Objectives: Tacrolimus is an effective immunosuppressant, safely administered in clinical practice by monitoring blood levels. In experimental transplants, many dosage regimes have been reported, often without such determinations. Anorexia and organ toxicity commonly occur. We report the toxic effects of tacrolimus in rabbits receiving intramuscular injections (1 mg/kg/d) and the subsequent dosage modifications that resulted in improved animal survival without toxic effects.

Materials and Methods: To obtain nontoxic drug concentrations in the blood, 3 dosage regimens were required. Drug concentrations were targeted using therapeutic human values as a guide (range, 5-20 ng/mL). First, a group of 12 Dutch Belted rabbits received vascularized femoral allografts and were treated with intramuscular dosages of tacrolimus (1 mg/kg/d) for 14 days. Subsequently, dosage reductions in 10 more rabbits, to 0.2 mg/kg/d for 14 days, were necessary. Finally, another group of 20 rabbits was treated with 0.08 mg/kg for 3 days, and then every other day thereafter. Weight loss > 30%, cardiopulmonary failure, and/or creatinine levels > 221 µmol/L were the criteria approved by our local Institutional Animal Care and Use Committee for euthanizing the animals. Treated animals were compared with 20 nonimmunosuppressed controls that underwent the same operation.

Results: At an intramuscular dosage of 1 mg/kg/d, the mean tacrolimus blood level was 90.7 ng/mL. Ten of the 12 animals in the original group died or required euthanasia. At necropsy, renal failure, cardiac abnormalities, and pulmonary edema were found. The tacrolimus dosage of 0.2 mg/kg/d produced a mean tacrolimus blood level of 17.6 ng/mL; however, 8 of the subsequent 10 rabbits died when given this dosage. Ultimately, the 0.08 mg/kg regimen in 20 rabbits permitted survival of 18 animals with a mean tacrolimus blood level of 6.8 ng/mL. None of 20 nonimmunosuppressed controls died after surgery.

Conclusions: For successful immunosuppression, Dutch-Belted rabbits require intramuscular tacrolimus dosages lower those required in other rabbit breeds. This has not been reported previously. The 0.08 mg/kg/d dosage combined with intermittent drug level monitoring permits survival without significant complications.

Key words: FK-506, Adverse effect, Adverse event, Immunosuppression, Bone allograft
White Japanese rabbits and rabbits of breeds not further specified (range, 0.1-1.6 mg/kg) (Table 1) [6-11]. Unacceptably high mortality resulted, necessitating adjustments in the drug dosage. We believe our experience will benefit other researchers using tacrolimus in Dutch-Belted and other rabbit models.

Materials and Methods

Animals
Dutch Belted (recipient) and New-Zealand White (donor) rabbits (*Oryctolagus cuniculus*; Myrtles Rabbitry, Thompson Station, TN, USA) were housed in an animal facility accredited by the Association for Assessment and Accreditation for Laboratory Animal Care, International. Rabbits were acclimated for at least 2 days prior to experimental manipulation to recover from the stress induced by transportation and the change in environment. All animals received water ad libitum and approximately 170 g standard laboratory rabbit chow daily (LabDiet, rabbit diet 5321, PMI Nutrition International, Richmond, IN, USA). Loose hay, apples, bananas, yogurt, and carrots were provided as nutritional supplements and enrichment to encourage the rabbits to eat, especially postoperatively. All procedures were approved by the institutional animal care and use committee, and the rabbits were housed in accordance with the Guide for the Care and Use of Laboratory Animals [12].

Young adult Dutch-Belted rabbits weighing between 2 and 2.5 kg received vascularized femoral allografts harvested from New-Zealand White rabbit donors. This represents an established model with a high histocompatibility barrier [13, 14]. The Dutch-Belted rabbits were initially anesthetized with ketamine (40 mg/kg), xylazine (7 mg/kg), and acepromazine (1 mg/kg), and then, following oral tracheal intubation, spontaneous breathing anesthesia was maintained with isoflurane and 75% oxygen. Intravenous lactated Ringer’s solution was administered as needed through a marginal ear vein. Ashaved front paw was used to measure intraoperative pulse oximetry. The lower abdominal wall and both hind limbs were clipped and prepared for surgery. Sterile precautions were maintained throughout the procedure.

Transplant
The recipient rabbit’s femur was exposed via a medial approach, and a 4-cm segment was resected. An axial fascial flap (which included the ipsilateral superficial inferior epigastric vessel pedicle) was raised from the abdominal wall through an additional 5-cm long paramedian incision. A 4-cm femoral allograft including the femoral nutrient vessel was transplanted, stabilized to the host femur with two figure-8 wires proximally, two 90° opposed intraosseous wires distally, and an intramedullary Kirschner wire. Next, the fascial flap was pulled through the femoral canal with the prepared pullout suture through a small trough in the proximal femur. The nutrient pedicle of the graft was microsurgically Anastomosed in an end-to-end fashion to the recipient’s femoral vessels distal to the origin of the deep femoral artery and vein. Preservation of the deep femoral vessels maintains adequate perfusion of the lower extremity [15, 16].

Aftercare
Postoperatively, rabbits wore a collar for 14 days to prevent self-mutilation of the surgical site. They were given intramuscular buprenorphine (0.2 mg/kg) for analgesia for 2 days and 0.1 mL/kg heparin for 3 days. Additionally, subcutaneous trimethoprim and sulfadiazine (0.2 mL) was given for 3 weeks for infection prophylaxis. Tacrolimus was initially administered to 12 animals at a dosage of 1 mg/kg/d intramuscularly for 14 days. The drug (prepared from a freeze-dried powder) was reconstituted with 0.9% sodium chloride solution in 10-mL vials. Later dosages (see below) were adapted to 0.2 mg/kg/d for 14 days (10 animals), and ultimately to 0.08 mg/kg/d for 3 days and then every other day until the 14th day after surgery (20 animals). Blood specimens for drug level monitoring were taken from the ear artery of the rabbits. In 20 more rabbits, the same procedure was performed with no postoperative immunosuppressive medication (nonimmunosuppressed control group).

Results
Mean blood loss (8-10 mL) during surgery and mean operative time (3 hours) were the same in all rabbit groups. Of 12 rabbits treated with tacrolimus at a dosage of 1 mg/kg/d intramuscularly for 14 days, 4 rabbits died on postoperative days 1, 2, 14, and 17, and according to the clinical endpoints, 4 rabbits had to be euthanized on postoperative days 7, 9, 13, and 15. Two additional rabbits died owing to convulsions on postoperative days 5 and 16. Drug monitoring in this group demonstrated high blood drug levels (mean, 90.7 ng/mL; range, 47.1-146.5 ng/mL) (Figure 1). Safe and therapeutic human tacrolimus levels are reported as being between 5 and 20 ng/mL. No such information is available in animals. Necropsy revealed serous effusions in the peritoneal and thoracic cavities, hemorrhagic and edematous
lungs, and pale kidneys in all of the animals in this first experimental group; in 4 animals, the liver was found to have a mottled aspect (Figure 2).

In all specimens, the histologic findings were (a) moderate to severe myocardial degeneration with variable but considerable inflammation (Figure 3); (b) hemorrhagic and diffusely congested, edematous lungs; (c) diffuse, severe cortical necrosis of the kidneys (Figure 4); and (d) centrolobular, multifocal severe hepatocyte necrosis. The findings suggest that the cause of death in these animals was severe myocarditis leading to heart failure and pulmonary edema. Additionally, the rabbits developed anorexia in the first week after surgery. Renal failure was reflected in elevated blood urea nitrogen (up to 37 mmol/L; normal range, 3.2-9.3 mmol/L) and creatinine levels (up to 283 µmol/L; normal range, 26.5-115 µmol/L). All of the above are consistent with the toxic effects of tacrolimus [6, 10, 17, 18]. We did not see these problems in the 20 control animals.

Knowing the blood levels at the highest dosage, we estimated that a reduction in the dosage to 0.2 mg/kg/d for 14 days would lower drug levels in the blood to concentrations comparable with those in the human therapeutic range (5-20 ng/mL). At this dosage, we continued to see problems in 8 of 10 rabbits, which resulted in 4 spontaneous postoperative deaths (on postoperative days 4, 6, 14, and 15) and 4 euthanasias on postoperative days 0, 10, 13, and 18. The blood levels decreased to a mean of 17.6 ng/mL (range, 11.9-24.6 ng/mL) between days 4 and 14 (Figure 5), and the histopathological changes became less severe, with the kidneys showing only slight tubular necrosis. Again, tacrolimus-induced heart failure with pulmonary edema was determined to be the cause of death. Careful review of the operative protocols identified no other treatment differences between the groups with 1 mg/kg and 0.2 mg/kg/d dosage.

Finally, in 20 rabbits, a tacrolimus dosage of 0.08
mg/kg/d was given intramuscularly in a loading phase for 3 consecutive days and then every other day until the 14th day after surgery. Only 1 rabbit had to be euthanized on day 14 owing to hypothermia and excessive weight loss. Another animal died of a seizure on the 18th postoperative day. The mean drug level after the loading phase was 6.8 ng/mL (range, 4.5-11 ng/mL) until the 14th postoperative day, when drug was discontinued according to the study protocol. In all tacrolimus-treated rabbits, postoperative anorexia was evident by reduced food intake and low fecal output. All of the survivors eventually recovered their initial weight, as did the controls, and survived for 16 weeks until termination of the study. All of the allotransplants healed and showed new periosteum formation originating from the graft, demonstrating viability and presumably adequate immunosuppression.

Discussion

The toxicity of tacrolimus has been demonstrated in several experimental animal species [10, 19-22]. It has been described in various rabbit breeds (except Dutch-Belted), with intravenous dosages as low as 0.2 mg/kg/d [6]; in dogs, at intravenous dosages of 0.3 mg/kg/d [22] and oral dosages of 1.0 mg/kg/d, with serum levels as low as 0.1 ng/mL [19, 21]; and in rats, at dosages of 1 mg/kg/wk intramuscularly and intravenously [18] and 4 mg/kg/d intramuscularly [17]. However, blood levels of tacrolimus after intramuscular injections in living rabbits (ie, not during isolated organ perfusions [7]) have not yet been reported.

Shirbacheh and associates provided a detailed study about the pharmacokinetics and tissue deposition rates of tacrolimus after intra-arterial administration, but the study did not include blood drug levels [23]. In the current study, intramuscular injection of tacrolimus at dosages higher than 0.08 mg/kg/d for 3 days and then qod after that resulted in deleterious effects within a few days. The authors of another study that used comparable dosages, length, and route of administration reported similar survival rates and no deleterious effects [9]. Unfortunately, that study provided no data about tacrolimus with regard to animal deaths, drug levels, or toxic effects [9]. The renal failure, convulsions, and cardiac hypertrophy that ultimately proved fatal to many of our rabbits mirror the clinical findings in humans and rabbits with tacrolimus overdose [6, 10, 24]. The gastrointestinal signs in our animals resemble those found in dogs [21, 22]. The non-immunosuppressed rabbits in our study showed no signs of organ failure, and none of the preceding pathological gross findings were present at 16 weeks.

Table 1 illustrates a variety of tacrolimus dosage regimens that have been used in experimental surgery. Relatively little data on rabbits exist. None of the studies in Table 1 used Dutch-Belted rabbits. Owing to possible different drug sensitivities among various species, we selected our initial dosage from the range of available data [6, 9-11]. Also, only short-term immunosuppression was necessary according to our study protocol, and the cumulative dosages and exposure times were comparatively low. As we treated only Dutch-Belted rabbits with tacrolimus, we cannot prove a similar susceptibility in other rabbit breeds, but our results certainly support cautious use of this drug in rabbits in general.

The pharmacokinetics of tacrolimus in rabbits is similar to that in humans, with a high percentage of intraerythrocyte binding and variable oral uptake [7, 8, 25]. Our data indicate variable tacrolimus concentrations after intramuscular administration as well (Figure 1). This variability, however, is acceptable at the nontoxic dosage of 0.08 mg/kg/d (Figure 5). The intramuscular route in rabbits is reliable and certainly is more convenient than intravenous injections given over several days, as the ear veins are easily subject to thromboses after repeated injections.

Extensive surgical procedures in rabbits can be problematic owing to difficulties in maintaining proper intravenous or inhalatory anesthesia levels and managing fluid and electrolyte levels, owing to rabbits’ low cardiac reserve, their low tolerance to opioids, and the difficulty in intraoperative monitoring. Their typical postoperative anorexia can complicate their recovery [11]. In our work using rabbit autografts, at 16 weeks after surgery, all of the rabbits weighed 102% of their initial weight; this also
applied to the nonimmunosuppressed rabbits (controls) receiving an allograft with the same procedure. In both groups, operative time and blood loss were similar.

The results of our data indicate 2 things. First, compared with other laboratory animals, rabbits are exceptionally sensitive to tacrolimus and require a substantially lower dosage. Second, blood level determination is tedious but provides good information about the concentration in that species. Based on animal and graft survival rates, we consider tacrolimus levels of 5 to 10 ng/mL in whole blood to be safe and effective. Of course, the actual effectiveness at this dosage can only be confirmed by investigating the reduction of T-cell activation. We will do this in a future study. Although future studies should involve larger populations with detailed blood chemistry workups and close monitoring of blood levels of tacrolimus for periods longer than 3 weeks, our current findings are important for others performing experimental transplant studies in a rabbit model.

Table 1. Overview of several studies with systemic administration of tacrolimus in different species. The data in rabbits are scarce, whereas there is a broad range of drug dosages in rats.

<table>
<thead>
<tr>
<th>Species</th>
<th>Author</th>
<th>Procedure</th>
<th>Dosage</th>
<th>Route of administration</th>
<th>Time period</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rabbit</td>
<td>Ikebe [9]</td>
<td>bone xenograft</td>
<td>1.6 mg/kg/d</td>
<td>IM</td>
<td>10 d</td>
<td>survival 16 wk</td>
</tr>
<tr>
<td>Rabbit</td>
<td>Nomoto [6]</td>
<td>no transplant</td>
<td>0.2 mg/kg/d</td>
<td>IV</td>
<td>28 d</td>
<td>left ventricular dilation, myocarditis after 28 d, not after 10 d</td>
</tr>
<tr>
<td>Rabbit</td>
<td>Piechoszewski [7, 8]</td>
<td>no transplant</td>
<td>0.5 mg/kg</td>
<td>IV</td>
<td>single dose</td>
<td>about 30 ng/mL after 24 h</td>
</tr>
<tr>
<td>Rabbit</td>
<td>Minamino [10]</td>
<td>no transplant</td>
<td>0.1-0.4 mg/kg/d</td>
<td>IV</td>
<td>4 wk</td>
<td></td>
</tr>
<tr>
<td>Rabbit</td>
<td>Hisamatsu [11]</td>
<td>trachea allograft</td>
<td>2.5 mg</td>
<td>local gel</td>
<td>single dose</td>
<td>peak at 3 d (about 17 ng/mL), not detectable after 21 d</td>
</tr>
<tr>
<td>Rat</td>
<td>Ochiai [26]</td>
<td>heart</td>
<td>0.1 mg/kg/d</td>
<td>IM</td>
<td>10 d</td>
<td></td>
</tr>
<tr>
<td>Rat</td>
<td>Peizer [1]</td>
<td>bone allograft</td>
<td>1 mg/kg/d</td>
<td>IM</td>
<td>14 d</td>
<td>survival 18 wk</td>
</tr>
<tr>
<td>Rat</td>
<td>Akar [18]</td>
<td>no transplant</td>
<td>1 mg/kg/d</td>
<td>IM</td>
<td>10 ng/mL after 3 mo</td>
<td></td>
</tr>
<tr>
<td>Rat</td>
<td>Gohda [27]</td>
<td>hind limb</td>
<td>5 mg/kg</td>
<td>IM</td>
<td>single dose</td>
<td>survival 14 d</td>
</tr>
<tr>
<td>Rat</td>
<td>Nakatani [17]</td>
<td>no transplant</td>
<td>4 mg/kg/d</td>
<td>IM</td>
<td>14 d</td>
<td>after 14 d: trough 18-28 ng/mL</td>
</tr>
<tr>
<td>Rat</td>
<td>Akst [28]</td>
<td>larynx</td>
<td>1.2 mg/kg/d</td>
<td>IM</td>
<td>5 d</td>
<td>survival 100 d</td>
</tr>
<tr>
<td>Rat</td>
<td>Inoue [29]</td>
<td>none</td>
<td>1 mg/kg/d</td>
<td>PO</td>
<td>28 d</td>
<td></td>
</tr>
<tr>
<td>Rat</td>
<td>Fukunaga [30]</td>
<td>bone xenograft</td>
<td>3 mg/kg/d</td>
<td>IM</td>
<td>14 d</td>
<td>survival 8 wk</td>
</tr>
<tr>
<td>Dog</td>
<td>Venkataramanan [31]</td>
<td>no transplant</td>
<td>0.49 mg/kg</td>
<td>IM</td>
<td>single dose</td>
<td>peak about 1 ng/mL</td>
</tr>
<tr>
<td>Dog</td>
<td>Wang [22]</td>
<td>kidney</td>
<td>0.3-0.6 mg/kg/d</td>
<td>PO</td>
<td>24 d</td>
<td>trough levels &lt; 10 ng/mL despite massive adverse effects</td>
</tr>
<tr>
<td>Minipig</td>
<td>Oike [32]</td>
<td>liver</td>
<td>0.1-0.4 mg/kg bid</td>
<td>IM</td>
<td>12 d</td>
<td>adjusted daily to trough levels of 7-20 ng/mL</td>
</tr>
</tbody>
</table>

IM, intramuscular; IV, intravenous; PO, by mouth; d, day; wk, week

References


