Optimizing the Safety Profile of Everolimus by Delayed Initiation in De Novo Heart Transplant Recipients: Results of the Prospective Randomized Study EVERHEART

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Background. Although everolimus potentially improves long-term heart transplantation (HTx) outcomes, its early postoperative safety profile had raised concerns and needs optimization. Methods. This 6-month, open-label, multicenter randomized trial was designed to compare the cumulative incidence of a primary composite safety endpoint comprising wound healing delays, pericardial effusion, pleural effusion needing drainage, and renal insufficiency events (estimated glomerular filtration rate ≤30/ mL/min per 1.73 m²) in de novo HTx recipients receiving immediate everolimus (EVR-I) (≤144 hours post-HTx) or delayed everolimus (EVR-D) (4-6 weeks post-HTx with mycophenolate mofetil as a bridge) with reduced-dose cyclosporine A. Cumulative incidence of biopsy-proven rejection ≥ 2R, rejection with hemodynamic compromise, graft loss, or death was the secondary composite efficacy endpoint. Results. Overall, 181 patients were randomized to the EVR-I (n = 89) or EVR-D (n = 92) arms. Incidence of primary safety endpoint was higher for EVR-I than EVR-D arm (44.9% vs 32.6%; P = 0.191), mainly driven by a higher rate of pericardial effusion (33.7% vs 19.6%; P = 0.04); wound healing delays, acute renal insufficiency events, and pleural effusion occurred at similar frequencies in the study arms. Efficacy failure was not significantly different in EVR-I arm versus EVR-D arm (37.1% vs 28.3%; P = 0.191). Three patients in the EVR-I arm and 1 in the EVR-D arm died. Incidence of clinically significant adverse events leading to discontinuation was higher in EVR-I arm versus EVR-D arm (P = 0.02). Conclusions. Compared with immediate initiation, delayed everolimus initiation appeared to provide a clinically relevant early safety benefit in de novo HTx recipients, without compromising efficacy.

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Materials and Methods

Study Design

EVERHEART was a 6-month, phase IIIB prospective, open-label, randomized, parallel-group study conducted across 12 HTx centers in Italy between September 2009 and December 2013. The open-label design was selected to allow therapeutic drug monitoring of everolimus and CsA. Patients were screened at transplant and randomized to 144 hours (days 1-5) after graft perfusion in a 1:1 ratio via a Web-based system to either immediate everolimus (EVR-I) or delayed everolimus (EVR-D) arms and stratified by baseline estimated glomerular filtration rate (eGFR) ≤60 or >60 mL/min per 1.73 m²; Modification of Diet in Renal Disease scale) and pretransplant diabetes status. In the EVR-D arm, MMF was replaced with everolimus at 4 to 6 weeks to prevent potentially adverse conditions. Patients were followed up at weeks 2, 4, and 6 and months 2, 3, and 6 (Figure 1A).

Study participants provided written informed consent, and study protocol was approved by each center’s institutional review board in accordance with the European Community Guidance on Good Clinical Practice guidelines and the Declaration of Helsinki.

Study Population

Male or female de novo HTx recipients aged 18 years or older were eligible. Main exclusion criteria at screening were intolerance to CsA/statins; seropositivity for human immunodeficiency virus, hepatitis C virus, or hepatitis B surface antigen; hepatitis C virus- and hepatitis B surface antigen-positive donors; and panel reactive antibodies greater than 30% as assessed by cytotoxicity assay. To be randomized, patients had to meet the following criteria between days 1 and 5 after graft perfusion: eGFR ≥40 mL/min per 1.73 m², platelet count ≥40,000/mm³, white blood cell count >4000/mm³; absence of a clinically significant systemic infection; and cold ischemia time less than 6 hours.

Treatment Regimen

Patients in the EVR-I arm received everolimus (trough level [C₀], 3-8 ng/mL [5-10 ng/mL by InnoFluor Certican® assay kit, Seradyn Inc.]0.75 mg bid orally at 12-hour intervals, with dose initiation of 144 hours or less after graft perfusion. The everolimus dose was reduced to 0.50 mg bid if platelet count was less than 100,000/mm³; white blood cell count, less than 5000/mm³; body mass index (BMI), greater than 30 kg/m²; body weight, less than 60 kg; eGFR, less than 60 mL/min per 1.73 m²; or if diabetes was present.

Patients in the EVR-D arm received the first MMF dose of 1 g or greater within 144 hours after graft perfusion; the target maintenance dose of 3 g daily was achieved by the first week. At 4 to 6 weeks, MMF was replaced with 0.5 mg bid everolimus. The dose was reduced to 0.25 mg bid if one of the conditions reported for the EVR-I arm was present. MMF was discontinued when everolimus C₀ reached 3 ng/mL (5 ng/mL, Certican assay kit kit).18

Patients in both arms received oral CsA (Neoral®) per the prespecified target trough levels ([C₀], Table S1, SDC, http://links.lww.com/TP/B489). During the first 4 to 8 weeks after transplant, CsA C₀ levels were lower in the EVR-I arm versus EVR-D arm. In the EVR-D arm, CsA C₀ levels were lowered when MMF was substituted by everolimus.

The use and choice of induction therapy, steroid downtitration protocol, and prophylactic/preemptive treatment of CMV infections were in accordance with the centers’ standard practice. Fluvastatin 40 mg was administered to all patients as a lipid-lowering agent, starting at week 4. At month 3, the dose was increased to 80 mg in patients with low-density lipoprotein of 100 mg/dL or greater.19,20 Acute rejection was monitored by endomyocardial biopsies and treated with intravenous pulse steroids if graded 2R or higher.

Primary Endpoints

The primary composite safety endpoint was the 6-month cumulative incidence of surgical wound healing delays, PLEs, PCEs, and acute renal insufficiency (eGFR ≤30 mL/min/1.73 m²). Any sign of wound dehiscence and need for surgical operations was noted. The primary composite efficacy endpoint was the 6-month cumulative incidence of graft rejection and mortality.ing operations was noted. The primary composite efficacy endpoint was the 6-month cumulative incidence of graft rejection and mortality.
revision of lymphoceles recorded at weeks 2, 4, and 6 and month 6 were considered as events concurring with the composite endpoint. Need for surgical drainage tubes for more than 7 days after surgery and/or PLE leading to drainage were considered as events concurring with the composite endpoint. Any PCE recorded on echocardiograms obtained at days 1 to 5, weeks 2 and 4, and months 3 and 6 and defined as at least moderate (ie, \( \geq 2.0 \) cm in diastole), with/without signs of hemodynamic compromise, or leading to drainage/prolonged hospitalization met the criteria as events concurring with the composite endpoint. Echocardiograms were reviewed and assigned as PCEs by treatment-blinded data monitoring committee members. Patients lost to follow-up without experiencing any of the above events were considered as failures.

**Secondary Endpoints and Adverse Events**

Secondary efficacy endpoint was assessed as the cumulative incidence of biopsy-proven acute rejection (BPAR) \( \geq 2R,^{21} \) rejection with hemodynamic compromise, graft loss, and death. Hemodynamic compromise was defined as the need of inotrope therapy or vasoactive treatment. The incidences of adverse events (AEs), serious AEs (SAEs), AEs leading to discontinuation, CMV infections and disease, and low-density lipoprotein levels of 100 ng/mL or greater were separately monitored.

**Sample Size and Statistical Analyses**

Based on the empirical incidence of 60% for the primary endpoint on introducing everolimus immediately after surgery (EVR-I arm),\(^{5-7,9} \) we hypothesized that the primary endpoint would occur in 40% of patients with delayed everolimus introduction. Thus, a recruitment target of 194 patients (97 per arm) was aimed to determine statistically significant superiority of EVR-D, with a 5% 2-tailed alpha error and 80% power.

All analyses were conducted on the safety population (intention-to-treat population including patients who could...
not be switched to everolimus). The hazards ratios (HRs) with 95% confidence intervals (CIs) for occurrence of primary and secondary endpoints were determined using the Cox regression model. Effects of potential confounders were analyzed between the composite safety endpoints and baseline clinical or demographic variables using odds ratios (ORs).

RESULTS

Study Population and Interventions

Overall, 201 patients were screened; of these, 182 were randomized to the EVR-I (n = 90) and EVR-D (n = 92) arms; of these, 1 patient assigned to the EVR-I arm did not receive any everolimus dose. Randomized patients corresponded to 93% of the planned population with a 77% power against the initial study assumption of 80%. In all, 175 patients completed the study (EVR-I, 85 and EVR-D, 90; Figure 1B). The baseline characteristics of randomized patients and donors were mostly balanced between the 2 arms (Table 1). The therapeutic switch from MMF to everolimus was successful in 87 (94.6%) patients. The median time-to-switch was 6.0 weeks (range, 3.4-14.1 weeks). The delayed approach was associated with a lower rate of study drug discontinuation (24.7%, EVR-I vs 9.8%, EVR-D; P = 0.008; Figure 1B).

Majority of patients in both arms received induction therapy (EVR-I, 77/89 [86.5%] vs EVR-D, 71/92 [77.2%]). Of these, 9 (11.7%) of 77 patients and 6 (8.5%) of 71 patients in EVR-I versus EVR-D arm received basiliximab, respectively. Alternatively, 45 (58.4%) of 77 patients and 39 (54.9%) of 71 patients in EVR-I and EVR-D arms received ATG, respectively. Antilymphocyte immunoglobulin was administered to 18 (23.4%) 77 patients and 22 (31.0%) 71 patients in EVR-I and EVR-D arms, respectively. Steroids were administered at comparable doses during induction and maintenance.

The recommended CsA levels were more frequently adhered to in the EVR-D versus EVR-I arms (Figure S1A-D, SDC, http://links.lww.com/TP/B489). At month 6, CsA C0 levels were above the protocol-defined target range in 43 (62.3%) of 69 patients and 35 (47.9%) of 73 patients in the EVR-I and EVR-D arms, respectively (P = 0.085; Figure S1D, SDC, http://links.lww.com/TP/B489).

Primary Safety Endpoint

The 6-month cumulative incidence of composite safety endpoints was 44.9% (n = 40) versus 32.6% (n = 30) in the EVR-I vs EVR-D arms, respectively (log-rank test: P = 0.1913; Figure 2A). This accounted for a 48% nonsignificant increase in relative risk in the EVR-I arm (HR, 1.48; 95% CI, 0.92-2.38; P = 0.104; Table 2). The difference in study arms was driven by the increased incidence of PCEs in the EVR-I arm, which had an 80% significant risk of PCEs (HR, 1.84; 95% CI 1.02-3.30; log-rank test: P = 0.041; Figure 2B). Furthermore, among the 30 patients with PCEs in the EVR-I arm, 10 (33.3%) needed pericardiocenteses, as opposed to only 1 (5.5%) of the 18 patients with PCEs in the EVR-D arm (P = 0.03). On the other hand, wound healing delays, episodes of renal insufficiency, and PLEs needing drainage were not significantly different between the 2 study groups (Table 2).

During the study follow-up, eGFR (Figure S2, SDC, http://links.lww.com/TP/B489) levels were comparable, indicating no impairment in renal function with the delayed approach. Nevertheless, baseline eGFR less than 60 mL/min per 1.73 m² was a significant risk factor for post-HTx renal insufficiency (HR, 3.33; 95% CI, 1.13-9.77; P = 0.029; Table 2).

### Table 1. Baseline demographics

| EVR-I (N = 89) | EVR-D (N = 92) |
|---------------|---------------|
| Age: mean (SD), y | 38.0 (12.0) | 39.2 (13.9) |
| Male sex, n (%) | 53 (59.6) | 61 (66.3) |
| Diabetes history |  |  |
| Yes | 2 (2.2) | 3 (3.3) |
| Unknown | 13 (14.6) | 11 (12.0) |
| Donor-recipient CMV status |  |  |
| Positive/positive | 41 (46.1) | 40 (43.5) |
| Positive/negative | 19 (21.3) | 11 (12.0) |
| Negative/positive | 11 (12.4) | 10 (10.9) |
| Negative/negative | 14 (15.7) | 12 (13.0) |
| Unknown | 4 (4.5) | 19 (20.7) |

#### Intraoperative and postoperative features

| EVR-I (N = 89) | EVR-D (N = 92) |
|---------------|---------------|
| Cold ischemia time: mean (SD), h | 2.9 (1.0) | 2.9 (0.9) |
| CPB duration: mean (SD), h | 2.6 (0.7) | 2.6 (0.8) |
| Conventional osteosynthesis, n (%) | 87 (97.8) | 87 (94.6) |
| Additional lateral reinforcement, n (%) | 1 (1.1) | 4 (4.3) |
| Surgical revision for bleeding, n (%) | 3 (3.4) | 2 (2.2) |
| Surgical drainage removal (days), mean (SD) | 4.71 (2.23) | 4.41 (1.39) |

* Includes idiopathic, postmyocarditis, peripartum, familial, antilymphocyte-induced, alcoholic, and ischemic dilated cardiomyopathy, and so on.
* Includes hypertrophic cardiomyopathies, idiopathic or restrictive cardiomyopathies.
* Includes endomyocardial fibrosis, valvular heart disease, congenital structural heart disease, or other cardiovascular diseases.
* A technique used for repairing sternum dehiscence.
* Calculated as date of surgical removal—date of transplant + 1.
* All characteristics tested not significantly different between study groups.

CPB, cardiopulmonary bypass.
None of the baseline and procedural variables was associated with significant risk of a composite safety endpoint, except for borderline effects of donor older than 35 years and female donors ($P = 0.063$; Figure 3A). Within the patient subgroups, the risk of safety endpoint was significantly lower in the EVR-D versus EVR-I arms for patients with BMI of 30 kg/m$^2$ or less, without diabetes, and with 35 year or younger donors (Figure 3B).

**Secondary Efficacy Endpoint**

Composite efficacy failure events were higher, but not significantly different, in the EVR-I arm (37.1% [n = 33], EVR-I vs 28.3% [n = 26], EVR-D; HR, 1.40; 95% CI: 0.84-2.35; log-rank test: $P = 0.191$; Table 3). During the 6-month period, 4 patients died: 3 in the EVR-I arm (1 each due to congestive heart failure/multiple organ failure, pulmonary infection, and arrhythmia), and 1 in the EVR-D arm (gastric adenocarcinoma).

**Other Safety Events**

At least 1 AE occurred in 86.5% (n = 77) and 80.4% (n = 74) of patients in the EVR-I and EVR-D arms, respectively. Overall, clinically significant AEs and AEs leading to drug discontinuation were significantly higher in the EVR-I arm than in the EVR-D arm (Table 4). Discontinuations due to SAEs were also significantly higher in the EVR-I versus EVR-D arm ($P = 0.048$). Incidence of patients requiring fluvastatin dose adjustment was comparable between the 2 arms at month 1 (46.1% [n = 41], EVR-I vs 41.8% [n = 38], EVR-D), month 3 (43.0% [n = 37], EVR-I vs 40.2% [n = 37], EVR-D), and month 6 (38.2% [n = 34], EVR-I vs 39.1% [n = 36], EVR-D). Risk of CMV disease/infection was significantly lower in the EVR-I versus EVR-D arms (OR, 0.414; 95% CI, 0.223-0.771; $P = 0.005$).

**DISCUSSION**

This multicenter, prospective, randomized, open-label study designed to systematically evaluate early postoperative complications in de novo HTx recipients provides suggestive evidence that the timing of everolimus initiation after HTx may influence the risk of AEs, mainly moderate PCEs and PCEs leading to pericardiocentesis. In addition,
FIGURE 3. A, Prognostic factors of primary composite endpoint by baseline characteristics. B, Forest plot depicting occurrence of composite safety endpoint by baseline characteristics between delayed and immediate everolimus initiation arms. *BMI, body mass index; **IC, ischemic cardiomyopathy; ***eGFR, estimated glomerular filtration rate (mL/min per 1.73 m²).

TABLE 3.
Incidence and risk of efficacy endpoints in safety population

|                     | EVR-I (N = 89) | EVR-D (N = 92) |
|---------------------|---------------|---------------|
| **Efficacy**        |               |               |
| At least occurrence| 33 (37.1)     | 26 (28.3)     |
| BPAR³               | 30 (33.7)     | 24 (26.1)     |
| Rejection with hemodynamic compromise | 4 (4.5) | 5 (5.4) |
| Graft loss           | 0 (0.0)       | 0 (0.0)       |
| Death                | 3 (3.4)       | 1 (1.1)       |

| **Cox proportional hazard model analysis** |
|------------------------------------------|

| Efficacy | EVR immediate vs delayed | eGFR ≤60 vs >60 mL/min per 1.73 m² | Diabetes presence vs absence |
|----------|--------------------------|-----------------------------------|-----------------------------|
| ≥1 Occurrence | HR 95% CI P | HR 95% CI P | HR 95% CI P |
| BPAR³ | 1.40 (0.84-2.35) 0.197 | 0.79 (0.46-1.35) | 0.383 (0.88-1.93) 0.740 |
| Rejection with hemodynamic compromise | 1.36 (0.80-2.33) 0.259 | 0.82 (0.47-1.44) | 0.488 (0.82-3.51-3.19) 0.654 |
| Death | 0.83 (0.22-3.09) 0.778 | 0.79 (0.20-3.16) | 0.739 (0.84-11-6.74) 0.871 |

³Defined as International Society for Heart and Lung Transplantation grade ≥2R.

P-values were determined by proportionality hazard assumption test.
everolimus initiation between 4 and 6 weeks after surgery appears to provide a better safety profile as compared with immediate initiation, retaining antirejection efficacy, whereas everolimus discontinuation due to AEs appears to be more frequent when the drug is initiated in the immediate postoperative period. On the other hand, CMV infection was more common with delayed rather than early everolimus initiation.

To date, 4 multicenter randomized studies have evaluated the efficacy of everolimus combined with CsA in de novo HTx recipients with major differences in everolimus dosing and administration strategies.5-7,9,23 PCEs and PLEs were reported more frequently in the everolimus arm in 2 of the 4 studies,7,9 which is consistent with the higher incidence in the EVR-I arm (44.9%, EVR-I arm vs 32.6%, EVR-D arm), for which everolimus was introduced within 5 days of HTx. Therefore, we suggest that delayed everolimus initiation could markedly reduce the incidence of critical AEs, like PCEs/PLEs, which can compromise safety in heart transplant recipients.

Previous studies in which everolimus was introduced with sCsA in the immediate postoperative period,7,9 impaired renal function appeared as a concern.6,7,9,10,21,28,29 Thus, if the goal of everolimus-based regimen initiated within 72 hours post-HTx is associated with a lower incidence of CMV infections compared with azathioprine- or MMF-based regimens. Indeed, the risk of CMV disease in our study was significantly higher in the EVR-D versus EVR-I arms, indicating that delayed initiation may not alleviate baseline CMV infection, most likely because preemptive strategy was usually preferred over prophylaxis in most centers, thereby exposing the patients to high risk of early reactivation during the initial phase of MMF intake.

Evidence from randomized trials and observational studies support the concept that everolimus improves long-term outcomes by reducing early CAV development and malignancy onset.7,9,10,21,28,29 Thus, if the goal of everolimus-based therapies is long-term benefit, a strategy that avoids early toxicity could improve patient management, reduce the chances of early discontinuation, and increase potential benefits for patients.

The main limitation of this study was a slightly underpowered sample size, resulting from dramatic decrease in the number of HTx procedures performed in Italy during the study period. Patient enrollment began in June 2009 and had to be repeatedly prolonged until December 2013. When the study was first designed, 320 to 350 HTx procedures/year were being performed in Italy.30 However, since initiation, the number of procedures dramatically reduced to less than 250 cases/year. To avoid excessive prolongation of study

| TABLE 4. Incidence of adverse and serious AEs |
|-----------------------------------------------|
| Safety parameters, n (%) | EVR-I (N = 89) | EVR-D (N = 92) | P |
|--------------------------|---------------|---------------|---|
| At least 1 SAE           | 35 (39.3)     | 31 (33.7)     | 0.443 |
| At least 1 clinically significant AE<sup>a</sup> | 41 (46.1) | 27 (29.3) | 0.020 |
| Death                    | 3 (3.4)       | 1 (1.1)       | 0.361 |
| Discontinued due to SAE(s) | 13 (14.6) | 5 (5.4) | 0.048 |
| Discontinued due to clinically significant AEs | 19 (21.3) | 7 (7.6) | 0.011 |
| Most frequently reported SAEs | At least one SAE (≥3% in any arm) | 35 (39.3) | 31 (33.7) | 0.443 |
| Pneumonia                | 4 (4.5)       | 1 (1.1)       | 0.206 |
| Acute renal failure      | 3 (3.4)       | 1 (1.1)       | 0.361 |
| CMV syndrome             | 0 (0.0)       | 3 (3.3)       | 0.246 |
| Most frequently reported AEs (≥5% in any arm) | At least 1 AE | 77 (86.5) | 74 (80.4) | 0.320 |
| Leukopenia               | 16 (18.0)     | 8 (8.7)       | 0.081 |
| Hypertension             | 10 (11.2)     | 12 (13.0)     | 0.821 |
| Anemia                   | 9 (10.1)      | 9 (9.8)       | 1.000 |
| Peripheral edema         | 7 (7.9)       | 3 (3.3)       | 0.207 |
| Hypertriglyceridemia     | 5 (5.6)       | 10 (10.9)     | 0.282 |
| Atrial fibulation        | 5 (5.6)       | 4 (4.3)       | 0.744 |
| Hypercholesterolemia     | 4 (4.5)       | 6 (6.5)       | 0.747 |
| Thrombocytopenia          | 3 (3.4)       | 5 (5.4)       | 0.721 |
| Pyrexia                  | 3 (3.4)       | 6 (6.5)       | 0.497 |
| Renal failure            | 2 (2.2)       | 5 (5.4)       | 0.444 |
| Infections and infestations (≥2% in any arm) | All infections (≥3% in any arm) | 46 (51.7) | 63 (68.5) | 0.005 |
| CMV infection            | 10 (11.2)     | 5 (5.4)       | 0.185 |
| Urinary tract infections | 2 (2.2)       | 2 (2.2)       | 1.00 |
| CMV syndrome             | 0 (0.0)       | 3 (3.3)       | 0.246 |
| Upper respiratory tract infection | 2 (2.2) | 0 (0.0) | 0.240 |

<sup>a</sup> Clinically significant AE was defined as adjustment/temporary interruption of study drug dose or permanent discontinuation of study drug.
period, we terminated enrollment at 93% of the planned population, accounting for a 77% power for the initial assumptions. The study power was additionally affected by the discrepancy between expected versus observed incidences of the primary endpoint.5–7 The use of CsA instead of tacrolimus, the current CNI of choice, may appear as another study limitation. However, little clinical evidence of tacrolimus and everolimus was available when the study was designed; moreover, the objective of our study was evaluation of everolimus-related safety profile, rather than the CNI combination therapy. Finally, it must be noted that as in most of randomized controlled studies, the current cohort of patients represents a healthier subgroup when compared with the overall population of patients undergoing HTx. It is possible that this selection may be driven by the need for patient stability between screening and randomization at day 5, with unstable patients unlikely to get enrolled.

In conclusion, in the context of an immunosuppressive regimen based on everolimus and reduced-dose CsA, delayed initiation of everolimus with MMF as a bridge appears to provide a better safety profile than immediate initiation, by reducing the incidence of PCEs, especially those requiring pericardiocentesis, and by improving overall drug tolerability, with less AE-driven discontinuations, without compromising antirejection efficacy.

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