



# Article Improved Kidney Allograft Function after Early Conversion of Fast IR-Tac Metabolizers to LCP-Tac

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Abstract: Fast tacrolimus (Tac) metabolism is associated with a more rapid decline of renal function after renal transplantation (RTx). Because the pharmacokinetics of LCP-Tac (LCPT) and immediaterelease Tac (IR-Tac) differ, we hypothesized that switching from IR-Tac to LCPT in kidney transplant recipients would improve the estimated glomerular filtration rate (eGFR), particularly in fast metabolizers. For proof of concept, we performed a pilot study including RTx patients who received de novo immunosuppression with IR-Tac. A Tac concentration-to-dose ratio (C/D ratio) < 1.05 ng/mL·1/mg defined fast metabolizers and  $\geq$ 1.05 ng/mL·1/mg slow metabolizers one month after RTx. Patients were switched to LCPT  $\geq$  1 month after transplantation and followed for 3 years. Fast metabolizers (n = 58) were switched to LCPT earlier than slow metabolizers (n = 22) after RTx (2.0 (1.0–253.1) vs. 13.2 (1.2–172.8) months, p = 0.005). Twelve months after the conversion to LCPT, Tac doses were reduced by about 65% in both groups. The C/D ratios at 12 months had increased from 0.66 (0.24–2.10) to 1.74 (0.42–5.43) in fast and from 1.15 (0.32–3.60) to 2.75 (1.08–5.90) in slow metabolizers. Fast metabolizers showed noticeable recovery of mean eGFR already one month after the conversion  $(48.5 \pm 17.6 \text{ vs. } 41.5 \pm 17.0 \text{ mL/min}/1.73 \text{ m}^2, p = 0.032)$  and at all subsequent time points, whereas the eGFR in slow metabolizers remained stable. Switching to LCPT increased Tac bioavailability, C/D ratio, and was associated with a noticeable recovery of renal function in fast metabolizers. Conversion to LCPT is safe and beneficial early after RTx.

Keywords: kidney transplantation; renal; LCPT; tacrolimus; metabolism; C/D ratio; conversion; switch

# 1. Introduction

During the last decades, tacrolimus (Tac) has become the most commonly prescribed calcineurin inhibitor (CNI) after solid organ transplantation [1,2]. Three different Tac formulations are available. Immediate-release Tac (IR-Tac) was the first formulation on the market and has to be administered twice-daily, resulting in two blood peak levels: a higher one in the morning, about 1.5 to 2 h after drug intake, and a lower one about 12 h later [3]. Secondly, extended-release Tac (ER-Tac) was developed to simplify administration by replacing croscarmellose in IR-Tac with ethylcellulose in ER-Tac. The formulation of ER-Tac allows for once-daily administration, resulting in higher adherence rates [4], but is associated with an early high Tac peak level after ingestion that is comparable to the first peak of IR-Tac [3]. Thirdly, LCP-Tac (LCPT) is a new formulation using the MeldDose<sup>®</sup> prolonged-release technology that provides absorption throughout the entire gastrointestinal tract. This results in lower and later peak concentrations approximately 4 to 6 h after drug intake [3].



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Several studies have reported that the treatment of fast metabolizers after RTx with IR- or ER-Tac is associated with adverse outcomes [5-10]. In 2014, we established the Tac concentration-to-dose ratio (C/D ratio) as a surrogate marker for the Tac metabolism rate [5]. Interestingly, a low C/D ratio is associated with higher peak-level concentrations and with a more rapid decline of kidney function compared to slow metabolizers (C/D ratio  $\geq$ 1.05 ng/mL·1/mg) as early as one month after RTx [5,11]. In two consecutive studies, we were able to confirm the decreased renal function of fast Tac metabolizers compared with slow metabolizers in separate IR- and ER-Tac cohorts in a 5-year follow-up [6,12]. The reasons for the impairment of renal function in the group of fast metabolizers have been reported as increased rates of acute rejections (AR) [6,10], BK-virus infections and nephropathy [5,13], CNI-nephrotoxicity (CNIT) [5,12], and higher incidences of interstitial fibrosis and tubular atrophy (IF/TA) [9]. Fast Tac metabolizers were also converted earlier and more frequently to other immunosuppressive drugs compared with slow metabolizers because of Tac-associated side effects [5,12]. For example, we found a strong association between the Tac metabolism rate and the degree of biopsy-proven CNIT [11]. Recently, two studies in French and German cohorts showed increased mortality rates even in fast Tac metabolizers [6,8]. In 2020, von Einsiedel et al. published a study on liver transplant recipients (LTR) and showed an improvement in renal function after the conversion from IR- and ER-Tac to LCPT [14]. Though in this study, the Tac dose was reduced by about 30%, the C/D ratio or bioavailability increased by about 50% twelve months after the conversion.

As bioavailability, and Tac peak levels in the blood especially, differ significantly between LCPT and IR-Tac treated patients, we hypothesize that the conversion of fast IR-Tac metabolizers to LCPT results in an improvement in kidney function.

### 2. Patients and Methods

# 2.1. Study Cohort

We retrospectively analyzed RTx recipients who underwent transplantation at the University Hospital of Münster between 2005 and 2019. All patients started immunosuppression with IR-Tac and remained on the medication for at least one month after RTx. Included patients were switched to LCPT one month after RTx or later and then followed up for three years. Target Tac trough level was 6–10 ng/mL for the first month. LCPT was introduced with a 30% reduction of preconversion daily IR-Tac [3]. Co-immunosuppression consisted of mycophenolate mofefil or mycophenolate sodium and prednisolone. Induction therapy was basiliximab or thymoglobulin, depending on immunological risk.

The C/D ratio was calculated as described before [13]. A C/D ratio < 1.05 ng/mL·1/mg defined fast metabolizers, and a value  $\geq$  1.05 ng/mL·1/mg defined slow metabolizers. IR-Tac doses, trough levels, and C/D ratios were given at month 1 (M1) before the switch, at the day of the switch (before switch), for LCPT at day 10 (D10), and month 1-36 (M1-36) after conversion. The estimated glomerular filtration rate (eGFR) was calculated using the CKD-EPI formula [15] at D10 and M1 after RTx, at the day of the switch (before switch), and D10, M1-36 after conversion.

General demographic data on recipients and information on transplantation were obtained from electronic health records, and data on donors were taken from Eurotransplant. The study was performed in accordance with the Declaration of Helsinki and approved by the local ethics committee (Ethik Kommission der Ärztekammer Westfalen-Lippe und der Medizinischen Fakultät der Westfälischen Wilhelms-Universität, No. 2014-381-f-N). Written informed consent with regard to recording their clinical data was given by all participants at the time of transplantation or inclusion into the study. Data were anonymized before analysis. RTx recipients younger than 18 years of age and pregnant women were excluded.

# 2.2. Statistics

Statistical analyses were performed using IBM SPSS<sup>®</sup> Statistics 27 for Windows (IBM Corporation, Somers, NY, USA) and SAS software, Version 9.4 TS1M7 of the SAS System for Windows (Copyright © 2021 SAS Institute Inc., Cary, NC, USA). All *p*-values and

confidence limits were two-sided and were intended to be exploratory, not confirmatory. Therefore, no adjustment for multiplicity was made. Exploratory *p*-values  $\leq$  0.05 were considered to be statistically noticeable.

In descriptive analysis, normally distributed continuous variables are reported as mean  $\pm$  standard deviation and non-normally distributed continuous variables as median (25% quantile–75% quantile, IQR). Absolute and relative frequencies are given for categorical variables. Metabolism groups were compared using Welch's *t*-tests for normally distributed data, Mann–Whitney U tests for skewed-distributed data, and Fisher's exact or chi-squared tests for categorical variables. A comparison of the eGFR changes within each metabolism group was performed using Wilcoxon's signed-rank tests. Boxplots were used for graphical representation.

In order to model renal function (eGFR) over time adjusted for dropouts, a multivariable linear mixed model was fitted. The main effects of the factors, which were time since switch (Day 10, month 1, 3, 6, 9, 12, 24, and 36 after switch), metabolizer group (fast/slow), and the interaction between time and group were included as influencing variables. Repeated measurements of each patient were modeled using SAS PROC MIXED by fitting a marginal linear mixed model with an unstructured variance–covariance matrix for the residuals with patient as subject and the order given by time. The empirical sandwich covariance estimator was applied. Missing values were treated as missing at random. Results are reported as least square estimates with the corresponding 95% confidence interval (CI) and *p*-values from the Wald test.

# 3. Results

# 3.1. Study Cohort

In total, we followed up 80 RTx recipients who started on IR-Tac and were switched to LCPT one month after RTx or later. A total of 58 patients were characterized as fast and 22 as slow metabolizers according to their corresponding Tac C/D ratio at one month after RTx (Figure 1). Baseline characteristics at RTx did not differ noticeably between the groups (Table 1). There were no noticeable differences between the groups in terms of underlying diseases leading to end-stage renal disease. Most of the patients had arterial hypertension at the time of RTx. In average, fast Tac metabolizers were converted earlier to LCPT compared to slow metabolizers (0.2 (1.0–253.1) vs. 13.2 (1.2–172.8) months after RTx; p = 0.005). The main reasons for a switch in both groups were large trough level variations and avoidance of adverse effects (Table 2).



**Figure 1.** Study design and patient enrolment. A total of 80 renal transplant recipients met the inclusion criteria. RTx recipients were defined as fast and slow Tac metabolizers one month after transplantation. All patients were switched to LCPT and observed in a 3-year follow-up. Abbreviations: RTx, renal transplantation; IR-Tac, immediate-release tacrolimus; C/D ratio, concentration-to-dose ratio.

	Fast Metabolizers $n = 58$	Slow Metabolizers $n = 22$	<i>p</i> -Value		
age (years)	$50.0\pm16.1$	$49.8 \pm 14.7$	0.957 <sup>a</sup>		
sex (m/f), n (%)	37 (63.8%)/21 (36.2%)	13 (59.1%)/9 (40.9%)	0.797 <sup>b</sup>		
weight (kg)	$80.8 \pm 18.3$	$76.4 \pm 15.1$	0.274 <sup>a</sup>		
height (m)	$1.77\pm0.09$	$1.75\pm0.09$	0.267 <sup>a</sup>		
$BMI (kg/m^2)$	$25.5\pm5.3$	$24.9\pm4.3$	0.570 <sup>a</sup>		
living donor transplantation	37 (63.8%)	16 (72.7%)	0.598 <sup>b</sup>		
ABO-i	13 (22.4%)	7 (31.8%)	0.399 <sup>b</sup>		
ESP transplantation	4 (6.9%)	1 (4.5%)	1 <sup>b</sup>		
DGF	6 (10.3%)	3 (13.6%)	0.700 <sup>b</sup>		
cold ischemic time (h)	$7.4 \pm 5.0$	$6.7 \pm 4.3$	0.539 <sup>a</sup>		
warm ischemic time (min)	$35.9\pm8.3$	$35.9\pm9.2$	0.968 <sup>a</sup>		
	Number of Transplan	tations			
1	51 (87.9%)	19 (86.4%)			
2	6 (10.3%)	3 (13.6%)	0.785 <sup>b</sup>		
3	1 (1.7%)	0	000		
	HLA MM				
0	13 (22.4%)	8 (36.4%)			
1–3	23 (39.7%)	7 (31.8%)	0.462 <sup>b</sup>		
4–6	22 (37.9%)	7 (31.8%)	0.10_		
PRA > 20%	5 (8.6%)	2 (9.1%)	1 <sup>b</sup>		
	CMV risk status	5			
low	13 (22.4%)	2 (9.1%)			
intermediate	34 (58.6%)	11(50.0%)	0.105 <sup>b</sup>		
high	11 (19.0%)	9 (40.9%)			
Donor Characteristics					
donor age (years)	$52.7 \pm 12.2$	$55.8\pm9.7$	0.246 <sup>a</sup>		
donor sex $(m/f)$ , n (%)	24 (41.4%)/34 (58.6%)	7 (31.8%)/15 (68.2%)	0.608 <sup>b</sup>		
Diagnosis of ESRD					
benign nephrosclerosis	4 (6.9%)	1 (4.5%)			
diabetic nephropathy	3 (5.2%)	1 (4.5%)			
glomerulonephritis	31 (53.4%)	14 (63.6%)			
chronic pyelonephritis	0	1 (4.5%)			
cystic nephropathy	14 (24.1%)	2 (9.1%)	0.345 <sup>b</sup>		
Alport syndrome	1 (1.7%)	0			
Mediterranean fever	0	1 (4.5%)			
congenital renal dysgenesis	4 (6.9%)	1 (4.5%)			
interstitial nephritis	1 (1.7%)	1 (4.5%)			
Comorbidities Before Transplantation					
arterial hypertension	56 (96.6%)	21 (95.5%)	1 <sup>b</sup>		
diabetes mellitus	8 (13.8%)	1 (4.5%)	0.432 <sup>b</sup>		

Table 1. Patients' characteristics.

Data presented as mean  $\pm$  standard deviation or absolute and relative frequencies. Abbreviations: BMI, body mass index; ABO-i, ABO incompatible transplantation; ESP, European Senior Program; DGF, delayed graft function; HLA MM, human leucocyte antigen mismatch; PRA, panel reactive antibodies; CMV, cytomegalovirus; ESRD, end-stage renal disease. Two-sided *p*-value from <sup>a</sup> Welch's *t*-test or <sup>b</sup> Fisher's exact test.

	Fast Metabolizers $n = 58$	Slow Metabolizers $n = 22$	<i>p</i> -Value
infection	1 (1.7%)	1 (4.5%)	
neurological disorder	2 (3.4%)	0	
acute rejection	1 (1.7%)	0	
CNIT	2 (3.4%)	2 (9.1%)	0.778
DGF	2 (3.4%)	1 (4.5%)	
diabetes mellitus	1 (1.7%)	0	
trough level			
variation/avoidance of adverse effects	49 (84.5%)	18 (81.8%)	

Table 2. Reasons for the switch from IR-Tac to LCPT.

IR-Tac, immediate-release tacrolimus; LCPT, LCP-tacrolimus; CNIT, calcineurin inhibitor toxicity; DGF, delayed graft function. The *p*-value is from the chi-squared test of independence.

#### 3.2. Immunosuppression

The immunosuppression is shown in Table 3. Prednisolone and mycophenolate doses did not differ between the metabolizer groups one month after RTx.

IR-Tac and LCPT doses of fast metabolizers were higher at all time points compared to slow metabolizers (all p < 0.016). After the recommended dose reduction of 30% at the day of conversion in both groups, the dose reduction in fast metabolizers was 31.7% on D10, 64.6% on M12, and 70.7% on M36. In slow metabolizers, the median Tac dose was reduced by 48.2% at D10, by 66.7% at M12, and by 70.4% at M36.

IR-Tac trough levels at month 1 and LCPT trough levels at time points D10, M1 and M6 were lower in fast metabolizers. Tac trough levels of fast metabolizers increased slightly after the conversion and achieved the level of the trough concentration "before conversion" approximately at the 12 months mark after conversion (M12). In the group of slow metabolizers, trough concentrations were reduced considerably during the 36-month follow-up.

The C/D ratios of fast metabolizers were lower compared to the C/D ratios of slow metabolizers at all time points. After conversion to LCPT, the median C/D ratio in fast metabolizers increased by 38.9% at D10, 163.6% at M12 and 180.3% after M36. The C/D ratios of slow metabolizers rose by 66.1% at D10 days, 139.1% at M12 and 130.4% after 36 months.

## 3.3. Renal Function

Compared with slow metabolizers, fast IR-Tac metabolizers showed a slightly but not noticeably reduced renal function one month after RTx (47.6  $\pm$  20.8 vs. 44.4  $\pm$  19.0 mL/ min/1.73 m<sup>2</sup>, respectively, *p* = 0.539, Figure 2). After conversion to LCPT, we observed similar eGFR values in both groups. In fast metabolizers, the mean eGFR increased from 41.5  $\pm$  17.0 (before switch) to 48.5  $\pm$  17.6 at M1, to 47.8  $\pm$  15.4 at M12, and to 48.9  $\pm$  19.0 mL/min/1.73 m<sup>2</sup> at M36 after the conversion. The mean eGFR values of slow metabolizers were 42.2  $\pm$  17.5 before the switch and 43.2  $\pm$  17.7 at M1, 43.7  $\pm$  17.9 at M12, and 43.8  $\pm$  19.2 mL/min/1.73 m<sup>2</sup> at M36 after the conversion.

	Fast Metabolizers n = 58	Slow Metabolizers $n = 22$	<i>p</i> -Value
Prednisolone Dose after 1 Month	17.5 (5–25)	15 (5–50)	0.680 <sup>a</sup>
Mycophenolate after 1 Month			
mycophenolate mofetil, <i>n</i> (%)	30 (51.7%)	13 (59.1%)	0 ( <b>0</b> 1 h
mycophenolate sodium, n (%)	28 (48.3%)	9 (40.9%)	0.621 °
mycophenolate mofetil dose (mg)	1000 (500-2000)	1000 (500-2000)	0.932 <sup>a</sup>
mycophenolate sodium dose (mg)	1440 (720–1440)	1080 (720–1440)	0.213 <sup>a</sup>
Tac Doses (mg)			
IR-Tac M1	12 (5–20)	7 (4–12)	<0.001 a
before switch (IR-Tac)	10.25 (3-18)	6.75 (1.5–17)	<0.001 <sup>a</sup>
D10 LCPT	7 (1.5–14)	3.5 (1.8 11)	<0.001 <sup>a</sup>
M1 LCPT	6 (1.5–13.5)	3 (1.5–11)	<0.001 <sup>a</sup>
M3 LCPT	4.75 (1.5–12)	3 (1–8)	0.002 <sup>a</sup>
M6 LCPT	4 (1.5–12)	2.5 (0.75-5)	0.001 <sup>a</sup>
M9 LCPT	4 (1.5–11)	2.5 (1-6)	0.001 <sup>a</sup>
M12 LCPT	3.63 (1.5-11)	2.25 (1-5)	0.001 <sup>a</sup>
M24 LCPT	3.38 (1–9)	2.13 (0.75-5.5)	0.008 <sup>a</sup>
M36 LCPT	3 (1–8.5)	2 (0.75–3.5)	0.016 <sup>a</sup>
Tac trough Levels (ng/mL)			
IR-Tac M1	6.8 (2.4–15.9)	8.7 (6.8–13.5)	<0.001 <sup>a</sup>
before switch (IR-Tac)	6.3 (2.4–12.3)	7.5 (3.9–13.5)	0.065 <sup>a</sup>
D10 LCPT	7.2 (1.6–14.7)	6.3 (4.1–9.9)	0.026 <sup>a</sup>
M1 LCPT	7.6 (1.5–19.5)	6.2 (3.7–12.9)	0.041 <sup>a</sup>
M3 LCPT	7.3 (3.8–18.1)	6.7 (4.9–9.9)	0.311 <sup>a</sup>
M6 LCPT	7.0 (2.7–11.4)	6.2 (4.0–10.4)	0.043 <sup>a</sup>
M9 LCPT	6.5 (3.4–10.7)	6.3 (4.2–9.2)	0.992 <sup>a</sup>
M12 LCPTT	6.2 (3.5–10.3)	6.05 (3.8-8.3)	0.224 <sup>a</sup>
M24 LCPT	6.2 (4.0-10.7)	6.2 (2.1–9.4)	0.254 <sup>a</sup>
M36 LCPT	5.5 (4.1-8.9)	5.40 (4.0-8.7)	0.698 <sup>a</sup>
Tac C/D Ratio (ng/mL·1/mg)			
IR-Tac M1	0.64 (0.24-1.01)	1.25 (1.08-3.38)	<0.001 <sup>a</sup>
before switch (IR-Tac)	0.66 (0.24-2.10)	1.15 (0.32-3.60)	0.001 <sup>a</sup>
D10 LCPT	1.08 (0.33-4.90)	1.91 (0.40-4.06)	0.002 <sup>a</sup>
M1 LCPT	1.24 (0.21-6.93)	2.23 (0.55-3.47)	0.010 <sup>a</sup>
M3 LCPT	1.52 (0.55-4.93)	2.33 (0.94-6.60)	0.004 <sup>a</sup>
M6 LCPT	1.58 (0.39-5.93)	2.65 (1.06-7.07)	0.007 <sup>a</sup>
M9 LCPT	1.63 (0.40-5.07)	3.23 (1.23-6.30)	<0.001 a
M12 LCPTT	1.74 (0.42-5.43)	2.75 (1.08-5.90)	0.007 <sup>a</sup>
M24 LCPT	1.81 (0.64-5.40)	2.58 (0.96-6.27)	0.083 <sup>a</sup>
M36 LCPT	1.85 (0.69-5.80)	2.65 (1.32-5.73)	0.026 <sup>a</sup>

# Table 3. Immunosuppression.

Data presented as median (25% quantile-75% quantile) or absolute and relative frequencies. Abbreviations: Tac, tacrolimus; C/D, concentration-to-dose; two-sided *p*-values from <sup>a</sup> Mann–Whitney U test or <sup>b</sup> Fisher 's exact test.





**Figure 2.** Boxplots of the renal function at different time points (eGFR, estimated glomerular filtration rate) within 3 years after kidney transplantation. *p*-values are from Mann–Whitney U tests comparing fast vs. slow metabolizers at each time point. There were no noticeable differences between the groups after conversion.

In addition, the differences in eGFR between the "before switch" time point and all other time points ( $\Delta$ eGFR) were analyzed (Figure 3). In fast metabolizers, the  $\Delta$ eGFR between "M1 before switch" and the date "before switch" differed noticeably (Figure 3A). Additionally, all  $\Delta$ eGFR values following the conversion also showed a noticeable increase compared with the value "before switch". In slow metabolizers, eGFR values did not differ noticeably between "before switch" and all other time points (Figure 3B).

In the multivariate analysis, a noticeable change of the eGFR over time pooled over both metabolism groups was observed (p = 0.003, Table S1). In this analysis, there was a noticeable eGFR increase in fast metabolizers at all time points compared to before the conversion. No noticeable changes in eGFR values ( $\Delta$ eGFR) in slow metabolizers compared to before the conversion were found. When comparing the changes in eGFR ( $\Delta$ eGFR) between fast and slow metabolizers (combination of main and interaction effects of tacrolimus metabolism group and time points), the fast group diverged the most from slow metabolizers within the first 10 days after the switch (4.15 (95% CI 0.54–7.76), p = 0.025). A further slight increase in the difference between the groups was found between D10 and M1 (1.84 (95% CI -1.39-5.07, p = 0.259).







Figure 3. Boxplots of the differences in estimated glomerular filtration rates ( $\Delta eGFR$ ) of fast and slow Tac metabolizers (all time points-date of the switch). p-values are from Wilcoxon-signed rank tests for the dependent comparisons of eGFR values at each time point with the eGRF values at the time of the switch. The C/D ratio cut-off to characterize the metabolizer group was 1.05 ng/mL·1/mg. Fast metabolizers developed a noticeable increase in ΔeGFR at D10 after conversion and all following time points (A).  $\Delta$ eGFR values of slow metabolizers remained stable during the 3 years after the switch (B).

In a further examination,  $\Delta eGFR$  values were compared between fast and slow metabolizers when a lower C/D ratio cut-off of 0.6 ng/mL $\cdot$ 1/mg was chosen (<0.6 = fast metabolizers;  $\geq 0.6 =$  slow metabolizers (Figure 4)). In this analysis, the increase in  $\Delta$ eGFR was even more pronounced in fast metabolizers (Figure 4A). Interestingly, slow metabolizers were also found to have higher ∆eGFR at "M1 before switch" and "M1 to M9 after switch" (Figure 4B).





**Figure 4.** Boxplots of the differences in estimated glomerular filtration rates ( $\Delta$ eGFR) of fast and slow Tac metabolizers (all time points—date of the switch). *p*-values are from Wilcoxon-signed rank tests for the dependent comparisons of eGFR values at each time point with the eGRF values at the time of the switch. The C/D ratio cut-off that characterized the metabolizer groups was changed to 0.6 ng/mL·1/mg. In this case, fast metabolizers developed an even more pronounced increase in  $\Delta$ eGFR at D10 after conversion and all following time points compared with fast metabolizers defined by a C/D ratio cut-off of 1.05 ng/mL·1/mg (**A**).  $\Delta$ eGFR of slow metabolizers also increased at M1–M9 after conversion to LCPT (**B**).

## 3.4. Complication Rates

No differences were observed in infection rates (CMV, BKV), BKVN, de novo development of post-transplant diabetes mellitus, CNIT, AR, or death between the groups at any time during the study (Table 4).

	Fast Metabolizers n = 58	Slow Metabolizers $n = 22$	<i>p</i> -Value
CMV infection			
before switch to LCPT after switch to LCPT (3 y-follow up)	8 (13.8%) 3 (5.2%)	2 (9.1%) 2 (9.1%)	0.719 0.612
BKV infection			
before switch to LCPT after switch to LCPT (3 y-follow up)	3 (5.2%) 1 (1.7%)	3 (13.6%) 0	0.338 1
BKV nephropathy			
before switch to LCPT after switch to LCPT (3 y-follow up)	1 (1.7%) 0	2 (9.1%) 0	0.182
CNIT			
before switch to LCPT after switch to LCPT (3 y-follow up) acute rejection	3 (5.2%) 1 (1.7%)	2 (9.1%) 0	0.612 1
before switch to LCPT after switch to LCPT (3 y-follow up)	12 (20.7%) 8 (13.8%)	9 (40.9%) 2 (9.1%)	0.089 0.719
death within 3 years after switch diabetes mellitus	2 (3.4%)	0	1
before switch to LCPT after switch to LCPT (3 y-follow up)	1 (1.7%) 0	1 (4.5%) 0	0.477

Table 4. Complications before and after switch.

CMV, cytomegalovirus; BKV, BK-virus; CNIT, calcineurin-inhibitor nephrotoxicity; LCPT, LCP-tacrolimus. *p*-values are from the Fisher's exact test.

## 4. Discussion

In this study, we observed that the marked decline in renal function frequently observed in fast Tac metabolizers treated with IR- or ER-Tac after transplantation was prevented after switching to LCPT (Figure 3) [6,12].

Previously, we found that fast Tac metabolism in IR-treated RTx patients is associated with higher Tac peak levels and higher rejection and BKVN rates, leading to worse renal function compared with IR-treated slow metabolizers [5,6,11,13]. In our current observations, the renal function (eGFR) of the fast metabolizers was about 3 mL/min/1.73 m<sup>2</sup> lower than that of the comparison group before switching (Figure 2), but due to smaller cohorts in this study, the eGFR did not differ noticeably (p = 0.539) between groups. Others observed higher IF/TA levels in fast metabolizers than in slow metabolizers within one year of RTx [9,16]. Consistently, this was associated with worse graft survival and even mortality in fast metabolizers [6,8]. Although mortality was mainly caused by cardiovascular events, we did not observe differences in arterial stiffness or lipid levels between fast and slow metabolizers [11,17].

A major reason for switching from IR-Tac to LCPT was the variability in intra-patient trough levels (IPV, Table 2). IPV is associated with worse outcomes for several reasons [18]. LCPT has been shown to at least decrease Tac fluctuation and swing, and once-daily dosing is usually associated with increased adherence, both factors contributing to IPV, although study data on IPV in LCPT-treated patients are lacking [18,19].

LCPT was shown to be non-inferior to IR-Tac in RTx recipients and demonstrated lower efficacy failure rates in blacks [20,21]. Notably, blacks are often CYP3A5 expressers encoding tacrolimus-metabolizing cytochrome P450 3A5 enzymes and are therefore predominantly fast metabolizers [22,23]. Moreover, the fraction of fast metabolizers that are in the therapeutic Tac trough range is considerably higher for LCPT than for IR-Tac, potentially reducing the rejection rate in fast metabolizers (at least by trend, LCPT group: 23.2% vs. 36.6% IR-Tac group) [24]. In contrast, others showed that the rate of BKVN was lower in LCPT-treated recipients than in IR-Tac-treated ones [25]. This is interesting because we found an association between fast Tac metabolism and BKVN as well as BKV infection [5,13]. However, other studies found comparable BKV infection rates between LCPT and IR-Tac in de novo RTx recipients [21]. There were also no differences in metabolic parameters such as lipids or HbA1c between LCPT- and IR-treated patients [21,25]. Since the total exposure to the drug (area under the curve) and the trough level of fast and slow metabolizers and of LCPT and IR-Tac-treated patients are generally comparable, other factors—and it is probably not Tac metabolites that are similar in patients with low and high C/D ratios—must be responsible for the observed differences [11,19,22,26,27].

An important difference in the Tac-pharmacokinetics (PK) of patients with a low and a high C/D ratio is the peak concentration, which is thought to play a role in the development of Tac-associated nephrotoxicity [11]. In addition, it is noteworthy that the potentially harmful peak concentration is much lower with LCPT than with IR-Tac or ER-Tac [3]. In contrast to IR-Tac, LCPT treatment causes only one and not two Tac peaks and is not affected by chronopharmacokinetic effects [28]. An important point in this regard is that the major absorption site of LCPT is more distal in the gut than that of IR-Tac. As CYP3A5 expression decreases in the distal intestine, the bioavailability of LCPT is better than that of IR-Tac (as clearly shown by the reduced dosage after conversion, Table 3) and the dependence on CYP3A4/5 drug interference decreases significantly [29,30]. This is important because CYP3A5\*1 is distinctly associated with fast Tac metabolism and CNIT [26]. Trofe-Clark et al. elegantly demonstrated in a study of stable RTx patients using a cross-over design that the PK of CYP3A5\*1 expressers treated with LCPT, as opposed to IR-Tac treated individuals, was comparable to the PK of non-expressers. All treated patients benefited from the reduction in peak Tac concentration regardless of the CYP3A expression status, although the effect was significantly more pronounced in CYP3A5\*1 expressers [22]. This may be one reason why fast metabolizers did not show pronounced eGFR loss during the 3-year follow-up period in the current study (Figure 2). Of note, fast metabolizers even showed a noticeable increase in renal function after the conversion to LCPT. These results are in line with a study on LTR that showed a noticeable recovery of eGFR 12 months after the conversion to LCPT (65.3 vs. 70.9 mL/min/1.73 m<sup>2</sup>; p < 0.001) [14].

As previously shown by us and others, eGFR was lower in fast IR- and ER-Tac metabolizers than in slow metabolizers already at M1 after RTx [6–8,12]. Since we did not perform PK analyses and assessed peak level, this is a limitation of our study. However, it should be remembered that the total absolute daily dose remained higher in fast metabolizers than in slow metabolizers, even under LCPT treatment; an effect that we also observed in our study (Table 3). Overall, it seems logical that the treatment with LCPT increases the C/D ratio when compared to IR-Tac calculated C/D ratios (Figure 3). This is in line with the literature [31,32].

Our study has several limitations. Because it is a single-center study with a limited number of patients, its power is limited. Further, due to the retrospective nature of our pilot study (first proof of concept), it can only be hypothesis-generating and encourage further prospective studies. Moreover, we did not determine the PK, peak, or IPV in our patients. Therefore, we can only make assumptions about the effects of LCPT on renal values. Fast metabolizers were mostly switched earlier to LCPT than slow metabolizers. This is not surprising, as the main reasons for switching were the side effects and IPV. Side effects and IPV are partly related to Tac dosage which are usually higher early after transplantation than later, due to different reasons, such as the intended trough level or decreasing steroid dosage [23]. However, steroid dosage was comparable in both groups (Table 3). Because we and others already saw the detrimental renal effects of Tac in the first months after RTx in fast metabolizers, this could be a reason for the more pronounced effect in the group with the low C/D ratio [5,10].

## 5. Conclusions

Our study revealed that switching from IR-Tac to LCPT increased the bioavailability of Tac, saved doses (and costs), and increased the C/D ratio. This may have mitigated the pronounced deterioration of renal function, particularly in fast metabolizers, which are

more vulnerable in this respect. Because patients with a low C/D ratio are at risk for a worse outcome after RTx, we suggest calculating the C/D ratio early after RTx in patients treated with IR-Tac. Switching to LCPT was safe and could be beneficial early after RTx.

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/jcm11051290/s1, Table S1: Immunosuppression; Table S2: Model-based estimates of eGFR.

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