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Findings from 4C-T Study demonstrate an increased cardiovascular burden in girls with end stage kidney disease and kidney transplantation

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Mortality in children with kidney failure is higher in girls than boys with cardiovascular complications representing the most common causes of death. Pulse wave velocity (PWV), a measure of vascular stiffness, predicts cardiovascular mortality in adults. Here, PWV in children with kidney failure undergoing kidney replacement therapy was investigated to determine sex differences and potential contributing factors. Two-hundred thirty-five children (80 girls; 34%) undergoing transplantation (150 pre-emptive, 85 with prior dialysis) having at least one PWV measurement pre- and/or post-transplantation from a prospective cohort were analyzed. Longitudinal analyses (median/maximum follow-up time of 6/9 years) were performed for PWV z-scores (PWVz) using linear mixed regression models and further stratified by the categories of time: pre-kidney replacement therapy and post-transplantation. PWVz significantly increased by 0.094 per year and was significantly higher in girls (PWVz +0.295) compared to boys, independent of the underlying kidney disease. During pre-kidney replacement therapy, an average estimated GFR decline of 4 ml/min/1.73 m² per year was associated with a PWVz increase of 0.16 in girls only. Higher diastolic blood pressure and low density

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lipoprotein were independently associated with higher PWVz during pre-kidney replacement therapy in both sexes. In girls post-transplantation, an estimated GFR decline of 4ml/min/1.73m² per year pre-kidney replacement therapy and a longer time (over 12 months) to transplantation were significantly associated with higher PWVz of 0.22 and of 0.57, respectively. PWVz increased further after transplantation and was positively associated with time on dialysis and diastolic blood pressure in both sexes. Thus, our findings demonstrate that girls with advanced chronic kidney disease are more susceptible to develop vascular stiffening compared to boys, this difference persist after transplantation and might contribute to higher mortality rates seen in girls with kidney failure.

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verall childhood mortality rates are declining.¹ In the general population, boys show higher mortality in most regions of the world,^{2,3} largely because of more accidents,¹ prematurity, respiratory distress during infancy,^{2,4} and sepsis occurring postpuberty.³ Inferior survival in girls is associated with poverty, marginalization, and a sociocultural preference for male offspring.² Mortality in children with end-

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functioning grafts, the proportion of cardiovascular mortality remains unchanged and is $\sim 20\%$ higher in girls.⁷ Cardiovascular events as the most common causes of death in children with end-stage kidney disease account for about one-third of deaths in children on dialysis and a quarter of

stage kidney disease is >30 times higher than that in the general

population.⁵ Data from the United States Renal Data System on

14,024 children on kidney replacement therapy (KRT) suggest a

higher mortality risk in girls (hazard ratio 1.36; 95% confidence

interval 1.25-1.50) because of their higher risk of cardiovascular

death.⁶ Despite declining overall mortality rates in children with

those undergoing transplantation.⁸ Data from the Australian and New Zealand Dialysis and Transplant Registry suggested an even higher mortality in pediatric kidney transplant recipients due to cardiovascular causes.9 The post-transplant mortality due to cardiovascular causes is higher than that related to nonfunctioning grafts.¹⁰

Early measures of arterial stiffness such as increased aortic pulse wave velocity (PWV) are highly predictive for cardiovascular events and mortality¹¹ and associated with a faster decline in estimated glomerular filtration rate (eGFR) in adults with chronic kidney disease (CKD).¹² Aortic PWV can be measured noninvasively and reproducibly in children.^{13,14} Higher PWV was demonstrated in children with CKD even after transplantation compared with their healthy peers.^{15–18}

Findings in adults indicate that the global survival advantage of females is lost in end-stage kidney disease,¹⁹ a phenomenon that is not sufficiently explained by disparities of access to transplantation due to higher levels of panel reactive antibodies in women²⁰ and pregnancy-induced incompatibility.²¹ In the pediatric population, girls are less likely to undergo preemptive transplantation^{6,22} and show poorer graft survival than boys, the latter being partly explained by receiving male donor organs.^{23,24} Our own data indicated a higher susceptibility of girls for cyclosporin A-associated hypertension,²⁵ which could contribute to poorer graft survival and increased cardiovascular mortality.

Here we aimed to study the course of arterial stiffness in children with end-stage kidney disease who underwent transplantation either preemptively or after prior dialysis to uncover potential sex differences.

METHODS

151 Study design, setting, and participants 152

The 4C-T (Cardiovascular Comorbidity in Children with Chronic 153 Kidney Disease – Transplantation) substudy is part of the 4C study,²⁶ a 154 prospective observational study. Seven hundred four pediatric patients 155 with CKD (age 6–17 years) with an eGFR below 60 ml/min per 1.73 m² 156 not yet receiving KRT were enrolled between 2009 and 2011. Ethical 157 aspects and details of the data acquisition were described previously.²⁶ 158 The median follow-up time was 6 years, with a maximum of 9 years. 159

160 Data sources/measurements

161 PWV was assessed annually using the oscillometric Vicorder device 162 Q12 (SMT medical, Würzburg, Germany), as described previously.^{13,14}

Every 6 months blood and urine samples, anthropometrics, casual Q13 blood pressure (BP), and medical history updates were obtained per standardized protocol. Laboratory measurements were performed centrally. eGFR was calculated using the Schwartz formula.²

Variables

Sex- and height-adjusted standardized scores (z scores) were calculated for PWV¹³ as the primary end point.

The following parameters were considered as covariates: eGFR decline, body mass index, BP, lipids, hemoglobin, sodium, potassium, calcium, phosphorus, bicarbonate, parathyroid hormone, uric acid, and urea. Kidney diseases were categorized as congenital anomalies of the kidney and urinary tract (CAKUT) or non-CAKUT. Supplementary Table S1 provides a more granular classification of primary renal diseases. Antihypertensive and immunosuppressive Q14 medications (including trough levels) were recorded. Systolic and diastolic BP (sex-, age-, and height-adjusted)²⁸ z scores as well as height and body mass index (sex- and age-adjusted)²⁹ z scores were calculated.

As ambulatory BP measurements were available only in a subgroup of patients, we provide data for the correlation between the BP and the ambulatory BP measurement in Supplementary Table S2.

Time variable

Time (in years) was assessed by the following variables: time since inclusion, time pre-KRT (time since inclusion but before KRT start), time post-transplantation (time since transplantation), time from eGFR \leq 30 ml/min per 1.73 m² to transplantation, and time on dialysis (see Supplementary Figure S1 for more details).

Healthy control cohort

Longitudinal PWV measurements in 307 healthy children (girls, n =145) from the REBIRTH active school study were used to assess possible sex differences in the physiological development of PWV. The study investigated cardiovascular parameters in healthy children during a school-based physical activity program³⁰ with 2 repetitive PWV measurements with an interval of 12.7 \pm 3.3 months between 2017 and 2018.

Analysis steps

The analyses for PWV z scores (PWVz) were performed in 3 analysis Q15 steps: (i) all data comprising the whole observation time and then divided into 2 separate analyses according to transplantation: (ii) "pre-KRT" and (iii) "post-transplantation" (Figure 1).

Step 1: all data. We included patients with at least 1 visit during the observation period representing the complete observation time (n = 235, Figure 2). This includes data before KRT, on dialysis, and after transplantation. A spline regression was fitted to the data to visualize the course of PWVz and the sex difference on the development of PWVz. Linear mixed regression models (mixed models) for PWVz were performed as follows: (i) adjusted for time since Q16 inclusion and kidney disease category to understand the development of PWVz over time; (ii) adjusted for sex, time since inclusion, and kidney disease category to understand the development of PWVz over time depending on sex; and (iii) adjusted for the interaction term |sex*KRT modality*time since inclusion| and kidney disease category to understand the development of PWVz stratified for each treatment modality depending on sex (Figure 1).

Step 2: pre-KRT data. All data before KRT start were included to study the development of PWVz during CKD progression. Patients with at least 1 visit pre-KRT were included (n = 230, Figure 2)

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Figure 1 Process flow of the analyses for pulse wave velocity *z* scores (PWVz). Step 1, as shown in the box, describes the analyses for the complete observation period including pre-kidney replacement therapy (KRT), on dialysis, and post-transplantation. Step 2, as marked in the light gray area, describes the analyses flow for PWVz during pre-KRT. Step 3, as marked in the dark gray area, describes the analyses flow for PWVz post-transplantation. CAKUT, congenital anomalies of kidney and urinary tract; CKD, chronic kidney disease; eGFR, estimated glomerular filtration rate.

in the mixed model for PWVz adjusted for pre-KRT time, sex, interaction term of |pre-KRT time*sex|, and kidney disease category to understand potential sex differences on the development of PWVz before KRT. This model was then set as the basic model for "pre-KRT" (Figure 1).

To further investigate the possible influencing factors on sex differences in the development of PWVz before KRT, we screened covariates that are assumed relevant for PWVz using the above predefined basic model. We included patients with at least 2 visits pre-KRT (n = 158) to assess the pre-KRT eGFR decline as one of potential covariates. Covariates showing significant associations with PWVz (P < 0.05) and/or eliminating the association (P > 0.05) between PWVz and the interaction term |pre-KRT time*sex| were included in the backward selection. Covariates that are highly correlated with each other (BP values and lipid levels) were grouped. If \geq 2 covariates from the same group were eligible, the one with the better model fit (lower Akaike information criterion) was selected.

To assess CKD progression, eGFR decline was calculated. Delta eGFR (Δ eGFR) for each patient *i* at visit *v* was calculated as the difference between eGFR at visit *v* and the previous visit *v* - 1 divided by the time *T* interval (in years) between both visits: Δ eGFR = [eGFR_{*iv*} - eGFR_{*i*(*v*-1)}] / [*T_{iv}* - *T_{i*(*v*-1)}] (Supplementary Figure S2). In the case of a missing eGFR value between 2 visits, Δ eGFR was interpolated.



Figure 2 | Inclusion flowchart of the study population. CKD, chronic kidney disease; KRT, kidney replacement therapy; PWV, pulse wave
velocity; Tx, transplantation.

Table 1 | Patients' characteristics at inclusion, at the last visit pre-KRT, and at 1 yr post-transplantation

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Disease and transplant modality		Girls $(n = 80)$ Boys $(n = 7)$			
Underlying disease					
CAKUT		33 (41)		98 (63)	
Non-CAKUT		47 (59)		57 (37)	
Transplantation					
Preemptive		48 (60)		102 (66)	
With prior dialysis		32 (40)		53 (34)	
		Girls ($n = 78$)		Boys ($n = 152$)	
At inclusion	n	Median (IQR)	n	Median (IQR)	
Age (yr)	78	12.2 (9.24 to 14.3)	152	11.6 (8.98 to 14.2)	
Height (cm)	78	139 (126 to 151)	152	138 (124 to 155)	
Height z score	78	-1.45 (-2.09 to -0.32)	152	-1.16 (-2.03 to -0.19	
BMI (kg/m ²)	78	16.6 (15.3 to 18.4)	152	17.5 (15.7 to 19.9)	
BMI z score	78	-0.62 (-1.08 to 0.18)	152	-0.08 (-0.79 to 0.84)	
Systolic BP (mm Hg) ^a	78	110 (100 to 114)	152	114 (106 to 123)	
Systolic BP z score	78	0.4 (-0.08 to 1.29)	152	1.09 (0.38 to 1.74)	
Diastolic BP (mm Hg)	78	65.5 (60.0 to 78.0)	152	69.0 (62.0 to 78.0)	
Diastolic BP z score ^a	78	0.41 (-0.16 to 1.30)	152	0.72 (0.18 to 1.31)	
eGFR (ml/min per 1.73 m ²)	78	21.5 (15.4 to 30.7)	152	19.3 (14.6 to 28.0)	
		Girls ($n = 78$)		Boys ($n = 152$)	
At last visit pre-KRT	n	Median (IQR)	n	Median (IQR)	
Age (vr)	78	13.5 (11.1 to 15.8)	152	13.7 (11.9 to 15.7)	
Time since inclusion (vr)	78	1.2 (0 to 3.1)	152	1.9 (0 to 3.1)	
Time from the last visit during	78	0.5(0.3 to 0.8)	152	0.6 (0.2 to 0.8)	
CKD to KRT start (vr)	70				
Height (cm) ^a	78	149 (135 to 157)	152	152 (137 to 165)	
Height z score	78	-1.23 (-2.12 to -0.43)	152	-0.96 (-2.14 to -0.22	
$BMI (kg/m^2)$	78	18.0 (16.1 to 19.1)	152	18.1 (16.6 to 20.5)	
BMI z score	78	-0.57 (-1.21 to 0.28)	152	-0.25 (-0.99 to 0.60)	
Systolic BP (mm Hg) ^a	78	112 (105 to 119)	152	120 (109 to 128)	
Systolic BP z score ^a	78	0.55(-0.12 to 1.39)	152	1.09 (0.26 to 1.90)	
Diastolic BP (mm Hg)	78	70.0 (63.0 to 80.0)	152	70.0 (65.0 to 80.0)	
Diastolic BP z score	78	0.46 (-0.01 to 1.48)	152	0.81 (0.13 to 1.43)	
eGER (ml/min per 1 73 m ²)	76	13.9 (11.5 to 16.4)	1/2	13.3 (10.9 to 17.0)	
Cholostorol (mg/dl)	70	19.5 (11.5 to 10.4)	150	160 (142 to 200)	
HDL (mg/dl) ^a	77	54.0 (44.0 to 64.0)	150	109 (142 to 200) 13.0 (26.5 to 53.0)	
IDL (mg/dl)	77	90.0 (91.0 to 123)	150	98.6 (60.5 to 122.3)	
Homoglobin (g/dl) ^a	77	10.6 (0.80 to 11.8)	145	11.2 (10.3 to 12.1)	
Ferritin (ug/l)	77	10.0 (9.80 to 11.8) 99.8 (52.3 to 181)	145	11.2 (10.3 to 12.1) 101 5 (51.0 to 204)	
Sodium (mmol/l) ^a	72	140 (136 to 142)	1/0	$140(137 \pm 0.142)$	
Potacsium (mmol/l)	77	140 (100 to 142)	145	4 50 (4 10 to 4 99)	
Calcium (mmol/l) ^a	77	2 30 (2 15 to 2 49)	1/7		
Phosphorus (mmol/l)	77	1.61 (2.13 to 2.40)	147	2.30 (2.20 to 2.32) 1.67 (1.51 +o. 1.07)	
Ricarbonate (mmol/l)	7/	21 0 (2.71 (0 2.72) 21 0 (10 0 to 22 0)	1/2	21 0 (100 +o 22 0)	
Parathyroid hormone (pmol/l)	75	16 3 (7 50 to 29 0)	170	21.0(19.0(0,23.0))	
Liric acid (mg/dl) ^a	75	6.12 (5.30 to 7.21)	150	24.7 (13.3 t0 40.9) 7 17 (6 10 +0 0 21)	
Lirea (mg/dl) ^a	76	60.7 (43.2 to 70.5)	1/10	73 0 (56 5 to 116)	
Antihynertensive use	70	54 of 78 (69)	140	104 of 152 (68)	
RAAS antagonists		37 (47)		54 (36)	
CCB		29 (37)		57 (38)	
β-Blockers		8 (10)		28 (18)	
Perinheral <i>a</i> -blockers		5 (6)		8 (5)	
		5 (0) 0 (0)		0 (<i>J</i>) 1 (1)	
		6 (9)		12 (0)	
Thiazide diuretics		4 (5)		2 (1)	
		Girls (n - 71)		Boys(n - 127)	
Transplantation		Median (IOP)		Modian (IOP)	
Age at transplantation (yr) Time from $aGEP \leq 20$ ml/min	71	15.2 (12.3 to 17.0)	128	14.7 (12.6 to 16.3	
nine non eur $h \ge 50$ mi/min	/1	2.3 (1.5 to 3.8)	128	2.5 (1.3 to 4.0)	
Time since inclusion to the function	71	24/14 + 44	100	76 /1 4 +- 4 1	
	/1		120	2.0 (1.4 to 4.1)	
Time on dialusis (ur)			53	1.2 (U.9 to 1.6)	
Time on dialysis (yr)	32	42 (50)		02 (CE)	
Time on dialysis (yr) Transplanted >1 yr after $aCER \leq 30$ m//win zer 1.72	32	42 (59)		82 (65)	

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Table 1 (Continued)

		Girls ($n = 71$)		Boys $(n = 127)$		
Transplantation	n	Median (IQR)	n Median (IQR)			
Transplanted ≤ 1 yr after eGFR ≤ 30 ml/min per 1.73 m ² or after dialysis start		29 (41)		45 (35)		
		Girls ($n = 53$)	Boys (n = 103)			
At 1 yr post-transplantation	n	Median (IQR)	n	Median (IQR)		
Age (yr)	53	15.7 (13.3 to 18.2)	103	15.3 (13.1 to 17.3)		
Time since inclusion (yr)	53	3.1 (2.2 to 5.1)	103	3.2 (2.1 to 5.0)		
Time post-transplantation (yr)	53	0.9 (0.7 to 1.1)	103	0.9 (0.8 to 1.2)		
Height (cm) ^a	53	153 (143 to 160)	103	160 (147 to 168)		
Height z score	53	-1.00 (-1.98 to -0.03)	103	-1.02 (-2.13 to -0.37		
BMI (kg/m ²)	53	20.3 (17.6 to 22.5)	103	21.1 (18.0 to 23.5)		
BMI z score	53	-0.12 (-0.93 to 0.95)	103	0.37 (-0.50 to 1.11)		
Systolic BP (mm Hg) ^a	53	115 (107 to 121)	103	120 (112 to 130)		
Systolic BP z score	53	0.73 (-0.02 to 1.51)	103	1.04 (0.21 to 1.76)		
Diastolic BP (mm Hg)	53	71 (63.0 to 77)	103	72 (65 to 79)		
Diastolic BP z score	53	0.54 (-0.14 to 1.14)	103	0.71 (0.13 to 1.33)		
eGFR (ml/min per 1.73 m ²) ^a	50	68.4 (50.3 to 82.3)	100	59.4 (46.7 to 74.0)		
Tacrolimus trough level (μg/l)	41	5.00 (4.00 to 7.00)	82	6.00 (5.00 to 8.00)		
Cyclosporin A trough level (µg/l)	11	100 (67.0 to 129)	16	104 (85.5 to 124)		
Steroid dosage (mg/d)	46	5.00 (3.00 to 5.00)	79	5.00 (4.00 to 7.50)		
Antihypertensive use		32 of 53 (60)	64 of 103 (62)			
RAAS antagonists		9 (17)	16 (15)			
CCB		25 (47)	47 (46)			
β-Blockers		7 (13)	29 (28)			
Peripheral α-blockers	0		4 (4)			
Central α-blockers	1 (2)		1 (1)			
Loop diuretics	1 (2)		6 (6)			
Thiazide diuretics	0		4 (4)			
Immunosuppresive use	53 of 53 (100)		103 of 103 (100)			
Steroid	46 (87)		79 (77)			
CNI		52 (98)		101 (98)		
MMF		47 (87)	82 (80)			
mTOR	6 (11)			13 (13)		

BMI, body mass index; BP, blood pressure; CAKUT, congenital anomalies of kidney and urinary tract; CCB, calcium channel blocker; CKD, chronic kidney disease;
CNI, calcineurin inhibitor; eGFR, estimated glomerular filtration rate; HDL, high-density lipoprotein; IQR, interquartile range; KRT, kidney replacement therapy; LDL, low-density lipoprotein; MMF, mycophenolate mofetil; mTOR, mechanistic target of rapamycin; RAAS, renin-angiotensin-aldosterone system.

/5 ^a*P* value of < 0.05.

476 Data are *n* (%) unless otherwise noted.

Step 3: post-transplantation data. Patients with at least 1 visit post-transplantation were included (n = 199, Figure 2). A mixed model for PWVz adjusted for post-transplantation time, sex, interaction term |post-transplantation time*sex|, kidney disease category, and time on dialysis was performed to understand the sex differences on the development of PWVz post-transplantation (Figure 1).

For further investigation, patients with visits at pre-KRT and post-transplantation were included to assess the eGFR slope pre-KRT (n = 195, Figure 2). Covariates were then screened using the above predefined basic model to identify the contributing factors. Cova-riates showing significant associations with PWVz (P < 0.05) and/or eliminating the association (P > 0.05) between PWVz and the var-iable "sex" were included in the backward selection. Similar to the analysis for pre-KRT, if ≥ 2 covariates were eligible but highly correlated with each other, the one with the lower Akaike infor-mation criterion was included.

We calculated individual eGFR slopes using the eGFR measurements pre-KRT to reflect the pace of the functional decline. The eGFR slope was computed as the function (regression coefficient, *B*) of the fixed effect of time pre-KRT (in years) for each patient *i* from linear regression of eGFR: eGFR_i = Intercept_i + B_i (time pre -KRT in years) (Supplementary Figure S2). eGFR slopes were defined as 0 in 20 cases undergoing dialysis and 17 who underwent preemptive transplantation. In all cases, final CKD stages were reached and KRT was initiated before a second measurement could be performed. Three children with only 1 eGFR measurement pre-KRT >12 months before the initiation of KRT were excluded. A sensitivity analysis for the final model including individual eGFR slopes computed from a single mixed model for eGFR pre-KRT was performed. To assess the time to transplantation, patients were grouped into shorter (\leq 12 months) or longer (>12 months) time to transplantation calculated as time to transplantation since eGFR dropped to \leq 30 ml/min per 1.73 m² (preemptive) or since dialysis start (after prior dialysis).

Additional analyses for PWVz by kidney diseases

As the underlying kidney diseases differ between sexes, additional analyses had to be performed. Patients with at least 1 visit were included (n = 235). Two mixed models for PWVz were performed:

(i) a mixed model adjusted for the interaction term |time since inclusion*kidney disease category| to understand whether the PWVz development differs between patients with CAKUT and those without CAKUT. The corrected means and 95% confidence intervals of PWVz adjusted for the respective models were calculated for CAKUT and non-CAKUT groups; and

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Figure 3 | Analyses of pulse wave velocity z scores (PWVz) over the complete observation period. (a) Spline regression fit and 95% confidence interval (upper panel) and mixed model for PWVz adjusted for time since inclusion and kidney disease (the table in the lower panel). (b) Spline regression fit and 95% confidence interval differentiated by sex, showing a red line for girls and a blue line for boys (upper 933 panel), and mixed model for PWVz adjusted for time since inclusion, kidney disease, and sex (the table in the lower panel). β , regression coefficient; CAKUT, congenital anomalies of kidney and urinary tract; KRT, kidney replacement therapy; Ref., reference; SE, standard error.

(ii) a mixed model adjusted for the interaction term |time since inclusion*sex and kidney disease category (girls-CAKUT, girlsnon-CAKUT, boys-CAKUT, and boys-non-CAKUT)| to understand how sex influences the PWVz course in each kidney disease category.

General statistical analysis

Data are given as median and interquartile range or absolute and relative frequencies. t tests were performed to test differences between sexes. Complete data analyses were performed, and covariates with missing >10% were not included in the covariate



Figure 4 | Regression fit of pulse wave velocity z score (PWVz) and 95% confidence interval in the cohort of healthy children differentiated by sex, showing a red line for girls and a blue line for boys, adjusted for the mixed model for PWVz as given in the table below the graph. β , regression coefficient; Ref., reference; SE, standard error.

selection. Supplementary Figure S3 provides the number of observations over time. The pattern of missing data accounting for the variables included in the final models is provided in Supplementary Table S3A and B. Spline regression and mixed models were performed as described above. In the mixed models, patient ID and center were included as random effects to model the between-subject variation and time since inclusion as a repeated effect to model within-subject variation.³¹ Statistical analysis was performed using SAS 9.4 (SAS Institute, Cary, NC). This manuscript was written according to the Strengthening the Reporting of Observational Studies in Epidemiology guidelines.³

RESULTS

Patient characteristics

Of 704 children, 338 underwent KRT. Four patients without any PWV measurements and 99 patients only receiving dialysis without subsequent transplantation were excluded. Two hundred thirty-five patients (girls, n = 80) undergoing kidney transplantation (preemptive, n = 150; with prior dialysis, n = 85) were included. Of those, 196 had observations before and after transplantation, 36 only before transplantation, and 3 only after transplantation (Figure 2).

eGFR at inclusion and at the last visit pre-KRT, age at inclusion and at transplantation, time from eGFR \leq 30 ml/ min per 1.73 m² to transplantation, and time on dialysis did not differ between sexes. At the last visit pre-KRT, girls showed significantly lower height, systolic BP, hemoglobin, sodium, calcium, uric acid, urea, and higher high-density lipoprotein (HDL) than did boys. Table 1 summarizes patient Q19 characteristics at study inclusion, pre-KRT, at transplantation, and 1-year post-transplantation.

PWV and the effect of time and sex

Figure 3a shows the spline regression slope fitted for PWVz over the complete observation period. PWVz increased by 0.095 per year since inclusion (P < 0.0001),

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Mixed model for PWVz with interaction term between KRT modality, sex, and time since inclusion (yr)

(patients, n = 235; observations, n = 1368)

Effect	β	SE	Р
Intercept	0.076	0.10	0.46
Effect of time since inclusion in:			
Girls pre-KRT (yr)	0.17	0.057	0.004
Girls on dialysis (yr)	0.20	0.073	0.006
Girls post-Tx (yr)	0.13	0.024	<0.0001
Boys pre-KRT (yr)	0.022	0.038	0.56
Boys on dialysis (yr)	0.084	0.053	0.11
Boys post-Tx (yr)	0.075	0.019	<0.0001
Non-CAKUT (ref.: CAKUT)	0.015	0.14	0.92



Figure 5 | Mixed model for pulse wave velocity z scores (PWVz) adjusted for the interaction term |time since inclusion*sex*kidney replacement therapy (KRT) modality (pre-KRT/dialysis/ transplantation [Tx]). The red shaded rows highlight the effect of

time pre-KRT and dialysis in girls. The blue shaded rows highlight the effect of time pre-KRT and dialysis in boys (the table in the upper panel) and the different slopes of estimates according to the given linear mixed model differentiated by sex and KRT modality. The yellow area shows the time pre-Tx (including pre-KRT and dialysis), and the green area shows the time post-Tx. The pink lines show the PWVz slopes for girls and blue lines for boys. β , regression coefficient; CAKUT, congenital anomalies of kidney and urinary tract; Ref., reference; SE, standard error.

independent of kidney disease (P = 0.64). Figure 3b visualizes the sex-adjusted PWVz. The mixed model demonstrated that PWVz was 0.295 higher in girls (P =0.045) than in boys.

We compared our study population with a cohort of 706 707 healthy children with comparable height. Our cohort of 708 healthy children demonstrated considerably lower PWVz (median -0.28; interquartile range -0.84 to 0.39) at study 709 710 inclusion (Supplementary Table S4). PWVz in healthy children did not increase with time (PWVz -0.048/yr; P = 0.27) and did not differ between girls and boys (Figure 4). 712

As the spline regression of PWVz indicated an interaction 713 between sex and time before transplantation, a mixed model 714 715 for PWVz adjusted for the interaction term time since inclusion*sex*KRT modality| was performed. Girls showed a 716 PWVz increase of 0.17 pre-KRT (P = 0.004) and of 0.20 717 during dialysis (P = 0.006) per year since inclusion. These 718 719 time effects during pre-KRT and dialysis were not present in 720 boys (Figure 5, upper panel). The lower panel of Figure 5 721 illustrates the different slopes of PWVz for girls and boys, depending on KRT modality and highlights the greater 722

Table 2 | Mixed models for PWVz "pre-KRT": basic model adjusted for pre-KRT time, interaction term [pre-KRT time*sex], sex, and kidney disease category; final model adjusted for the covariates included in the basic model, delta eGFR, interaction term |delta eGFR*sex|, diastolic BP z score, and LDL

	Ba: (A	sic mod IC: 222	lel 1)	Final model (AIC: 1340)				
	230 650 c) patien bserva	its, tions	156 patients, 410 observations				
Variable	β	SE	Р	β	SE	Р		
Intercept	0.14	0.14	0.33	-0.89	0.25	0.0005		
Pre-KRT time	0.027	0.039	0.48	0.0003	0.048	0.99		
Pre-KRT time*girls	0.15	0.072	0.039	0.13	0.093	0.18		
Pre-KRT time*boys		Ref.			Ref.			
Girls (ref.: boys)	-0.018	0.22	0.94	0.028	0.28	0.92		
Non-CAKUT	0.042	0.20	0.83	-0.11	0.20	0.59		
Delta eGFR		_		0.002	0.012	0.89		
Delta eGFR/ year*girls		_		-0.040 ^ª	0.017	0.017		
Delta eGFR/ year*boys		—			Ref.			
Diastolic BP z score		_		0.47	0.064	< 0.0001		
I DI		_		0.007	0.002	0 0001		

AIC, Akaike information criterion; β , regression coefficient; BP, blood pressure; CAKUT, congenital anomalies of kidney and urinary tract; eGFR, estimated glomerular filtration rate; KRT, kidney replacement therapy; LDL, low-density lipoprotein; PWVz, pulse wave velocity z score; Ref., reference; SE, standard error.

^aA delta eGFR (a decline of eGFR) of -1 ml/min per 1.73 m² per year was associated with an increase of 0.04 PWVz in girls compared with boys. 029 Bold data indicate xxx.

progression of PWVz in girls pretransplantation. This indicated the need of separating the analyses according to KRT, that is, "pre-KRT" and "post-transplantation." A separate analysis for dialysis was not possible because of the low number of observations (only 25 of 62 patients had ≥ 2 visits).

The effect of sex on PWV pre-KRT

We analyzed 230 patients. A higher PWVz increase of 0.15 per year was shown in girls than in boys (P = 0.039; Table 2, basic model). One hundred fifty-eight patients were included in the covariate screening for the final model (Supplementary Table S5).

The final model revealed that Δ eGFR was a strong predictor for PWVz in girls. An eGFR decline of -4 ml/min per 1.73 m² per year pre-KRT was associated with a higher PWVz of 0.16 in girls (P = 0.017) compared with boys. A higher diastolic BP z score and higher low-density lipoprotein (LDL) were associated with a higher PWVz in both sexes (Table 2, final model). Supplementary Figure S4 illustrates the sex difference on the effect estimate of the influencing factors on PWVz as a result from the respective model stratified by sex (Supplementary Table S6).

The effect of sex on PWV post-transplantation

We analyzed 199 patients. PWVz for girls was 0.44 higher than that for boys (P = 0.02). PWVz increased by 0.12 per year post-transplantation (P = 0.003) and by 0.25 per year on

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Table 3 | Mixed models for PWVz "post-transplantation": basic model adjusted for post-transplantation time, sex, kidney disease category, and time on dialysis; prefinal model adjusted for covariates included in the basic model, PWVz at the last visit during pre-KRT, diastolic BP z score, and cholesterol; and final model adjusted for covariates included in the prefinal model, eGFR slope, interaction term |eGFR slope*sex|, and interaction term |time to transplantation*sex|

	Basic model (AIC: 1900)		Prefinal model (AIC: 1620) 190 patients, 540 observations			Final model (AIC: 1606.4) 188 patients, 533 observations			
Variable	199 patients, 613 observations								
	β	SE	Р	β	SE	Р	β	SE	Р
Intercept	0.0001	0.14	0.99	-0.76	0.28	0.007	-0.85	0.31	0.006
Post-transplantation time	0.12	0.04	0.003	0.13	0.03	< 0.0001	0.14	0.033	< 0.000
Post-transplantation time*girls	-0.008	0.06	0.89		_			_	
Post-transplantation time*boys		Ref.							
Girls (ref.: boys)	0.44	0.19	0.024	0.38	0.14	0.010	-0.048	0.27	0.86
Non-CAKUT (ref.: CAKUT)	-0.02	0.15	0.92	-0.011	0.14	0.94	0.032	0.14	0.82
Time on dialysis	0.25	0.09	0.006	0.19	0.09	0.032	0.19	0.085	0.024
PWVz at the last visit pre-KRT		_		0.22	0.04	< 0.0001	0.20	0.040	< 0.000
Diastolic BP z score		_		0.36	0.06	< 0.0001	0.36	0.055	< 0.000
Cholesterol		_		0.003	0.001	0.059	0.002	0.001	0.10
eGFR slope pre-KRT		—			-		0.012	0.014	0.42
eGFR slope pre-KRT*girls		—			_		-0.054ª	0.026	0.039
eGFR slope pre-KRT*boys		—			—			Ref.	
Girls–longer time to transplantation ^b		—			—		0.57	0.24	0.017
Girls-shorter time to transplantation ^b		—			—			Ref.	
Boys–longer time to transplantation ^b		—			—		0.29	0.18	0.10
Boys–shorter time to transplantation ^b								Ref.	

AIC, Akaike information criterion; β, regression coefficient; BP, blood pressure; CAKUT, congenital anomalies of kidney and urinary tract; eGFR, estimated glomerular filtration
rate; KRT, kidney replacement therapy; PWVz, pulse wave velocity z score; Ref., reference; SE, standard error.

^aAn eGFR slope of -1 ml/min per 1.73 m² per year (a declining slope) at pre-KRT was associated with a higher PWVz of 0.054 in girls after transplantation as compared with boys.

808dialysis (P = 0.006) for both sexes. An interaction between809time and sex was not detected (Table 3, basic model;810Supplementary Table S7 shows the basic model separated by811the transplantation type, i.e., preemptive and after prior812dialysis).

We further analyzed 195 patients and screened for po-813 tential covariates using the basic model (Supplementary 814 815 Table S8). PWVz increased by 0.13 per year posttransplantation (P < 0.0001) and by 0.19 per year on dial-816 ysis (P = 0.03). A 1-unit higher PWVz at the last pre-KRT 817 visit was associated with a 0.22 increase in post-818 819 transplantation PWVz (P < 0.0001); furthermore, a 1-unit 820 increase in diastolic BP z score was associated with a posttransplantation PWVz increase of 0.36 (P < 0.0001). 821 822 Importantly, the association of female sex and higher PWVz persisted (P = 0.01; Table 3, prefinal model). 823

The previously observed effect of the eGFR decline on 824 PWVz before KRT in girls was further elucidated by intro-825 ducing 2 interaction terms: |sex*eGFR slope| and |sex*time to 826 827 transplantation (Table 3, final model). Although the global sex effect disappeared (P = 0.86), the eGFR slope pre-KRT 828 and a longer time to transplantation (>12 months after 829 eGFR dropped to or below 30 ml/min per 1.73 m² or after 830 dialysis start) revealed significant associations with higher 831 832 PWVz in girls. An eGFR decline, for example, of -4 ml/min per 1.73 m² per year pre-KRT was associated with a PWVz 833 increase of 0.22 after transplantation in girls (P = 0.039). A 834

longer time to transplantation (>12 months) was associated with a higher PWVz of 0.57 in girls (P = 0.017; Table 3, final model). The associations of other contributing factors with PWVz persisted (Table 3, prefinal and final models). A sensitivity analysis for the PWVz post-transplantation model including individual eGFR slopes computed from the single mixed model is provided in Supplementary Table S9 showing similar findings. A description of PWVz and absolute PWVz at different time points is given in Supplementary Table S10.

Sex differences are independent of the underlying disease

As expected, the distribution of the underlying kidney disease differed between the sexes with a higher proportion of CAKUT in boys (63%) and non-CAKUT in girls (59%). We explored the potential effect of the difference in disease distribution. An additional model showed that PWVz did not differ between the categories of underlying diseases, as demonstrated by the corrected means showing no differences between the categories of CAKUT (mean 0.39; 95% confidence interval 0.20-0.58) and non-CAKUT (mean 0.40; 95% Q20 confidence interval 0.19–0.60; Supplementary Figure S5A). We then explored the potential differences between the combinations of the sex and kidney disease category (girls-CAKUT, girls-non-CAKUT, boys-CAKUT, and boys-non-CAKUT). The mixed model adjusted for the interaction term time since inclusion*category for sex and kidney disease also showed that the development of PWVz did not differ between

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the 4 categories (Supplementary Figure S5B). This demonstrated that the higher PWVz in girls was independent of the kidney disease distribution.

DISCUSSION

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Our study characterized the evolution of vascular stiffness in 896 897 girls and boys with progressing CKD and subsequent trans-898 plantation. Girls with advanced CKD showed more pro-899 nounced arterial stiffening than did boys. This is in contrast to the physiological development as demonstrated in a cohort 900 of healthy children. The faster progression of arterial stiff-901 902 ening in girls occurred before transplantation, reflecting a 903 higher vulnerability of girls' vascular system toward the magnitude as well as the duration of the exposure to an 904 905 impaired renal function. Our key finding is that time acts 906 differently on the cardiovascular burden between boys and 907 girls during CKD. Importantly, this was independent of the 908 underlying kidney disease.

A higher susceptibility of females with CKD to develop 909 910 arterial stiffness in conjunction with renal disease progression has not been described to date. Studies in adults so far have 911 shown more severe arterial stiffness in women than in men³³ 912 913 and an association between arterial stiffness and eGFR decline without sex differences.³⁴ A tendency toward faster decline in 914 renal function in girls than in boys has been demonstrated 915 especially before puberty.²² Although in the general popula-916 917 Q21 tion and in CKD eGFR declines at a faster rate in males, a 918 meta-analysis of a large number of postmenopausal women suggested a faster decline in women.^{35–38} This could be 919 explained by the absence of estrogens' nephroprotective ef-920 fect.^{39,40} In light of lower levels of sex hormones during pu-921 922 berty in children with CKD contributing to their well-known delayed puberty,⁴¹ one could speculate that girls in our cohort 923 are less protected by estrogen and renal function decline 924 925 should be similar between the sexes. However, this is not the case, suggesting sex hormones alone do not explain the dif-926 927 ferences seen.

Although the physiological development of PWV over time 928 did not differ between girls and boys, we did see a significant 929 difference in children with CKD. This difference occurred 930 before transplantation and was associated with a longer time 931 932 to transplantation, indicating that girls are particularly 933 vulnerable during the final stages of CKD progression. 934 Importantly, we can show that sex difference in PWV exists irrespective of the underlying kidney diseases, which are 935 differently distributed between girls and boys. 936

937 Factors associated with bone and mineral metabolism 938 (parathyroid hormone, vitamin D, and calcium, phosphorus, and their product itself)⁴²⁻⁴⁴ are known contributors to the 939 arterial stiffness increase during CKD progression. None of 940 941 these factors explained the observed sex differences in PWV. 942 In fact, serum calcium and parathyroid hormone were higher 943 in boys. This, however, does not exclude the possibility that 944 girls may develop a more pronounced PWV increase for a given calcium or parathyroid hormone level. Another 945 946 important factor in mineral metabolism is fibroblast growth factor 23,⁴⁵ which was measured only at baseline in our ^{Q2} cohort. Postmenopausal women without estrogen substitution show higher fibroblast growth factor 23 levels than do women with estrogen substitution or men.⁴⁶ It is conceivable that the fibroblast growth factor 23 pathway is more active in prepubertal girls or girls with an altered estrogen metabolism because of their uremic state. Similarly, osteoprotegerin, a cytokine that regulates bone resorption, is associated with cardiovascular events in patients with CKD and on hemodialysis.^{47–49} In the CRIC study, higher osteoprotegerin levels ^{Q23} were associated with increased PWV and females had ~10% higher osteoprotegerin levels than did male patients with CKD.⁵⁰

Cholesterol and its subclasses LDL and HDL are known to influence PWV and predict cardiovascular risk.51,52 In the general population, LDL is associated with an increased risk and HDL with a decreased risk. In children with CKD,^{53,54} increased HDL promotes endothelial dysfunction and is associated with vascular damage, possibly because of a uremia-associated altered HDL functionality.53 Our data showed higher HDL levels in girls, another factor that could explain the accelerated vascular damage in girls. This assumption is further supported by the performance of HDL in the model building process. HDL being in the causal pathway between the progression of CKD and PWV could explain why the introduction of HDL together with the interaction term "\DeGFR/year*girls" did not reveal a significant result. LDL, however, showed an association with higher PWV in both sexes, but did not explain the sex differences. Our data highlight the importance of both cholesterol subclasses in pediatric patients with CKD and the need for early preventive strategies, especially because in adults with CKD and on dialysis, LDL lowering with atorvastatin and ezetimibe was successful in reducing atherosclerotic events.⁵⁵

PWV increased further after transplantation. In previous 981 studies, PWV tended to stabilize or slightly decrease during 982 the first year after kidney transplantation^{18,56-58} but longer 983 observations in adults revealed an increase in PWV.⁵⁹ The 984 increase in PWV post-transplantation with time likely reflects 985 an ongoing damage even after transplantation in addition to 986 the "preexisting" burden from the time pre-KRT. This implies 987 the need of a better cardiovascular monitoring and cardio-988 vascular disease prevention, especially before the onset of 989 KRT. Our data also highlight the clinical importance of an 990 even faster access for girls to transplantation, especially in 991 light of studies showing a slower access to preemptive 992 transplantation.^{6,22} A longer time on dialysis was associated 993 with increased PWV, which is in line with our previous 994 finding showing the advantage of preemptive transplantation 995 compared with dialysis.¹⁸ The observed association between 996 higher BP and higher PWV in both sexes is in accordance 997 with previous findings in the general pediatric popula-998 tion,^{60,61} in various patient groups,^{15,17,62} and after kidney 999 transplantation.^{18,63} Uncontrolled or untreated hypertension 1000 is present in 30% to 40% of pediatric kidney allograft re-1001 cipients.^{25,63} Notably, we previously showed that girls are 1002

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more susceptible to cyclosporine A-associated hypertension
than boys.²⁵

1005 Patients were allocated to dialysis or preemptive trans-1006 plantation on the basis of clinical decisions. Bias by indication was overcome by adjusting for all factors that potentially in-1007 1008 fluence the treatment decision (e.g., kidney disease, center, 1009 time, and kidney function parameter). As not all patients' data 1010 were available for the final model because of the timing of 1011 examinations, there was a potential selection bias. However, 1012 as there were no differences in PWV or sex distribution be-1013 tween patients who were or were not included in the final 1014 models (pre-KRT: inclusion, n = 156; noninclusion, n = 741015 and post-transplantation: inclusion, n = 187; noninclusion, n = 43), this should have not influenced the results of the 1016 1017 comparison between sexes. The majority of our study pop-1018 ulation is Caucasian (88%) and so was the population from which PWV reference values were calculated.¹³ This might 1019 1020 limit the generalizability of our finding.

1022 Conclusion

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1023 The observed higher susceptibility of girls for cardiovascular 1024 organ damage in conjunction with kidney disease progression 1025 highlights the importance of a closer attention to cardiovas-1026 cular and kidney function parameters early in the disease 1027 course in female patients. Importantly, girls are more 1028 vulnerable toward eGFR decline and when exposed to a 1029 longer waiting time to transplantation. Early interventions 1030 and a faster access of girls to transplantation are crucial to 1031 tackle the sex differences in cardiovascular and mortality risk. 1032 Strict BP control and management of dyslipidemia are of 1033 importance for both sexes. 1034

1035 1026 DISCLOSURE

All the authors declared no competing interests.

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1048 SUPPLEMENTARY MATERIAL

1049 Supplementary File (PDF)

- 1050 Figure S1. Assessment of time variables.
- 1051Figure S2. Assessment of changes in estimated glomerular filtration1052rate (eGFR) during pre-kidney replacement therapy (KRT).
- 1053 Figure S3. Number of observations over time since inclusion,
- 1054 differentiated by the modality of kidney replacement therapy at each visit since the inclusion.
- Figure S4. Effect estimates and 95% confidence interval for factors associated with PWV z scores (PWVz) during pre-kidney replacement therapy (KRT).

- Figure S5. Additional mixed model for with PWV z scores (PWVz) 1059 adjusted for kidney underlying disease category. 1060 Table S1. Sub-classifications of primary kidney diseases. 1061 Table S2. Correlation between Casual blood pressure (BP) and 1062 ambulatory BP measurement (ABPM). 1063 Table S3. Matrix of missing data for each final model of pre-kidney 1064 replacement therapy (KRT; A) and post-transplantation (B). 1065 Table S4. Comparison of baseline characteristics between the study population (at first visit pre-kidney replacement therapy [KRT]) and 1066 healthy children cohort (at study inclusion). 1067 Table S5. Covariates screening based on basic model for pulse wave 1068 velocity z scores (PWVz) "pre-kidney replacement therapy (KRT)". 1069 Table S6. Final models for pulse wave velocity z scores (PWVz) "pre-1070 kidney replacement therapy (KRT)" separated by sex. 1071 Table S7. Basic mixed models for pulse wave velocity z scores (PWVz) 1072 "post-transplantation" separated by transplantation type, i.e., preemptive transplantation and transplantation after prior dialysis. 1073 Table S8. Covariates screening based on basic model for pulse wave 1074 velocity z scores (PWVz) "post-transplantation". 1075 Table S9. Sensitivity analysis for the effect of the pre-kidney 1076 replacement therapy (KRT) estimated glomerular filtration rate (eGFR) 1077 slopes calculated from single mixed model on post-transplantation 1078 pulse wave velocity z scores (PWVz). Table S10. Pulse wave velocity z scores (PWVz) and pulse wave 1079 velocity (PWV; m/s) at different time points for all patients and 1080 differentiated by sex. 1081 1082 1083 REFERENCES 1. Viner RM, Coffey C, Mathers C, et al. 50-Year mortality trends in children 1084 and young people: a study of 50 low-income, middle-income, and high-1085 income countries. Lancet. 2011;377:1162-1174. 1086 Sawyer CC. Child mortality estimation: estimating sex differences in 2. childhood mortality since the 1970s. PLoS Med. 2012;9:e1001287. 1087 Ghuman AK, Newth CJ, Khemani RG. Impact of gender on sepsis 3 1088 mortality and severity of illness for prepubertal and postpubertal 1089 children. J Pediatr. 2013;163:835-840.e831. 4. Bhaumik U, Aitken I, Kawachi I, et al. Narrowing of sex differences in 1090 infant mortality in Massachusetts. J Perinatol. 2004;24:94-99. 1091 Groothoff JW, Gruppen MP, Offringa M, et al. Mortality and causes of 5. 1092 death of end-stage renal disease in children: a Dutch cohort study. Kidney Int. 2002:61:621-629. 1093 Ahearn P, Johansen KL, McCulloch CE, et al. Sex disparities in risk of 1094 mortality among children with ESRD. Am J Kidney Dis. 2019;73:156-162. 1095 7. Laskin BL, Mitsnefes MM, Dahhou M, et al. The mortality risk with graft function has decreased among children receiving a first kidney 1096 transplant in the United States. Kidney Int. 2015;87:575-583. 1097 Mitsnefes MM. Cardiovascular disease in children with chronic kidney 8. 1098 disease. J Am Soc Nephrol. 2012;23:578-585. Francis A, Johnson DW, Melk A, et al. Survival after kidney 1099 transplantation during childhood and adolescence. Clin J Am Soc 1100 Nephrol. 2020;15:392-400. 1101
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