FOXN1$^{\text{GFP/w}}$ Reporter hESCs Enable Identification of Integrin-β4, HLA-DR, and EpCAM as Markers of Human PSC-Derived FOXN1$^+$ Thymic Epithelial Progenitors

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SUMMARY

Thymic epithelial cells (TECs) play a critical role in T cell maturation and tolerance induction. The generation of TECs from in vitro differentiation of human pluripotent stem cells (PSCs) provides a platform on which to study the mechanisms of this interaction and has implications for immune reconstitution. To facilitate analysis of PSC-derived TECs, we generated hESC reporter lines in which sequences encoding GFP were targeted to FOXN1, a gene required for TEC development. Using this FOXN1GFP/w line as a readout, we developed a reproducible protocol for generating FOXN1-GFP+ thymic endoderm cells. Transcriptional profiling and flow cytometry identified integrin-β4 (ITGB4, CD104) and HLA-DR as markers that could be used in combination with EpCAM to selectively purify FOXN1+ TEC progenitors from differentiating cultures of unmanipulated PSCs. Human FOXN1+ TEC progenitors generated from PSCs facilitate the study of thymus biology and are a valuable resource for future applications in regenerative medicine.

INTRODUCTION

T cells undergo most of their development in the thymus, the primary lymphoid organ that regulates their differentiation and maturation from blood-borne bone-marrow-derived precursors and appropriate selection for the induction of self-tolerance (Anderson et al., 2007). Thymic function is critically dependent on thymic epithelial cells (TECs), the most abundant cellular constituent of the stromal microenvironment. TECs are classified as two morphologically and functionally distinct subsets based on their localization to the thymic cortex (cTECs) or medulla (mTECs). TEC development and identity require the forkhead-box transcription factor Foxn1, which, in the mouse, demarcates the prospective thymic primordium within the third pharyngeal pouch at embryonic day 11.5 (E11.5). Loss of Foxn1 results in a “nude” phenotype in mice and rats (Nehls et al., 1994, 1996) and in humans (Pignata et al., 1996), characterized by congenital hairlessness and defective TEC differentiation, the latter of which results in the absence of functional T cells and severe immunodeficiency.

T cell insufficiency is associated with other congenital thymic hypoplasias, such as DiGeorge syndrome (Jerome and Papaioannou, 2001), age-related thymus atrophy, or cytoblastic therapy-induced thymic involution. In the last case, patients undergoing high-dose chemotherapy often experience chronic immunosuppression, predisposing them to a host of opportunistic infections. In all of these instances, replenishment of the thymic epithelial compartment might provide an avenue to augment thymus function and boost T cell output.

The derivation of tissues by in vitro differentiation of pluripotent stem cells (PSCs) has been advanced as a platform for use in the emerging fields of cell therapy and regenerative medicine. PSCs possess the capacity to give rise to the three embryonic germ layers, including the definitive endoderm, the precursor of thymic epithelium. PSC differentiation protocols that promote definitive endoderm formation (D’Amour et al., 2005; Kubo et al., 2004; Tada et al., 2005; Yasunaga et al., 2005) and the subsequent generation of posterior foregut derivatives, including pancreatic cells (D’Amour et al., 2006; Kroon et al., 2008) and hepatocytes (Cai et al., 2007; Gouon-Evans et al., 2006), are well established. Similarly, strategies designed to generate anterior foregut derivatives, such as the lung, thyroid, and thymus, have also been reported (Green et al., 2011; Lai and Jin, 2009; Longmire et al.,...
Figure 1. Targeted FOXN1<sup>GFP</sup> hESCs Facilitate Monitoring of Thymic Differentiation toward FOXN1-GFP<sup>+</sup> Cells

(A) Targeting strategy used to insert sequences encoding GFP into exon 2 (e2) of the FOXN1 gene locus via homologous recombination. The selection cassette included a neomycin resistance gene (neo<sup>a</sup>) flanked by loxP sites (black triangles). SwaI restriction endonuclease was used to linearize the vector prior to electroporation into hESCs. The locations of PCR primer pairs (a, b, and c) and a GFP probe (blue line) used to characterize targeted clones are also indicated.

(B) PCR analyses of HES3-derived FOXN1<sup>GFP</sup> hESCs with primer pairs (a and b) spanning the junctions between the targeting vector and genomic DNA generated bands of 9.6 kb (5' end) and 3.9 kb (3' end) from targeted FOXN1<sup>GFPneoR/w</sup>, but not from wild-type hESCs. PCR analysis with primer pair c flanking the neo<sup>a</sup> gene generated bands of 3.7 kb compared to 5.4 kb, indicating excision of the neo<sup>a</sup> cassette. Southern blot analysis of HindIII-digested FOXN1<sup>GFP</sup> genomic DNA using the GFP probe detected a single 4.2 kb fragment, demonstrating integration of a single copy of the FOXN1-targeting vector.

(C) Heatmap summarizing the results of flow cytometry experiments assessing the percentage of GFP<sup>+</sup> cells from days 30–35 embryoid bodies generated with the indicated BMP4 and Activin A concentrations. Data were derived from three independent experiments. FACS profiles representing selected conditions are shown on the right.

(D) Timecourse of KGF addition to the culture. Graft aggregation is shown starting from Day 0 to Day 14 onwards.

(E) KGF Addition Timecourse

(legend continued on next page)
Recently, two groups reported methods for the generation of thymic endoderm from human pluripotent stem cells (Parent et al., 2013; Sun et al., 2013). Importantly, these studies showed that differentiated mixed cultures containing thymic progenitors could mature in vivo to form grafts capable of supporting T cell development in nude (Foxn1−/−) or other immunocompromised mice. However, because of the lack of appropriate surface markers, it was not possible to dissect the contributions or requirements of the various cell types present in the differentiation cultures at the time of transplantation. This deficit could be partly remedied by the availability of FOXN1 reporter lines or surface markers that allow further fractionation of cultures containing FOXN1+ cells.

We generated FOXN1GFP/w human embryonic stem cell (hESC) reporter lines that were used to develop a robust serum-free protocol for the generation of FOXN1+ thymic endodermal progenitors. We found that high levels of Activin A and KGF efficiently induced the differentiation of FOXN1+ cells and that these cells expressed genes involved in endoderm and thymus development. Transcriptional profiling of purified FOXN1-GFP+ cells allowed the identification of several combinations of cell-surface markers that could selectively isolate FOXN1+ TEC progenitor populations derived from unmodified PSC lines. Collectively, these reagents and findings represent a valuable resource for the further investigation of thymic development from pluripotent stem cells.

**RESULTS**

**FOXN1GFP/w hESCs Facilitate Analysis of Thymic Differentiation In Vitro**

To facilitate analysis of thymic differentiation of PSCs, we used homologous recombination to target a GFP reporter gene to the endogenous FOXN1 locus in MEL1 (Millipore) and HES3 (Richards et al., 2002) hESCs (Figures 1A and 1B; Figure S1A available online). Targeted FOXN1GFP/w hESCs retained expression of pluripotent stem cell markers, formed multilineage teratomas when transplanted into immunodeficient NOD/SCID mice, and maintained normal karyotypes (Figures S1B–S1D). In order to identify conditions that favored the formation of FOXN1+ endodermal cells, we differentiated FOXN1GFP/w hESCs in serum-free media as spin embryoid bodies (EBs) (Ng et al., 2008) and cross-titrated bone morphogenetic protein-4 (BMP4) and Activin A, growth factors that activate signaling pathways required for the generation and patterning of mesendoderm and have been previously shown to promote PSC differentiation in vitro (D’Amour et al., 2005; Davis et al., 2008; Kubo et al., 2004; Tada et al., 2005; Yasunaga et al., 2005). This heatmap analysis employed an iterative process (Elliott et al., 2011) and identified combinations of BMP4 and Activin A that promoted the appearance of FOXN1-GFP+ cells, which also coexpressed with the endodermal marker EpCAM (Figure 1C; Figures S2A–S2C). Based on these results and findings from experiments using different batches of Activin A (data not shown), for all subsequent experiments, we differentiated hESCs for 5 days in serum-free medium supplemented with 150 ng ml−1 Activin A alone (Figure 1D). These high Activin conditions were similar to those used in previously published differentiation protocols where definitive endoderm derivatives were the desired outcome (D’Amour et al., 2006; Green et al., 2011). Flow cytometric analysis of differentiating MIXL1GFP/w hESCs, which can be used to monitor mesendoderm induction (Davis et al., 2008), indicated that at differentiation day 4, over 99% of cells were MIXL1-GFP+, arguing against the possibility that FOXN1-GFP+ cells represented ectodermally derived keratinocytes (Figure S3A). Time course quantitative real-time PCR analysis of differentiation cultures demonstrated that from day 12, cells progressively upregulated expression of the pharyngeal pouch markers HOXA3 and PA2X9, as well as the thymic markers INVOLUCRIN, FGFR2 (the receptor for KGF), KERATIN-10, and FOXN1 itself. Expression of the general endodermal markers FOXA2 and SOX17 declined from a peak at day 12, consistent with expansion of a specialized anterior endoderm population that does not express these markers at high levels (Figure S3B).

Keratinocyte growth factor (KGF or FGF7), a factor previously implicated in thymic development, was required to expand the FOXN1-GFP+ population once it emerged in culture, with the timing of the KGF supplementation affecting the percentage of GFP+ cells obtained (Figures 1D and 1E). KGF is expressed in the thymus by the surrounding mesenchyme and by different thymocyte subsets (Rossi et al., 2007) and exerts a paracrine effect by binding to the Fgfr2IIb receptor, which, within the thymus, is expressed exclusively by TECs (Erickson et al., 2002). Moreover, exogenous administration of KGF has been shown to increase thymic cellularity by enhancing the proliferation and function of TECs (Erickson et al., 2002; Min et al., 2007; Rossi et al., 2007). After the addition of KGF, GFP+
cells could be visualized using fluorescence microscopy from approximately day 25 of differentiation. Flow cytometric analysis of cells differentiated as outlined in Figure 1D indicated that the first FOXN1-GFP+ cells emerged from a PDGFRα- and EpCAM+ population as early as differentiation day 15 and increased in frequency thereafter (up to 38% GFP+ at day 35) (Figure 2A). MEL1-derived FOXN1-GFP EPSCs differentiated with similar kinetics to the HES3-derived line, suggesting our protocol is applicable to other PSC lines (Figure 2A). Quantitative real-time PCR analysis of cells purified on the basis of GFP and EpCAM expression showed that FOXN1 transcripts were confined to the GFP+ population (Figure 2B), confirming the fidelity of the GFP reporter gene. This conclusion was supported by intracellular flow cytometric analysis of FOXN1 protein expression in purified GFP+ and GFP- cells (Figure S3C). Prolonged culture of aggregates of fluorescence-activated cell sorting (FACS)-purified GFP+EpCAM+ cells indicated that GFP expression was retained for up to 3 weeks (Figure 2B).

Gene Expression Analysis of hESC-Derived FOXN1+ TEC Progenitors

We next performed microarray analysis to survey genes expressed in the GFP+ fraction and to search for markers that might identify FOXN1+ cells in differentiating cultures derived from unmanipulated human PSCs. We first compared the global gene expression within the GFP+ and GFP- EpCAM+ epithelial subpopulations. This analysis identified 115 genes that were upregulated in the GFP+ EpCAM+ fraction compared to the GFP- EpCAM+ fraction (Figure 3A; see Tables S1 and S2 for the full list of genes differentially upregulated by 3-fold), and many of these have been previously associated with thymus development and/or FOXN1 expression. These included the canonical WNT-signaling molecules WNT4 and WNT3 and the WNT regulator KRM2, the NOTCH ligands DLL1, thought to be required for crosstalk in driving T-lymphopoiesis, and JAG2 (Candi et al., 2007), a downstream effector of the ΔNp63 isofrom of the TP63 transcription factor, required for the proliferation of TECs and other stratified epithelia, human TEC type I cytokeratins KRT14, KRT15, and KRT16, and the chemokine CXCL14 whose transcript is expressed in fetal TECs and may aid in the recruitment of T cell precursors (Liu et al., 2005). Conversely, 213 transcripts were preferentially upregulated in the GFP- EpCAM+ population (Figure 3A). These included HAND1 and ISL1, genes expressed during cardiac morphogenesis and in neural crest derivatives, although ISL1 is also expressed in pancreatic endoderm, other definitive endodermal markers, such as APOA1, SOX9, and TGFβ1, and human TEC type II cytokeratins KRT6, KRT7, and KRT8. Interestingly, the chemokine CXCL12 was also upregulated within the GFP+ EpCAM+ population. CXCL12 is expressed within TECs and bone marrow niches, where it acts as a chemoattractant for T cell precursors, monocytes, and hematopoietic stem cells (HSCs) (Greenbaum et al., 2013; Liu et al., 2005). Overall, the GFP+ EpCAM+ population expressed genes associated with gene ontology (GO) categories including “blood circulation” (p value < 0.002), “circulatory system process” (p value < 0.002), “cell migration” (p value < 0.003), and “cell motility” (p value < 0.006) (Figure S4A). In contrast, GFP+EpCAM+ cells were enriched for genes associated with FOXN1 expression, thymus development, or epithelial-related GO categories (Figure S4A). Although it is not possible to infer from gene expression data which genes are direct targets of FOXN1, the ability to generate and purify a large number of FOXN1+ cells provides an opportunity to examine this issue using techniques such as chromatin immunoprecipitation (ChIP).

Last, we also compared gene expression of the GFP+ EpCAM+ and GFP-EpCAM- populations (Figure S4B). Included among the transcripts preferentially expressed within GFP+EpCAM+ cells were genes with a role in thymus development (KRM2, JAG2, WNT4, BCL11B, EFN1, and EYA2), epithelial morphogenesis (KRT5, TP63, EpCAM, KRT14, ECAD, ITGB4, and EGFR), and endoderm specification (ALB and AFP). Keratin-5 and Keratin-8 have been detected within the human thymus as markers of the medulla and cortex, respectively (Laster et al., 1986; Shezen et al., 1995). Our microarray analysis demonstrated high coexpression of Keratin-5 and its heterodimerization partner Keratin-14 within FOXN1+ (GFP+EpCAM+) cells (Figure 3B). The former result was confirmed by quantitative real-time PCR (Figure S4C). Interestingly, although also detected in GFP+EpCAM+ cells, Keratin-8 is preferentially expressed by the GFP-EpCAM- subpopulation (Figure 3A, Figure S4C). These results might suggest that the FOXN1+ cells generated in our protocol more closely reflect a medullary phenotype. Finally, FOXN1+ cells were devoid of genes that mark the parathyroid gland, including GCM2, PTH, and CaSR (Figures S4D and S4E), consistent with the observation that these two developmentally related tissues can be delineated on the basis of FOXN1 and GCM2 expression (Gordon et al., 2001). Similarly, other genes associated with the demarcation of thymic versus parathyroid primordia (BMP4, BMPR1A, BMP2, and NOGG) were not restricted to specific subpopulations (Figure S4E).

Examination of microarray data indicated that a number of transcription factors, signaling molecules, and structural proteins implicated in TEC development were preferentially expressed in GFP+EpCAM+ cells (Figure 3B). We were particularly interested in identifying transmembrane proteins for which well-characterized cell-surface antibodies were readily available. Such antibodies could then

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Figure 2. Activin A and KGF Induce GFP Expression in Differentiating FOXN1<sup>GFP/w</sup> hESCs

(A) Flow cytometric analysis of differentiating HES3-derived FOXN1<sup>GFP/w</sup> hESCs shows FOXN1-GFP<sup>+</sup> cells emerge at day 15 and subsequently increase thereafter. FOXN1-GFP<sup>+</sup> cells coexpress the thymic epithelial marker EpCAM, but not the mesodermal marker PDGFR<sub><sup>α</sup></sub>. Flow cytometric analysis of a second independent MEL1-derived FOXN1<sup>GFP/w</sup> hESC line demonstrated that the time course of GFP and EpCAM induction was comparable between HES3- and MEL1-derived lines.

(B) Quantitative real-time PCR analysis showed that FOXN1 expression was confined to the GFP<sup>+</sup>EpCAM<sup>+</sup> fraction, with expression absent from the GFP<sup>+</sup>EpCAM<sup>-</sup> and GFP<sup>-</sup>EpCAM<sup>+</sup> subpopulations. Aggregated GFP<sup>+</sup>EpCAM<sup>+</sup> cells retained GFP expression for up to 3 weeks.
be employed to isolate FOXN1+ cells from differentiating cultures of genetically unmodified PSC lines. One such gene upregulated in GFP+EpCAM+ cells was integrin-α6 (ITGB4 or CD104) (Figure 3B). The ITGB4 subunit commonly associates with the integrin-α6 (ITGA6) subunit to form an integrin α6β4 heterodimer that functions as a receptor for laminins within the extracellular matrix (Giannotti, 1996). ITGB4 is expressed in normal human TECs, where it is thought to activate intracellular signaling pathways that regulate production of the thymic cytokine IL-6, which in turn affects TEC differentiation and thymocyte homing (Mainiero et al., 2003). Moreover, ITGB4 is also expressed by cultured human TEC lines (Beaudette-Zlatanova et al., 2011). Time course analysis showed low expression of ITGB4 in early differentiating embryoid bodies, which preceded the appearance of FOXN1-GFP+ cells, and was associated with a population that concomitantly upregulated EpCAM expression (Figure 4A). Indeed as differentiation proceeded, by day 15 the majority of EpCAM+ cells coexpressed ITGB4, consistent with the expression of these markers on a broad range of epithelial tissues (Figure 4A). EpCAM+, FOXN1-GFP+ cells appearing from differentiation day 20 onward expressed high levels of ITGB4 (ITGB4hi). Furthermore, as the number of FOXN1-GFP+ cells increased in response to KGF treatment, GFP expression became more clearly associated with the ITGB4hiEpCAM+ subpopulation. Quantitative real-time PCR analysis of FACS-purified day 34 cells derived from the differentiation of FOXN1 GFP/h ESCs confirmed that FOXN1 gene expression was largely confined to the ITGB4hiEpCAM+ population and that expression levels were comparable to those seen in cells sorted on the basis of GFP expression alone (Figure 4B). Negligible FOXN1 expression was observed in sorted populations that expressed intermediate or low levels of ITGB4 or were EpCAM-.

To explore the utility of these cell-surface makers for monitoring developing TECs, we examined other human pluripotent stem cell lines, including MEL1-derived FOXN1 GFP/h ESCs, genetically unmodified H9 hESCs (Thomson et al., 1998) and DF19-9-7T human iPSCs (Yu et al., 2009) (Figure 4B). All human PSC lines generated ITGB4hiEpCAM+ cells with parallel kinetics, although the proportion of this population varied between different cell lines. Cell populations that expressed ITGB4 and EpCAM but were not ITGB4hi (designated ITGB4 EpCAM+) varied in composition between different PSC lines. Nevertheless, we found FOXN1 expression to be solely confined to those EpCAM+ cells that were ITGB4hi (Figures 4B and 4C). Overall, flow cytometric analysis of HES3 and MEL FOXN1 GFP/h cultures indicated that FOXN1-GFP+ cells comprised over 80% of the ITGB4hiEpCAM+ fraction (Figure 4F). Conversely, 95% of the GFP+ population was found to be ITGB4hiEpCAM+, testifying to the close association between these markers and FOXN1 expression (Figure 4G).

In addition to those markers identified by transcriptional profiling, we examined the expression of the TEC-associated receptor, HLA-DR: a human major histocompatibility complex class II (MHCII) cell-surface receptor encoded by the human leukocyte antigen (HLA) complex (Walsh et al., 2003). Within the thymus, the primary function of
HLA-DR, similar to other MHCII genes, is to present peptide antigens during positive and negative T cell selection in the thymic cortex and medulla (Yang et al., 2006). TECs are unique in that they constitutively express MHCII without the need for interferon-γ (IFN-γ) stimulation (Drukker et al., 2002; Yang et al., 2006), consistent with the fact
that IFN-γ-deficient mice undergo normal thymic development (Dalton et al., 1993). To explore whether HLA-DR could also identify FOXN1-GFP + cells, we performed flow cytometry to monitor the expression of HLA-DR and EpCAM over a 40 day differentiation time course (Figure 5A). Similar to previous results (Drukker et al., 2002), HLA-DR was not expressed on the surface of undifferentiated hESCs (data not shown) or during the early stages of embryoid body formation. Expression was first observed from approximately day 15 and increased thereafter. By day 40, we detected a distinct population of HLA-DR + EpCAM + cells that contained the majority of FOXN1-GFP + cells (Figure 5A). Furthermore, all HLA-DR + cells were also EpCAM +, suggesting that HLA-DR is restricted to only thymic epithelial cells in our culture system. Consistent with past studies, HLA-DR expression on FOXN1-GFP + TEC progenitors was not upregulated in response to IFN-γ stimulation. Conversely, as expected, IFN-γ treatment induced expression of HLA-DR in the non-GFP population (Figure S4H). Finally, quantitative real-time PCR analysis was performed on day 30 differentiated cells purified on the basis of HLA-DR and EpCAM expression (Figure 5B). This analysis showed that FOXN1 transcripts were greatly enriched in the HLA-DR + EpCAM + population differentiated from HES3-derived FOXN1GFP/w cells and were present at levels similar to those observed in FOXN1-GFP + cells. FOXN1 expression in differentiating MEL1-derived FOXN1GFP/w cells and H9 hESCs was also enriched in the HLA-DR + EpCAM + population. Colored boxes represent the gating strategy used to isolate subpopulations for flow cytometric sorting. H, HLA-DR; E, EpCAM.

Figure 5. HLA-DR and EpCAM Identify PSC-Derived FOXN1+ Thymic Epithelial Progenitor Cells

(A) Time course analysis of differentiating HES3-derived FOXN1GFP/w cells demonstrated that FOXN1-GFP + cells could be identified on the basis of coexpression of HLA-DR and EpCAM. EpCAM + cells are shown gated in light blue, whereas GFP + cells are shown gated in green. Numbers indicate the percentage of cells in the corresponding quadrants.

(B) Quantitative real-time PCR analysis confirmed that FOXN1 transcripts were substantially enriched in the HLA-DR + EpCAM + population differentiated from HES3-derived FOXN1GFP/w cells and were present at levels similar to those observed in FOXN1-GFP + cells. FOXN1 expression in differentiating MEL1-derived FOXN1GFP/w cells and H9 hESCs was also enriched in the HLA-DR + EpCAM + population. Colored boxes represent the gating strategy used to isolate subpopulations for flow cytometric sorting. H, HLA-DR; E, EpCAM.

(C) Histograms demonstrating the summary of raw (left panel) and normalized (right panel) data from individual quantitative real-time PCR experiments shown in Figure 5B (n = 3). Data are shown as the mean ± SEM. p values were calculated using a two-tailed Student’s t test.
fraction, with expression levels comparable to those seen in cells sorted on the basis of GFP alone. The veracity of this result was confirmed in similar analyses of HLA-DR+ EpCAM+ cells isolated from differentiating cultures of MEL1 FOXN1+GFP+ and H9 hESC lines (Figures 5B and 5C). Similar to results obtained for cells isolated on the basis of ITGB4 and EpCAM expression, flow cytometric analysis indicated FOXN1-GFP+ cells comprised over 70% of cells within the HLA-DR+EpCAM+ fraction (Figure S4I). The clear association of FOXN1 expression with HLA-DR further substantiates the thymic identity of cells generated using our protocol.

The upregulation of HLA-DR in conjunction with gene expression analyses indicating the presence of NOTCH signaling receptors and ligands prompted us to test whether FOXN1-GFP+ cells generated with our differentiation protocol might be competent to promote the development of T lineage cells from human CD34+ hematopoietic progenitors. To test this hypothesis, we cocultivated purified FOXN1-GFP+ with human umbilical cord blood-derived CD34+CD7+ T lineage progenitors, a cell type previously shown to differentiate into CD4+ or CD8+ T cells (Awong et al., 2009; La Motte-Mohs et al., 2005). Over a 4 week coculture period, we analyzed the hematopoietic component for expression of CD1a, CD5, CD4, CD8, and CD3, markers of immature T lineage progenitors, differentiating pro-T-cells and mature T cells. Although we observed the induction of CD1a, CD5, CD4, and CD8 expression, the differentiation step was not specifically dependent on the presence of FOXN1-GFP+ cells (Figures S5A–S5C and S5E). Moreover, we did not observe convincing expression of the T-lineage-specific marker CD3 (Figure S5D). Rather, a significant proportion of CD34+ cells differentiated into cells with a myeloid phenotype (CD45+CD14+) (Figure S5E). Hence, although our human PSC-derived cells possess a TEC-like phenotype, we believe that they represent a stage that is too immature to drive T cell differentiation. Further investigations into the growth factors necessary for the commitment of FOXN1+ TEC progenitors into mature TECs in our culture system are therefore required.

**DISCUSSION**

We devised a simple method for generating and isolating FOXN1+ thymic epithelial progenitor cells differentiated from human pluripotent stem cells. We inserted a GFP reporter gene into the locus encoding the key thymic transcription factor FOXN1, to facilitate monitoring of the early steps involved in human thymus specification. This reporter gene faithfully identified cells expressing FOXN1 and enabled isolation of FOXN1+ thymic progenitors from heterogeneous differentiation cultures. Although targeted lines represent an extremely useful research tool, genetically manipulated lines are undesirable for potential future clinical use. Moreover, the technical requirements for genetic modification make the application of this approach impractical for the ever-expanding number of available pluripotent stem cell lines. With these points in mind, we interrogated the expression profile of FACSPurified FOXN1-GFP+ cells through microarray analysis with the aim of identifying genes that could serve as a surrogate marker of FOXN1 expression. From this screen, we identified ITGB4 as a cell-surface marker that, in conjunction with EpCAM, could be used to isolate FOXN1+ cells from genetically unmodified hESCs and from human iPSCs. Similarly, our analysis showed that HLA-DR and EpCAM were also specific markers of FOXN1+ TEC progenitors. Taken together, our study provides a model system to investigate the signaling pathways required for TEC commitment and differentiation.

Gene expression analysis revealed that cells differentiated for up to 30 days expressed markers implicated in endoderm specification and TEC development but lacked the expression of important late-stage TEC markers associated with functional maturation. In particular, relative to human pediatric thymic stroma (data not shown), in vitro-derived FOXN1+ TEC progenitors showed low expression of autoimmune regulator (AIRE), a gene critical for intrathymic expression of tissue-restricted antigens, which in turn is required to induce tolerance to peripheral antigens (Anderson et al., 2007). In addition, CD80, a costimulatory molecule expressed on functionally mature mTECs (Derbinski et al., 2005), was also not enriched on FOXN1+ cells. These data indicated that the thymic endoderm produced using our culture conditions most likely represented an early stage of ontogeny in which overt thymic functional characteristics were yet to be acquired.

The latter stages of thymic epithelial development are dependent on the presence of hematopoietic cells that, in the mouse, infiltrate the thymic primordium at embryonic day 12, soon after the onset of FOXN1 expression. Subsequent to this, the thymic endoderm and lymphoid progenitors engage in signaling crosstalk that ultimately yields functionally mature TECs and educated T cells. The Notch-Delta system is a central signaling mechanism implicated in directing this crosstalk (Mohtashami et al., 2010). In the thymus, Notch1 expressed on hematopoietic progenitors interacts with Dll1 or Dll4 on TECs to either maintain or induce T-lineage commitment and differentiation. Because of this, we tested whether coculture of FOXN1+ thymic endoderm with hematopoietic progenitor cells could provide an environment whereby both cell types could undergo further maturation. However, under the conditions used, we saw no evidence of ongoing
Tcell differentiation. Moreover, analysis of these cocultures suggested that the viability of thymic endoderm cells was compromised, suggesting that further work will be required to define conditions that are permissive for the continued growth and development of both hematopoietic and endodermal cell types.

Recently, Parent et al. (2013) and Sun et al. (2013) reported methods for the generation of thymic endoderm from pluripotent stem cells. Following induction of definitive endoderm by Activin A treatment, both groups could promote the appearance of anterior foregut and subsequently pharyngeal endoderm by carefully choreographed manipulation of WNT, retinoic acid, and BMP4 signaling, mirroring the inductive and repressive actions of these pathways during early thymic ontogeny. Both studies showed that mixed differentiation cultures containing thymic progenitors could instruct T cell development in xenotransplantation models. In our differentiation protocol, the activities of the WNT, retinoic acid, and BMP4 pathways were not deliberately manipulated with exogenous growth factors or inhibitors, potentially explaining the inability of our FOXN1+ cells to upregulate markers of late-stage TEC differentiation or to support T cell development. Having said this, we did not formally test whether heterogeneous non-FACS-purified populations containing FOXN1+ cells could differentiate further following transplantation or indeed support T cell development in vitro. Consequently, the potential functionality and differentiation status of the cells generated using our simple method is difficult to directly compare with cells described by Parent et al. (2013) and Sun et al. (2013), where mixed cultures were used for in vivo studies.

The cell lines and surface markers reported in this study provide a facile platform on which to begin the process of identifying key events and critical cell populations required to derive fully functional TECs from PSCs. These FOXN1GFP/w reporter lines represent a unique research tool to analyze human thymus development, whereas the cell-surface marker combinations recognized by antibodies against ITGB4, HLA-DR, and EpCAM can be used to purify TEC progenitors from cultures of differentiating unmanipulated PSCs. Although the exogenous growth factors required for ongoing differentiation of FOXN1+ progenitor cells into functionally mature TECs remain to be established, our study contributes critical tools for efforts to generate this clinically important cell type.

EXPERIMENTAL PROCEDURES

Construction of the FOXN1-Targeting Vector

The FOXN1-targeting vector was assembled using standard cloning techniques and Gene Bridges Red/ET Recombination technology (see the Supplemental Experimental Procedures). The final 18.2 kb FOXN1-targeting vector comprised a 9.6 kb 5’ homology arm, a GFP-coding sequence, a loxp-flanked neomycin resistance gene driven by the mouse phosphoglycerate kinase (PGK) gene promoter, a 3.9 kb 3’ homology arm, and a pBR322-based plasmid backbone. The targeting vector was linearized by Swal digestion prior to electroporation.

Generation, Identification, and Characterization of Targeted FOXN1GFP/w hESCs

The FOXN1-targeting vector was electroporated into hESCs as previously described (Costa et al., 2007) and as detailed in the Supplemental Experimental Procedures. The loxp-flanked neoR cassette was removed by Cre recombinate-mediated excision as previously described (Davis et al., 2008). For one of the two independently derived targeted hESC lines (HES3 FOXN1GFP/w and MELI FOXN1GFP/w), Southern blot analysis with a labeled GFP probe (Figure 1A) was used to confirm the presence of a single-integration event (Figure 1B). Karyotype analysis was performed by the Cytogenetics Department at Southern Cross Pathology, Monash Medical Centre. Teratoma formation and analysis were performed as previously described (Costa et al., 2005). Animal experiments were conducted under the approval of the Monash University Animal Ethics Committee (number S0B1/SIS/2009/07).

hESC and iPSC Culture and Differentiation

hESC lines (HES3, MEL1, and H9 [Thomson et al., 1998]) and the induced pluripotent stem cell (iPSC) line DF19-9-7T (Yu et al., 2009) were maintained as previously described (Costa et al., 2008). hESCs and iPSCs were differentiated as spin embryoid bodies in serum-free media as previously described (Ng et al., 2008). Briefly, 1 day prior to differentiation, cells were passaged onto a new tissue culture flask seeded with low density (1 x 10^4 cells/cm^2) mouse embryonic fibroblasts. At day 0, cells were harvested and deposited into each well (3 x 10^5 cells/well) of a 96-well round-bottom nonadherent plate (Nunc) and briefly centrifuged to promote cell aggregation.

Heatmap experiments were performed using APEL medium (Ng et al., 2008), supplemented with 0–320 ng ml^-1 Activin A (R&D Systems) and 0–320 ng ml^-1 bone morphogenetic protein-4 (BMP4; R&D Systems) in 96-well round-bottom low-attachment plates (Costar) (Elliott et al., 2011). Cytokines were removed on day 5; at day 7, embryoid bodies were transferred to 96-well flat-bottom adherent tissue culture-treated plates (BD Falcon) containing APEL medium lacking PVA (APEL medium) (Ng et al., 2008). After 30–35 days of differentiation, embryoid bodies were dissociated with TrypLE Select (Invitrogen), stained with the relevant antibodies, and analyzed with FACSDIVA software (BD Biosciences) on either the BD LSRII or BD LSRRostra cell analyzers (BD Biosciences), fitted with a BD high-throughput sampler (HTS) module.

For thymic endoderm differentiation assays, cells were initially cultured in APEL or BPEL medium (Ng et al., 2008), supplemented with 150 ng ml^-1 human Activin A. On day 5, Activin A-containing APEL or BPEL medium was replaced with APEL or BPEL medium alone. On day 7, EBs were transferred to 96-well flat-bottom adherent plates (BD Falcon) in AEL or BEL medium (media lacking...
FOXN1-GFP hESCs Identify Cell-Surface TEC Markers

PVA) (Ng et al., 2008). AEL or BEL medium supplemented with 40 ng ml⁻¹ human keratinocyte growth factor (KGF; R&D Systems) was used to replenish the cultures on days 14, 21, 28, and 35. Embryoid bodies were harvested for analysis by flow cytometry at the times indicated. For experiments using MIXL1-GFP/w hESCs (Davis et al., 2008), cells were differentiated in BPEL medium supplemented either with 150 ng ml⁻¹ Activin A, 20 ng ml⁻¹ human BMP4, and 100 ng ml⁻¹ Activin A, or with 100 ng ml⁻¹ FGF2 (Peprotech). In all instances, hESC and iPSC cultures and differentiations were maintained at 37°C in a 5% CO₂/air environment.

Flow Cytometric Analysis and Sorting

For analysis and sorting of live cells, hESCs and embryoid bodies were dissociated to a single-cell suspension using TrypLESelect (Invitrogen), filtered through a 35 μM cell-strainer cap (BD Falcon), and labeled with the appropriate antibodies (see the Supplemental Experimental Procedures) as previously described (Davis et al., 2008). Flow cytometry was performed using a FACScalibur, FACS Diva, or Influx Cell Sorter (all from BD Biosciences). Flow cytometric gates were set using unmodified hESCs or FOXN1-GFP-targeted cells labeled with an appropriate isotype control antibody. Live cells were identified on the basis of forward scatter, side scatter, and propidium iodide (PI) exclusion.

Intracellular flow cytometry using mouse anti-human OCT4 (clone C-10; Santa Cruz Biotechnology) and mouse anti-human FOXN1 (clone E-3; Santa Cruz Biotechnology) antibodies was performed essentially as described by Davis et al. (2008) and as detailed in the Supplemental Experimental Procedures.

Culture of Sorted Cell Aggregates

Cells purified by flow cytometry were aggregated by centrifugation for 5 min at 1,500 rpm, using the spin EB protocol (5–10 × 10⁶ cells/well) in BPEL medium supplemented with 5 μM Y27632 ROCK inhibitor, as previously described (Goulburn et al., 2011). Aggregated cell clusters were cultured on gelatin-coated wells of a 96-well flat-bottom adherent plate in BEL medium supplemented with 40 ng ml⁻¹ KGF.

Quantitative Real-Time PCR and Microarray Gene Expression Analysis

Total RNA was prepared using a High Pure RNA Isolation Kit (Roche), in accordance with the manufacturer's instructions. Quantitative real-time PCR was performed essentially as described by Pick et al. (2007), with probes detailed in the Supplemental Experimental Procedures. For microarray analysis, RNA samples were amplified, labeled, and hybridized to the Human WG-6 (v. 3.0) BeadChip and the Human HT-12 (v. 3.0) BeadChip (Illumina) at the Australian Genome Research Facility (The Walter and Eliza Hall Institute of Medical Research). Data were analyzed using GenomeStudio Gene Expression Module (v. 1.5.4) (Illumina) using average normalization across all samples, with additional analysis performed using GeneSpring GX software (Agilent Technologies), as previously described (Goulburn et al., 2011).

IFN-γ Stimulation Assay

Human IFN-γ (R&D Systems) at 10 ng ml⁻¹ was added to day 30 thymic differentiation cultures in BEL medium. After 72 hr of IFN-γ stimulation, differentiated cells were harvested and analyzed for HLA-DR expression by flow cytometry.

CD34⁺CD7⁺ proT-Cell and FOXN1-GFP⁺ TEC Progenitor Coculture Assays

Unbilical cord blood (UCB) mononuclear cells were obtained and processed as previously described (La Motte-Mohs et al., 2005). They were pre-enriched into lineage-negative (Lin⁻) fractions, sorted into CD34⁺ hematopoietic stem cells, and cultured on OP9-DL1 cells for 9–10 days, as previously described (Awong et al., 2009). CD45⁻CD34⁺CD7⁺ proT-cells were FACs purified and either cultured alone (1 × 10⁴ cells/well) or with FOXN1-GFP⁺ (3.5 × 10³ cells/well) or GFP⁺ (3.5 × 10³ cells/well) cells in 96-well round-bottom low-attachment plates (Costar), containing APEL medium supplemented with 20% FBS, rhFLT3L (5 ng ml⁻¹, Peprotech), rhIL7 (5 ng ml⁻¹, Peprotech), and rhSCF (30 ng ml⁻¹, Peprotech). A half-media change was performed every 3–4 days. Cultures were analyzed by flow cytometry using the indicated antibodies and as detailed in the Supplemental Experimental Procedures.

ACCESSION NUMBERS

The GEO accession number for the microarray data reported in this paper is GSE56373.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, five figures, and two tables and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2014.04.009.

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FOXN1<sup>GFP/w</sup> Reporter hESCs Enable Identification of Integrin-β4, HLA-DR, and EpCAM as Markers of Human PSC-Derived FOXN1<sup>+</sup> Thymic Epithelial Progenitors

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Supplemental Experimental Procedures

**FOXN1 targeting vector construction:** The FOXN1 targeting vector was assembled using standard cloning techniques and Red/ET Recombination technology (Gene Bridges). DNA fragments representing the 5' and 3' homology arms were derived from bacterial artificial chromosome (BAC) RP11-915B21 spanning the human FOXN1 genomic locus (Roswell Park Cancer Institute). *Ascl* and *ClaI* restriction endonuclease sites were introduced during recombineering steps to facilitate subcloning of a *GFP-PGK-neo*R reporter/selection cassette into a pBR322-based targeting vector backbone, while *SwaI* restriction sites were introduced to allow linearization of the construct.

**Generation of FOXN1\(^{\text{GFP/w}}\) hESC Reporter lines:** \(10^7\) HES3 (Richards et al., 2002) or MEL1 (Millipore catalogue no. SCC020) hESCs were electroporated with 25µg of the FOXN1 targeting vector and subjected to G418 (Geneticin, Gibco) selection. Resultant G418 resistant colonies were screened for homologous recombination events using a PCR-based strategy incorporating the forward primer *Neo4* (5'-cagaccaatagatgttaga-3') and the reverse primer *FOXN1 26063r* (5'-gccctgactttgacctctgaacc-3') (*primer pair b, Figures 1A and 1B*). Six (of 134) MEL1 and three (of 144) HES3 correctly targeted clones were obtained. Homologous integration of the 5' homology arm was confirmed by PCR amplification using the forward primer *FOXN1 10546f* (5'-cattcccagaccccgagtttccttg-3') and the reverse primer *GFP Reverse2* (5'-ccggtgaacagctcctcgcccttgc-3') (*primer pair a, Figures 1A and 1B; Figure S1A*). The loxP-flanked neo\(^{R}\) cassette was removed by Cre recombinase-mediated excision as previously described (Davis et al., 2008). Sub-clones that had excised the neo\(^{R}\) gene were identified using the PCR primer *GFP1* (5'-gtgtgctgccgacaaccactac-3') and the reverse primer *FOXN1 26063r*, which generated a product that spanned the site formally occupied by the neo\(^{R}\) cassette (*primer pair c, Figures 1A and 1B; Figure S1A*). Correctly targeted clones that had excised the selectable marker were single cell cloned as previously described (Davis et al., 2008).

**Antibodies used in this study:** Primary antibodies used were mouse anti-human E-CADHERIN (Invitrogen), mouse anti-human SSEA-4 (Chemicon International), mouse anti-human TRA-1-60 (Chemicon International), mouse anti-human PDGFR\(\alpha\) (BD Pharmingen), mouse anti-human CD104 (BD Pharmingen), fluorescein isothiocyanate (FITC)-conjugated mouse anti-human CD9 (BD Pharmingen), phycoerythrin (PE)-conjugated mouse anti-human EpCAM (BD Biosciences), and allophycocyanin (APC)-conjugated mouse anti-human HLA-DR (BD Pharmingen). Unconjugated primary antibodies were detected with either APC- or PE-conjugated goat anti-mouse IgG (BD Biosciences).

**Intracellular Flow Cytometry:** Following dissociation and filtration, cells were fixed and permeabilized with BD Cytofix/Cytoperm (BD Biosciences) at 4°C for 20 min. Cells were washed and stained with primary antibody or an isotype control diluted in 1X BD Perm/Wash (BD Biosciences) at 4°C for 30 min. Unconjugated OCT4 and FOXN1 primary antibodies were detected with APC-conjugated goat anti-mouse IgG (BD Pharmingen). Cells were incubated with secondary antibodies at 4°C for 30 min in the dark. Antibody staining was quantified using the BD FACSCalibur. Forward and side scatter were used to identify cells that were viable immediately prior to the initial fixation process. Flow cytometric analysis was performed using Flowlogic Software (Inivai Technologies).
Quantitative Real-Time Polymerase Chain Reaction (q-PCR): Total RNA was prepared using High Pure RNA Isolation Kit (Roche), according to manufacturer’s instructions. First-strand cDNA was reverse-transcribed from RNA samples with random hexamer priming using SuperScript III reagents (Invitrogen), as previously described (Pick et al., 2007). Q-PCR was performed using TaqMan Gene Expression Assays with TaqMan Universal PCR Master Mix (Applied Biosystems) on the 7500 Fast Real Time PCR System (Applied Biosystems) under standard cycling parameters according to manufacturer’s instructions. The TaqMan Gene Expression Assays employed in this study were FOXN1 (Hs00186096_m1), GAPDH (Hs99999905_m1), FOXA2 (Hs00232764_m1), SOX17 (Hs00751752_s1), HOXA3 (Hs00601076_m1), PAX9 (Hs00196354_m1), FGFR2 (Hs01552926_m1), INVOLUCRIN (Hs00902520_m1), KERATIN-10 (Hs00166289_m1), KERATIN-5 (Hs00361185_m1), KERATIN-8 (Hs01670053_m1) and GCM2 (Hs00171702_m1). The comparative cycle threshold (Ct) method was used to analyze data, with gene expression levels normalized against input determined by reference to the GAPDH housekeeping gene, with an arbitrary multiplier of 10,000 for sample normalization, as previously published (Pick et al., 2007).

FOXN1-GFP⁺ TEC progenitor - CD34⁺CD7⁺ proT-cell co-culture assays: Sorted CD34⁺ cells from Lin⁻ UCB were cultured on OP9-DL1 cells for 9-10 days in OP9 Medium containing rhFLT3L (5 ng ml⁻¹, Peprotech), rhIL7 (5 ng ml⁻¹, Peprotech) and rhSCF (30 ng ml⁻¹, Peprotech) (Awong et al., 2009; La Motte-Mohs et al., 2005). FACS purified CD45⁻CD34⁺CD7⁺ proT-cells were either cultured alone (1 x 10³ cells/well) or with FOXN1-GFP⁺ (3.5 x 10³ cells/well) or GFP⁺ (3.5 x 10³ cells/well) cells in 96-well round-bottom low attachment plates (Costar) containing APEL Medium supplemented with 20% FBS, rhFLT3L (5 ng ml⁻¹), rhIL7 (5 ng ml⁻¹) and rhSCF (30 ng ml⁻¹). A half-media change was performed every 3-4 days. Differentiated proT-cells were analyzed for the expression of CD34, CD1a, CD5, CD4, CD8, CD3, CD45 and CD14 cell surface markers at weekly intervals. The following antibodies (all supplied by BD Pharmingen) were employed for analyses: APC-Cy7- and APC-conjugated mouse anti-human CD45, PE-Cy7- and APC-conjugated mouse anti-human CD34, PE-conjugated mouse anti-human CD7, PE-conjugated mouse anti-human CD1a, Biotin-conjugated mouse anti-human CD5, APC-conjugated mouse anti-human CD4, APC- and PE-conjugated mouse anti-human CD8, APC-conjugated mouse anti-human CD3 and PE-conjugated mouse anti-human CD14. The biotin-conjugated CD5 was detected with a Streptavidin-PE-conjugated secondary antibody.
References for Supplemental Material

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