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Jinyu Liu Huazhong University of Science and Technology

Dong Liu Huazhong University of Science and Technology

Juan Li Huazhong University of Science and Technology

Lan Zhu Huazhong University of Science and Technology

Chengliang Zhang Huazhong University of Science and Technology

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Authors

Jinyu Liu, Dong Liu, Juan Li, Lan Zhu, Chengliang Zhang, Kai Lei, Qiling Xu, and Ruxu You



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RESEARCH ARTICLE

Efficacy and Safety of Everolimus for Maintenance Immunosuppression of Kidney Transplantation: A Meta-Analysis of Randomized Controlled Trials

Jinyu Liu¹, Dong Liu¹, Juan Li¹, Lan Zhu², Chengliang Zhang¹, Kai Lei¹, Qiling Xu³, Ruxu You⁴*

 Department of Pharmacy, Tongji Hospital, Tongji Medical College, Huazhong University of Science and Technology, Wuhan, Hubei, China, 2 Institute of Organ Transplantation, Tongji Hospital, Tongji Medical College, Huazhong University of Science and Technology, Wuhan, Hubei, China, 3 Department of Biotechnology and Molecules, Assumption College, Worcester, Massachusetts, United States of America, 4 Department of Pharmacy, Union Hospital, Tongji Medical College, Huazhong University of Science and Technology, Wuhan, Hubei, China

* youruxu2008@163.com

Abstract

Background

Conversion to everolimus is often used in kidney transplantation to overcome calcineurin inhibitor (CNI) nephrotoxicity but there is conflicting evidence for this approach.

Objectives

To investigate the benefits and harm from randomized clinical trials (RCTs) involving the conversion from CNI to everolimus after kidney transplantation.

Methods

Databases were searched up to March 2016. Two reviewers independently assessed trials for eligibility and quality, and extracted data. Results are expressed as risk ratio (RR) or mean difference (MD) with 95% confidence intervals (CI).

Results

Eleven RCTs, with a total of 1,633 patients, met the final inclusion criteria. Patients converted to everolimus had improved renal function at 1 year posttransplant with an estimated glomerular filtration rate (eGFR) of 5.36 mL/min per 1.73 m² greater than patients remaining on CNI (p = 0.0005) and the longer-term results (> 1 year) of renal function was identical to that of 1 year. There was not a substantial difference in graft loss, mortality, and the occurrence of adverse events (AEs) or serious AEs. However, the risks of acute rejection and trial termination due to AEs with everolimus are respectively 1.82 and 2.63 times greater than patients staying on CNI at 1 year posttransplant (p = 0.02, p = 0.03, respectively). Further, those patients who converted to everolimus had a substantially greater risk of anemia,

hyperlipidemia, hypercholesterolemia, hypokalemia, proteinuria, stomatitis, mouth ulceration, and acne.

Conclusions

Conversion from CNI to everolimus after kidney transplantation is associated with improved renal function in the first 5 years posttransplant but increases the risk of acute rejection at 1 year posttransplant and may not be well endured.

Introduction

Kidney transplantation is the treatment of choice for most patients with end-stage renal disease. Strategies to increase donor organ availability and to prolong the transplanted kidney's survival have become priorities in kidney transplantation. Immunosuppressive therapy is essential and significant in this respect, nevertheless, but choosing the best suitable immunosuppressive therapy is still fairly complex. The calcineurin inhibitors (CNIs) are the principal components of immunosuppressive therapy after kidney transplantation and have made a major contribution to current long-term transplant outcomes[1, 2]. Meanwhile, tacrolimus has been recommended as a first-line agent for kidney transplantation recipients in kidney disease improving global outcomes (KDIGO) in 2009 [3]. However, CNIs are associated with a number of potentially serious side effects, including nephrotoxicity, diabetes, hypertension, and neurotoxicity that contribute to morbidity and mortality after transplantation [4-8]. Upon the fifth year, 90% of grafts revealed evidence of CNI-related lesions, and whereas early acute nephrotoxicity is typically reversible with CNI minimization; chronic lesions cannot be altered once initiated [5]. Furthermore, combined therapy with tacrolimus and mycophenolic acid may be associated with a particularly higher risk of BK infection [9], which is the significant and dangerous factor in the failure of the transplanted kidney. Therefore, diminishing or even eliminating CNI has become the focus of further optimization of immunosuppressive therapy in renal transplantation.

Everolimus (EVR), a mammalian target of the rapamycin inhibitor (mTORi), works 100 times greater than ciclosporin. It is part of a distinguished group of immunosuppressive drugs that have a divergent mode of action to that of CNIs even though they bind to the identical intracellular immunophilin as tacrolimus, namely, FKBP12. The mTORi/FKBP12 complex binds to and hinders the TORC1 complex, preventing proliferation of numerous cell types, such as T-lymphocytes, alloprimed B cells, and CD8 TAb-supp cells [10, 11]. The side effect profile of EVR is different from that of CNI and includes impaired wound healing, mouth ulcers, stomatitis, arthralgia, hyperlipidemia, and anemia [12-15]. The KDIGO instructions suggest that EVR should not be initiated prior to surgical wounds healing [3]. However, EVR may improve renal function and reduce the occurrence of malignancy, which makes it an attractive alternative to CNIs for maintenance therapy after kidney transplantation [16, 17], even though it does cause glomerular disease in some patients resulting in marked proteinuria [18]. A number of randomized controlled trials (RCTs) have examined the potential benefits of introducing EVR after kidney transplantation using a variety of protocols that differ with respect to the timing and mode of conversion to EVR, whether CNIs are eliminated or diminished, and in the level of baseline renal function at the time of EVR introduction. Such studies have provided conflicting results on the efficacy and side effect profile of EVR.

Up to now, there is no published work that systematically reviews EVR used as a maintenance therapy in renal transplant recipients. We undertook the meta-analysis of randomized controlled trials (RCTs) to systematically identify and summarize the current available evidence of the short- and long-term benefits and harm of conversion from CNI to EVR-based maintenance immunosuppression for kidney transplant recipients.

Materials and Methods

Information Sources and Search Strategies

A systematic literature search was performed using Pubmed/MEDLINE, OVID/EMBASE, and Cochrane Library covering the period from the inception of the database until March 2016 using a predefined algorithm (<u>S1 Table</u>) without language restrictions. References included in pertinent systematic reviews were also screened.

Eligibility Criteria

Every RCT examining the conversion from CNI to EVR-based maintenance immunosuppression in adult isolated kidney transplantation was assessed. Research was judged to be suitable if they examined sudden or slow conversion to EVR, in the initial or later kidney transplant recipients, regardless of the time posttransplantation and baseline renal function. Trials that were deemed suitable included those in which the intervention (conversion to EVR) and reference (CNI continuation) groups were administered more maintenance immunosuppression comprising antimetabolites (mycophenolate or azathioprine) and steroids. Observational and uncontrolled studies, those examining children, those in which the participants obtained other solid organs plus a renal transplantation, and animal trials were eliminated.

Data Extraction

Data removal was done separately by 2 investigators (Jinyu Liu and Ruxu You) and discrepancies were resolved by a third investigator (Juan Li). We documented data on trial characteristics and demographics, such as authors' names, year published, journal's name, sample size per-arm, population characteristics, indications, CNI dose, EVR characteristics (bolus and maintenance dose, mode of administration, timing of intervention, sudden stopping or tapering of prior immunosuppression) and length of the trial. The main meta-analysis result was renal function (estimated Glomerular Filtration Rate, eGFR) and secondary results were acute rejection, graft loss, patient survival and AEs. All of the results were investigated 1 year posttransplantation; long-term results, if available, were also investigated.

Assessment of Methodological Quality

The methodological quality of the included trials and the risk of bias were evaluated with elements from the Cochrane Collaboration's tool for determining the risk of bias [19, 20]. The domains implemented in the current systematic review concerned randomization and allocation concealment (selection bias), blinding (performance and detection bias), loss to followup, and keeping to the intention-to-treat principle (attrition bias), selective reporting (reporting bias), and other biases. We decided to put forth the meta-analysis of every one of the studies while offering a synopsis of the risk of bias across trials.

Data Synthesis and Statistical Analysis

For every study we recovered, we computed the crude risk ratio (RR) estimates and corresponding 95% confidence intervals (CI) for the evaluated binary results (acute rejection, graft

loss, patient survival, and AEs), and the mean difference (MD) and corresponding standard deviation (SD) for the evaluated continuous outcomes (GFR). The existence of statistically significant heterogeneity was evaluated using the Cochran Q test and the degree of the observed heterogeneity was evaluated by the I² (ranging from 0% to 100%). A Cochran's Q p < 0.10 was considered to show significant heterogeneity, and analysis was undertaken using the random effects model. Otherwise, the fixed effects model was used. Summary effect estimates for renal function, acute rejection, graft loss, and patient survival were set on intention-to-treat (ITT) groups as identified in the study publications; further, all qualified trials documented AEs in the safety population and, therefore, summary effect sizes for AEs were approximated using safety populations as stated in the study publications.

Pre-determined subgroup analyses were done to investigate origins of heterogeneity; trials were stratified by time of conversion to EVR (early versus late conversion, defined as ≤ 3 months after transplantation) or by the duration to follow up. Sensitivity analyses were done based on to the protocol examining GFR or acute rejection. To additionally determine moderators of the noted effect estimates, a meta-regression analysis was performed to account for baseline renal function. Publication bias was investigated using funnel plots. Where information was missing we contacted the researchers to seek the pertinent information. Analyses were done in RevMan 5.3 (Cochrane Collaboration, 2014) and STATA 10 (STATA Corp., College Station, TX). All p values were 2-tailed. The study is documented based on the favored reporting items for systematic reviews and meta-analyses checklist [21].

Results

Description of Studies

We identified 3,129 possibly relevant references (PubMed, 673; Ovid, 1801; Cochrane Library, 643; and other sources, 12). After examining the titles and abstracts and eliminating identical publications, 67 possibly qualified articles were determined. Fifteen reports [22–36] of 11 trials, including a total of 1,633 randomized patients, were chosen for inclusion in the meta-analysis (Fig 1). One of these trials was available in abstract form only by Wolfgang et al 2012 [31]. Nine trials compared EVR to cyclosporine, and 2 trials compared EVR to CNI (cyclosporine or tacrolimus).

All of the incorporated studies were designed to determine the safety and efficacy of conversion from CNI to EVR in adult kidney transplant patients. Design characteristics, immunosuppression regimens, and reported results for every study are summarized in Table 1. The median sample size was 96 participants (interquartile range, 13–337), and the median treatment duration was 19.5 months (with a minimum of 6 months and a maximum of 60 months). All of the trials stated renal function, acute rejection, graft loss, patient survival, and AEs. GFR was estimated by Nankivell formula in 5 trails [24–27, 29–33], modification of diet in renal disease (MDRD) formula in 3 trails [22, 34, 35], and was unclear in 3 studies [23, 28, 36]. Early conversion to EVR (defined as \leq 3 months after transplantation) was evaluated in 5 studies [22, 23, 25, 28, 30, 31], whereas 6 studies evaluated the late conversion to EVR [24, 26, 27, 31, 32–36]. There was a difference in baseline renal function, both within and between the trials, but most of the patients had mild or moderate renal dysfunction at the time of randomization.

Risk of Bias in Included Studies

The Cochrane Collaboration tool was implemented to evaluate the risk of bias (Fig 2 and S1 Fig). All trials were randomized, but 3 trials gave no indication of the allocation method used. Ten trials were open-label and only 1 trial reported blinding investigators [35]. Attrition was



Fig 1. Flow chart of included studies in the meta-analysis.

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adequately reported in 10 trials and was generally low (< 20%). Intention-to-treat analysis was confirmed for 6 trials and was unclear for the remaining 5 trials.

Assessed Outcomes and Evidence Synthesis

Renal function. Renal function was reported at 1, 2, 3, and 5 years after transplantation by 8, 5, 3 and 2 included studies, respectively. In the ITT analysis, patients converted to EVR had significantly better renal function at 1, 2, 3, and 5 years after transplantation compared to patients remaining on CNI [(MD, 5.36; 95% CI, 2.32–8.39; I², 51%), (MD, 6.91; 95% CI, 3.04– 10.79; I², 32%), (MD, 6.65; 95% CI, 1.61–11.70; I², 53%), and (MD, 6.50; 95% CI, 2.38–10.63; I², 0%); Fig 3A]. When studies were stratified according to the time of conversion to EVR (early versus late conversion, defined as \leq 3 months after transplantation), there was a statistically significant trend toward a more favorable GFR difference between EVR and CNI groups in the late conversion trials (MD, 9.42; 95% CI, 6.13–12.70; I², 0%) compared to the early conversion trials (MD, 3.65; 95% CI, 0.70–6.60; I², 27%), with reduction in heterogeneity (Fig 3B).

Table 1. Characteristics of included studies.

Study ID	Study	Time to		Immur	nosuppressive		follow						
	design	conversion since transplant	Method of switching	Pre- conversion	Intervention	Comparator	CNI in the interventionarm after switching	GFR	Graft loss	Mortality	Acute rejection	AEs	up (m)
Wolfgang 2012 [31]	a prospective open-label multicenter RCT (N = 337)	3 m	Unclear	CsA, EC-MPS, steroids	EVR: Target level (5– 10ng/ml), EC-MPS, steroids (N = 171)	CsA, EC-MPS, steroids (N = 166)	Unclear	V	V	V	V	V	12
Thibault 2016 [22]	a prospective multicenter open-label RCT (N = 13)	within 24 h	No conversion	No	EVR: Bolus (6mg/d), Target level (6–10ng/ml), EC-MPS, steroids (N = 7)	CsA, EC-MPS, steroids (N = 6)	None	V	V	V	V	V	6
Seckinger 2008 [35]	a prospective single-center RCT (N = 39)	6 m	Tapered	CsA, MPA, steroids	EVR: Target level (6– 10ng/ml), MPA, steroids (N = 20)	CsA, MPA, steroids (N = 19)	CsA	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	15
Budde 2011 [<u>33]</u>	a prospective multicentrer parallel- group RCT (N = 300)	4.5 m	Tapered	CsA, EC-MP, steroids	EVR: Bolus (1.5mg/d), Target level (6–10ng/ml), EC-MPS, steroids (N = 154)	CsA, EC-MPS, steroids (N = 146)	CsA	V	V	\checkmark	\checkmark	\checkmark	12
Mjörnstedt 2012 [30]	an open- label multicenter RCT (N = 202)	7 w	Abrupt	CsA, EC-MP, steroids	EVR: Bolus (3mg/d), Target level (6–10ng/ml), EC-MPS, steroids (N = 102)	CsA, EC-MPS, steroids (N = 100)	None	V	V	V	V	V	12
Rostaling 2015 [23]	a prospective open-label multicenter RCT (N = 194)	3 m	Tapered	CsA, EC-MPS, steroids	EVR: Bolus (3mg/d), Target level (6–10ng/ml), EC-MPS, steroids (N = 96)	CsA, EC-MPS, steroids (N = 98)	CsA	V	V	V	V	V	12
Chadban 2014 [28]	a prospective multinational open-label RCT (N = 96)	2 w	Tapered	CsA, MPA, steroids	EVR: Target level (8– 12ng/ml), steroids (N = 48)	CsA, MPA, steroids (N = 47)	CsA	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	12
Budde 2015 [27]	an open- label prospective parallel- group RCT (N = 93)	> 6m	Tapered	CsA or TaC, EC-MPS, steroids	EVR: Bolus (1.5mg/d), Target level (6–10ng/ml), EC-MPS, steroids (N = 46)	CsA or TaC, EC-MPS, steroids (N = 47)	CsA or TaC	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	12
Bemelman 2009 [34]	a prospective multicenter open-label RCT (N = 77)	6 m	Tapered	CsA, MPS, steroids	EVR: Target AUC12(150 mg·h/L), EC-MPS, steroids (N = 38)	CsA, MPS, steroids (N = 39)	CsA	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	24

(Continued)

Table 1. (Continued)

Study ID	Study	Time to		Immur	nosuppressive	dosing regime			Ou	itcomes re	ported		follow
	design	conversion since transplant	Method of switching	Pre- conversion	Intervention	Comparator	CNI in the interventionarm after switching	GFR	Graft loss	Mortality	Acute rejection	AEs	up (m)
Holdaas 2011 [32]	an open- label multicenter RCT (N = 250)	6 m	Tapered	CsA or TaC, MPA, Aza, steroids	EVR: Target level (6– 10ng/mL), MPA, Aza, steroids (N = 127)	CsA or TaC, MPA, Aza, steroids (N = 123)	CsA or TaC	V	\checkmark	V	V	V	24
Kihm 2009 [36]	a prospective single-center RCT (N = 32)	12 m	Tapered	CsA, MPA, steroids	EVR: Bolus (3mg/d), Target level (6–10ng/ml), MPA, steroids (N = 17)	CsA, MPA, steroids (N = 15)	CsA	V	V	V	~	\checkmark	24
Budde 2012 [29]	a prospective multicentrer parallel- group RCT (N = 300)	4.5 m	Tapered	CsA, EC-MPS, steroids	EVR: Bolus (1.5mg/d), Target level (6–10ng/ml), EC-MPS, steroids (N = 154)	CsA, EC-MPS, steroids (N = 146)	CsA	V	V	~	~	\checkmark	36
Mjörnstedt 2015 [25]	an open- label multicenter RCT (N = 202)	7 w	Abrupt	CsA, EC-MPS, steroids	EVR:Bolus (3mg/d), Target level (6–10ng/ml), EC-MP, steroids (N = 102)	CsA, EC-MPS, steroids (N = 100)	None	V	V	V	~	\checkmark	36
Budde (ZEUS) 2015 [24]	a prospective multicentrer parallel- group RCT (N = 300)	4.5 m	Tapered	CsA, EC-MPS, steroids	EVR: Bolus (1.5mg/d), Target level (6–10ng/ml), EC-MPS, steroids (N = 154)	CsAEC-MPS steroids (N = 146)	CsA	V	V	~	~	V	60
Budde (APOLLO) 2015 [26]	an open- label prospective parallel- group RCT (N = 93)	>6 m	Tapered	CsA or TaC, EC-MPS, steroids	EVR: Bolus (1.5mg/d), Target level (6–10ng/ml), EC-MPS, steroids (N = 46)	CsA or TaC, EC-MPS, steroids (N = 47)	CsA or TaC	\checkmark	V	√	√	\checkmark	60

CsA, cyclosporine A; EVR, everolimus; TaC, tacrolimus; Aza, azathioprine; MPA, mycophenolic acid; CNI, calcineurin inhibitor; EC-MPS, enteric-coated mycophenolate sodium; GFR, glomerular filtration rate; AEs, adverse effects; RCT, randomized controlled trail; h, hour; m, month; w, week.

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To additionally confront heterogeneity for renal function at 1 year after conversion to EVR, sensitivity analyses were done using the estimating procedures of GFR. We performed sensitivity analysis excluding 2 trials [22, 35] that reported GFR estimates based on MDRD and a similar effect to the original meta analysis was observed (MD, 5.18; 95% CI, 2.07–8.30; I², 56%). We performed additional sensitivity analysis excluding 2 trials [19, 24] that did not specify just which methods used to estimate GFR and a similar effect to the original meta analysis was observed (MD, 6.30; 95% CI, 3.06–9.53; I², 42%). Further, meta-regression analyses taking into consideration the baseline GFR calculations revealed that the baseline renal function did not have a substantial impact on the difference in renal function between EVR and CNI groups 1 year posttransplantation.



Fig 2. Quality of included studies in the meta-analysis.

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PLOS ONE

Acute rejection. Every one of the included studies added to the meta-analysis examining the link between conversion from CNI to EVR and acute renal allograft rejection. Every one of the trials implemented the description of biopsy-proven acute rejection (BPAR), excluding the trial by Seckinger et al [35]. In contrast to CNI maintenance, conversion to EVR was linked with a greater risk of documented acute rejection 1 year posttransplantation (RR, 1.82; 95% CI, 1.11–2.99; I², 43%; Fig 4A). Analysis based on a definition of BPAR showed similar findings (RR, 1.91; 95% CI, 1.12-3.25; I², 50% for patients converted to EVR). However, patients converted to EVR had a similar risk of acute rejection compared with the patients remaining on CNI up to 2, 3, and 5 years after transplantation [(RR, 1.55; 95% CI, 0.35–6.82; I², 14%), (RR, 1.62; 95% CI, 0.67–3.92; I², 71%), and (RR, 1.85; 95% CI, 0.94–3.65; I², 0%); Fig 4A]. Moreover, we performed subgroup analysis between early conversion (between 0~3 months) and late conversion (> 3 months), and the subgroup analysis showed that early conversion to EVR compared to CNI maintenance was associated with a significantly higher risk of acute rejection up to 1 year after transplantation. In contrast, late conversion to EVR compared to CNI maintenance had a higher risk of acute rejection than CNI maintenance and it did not reach statistical significance (Fig 4B). In total, subgroup analyses did not show statistical heterogeneity among the 2 subgroups ($I^2 = 0\%$, p = 0.59).

Renal allograft loss and mortality. There were 3 trials [23, 28, 31] that added to the meta-analysis examining the link between conversion from CNI to EVR and renal allograft loss while in the 5 other trials [22, 27, 30, 33, 35], no renal allografts were lost up to the first year posttransplantation. Patients converted to EVR had a similar risk of graft loss compared with the patients remaining on CNI (RR, 1.43; 95% CI, 0.44–4.68; I², 37%; Fig 5). Overall, 6 (0.93%) patients in the EVR group and 4 (0.93%) patients in the CNI group lost their graft within the first year after transplantation. There was no significant difference for long-term graft loss as well.

Mortality up to 1 year posttransplantation was reported in 8 studies. Altogether, 4 (0.62%) patients in the EVR group and 6 (0.95%) patients in the CNI group died within the first year after transplantation. There were no distinctions in mortality among patients who converted

А	Eve	erolim	IS		CNI			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Tota	Weight	IV. Random, 95% C	I IV. Random. 95% CI
1.1.1 One year							-		
Chadban 2014	65.1	15.4	49	67.1	18.2	47	11.7%	-2.00 [-8.76, 4.76]	-
Klemens 2011	71.8	16.5	143	61.9	15.9	137	19.4%	9.90 [6.10, 13.70]	
Klemens 2015	63.8	19.8	44	58.2	16.6	44	10.1%	5.60 [-2.03, 13.23]	
Mjörnstedt 2012	73.9	17.1	102	71.3	17.6	100	16.4%	2.60 [-2.19, 7.39]	
Rostaing 2015	60.1	20	96	53.5	16.9	98	15.3%	6.60 [1.38, 11.82]	· · · · · · · · · · · · · · · · · · ·
Seckinger 2008	70.1	17	17	54.8	17.8	10	4.2%	15.30 [1.62, 28.98]	
Thibault 2016	54.4	10.5	174	55.7	15.7	400	3.7%	-1.30 [-16.08, 13.48]	+
Wolfgang 2012 Subtotal (95% CI)	68.6	17.6	1/1	63	18.2	166	100.0%	5.60 [1.78, 9.42]	· · · · · · · · · · · · · · · · · · ·
Hotorogonoity: Tau2 = 9.62	Chi2 -	14 16	df = 7/0	2 - 0.04	5): 12 -	51%	100.0%	5.50 [2.52, 6.59]	
Test for overall effect: 7 = 1	3 46 (P =	0.000	ui - 7 (r 5)	0.03), i [_] =	5170			
	0.40 (i =	0.000	5)						
1.1.2 Two years									
Bemelman 2009	55	20	38	44	15	39	17.5%	11.00 [3.09, 18.91]	
Holdaas 2011	48	22	110	46	20.4	112	27.7%	2.00 [-3.58, 7.58]	+
Kihm 2009	96.9	22.6	13	73.2	37.3	11	2.3%	23.70 [-1.53, 48.93]	
Klemens (APOLLO) 2015	63.9	20.2	44	57.4	16.1	44	18.4%	6.50 [-1.13, 14.13]	-
Klemens 2012	69.2	19	123	61.3	17.2	118	34.1%	7.90 [3.33, 12.47]	
Subtotal (95% CI)			328			324	100.0%	6.91 [3.04, 10.79]	•
Heterogeneity: Tau ² = 6.01	; Chi² = 5	5.87, d	f = 4 (P	= 0.21)	; 2 = 3	32%			
Test for overall effect: Z = 3	3.50 (P =	0.000	5)						
1.1.3 Three years									-
Klemens (APOLLO) 2015	63.8	19.8	44	58.2	15.6	44	26.6%	5.60 [-1.85, 13.05]	
Klemens 2012	72.2	20	116	61.6	16.7	111	40.3%	10.60 [5.81, 15.39]	
Mjornstedt 2015	60.1	19.7	89	57.4	20.2	79	33.1%	2.70 [-3.35, 8.75]	▲
Subtotal (95% CI)	O. Chiz -	4.94	249 df = 0 //	4		234	100.0%	0.05 [1.01, 11.70]	•
Test for everall effect: 7 = 10.5	0; Chr =	4.24,	ui = ∠ (i	= 0.14	2); 1- =	53%			
	2.36 (F =	0.010	,						
1.1.4 Five years									
Klemens (APOLLO) 2015	65	22.1	44	57.9	16.5	44	25.6%	7.10 [-1.05, 15,25]	
Klemens (ZEUS) 2015	66.7	17.4	97	60.4	16.8	100	74.4%	6.30 [1.52, 11.08]	
Subtotal (95% CI)			141			144	100.0%	6.50 [2.38, 10.63]	•
Heterogeneity: Tau ² = 0.00	; Chi ² = (0.03, d	f = 1 (P	= 0.87)	; 2 = (0%			
Test for overall effect: Z = 3	3.09 (P =	0.002)						
									-100 -50 0 50 100
									Everolimus CNI
Test for subaroup difference	es: Chi ²	= 0.48	. df = 3	(P = 0.9)	92), I² :	= 0%			
D									
D	Everoli	imus		CN	11			Mean Difference	Mean Difference
Study or Subgroup M	ean S	SD To	otal Me	ean S	SD T	otal	Weight	IV, Random, 95% CI	IV. Random, 95% Cl
1.2.1 Early conversion									
Chadban 2014	65.1 15	.4	49 6	7.1 18	3.2	47	11.7%	-2.00 [-8.76, 4.76]	T
Mjörnstedt 2012	73.9 17	.1 1	02 7	1.3 17	7.6	100	16.4%	2.60 [-2.19, 7.39]	T.
Rostaing 2015	60.1 2	20	96 5	3.5 16	5.9	98	15.3%	6.60 [1.38, 11.82]	
Thibault 2016	54.4 10	.5	7 5	5.7 15	5.7	6	3.7%	-1.30 [-16.08, 13.48]	
Wolfgang 2012	68.6 17	.6 1	71	63 18	3.2	166	19.3%	5.60 [1.78, 9.42]	
Subtotal (95% CI)		4	25			417	66.3%	3.65 [0.70, 6.60]	•
Heterogeneity: Tau ² = 3.0	2; Chi ² =	5.48,	df = 4 (P = 0.2	24); l²	= 27%	b		
Test for overall effect: Z =	2.42 (P	= 0.02	2)						
4001-4									
1.2.2 Late conversion		_							-
Klemens 2011	71.8 16	.5 1	46 6	1.9 15	5.9	137	19.4%	9.90 [6.12, 13.68]	
Klemens 2015	63.8 19	.8	44 5	8.2 16	5.6	44	10.1%	5.60 [-2.03, 13.23]	
Seckinger 2008	70.1 1	17	17 5	4.8 17	7.8	10	4.2%	15.30 [1.62, 28.98]	
Subtotal (95% CI)		2	07	-		191	33.7%	9.42 [6.13, 12.70]	•
Heterogeneity: Tau ² = 0.0	0; Chi ² =	1.73,	df = 2 (P = 0.4	12); l²	= 0%			
Test for overall effect: Z =	5.62 (P	< 0.00	0001)						
Total (05% CI)			22		3	600	100 09/	E 26 [2 20 0 40]	A
I otal (95% CI)		6	52			000	100.0%	5.30 [2.32, 8.40]	▼

Fig 3. A. Everolimus versus calcineurin inhibitor; mean GFR up to 1–5 years after transplantation stratified by study duration. B. Everolimus versus calcineurin inhibitor; mean GFR up to 1 year after transplantation stratified by time to conversion.

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Everolimus CNI

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Test for overall effect: Z = 3.46 (P = 0.0005)

Heterogeneity: Tau² = 8.65; Chi² = 14.21, df = 7 (P = 0.05); I² = 51%

Test for subgroup differences: $Chi^2 = 6.55$, df = 1 (P = 0.01), $I^2 = 84.7\%$

100

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А	Everolii	nus	CNI			Risk Ratio		Ri	sk Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H. Random, 95% C	I	M-H, Ra	andom, 95% (CI	
1.6.1 One year					-						
Chadban 2014	15	49	6	47	17.3%	2.40 [1.02, 5.66]					
Klemens 2011	15	154	5	146	14.8%	2.84 [1.06, 7.63]				-	
Klemens 2015	0	46	0	47		Not estimable					
Mjörnstedt 2012	15	102	14	100	21.6%	1.05 [0.54, 2.06]					
Rostaing 2015	24	96	5	98	16.0%	4.90 [1.95, 12.31]					
Seckinger 2008	2	20	2	19	5.9%	0.95 [0.15, 6.08]					
Thibault 2016	1	7	1	6	3.4%	0.86 [0.07, 10.96]			-		
Wolfgang 2012	16	171	13	166	21.0%	1.19 [0.59, 2.41]					
Subtotal (95% CI)		645		629	100.0%	1.82 [1.11, 2.99]			•		
Total events	88		46								
Heterogeneity: Tau ² = 0.18;	Chi ² = 10.	46, df =	6 (P = 0.	11); l ² :	= 43%						
Test for overall effect: Z = 2.	38 (P = 0.	02)	,	,.							
		,									
1.6.2 Two years											
Bemelman 2009	0	38	1	39	19.9%	0.34 [0.01, 8.14]		-		-	
Holdaas 2011	7	127	3	123	80.1%	2.26 [0.60, 8.54]				_	
Kihm 2009	0	17	0	15		Not estimable					
Subtotal (95% CI)		182		177	100.0%	1.55 [0.35, 6.82]		-			
Total events	7		4								
Heterogeneity: Tau ² = 0.25;	Chi ² = 1.1	6, df = '	1 (P = 0.2	8); l ² =	14%						
Test for overall effect: Z = 0.	58 (P = 0.	56)		,.							
	,	,									
1.6.3 Three years											
Klemens 2012	20	154	7	146	42.7%	2.71 [1.18, 6.21]					
Mjörnstedt 2015	27	102	24	100	57.3%	1.10 [0.69, 1.77]					
Subtotal (95% CI)		256		246	100.0%	1.62 [0.67, 3.92]					
Total events	47		31								
Heterogeneity: Tau ² = 0.30;	Chi ² = 3.4	8, df = 1	1 (P = 0.0	6); l ² =	71%						
Test for overall effect: Z = 1.	07 (P = 0.	28)									
1.6.4 Five years											
Klemens (APOLLO) 2015	1	46	0	47	4.6%	3.06 [0.13, 73.33]					
Klemens (ZEUS) 2015	21	154	11	146	95.4%	1.81 [0.90, 3.62]					
Subtotal (95% CI)		200		193	100.0%	1.85 [0.94, 3.65]					
Total events	22		11								
Heterogeneity: Tau ² = 0.00;	Chi ² = 0.1	0, df = '	1 (P = 0.7	5); l² =	0%						
Test for overall effect: Z = 1.	79 (P = 0.	07)									
							0.01	0.1	1	10	100
							0.01	Everolim	us CNI	10	100
									10 0111		

Test for subaroup differences: $Chi^2 = 0.10$. df = 3 (P = 0.99). $I^2 = 0\%$

В	Everoli	nus	CNI			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H. Random. 95% Cl	M-H, Random, 95% Cl
1.7.1 Early conversion	n						
Chadban 2014	15	49	6	47	17.3%	2.40 [1.02, 5.66]	
Mjörnstedt 2012	15	102	14	100	21.6%	1.05 [0.54, 2.06]	
Rostaing 2015	24	96	5	98	16.0%	4.90 [1.95, 12.31]	
Thibault 2016	1	7	1	6	3.4%	0.86 [0.07, 10.96]	
Wolfgang 2012	16	171	13	166	21.0%	1.19 [0.59, 2.41]	
Subtotal (95% CI)		425		417	79.3%	1.77 [0.96, 3.28]	-
Total events	71		39				
Heterogeneity: Tau ² = 0	0.25; Chi ²	= 9.03,	df = 4 (P	= 0.06)	; I ² = 56%		
Test for overall effect: 2	z = 1.83 (F	P = 0.07)				

Fig 4. A. Everolimus versus calcineurin inhibitor; any rejection up to 5 years after transplantation stratified by study duration. B. Everolimus versus calcineurin inhibitor; any rejection up to 1 year after transplantation stratified by time to conversion.

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	Everolin	nus	CNI			Risk Ratio		Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C		M-H, Fixe	ed, 95% Cl		
1.5.1 one year											
Chadban 2014	0	49	2	47	56.0%	0.19 [0.01, 3.90]	←	-			
Klemens 2011	0	154	0	146		Not estimable					
Klemens 2015	0	46	0	47		Not estimable					
Mjörnstedt 2012	0	102	0	100		Not estimable					
Rostaing 2015	5	96	1	98	21.7%	5.10 [0.61, 42.89]					
Seckinger 2008	0	20	0	19		Not estimable					
Thibault 2016	0	7	0	6		Not estimable					
Wolfgang 2012	1	171	1	166	22.3%	0.97 [0.06, 15.39]					
Subtotal (95% CI)		645		629	100.0%	1.43 [0.44, 4.68]					
Total events	6		4								
Heterogeneity: Chi ² = 3.16, d	lf = 2 (P =	0.21);	l² = 37%								
Test for overall effect: Z = 0.6	60 (P = 0.	55)									
1.5.2 Two years											
Bemelman 2009	0	38	0	39		Not estimable		_			
Holdaas 2011	4	127	8	123	100.0%	0.48 [0.15, 1.57]			_		
Kihm 2009	0	17	0	15		Not estimable					
Subtotal (95% CI)		182		177	100.0%	0.48 [0.15, 1.57]					
Total events	4		8								
Heterogeneity: Not applicable	e										
Test for overall effect: Z = 1.2	21 (P = 0.:	23)									
1.5.3 Three years								_			
Klemens 2012	2	154	2	146	100.0%	0.95 [0.14, 6.64]					
Mjörnstedt 2015	0	102	0	100		Not estimable					
Subtotal (95% CI)		256		246	100.0%	0.95 [0.14, 6.64]					
Total events	2		2								
Heterogeneity: Not applicable	е										
Test for overall effect: Z = 0.0	05 (P = 0.9	96)									
1.5.4 Five years											
Klemens (APOLLO) 2015	3	46	1	47	24.3%	3.07 [0.33, 28.40]					
Klemens (ZEUS) 2015	4	154	3	146	75.7%	1.26 [0.29, 5.55]					
Subtotal (95% CI)		200		193	100.0%	1.70 [0.51, 5.70]					
Total events	7		4								
Heterogeneity: Chi ² = 0.42, d	lf = 1 (P =	0.52);	l² = 0%								
Test for overall effect: Z = 0.8	36 (P = 0.3	39)									
							L				1
							0.01	0.1	1	10	100
								Everolimus	CNI	ener (The	
lest for subgroup differences	s: Chi ² = 2	.56, df	= 3 (P = (J.47). I	² = 0%						

Fig 5. Everolimus versus calcineurin inhibitor; graft loss within 5 years after transplantation stratified by study duration.

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to EVR and those who continued on CNI (RR, 0.70; 95% CI, 0.22–2.18; I^2 , 0%; Fig 6A). Risk ratios (RRs) and heterogeneity were alike when studies were stratified based on the time of conversion to EVR (Fig 6B). There was no significant difference for long-term mortality as well.

Treatment failure. Five studies [27, 28, 30, 31, 33] reported treatment failure 1 year post-transplantation. Overall, 64 (18.19%) patients in the EVR group and 29 (8.43%) patients in the CNI group failed within the first year after transplantation. There were no differences in

А	Everoli	mus	CN			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	M-H, Fixed, 95% Cl
1.3.1 One year							
Chadban 2014	0	49	1	47	21.6%	0.32 [0.01, 7.66]	
Klemens 2011	0	154	1	146	21.7%	0.32 [0.01, 7.70]	
Klemens 2015	1	46	1	47	13.9%	1.02 [0.07, 15.85]	
Mjornstedt 2012	2	102	1	100	14.2%	1.96 [0.18, 21.28]	
Rostaing 2015 Sockinger 2008	0	96	0	98		Not estimable	
Thibault 2016	0	20	0	6		Not estimable	
Wolfgang 2012	1	171	2	166	28.6%	0.49 [0.04, 5.30]	
Subtotal (95% CI)		522	-	506	100.0%	0.70 [0.22, 2.18]	
Total events	4		6				
Heterogeneity: Chi ² = 1.35, c	if = 4 (P =	= 0.85);	l² = 0%				
Test for overall effect: Z = 0.	62 (P = 0.	.54)					
4.2.2.7							
1.3.2 Two years		~~~	0	20			
Bemeiman 2009	2	127	0	122	100.0%	6 79 10 25 120 041	
Kihm 2009	0	127	0	123	100.0%	0.70 [0.35, 129.94] Not estimable	-
Subtotal (95% CI)	0	182	0	177	100.0%	6.78 [0.35, 129,94]	
Total events	3		0				
Heterogeneity: Not applicabl	е						
Test for overall effect: Z = 1.	27 (P = 0.	.20)					
1.3.3 Three years							
Klemens 2012	3	154	3	146	37.9%	0.95 [0.19, 4.62]	
Mjörnstedt 2015	2	102	5	100	62.1%	0.39 [0.08, 1.97]	
Subtotal (95% CI)		256		246	100.0%	0.60 [0.20, 1.82]	
Hotorogonoity: Chi2 = 0.59	C +f - 1 /D -	0 44).	0/12 - 0%				
Test for overall effect: $7 = 0.59$, 0	P = 1 (P = 0)	37)	1 0%				
	00 (1 = 0.	.01)					
1.3.4 Five years							
Klemens (APOLLO) 2015	2	46	4	47	56.2%	0.51 [0.10, 2.65]	
Klemens (ZEUS) 2015	4	154	3	146	43.8%	1.26 [0.29, 5.55]	
Subtotal (95% CI)		200		193	100.0%	0.84 [0.29, 2.46]	
Total events	6	-	7				
Heterogeneity: Chi ² = 0.64, c	if = 1 (P =	: 0.42);	l² = 0%				
Test for overall effect: $Z = 0.3$	32(P=0.	.75)					
							0.01 0.1 1 10 100
Test for subaroup difference	s: Chi ² = 2	2.32. df	= 3 (P =).51). P	2 = 0%		Everolimus CNI
В ел	verolimu	s	CNI			Risk Ratio	Risk Ratio
Study or Subgroup Ev	ents To	otal E	vents T	otal V	Veight	M-H, Fixed. 95% CI	M-H, Fixed, 95% CI
1.4.1 Early conversion							
Chadban 2014	0	49	1	47	21.6%	0.32 [0.01, 7.66]	
Mjörnstedt 2012	2	102	1	100	14.2%	1.96 [0.18, 21.28]	
Rostaing 2015	0	96	0	98		Not estimable	
Thibault 2016	0	7	0	6		Not estimable	
Wolfgang 2012	1 1	171	2	166	28.6%	0.49 [0.04, 5.30]	
Subtotal (95% CI)	4	425		417	64.4%	0.76 [0.19, 3.03]	
Total events	3		4				
Heterogeneity: Chi ² = 1.03,	df = 2 (P	P = 0.60); $ ^2 = 0\%$	•			
Test for overall effect: Z = 0).40 (P =	0.69)					
1.4.2 Late conversion							
Klomone 2014	0	154	4	146	21 70/	0 22 [0 04 7 70]	
Kiemens 2011	1	104	1	40	12 0%	1.02 [0.07, 15, 95]	
Seckinger 2009	0	20	0	4/	13.9%	Not estimable	
Subtotal (95% CI)		20	0	212	35.6%	0.59 [0.08 4 38]	
Total events	1		2		00.070	0.00 [0.00, 4.00]	
Heterogeneity: Chi ² = 0.30	df = 1 (P	P = 0.58); $ ^2 = 0\%$				
Test for overall effect: Z = ().51 (P =	0.61)	,, - 07				
Total (95% CI)	6	645		629 1	00.0%	0.70 [0.22, 2.18]	

0.01

0.1

1 Everolimus CNI

Total events 4 6 Heterogeneity: Chi² = 1.35, df = 4 (P = 0.85); l² = 0% Test for overall effect: Z = 0.62 (P = 0.54) Test for subgroup differences: Chi² = 0.04, df = 1 (P = 0.84), l² = 0%

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Fig 6. A. Everolimus versus calcineurin inhibitor; mortality up to 1–5 years after transplantation stratified by study duration. B. Everolimus versus calcineurin inhibitor; mortality up to 1 year after transplantation stratified by time to conversion.

doi:10.1371/journal.pone.0170246.g006

treatment failure between patients converted to EVR and those remaining on CNI (RR, 1.74; 95% CI, 0.79–3.86; I², 67%).

Adverse events. Every one of the trials involved in the meta-analysis documented adverse events (AEs), although there were differences between studies in the nature and incidence of the reported adverse events. The main AEs involved in infection, blood and lymphatic system disorders, gastrointestinal disorders, metabolism and nutrition disorders, and the body as a whole-general disorders. Overall, 427 (94.05%) patients in the EVR group and 403 (90.77%) patients in the CNI group suffered 1 or more AEs within the first year after transplantation. There were no differences in the incidence of AEs or serious AEs between patients converted to EVR and those remaining on CNI [(RR, 1.04; 95% CI, 1.00–1.08; I², 0%), and (RR, 1.07; 95% CI, 0.96–1.21; I², 18%); Fig 7A and 7B]. However, the risk of study discontinuation due to AEs up to 1 year after transplantation was greater in patients converted to EVR than in patients remaining on CNI (RR, 2.63; 95% CI, 1.13–6.15; I², 69%).

А	Everolin	nus	CNI			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	M-H, Fixed, 95% Cl
Chadban 2014	49	49	47	47	11.9%	1.00 [0.96, 1.04]	• •
Klemens 2011	143	154	131	146	32.9%	1.03 [0.96, 1.11]	•
Klemens 2015	44	46	44	47	10.7%	1.02 [0.93, 1.13]	• •
Mjörnstedt 2012	97	102	90	100	22.3%	1.06 [0.98, 1.14]	•
Rostaing 2015	87	96	85	98	20.6%	1.04 [0.94, 1.16]	• •
Thibault 2016	7	7	6	6	1.7%	1.00 [0.76, 1.31]	+
Total (95% CI)		454		444	100.0%	1.04 [1.00, 1.08]	
Total events	427		403				
Heterogeneity: Chi ² = 3	.33, df = 5	5 (P = 0	.65); l ² = (0%			
Test for overall effect: 2	z = 1.82 (F	P = 0.07	")				Everolimus CNI

В	Everolin	nus	CNI			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	M-H, Fixed, 95% Cl
Chadban 2014	33	49	31	47	11.4%	1.02 [0.77, 1.36]	+
Klemens 2011	47	154	50	146	18.6%	0.89 [0.64, 1.24]	-
Klemens 2015	12	46	11	47	3.9%	1.11 [0.55, 2.27]	
Mjörnstedt 2012	55	102	38	100	13.9%	1.42 [1.04, 1.93]	-
Rostaing 2015	46	96	39	98	14.0%	1.20 [0.87, 1.66]	+
Thibault 2016	5	7	2	6	0.8%	2.14 [0.63, 7.29]	
Wolfgang 2012	103	171	102	166	37.4%	0.98 [0.83, 1.16]	• • •
Total (95% CI)		625		610	100.0%	1.07 [0.96, 1.21]	•
Total events	301		273				
Heterogeneity: Chi ² = 7	.33, df = 6	6 (P = 0	.29); 2 =	18%			
Test for overall effect: 2	Z = 1.22 (F	P = 0.22	2)				Everolimus CNI

Fig 7. A. Everolimus versus calcineurin inhibitor; any adverse events up to 1 year after transplantation; B. Everolimus versus calcineurin inhibitor; any serious adverse events up to 1 year after transplantation.

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Fig 8. Pooled risk ratio estimates and 95% confidence intervals for adverse events up to 1 year after transplantation.

doi:10.1371/journal.pone.0170246.g008

Reported adverse events, along with RR estimates and 95% CI up to 1 year after transplantation, are summarized in Fig 8 and S2 Table. Compared with patients on CNI continuation, those converted to EVR had a substantially higher risk of anemia, hyperlipidemia, hypercholesterolemia, hypokalemia, proteinuria, stomatitis, mouth ulceration, acne and a nonstatistically significant trend toward a higher risk of thrombocytopenia. Nevertheless, patients converted to EVR had a lower risk of cytomegalovirus, BK virus, and urinary tract infection, diabetes mellitus, hypertriglyceridemia, malignancy, and cough; however, it did not reach statistical significance (Table 2).

As for long-term safety, we did not perform meta-analysis because there were too few trials within the same duration. The occurrence of AEs and serious AEs was similar in the EVR group versus the CNI group. The most recurrent AEs (> 10%) were anemia, diarrhea, peripheral edema, proteinuria, mouth ulceration, pyrexia, urinary tract infection, upper respiratory

Adverse events	Incidence of adver	p-Value	
	Everolimus	CNI	
anemia	13.92	8.52	0.004
hyperlipidemia	10.15	5.23	0.006
hypercholesterolaemia	11.11	5.59	0.01
hypokalaemia	6.25	1.63	0.02
proteinuria	8.55	3.53	0.007
stomatitis	18.04	1.94	<0.00001
mouth ulceration	15.38	0.75	<0.00001
acne	10.25	1.63	0.0005
thrombocytopenia	5.12	2.23	0.29
cytomegalovirus	8.90	11.59	0.11
BK virus	2.85	4.88	0.14
urinary tract infection	21.48	25.61	0.27
diabetes mellitus	4.92	8.29	0.16
hypertriglyceridaemia	2.16	2.51	0.77
malignancy	5.62	7.08	0.51
cough	6.97	7.35	0.90

Table 2. Incidence of the most common adverse events up to 1 year after transplant.

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tract infection, increased serum creatinine, cough, rash, and acne; the first 5 were substantially elevated in the EVR group than the CNI group (p < 0.05). In addition, leukocyte and platelet counts were decreased and liver enzymes (aspartate aminotransferase and alanine aminotransferase) and lipids (total cholesterol and triglycerides) were substantially elevated in the EVR group versus the CNI group (p < 0.05). No AEs were documented less commonly in the EVR group with a statistical difference.

Discussion

The findings from this meta-analysis establish that conversion from CNI to EVR-based maintenance immunosuppression after kidney transplantation is associated with an increased eGFR (about 10%, p < 0.01) up to 5 years after transplantation. There were no substantial differences in efficacy results between the 2 groups except that recipients who converted to EVR had a higher risk of acute graft rejection up to 1 year after transplantation. In terms of safety and tolerability, results were subordinate in the EVR group in contrast to the CNI group.

In renal transplantation, chronic allograft nephropathy (CAN) continues to be the primary reason for graft loss[37, 38]. CAN is characterized by a gradual decline of renal function with interstitial fibrosis and tubular atrophy, which causes proteinuria, high blood pressure, and a gradual escalation in serum creatinine. GFR is the most commonly utilized marker for the decline in renal function. Further, scientists have proposed that early low-grade proteinuria could offer a different marker to GFR for future risk [39–41]. Two previous meta-analysis [14, 42] evaluated mTORi for primary immunosuppression of kidney transplant recipients and concluded that mTORi showed significantly lower serum creatinine and more improved eGFR than those treated with a CNI. In the present meta-analysis, every one of the RCTs were available for analysis on renal function with GFR as the main endpoint. Our analysis conducted that eGFR was significantly elevated in the EVR group versus the CNI controls: a mean difference of 5.36mL/min per 1.73m² (p = 0.0005) at 1 year and 6.50 mL/min per 1.73m² (p = 0.002) 5 years after transplantation. Given the observed marked heterogeneity (I², 51%) for trials reporting on the effect of EVR conversion on renal function at 1 year after

transplantation, caution is required with respect to the magnitude of the overall estimate for this outcome. Subgroup and sensitivity analyses reinforced the overall conclusion that conversion to EVR was associated with improved eGFR and eliminated heterogeneity as well. Nevertheless, those patients who converted to EVR had a substantially greater risk of proteinuria (8.55% vs 3.53%, p = 0.007) at 1 year after the transplant and 2 patients developed proteinuria >1,000 mg/day post-conversion to everolimus. Offset against the insistent eGFR advantage of conversion to everolimus, this risk would seem to be allowable but because patients with elevated urine protein (e.g.> 150 or > 500 g/day) at the time of conversion seem to be more susceptible to gradual proteinuria after conversion to EVR [43], the transition should be avoided or done with care.

It has been reported that conversion from CNI to mTORi increases the risk of acute rejection [16, 22]. The present meta-analysis showed that conversion to EVR is associated with a higher risk of acute rejection at 1 year after transplantation. However, there was a similar risk of long-term acute rejection (> 1 year). Since the risk for acute rejection is highest in the first 3 months after transplantation [3], we did subgroup analysis between early conversion (between 0 and 3 months) and late conversion (> 3 months). The subgroup analysis showed that early conversion to EVR compared to CNI maintenance was associated with a significantly higher risk of acute rejection up to 1 year after transplantation; in contrast, although late conversion had a higher risk of acute rejection, it did not reach statistical significance. The meta-analysis showed that late versus early conversion to EVR was associated with a trend toward better renal function and a lower risk of acute rejection at 1 year. According to the evidence, late transition may be the better advice. In the 8 trials, only 1 trial abruptly discontinued CNI, so the present analysis did not show a difference in acute rejection between the trials evaluating abrupt and tapered discontinuation of CNI. However, experience in liver transplantation suggests that tapered discontinuation is preferable, especially when EVR conversion is introduced within the first few months of transplantation [44].

Mammalian target of rapamycin inhibitors are associated with a number of well-described side effects [4, 13, 18] that may limit the ability of patients to tolerate them. Our analysis confirmed this, indicating that the risk of study discontinuation due to AEs after EVR conversion was 2.63 times that of the patients maintained on CNIs. Overall, there was a substantial risk of adverse events after conversion to EVR, and this represents a significant barrier for its utility in preserving renal function after kidney transplantation. It was reported that EVR may lead an increased risk of bone marrow suppression [42]. Our analysis showed that conversion to EVR had a significantly higher risk of anemia, and a higher risk of thrombocytopenia, and leucopenia with no significant difference. Therefore, the real harm of EVR-induced thrombocytopenia and leukopenia is unclear from these trials. In addition, our analyses showed that conversion to EVR is associated with an increased risk of hyperlipidemia and hypercholesterolemia, although the requirements for a new statin therapy was not different to patients maintained on CNI. Conversion to EVR significantly increased the risk of stomatitis by 9.30 times (p < 0.00001) and mouth ulceration by 20.51 times (p < 0.00001) than CNI, but the rate of infection was similar to that of patients on CNI maintenance. A significant drawback of treatment with mTORi is the development of proteinuria which may reach the nephrotic range, especially after exposure to high everolimus concentrations [45]; our analysis confirmed the association between EVR and proteinuria, although the reported proteinuria levels were usually mild or less often moderate. We were unable to confirm any benefit in reduced malignancy in these 11 randomized trials, although this has been suggested by other data [46, 47]. The main limitation of this review is that of the contributory trials. Additionally, where graft-focused outcomes were well reported, many of the potentially informative patient-focused adverse outcomes were not reported, not defined, or inconsistently reported.

A longitudinal cohort study included 9,353 kidney transplant recipients showed that mTORi use was associated with a higher risk of all-cause mortality but not allograft loss [48]. The majority of RCTs included in our meta-analysis were not powered to detect a difference in graft or patient survivals, and the true effect of EVR conversion on graft and patient out-comes is still to some extent uncertain. This is especially the case for long-term outcomes, although there are 6 trials included in the meta-analysis reported outcomes beyond 1 year, they scatter in different time periods. As to long-term safety, there are too few studies to perform a meta-analysis. This review focuses on the requirement for follow-up of clinical studies in transplantation to go well beyond the immediate term to 5 years and beyond. This has been shown in renal transplantation in which a coordinated effort between the researchers and a registry of long-term patient follow-up of up to 15 years posttransplantation [49].

There are further possible restrictions in the current meta-analysis. First, we did not incorporate any unpublished evidence, so there may be publication bias in the meta-analysis. Second, even though randomized evidence is safeguarded from selection bias, performance and detection biases could be possible confounders. An approach toward addressing this would be to exempt open-label trials. Ultimately, the majority of the incorporated trials were, by necessity, open-label trials, and so trial exclusion was not a viable choice. Lastly, a group of study parameters could possibly impede our study outcomes. The variation in baseline renal function between studies, and the unmistakable patient characteristics inside separate studies may have added to the noted heterogeneity.

Conclusion

The currently available randomized evidence indicates that conversion from CNI to EVR after kidney transplantation is associated with short and long-term improvements in GFR in a number of studies, but these findings must be balanced against the greater rates of proteinuria, discontinuation due to AEs, and higher incidences of acute rejection up to 1 year posttransplant with the use of EVR as compared with CNI. Moreover, late conversion to EVR might be associated with a trend toward better renal function and a lower risk of acute rejection up to 1 year posttransplantion than early conversion. Additionally, in deciding the optimal immunosuppression strategy for their patients, clinicians should be alert to the increased risk of bone marrow suppression, dyslipidemia, mouth ulceration, and stomatitis after conversion to EVR which, however, has no effect on graft and patient survival.

Supporting Information

S1 Fig. Risk of bias summary: each risk of bias item for each included study. (TIF)

S2 Fig. Funnel plot of pooled efficacy and safety of everolimus comparing with calcineurin inhibitor.

(TIF)

S1 Table. Literature search algorithm. (DOCX)

S2 Table. Effect estimate of all outcomes. (DOCX)

S3 Table. PRISMA checklist. (DOC)

S1 Text. Abbreviations page. (DOCX)S2 Text. Flow diagram. (DOC)

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Author Contributions

Conceptualization: JYL DL JL LZ CLZ KL QLX RXY.

Data curation: JYL JL CLZ.

Formal analysis: JYL KL QLX.

Funding acquisition: JYL.

Investigation: JYL JL RXY.

Methodology: JYL DL RXY.

Project administration: JYL DL JL.

Resources: JYL CLZ RXY.

Software: JYL KL QLX.

Supervision: DL LZ RXY.

Validation: JYL LZ RXY.

Visualization: JYL RXY.

Writing - original draft: JYL.

Writing - review & editing: QLX RXY.

References

- Liu JY, You RX, Guo M, Zeng L, Zhou P, Zhu L, et al. Tacrolimus versus cyclosporine as primary immunosuppressant after renal transplantation: A Meta-Analysis and economics evaluation. Am J Ther. 2016; 23(3): e810–e824. doi: 10.1097/MJT.00000000000150 PMID: 25299636
- Riella LV, Gabardi S, Azzi J. Belatacept and Long-Term outcomes in kidney transplantation. N Engl J Med. 2016; 374(26): 2599–2600.
- SURGEONS OJOT. Kidney disease improving global outcomes (KDIGO) clinical practise guideline for the care of kidney transplantation receipts. American Journal of tansplantation. 2009; 9(Suppl 3): 131– 155.
- Almeida C, Silveira M, de Araújo V, de Lemos L, de Oliveira Costa J, Reis C, et al. Safety of immunosuppressive drugs used as maintenance therapy in kidney transplantation: A systematic review and Meta-Analysis. Pharmaceuticals. 2013; 6(10): 1170–1194. doi: 10.3390/ph6101170 PMID: 24275847
- Nankivell BJ, Borrows RJ, Fung CLS, O'Connell PJ, Chapman JR, Allen RDM. Calcineurin inhibitor nephrotoxicity: Longitudinal assessment by protocol histology. Transplantation. 2004; 78(4): 557–565. PMID: 15446315
- Nankivell BJ, P'Ng CH, O'Connell PJ, Chapman JR. Calcineurin inhibitor nephrotoxicity through the lens of longitudinal histology. Transplantation. 2016; 100(8): 1723–1731. doi: <u>10.1097/TP</u>. 00000000001243 PMID: 27306529

- Borda B, Lengyel C, Várkonyi T, Kemény É, Ottlakán A, Kubik A, et al. Side effects of the calcineurin inhibitor, such as new-onset diabetes after kidney transplantation. Acta Physiologica Hungarica. 2014; 101(3): 388–394. doi: 10.1556/APhysiol.101.2014.3.13 PMID: 25183511
- Chapman JR. Chronic calcineurin inhibitor Nephrotoxicity-Lest we forget. American Journal of Transplantation. 2011; 11(4): 693–697. doi: 10.1111/j.1600-6143.2011.03504.x PMID: 21446974
- Suwelack B, Malyar V, Koch M, Sester M, Sommerer C. The influence of immunosuppressive agents on BK virus risk following kidney transplantation, and implications for choice of regimen. Transplant Rev (Orlando). 2012; 26(3): 201–211.
- Sanchez-Fructuoso AI. Everolimus: An update on the mechanism of action, pharmacokinetics and recent clinical trials. Expert Opin Drug Metab Toxicol. 2008; 4(6): 807–819. doi: 10.1517/17425255.4.6. 807 PMID: 18611120
- Avila CL, Zimmerer JM, Elzein SM, Pham TA, Abdel-Rasoul M, Bumgardner GL. MTOR inhibition suppresses posttransplant alloantibody production through direct inhibition of alloprimed b cells and sparing of CD8+ Antibody-Suppressing t cells. Transplantation. 2016; 100(9): 1898–1906. doi: <u>10.1097/TP.</u>00000000001291 PMID: 27362313
- Zaza G, Tomei P, Ria P, Granata S, Boschiero L, Lupo A. Systemic and nonrenal adverse effects occurring in renal transplant patients treated with mTOR inhibitors. Clin Dev Immunol. 2013; 2013: 403280. doi: 10.1155/2013/403280 PMID: 24151517
- Campistol JM, de Fijter JW, Flechner SM, Langone A, Morelon E, Stockfleth E. MTOR inhibitor-associated dermatologic and mucosal problems. Clin Transplant. 2010; 24(2): 149–156. doi: 10.1111/j.1399-0012.2010.01232.x PMID: 20236129
- Bin W, Feng-bo W, J L, Lei Y, Yao T. Meta-analysis of immunosuppressive therapy with target-of-rapamycin inhibitor instead of calcineurin inhibitor after kidney transplantation operation. China Pharmacy. 2011; 22(34): 3229–3231.
- Glover TE, Watson CJE, Gibbs P, Bradley JA, Ntzani EE, Kosmoliaptsis V. Conversion from calcineurin to mammalian target of rapamycin inhibitors in liver transplantation. Transplantation. 2016; 100(3): 621–629. doi: 10.1097/TP.00000000001006 PMID: 26636736
- Lim WH, Eris J, Kanellis J, Pussell B, Wiid Z, Witcombe D, et al. A systematic review of conversion from calcineurin inhibitor to mammalian target of rapamycin inhibitors for maintenance immunosuppression in kidney transplant recipients. Am J Transplant. 2014; 14(9): 2106–2119. doi: <u>10.1111/ajt.12795</u> PMID: 25088685
- Lieberthal W, Levine JS. The role of the mammalian target of rapamycin (mTOR) in renal disease. J Am Soc Nephrol. 2009; 20(12): 2493–2502. doi: 10.1681/ASN.2008111186 PMID: 19875810
- Diekmann F, Andres A, Oppenheimer F. MTOR inhibitor-associated proteinuria in kidney transplant recipients. Transplant Rev (Orlando). 2012; 26(1): 27–29.
- Higgins JPT, Altman DG, Gotzsche PC, Jueni P, Moher D, Oxman AD, et al. The Cochrane Collaboration's tool for assessing risk of bias in randomised trials. Bmj-Brit Med J. 2011; 343: d5928.
- Savovic J, Weeks L, Sterne JAC, Turner L, Altman DG, Moher D, et al. Evaluation of the Cochrane Collaboration's tool for assessing the risk of bias in randomized trials: Focus groups, online survey, proposed recommendations and their implementation. Systematic reviews. 2014; 3: 37. doi: 10.1186/2046-4053-3-37 PMID: 24731537
- Liberati A, Altman DG, Tetzlaff J, Mulrow C, Gotzsche PC, Ioannidis JP, et al. The PRISMA statement for reporting systematic reviews and meta-analyses of studies that evaluate healthcare interventions: Explanation and elaboration. BMJ. 2009; 339: b2700. doi: 10.1136/bmj.b2700 PMID: 19622552
- 22. Thibault G, Paintaud G, Legendre C, Merville P, Coulon M, Chasseuil E, et al. CD25 blockade in kidney transplant patients randomized to standard-dose or high-dose basiliximab with cyclosporine, or high-dose basiliximab in a calcineurin inhibitor-free regimen. Transpl Int. 2016; 29(2): 184–195. doi: 10. 1111/tri.12688 PMID: 26369526
- Rostaing L, Hertig A, Albano L, Anglicheau D, Durrbach A, Vuiblet V, et al. Fibrosis progression according to Epithelial-Mesenchymal transition profile: A randomized trial of everolimus versus CsA. Am J Transplant. 2015; 15(5): 1303–1312. doi: 10.1111/ajt.13132 PMID: 25808994
- Budde K, Lehner F, Sommerer C, Reinke P, Arns W, Eisenberger U, et al. Five-Year outcomes in kidney transplant patients converted from cyclosporine to everolimus: The randomized ZEUS study. Am J Transplant. 2015; 15(1): 119–128. doi: 10.1111/ajt.12952 PMID: 25521535
- 25. Mjornstedt L, Schwartz SS, von Zur MB, Jespersen B, Hansen JM, Bistrup C, et al. Renal function three years after early conversion from a calcineurin inhibitor to everolimus: Results from a randomized trial in kidney transplantation. Transpl Int. 2015; 28(1): 42–51. doi: 10.1111/tri.12437 PMID: 25176389

- 26. Budde K, Sommerer C, Rath T, Reinke P, Haller H, Witzke O, et al. Renal function to 5 years after late conversion of kidney transplant patients to everolimus: A randomized trial. J Nephrol. 2015; 28(1): 115–123. doi: 10.1007/s40620-014-0134-4 PMID: 25192833
- Budde K, Rath T, Sommerer C, Haller H, Reinke P, Witzke O, et al. Renal, efficacy and safety outcomes following late conversion of kidney transplant patients from calcineurin inhibitor therapy to everolimus: The randomized APOLLO study. Clin Nephrol. 2015; 83 (01): 11–21.
- Chadban SJ, Eris JM, Kanellis J, Pilmore H, Lee PC, Lim SK, et al. A randomized, controlled trial of everolimus-based dual immunosuppression versus standard of care in de novo kidney transplant recipients. Transpl Int. 2014; 27(3): 302–311. doi: 10.1111/tri.12252 PMID: 24279685
- Budde K, Lehner F, Sommerer C, Arns W, Reinke P, Eisenberger U, et al. Conversion From Cyclosporine to Everolimus at 4.5 Months Posttransplant: 3-Year Results From the Randomized ZEUS Study. Am J Transplant. 2012; 12(6): 1528–1540. doi: 10.1111/j.1600-6143.2012.03994.x PMID: 22642473
- Mjörnstedt L, Sørensen SS, von Zur Mühlen B, Jespersen B, Hansen JM, Bistrup C, et al. Improved renal function after early conversion from a calcineurin inhibitor to everolimus: A randomized trial in kidney transplantation. Am J Transplant. 2012; 12(10): 2744–2753. doi: 10.1111/j.1600-6143.2012. 04162.x PMID: 22812414
- Arns W. SCWO H. H. SMJJ K. Efficacy and Safety of Three Different Treatment Regimens in de. Transplantation. 2012; 94(Suppl 10): S995.
- Hallvard Holdaas LRDS, Erik H. Strom AJKM, Connell APJO, Investigators OBOT. Conversion of Long-Term kidney transplant recipients from calcineurin inhibitor therapy to everolimus: ARandomized, multicenter, 24-Month study. Transplantation. 2011; 92(4): 410–418. doi: 10.1097/TP. 0b013e318224c12d PMID: 21697773
- **33.** Budde K, Becker T, Arns W, Sommerer C, Reinke P, Eisenberger U, et al. Everolimus-based, calcineurin-inhibitor-free regimen in recipients of de-novo kidney transplants: An open-label, randomised, controlled trial. The Lancet. 2011; 377(9768): 837–847.
- Bemelman FJ, de Maar EF, Press RR, van Kan HJ, Ten Berge IJ, Homan Van Der Heide JJ, et al. Minimization of maintenance immunosuppression early after renal transplantation: An interim analysis. Transplantation. 2009; 88(3): 421–428. doi: 10.1097/TP.0b013e3181af1df6 PMID: 19667948
- Seckinger J, Sommerer C, Hinkel U, Hoffmann O, Zeier M, Schwenger V. Switch of immunosuppression from cyclosporine a to everolimus: Impact on pulse wave velocity in stable de-novo renal allograft recipients. J Hypertens. 2008; 26(11): 2213–2219. doi: <u>10.1097/HJH.0b013e32830ef940</u> PMID: <u>18854763</u>
- Kihm LP, Hinkel U, Michael K, Sommerer C, Seckinger J, Morath C, et al. Contrast enhanced sonography shows superior microvascular renal allograft perfusion in patients switched from cyclosporine a to everolimus. Transplantation. 2009; 88(2): 261–265. doi: <u>10.1097/TP.0b013e3181ac8cdd</u> PMID: <u>19623023</u>
- Grinyo JM, Saval N, Campistol JM. Clinical assessment and determinants of chronic allograft nephropathy in maintenance renal transplant patients. Nephrology Dialysis Transplantation. 2011; 26(11): 3750– 3755.
- CHAPMAN JR. Longitudinal analysis of chronic allograft nephropathy: Clinicopathologic correlations. Kidney International. 2005; 68(Suppl 99): 108–112.
- Cantarovich F, Martinez F, Heguilen R, Thervet E, Mamzer-Bruneel MF, Gonzalez C, et al. Proteinuria >0.5 g/d, a prevalent prognostic factor for patient and graft survival in kidney transplantation. Clinical Transplantation. 2010; 24(2): 175–180. doi: 10.1111/j.1399-0012.2009.01043.x PMID: 19624694
- **40.** Sancho CA, Pallardo ML, Avila BA, Gavela ME, Beltran CS, Crespo AJ. Very low-grade proteinuria at 3 months posttransplantation is an earlier marker of graft survival. Transplant Proc. 2009; 41(6):
- Halimi JM, Laouad I, Buchler M, Al-Najjar A, Chatelet V, Houssaini TS, et al. Early low-grade proteinuria: Causes, short-term evolution and long-term consequences in renal transplantation. American Journal of Transplantation. 2005; 5(9): 2281–2288. doi: 10.1111/j.1600-6143.2005.01020.x PMID: 16095510
- Webster AC, Lee VWS, Chapman JR, Craig JC. Target of rapamycin inhibitors (Sirolimus and everolimus) for primary immunosuppression of kidney transplant recipients: A systematic review and Meta-Analysis of randomized trials. Transplantation. 2006; 81(9): 1234–1248. doi: 10.1097/01.tp. 0000219703.39149.85 PMID: 16699448
- Sanchez-Fructuoso AI, Ruiz JC, Calvo N, Rodrigo E, Perez-Flores I, Gomez-Alamillo C, et al. Everolimus as primary immunosuppression in kidney transplantation: Experience in conversion from calcineurin inhibitors. Transplantation. 2012; 93(4): 398–405. doi: <u>10.1097/TP.0b013e31823ffd0e</u> PMID: 22245871
- 44. Fischer L, Klempnauer J, Beckebaum S, Metselaar HJ, Neuhaus P, Schemmer P, et al. A randomized, controlled study to assess the conversion from calcineurin-inhibitors to everolimus after liver

transplantation—PROTECT. Am J Transplant. 2012; 12(7): 1855–1865. doi: 10.1111/j.1600-6143. 2012.04049.x PMID: 22494671

- Loriga G, Ciccarese M, Pala PG, Satta RP, Fanelli V, Manca ML, et al. De novo Everolimus-Based therapy in renal transplant recipients: Effect on proteinuria and renal prognosis. Transpl P. 2010; 42(4): 1297–1302.
- 46. Opelz G, Unterrainer C, Susal C, Dohler B. Immunosuppression with mammalian target of rapamycin inhibitor and incidence of post-transplant cancer in kidney transplant recipients. Nephrol Dial Transplant. 2016; 31(8): 1360–1367. doi: 10.1093/ndt/gfw088 PMID: 27190384
- 47. Miura M, Yanai M, Fukasawa Y, Higashiyama H, Ito Y. De novo proteinuria with pathological evidence of glomerulonephritis after everolimus induction. Nephrology. 2014; 19(Suppl 3): S57–S59.
- Badve SV, Pascoe EM, Burke M, Clayton PA, Campbell SB, Hawley CM, et al. Mammalian target of rapamycin inhibitors and clinical outcomes in adult kidney transplant recipients. Clin J Am Soc Nephrol. 2016 Jul 21. pii: CJN.00190116.
- Gallagher MP, Hall B, Craig J, Berry G, Tiller DJ, Eris J. A randomized controlled trial of cyclosporine withdrawal in renal-transplant recipients: 15-Year results. Transplantation. 2004; 78(11): 1653–1660. PMID: 15591955