acute rejection or cAMR. However, although allograft rejection could be a direct cause of allograft failure in some patients, the probability for allograft failure can increase when it develops in kidney with decreased nephron mass such as kidney from elderly DDs. Another possible reason is that AKI can induce the activation of innate immune system. After transplant, it can be a crosstalk to the activation of allograft rejection. Taken together, the main reason of the adverse allograft outcome in AKI-elderly-DDKT subgroup might be the low nephron mass at baseline. Hence, additional insult such as acute kidney injury before transplant and allo-immune responses after transplant may accelerate allograft failure.

In contrast to death-censored allograft survival rate, patient survival rate did not differ significantly between young-DDKT and elderly-DDKT groups or between AKI-DDKT and non-AKI-DDKT subgroups. It might be because patient death rate was so low that it cannot draw any conclusion about this issue. Another possible reason was that recipient age had such a strong effect on patient survival rate that the effect of other factors could not appear.

Our study has some limitations as suggested in our previous reports using this cohort. First, because it was a retrospective study, our study had a possibility of selection bias. However, we analyzed a large number of KTRs from multiple centers. In addition, we also adjusted our results considering transplant centers and transplant year in the multivariable analysis. Second, we did not have detailed data or clinical outcomes of contralateral kidneys transferred to another transplant center according to the kidney allocation rule in Korea, which might have induced a bias during analysis. Well-designed prospective multi-center studies are required to overcome these issues. Lastly, we did not suggest any possible strategies to prevent or early detect AKI in elderly-DDs. In the recent, procurement biopsies are suggested as strategies to better detect grafts at high risk of worse outcomes for protecting against AKI in elderly-DDs to prevent its adverse effect on post-transplant allograft outcomes. Some research reported that procurement biopsies were performed in approximately one-half of all DD kidneys and 85% of expanded criteria donor/donor kidneys with high kidney donor profile index score in the United States, it was useful to predict the post-transplant allograft outcome.

In conclusion, our study suggests that AKI occurred to elderly-DDs has a synergistically adverse effect on long-term allograft outcomes after KT in corresponding recipients. Therefore, careful monitoring and strategies against occurrence of AKI in DDs, especially in elderly-DDs, to prevent its adverse effect on post-transplant allograft outcomes.

Materials and Methods

Study population. A total of 709 KTRs receiving kidneys from 602 DDs at four transplant centers between October 1996 and December 2017 were enrolled. Young and elderly DDs were defined according to the age at the time of donation (<60 years vs. ≥ 60 years). AKI in DDs was defined according to the Kidney Disease: Improving Global Outcomes (KDIGO) criteria as described in previous reports. Patient distribution is presented in Fig. 4.

KTRs were classified into young-DDKT and elderly-DDKT groups. Each group was subdivided into non-AKI-DDKT and AKI-DDKT subgroups according to whether their corresponding DDs were young or elderly and whether they were diagnosed to AKI or not. Out of 602 DDs, 522 DDs were young DDs and the remaining 80 were Elderly-DDs. In the group of young DDs, AKI was diagnosed in 276 (52.9%). In the group of elderly DDs, 46 (57.5%) donors had AKI. Finally, 598 cases of DDKT were classified into young-DDKT group and 111 cases were classified into elderly-DDKT. Within the young-DDKT group, 264 cases were classified as...
the non-AKI-Young-DDKT subgroup and 334 cases were classified in the AKI-Young-DDKT subgroup. Within the elderly-DDKT group, 47 cases were classified as the non-AKI-Elderly-DDKT subgroup, and 64 cases were classified as the AKI-Elderly-DDKT subgroup.

Clinical parameters and outcomes. We analyzed the medical records of both DDs and their corresponding DDKT recipients in the study population retrospectively. We collected baseline donor data including age, sex, body mass index (BMI) (kg/m²), history of diabetes mellitus (DM) and hypertension (HTN), cause of death due to cerebrovascular accident (CVA), and kidney donor profile index score (KDPI)37–39. Data of baseline and followed-up estimated glomerular filtration rate (eGFR) were calculated using the Chronic Kidney Disease Epidemiology Collaboration (CKD-EPI) equation to assess kidney function of DDs daily between the admission date and the date of KT. Baseline eGFR was the value obtained when DDs were admitted to the hospital for another cause and determined to brain death.

In addition, we collected the baseline recipient data: age, sex, BMI, dialysis type, dialysis vintage before KT, frequency of previous KTs, cause of end-stage renal disease, history of DM and HTN, cold ischemic time, number of human leukocyte antigen (HLA) mismatches, immunosuppressant type for induction, and maintenance and percentage of panel-reactive antibodies (PRAs).

BPAR was diagnosed by Banff classification40. DGF was defined as the need for dialysis within the first week after KT because of unrecovered allograft function41. Death-censored allograft survival rate was defined as the rate from KT to the return to dialysis excepting for allograft loss by patient death. Patient survival rate was defined as the rate from KT to patient death by any cause.

The primary outcome of this study was to investigate death-censored allograft survival rate of KTRs by young-DDs or elderly-DDs with AKI. Death-censored allograft survival was compared between the non-AKI-DDKT and AKI-DDKT groups with KT from young DDs or elderly DDs, respectively. The interaction between donor age and AKI was also analyzed. Secondary outcomes were to investigate the incidence of DGF, BPAR, the change of allograft function during the first year after KT, patient survival rate between young-DDKT and elderly-DDKT, and patient survival rate between non-AKI-DDKT and AKI-DDKT subgroups in young-DDKT and elderly-DDKT groups. Clinical outcomes were also analyzed according to AKI severity in each group analysis. The cause of allograft failure included biopsy-proven acute rejection (both T-cell mediated rejection and antibody-mediated rejection (AMR)), biopsy-proven chronic AMR (cAMR), chronic allograft dysfunction, biopsy-proven BK virus-associated nephropathy (BKVN), and biopsy-proven recurrent primary glomerulonephritis. Chronic allograft dysfunction was defined when allograft findings showed non-specific chronic tissue injury without evidence of rejection or when allograft biopsy was not done within one year of allograft failure and the allograft function showed a gradual deterioration several years before the allograft failure.

This study was approved by Institutional Review Boards of Seoul St. Mary’s Hospital (XC15RIMI0061K), Uijeongbu St. Mary’s Hospital (XC15RIMI0061U), Keimyung University School of Medicine, Dongsan Medical Center (2019-09-053), and Daejeon St. Mary’s Hospital (XC15RIMI0061K). The requirements for informed consent were waived by the Institutional Review Boards of above four centers because it was explained to all donor’s family and all recipients prior to KT that personal data associated with the donor and recipient’s clinical course was used and the information identifying the individual were protected. Our study did not contain any distinguishable personal identification information as a retrospective medical record study. Furthermore, all methods were performed in accordance with the relevant guidelines and regulations. The above four transplant centers never underwent transplantation with kidneys procured from prisoners, and this study did not include them as a study population.
Statistical analysis. Continuous variables with normal distributions were expressed as mean with standard deviation and analyzed by Student’s t-test. While those with non-normal distributions were expressed as median with interquartile range and analyzed by Mann-Whitney test. Categorical variables were expressed as number with percentage and analyzed by Chi-square test or Fisher’s exact test. Death-censored graft survival and patient survival rates were investigated by Kaplan-Meier curves and log-rank tests. All missing data were treated by censoring since the last follow-up date. Cox proportional hazards regression model was used to investigate an independent risk factor for allograft failure. Transplant years (1996~2005 vs. 2006~2010 vs. 2011~2017), transplant centers, recipient age, gender, donor gender, DGF, and BPAR were contained as confounding factors.

Data availability

All data generated or analyzed during this study are included in this published article and its Supplementary Information files. The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

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