Cross-sectional analysis of immunosuppressive regimens focused on everolimus after liver transplantation in a Korean high-volume transplantation center

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Background: The mammalian target of the rapamycin inhibitor has dual inhibitory effects on cell growth and angiogenesis. This study aimed to analyze the usage of everolimus on actual immunosuppression (IS) regimens through a cross-sectional study in a high-volume liver transplantation (LT) center.

Methods: Our institutional LT database was searched for adult patients who underwent primary LT surgery between January 2010 and December 2016. We identified 2,093 LT recipients with observation periods of 1 to 8 years.

Results: We divided the 2,093 recipients into three groups according to the posttransplant follow-up period as follows: group A (12–36 months; n=680), group B (37–60 months; n=560), and group C (>60 months; n=853). The individual IS agents were tacrolimus in 1,807 patients (86.3%), cyclosporine in 169 patients (8.1%), mycophenolate mofetil (MMF) in 1,310 patients (62.6%), and everolimus in 115 patients (5.5%). The most common IS regimens were tacrolimus-MMF combination and tacrolimus monotherapy, regardless of the posttransplant period. Patients with pretransplant malignancies were administered everolimus more frequently than those without pretransplant malignancies (P<0.001). In 102 patients with hepatocellular carcinoma recurrence or de novo malignancies, IS regimens included everolimus-tacrolimus in 41 patients (40.2%), tacrolimus-MMF in 27 patients (26.4%), tacrolimus in 20 patients (19.6%), MMF in 10 patients (9.8%), cyclosporine in three patients (2.9%), and cyclosporine-MMF in one patient (1.0%).

Conclusions: Administration of everolimus after LT has been gradually increasing with the expansion of indications in our institutional practice. Currently, the role of everolimus is minimal and not comparable to that of tacrolimus, but it has a unique position in the field of IS after LT.

Keywords: Tacrolimus; Mycophenolate mofetil; Everolimus; Malignancy; Hepatocellular carcinoma

INTRODUCTION

Immunosuppression (IS) is essential after liver transplantation (LT), with the exception of the development of operational tolerance. To date, every available immunosuppressive agent (ISA) has been administered after LT. Most IS protocols for LT include calcineurin inhibitors (CNI), mycophenolate mofetil (MMF), and steroids [1-3]. Steroids are usually tapered off early or in-
everolimus after liver transplantation

HIGHLIGHTS

∙ A cross-sectional study in a high-volume center revealed that the most common immunosuppressive regimens were tacrolimus–mycophenolate mofetil combination and tacrolimus monotherapy.

∙ Everolimus use has been gradually increasing with expansion of indications in our institutional practice.

The mammalian target of rapamycin inhibitor (mTORi) is an ISA that interrupts the interleukin-2 proliferative response cascade and has dual inhibitory effects on cell growth and angiogenesis. mTORi is increasingly administered as indicated for renal dysfunction or malignancies. Everolimus has been financially covered by the Korean social health insurance since early 2016 and is reported to be beneficial in reducing the risk of HCC recurrence after LT as well as being effective after overt HCC recurrence [4-6]. However, the real-world status of everolimus administration after LT in Korea has not been well reported [1,3,5,7,8]. The purpose of this study was to analyze the use of everolimus on actual IS regimens through a cross-sectional study in a Korean high-volume LT center.

METHODS

Study Design and Patient Selection

This cross-sectional study analyzed actual ISA regimens in adult LT recipients, with a focus on everolimus. We set the study period for cross-sectional review as 3 months from January 2018 to March 2018. The LT database at our institution was searched to identify adult patients who underwent primary LT during between January 2010 and December 2016. The inclusion criteria were patient survival for at least 12 months following LT and until the end of December 2018, recipient age ≥18 years at LT surgery, and regular visits to our institution’s outpatient clinic. Finally, we identified 2,093 LT recipients with an observation period of between 1 to 8 years. Our study protocol was approved by the Institutional Review Board of our institution.

Institutional IS Regimen Protocols

Primary IS protocols used for adult LT recipients at our institution consisted of interleukin-2 receptor inhibitor, intraoperative steroid bolus (5–10 mg/kg), intravenous or oral CNI and corticosteroid recycling beginning on day 1, and adjunctive MMF for patients showing CNI-associated adverse effects or for IS augmentation. For CNI-sparing to control CNI-associated adverse effects, tacrolimus and cyclosporine were occasionally exchanged. There was no differences in IS regimens between living-donor and deceased-donor LTs. Corticosteroids were rapidly tapered off within the first 3 months post LT [3,9,10].

With regards to mTORi, we currently use only everolimus in our institution as it is financially covered for LT recipients by the Korean social health insurance program. The main indications for everolimus at our institution include HCC recurrence, de novo malignancy, and renal dysfunction. Intentional weaning from all ISAs is, to date, not considered at our institution, except for in cases of life-threatening severe infection or terminal status due to advanced malignancies. As everolimus was not included as a primary ISA at our institution during the study period, it was not commonly used prior to HCC recurrence or the development of de novo malignancies. If recurrence of HCC occurred, the IS regimen was switched towards everolimus-based regimens if the adverse side effects were tolerable.

Stratification of LT Recipients According to Posttransplant Periods

As the peritransplant condition of recipients was diverse, the IS regimens were also highly variable. Thus, we did not include IS regimens during the first 1 year in the present analysis.

We divided 2,093 recipients into three groups according to the posttransplant follow-up period, as follows: 2nd and 3rd years (12-36 months, group A: n=680), 4th and 5th years (37-60 months, group B: n=560) and beyond the 5th year (>60 months, group C: n=853). These
RESULTS

Stratification of Patients According to Posttransplant Periods

The numbers of LT recipients according to the posttransplant periods were 680 (32.5%) in group A, 560 (26.8%) in group B and 853 (40.1%) in group C. The clinical profiles are summarized in Table 1. Pretransplant malignancies were diagnosed in 980 patients (46.8%).

Patterns of IS Regimens According to Posttransplant Periods

Individual ISAs used in all patients were tacrolimus in 1,807 patients (86.3%), cyclosporine in 169 patients (8.1%), MMF in 1,310 patients (62.6%), and everolimus in 115 patients (5.5%). In group A, the ISA regimens included tacrolimus in 652 patients (95.9%), cyclosporine in 18 patients (2.6%), MMF in 387 patients (56.9%), and everolimus in 64 patients (9.4%). The most common IS regimens were tacrolimus–MMF combination (n=358,

Table 1. Comparison of recipient profiles according to the posttransplant periods

| Variable                         | Group A (12–36 mon) | Group B (37–60 mon) | Group C (>60 mon) | Total       |
|----------------------------------|---------------------|---------------------|-------------------|-------------|
| No. of cases                     | 680 (32.5)          | 560 (26.8)          | 853 (40.1)        | 2,093       |
| Age at LT (yr)                   | 53.3±9.1            | 52.4±9.1            | 51.6±8.3          | 52.3±8.7    |
| Age at study (yr)                | 55.8±9.0            | 59.3±9.0            | 58.7±8.2          | 57.1±8.7    |
| Sex                              | Male 500 (73.5)     | 409 (73.0)          | 648 (76.1)        | 1,556 (74.4) |
|                                  | Female 180 (26.5)   | 151 (27.0)          | 536 (25.6)        |             |
| Primary liver disease            |                     |                     |                   |             |
| Hepatitis B                      | 357 (52.5)          | 329 (60.5)          | 570 (66.8)        | 1,266 (60.5) |
| Hepatitis C                      | 27 (4.0)            | 39 (7.0)            | 51 (6.0)          | 117 (5.6)   |
| Alcoholic disease                | 179 (26.3)          | 107 (19.1)          | 118 (17.9)        | 404 (19.4)  |
| Others                           | 117 (17.2)          | 75 (13.4)           | 114 (13.4)        | 306 (14.6)  |
| Type of LT (n)                   |                     |                     |                   |             |
| Deceased-donor                   | 94 (13.8)           | 58 (10.4)           | 108 (12.7)        | 260 (12.4)  |
| Living-donor                     | 586 (86.2)          | 502 (89.6)          | 745 (87.3)        | 1,833 (87.6) |
| Pretransplant malignancy (n)     |                     |                     |                   |             |
| Present                          | 295 (43.4)          | 262 (46.8)          | 423 (49.6)        | 980 (46.8)  |
| Absent                           | 385 (56.6)          | 298 (53.2)          | 430 (50.4)        | 1,113 (53.2) |
| De novo malignancy (n)           |                     |                     |                   |             |
| Present                          | 8 (1.2)             | 24 (4.3)            | 37 (4.3)          | 69 (3.3)    |
| Absent                           | 672 (98.8)          | 536 (95.7)          | 816 (95.7)        | 2,024 (96.7) |

Values are presented as number (%) or mean±standard deviation.

LT, liver transplantation.
Patterns of IS Regimens According to Pretransplant Malignancy

In the 1,113 patients without pretransplant malignancies, the ISA regimens included tacrolimus in 962 patients (86.4%), cyclosporine in 86 patients (7.7%), MMF in 699 patients (62.8%), and everolimus in 28 patients (2.5%). The most common IS regimens were tacrolimus–MMF combination (n=585, 52.6%) and tacrolimus monotherapy (n=349, 31.4%). In the 980 patients with pretransplant malignancies, the ISA regimens included tacrolimus in 845 patients (86.2%), cyclosporine in 83 patients (8.5%), MMF in 611 patients (62.3%), and everolimus in 87 patients (8.9%). The most common IS regimens were tacrolimus–MMF combination (n=509, 51.9%) and tacrolimus monotherapy (n=250, 25.5%) (Fig. 2). There was no difference in ISA use between the two groups, with the exception of everolimus (P < 0.001). The proportion of recipients who received everolimus was 8.9% and 2.5% in recipients with and without pretransplant malignancies, respectively.
Patterns of IS Regimens According to Posttransplant Malignancy

At the end of study period, the number of living patients with HCC recurrence or de novo malignancies were 52 and 67, respectively. With exclusion of the overlapped patients (n=17), a total of 102 patients were categorized as the posttransplant malignancy group. The IS regimens included everolimus–tacrolimus combination in 41 patients (40.2%), tacrolimus–MMF combination in 27 patients (26.4%), tacrolimus monotherapy in 20 patients (19.6%), MMF monotherapy in 10 patients (9.8%), cyclosporine monotherapy in three patients (2.9%), and cyclosporine–MMF combination in one patient (1.0%). Thus, 41 recipients (40.2%) were switched to everolimus due to HCC recurrence or the development of de novo malignancies [5]. Of the 115 patients treated with everolimus, 74 patients (64.3%) were administered everolimus to prevent HCC recurrence and de novo malignancy development or to inducing CNI-sparing effect.

Institutional Experience of Switching to mTORi in Recipients with HCC Recurrence

Of the 232 recipients who showed posttransplant HCC recurrence prior to December 2015, mTORi was administered to 51 recipients. However, 12 recipients (23.5%) dropped out within first 3 months due to adverse effects or other causes, including excessive financial burdens. However, subgroup analysis revealed that mTORi administration was stopped in only four of these patients as the severe adverse effects were not controlled despite dosage reduction. In the remaining 39 patients, mTORi was maintained to date or until patient death. After the implementation of insurance coverage for everolimus in Korea, the dropout rate of everolimus was lowered to two out of 30 recipients (6.7%). Everolimus was used alone or in combination with minimal CNI or MMF. None of the everolimus recipients experienced overt acute rejection that required high-dose steroid therapy [5].

DISCUSSION

The recent introduction of mTOR inhibitors in Korea has resulted in noticeable changes to IS regimens. Everolimus coverage was introduced to Korean social health insurance in early 2016. Thereafter, the rate at which mTORi administration was terminated decreased significantly. So far, we do not consider this agent to be a primary ISA, thus it is not, as of yet, frequently used in our institution. Everolimus can be administered following the first 1 month post LT, primarily due to the risk of hepatic artery thrombosis and issues with regards to wound healing. Korean social health insurance policy forcefully suggests administrating tacrolimus and everolimus concurrently.

In our institution, the first indication that everolimus should be prescribed is HCC recurrence or the development of de novo malignancies. This is due to the fact that mTORi are known to be the only ISA with anti-tumor effects [11-16]. Our study revealed that the overall use of everolimus is not high, but its use is gradually increasing as the criteria for its use are expanded. Due to the social health insurance policy and its relatively low immunosuppressive potency, it is usually administered concurrently with tacrolimus or MMF. Everolimus monotherapy has rarely been adopted, especially in patients with overt presentation of HCC recurrence or de novo malignancies. We intend to concurrently use mTORi and sorafenib with the expectation of the synergistic effect against HCC recurrence. However, their beneficial effects were not demonstrated in our precedent clinical study [5].

The second indication for everolimus use in our institution is renal dysfunction because of its CNI-sparing effect. It is known to be beneficial in effectively decreasing the concentration of CNIs. The H2304 study is a multicenter, open-label, randomized control study to evaluate the efficacy and safety of everolimus to eliminate or reduce tacrolimus in de novo LT recipients [6,17,18]. This study showed clear separation and clinically relevant reductions in tacrolimus exposure in the everolimus–reduced tacrolimus group, which resulted in improvements in renal function. However, the everolimus monotherapy following tacrolimus group was stopped early due to a high incidence of acute rejection. It is a critically important finding that everolimus is not sufficiently potent to pre-
vent acute rejection, especially during the first post-
transplant year.

In the current Korean setting, CNI–spared effect of
everolimus does not appear critical because many Korean
LT centers accumulated experience on reduced CNI–MMF
combination or MMF monotherapy to cope with CNI–in-
duced renal dysfunction [3,9,10]. At this point, we em-
phasize that MMF dosing requires mycophenolic acid
(MPA) therapeutic drug monitoring because a consid-
erable proportion of LT recipients are poor absorbers of
MMF. Poor absorption of MMF is not an eligible in-
dication for MMF monotherapy. We arbitrarily defined
poor MMF absorption as a 12 hour trough level of MPA
of <1.0 mg/L after administration of MMF 500 mg twice
per day [9]. Since the potency of MMF for IS is not suf-
ciently high [2], MMF monotherapy was rarely used
within the first year of LT at our institution. After in-
troduction of everolimus, a triple combination of MMF
and everolimus with very low-dose tacrolimus was se-
lectively used in recipients showing progressive renal
dysfunction during the first year post LT [19-22].

Considering that most of these recipients undergoing
MMF monotherapy had renal dysfunction, meticulous ad-
justment of MMF dosage is necessary because renal dys-
function interferes with metabolism and excretion of MPA
[1-3,9,10].

The third indication for everolimus use in our in-
stitution is refractory rejection. The primary indications
for the use of mTORi after LT are reported, in decreas-
ing order, as follows: renal failure, prevention of HCC
recurrence, recurrent HCC, de novo malignancies, dia-
abetes, hypertension, neurotoxicity, and others [23].

However, the common indications for switching to ever-
olimus are reported to be refractory rejection, extended
HCC, de novo malignancies, severe neurotoxicity, HCC
recurrence, renal insufficiency, and others [24]. After
we experienced a few cases showing marked therapeutic
effect on refractory rejection, we have carefully added
everolimus to tacrolimus in patients presenting with un-
explained hepatic dysfunction.

Everolimus can induce various adverse side–effects. In
the H2304 study, the incidence of severe adverse side–effects was 29.7% in the everolimus–reduced ta-
crolimus group, which is higher than the 21.5% reported
in the tacrolimus control group [18]. There are some
other potential conflicts of interest relevant to this article were
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Conflict of Interest
Shin Hwang is an editorial board member of the journal
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