Bioengineering thymus organoids to restore thymic function and induce donor-specific immune tolerance to allografts

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Abstract

One of the major obstacles in organ transplantation is to establish immune tolerance of allografts. While immunosuppressive drugs can prevent graft rejection to a certain degree, their efficacies are limited, transient, and associated with severe side effects. Induction of thymic central tolerance to allografts remains challenging, largely due to the difficulty of maintaining donor thymic epithelial cells (TECs) in vitro to allow successful bioengineering. Here, we show that 3-D scaffolds generated from decellularized mouse thymus can support TEC survival in culture, and maintain their unique molecular properties. When transplanted into athymic nude mice, the bioengineered thymus organoids effectively promoted homing of lymphocyte progenitors and supported thymopoiesis. Nude mice transplanted with thymus organoids promptly rejected skin allografts, and were able to mount antigen-specific humoral responses against ovalbumin upon immunization. Notably, tolerance to skin allografts was achieved by transplanting thymus organoids constructed with either TECs co-expressing both syngeneic and allogenic MHCs, or mixtures of donor and recipient TECs. Our results demonstrate the technical feasibility of restoring thymic function with bioengineered thymus organoids and highlight the clinical implications of this thymus reconstruction technique in organ transplantation and regenerative medicine.
Introduction

The primary function of the thymus is to continuously generate a diverse population of T-cells that can elicit adaptive immune responses against invading pathogens, while promoting self-tolerance [1]. The thymus is a rather vulnerable organ as many factors, including environmental insults, aging, genetic composition, virus infection, irradiation, and anti-cancer drug treatments, can all irreversibly compromise its function[2, 3]. Impaired immune surveillance consequent to thymic dysfunction leads to diseases ranging from autoimmunity to immunodeficiency and malignancy[4].

The thymus is organized into two morphologically and functionally distinct compartments: the cortex and the medulla, which house two distinct populations of thymic epithelial cells (TECs): the cortical TECs (cTECs) and the medullary TECs (mTECs) [5-8]. Other thymic stromal cells (TSCs) include thymic fibroblasts, endothelial cells, as well as antigen presenting cells like macrophages and dendritic cells. Together, this network of thymic cells provides both homing signals for the immigration of lymphocyte progenitors originated from the bone marrow (BM), and trophic factors necessary for the differentiation and maturation of thymocytes [9].

While numerous efforts have been made to correct thymic defects, manipulating the thymus, either in vitro or in vivo, proves to be challenging. This is mainly attributed to the unique architecture of the thymic stroma that is essential for the maturation, survival and function of TECs. Unlike epithelial cells of other visceral organs, which form a two-dimensional (2-D) sheet-like structure on the basement membrane to create borders within and between organs[10], TECs form a sponge-like three dimensional (3-D) network that is essential for their
TECs cultured on irradiated 3T3 feeders (a 2-D environment) are unable to support T-cell differentiation from lymphocyte progenitors, but start to express markers of terminally differentiated epithelial cells[12]. Recently, TEC stem cells derived from early embryos were shown to differentiate into skin cells when cultured in 2-D environment[13]. Indeed, the expression of key genes for the specification and proliferation of TECs (e.g. FoxN1, DLL-4, CLL-22 and Tbata) are shown to be dependent on the 3-D organization of the thymic stroma, further indicating that the unique microenvironment of the thymus is essential to maintain the unique property of TECs to support T lymphopoiesis[14].

Over the years, substantial progress has been made to reproduce the thymic microenvironment. Matrigel and other collagen-based synthetic matrices were shown to be able to support limited differentiation of lymphocyte progenitors into T-cells [15, 16]. TECs cultured in artificial 3-D matrix are viable and can partially support thymocyte development. Recently, Kyewski and colleagues developed a co-culture system, in which mTECs were layered on top of a 3-D artificial matrix embedded with human skin-derived dermal fibroblasts. Under such conditions, mTECs can retain some of their key features (e.g. expression of FoxN1, Aire and tissue-specific antigens) [17]. In a similar approach, Chung et al. mixed TECs and thymic mesenchyme, both isolated from postnatal human thymi, with CD34+ cells from cord blood to form implantable thymic units [18]. The thymic microenvironments of these thymic re-aggregates can support thymopoiesis in vitro and are able to generate a complex T-cell repertoire when transplanted in NOD.scid gamma (NSG) humanized mice in vivo. However, to date, none of these approaches has been able to fully recapitulate the function of a thymus.

Recently, significant advances have been made in “cell-scaffold” technology[19]. This groundbreaking technology uses a detergent-perfusion based approach that allows the clearance
of the cellular constituent of almost any organ of any scale, while retaining its original 3-D architecture and extracellular matrix (ECM) components [20, 21]. Repopulating the decellularized natural scaffolds with tissue-residing mature cells or progenitor/stem cells can promote its recellularization, and partially recover organ function [22]. To date, these “cell-scaffolds” have been primarily applied to manufacture and implant relatively simple organs, such as tissue engineered vascular grafts and skin, with some success [23-25]. Regeneration of complex organs such as liver, heart, lung, and kidney has also been attempted in animal models [21, 26-29]. While limited, encouraging functional regeneration of the engineered organs was observed. Furthermore, a successful clinical implantation of reconstructed decellularized trachea underlines the clinical potential of this technology [30].

Here, we show that thymus organoids reconstructed with the “cell-scaffold” technology can support thymopoiesis in vivo to establish both humoral and cellular adaptive immunity in athymic nude mice. Additionally, they also induce central immune tolerance to allo-skin grafts.

Results

Bioengineering thymus organoids with decellularized thymus scaffolds.

To investigate the possibility of reconstructing viable thymus organoids with TECs, we developed a thymus decellularization protocol improvised from a previous approach described for embryoid bodies [22, 31, 32]. This allowed us to remove all the cellular elements of a mouse thymus, while maintaining all the major ECM components (Fig. 1a-1d). Scanning electron microscopy (SEM) analysis of the cross-section images of the acellular thymic scaffolds revealed the preservation of ECM micro structures (e.g. grooves, ridges, and the fibrillar meshwork), indicating that the 3-D architecture of thymic stromal ECM is largely intact following the decellularizing treatment (Fig. 1e).
CD45- TSCs, including TECs, endothelial cells and thymic fibroblasts (Supplementary Fig. S1a), were harvested from 3-4 weeks old mice and were injected into the decellularized scaffolds and cultured in the top chamber of transwells. Since previous studies have shown that cross talk between TECs and the developing thymocytes are essential for their mutual survival and proliferation [2], lineage marker negative (Lin\(^-\)) progenitor cells from BM were co-introduced into the thymus scaffolds to mimic the immigration of lymphocyte progenitors (Fig. 2a, and Supplementary Fig. S1b - S1c). In striking contrast to the rapid loss of adult TSCs in hanging drop culture, or in porous disc on a gel foam routinely used for \textit{in vitro} fetal organ cultures, TSCs remained viable for more than 3 weeks in the 3D-thymic scaffolds (Fig. 2b-2c and Supplementary video SV1-2). Of note, 7 days after thymus reconstruction, some of the injected stromal cells began to assume a fibroblast-like morphology, suggesting that these cells successfully colonized the 3D ECM (Fig. 2b). Immunohistochemical analysis of reconstructed thymus organoids cultured \textit{in vitro} showed the presence of both TECs and CD45\(^+\) lymphoid cells (Fig. 2c, \textit{left} panel; Supplementary Fig. S2 and Supplementary video SV3). Thymic “nurse” cells, the subset of cTECs that envelop multiple CD4\(^+\)CD8\(^+\) double positive thymocytes within its intracellular vesicles to support their TCR selection and survival, were also present, suggesting that the reconstructed thymus organoid can at least retain some of its supportive properties of T lymphopoiesis \textit{in vitro} (Fig. 2c, yellow arrow in \textit{right} panel). Furthermore, Ki67\(^+\)Epcam\(^+\) TECs within the 3-D scaffolds were also observed, suggesting their proliferative potential (Fig. 2c, red arrow in the \textit{right} panel). These data are consistent with previous findings that there exist progenitor cells of the thymic epithelia in the adult thymus[33]. In addition, other thymic stromal cells, such as CD31\(^+\) endothelial cells and thymic fibroblasts were also present,
suggesting that the 3-D scaffold microenvironment is suitable for the survival of various TSC types (Fig. 2d).

Under 2-D culture conditions, TECs rapidly lose their signature gene expression and up-regulate the expression of markers of terminally differentiated skin epithelial cells[17]. Simultaneously, they also lose their capability to positively and negatively select developing thymocytes. In contrast, transcription of TEC-specific genes was readily detectable in the reconstructed thymus organoids cultured for 7-days in vitro (Fig. 2e). These included the widely used thymic epithelium marker EpCAM, and the transcription factor Foxn1, which regulates the expression of a number of key factors in TECs (e.g. Dll4) that are essential for thymopoiesis. Transcripts of Ccl25, a downstream target of Foxn1 that promote the homing of hematopietic progenitors in the thymus, were also present in the thymus organoids[34]. In addition, both Krt5 and Krt8, the cytokeratins predominantly present in mTECs and cTECs, respectively, were expressed, suggesting that both cTECs and mTECs were present. Interestingly, factors that can modulate TEC proliferation (e.g. Trp63 and Tbata) were also expressed in cells of the bioengineered thymus.

Another signature feature of mTECs is their capability to ectopically express tissue specific antigens (TSAs), believed to be a key part of the central mechanism to distinguish self from non-self and establish immune tolerance to peripheral tissues and organs[35, 36]. mRNA transcripts of Aire, one of the key regulators of thymic TSA expression, were present in TSCs of the thymus organoids, in conjunction with both Aire-dependent (e.g. Ins2)[32] and Aire-independent (e.g. Ica1)[37] TSAs (Fig. 2f). The persistence of TEC marker expression in the thymus organoids after extensive in vitro culture (Fig. 2f) also suggests that the thymic scaffold microenvironments might support the survival of thymic epithelial progenitor cells (TEPCs).
While embryonic TEPCs are much better defined (MTS20\textsuperscript{+}MTS24\textsuperscript{+}Plet-1\textsuperscript{+})\cite{7, 8, 38, 39}, markers of their adult counterparts remain unclear. Recently, Wong et al. identified a subset of TECs (EpCAM\textsuperscript{+} MHCII\textsuperscript{lo} UEA\textsuperscript{lo}, designated as TEC\textsuperscript{lo}) in adult mouse that display properties of TEPCs, including self-renewal and capability to differentiate into multiple TEC lineages\cite{40}. We examined the presence of TEC\textsuperscript{lo} cells in the reconstructed thymus organoids after 12 days of \textit{in vitro} culture, and found comparable frequencies of the TEC\textsuperscript{lo} population as those of the initial input (Supplementary Fig. 3). Taken together, these findings suggest that the 3-D scaffold environment of the decellularized thymus can support the long-term survival of TECs \textit{in vitro}, and enable them to retain the thymic specific patterns of molecule expression that is essential for T-cell development.

To examine whether the ECM microenvironment of the reconstructed thymus organoids can support T-cell lineage determination and differentiation \textit{in vitro}, Lin\textsuperscript{−} BM progenitors were isolated from C57BL/6 mice (of CD45.2 allelic type) and mixed with TSCs harvested from C57BL/6.CD45.1 congenic mice (designated as B6.CD45.1 hereinafter) at 1:1 ratio. The reconstructed thymus organoids were cultured \textit{in vitro} for 9 days, in the presence of recombinant interleukin 7 (IL-7), and the differentiation of CD45.2\textsuperscript{+} BM progenitors into T-cell lineages was examined with flow cytometry (FCM). CD3 was expressed in approximately 10\% of the CD45.2\textsuperscript{+} cells, which include both double positive (DP, CD4\textsuperscript{+}CD8\textsuperscript{+}) and single positive (SP, CD4\textsuperscript{+}CD8\textsuperscript{−} or CD4\textsuperscript{−}CD8\textsuperscript{+}) thymocytes (Supplementary Fig. S4). Even though further optimization of the conditions is needed, the results suggest that the reconstructed thymus organoids can support the development of T-cells \textit{in vitro}.

\textit{The bioengineered thymus can support T lymphopoiesis in vivo}
The capability of the bioengineered thymus to support effective thymocyte development and maturation in vivo was examined with transplantation experiments. Thymus organoids reconstructed with mixtures of TSCs and Lin- BM progenitors at 1:1 ratio, both harvested from B6.CD45.1 mice, were transplanted underneath the kidney capsules of B6.nude athymic recipients (designated as Tot.B6.nude for thymus organoid transplanted B6.nude mice hereinafter). Homing of hematopoietic progenitors to the thymus is an intermittent, gated process, alternating between ~1 week of receptive period and ~3 weeks of refractory period [41, 42]. The complement of BM progenitors was used to ensure the continuity of cross talk between TECs and the developing thymocytes that is essential for the survival of TECs, at the early post transplantation stage. The origins of the T-cells in the periphery were identified by FCM analysis of the CD45 congeneric markers (i.e., CD45.1 and CD45.2 for donor and recipient origin, respectively).

From 8-weeks post-op, populations of CD3+CD4+ and CD3+CD8+ T-cells can be clearly detected in circulation, and gradually increased over time (Fig. 3a and 3b). FCM analyses showed that secondary lymphoid organs (e.g., spleen and lymph nodes) were populated with TCRαβ+ T-cells. Both CD4+ and CD8+ T-cells were present, and many displayed similar ratios as immunocompetent naïve B6 mice (Fig. 3c). To ascertain the diversities of T-cell repertoires in the Tot.B6.nude mice, we first analyzed the distribution of TCR Vβ subtypes in T-cells isolated from the spleens, with a panel of antibodies specific to each Vβ subtype. Similar diversities of Vβ usage were observed between the naïve B6 mice and the Tot.B6.nude mice (Supplementary Fig. S5). To further assess the diversity of T-cells generated from the reconstructed thymus organoids, we performed next-generation-sequencing-spectratyping (NGS-S) analysis, which employs high coverage Roche/454 sequencing of TCR (β)-chain amplicons[43]. Broad spectra of
Vβ gene families were observed, suggesting that the T-cell repertoire was quite diverse (Fig. 3d and Supplementary Fig. S6). Notably, more than 99% of the T-cells were CD45.1−CD45.2+, indicating their recipient origin (Fig. 3e). Taken together, these results suggested that the bioengineered thymus organoid constructed from adult TSCs could effectively attract the homing of lymphocyte progenitors from the recipient’s BM, and supported the development of a diverse T-cell repertoire.

The majority of the CD8+ T lymphocytes in the spleens of Tot.B6.nude mice displayed a CD62L<sup>high</sup>CD69<sup>low</sup> phenotype, suggesting that they were naïve cells, whereas the percentages of CD4+ T-cells with activated phenotype was higher than naïve B6 mice (Fig. 3f). This might be the natural response of newly generated T-cells under lymphopenic conditions [44]. Nevertheless, none of the thymus-reconstructed mice displayed any pathological sign of autoimmunity, nor did we observe any lymphocytic infiltration in various solid organs with histology analysis (e.g., liver, pancreas and heart, etc. Data not shown). The percentages of regulatory T-cells (Treg) in the spleens of transplanted nude mice were similar to those of naïve B6 controls (Fig. 3g), consistent with a self-tolerant T-cell repertoire. Furthermore, the CD4+CD25+Foxp3+ Tregs were predominantly positive for Helios, a marker for naturally occurring Treg subset[45], suggesting that they most likely originated from the transplanted thymus organoids (Fig. 3h).

Immunohistochemical examination of the thymus organoid grafts 16-weeks post-transplantation showed the presence of EpCAM+ TECs, as well as CD4+ and/or CD8+ thymocytes underneath the kidney capsules (Fig. 3i). These results further suggested that the thymus organoid grafts can support the survival and function of the TECs, but also promote the homing and differentiation of lymphocyte progenitors <i>in vivo</i>. 

**Effective cellular and humoral adaptive immunity mediated by T-cells matured in bioengineered thymus organoids**

Proliferation under various stimuli has been widely used as a tool to assess the functionality of T lymphocytes. To demonstrate that T-cells derived from the reconstructed thymus organoids are functionally competent, we labeled them with carboxyfluorescein diacetate succinimidyl ester (CFSE) and stimulated them with anti-CD3 antibodies. Similar to T-cells of naïve B6 mice, a significant percentage of T-cells underwent division, as indicated by dilution of CFSE signals (Fig. 4a). To further test the function of T-cells derived from the reconstructed thymi, we performed mixed leukocyte reaction (MLR) experiments to evaluate their responses to alloantigens. Proliferation responses similar to those of wild-type B6 mouse were observed, indicating that these T-cells were capable to react to alloantigens (Fig. 4b). Taken together, these results demonstrated that T-cells matured in the transplanted thymus organoids were capable to response to TCR stimulation.

To demonstrate that T-cells derived from the bioengineered thymus can effectively mediate cellular immune response in vivo, we performed allo-skin transplantation experiments, to examine whether the recipients can reject skin allografts. Allogeneic skin grafts harvested from CBA/J mice (H-2^k^) were transplanted to the back of Tot.B6.nude mice (n=3). Syngeneic skin grafts from B6 mice (H-2^b^) were co-transplanted as controls. While the syngeneic skin grafts were well tolerated, the allogeneic skins were rejected within 2-3 weeks, with kinetics similar to naïve B6 recipient controls (n=4) (Fig. 4c). In contrast, allogeneic skin grafts on B6.nude mice that had been transplanted with decellularized empty scaffolds were viable for more than 8-weeks (Fig. 4d-e). These results demonstrated the capability of T-cells in the
Tot.B6.nude mice to efficiently mediate rejection of allogeneic skin grafts, while being unresponsive to self-tissues.

One of the essential roles of T-cells in adaptive immunity is their helper function for humoral immunity, namely to mediate immunoglobulin (Ig) class switch in antibody producing B-cells. To examine whether T-cells from the bioengineered thymus can provide helper function to humoral responses, we immunized thymus recipients with chicken ovalbumin (OVA), a T-cell-dependent antigen commonly used for studying antigen-specific immune responses in mice. As expected, high titers of anti-OVA antibodies were detectable by ELISA in sera of B6 mice 4-6 weeks post-immunization (Fig. 4f). Comparable levels of anti-OVA immunoglobulins of IgG isotype (i.e. IgG2b and IgG3) were found in serum samples harvested from the Tot.B6.nude mice, whereas seroreactivities against OVA remained at the background levels in immunized B6.nude mice transplanted with empty thymus scaffolds (Fig. 4f). These results suggest that T-cells generated from the thymus organoids can support B-lymphocyte Ig class switch to mount efficient humoral responses.

To further demonstrate that T-cells matured in the thymus organoids can elicit antigen-specific adaptive immune responses, Tot.B6.nude mice were immunized with OVA peptide (AVHAHAHAEINEAGSIINFEKL), which contains an H-2Kb restricted, cytotoxic T lymphocyte (CTL) epitope (the underlined region). 4-weeks after the initial vaccination, splenocytes were challenged with the OVA peptide and the presence of T-cells secreting proinflammatory cytokine interferon γ (IFN-γ) was evaluated with the Enzyme-Linked ImmunoSpot (ELISPot) assay. As shown in Figure 4g, significant numbers of OVA peptide responding cells were present, suggesting that Tot.B6.nude mice can effectively mobilize pro-inflammatory response upon antigen challenge.
Induction of allo-skin tolerance with bioengineered thymus organoids constructed with TECs of F1 hybrid of donor and recipient mice

Achieving donor-specific immune unresponsiveness, without the need for pharmacologic immunosuppression, remains a major goal of transplantation immunological research. To prove the principle that transplantation of bioengineered thymus organoids expressing both donor and recipient MHCs can establish central tolerance to donor antigens, we reconstructed the acellular thymus scaffolds with TSCs harvested from the F1 offspring (B6.H-2b/g7) of a cross between B6 (H-2b) and B6.H-2g7 congenic mice, and transplanted them to the B6.nude recipients (H-2b). The B6.H-2g7 mouse is a congenic line in which a 19 cM segment of Chr 17 including the major histocompatibility complex (MHC) of the B6 mouse (of the H-2b haplotype) were replaced with that of the non-obese diabetes (NOD) mouse line (of the H-2g7 haplotype). It was established through multiple rounds of backcrossing of the NODxB6 F1 mice to the parental B6 line. Once a substantial population of T-cells became detectable (12-16 weeks) in the peripheral bloods of the recipients, skin grafts harvested from both the syngeneic B6 (H-2b) and the allogeneic B6.H-2g7 congenic mice were transplanted on their backs. To demonstrate that the thymus organoid transplanted recipients retained their capabilities to reject third party alloantigens, skin grafts harvested from CBA/J (H-2k) mice were also transplanted (Fig. 5a). Successful engraftments of skin transplants from both the syngeneic B6 and the allogeneic B6.H-2g7 mice were observed, whereas the third party CBA/J skin grafts were rejected within 2-3 weeks (Fig. 5b and 5c). Immune unresponsiveness to H-2g7 alloantigens in the recipients was further demonstrated in MLR assays (Fig. 5d). Taken together, these results suggested that transplantation of bioengineered thymus organoids co-expressing both syngeneic and allogeneic MHCs can effectively establish donor-specific immune tolerance.
Induction of allo-skin tolerance with bioengineered thymus organoids constructed with mixture of TECs from both the donor and the recipient.

While reconstructing thymus organoids with TECs co-expressing both donor- and recipient-MHCs can effectively induce tolerance to donor MHC-expressing grafts, it is not clinically feasible to transfer donor MHC genes to the recipient’s TECs at such high efficiency (100% in the case of F1 TECs), using currently available gene engineering techniques. Moreover, epitopes derived from mismatched genes other than the MHCs in the allogeneic donor organ(s) can also contribute to its rejection. One possible way to overcome these obstacles is to incorporate the donor TECs in the thymus organoids, together with the recipient’s TECs. To test this hypothesis, we performed the experiments schematically illustrated in Figure 6a. TECs harvested from B6 (H-2b) and CBA/J (H-2k) were mixed at 1:1 ratio and co-injected with B6 BM progenitors to the decellularized thymus scaffolds. B6.nude mice reconstructed with the thymus organoids were challenged with allogeneic CBA/J skins, as well as skin grafts from the third party Balb/C mice (H-2d). Prolonged survival of the CBA/J skin allografts was observed (Fig. 6b). Consistently, results of MLR experiments revealed that when challenged with CBA/J APCs, the levels of T-cell proliferation were significant lower than those stimulated with Balb/C APCs. These findings further suggested that including donor TECs in the reconstruction of thymus organoids might be a clinical applicable means to induce donor-specific immune tolerance.

Discussion

We provided functional evidence that bioengineered thymus organoids made of decellularized thymic scaffolds populated with TECs and Lin- BM progenitors, can re-assert thymic T-cell generation in athymic nude mice. While attempts to decellularize the thymus glands and repopulate them with epithelial cells have been reported previously [46, 47], to our
knowledge, we show for the first time that the decellularized thymus scaffolds can support the survival of TECs \textit{in vitro}, and that the bioengineered thymus organoids can successfully support thymopoiesis and T-cell generation \textit{in vivo}.

Induction of donor-specific immune tolerance remains a major challenge in solid organ transplantation. While the rates of acute rejection have decreased significantly over the past 10 years as a result of new immunosuppressive drugs, long-term allograft survival has not correspondingly improved. Conventional pharmacological drugs (e.g., calcineurin inhibitors and corticosteroids) can cause severe side effects including increased risk of cardiovascular death, diabetes and kidney disease. Even with the development of modern immunomodulatory protocols such as depletion of mature T cells with antibodies, blockade of costimulatory molecules (e.g. CD28 and CD40L), and promotion of tolerogenic dendritic cells and regulatory T cells, issues commonly associated with immune suppression (e.g. high risks of opportunistic microbe infection and tumor development) remain unresolved. Notably, these approaches aim to eliminate/suppress mature donor-reactive T-cells in the periphery; none is designed to address the source of the alloreactive T-cells: the thymus. In addition, some of these treatments might hinder the recovery of thymus function, which is key to re-establish host defense. Our study demonstrated the technical feasibility of establishing functional donor-specific immune tolerance at the central level, by introducing allogenic donor TECs into bioengineered thymus organoids to obviate the need for immunosuppressive drugs to maintain allograft survival.

Targeting the thymus as a means to achieve immune tolerance in organ transplantation is not a novel concept. Komori and colleagues have shown recently that B6 thymus fragments transplanted into the lymph nodes of Balb/C nude mice can support thymopoiesis and induce immune tolerance of B6 skin grafts[48]. Other approaches undertaken to introduce allogeneic
antigens in the thymus include directly injecting donor antigen presenting cells (APCs) or alloantigen-expressing viral vectors, or establishing mixed chimerism through thymus irradiation in conjunction with bone marrow transplantation [49-52]. However, most of these approaches predominantly affect APCs, especially thymic dendritic cells (DCs), which are replenished by fresh immigrating cells of the recipients in less than 2 weeks. This intrinsic property of fast turnover of thymic APCs could compromise the long-term efficacy of treatment. In contrast, TECs are an integral cell population of the thymic stroma, whose homeostasis is largely maintained through self-renewal. Although their capability to directly present antigens to developing thymocytes is still under debate, recent studies have shown that thymic DCs can readily acquire antigens from mTECs and present them for negative selection. Thus, the presence of donor antigen-expressing TECs in the bioengineered thymus organoids might ensure the passage of alloantigens to thymic DCs (of the hosts) for negative selection of donor-reactive T cells, facilitating the establishment of long-term donor-specific immune tolerance.

One of the major challenges to manipulate postnatal TECs for clinical application is their dependency on a properly configured 3-D ECM microenvironment for survival and proliferation. Early studies have shown that thymus fragments cultured in 3-D scaffolds constructed with biocompatible inorganic materials can support the differentiation of hematopoietic progenitor cells into CD3+ T-cells [53]. The findings that TECs injected into the thymus scaffolds could retain their specific molecular properties for up to 8-weeks in vitro strongly suggested that the 3-D ECM environment of the decellularized thymus provided a suitable and essential niche for the long-term survival of adult TECs. Additionally, Lin- BM progenitors co-injected with the TECs can differentiate into CD4⁻CD8⁺ DP as well as CD4⁺CD8⁺, or CD4⁺CD8⁺ SP thymocytes,
suggesting the possibility that with further optimization, the thymus scaffolds could serve as a suitable in vitro culture microenvironment to study T cell development.

Despite the findings that the bioengineered thymus organoids can effectively support T lymphopoiesis and restore thymus functions for both cellular and humoral immunity, the total numbers of T-cells in the spleens of transplanted thymic nude mice were less than 10% of those of naïve B6 mice. Moreover, the T-cell repertoire in the thymus recipients was much less complex than that of naïve mice. Many factors might contribute to this inefficiency. For example, thymopoiesis is a well-controlled developmental process, which depends on cross talk between the developing thymocytes and subsets of TECs in a spatial and temporal manner. Expansion and positive selection of thymocytes occur in the cortical region, whereas final maturation and negative selection of the SP thymocytes predominantly takes place in the medulla. Our current top-down approach (breaking up and reassembling) is limited in part by the lack of mechanical control to sort the dissociated cTECs and mTECs into separated functional units. Such lack of proper organization and compartmentalization of TECs might affect the extent of interactions between the developing thymocytes and the thymic stroma, skewing the V(D)J recombination events to favor the selection of certain TCRs. Alternatively, deviant thymopoiesis might lead to inefficient output of naïve T-cells, resulting in impaired proliferation and over representation of certain early emigrants in the lymphopenic immune environment of the nude mice.

Second, the engraftment site (i.e. kidney capsules) might not be optimal for the survival and function of the thymus organoids. In our studies, intact thymi of 2-week old mice had largely degenerated 2-4 weeks after transplantation underneath the kidney capsules (data not shown). One of the major obstacles might be the inefficiency of angiogenesis. To promote graft
vascularization, Seach et al. embedded thymus fragments in housing chambers consisting of silicone tubing and implanted the device in the vicinity of epigastric blood vessels of nude mice [54]. While limited, successful T-cell development was observed, and mice were able to reject MHC miss-matched skin grafts. Recently, Chung and colleagues co-cultured human postnatal TECs and thymic mesenchymal cells that were transduced with vascular endothelial growth factor (VEGF)-expressing lentiviral vectors, and showed that the engineered human thymic aggregates can support thymopoiesis both in vitro and in vivo [18]. Refining our thymus bioengineering protocol with these angiogenesis-promoting techniques might enable us to improve survival and the efficacy of the transplanted thymus organoids to support T lymphopoiesis.

To closely mimic the clinical situation, we did not use 16-day prenatal mouse embryos to obtain TSCs, but harvested them from 2-4 week old young mice. While it was previously demonstrated that a single stem cell from the embryonal region of the thymus can recapitulate the developmental process of the whole thymus organ, and TEC stem cell lines with the potency to differentiate into various TEC subsets have been isolated from 16-day prenatal embryos, the presence and prevalence of multipotent TEC progenitors in a postnatal thymus remains a matter of debate. Consistently, the levels of T-cell development in nude mice transplanted with bioengineered thymus organoids were similar to those transplanted with intact, thymocyte-depleted thymi of 2-4 week old mice. Thus, the low T-cell numbers in the thymus-transferred nude mice might reflect the intrinsic limitation of the postnatal thymus to fully reconstruct the T-cell repertoire.

Such limitation of postnatal donor TECs (e.g. numbers, regeneration capabilities, properties) could potentially compromise the efficacy of establishing donor-specific tolerance,
when the bioengineered thymus is co-transplanted with the solid organ into a transiently immunosuppressed recipient. Especially if the donor is of old age, there might not be enough donor TECs available to completely reconstitute a functional hybrid thymus. While the factors that can support the survival and proliferation of TEC progenitors remain unknown, a number of cytokines and growth factors [e.g. keratinocyte growth factor (KGF), GH, and IL-22] have been implicated in maintaining the integrity of thymic stroma, especially the unique characteristics of TECs [55-58]. Recent studies showed that forced expression of the TEC-specific transcription factor FoxN1 could rejuvenate the involuted thymus and increase naïve T cell output in aged mice, suggesting the feasibility of prolonging the function of adult TECs by genetically modulating the expression of key lineage-determining factors in TEC biology [59, 60]. Alternatively, donor TECs can be generated from embryonic stem cells (ESCs)[61, 62], induced pluripotent stem cells (iPSCs) [63], or genetically reprogrammed embryonic fibroblasts [64].

Aided by advances in targeted gene delivery technologies [65, 66], we might be able to fine-tune the antigen presentation properties of TECs and bioengineer individualized thymus organoids to achieve long-term donor-specific immune unresponsiveness in clinical transplantation, or to regain self-tolerance to specific tissues in treating autoimmune disorders with known major autoantigen(s). At present, thymus scaffolds can be prepared from thymus glands harvested from cadavers or patients undergoing cardiothoracic surgery in which the thymic samples are removed as waste tissue. As shown in our study, scaffolds prepared from mouse thymi can be stored at 4°C for up to one month before use; it is likely that human thymus scaffolds can be preserved in a similar fashion. Since there is no cellular component in the decellularized thymus scaffold, we believe that allogeneic (or even xenogeneic) rejection won’t be a concern. The thymus organoid can be reconstructed with TECs harvested from the donor
and a properly preserved decellularized thymus scaffold from a third party. Indeed, even with the current technology, our preliminary experiments suggest that this thymus bioengineering approach is applicable to nonhuman primates (Fan et al., unpublished observations).
Materials and methods

Mice

All animal protocols were reviewed and approved by the Institutional Animal Care and Use Committee of the Allegheny Health Network (AHN)/Allegheny Singer Research Institute (ASRI). All animals were housed in specific pathogen free environment. The following strains of mice were purchased from the Jackson Laboratory (Bar Harbor, ME): CBA/J, B6 (C57BL/6J), B6.H-2^g7 and C57BL/6J.CD45.1 (B6.CD45.1) congenic strain. Athymic B6.nude mice were obtained either from the Jackson Laboratory or from Taconic (Germantown, NY); similar results were obtained from both nude mouse strains.

Skin Transplantation

Tail-skin graft was excised from euthanized donor mouse (about 1 cm long and 0.5 cm wide), and was placed on a bed prepared by removing an area on the back dermis of the either B6.nude or Tot.B6.nude recipient. The graft was sutured, covered with gauze, and was wrapped with sterile bandage. Skin graft survival was monitored daily and rejection was defined as graft necrosis of more than 80%.

Decellularization of mouse thymus

Decellularization was carried out by chemical detergent washing similar to previous study [22, 31]. Briefly, thymi of 3-4 week old mice were stored in -80°C until decellularization was initiated. Thymi were thawed in a 37°C water bath. This free-thaw process was repeated three times. Next, ionic detergent, 0.1% SDS (Invitrogen) in deionized water was added to the thymi and placed on a 3-D rotator (Lab Line, Thermo Scientific) for continuous rotation until the tissues became translucent and white in color (24 hours). Thymi were subsequently washed in PBS for three times, each for 15 minutes, followed by 30 minutes in 1% Triton X-100 (Sigma
Aldrich). This was followed by three more PBS washes. A final wash step of PBS with Pen/Strep (100U/ml) was added and the scaffolds were rotated for additional 48 hours. The decellularized thymus scaffolds were stored in PBS at 4°C for up to a month, and were switched to RPMI-10 culture medium (RPMI-1640 with 10% fetal bovine serum (FBS), 100 U/mL Penicillin, 100 ug/mL Streptomycin, 2mM L-glutamine, 10mM HEPES) 24 hours before use.

**Thymus Organoid Reconstruction**

To reconstruct the thymus organoids, thymic tissue was harvested from 2-3 week old B6 mice, unless specified otherwise, and separated into single cells with collagenase digestion as previously described[32, 67]. In brief, thymic tissue (n=3-4) was pooled and needle dissected into small pieces of approximately 1 mm³, and digested with the Liberase TM solution [0.025mg/ml Liberase TM (Roche Applied Science, Indianapolis), 0.2mg/ml DNase I (Roche Applied Science), and 10mM HEPES in RPMI-1640 (Life Technologies, Carlsbad, CA)] at 37°C for a total of 18 minutes (3 rounds of 6 minutes each). All fractions were pooled, incubated with magnetic bead-conjugated anti-CD45 antibodies, and subjected to negative selection of CD45⁻ thymic stromal cells (e.g. TECs and thymic fibroblasts) with MACS separation technology (Miltenyi Biotec, Auburn, CA). Approximately 0.25-0.5 million stromal cells were routinely obtained per thymus.

Lin⁻ progenitors were enriched from bone marrow cells of 2-3 week old B6.CD45.1 congenic mice (unless specified otherwise) with mouse Lineage cell depletion kit (Miltenyi Biotec, Auburn, CA), following the manufacturer’s instructions. Lin⁻ progenitors and TSCs were mixed at 1:1 ratio and resuspended in RPMI-10 solution at a concentration of 5x10⁷/ml. 10 µL of cell mix (~0.5x10⁶ cells) was injected into the decellularized thymus scaffolds using syringes with 32G needles or pulled glass needles under the dissection scope. The reconstructed thymus
organoids were cultured in the top chamber of a 3.5cm transwell in RPMI-10 medium before being transplanted underneath the kidney capsules of 10-16 week old B6.nude mice.

To evaluate T-cell development in the reconstructed thymus organoids, BM progenitors harvested from the B6 mice were mixed with TSCs isolated from C57BL/6.CD45.1 thymi at 1:1 ratio. The reconstructed thymus organoids were cultured in the upper chambers of 24-well transwell plantes (one thymus per well; Corning, Tewksbury, MA) in complete RPMI-10 medium supplemented with 2 ng/ml Interleukin-7 (IL-7, Miltenyi Biotec) for 9-days. At the end of the culture, the thymus organoids were digested with Dispase I (0.6 U/ml, Roche Diagnostic, Indianapolis, IN) for 15 minutes at 37°C. The isolated cells were stained with anti-CD45.2, anti-CD3, anti-CD4 and anti-CD8 antibodies for FCM analysis.

**Stimulation of peripheral T cells**

T-cells isolated from the spleens and/or lymph nodes of Tot.B6.nude mice were labeled with CFSE, and were subjected to stimulation with either anti-CD3 antibodies or allogeneic cells isolated from the spleens of CBA/J or Balb/C mice. Proliferation of T-cells under different stimulatory conditions was evaluated by the dilution of the intensities of CFSE signals with FCM analyses, as previously described [68, 69]. Briefly, stock CFSE (5mM) was diluted at 1:1 ratio in HBSS. T-cells were resuspended at a concentration of 10x10^6 cells/ml, mixed with equal volumes of diluted CFSE, and incubated at 37°C for 10 min. For assays with anti-CD3 antibody mediated stimulation, 5x10^5 of CFSE-labeled cells in 100ul were added to wells of 96-well plates pre-coated with anti-CD3 (clone 500A2) antibodies, and incubated for 72 hours. Cells were labeled with anti-CD4 and -CD8 antibodies, followed by staining with LIVE/DEAD Fixable Violet dead cell staining reagent for FCM analyses. For assays with allogeneic antigen presenting cells as stimulants, 2.5x10^5 of CFSE-labeled cells in 100ul were mixed with 2.5x10^5
T-cell depleted, mitomycin C (Sigma) treated splenocytes harvested from CBA/J mice, added to wells of 96-well plates, and cultured for 4-7 days. Cells were labeled as described above for FCM analyses. In addition, cells were stained with anti-H2-K\(^k\) and anti-H2-K\(^b\) antibodies to label the stimulator (CBA/J) and responder (Tot.B6.nude) populations, respectively. Unless specified otherwise, all the experiments were run in triplicate and repeated at least three times.

**Histology and Immunohistochemistry**

Reconstructed thymus organoids and kidneys were fixed in 4% paraformaldehyde for 3 h at 4°C, and placed in 30% sucrose overnight. Cryosections of 7 μm thick were cut and stained with primary antibodies. Antibodies used in the study: Epcam (g8.8), B220, CD45, CD4 and CD8 (BD Biosciences, Franklin Lakes, NJ), rabbit anti-laminin, rabbit anti-collagen I, rabbit anti-collagen IV, rabbit anti-fibronectin (Abcam, 1:200).

**Detection of humoral responses against ovalbumin and peptides**

Tot.B6.nude mice (n=5) were immunized with subcutaneous injection of 50μg of the ovalbumin protein (OVA, Invivogen, San Diego, California, USA) emulsified in complete Freund’s adjuvant (CFA) at the tail bases, followed by intraperitoneal injection of 25μg of OVA in incomplete Freund’s adjuvant (IFA) to boost the immune responses one week later. Wild type B6 mice (n=5), as well as B6.nude mice transplanted with empty thymic scaffolds (n=3) were also immunized similarly, as positive and negative controls, respectively.

Sera were harvested 6 weeks post-immunization, and the presence of anti-OVA immunoglobulin was examined with the mouse monoclonal antibody isotyping reagents kit (Sigma-Aldrich, St. Louis, USA), following manufacturer suggested protocol. Briefly, wells of 96-well ELISA plate were coated with 1μg of OVA in 100μL of PBS for 2 hours at 37°C, and incubated with 100μL of diluted sera samples (1:50 dilution) overnight at 4°C. Monoclonal
antibody specific to different immunoglobulin isotypes (1:1000 dilution) was added to each of the wells, and incubated for 2 hours at room temperature. After wash, 100ul of the peroxidase labeled goat anti-mouse IgG antibody (1:5000) was added to each well and incubated at room temperature for 30 minutes. After wash, TMB substrate (BioLegned, San Diego, CA) was added to each well and the absorbance was measured at 450 nm (minus 570 nm for wavelength correction) with microplate reader (Molecular Devices). The experiments were run in triplicate and repeated three times.

**ELISPOT analysis**

ELISPOT was performed using the BD mouse IFN-γ set (BD Biosciences), as previously described[70]. Briefly, splenocytes were isolated from OVA peptide immunized Tot.B6.nude (n=4), or B6 controls (n=4), suspended at 4.5x10^6/ml concentration in RMPI-10, and cultured for 24 hours at 37°C, in the presence of 50µg/ml OVA peptides (AVHAAHAEIENAGSIINFEKL). Unattached cells were harvested, washed, counted and resuspended at 3x10^6 cells/ml in RPMI-10, supplemented with 50µg/ml OVA peptides. 1x10^6 cells were seeded to one well of 96-well ELISPOT plate (MSIPS4W 10, EMD Millipore, Thermo Fisher Scientific, Waltham, MA), pre-coated with 0.5µg of anti-IFNγ capture antibody (BD Biosciences), and cultured overnight at 37°C. At least 3x10^6 cells (10 wells) were analyzed for each sample. The wells were subsequently stained with HRP-conjugated, detection antibodies. IFNγ-producing cells were visualized with AEC substrate set (BD Biosciences), and analyzed with CTL-ImmunoSpot S6 Micro Analyzer (Cellular Technology, Ltd., Shaker Heights, OH). The assays were repeated at least three times.

**Detection of live cells in the reconstructed thymus organoids**
Live cells in the thymus organoids cultured \textit{in vitro} were detected with LIVE/DEAD viability kit for mammalian cells (Lifetechnologies, Grand Island, NY), following the fluorescence microscopy protocol suggested by the manufacturer. The kit discriminates live from dead cells by simultaneously staining with green-fluorescent calcein-AM, an indicator of intracellular esterase activity, and red-fluorescent ethidium homodimer-1 (EthD-1), an indicator of loss of plasma membrane integrity. Briefly, reconstructed thymus organoids were washed extensively with Dulbecco’s phosphate-buffered saline (D-PBS). Staining solution was prepared by diluting stock solutions of both EthD-1 and calcein-AM in D-PBS to a final concentration of 4\textmu M and 2\textmu M, respectively. The thymus organoids were incubated in the staining solution for 30 minutes at room temperature, and examined under the inverted laser scanning fluorescence microscope (FV1000, Olympus).

In some experiments, TSCs were first labeled with carboxyfluorescein diacetate, succinimidyl ester (CFSE) before being injected into the acellular thymus scaffolds. The presence and distribution of CFSE+ cells in the reconstructed thymus organoids cultured \textit{in vitro} were followed under the fluorescence microscope for up to 4-weeks.

\textbf{RNA analysis}

Total RNA was extracted from 250,000 TSCs and reconstructed thymus organoids cultured \textit{in vitro} with RNeasy micro kit, according to the manufacturer’s protocol (Qiagen). Following DNase I treatment (Ambion), RNA samples were reverse-transcribed into cDNAs with Superscript III cDNA kit, with random hexamers as primers for the RT reaction (Invitrogen). Semi-quantitative PCR was performed as previously described\cite{71}. cDNA samples were diluted serially (1:5) and equal volumes of dilutents were used as templates for PCR.
amplification of the gene of interest. Sequences of primer pairs and conditions are listed in the table below.

<table>
<thead>
<tr>
<th>Genes</th>
<th>Primers</th>
<th>Annealing Temperature (°C)</th>
<th>Number of Cycles</th>
<th>PCR size (base pairs)</th>
</tr>
</thead>
</table>
| Ins2  | F 5'-CGC CGT GAA GTG GAG GAC-3'  
R 5'-TCTACAATGCCCAGCCTTCTG-3' | 62 | 40 | 115 |
| Ica1  | F 5'-TGAGTCTCGCAACCTCAAACAGGA-3'  
R 5'-AAACAGGGCCCTTGAACCCCTCATT-3' | 58 | 42 | 141 |
| Aire  | F 5'-AATCTCCCGCTGCAAATCCTGCTCT-3'  
R 5'-ACTGCAGGATGCCGTCAAATGAGT-3' | 60 | 45 | 199 |
| Foxn1 | F 5'-TGACGGGAGCAGCTTCTTTAC-3'  
R 5'-GACAGGTTATGCGAAGACAGA-3' | 60 | 42 | 296 |
| CCL25 | F 5'-GAGTGCCACCTAGGTCATC-3'  
R 5'-CCACGCTGCTTACCTCCTGTA-3' | 60 | 40 | 87 |
| Tbata | F 5'-TGACTGACCCACCTTATCTAC-3'  
R 5'-GGGAAACCCCTTGGATTTCT-3' | 58 | 40 | 353 |
| Trp63 | F 5'-GCCTCATTCATGCTGAGACTATTT-3'  
R 5'-CGCTCAATTCTCCTCTCTTCTTCT-3' | 60 | 42 | 352 |
| EpCAM | F 5'-AGAAATACGTCATTTGCTCAAAACT-3'  
R 5'-GTTCGATCGCCACCTTC-3' | 58 | 40 | 110 |
| Krt5  | F 5'-CAGGCGAACAGACCATAAA-3'  
R 5'-CTGCTGACGTCTCATCTT-3' | 58 | 40 | 332 |
| Krt8  | F 5'-CGCTCTGGTGCTCTATG-3'  
R 5'-CTTGGTCTGCGGCACTCTTAAAT-3' | 58 | 40 | 332 |
| 18S   | F 5'-AAACGCGTACACACATCAAAG-3'  
R 5'-CCTCCAATGGATCGGCCTT-3' | 60 | 28 | 113 |

**Flow Cytometry**

Flow cytometric analysis was performed on the BD FACSCalibur flow cytometer (BD Biosciences, San Jose, CA) and analyzed with either the CellQuest Pro software (BD Biosciences), or Flowjo. Single cell suspensions were prepared from spleen, subjected to erythrocyte depletion in red blood cell lysis buffer (Sigma-Aldrich, St. Louis, MO), blocked with anti-CD16/32 antibody and then stained with the other antibodies. The following antibodies were purchased from BD Biosciences: anti-CD16/32 (2.4G2), anti-CD4 (H129.9), anti-CD45 (30-
F11), anti-CD3 (145-2C11), anti-CD45.1 (A20), anti-CD45.2 (104), anti-Epcam (g8.8), anti-CD62L (MEL-14), anti-CD69 (H1.2F3), and anti-CD44 (IM7). Anti-CD25 (7D4) antibody was purchased from Miltenyi Biotec (Auburn, CA). Staining buffer: phosphate buffered saline (PBS, calcium and magnesium free, Invitrogen) supplemented with 1% bovine serum albumin (BSA, Sigma-Aldrich) and 0.1% sodium azide (Sigma-Aldrich). Intracellular staining of the Foxp3 and Helios protein was performed with commercial kit purchased from eBiosciences (San Diego, CA), following manufacturer’s suggested protocol.

**Scanning Electron Microscopy (SEM)**

Native and decellularized thymi were fixed in 2.5% glutaraldehyde in 0.1 M PBS (pH 7.4) for 60 min. The samples were washed thoroughly in 3 changes 0.1 M PBS for 15 min each. Next, the samples were fixed in 1% OsO4 in 0.1 M PBS for 60 min. This was followed by another 3 changes of PBS washing steps for 15 min each. The samples were then dehydrated in gradient series of alcohol for 15 min each. Additionally, samples were critical point dried and coated with Au/Pd using a Cressington Coater 108A sputter coater. Electron microscope images were taken using a Jeol JSM-6335F field emission SEM.

**Statistical analysis.**

All values are expressed as the mean ± SEM unless otherwise specified. In mouse studies, statistical significance was determined using nonparametric Mann-Whitney test. All statistical analyses were carried out with the GraphPad Prism 4.0 Software. In all experiments, differences were considered significant when \( p < 0.05 \).
Acknowledgements

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Supplementary Materials

Fig. S1. Reconstruction of thymus organoids with decellularized 3-D thymus scaffolds.

Fig. S2. Fluorescence microscopy images of reconstructed thymus organoids.

Fig. S3. Frequencies of TEPCs in reconstructed thymus organoids cultured in vitro.

Fig. S4. Reconstructed thymus organoids support T-cell development in vitro.

Fig. S5. FCM analyses of the diverse distribution of T-cell receptor (TCR) Vβ genes.

Fig. S6. Diversity of TCR Vβ genes in Tot.B6.nude mice determined by next-generation-sequencing-spectratyping (NGS-S).

Video S1. 3-D animation of the reconstructed thymus organoid.

Video S2. Representative 3-D composition of images of Epcam+ TECs in thymus organoids.

Video S3. Representative composite images of CD4+ and/or CD8+ T-cells in reconstructed thymus organoid.
References


Figure Legends

**Figure 1. Preservation of 3-D ECM architecture in decellularized mouse thymus scaffolds.**

*a.* Mouse thymus was decellularized with detergent and preserved in PBS. D, decellularized thymus (arrow); N, naïve thymus. 

*b.* H&E images of 7μm paraffin sections of decellularized thymus scaffolds (D, *left panel*) and naïve thymus (N, *right panel*). No remnant cell is detected after the completion of decellularization. White scale bar, 50um.

*c.* *Left panel,* Picogreen analysis of DNA contents in decellularized thymus scaffolds (D, n=3), showing the removal of up to 99% of DNA materials, in comparison to the naïve thymi (N, n=3). *Right panel,* glycosaminoglycan (GAG) content in the decellularized thymus scaffolds. About 7.5% of sulfated GAG contents are preserved in the scaffolds.

*d.* Immunohistochemical analysis of the preservation of the extracellular matrix (ECM) components (red) in the thymus scaffolds. Cryosections of naïve thymus (N, *upper panels*) and decellularized thymus scaffolds (D. *lower panels*) were stained with antibodies against ECM proteins (*red.* Collagen I, Collagen IV, Fibronectin, and Laminin), and counter-stained with DAPI for nucleus (blue).

*e.* Ultrastructure characterization of native and decellularized thymus. *Left panel,* representative SEM image of native thymus (N), showing distinct individual cells. *Right panel,* SEM image of decellularized thymus (D) shows the preservation of 3D meshwork within the parenchymal space composed of variety of fibers, including large bundle of Type I collagen (yellow arrowhead) associated with a variety of smaller fibers (white arrowhead). No cell is detected throughout all tissue layers, indicating complete removal of the cellular components.

**Figure 2. Reconstruction of thymus organoids from decellularized thymus scaffolds.**

*a.* Light microscopic images of a decellularized thymus scaffold (*left panel*) and a reconstructed
thymus organoid (with CD45− thymic stromal cells and bone marrow cells of Lin− population at 1:1 ratio) cultured overnight *in vitro* (*right* panel). *b.* Fluorescent microscopic images of TSCs cultured either as “hanging drop” overnight (*left* panel) or in the 3-D scaffold for 7 days (*right* panel). Live cells were discriminated from the dead cells by their intracellular esterase activities to generate green fluorescent calcein-AM (green) and their capabilities to exclude the red-fluorescent ethidium homodimer-1 (EthD-1, red) from entering the nucleus. *c-d.* Representative immunohistochemical images of reconstructed thymus organoids cultured *in vitro* for 7 (*d*) or 21 (*c*) days. *c.* Cryosections were stained with antibodies against Epcam (green), counter-stained with either anti-CD45 (red, *left* panel), or anti- Ki67 (red, *right* panel) antibodies. In the left panel, white arrows show the presence of close interactions between the CD45+ thymocytes and the Epcam+ TECs. In the right panel, the yellow arrows show the presence of multi-cellular thymus nurse cell complex, whereas the red arrow shows a Ki67+Epcam+ TEC. *d.* Cryosections were stained with endothelial cell-specific anti-CD31 antibodies (red, left panel) and fibroblast-specific antibodies (red, Fibro, right panel). Both sections are counter stained with the anti-Epcam antibodies (green) and the Hoechst 33342 dye (blue) for TECs and nuclei, respectively. *e.* Semiquantitative RT-PCR analyses of CD45− thymic stroma specific gene expression in TSCs, reconstructed thymus organoids cultured *in vitro* for 0 and 7 days (day 0 and day 7, respectively). Sample dilutions: undiluted, 1/4, 1/16 and 1/64. *f.* RT-PCR analyses of tissue-specific antigen transcription in reconstructed thymus organoids cultured *in vitro* for 0, 7, 28 and 56 days. All the experiments were repeated at least once with similar results.

**Figure 3. The reconstructed thymus organoids support T lymphopoiesis in athymic hosts *in vivo.* a.** Representative flow cytometric (FCM) profiles for both CD3+CD4+ and CD3+CD8+ T
cells in the blood circulation of B6.nude mice transplanted with reconstructed thymus organoids (Tot.B6.nude, 16-weeks post transplantation, lower panels), in comparison to the profiles of either the wildtype (WT) C57BL/6 (B6, 16-weeks old, top panels) or the athymic B6.nude (16-weeks old, middle panels) mice. Numbers indicate the frequencies of cells within the indicated areas. b. Progression of T-cell development in Tot.B6.nude mice. Percentages of CD3+CD45+ T-cells in peripheral blood leukocytes (PBLs) of Tot.B6.nude mice (n=4-15) were analyzed by FCM every 4-weeks post thymus organoid transplantation. Dashed line shows the percentage of T-cells in PBLs of 8-week old, wild type B6 mice (33.5±2.2%, n=5). Data were presented as mean ± SEM. c. Analyses of T lymphocytes in Tot.B6.nude mice. Splenocytes and lymph node cells were stained for CD3, CD45, CD4 and CD8, and were gated on the CD45+CD3+ population. Left panels, representative FCM profiles for CD4+ and CD8+ T-cells in the spleens (SPL, upper panels) and lymph nodes (LN, lower panels) of Tot.B6.nude mice (n=5, 20-weeks post-op) and B6 controls (n=3, 8-weeks old). Right panels, the total numbers of CD4 and CD8 T-cells in the spleens and lymph nodes of Tot.B6.nude (n=5) and B6 (n=3) mice. Data were presented as mean ± SEM. d. Spectratyping analysis of the CDR3 Vβ regions of mouse T cells isolated from the spleens of Tot.B6.nude (n=3) and WT B6 (n=3). e. FCM analyses of the origins of peripheral T-cells in Tot.B6.nude mice (n=5). CD4 T-cells harvested from lymph nodes of WT B6 (n=3), B6.CD45.1 congenic (n=3) and Tot.B6.nude mice were stained with CD45.1 and CD45.2 congenic markers. f. FCM analyses of activation status of peripheral CD4 and CD8 T-cells. T-cells harvested from spleens of WT B6 (open bar, n=5) and Tot.B6.nude (filled bar, n=5) mice were stained for CD62L and CD69. Left panel, representative dot blots. Numbers in the representative FCM profiles indicate the frequencies of cells within the indicated areas. Right panel, percentages of CD8+ (left columns) and CD4+ (right columns) T-cells displayed naïve
(CD62L+CD69-, top panels) or activated (CD62L-CD69+, lower panels) phenotypes. Data were presented as mean ± SEM. g. FCM analyses of the frequencies of Foxp3+ regulatory cells in the CD4 T-cell population. T-cells were harvested from spleens of WT B6 (n=5) and Tot.B6.nude (n=5) mice were stained intracellularly for Foxp3. Left panel, representative dot blots of CD4+Foxp3+ Tregs. Numbers in the representative FCM profiles indicate the frequencies of cells within the indicated areas. Right panel, numbers of CD4+Foxp3+ T-cells in the spleens of Tot.B6.nude mice (filled bar, n=5) and WT B6 controls (open bar, n=5). Data were presented as mean ± SEM. h. FCM analyses of CD4+Foxp3+ regulatory cells in the spleens of WT B6 (n=5) and Tot.B6.nude (n=5) mice. Representative dot blots showing CD25 (left panels) and Helios (right panels) expression in CD4+Foxp3+ Tregs. Numbers in the representative FCM profiles indicate the frequencies of cells within the indicated areas. Data were presented as mean ± SEM. i. Immunohistochemical analyses of reconstructed thymus organoids. Representative 7um cryosections of thymus organoid grafts, harvested from the kidneys of Tot.B6.nude mice (16-weeks post-op), were stained for EpCAM (left top panel, red), CD4 (left lower panel, red) and CD8 (right lower panel, red), and counter-stained with Hoechst 33342 (blue). Areas of the reconstructed thymus organoid graft (Thy) and the kidney (Kid) were separated by white dotted lines. Right top panel, a representative cryosection of a control WT B6 thymus stained with EpCAM (red). All the experiments were repeated at least once with similar results.

Figure 4. The reconstructed thymus organoids can support T-cell mediated immunity in athymic hosts in vivo. a. Proliferation of T cells in response to TCR stimulation. T-cells harvested from the spleens of Tot.B6.nude (n=4, right panels) and WT B6 mice (n=4, left panels) were labeled with CFSE and cultured for 7 days in the presence or absence of activating CD3
antibodies. Cells were stained for CD3 and B220, gated on the CD3+B220- populations were
analyzed by FCM for CFSE levels. All assays were run in triplicate. b. Proliferation of T cells in
response to alloantigens. T-cells enriched from Tot.B6.nude (n=4) and WT B6 mice (n=4) were
labeled with CFSE and cultured for 7 days in the presence (lower panels) and absence (upper
panels) of T-cell depleted allogeneic antigen presenting cells (APCs) isolated from CBA/J mice.
All proliferation assays were run in triplicate. c. Survival of allogeneic (CBA/J) skin grafts in
WT B6 (n=3), B6.nude (n=3) and Tot.B6.nude (n=3) mice, and syngeneic (B6) skin grafts in
Tot.B6.nude (n=3). d. Representative photographic images of allogeneic skin grafts in B6,
B6.nude and Tot.B6.nude mice at day 15 post transplantation. e. Representative histological
images (H&E) of paraffin sections of skin grafts harvested from Tot.B6.nude (n=4), B6.nude
(n=3) and WT B6 (n=3). Arrows indicate the infiltration of immune cells in the injected skin
grafts. f. Analyses of seroreactivities against ovalbumin (OVA) in naïve B6 mice (n=5, open
bar), B6 mice immunized with OVA (n=5, black bar), B6.nude immunized with OVA (n=3, grey
bar), and Tot.B6.nude immunized with OVA (n=4, shaded bar). Levels of OVA-reactive
immunoglobulins of various classes and subclasses were determined with ELISA-based,
colorimetric assay. NS, not significant; * p<0.05, nonparametric Mann-Whitney test. The assays
were repeated twice with similar results. g. ELISpot analyses of IFN-γ expressing T-cells.
Splenocytes harvested from immunized Tot.B6.nude mice (n=4) were challenged with OVA
peptides (AVHAHAEGNEGSIINFEKL). Left panels, representative ELISpot images (in
triplicate) of splenocytes cultured in presence (left columns) or absence (right columns) of OVA
peptides: top rows, immunized Tot.B6.nude mice (n=4); middle rows, naïve Tot.B6.nude (n=3);
lower rows, immunized WT B6 control (n=4). Right panels, numbers of IFN-γ producing spots
in the presence of medium (M, open bar) or OVA peptide (P, filled bar). Data were presented as
mean ± SEM. The results were obtained from three independent experiments. All assays were run at least in triplicate.

**Figure 5. Establishing donor-specific immune tolerance in mouse with reconstructed thymus organoids.**

**a.** Schematic outline of strategies to induce immune tolerance of allografts with reconstructed thymus organoid transplantation. Thymus organoids were constructed with TSCs harvested from F1 offspring of B6 (H-2b) and congenic B6.H2g7 (H-2g7) mice and transplanted to athymic B6.nude mice to generate Tot.B6.nude recipients. Once successful T-lymphopoiesis was demonstrated through FCM analyses of peripheral lymphocytes (12-16 weeks post thymus transplantation), tail skin grafts harvested from WT B6 (syngeneic), congenic B6.H2g7 (allogeneic) and CBA/J (H-2k, third party allogeneic) mice were transplanted to the recipients, and were monitored for their survival. **b.** Representative photographic images of syngeneic (B6) and allogeneic (CBA and B6.H2g7) skin grafts of Tot.B6.nude recipients (n=4) at day 1 and day 26. Arrows indicate the rejected graft. **c.** Representative histological images (H&E) of syngeneic (B6), allogeneic (B6. H2g7) and third party allogeneic (CBA) skin grafts harvested from Tot.B6.nude recipients (n=4) at 26 days post skin transfer. Higher magnified images of areas in the red boxes are shown in the lower panels. White arrows show areas with lymphocytic infiltration. **d.** Proliferation of T cells of Tot.B6.nude mice (n=3) in response to alloantigens. T-cells enriched from Tot.B6.nude (n=3) were labeled with CFSE and cultured for 7 days in the presence of T-cell depleted syngeneic (B6), allogeneic (B6.H2g7), and third party allogeneic (CBA) APCs. Representative FCM results of CD4+ (*left* panels) and CD8+ (*right* panels) T cells.
are shown. N/A, no APCs were added to the culture. All proliferation assays were run in triplicate.

**Figure 6. Induction of donor-specific immune tolerance with thymus organoids reconstructed of both donor and recipient TECs.**

**a.** The schematic drawing shows the strategy of the experiment. TECs were isolated from B6 and CBA/J mice and mixed at 1:1 ratio. The thymus organoids were reconstructed with B6 Lin- bone marrow progenitors and TECs at 1:1 ratio (2x10^5 each), and transplanted underneath the kidney capsules of B6.nude mice. Ten weeks post-op, skin grafts harvested from CBA/J and Balb/C (third party allografts) mice were transplanted to the Tot.B6.nude mice (n=4). Graft survival was monitored for up to 4 weeks. **b.** Representative photographic images of skin grafts (outlined) on Tot.B6.nude at 22 days post-op. While the third party allograft (left panel, Balb/C) is largely rejected, the CBA/J allograft (right panel) is well tolerated. **c-d.** Mixed lymphocyte reaction (MLR) experiments, showing the proliferation responses of CD3+ T-cells of the recipient Tot.B6.nude mice (n=4) in the presence of syngeneic or allogeneic APCs. **c.** Representative CFSE dilution histogram: **Left panel,** syngeneic B6 APCs; **middle panel,** CBA/J allogeneic APCs; **right panel,** third party allogeneic APCs. **d.** Percentages of proliferating CD3+ T-cells. * p<0.05, nonparametric Mann-Whitney test.