A Double-Blind, Double-Dummy, Flexible-Design Randomized Multicenter Trial: Early Safety of Single- Versus Divided-Dose Rabbit Anti-Thymocyte Globulin Induction in Renal Transplantation

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A previous nonblinded, randomized, single-center renal transplantation trial of single-dose rabbit antithymocyte globulin induction (SD-rATG) showed improved efficacy compared with conventional divided-dose (DD-rATG) administration. The present multicenter, double-blind/double-dummy STAT trial (Single dose vs. Traditional Administration of Thymoglobulin) evaluated SD-rATG versus DD-rATG induction for noninferiority in early (7-day) safety and tolerability. Ninety-five patients (randomized 1:1) received 6 mg/kg SD-rATG or 1.5 mg/kg/dose DD-rATG, with tacrolimus-mycophenolate maintenance immunosuppression. The primary end point was a composite of fever, hypoxia, hypotension, cardiac complications, and delayed graft function. Secondary end points included 12-month patient survival, graft survival, and rejection. Target enrollment was 165 patients with an interim analysis scheduled after 80 patients. Interim analysis showed primary end point noninferiority of SD-rATG induction (p = 0.6), and a conditional probability of <1.73% of continued enrollment producing a significant difference (futility analysis), leading to early trial termination. Final analysis (95 patients) showed no differences in occurrence of primary end point events (p = 0.58) or patients with no, one, or more than one event (p = 0.81), or rejection, graft, or patient survival (p = 0.78, 0.47, and 0.35, respectively). In this rigorously blinded trial in adult renal transplantation, we have shown SD-rATG induction to be noninferior to DD-rATG induction in early tolerability and equivalent in 12-month safety. (Clinical Trials.gov #NCT00906204.)

Abbreviations: BID, twice daily; CMV, cytomegalovirus; DD-rATG, divided-dose rabbit anti–thymocyte globulin; DGF, delayed graft function; FIO2, fraction of inspired oxygen; MMF, mycophenolate mofetil; MPA, mycophenolic acid; PO, orally; PRA, panel reactive antibody; rATG, rabbit anti–thymocyte globulin; SD-rATG, single-dose rabbit anti–thymocyte globulin; SEM, standard error of the mean

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Introduction

A continuing challenge in renal transplantation is finding an optimal immunosuppressive strategy that minimizes early graft dysfunction and rejection while maintaining appropriate safety. One strategy is the use of induction therapy, the administration of an agent at transplantation that will effect profound immunosuppression and reduce reperfusion-associated inflammation. Induction agents include lymphocyte-depleting polyclonal horse and rabbit anti–thymocyte globulin (rATG), alemtuzumab (anti CD-52 cytolytic antibody), and interleukin-2 receptor blockers. Although rATG was developed as an agent to deplete T cells, its manufacture results in the generation of multiple antibodies against a myriad of distinct epitopes. While studies in both primates and humans have shown that rATG does deplete T cells (1,2), several other studies have identified rATG antibodies that might prevent injury due not only to rejection but also to inflammation associated with brain death and/or reperfusion (3–6).

Because polyclonal horse and rabbit antithymocyte sera are associated with neutropenia and thrombocytopenia
and with cardiopulmonary instability, the prescribing information for rATG (in treating kidney rejection) recommends administering a series of small doses spaced at 1- or 2-day intervals (divided-dose rATG [DD-rATG]), along with pretreatment that includes corticosteroids (7). However, in primates, an intensive administration schedule of fewer, larger doses conferred more-comprehensive lymphocyte depletion than did a less-intensive regimen, both in the bloodstream and in secondary lymphoid structures (2).

Improved early renal function with deceased donor kidneys was reported when rATG administration was initiated before reperfusion (8), and in nonrandomized studies, single-dose rATG (SD-rATG) induction appeared to enable calcineurin inhibitor maintenance minimization and even complete withdrawal (9–11). In a blinded single-center trial that compared induction with rabbit versus equine DD-rATG, there was less rejection and superior graft survival in the rATG group after 10 years (12). Recent publications suggest that SD-rATG, compared with basiliximab or more-conventional divided-dose administration of the same rATG amount, may reduce the frequency of delayed graft function (DGF) or improve recovery of renal function in recipients of deceased-donor kidneys (1,13,14). If SD-rATG and DD-rATG are equally safe, these studies suggest that the SD-rATG confers greater benefit and may be a superior regimen.

The beneficial properties associated with rATG as both an induction agent and a treatment for rejection have led to its widespread application in solid organ transplantation; however, its optimal dosing has not been adequately investigated in randomized double-blind trials. The paucity of such trials addressing this issue is likely due to the perceived difficulty of maintaining blinding when one arm of the trial is expected to reveal itself through the frequency of obvious side effects (i.e. fevers, hypotension, etc.) (7,12,15). We believed that the difficulty of maintaining blinding could be overcome and hypothesized that SD-rATG induction is not inferior to DD-rATG induction. Here, we report the early tolerability and 12-month safety of SD-rATG induction at renal transplantation in a prospective randomized double-blind double-dummy multicenter trial that included the flexible design provision of a mid-point interim analysis to evaluate trial continuation futility (ClinicalTrials.gov NCT00906204, registered May 19, 2009).

**Methods**

**Study design**

This prospective randomized double-blind double-dummy multicenter trial in renal transplant recipients was designed to establish the noninferiority of 6 mg/kg SD-rATG induction compared with DD-rATG induction (four daily doses of 1.5 mg/kg) in early tolerability and 12-month safety. The study investigates the off-label use of Thymoglobulin® (anti-thymocyte globulin [rabbit]) (Sanofi Genzyme, Cambridge, MA) for immunosuppression induction. Patients were assessed and data were collected daily for up to 7 days after transplantation and at regularly scheduled clinic visits on days 21, 42, 90, 180, 270, and 365. The primary author designed the trial and supervised data collection and analysis (R.B.S.). The authors vouch for the data analysis and manuscript content (Figure 1).

**Inclusion, exclusion, randomization, and enrollment**

Potential study subjects undergoing renal transplantation were approached and evaluated for enrollment by nurse research coordinators if they were capable of giving written informed consent and appeared to meet study criteria (Figure 1). Excluded were multorgan transplant recipients or those planned for pancreas transplantation and patients with recent myocardial infarction (<6 months) or unstable cardiovascular disease, malignancy within 5 years (except nonmetastatic basal or squamous cell skin carcinoma, or successfully treated carcinoma in situ of the cervix), hepatitis B or C virus or HIV infection, or active liver disease. Also excluded were patients who were pregnant or breastfeeding, had previous treatment with rATG or hypersensitivity or extensive exposure to rabbits, or who had any condition that in the investigator’s opinion might compromise study participation.

Study participants were randomly allocated (1:1) by using StudyTRAX, a web-based online randomization and clinical data collection system (ScienceTRAX, Macon, GA). Subjects were stratified on donor type (deceased vs. living), nonwhite/Asian, and extended criteria donor Nyberg donor quality score A, B, or C.

**Preparation of blinded and double-dummy induction therapy**

Each patient’s study group assignment was generated automatically by the StudyTRAX program and communicated via e-mail only to the investigational pharmacy staff at each study site. Patients, caregivers, and those assessing outcomes remained blinded to study group assignments. The rATG was prepared and administered according to a double-blind double-dummy protocol (Table 1). All rATG or placebo infusions were labeled rATG/Placebo whether they contained rATG or saline.

For each treatment group, the day 0/1 rATG infusion was divided into two equal preparations: 3 mg/kg for the experimental (SD-rATG) group and 0.75 mg/kg for the control (DD-rATG) group. Each of these preparations was infused over 12–14 h, the second preparation being infused immediately after the first for a total of 6 mg/kg in the SD-rATG group and a total of 1.5 mg/kg in the DD-rATG group (Table 1). Subsequent rATG infusions in the DD-rATG group (1.5 mg/kg) were administered over 4–12 h on the 3 days after transplantation. SD-rATG recipients received equivalent administrations of a double-blind double-dummy placebo (normal saline) preparation.

Identical courses of methylprednisolone, diphenhydramine, and acetaminophen were administered with rATG/placebo to mitigate rATG side effects and preserve blinding. The first-day rATG infusion consisted of methylprednisolone 3 mg/kg IV every 6 h, administered to a maximum dose of 1.2 g in 24 h. Subsequent rATG/placebo infusions consisted of methylprednisolone 3 mg/kg IV administered during each rATG or placebo infusion. The total exposure to methylprednisolone for all study patients was 15 mg/kg, not to exceed 1.5 g total.

**Maintenance immunosuppression therapy**

Patients were maintained on tacrolimus and mycophenolate mofetil (MMF) or mycophenolic acid (MPA) (Table 2). Target tacrolimus trough blood levels were 8–12 ng/mL for months 0-3 and 6-10 ng/mL after 3 months. MMF/MPA agents were initiated on postoperative day 1 (MMF/MPA...
dosage targets: 1000/720 mg orally (PO) twice daily (BID) and adjusted according to the study protocol for leukopenia and/or renal function. To increase enrollment, the study protocol was amended after 48 patients to allow long-term administration of corticosteroids. Because all patients received steroids during assessment for the primary end point (posttransplantation days 0–7), this amendment did not affect the primary end point.

Prophylaxis against infection

Standard antifungal, antibacterial, and Pneumocystis jiroveci pneumonia prophylaxis was administered based on institutional standards of care:

- Antiviral—valganciclovir or equivalent
- Antifungal—clotrimazole or fluconazole
- Pneumocystis pneumonia—sulfamethoxazole-trimethoprim, dapsone, or aerosolized pentamidine

End points

The composite primary end point was assessed in hospitalized patients over days 0–7 after transplantation (Figure 2). The five primary composite end point components were (a) fever (≥38.9°C), (b) hypotension (systolic blood pressure <90 mmHg requiring vasopressor treatment), (c) hypoxia (increase in FiO₂ to >60% after rATG initiation or after transplantation, FiO₂ ≥50%, or nasal cannula delivering ≥3 L, either singly or combined for >12 of 24 h), (d) cardiac events (myocardial infarction or significant dysrhythmia), and (e) DGF (dialysis within 7 days of transplantation).

For the interim analysis, patients not dialedyzed but exhibiting slow-to-function grafts were classified as having DGF. Only patients dialedyzed within 7 days were included as having DGF in the final analysis. The hypothesis tested was that SD-rATG is statistically noninferior to DD-rATG in frequency of the composite primary early tolerability end point. The overall level of significance (α level) for this study is α = 0.05.
Divided-dose rATG group assuming an overall significance cannot difference in the rates of early rATG-associated adverse events, clinically significant (1). In this analysis, we determined that 75 subjects per analyzed by using either the Pearson \( \chi^2 \) or Fisher exact test; survival was analyzed by Kaplan–Meier estimation, and repeated measures by using a general linear model. Differences in means were tested by t-test, and values of \( p < 0.05 \) were considered significant.

**Planned noninferiority and futility analyses at study mid-point**
The Data and Safety Monitoring Committee continually assessed the safety of study subjects. Noninferiority and futility were analyzed after 80 patients by using the one-tailed Fisher exact test and a conditional power analysis of primary event rate trends, respectively, with early termination of the trial to be considered if conditional power to achieve a significant difference with continued enrollment fell below 15%.

**Results**

**Patient enrollment and recipient and donor characteristics**
Four transplant centers in the United States enrolled 97 patients between March 30, 2010, and March 25, 2014; 45 were randomized to receive SD-rATG, and 52 were randomized to receive DD-rATG (Figure 1). One patient in each group did not receive the allocated rATG induction regimen, leaving 95 patients for primary composite end point analysis. (Figure 1). There were no significant differences in donor characteristics (Table 3). SD-rATG recipients had higher panel reactive HLA class 2 antibody (PRA) levels, both peak and at transplantation.

**rATG and steroid exposure, and posttransplantation hyperglycemia**
The dosing schedule for rATG/placebo and concomitant medications is shown in Table 1. Total induction rATG exposure was similar between groups: SD-rATG 5.8 ± 0.7 versus DD-rATG 5.8 ± 0.6 mg/kg (\( p = 0.99 \)). During the first year, 15 (34%) of 44 SD-rATG patients and 15 (29%) of 51 DD-rATG patients received

| Table 1: Rabbit anti-thymocyte globulin (rATG) and concomitant medication dosing |
|---------------------------------------------------|
| **Single-dose rATG group**                        |
| rATG                                             |
| Methylprednisolone 6 mg/kg over 24 h              |
| Methylprednisolone 3 mg/kg IV q 6 h (12 mg/kg total) |
| Diphenhydramine 25 mg IV or 50 mg PO pre-med      |
| Acetaminophen 650 mg PO q 6 h                     |
| **Divided-dose rATG group**                      |
| rATG                                             |
| Methylprednisolone 1.5 mg/kg over 24 h           |
| Methylprednisolone 3 mg/kg IV q 6 h (12 mg/kg total) |
| Diphenhydramine 25 mg IV or 50 mg PO pre-med      |
| Acetaminophen 650 mg PO q 6 h                     |
| **SD-rATG**                                       |
| **DD-rATG**                                       |
| **p-value**                                       |
| Tacrolimus (ng/mL)                               |
| 1–3                                               |
| 1–3                                               |
| 6–12                                              |
| 6–12                                              |
| Methylprednisolone 1 mg/kg at time of rATG/placebo infusion |
| Diphenhydramine 25 mg IV or 50 mg PO pre-med      |
| Acetaminophen 650 mg PO pre-med and PRN           |
| SD-rATG, single-dose rabbit anti-thymocyte globulin; DD-rATG, double-dose rabbit anti-thymocyte globulin; MMF, mycophenolate mofetil; MPA, mycophenolic acid. \(^1\)MMF expressed as MMF equivalent. |
long-term prednisone therapy \( (p = 0.66) \). Average total prednisone exposure was SD-rATG \( 2.5 \pm 1.0 \) g and DD-rATG \( 2.7 \pm 1.3 \) g \( (p = 0.69) \). Nine patients began long-term steroid use in association with rejection treatment (five patients SD-rATG and four patients DD-rATG).

The prevalence of pretransplantation diabetes (with or without insulin dependence) was not different between groups \( (p = 1, \text{ Table } 3) \). Of patients without pretransplantation diabetes, 6 (18%) of 34 SD-rATG and 8 (20%) of 40 DD-rATG patients received insulin within 4 days of transplantation \( (p = 1.0) \).

**Maintenance immunosuppression**

Mean immunosuppressant trough levels and MMF-MPA exposure accorded with study design and were not statistically different, with an equal and significant reduction
in tacrolimus levels over time for both groups (Δ tacrolimus: all patients, p < 0.001; SD-rATG vs. DD-rATG, p = 0.70) (Table 2).

**Interim composite primary end point analysis**

A planned primary end point analysis (early safety and tolerability of SD-rATG) including a futility evaluation by conditional power analysis was performed at the enrollment mid-point. Rates for each of the five primary end point component events in the SD-rATG group were shown to be noninferior to their rates in the DD-rATG group by one-tailed Fisher exact test, as was the case when these rates were composited for analysis (p = 0.51) (Table 4). These analyses were performed without equivalence margin adjustment to add stringency in evaluating the advisability of early trial termination.

The conditional power analysis showed only a <1.73% probability that trial continuation would alter the conclusion that induction using SD-rATG is not inferior to DD rATG. All patients not withdrawn from the trial (SD-rATG 42 of 44 patients and DD-rATG 49 of 51 patients) were followed for 12 months (Figure 1).

**Final composite primary end point analysis**

The hypothesis in this noninferiority trial is that there is not a statistically greater frequency of occurrence of the primary end point among the SD-rATG patients. There was no significant difference in the rate of primary end point events between the two induction groups in any of the individual event rates or, using the Pearson χ² test, the overall proportions of all five events (Figure 2A). Even considering multiple safety end points per patient, there was no significant difference between the groups (p = 0.81).

**Twelve-month secondary end points: patient survival, graft survival, rejection, renal function**

There were no significant differences in these 12-month safety end points (Figures 2B–E). Five patients in each group experienced biopsy-proven acute rejection during the first year, all except one within the first month. There were three Banff IA and two Banff IIA events among SD-rATG patients and one Banff IA and four Banff IIA events in the DD-rATG group. Of the 10 rejection episodes, seven were steroid sensitive (four SD-rATG patients and three DD-rATG patients) and three were steroid resistant (one S-rATG patient and two DD-rATG patients).

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**Table 3: Recipient and donor characteristics**

|                        | Single-dose rATG (n = 44) | Divided-dose rATG (n = 51) |
|------------------------|---------------------------|---------------------------|
| **Recipients**         |                           |                           |
| Age (years)            | 48.0 ± 11.8               | 49.0 ± 12.4               |
| Body mass index (kg/m²)| 28.2 ± 4.6                | 27.9 ± 5.2                |
| Body mass index >30 kg/m² | 15 (34%)           | 15 (29%)                   |
| Males                  | 29 (66%)                  | 38 (75%)                   |
| Pretransplantation diabetes (types 1 and 2) | 10 (23%) | 11 (22%) |
| Pretransplantation diabetes (insulin dependent) | 5 (11%) | 4 (8%) |
| Pretransplantation diabetes (not insulin dependent) | 5 (11%) | 7 (14%) |
| Living donor           | 26 (59%)                  | 29 (57%)                   |
| Living donor related/unrelated | 14/12       | 11/18                     |
| Deceased donor         | 18 (41%)                  | 22 (43%)                   |
| Deceased donor cold ischemia (h) | 17.1 ± 7.6        | 18.8 ± 7.4                |
| PRA class 1 peak (%)   | 7.2 ± 16.4                | 4.4 ± 10.1                 |
| PRA HLA class 1 at transplantation (%) | 4.3 ± 12.2 | 2.7 ± 8.1 |
| PRA class 2 peak (%)   | 13.3 ± 25                 | 5.3 ± 12.7                 |
| PRA HLA class 2 at transplantation (%) | 7.6 ± 17     | 1.3 ± 4.8 |
| Donor/recipient height ratio | 1.0 ± 0.1      | 1.0 ± 0.1 |
| Antigen mismatch       | 4.6 ± 1.5                 | 4.1 ± 1.8                  |
| Race (nonwhite/Asian)  | 12 (27%)                  | 11 (22%)                   |
| CMV serostatus (D+/R– or D+/R+) | 25 (58%)     | 29 (59%) |
| **Donors**             |                           |                           |
| Age at procurement (years) | 39.5 ± 13.1             | 40.6 ± 14.2               |
| Males                  | 19 (43%)                  | 24 (47%)                   |
| Body mass index (kg/m²)| 27.8 ± 7.1                | 29.0 ± 7.2                 |
| Deceased donor final creatinine (mg/dL) | 1.2 ± 0.8     | 1.0 ± 0.5                  |
| Deceased donors <18 years of age | 1 (2%)            | 1 (2%)                     |
| Donation after cardiac death | 3 (7%)                | 3 (6%)                     |

rATG, rabbit anti–thymocyte globulin; PRA, panel reactive antibody; CMV, cytomegalovirus; D, donor; R, recipient.

*p = 0.05.

†p = 0.01.
Incomplete induction events

The administration of rATG deviated from the targeted rATG dose in three single-dose patients (7%) and six divided-dose patients (12%) ($p = 0.50$). Among SD-rATG patients, one patient’s infusion was halted due to a severe rATG reaction, one patient’s infusion was mistakenly terminated when 90% complete, and one patient was administered alemtuzumab instead of rATG. Among DD-rATG recipients, one patient’s fourth rATG dose was delayed 2 days due to neutropenia, one received only two of four doses due to thrombocytopenia, the fourth dose was discontinued in two patients (one for pancytopenia and one for a possible allergic reaction [not protocol violations]), one patient received only half of the first rATG dose due to a nursing error (protocol violation), and one patient received no rATG and was not transplanted due to poor donor kidney quality.

Frequency of noninfectious and infectious adverse events at 12 months

The frequency of clinically relevant adverse events, including serious adverse events, was not statistically different between the induction groups at 12 months (Table 5). A detailed breakdown of all serious adverse events and other adverse events with a frequency above 5% is presented in the supplemental data set. There was a trend toward more frequent leukopenia requiring treatment in the DD-rATG group ($p = 0.13$).

Hematologic effects of rATG induction

There was a similar severity of leukopenia between the rATG groups during the first 12 months after transplantation (Figure 3A) and an equivalent reduction in the absolute lymphocyte count in both groups, with a similar rate of recovery (Figure 3B). In both rATG groups, absolute neutrophil counts were significantly reduced immediately after induction, with a more rapid early recovery in the SD-rATG group (Figure 3C). Postinduction absolute monocyte counts and hemoglobin levels were similar between induction groups (Figures 3D and E). In the SD-rATG group, there was a greater immediate reduction in platelet counts with a more robust recovery; average platelet count on days $1–4$ was $123 ± 5103/μm^3$ on SEM) for SD-rATG patients versus $145 ± 6.0$ for DD-rATG patients ($p = 0.02$) (Figure 3F).

Discussion

While rATG is US Food and Drug Administration approved only for treating rejection in adult renal transplant recipients, it is nonetheless the most frequently used agent for induction immunosuppression in the United States (16). Dosing for induction usually mimics the recommendation for treating acute rejection (i.e. daily doses of 1.5 mg/kg) but with a shortened duration of 4–7 days. It is important to note that this dosing practice was not derived from systematic comparison of possible alternatives. There has never been a multicenter blinded double-dummy trial of rATG induction, in large part because investigators have believed that group concealment could not be maintained.

Conclusions drawn from nonblinded trials of rATG induction might reflect an impact of investigator bias in managing peri-induction adverse events. For example, an investigator might adjust maintenance immunosuppression to compensate for an induction regimen believed to be more (or less) immunosuppressive, which could distort the conclusions reached by the trial (e.g. rejection or rejection graft function; SD-rATG, single-dose rabbit anti–thymocyte globulin; DD-rATG, divided-dose rabbit anti–thymocyte globulin.

**Table 4:** Primary end point events at trial interim analysis

| Event                  | SD-rATG, with/without (%) with | DD-rATG, with/without (%) with | p-value |
|------------------------|--------------------------------|--------------------------------|---------|
| Fever                  | 5/34 (13%)                     | 4/41 (9%)                       | 0.41    |
| Hypotension            | 3/36 (8%)                      | 1/44 (2%)                       | 0.27    |
| Hypoxia                | 3/36 (8%)                      | 3/42 (7%)                       | 0.59    |
| Cardiac events         | 3/36 (8%)                      | 1/44 (2%)                       | 0.26    |
| DGF                    | 4/35 (10%)                     | 9/36 (25%)                      | 0.94    |
| Patients with event(s)| 13/26 (33%)                    | 14/31 (31%)                     | 0.51    |

DGF, delayed graft function; SD-rATG, single-dose rabbit anti–thymocyte globulin; DD-rATG, divided-dose rabbit anti–thymocyte globulin.

**Table 5:** Adverse events during first 12 months after transplantation

| Patients with, No. | Single-dose rATG (n = 44) | Divided-dose rATG (n = 51) | p-value |
|--------------------|---------------------------|-----------------------------|---------|
| Any event          | 44 (100%)                 | 49 (96%)                    | 0.5     |
| Serious events     | 23 (52%)                  | 21 (41%)                    | 0.31    |
| Leukopenia         | 12 (27%)                  | 22 (43%)                    | 0.13    |
| Anemia             | 14 (32%)                  | 14 (27%)                    | 0.66    |
| Thrombocytopenia   | 14 (32%)                  | 10 (20%)                    | 0.24    |
| Serum sickness     | 0                         | 1 (2%)                      | 1       |
| All infections     | 23 (52%)                  | 20 (39%)                    | 0.22    |
| Urinary tract      | 8 (18%)                   | 7 (14%)                     | 0.58    |
| Bacterial          | 7 (16%)                   | 5 (10%)                     | 0.54    |
| CMV                | 1 (2%)                    | 1 (2%)                      | 1       |
| BK viremia         | 6 (14%)                   | 6 (12%)                     | 1       |
| BK nephropathy     | 0                         | 1 (2%)                      | 1       |
| Fungal             | 1 (2%)                    | 0                            | 1       |
| Cancer             | 1 (2%)                    | 2 (4%)                      | 1       |
| Basal cell carcinoma | 1 (2%)                    | 0                            | 1       |
| Chronic            | 0                         | 1 (2%)                      | 1       |
| Lymphocytic leukemia | 0                         | 1 (2%)                      | 1       |
| Prostatic cancer   | 0                         | 1 (2%)                      | 1       |

rATG, rabbit anti–thymocyte globulin; PRA, panel reactive antibody; CMV, cytomegalovirus; D, donor; R, recipient.
infection rates, etc.). Clinical investigators often address this possibility by comparing immunosuppressant blood levels between treatment arms, but simply comparing average levels at widely spaced intervals may obscure significant effects of adjustments made across shorter time frames. This problem can be addressed with sophisticated numerical modeling, but even modern computing power and software cannot adequately compensate for the small enrollment numbers typically available to a transplantation clinical trial.

We included a planned interim analysis so that the trial could be modified in light of findings at the enrollment midpoint-point. This approach, which potentially reduces expense, has become recognized as an effective strategy for increasing the number of clinical trials and accelerating the introduction of novel agents into clinical use (17). This approach especially benefits so-called orphan conditions, which includes transplantation. Planning an interim analysis is beneficial whenever the anticipated effect size of an experimental therapeutic innovation has
substantial uncertainty, which is often the case with organ transplantation. In our case, we elected early trial termination due to the clear demonstration of noninferiority and a very low probability of that finding changing with continued enrollment.

We previously reported in a randomized single-center but nonblinded trial with early steroid withdrawal and tacrolimus/sirolimus maintenance that SD-rATG induction compared with DD-rATG was associated with improved renal function, fewer infections, and improved patient survival (1,14). However, despite multiple reports demonstrating reasonable safety and possible medical advantages to SD-rATG induction in renal transplantation, there remains a concern in the transplant community as to the early safety of this approach (9,10,13,18,19). The present multicenter study was designed to rigorously address these remaining safety concerns using a primary composite end point based on either the physiologic consequences of a severe rATG reaction (fever, hypotension, hypoxia) or the possible severe sequelae (cardiac events and DGF) (20–24). This prospectively randomized double-blind trial also used a double-dummy approach to help guarantee group concealment and eliminate clinician bias in interpreting and managing complications during and after rATG induction.

For the first time in a double-blind double-dummy clinical trial, we have demonstrated the noninferiority of SD-rATG induction. Additionally, the secondary 12-month safety end point of patient survival, death-censored graft survival, rejection-free patient survival, and renal allograft function were nearly identical. The frequency of serious infectious and noninfectious adverse events and the hematologic impact of SD-rATG induction also were not significantly different between the induction groups.

There are practical advantages to completing rATG induction within 24 h of transplantation. First, patients receiving the single dose are intensively monitored for early complications for the entire dosing period. In this trial, it is interesting to note that there were fewer errors of administration in the SD-rATG group. Second, SD-rATG may have hematologic advantages over divided dosing. Patients in the SD-rATG group experienced fewer dose reductions due to thrombocytopenia, neutropenia, or fevers. Third, SD-rATG induction may allow for earlier discharge from the hospital, a hypothesis not testable in this trial due to the double-dummy blinding strategy, which required four infusions for all patients.

The main limitation of this trial is that it is not sufficiently powered, due to both numbers and length of follow-up, to address the longer-term secondary end points. It is noteworthy that in our previous single-center trial with an enrollment of 178, improved SD-rATG safety (patient survival and infectious complications) was noted after an average follow-up of nearly 5 years but was not apparent after only 12 months (1,14). In our current trial, the relative paucity of deceased donors and limited follow-up precluded assessing possible long-term efficacy benefits to SD-rATG induction (e.g. superior renal function). Additionally, the trial findings cannot reasonably be thought to apply to higher cardiac risk patients, as such were excluded from enrollment in this study.

With the added certainty in the safety of SD-rATG, an exciting and highly informative next step would be a prospective randomized double-blind double-dummy multicenter trial designed to address the potential benefits of induction with SD-rATG in comparison to alemtuzumab and/or basiliximab.

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Disclosure

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Single- Versus Divided-Dose rATG Induction

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Supporting Information

Additional Supporting Information may be found in the online version of this article.

Data S1: Supplemental data set.