**Mixed chimerism induced without lethal conditioning prevents T cell– and anti-Galα1,3Gal–mediated graft rejection**

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Galα1,3Gal–reactive (Gal-reactive) antibodies are a major impediment to pig-to-human xenotransplantation. We investigated the potential to induce tolerance of anti-Gal–producing cells and prevent rejection of vascularized grafts in the combination of α1,3-galactosyltransferase wild-type (GalT+/+) and deficient (GalT−/−) mice. Allogeneic (H-2 mismatched) GalT+/+ bone marrow transplantation (BMT) to GalT−/− mice conditioned with a nonmyeloablative regimen, consisting of depleting CD4 and CD8 mAb’s and 3 Gy whole-body irradiation and 7 Gy thymic irradiation, led to lasting multilineage H-2<sup>b</sup>GalT+/+ + H-2<sup>d</sup>GalT−/− mixed chimerism. Induction of mixed chimerism was associated with a rapid reduction of serum anti-Gal naturally occurring antibody levels. Anti-Gal–producing cells were undetectable by 2 weeks after BMT, suggesting that anti-Gal–producing cells preexisting at the time of BMT are rapidly tolerated. Even after immunization with Gal-bearing xenogeneic cells, mixed chimeras were devoid of anti-Gal–producing cells and permanently accepted donor-type GalT+/+ heart grafts (>150 days), whereas non-BMT control animals rejected these hearts within 1–7 days. B cells bearing receptors for Gal were completely absent from the spleens of mixed chimeras, suggesting that clonal deletion and/or receptor editing may maintain B-cell tolerance to Gal. These findings demonstrate the principle that induction of mixed hematopoietic chimerism with a potentially relevant nonmyeloablative regimen can simultaneously lead to tolerance among both T cells and Gal-reactive B cells, thus preventing vascularized xenograft rejection.

state of mixed chimerism that is associated with specific tolerance of anti-Gal NAb–producing B cells (19). However, lethal irradiation is not a conditioning treatment that would be considered reasonable for use in humans needing organ transplantation. We now demonstrate that mixed chimerism, with vascularized GalT<sup>−/−</sup> donor heart graft acceptance, can be induced in GalT<sup>−/−</sup> mice using a more clinically relevant, less toxic, nonmyeloablative conditioning regimen, which does not include specific treatments to remove preexisting host anti-Gal–producing cells. Anti-Gal–producing cells were undetectable by 2 weeks after BMT, suggesting that anti-Gal–producing cells preexisting in the GalT<sup>−/−</sup> recipients at the time of BMT are rapidly tolerated by the induction of mixed chimerism. In addition, we provide data suggesting that a state of B-cell tolerance to Gal may be maintained by clonal deletion and/or receptor editing in mixed chimeras.

**Methods**

**Animals.** GalT<sup>−/−</sup> (H-2<sup>d</sup>) mice and GalT<sup>+/−</sup> (H-2<sup>b</sup>sd and H-2<sup>d</sup>) mice were derived from hybrid (129SV × DBA/2 × C57BL/6) animals (20). All mice used in this study were confirmed by flow cytometric (FCM) analysis to express homozygous levels of the Ly-2.2 allele. C.B.17 scid/scid (C.B.-17 scid) mice were purchased from the Department of Radiation Oncology, Massachusetts General Hospital (Boston, Massachusetts, USA). All mice were maintained in a specific pathogen-free microisolator environment.

**Conditioning and BMT.** Age-matched (8- to 12-week-old) GalT<sup>+/+</sup> (H-2<sup>b</sup>) recipient mice were intraperitoneally injected with 1.8 mg and 1.4 mg of rat IgG<sub>2b</sub> anti-mouse CD4 mAb GK1.5 (21) and anti-mouse CD8 mAb 2.43 (anti–Ly-2.2 mAb) (22), respectively, on day –5 of BMT. On day 0, 3 Gy whole-body irradiation and 7 Gy selective thymic irradiation were given to mAb-treated animals, as described (23). Bone marrow cells (BMCs) from GalT<sup>+/+</sup> (H-2<sup>b</sup>sd) donors were depleted of T cells, using anti-CD4 and anti-CD8 mAb’s and rabbit complement as described, and were administered intravenously on day 0 (24).

**FCM analysis of chimerism.** Chimerism was evaluated by 2-color FCM analysis of peripheral white blood cells (WBCs) on a FACScan cytometer (Becton Dickinson Immunocytometry Systems, Mountain View, California, USA) as described (25). Cells were stained with FITC-conjugated anti-CD4 (PharMingen, San Diego, California, USA), anti-CD8 (Caltag Laboratories Inc., San Francisco, California, USA; and PharMingen), B220 (PharMingen), and anti-MAC1 (Caltag Laboratories Inc.) mAb’s, together with biotinylated anti–donor mouse H-2K<sub>b</sub> mAb 5F1 (26). The biotinylated mAb was viewed with phycoerythrin-streptavidin (PE-streptavidin). Forward angle and 90° light scatter properties were used to distinguish lymphocytes, granulocytes, and monocytes in WBCs. The percentage of donor cells (staining with 5F1) was calculated separately for each cell population.

**FCM analysis of anti-Gal and anti-rabbit red blood cell antibodies.** Indirect immunofluorescence stainings of C.B.-17 scid mouse spleen cells and BMCs (which express the Gal epitope and lack surface Ig<sup>+</sup> B cells) or rabbit erythrocytes were used to detect anti-Gal NAb’s and anti-rabbit erythrocyte antibodies, respectively. One million cells were incubated with 10 μL of serially diluted mouse serum, washed, and then incubated with FITC-conjugated rat anti-mouse IgG mAb (PharMingen). The specificity for anti-Gal NAb in sera of GalT<sup>+/−</sup> mice, as detected by staining of scid mouse cells in this assay, has been verified by anti-Gal NAb–specific ELISA assay, which revealed a strong correlation between data obtained by these 2 methods (r > 0.90) (19).

**ELISA assays.** Anti-Gal levels in sera were quantified by ELISA according to procedures described previously (19). Total immunoglobulin levels in sera were also quantified by ELISA. ELISA plates were coated with 5 μg/mL of goat anti-mouse IgG or IgM (Southern Biotechnology Associates Inc., Birmingham, Alabama, USA). Diluted serum samples were incubated in the plates, and bound antibodies were detected using horse-radish peroxidase–conjugated goat anti-mouse IgG and substrate.

**Data analysis.** All data are expressed as mean ± SEM. Statistical significance was determined by Student’s t test.
immunoglobulin (250 ng/mL; Southern Biotechnology Associates Inc.). Color development was achieved using 0.1 mg/mL o-phenylenediamine dihydrochloride (OPD; Sigma Chemical Co., St. Louis, Missouri, USA) in substrate buffer. The OPD reaction was stopped using 3 M NH₂SO₄, and absorbance at 492 nm was measured.

Enzyme-linked immunospot for detecting anti-Gal–producing cells. The enzyme-linked immunospot (ELISPOT) assay was performed as described previously (19). Briefly, nitrocellulose membranes from a 96-well filtration plate (Millipore Corp., Bedford, Massachusetts, USA) were coated with 5 μg/mL of synthetic Galβ1-3Galβ1-4GlcNAc conjugated to BSA (Gal-BSA; Alberta Research Council, Alberta, Alberta, Canada). Non-specific binding sites were blocked with 0.4% BSA in culture medium. Serial dilutions of spleen, bone marrow, or peritoneal cell suspension were added to triplicate wells. After a 24-hour culture at 37°C, bound antibodies were detected using horseradish peroxidase–conjugated goat anti-mouse IgM plus IgG antibodies (Southern Biotechnology Associates Inc.), followed by color development with 3-amino-9-ethyl carbazole (Sigma Chemical Co.).

FCM analysis and cell sorting of B cells bearing receptors for Gal. One million cells per 100 μL were incubated with 0.5 μg/100 μL FITC-conjugated Gal-BSA (Alberta Research Council) or 0.5 μg/100 μL control FITC-conjugated BSA (Fisher Scientific Co., Pittsburgh, Pennsylvania, USA) in FCM medium for 2 hours at 4°C, together with biotinylated anti-mouse IgM mAb (PharMingen) (for 2-color FCM) or PE-conjugated anti-CD19 (PharMingen) and biotinylated anti-donor mouse H-2Kb 5F1 mAb’s (for 3-color FCM). FITC conjugation of Gal-BSA and BSA was performed with the QuickTag FITC Conjugation Kit (Boehringer Mannheim Biochemicals, Indianapolis, Indiana, USA) according to the manufacturer’s instructions. Biotinylated mAb’s were viewed with PE-streptavidin (for 2-color FCM) or CyChrome-streptavidin (for 3-color FCM). Based on Gal-BSA binding and IgM expression, splenic Gal-BSA+/IgM+ and Gal-BSA-/IgM+ populations were sorted under sterile conditions using a MoFlo high-speed cell sorter (Cytomation, Fort Collins, Colorado, USA). Sorted cells were immediately resuspended in culture medium and applied to ELISPOT plates precoated with Gal-BSA to determine the frequency of anti-Gal (IgM)–producing cells.

Heterotopic heart transplantation. Cervical heterotopic heart transplantation was performed using the cuff technique modified from a method described previously (27). Briefly, the recipients were initially prepared before the donor heart harvest to minimize the graft ischemic time. The right external jugular vein and the right common carotid artery were dissected free, mobilized as far as possible, and fixed to the appropriate cuffs. The cuffs were composed of polyethylene tubes.
tored by daily inspection and palpation. Rejection was determined by the cessation of beating of the graft and was confirmed by histology.

**Histological studies.** Formalin-fixed grafted heart tissue sections were stained with hematoxylin and eosin (H&E) and examined microscopically. Fresh frozen grafted heart tissue sections were stained with fluorescein-labeled anti-mouse IgG and IgM (Sigma Chemical Co.) and C3 (Cappel Research Products, Durham, North Carolina, USA) and were examined by fluorescence microscopy.

**Statistics.** The results were statistically analyzed by the unpaired or paired Student’s *t* test of means or the log rank test when appropriate. A *P* value less than 0.05 was considered to be statistically significant.

**Results**

Lasting multilineage GalT+/+→GalT−/− mixed chimera can be induced by nonmyeloablative conditioning. To determine whether GalT+/+ pluripotent hematopoietic stem cell engraftment could be achieved in GalT−/− recipients treated with a nonmyeloablative conditioning regimen that permits the development of allogeneic chimerism and T-cell tolerance across MHC barriers (23), we administered varying doses (20×106, 7.5×106, and 1×106) of T cell–depleted GalT+/+ BMCs (H2b1) to GalT−/− recipients (H-2d) conditioned with depleting anti-CD4 plus anti-CD8 mAb’s on day −5, and with 3 Gy whole-body irradiation and 7 Gy thymic irradiation on day 0. In addition to the Gal epitope, the donor strain expressed a full MHC haplotype (H-2b) not shared by the recipients. Untreated GalT−/− and GalT+/+ mice, and conditioned GalT+/+ and GalT−/− mice not receiving BMT, served as controls. In all recipients of 20×106 and 7.5×106 BMCs (n = 5 per group), and in 3 of 5 recipients of 1×106 BMCs, lasting mixed chimera was observed in peripheral blood B cells, CD4+ and CD8+ T cells, monocytes, and granulocytes at all times (Figure 1). Erythrocyte chimerism could not be detected because of low levels of Gal and H-2 expression on erythrocytes. Our data indicate that lasting multilineage GalT+/+→GalT−/− chimera could be induced with nonmyeloablative conditioning when appropriate doses of BMCs are transplanted.

**Loss of anti-Gal NAb’s in mixed GalT+/+→GalT−/− chimeras.**

The kinetics of serum anti-Gal NAb levels in each group are shown in Figure 2. In the control untreated GalT+/+ mice, serum anti-Gal NAb levels showed a gradual increase during the observation period, consistent with the age-related increase in mouse NAb described previously (28). In the control non-BMT GalT−/− mice that received conditioning treatment, serum anti-Gal NAb levels increased further at 2 weeks (*P* < 0.05 before conditioning vs. 2 weeks after conditioning) and remained high throughout the follow-up period. This increase of anti-Gal NAb levels may be due to loss of regulation by T cells, as similar results were observed in GalT−/− mice treated only with depleting anti-CD4 plus anti-CD8 mAb’s (H. Ohdan et al., manuscript in prepa-
ration). In contrast, \( \text{GalT}^{+/+} \) to \( \text{GalT}^{-/-} \) mixed chimeras had significantly reduced serum levels of anti-Gal NAb by 2 weeks after BMT. These declined further over time, eventually becoming undetectable above the level observed in control \( \text{GalT}^{+/+} \) mice.

Absence of anti-Gal–producing cells in mixed chimeras. The observation that mixed \( \text{GalT}^{+/+} \) to \( \text{GalT}^{-/-} \) chimerism led to a specific reduction in anti-Gal NAb’s suggested that \( \text{GalT}^{-/+} \) hematopoietic chimerism led to tolerance of anti-Gal NAb–producing cells. However, to rule out the possibility that anti-Gal NAb’s were merely absorbed by the Gal epitope expressed on engrafted \( \text{GalT}^{+/+} \) cells, the presence of anti-Gal–producing cells was assessed by ELISPOT assay. Eighteen to 19 weeks after BMT, chimeras were immunized by intraperitoneal injection of \( 1 \times 10^9 \) rabbit erythrocytes, which express large amounts of Gal (29). Eight days after immunization, anti-Gal (IgM and IgG)–producing cells were quantified in recipient spleen cells, BMCs, and peritoneal cavity cells. As is shown in Figure 3a, cells producing anti-Gal were undetectable in all 3 tissues of mixed chimeras, whereas large numbers of these cells were detected predominantly in the spleen of both untreated and conditioned \( \text{GalT}^{-/-} \) mice. The results in mixed chimeras resembled those from control \( \text{GalT}^{+/+} \) mice, which also lacked cells producing anti-Gal. In artificial mixtures of spleen cells from \( \text{GalT}^{+/+} \) and \( \text{GalT}^{-/-} \) mice immunized with rabbit erythrocytes, a linear relationship between the frequency of anti-Gal–producing cells and the percentages of \( \text{GalT}^{-/-} \) cells in the mixtures (\( r^2 = 0.9881 \)) was observed (data not shown), ruling out absorption of anti-Gal by \( \text{GalT}^{+/+} \) cells of mixed chimeras in the ELISPOT assay.

Consistent with the absence of anti-Gal–producing cells in mixed chimeras, serum levels of anti-Gal (both IgM and IgG) in mixed chimeras were undetectable, even after immunization with rabbit erythrocytes (Figure 3b). In mixed chimeras and conditioned \( \text{GalT}^{-/-} \) mice, total serum IgM and IgG levels were maintained at normal levels at all time points (data not shown). In all mixed chimeras, serum levels of anti-rabbit erythrocyte IgM antibody were elevated after immunization with rabbit erythrocytes, to levels similar to those detected in normal \( \text{GalT}^{+/+} \) control mice (data not shown).

Figure 4
Absence of anti-Gal–producing cells in mixed chimeras 2 weeks after BMT (ELISPOT detection of anti-Gal IgM/IgG–producing cells). Spleen cells from conditioned \( \text{GalT}^{-/-} \) mice receiving \( 2 \times 10^6 \) \( \text{GalT}^{+/+} \) BMCs (mixed chimeras), from conditioned \( \text{GalT}^{-/-} \) mice receiving \( 2 \times 10^6 \) 30 Gy–irradiated \( \text{GalT}^{+/+} \) BMCs (nonchimeras), and from normal \( \text{GalT}^{+/+} \) mice were used in ELISPOT assay. The frequency of anti-Gal–producing cells was determined as the average of red plaque numbers in triplicate wells of \( 8 \times 10^5 \) cells. Average values ± SEM for the individual groups are shown. Each point represents an individual mouse. * \( P < 0.05 \) compared with conditioned \( \text{GalT}^{-/-} \) mice receiving irradiated BMCs.

Figure 5
B cells bearing receptors for Gal detected by Gal-BSA comprise the anti-Gal–producing population in the spleens of \( \text{GalT}^{-/-} \) mice. Spleen cells were prepared from 5 normal \( \text{GalT}^{-/-} \) mice (12 weeks of age) 8 days after immunization by intraperitoneal injection of \( 1 \times 10^9 \) rabbit erythrocytes. The pooled cells were stained with FITC-conjugated Gal-BSA or control FITC-conjugated BSA, together with biotinylated anti-mouse IgM mAb and PE-streptavidin. The populations of Gal-BSA–binding and –nonbinding B cells (IgM+) were sorted as described in Methods. (a) FCM results of Gal-BSA–binding spleen cells. Sorted cells were reanalyzed for purity; 30,000 cells were analyzed for each contour plot. Percentages given are of total spleen cells. (b) ELISPOT detection of anti-Gal (IgM)–producing cells. The frequencies of anti-Gal–producing cells were determined for unsorted and sorted cells by ELISPOT assay. Numbers refer to the total cells seeded per well (\( \times 10^3 \)). Anti-Gal–producing cells were greatly enriched in the sorted Gal-BSA–binding B-cell population. The calculated frequencies of anti-Gal–producing cells were 0.1/\( 10^3 \), 0.005/\( 10^3 \), and 56/\( 10^3 \) in the unsorted, sorted Gal-BSA/IgM+, and Gal-BSA/IgM+ populations, respectively.
2Kb–negative cells were selected by gating and analyzed for the frequency of CD19+ cells staining with control FITC-conjugated BSA from the peripheral B cells was calculated by subtracting the percentage of CyChrome-streptavidin. (Derived cells were collected for each sample. (Chimeras. To ensure statistical significance, data on 100,000 host-derived spleen cells are shown. Average values ± SEM. (Normal mice that did not receive BMT, as indicated by the experiment presented in Figure 4, in which conditioned GalT–/– mice received irradiated GalT+/– BMCs had measurable anti-Gal–producing spleen cells. The frequency of anti-Gal–producing cells in the control conditioned GalT–/– mice receiving irradiated GalT+/– BMCs was similar to that of untreated normal GalT–/– mice (data not shown). Thus, mixed hematopoietic chimeraism is necessary for the induction of B-cell tolerance, as opposed to antigen exposure alone. Anti-Gal–producing cells were tolerant in mixed chimeras by 2 weeks after BMT.

Absence of B cells bearing receptors for Gal in the spleens of mixed chimeras. By surface staining with FITC-conjugated Gal-BSA, B cells bearing surface IgM (sIgM) receptors that can recognize Gal epitopes could be identified. In normal GalT+/+ mice immunized intraperitoneal with 1 × 10⁶ rabbit erythrocytes, Gal-BSA–binding B cells were detected in the spleen (a primary site of anti-Gal production, as demonstrated in Figure 3a) (Figure 5a). The combined FCM sorting and ELISPOT assay revealed that anti-Gal (IgM)–producing cells were greatly enriched in the sorted Gal-BSA+/IgM+ population, but were undetectable in the sorted Gal-BSA+/IgM– population (Figure 5b), demonstrating that the Gal-BSA–binding spleen cells included all anti-Gal–producing cells.

To address the question of how the specific tolerance of Gal-reactive B cells was maintained in mixed chimeras, we looked for the presence of B cells bearing receptors for Gal in long-term mixed chimeras. Spleen cells from mixed chimeras were analyzed for Gal-BSA–binding B cells 22 weeks after BMT and 8 days after immunization by intraperitoneal injection of 1 × 10⁹ rabbit erythrocytes. Gal-BSA–binding spleen cells were detected in control conditioned GalT–/– mice that did not receive BMT; they were undetectable in GalT+/– mice (Figure 6b). In mixed chimeras, the frequency of Gal-BSA–binding splenic B cells among host-derived cells (GalT–/–) was markedly lower than in conditioned GalT+/– mice, and resembled that of GalT+/+ mice (Figure 6c). Thus, B cells bearing receptors for Gal were absent from the major site of anti-Gal production in mixed chimeras.

Permanent acceptance of hearts from GalT+/+ mice in mixed chimeras. To evaluate T-cell and antibody tolerance in mixed GalT+/+ (H-2bxd)→GalT–/– (H-2d) chimeras, we transplanted hearts from donor-type GalT+/+ (H-2bxd) mice to chimeras 19–20 weeks after BMT. To enhance anti-Gal production, recipient animals were immunized by intraperitoneal injection of 1 × 10⁹ rabbit erythrocytes 8 or 9 days before heart transplantation. All untreated and conditioned control H-2d GalT–/– mice (n = 6 and n = 4, respectively) rejected H-2bxd GalT–/– hearts within 1–7 days (Figure 7). Upon histological and immunohisto-

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Figure 6
Absence of B cells bearing receptors for Gal in the spleens of mixed chimeras (22 weeks after BMT). Spleen cells were prepared from conditioned GalT–/– mice (H-2b), normal GalT+/+ mice (H-2d), and mixed chimeras (H-2bxd→H-2d) 8 days after immunization with rabbit erythrocytes. The cells were stained with FITC-conjugated Gal-BSA or control FITC-conjugated BSA together with PE-conjugated anti-CD19 mAb and biotinylated anti-donor mouse H-2Kb 5F1 mAb + CyChrome-streptavidin. (a) In mixed chimeras, the host-derived H-2Kb–negative cells were selected by gating and analyzed for the frequency of Gal-BSA–binding B cells. The percentage of host-derived cells in the spleens of mixed chimeras is indicated as average value ± SEM. (b) Representative contour plots obtained by FCM analysis show an absence of Gal-BSA–binding B cells in the spleens of mixed chimeras. To ensure statistical significance, data on 100,000 host-derived cells were collected for each sample. (c) The frequency of Gal-BSA–binding B cells was calculated by subtracting the percentage of CD19+ cells staining with control FITC-conjugated BSA from the percentage of CD19+ cells staining with FITC-conjugated Gal-BSA. Percentages of total host-derived spleen cells are shown. Average values ± SEM for the individual groups are shown (*P < 0.01; NS = no statistical difference). Number of animals in each group: conditioned GalT–/– mice that did not receive BMT, n = 4; mixed chimeras, n = 4; normal GalT+/+ mice, n = 4.

shown). Thus, the absence of a response to Gal, and the preserved normal response to other rabbit erythrocyte antigens in mixed GalT+/+→GalT–/– chimeras, confirms the presence of specific tolerance of Gal-reactive B cells.

Tolerance was apparent as early as 2 weeks after BMT, as indicated by the experiment presented in Figure 4, in which conditioned GalT–/– mice received 20 × 10⁶ GalT+/+ BMCs that were either untreated or irradiated with 30 Gy. The use of irradiated GalT+/+ BMC provided an appropriate control in which mixed chimerism could not be induced, but similar antigenic exposure occurred.

As expected, mixed chimerism was observed in all recipients of nonirradiated BMCs, but not in the recipients of irradiated BMCs. Despite the presence of host B cells in mixed chimeras at 2 weeks (mean: 42.7 ± 4.7% of splenic CD19+ cells), cells producing anti-Gal (IgM or IgG) were undetectable in the spleens of mixed chimeras by ELISPOT assay. In contrast, most conditioned GalT–/– recipients of irradiated GalT+/+ BMCs had measurable anti-Gal–producing spleen cells. The frequency of anti-Gal–producing cells in the control conditioned GalT–/– mice receiving irradiated GalT+/+ BMCs was similar to that of untreated normal GalT–/– mice (data not shown). Thus, mixed hematopoietic chimeraism is necessary for the induction of B-cell tolerance, as opposed to antigen exposure alone. Anti-Gal–producing cells were tolerant in mixed chimeras by 2 weeks after BMT.
Permanent acceptance of GalT+/+ donor-type hearts in GalT+/−→GalT−/− mixed chimeras. The hearts from donor-type GalT+/+ (H-2b) mice were heterotopically transplanted into mixed chimeras (n = 6; including 2 chimera transplanted with 20 × 10⁶, 7.5 × 10⁶, or 1 × 10⁶ GalT+/+ BMCs) 19–20 weeks after BMT, as well as into conditioned control GalT+/− (H-2b) (n = 6) and GalT−/− mice (H-2b) (n = 5) and untreated control GalT−/− mice (H-2b) (n = 4). As an H-2–identical control, hearts from GalT+/− (H-2b) mice were transplanted into GalT−/− (H-2b) mice (n = 3). To enhance anti-Gal NAb production, all recipient animals were immunized by intraperitoneal injection of 1 × 10⁹ rabbit erythrocytes 8 or 9 days before heart transplantation. Survival curves of the grafted hearts are shown. P < 0.005 normal GalT+/+, conditioned GalT+/−, or conditioned GalT+/− vs. mixed chimeras. P < 0.005 normal GalT+/− or conditioned GalT+/− vs. conditioned GalT−/−.

Discussion

We demonstrate here the simultaneous induction of T-cell tolerance and specific tolerance of Gal-reactive B cells in GalT+/→GalT−/− mixed chimeras prepared with a relatively nontoxic, nonmyeloablative regimen. These results suggest that the successful induction of mixed hematopoietic chimerism with nonmyeloablative conditioning in the pig-to-human (GalT+/→GalT−/−) xenogeneic combination could lead to tolerance among Gal-reactive B cells, as well as T cells recognizing histocompatibility antigens. Although the pathophysiological consequence of losing anti-Gal antibodies needs to be determined (30), it seems probable that induction of specific tolerance will be associated with less risk of infection than the high level of chronic immunosuppression that would be required to prevent xenograft rejection.

For success to be achieved with this approach, the possible role that preexisting anti-Gal NAb’s may play in resisting GalT+/− xenogeneic marrow engraftment must first be addressed. In this study, GalT−/− recipient mice with relatively high levels of preexisting anti-Gal NAb’s before BMT developed reduced GalT+/+ hematopoietic chimerism compared with GalT−/− recipients with lower levels of preexisting anti-Gal NAb’s (data not shown). However, once GalT+/−→GalT−/− mixed chimerism was achieved in GalT−/− mice, it was stable, and, even if present at low levels (7–10% donor WBCs), it was associated with both B-cell and T-cell tolerance. Our findings suggest that anti-Gal NAb’s may have inhibitory effects on the engraftment of GalT+/+ BMCs, but that this degree of resistance can be overcome when sufficient GalT+/+ BMCs are administered. This speculation is also supported by our previous findings that the ability of murine sera to inhibit engraftment of rat BMCs was correlated with their cytotoxic anti-rat NAb content, and that their inhibitory effect could be overcome by administration of large numbers of rat BMCs (28, 31, 32). Consistent with these results, we have confirmed the feasibility of inducing mixed chimerism in GalT+/+ mice with high levels of anti-Gal due to immunization with rabbit erythrocytes before conditioning. When high-dose BMT with GalT+/+ BMCs was given to such mice, chimerism and tolerance were achieved (H. Ohdan, K. Swenson, and M. Sykes, manuscript in preparation). If this approach were applied to the pig-to-human combination, previously reported strategies for reducing initial anti-Gal NAb and/or complement levels should be used to facilitate initial BMC engraftment. Despite such efforts, however, induction of persistent mixed chimerism has not yet been achieved in a pig-to-primate combination. In addition to the vigorous immune response to xenogenic, nonimmune physiological factors (e.g., the failure of crucial growth factors, adhesion molecular interactions, and cytokines to function across species barriers) also limit the achievement of chimerism between discordant species (33). Some of these problems might be alleviated by the use of donor-specific growth factors and/or cytokines at the time of BMT (34), and others may require genetic engineering of donor pigs.

Our previous results involving induction of mixed chimerism in lethally irradiated mice demonstrated that newly developing anti-Gal–producing B cells could be
tolerized by the presence of GalT+/+ hematopoietic cells (19). In contrast to lethal irradiation, the conditioning regimen used here would not be expected to eliminate preexisting anti-Gal–producing B cells (35, 36). Consistent with this possibility, we observed increasing anti-Gal NAb levels in sera of GalT+/+ mice receiving non-myeloablative conditioning without BMT. Thus, anti-Gal NAb–producing cells were present at high levels after conditioning. In contrast, anti-Gal–producing cells were already undetectable by ELISPOT assay in mixed chimeras as early as 2 weeks after BMT, suggesting that mixed chimerism induced tolerance among anti-Gal–producing cells that preexisted at the time of BMT. The cells that produce anti-Gal in GalT+/+ mice have not been defined. However, if it is assumed that their half-life is similar to that of other murine mature B cells (6 weeks) (37) or plasma cells (as long-lived as memory B cells) (36), then it is likely that preexisting anti-Gal NAb–producing mature B or plasma cells are tolerated by the induction of mixed chimerism. Although mature B cells are generally thought to be less sensitive than newly developing cells to tolerance induction by self-antigens or foreign antigens (38), chronic exposure to antigenic determinants present on cell surfaces has been reported to eliminate mature B cells in immunoglobulin-transgenic mice (39, 40). Based on these results, it is likely that preexisting anti-Gal–producing cells are tolerated through antigen-receptor cross-linking in mixed chimeras. However, an alternative explanation for the rapid development of tolerance in mixed chimeras is that anti-Gal–producing B cells or plasma cells may have rapid turnover. Studies are in progress to distinguish these possibilities.

In mixed chimeras, Gal epitopes may be recognized as self-antigens during B-cell maturation. The known mechanisms mediating tolerance of self-reactive B cells include clonal deletion (i.e., the physical elimination of autoreactive B-cell clones) (38, 40–42), anergy (i.e., the functional inactivation of autoreactive B cells) (43, 44), and receptor editing (i.e., the modification of B-cell receptors of autoreactive cells) (45–47). To investigate the possibility that these mechanisms are also involved in maintaining Gal-reactive B-cell tolerance in mixed chimeras, we assayed the presence of B cells with receptors (sIgM) recognizing Gal epitopes. Using FITC-conjugated Gal-BSA, we could directly identify B cells bearing anti-Gal receptors in the spleens of normal GalT+/+ mice (Figure 5a). Because anti-Gal–producing cells express sIgM (H. Ohdan and M. Sykes, unpublished data), the specificity of the Gal-BSA ligand for the corresponding receptors on B cells was demonstrated by showing enrichment of anti-Gal–producing cells among Gal-binding B cells and a complete absence of anti-Gal–producing cells among non-Gal-binding B cells, using combined FCM sorting and ELISPOT assay (Figure 5b). In mixed chimeras, B cells with receptors recognizing Gal were completely absent in the spleens (Figure 6), suggesting the possibility of clonal deletion and/or receptor editing of Gal-reactive B cells as a mechanism of tolerance.

In the present study, indefinite acceptance of vascularized GalT+/+ heart grafts was demonstrated in GalT+/+→GalT+/+ mixed chimeras, whereas rapid vascular rejection was observed in untreated and conditioned control GalT+/+ mice. Although the kinetics of GalT+/+ heart rejection in GalT+/+ mice were delayed...
compared with the typical HAR that usually occurs within minutes to hours in pig-to-primate combinations (12, 48), the histological and immunofluorescence data are consistent with a role for anti-Gal in rejection. The role of antibodies in this process is further substantiated by the more delayed (cell-mediated) rejection observed in GalT+/+ to GalT−/- mice with similar histoincompatibilities. The slower time course of GalT+/+ allogeneic mouse heart rejection observed in GalT−/- mice, compared with pig-to-primate transplantation, may be explained, first, by a lower inherent ability of mice to fix complement by the classical pathway and, second, by the intraspecies compatibility of complement regulatory proteins in our model. Using the GalT+/+ to GalT+/− mouse heart transplant model, other reports also demonstrate the absence of HAR (49) or the presence of DXR-like rejection (with graft survival of 8–12 days) (50). The more rapid rejection of allografts in control mice in our studies may be due to differences in the anatomical site of heart grafting, to the presence of MHC alloantigens in our donors, and, most probably, to the more advanced age of the mice in our studies, which was associated with high levels of anti-Gal antibodies at the time of heart grafting. Despite this rapid rejection in control mice, GalT+/−/− heart allografts in mixed chimeras were free from all types of rejection, indicating the presence of tolerance at the level of both T and B cells. This is, to our knowledge, the first direct demonstration that the induction of mixed chimerism can simultaneously prevent both T cell- and antibody-mediated rejection of vascularized solid-organ grafts. Taken together with our previous results demonstrating the induction of T-cell tolerance across rat-to-mouse and pig-to-mouse species barriers induced by a regimen similar to that used in the present studies (51, 52), it could be expected that the successful induction of mixed hematopoietic chimerism in the pig-to-human xenogeneic combination would similarly result in T- and B-cell tolerance and acceptance of vascularized organ xenografts.

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