The decision to develop into $\alpha\beta$ or $\gamma\delta$ T cells is pre-programmed in distinct subpopulations of DN1 thymocytes

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**Abstract**

T cells are divided into the αβ and γδ lineages. It is currently thought that these lineages differentiate in the thymus from uncommitted progenitor thymocytes. However, in this study we show that the decision to differentiate into one or either lineage is in fact already fixed in these apparent uncommitted progenitors. Using single-cell RNA sequencing, we reassemble *de novo* a model of early T cell development based on the transcriptional profiles of individual CD4⁻CD8⁻ double negative and γδ thymocytes. We show that the earliest thymocyte stage, known as CD4⁻CD8⁻ double negative 1 (DN1), is actually comprised of a mixture of transcriptionally distinct subpopulations. Although not yet expressing definitive markers of αβ and γδ lineages, such as the lineage-defining T cell receptors, specific DN subpopulations exhibit restricted developmental potential for either αβ or γδ lineage. Furthermore, specific γδ-primed DN1 subpopulations preferentially develop into IL-17 or IFNγ-producing γδ T cells. Thus, T cell lineage decisions are hardwired from the earliest stages of T cell development.
Introduction

T cells are divided into two principal lineages that are distinguished by the composition by their T cell receptors (TCRs) (Carpenter & Bosselut, 2010). Those that express an \( \alpha \beta \) TCR include conventional CD4\(^+\) and CD8\(^+\) T cells and these can be further divided into diverse subsets that have distinct immunologic functions (Harrington et al., 2005; Rothenberg, Moore, & Yui, 2008). T cells that express a \( \gamma \delta \) TCR are mainly divided into two subtypes defined by their secretion of IL-17A or IFN\(\gamma\). IL-17A-producing \( \gamma \delta \) T cells are important for immunity to intracellular bacteria, while IFN\(\gamma\)-producing \( \gamma \delta \) T cells are well-known for their anti-tumor function (Bai et al., 2017; Gao et al., 2016; Ribot, Lopes, & Silva-Santos, 2021).

T cell development occurs in the thymus, and it is thought that \( \alpha \beta \) and \( \gamma \delta \) T cells both arise from common progenitors that are seeded from the bone marrow (Ciofani & Zuniga-Pflucker, 2010; Taghon, Yui, Pant, Diamond, & Rothenberg, 2006). In mice, the earliest stages of T cell development are termed double-negative (DN) because they lack CD4 and CD8 expression. These are further subdivided into DN1 (CD44\(^+\)CD25\(^-\)), DN2 (CD44\(^+\)CD25\(^+\)), DN3 (CD44\(^-\)CD25\(^+\)) and DN4 (CD44CD25\(^-\)) stages. \( \alpha \beta \) and \( \gamma \delta \) development is thought to bifurcate at the DN2b>DN3 transition, when TCRB/G/D gene rearrangement occurs to generate a functional pre-TCR or \( \gamma \delta \) TCR (Ciofani, Knowles, Wiest, von Boehmer, & Zuniga-Pflucker, 2006; Fehling, Krotkova, Saint-Ruf, & von Boehmer, 1995; Kreslavsky, Gleimer, Garbe, & von Boehmer, 2010; Oh, Gray, & Chong, 2021). Evidence also suggests that effector outcomes of \( \gamma \delta \) T cells may also be determined in the thymus rather than in the periphery because the expression of IL-17A and IFN\(\gamma\) by mature cells correlates with \( V\gamma \) chain usage. \( V\gamma 2^+ \gamma \delta \) T cells tend to produce IL-17A while \( V\gamma 1^+ \gamma \delta \) T cells tend to produce IFN\(\gamma\) (Fahl, Coffey, & Wiest, 2014; Narayan et al., 2012).

The prevailing model suggests that TCR signaling plays a deterministic role in lineage commitment and subsequent \( \gamma \delta \) effector differentiation (Munoz-Ruiz, Sumaria, Pennington, & Silva-Santos, 2017). Strong signals have been shown to activate ERK and induce EGR-ID3 and this appears to drive \( \gamma \delta \) differentiation, whereas weak signals appear to promote an \( \alpha \beta \) fate (Hayes, Li, & Love, 2005; Lauritsen et al., 2009; Munoz-Ruiz et al., 2017).
Additionally, weak TCR signaling may promote an IL-17A phenotype in γδ T cells, while strong signaling promotes an IFNγ phenotype (Sumaria, Grandjean, Silva-Santos, & Pennington, 2017).

However, the role of the TCR remains debatable because a lack of Notch signaling at the DN2 stage compromises progression to DN3 and impairs αβ development but not γδ development (Tanigaki et al., 2004; Wolfer, Wilson, Nemir, MacDonald, & Radtke, 2002). How can the stage prior to the apparent γδ branching be compromised and yet γδ T cell development still proceed? Evidence suggest that this Notch-independence is at least partially due to the transcription factor ID3 because a weak TCR signal in cooperation with Notch signaling is sufficient to inhibit E-protein activity (Lauritsen et al., 2009). However, ID3 activity could explain the development of only a subset of γδ T cells. This murkiness surrounding the αβ versus γδ decision has arisen partly due to a lack of markers to define the development of thymocytes that have been selected to mature into γδ T cells (Pellicci, Koay, & Berzins, 2020). Together, these ambiguities raise the question as to whether the early stages of T cell development are really bipotent homogenous populations that can give rise to both αβ and γδ T cells. Emerging evidence in other developmental pathways suggests that progenitors can be heterogeneous with biases in lineage potential (Naik et al., 2013). In this study, we perform single-cell RNA sequencing (scRNA-seq) of early thymocytes from mouse thymus to determine cellular heterogeneity at single-cell resolution and to assemble a model of T cell development de novo in order to address when the αβ versus γδ lineage decision is actually made.

**Results**

**Differences in the kinetics of αβ versus γδ T cell development from DN1 thymocytes**

To better understand the development of the γδ lineage, we began by investigating the kinetics of αβ versus γδ T cell development in OP9-DL1 cultures, which supports the development of hematopoietic progenitors *in vitro* into either γδ T cells or αβ cells up to the CD4+CD8+ double positive (DP) stage (Holmes & Zuniga-Pflucker, 2009). Total DN1 (CD44+CD25−Lin−) and DN2 (CD44+CD25+Lin−) thymocytes were sorted from C57BL/6 mice.
and cocultured on OP9-DL1 cells. The cultures were then analyzed after 9-23d. DP cells were detected by CD8α (and CD4) expression while γδ cells were detected by TCRγδ expression (Supplementary Fig. 1a, b).

While DN1 and DN2 thymocytes gave rise to both lineages, surprisingly, there was a significant difference in the kinetics in the development of DN1 thymocytes into γδ cells compared to αβ cells. Starting from DN1 thymocytes, γδ cells were already present at 9d, with the percentage and cell number peaking at 12d, waning and then increasing again at 20d (Fig. 1a, b). In contrast, αβ cells only started appearing after 14d and peaked at 20d. The earlier γδ peak suggests that either γδ development is either simply more rapid than αβ development or that there may be different developmental pathways for the two lineages, with γδ cells potentially produced in two waves.

Unlike the discordant kinetics of the DN1 cultures, culturing DN2 thymocytes produced both αβ and γδ lineage cells from 9d and in similar proportions throughout (Fig. 1a, b). γδ cells were produced at a consistently low frequency of 2-6%, while between one-third to half the cells were αβ cells at each timepoint. Given that DN2 thymocytes are only one stage along from DN1, the consistent rate of both αβ and γδ production argues against γδ development being simply more rapid.

If the αβ and γδ lineages develop independently, we might expect to see each lineage being derived from distinct DN1 thymocytes. To investigate this, we tagged individual DN1 thymocytes with unique genetic heritable barcodes (Naik et al., 2013) and tracked the propagation of these barcodes in OP9-DL1 cultures. If an αβ cell and γδ cell inherits the same barcode sequences, it means that they were derived from the same progenitor. CD8α+ and TCRγδ+ cells were sorted after 14d and 20d of culture and analyzed for barcode composition (Supplementary Fig. 2a). While there was some overlap in barcode usage between the four populations, the barcode profiles were largely non-overlapping, with Pearson’s correlations of r<0.6 for the first experiment (Fig. 1d, e) and even lower in the second experiment (Supplementary Fig. 2c, d). Together with the kinetic analysis, this suggest that the DN1
thymocytes may in fact be a heterogeneous mixture consisting of distinct subpopulations that developed into either $\alpha\beta$ cells or $\gamma\delta$ cells.

**Fig 1.** Potential heterogeneity in the lineage outcomes of DN1 thymocytes. (a) Total DN1 (CD44+CD25−Lin−) and DN2 (CD44+CD25+Lin−) thymocytes were sorted from the thymus of
C57BL/6 mice then analyzed for $\alpha\beta$ versus $\gamma\delta$ lineage differentiation in OP9-DL1 cultures. The cultures were analyzed at the indicated time points by flow cytometry. $\alpha\beta$ lineage cells were identified as CD8$\alpha^{+}$ while $\gamma\delta$ lineage cells were identified as TCR$\gamma\delta^{+}$. CD4 expression was also analyzed but was concomitant with CD8$\alpha$ expression (not shown). Lin = CD4, CD8, B220, CD11b and CD11c. Representative flow cytometric plots are shown. (b) The mean percentages ± S.E.M. of 4 to 7 replicates are shown. (c) The mean cell numbers ± S.E.M. are shown. (d, e) Barcoded total DN1 thymocytes were differentiated on OP9-DL1 cells. $\alpha\beta$ and $\gamma\delta$ cells were then sorted from the culture at 14d and remaining culture was sorted at 20d. The barcode composition of the four populations was then analyzed. Heatmaps are comparing by individual barcodes (rows) across the four populations the (column) are shown. Heatmap with normalization of barcodes within each population is shown in (d), while normalization of each barcode usage across all four populations is shown in (e). Indicated under (d) are Pearson’s correlation ($r$) analysis between each pair of populations. Shown is one replicate of two independent experiments. The second replicate is shown in Figure S2C-D and the corresponding t-SNE clustering to group the barcodes information is shown in the Supplementary Figure 2b.

A comprehensive transcriptional map of early T cell development at single-cell resolution

To better understand the potential heterogeneity of the early stages of T cell development, we next characterized the transcriptional landscape of DN and $\gamma\delta$ thymocytes at single-cell resolution. Cells were sorted from the thymus of C57BL/6 mice for analysis by 10X scRNA-seq over three independent runs (Fig. 2a, b and Supplementary Table 1). The first run consisted of total DN and TCR$\gamma\delta^{+}$ cells, the second consisted only of DN1 and DN2 cells and the third involved sorting DN1+2, DN3 and TCR$\gamma\delta^{+}$ cells separately and mixing back together post-sort at a ratio of 55% to 30% to 15%, respectively (Fig. 2a and Supplementary Table 1). This was to ensure that a sufficient number of DN1 and DN2 cells were captured for high-resolution analysis that would not have been achieved by analyzing total DN cells.
Fig. 2. Single-cell RNA sequencing analysis of early T cell development. (a) Shown are the gating strategies used to sort DN and gd thymocytes from C57BL/6 mice for 10X scRNA-seq. Three separate runs were completed. The first run was total DN and TCRγδ+ cells. The second run consisted
only of DN1 and DN2 cells. The third run involved sorting DN1 plus DN2, DN3 and TCRγδ+ cells separately, which were then mixed back together post-sort at a ratio of 55% to 30% to 15% respectively. Dump = CD4, CD8, B220, CD11b, CD11c, TCRβ and TCRγδ. (b) Following processing of the 10X data on CellRanger, each dataset was analyzed for clustering based on the first 12 principal components in Seurat. Shown is the t-SNE plot of each run color-coded to DN developmental stage, TCRγδ+ thymocytes or other (non-thymocytes). (c) The three datasets, totaling 22,094 cells, were integrated with SCTransform, then the cells were clustered with Seurat. This yielded 30 distinct clusters (left plot), which were then annotated to DN developmental stage, TCRγδ+ thymocytes or other (non-thymocytes) (right plot). (d) Violin plots showing expression of selected markers used to assign individual clusters to DN development stage.

A total of 22,094 high quality cells passed quality control checks across the three 10X datasets. The three datasets were first integrated to assemble a global view of early T cell development. To recover biological distinction from these different replicates and minimize batch-associated variability, the pooled data was normalized using SCTransform. Following dimensional reduction, unsupervised clustering was performed using the first 12 principal components (PC). This identified 30 distinct clusters (Fig. 2c), which were assigned to a DN stage or mature γδ thymocytes based on the expression of canonical markers (Supplementary Fig. 1a, b). High Cd44 and Il7r expression but low T lineage gene expression, including Il2ra, Tcf7, Cd24a, Notch1 and Bcl11b, identified DN1 cells (Fig. 2d). DN2 cells were identified by upregulation of T lineage genes and downregulation of Il7r. DN3 cells were identified by Ptcra and further upregulation of T lineage genes. Low Cd8b1 and loss of Il2ra distinguished DN4 from DN3 cells (Fig. 2d). High Trdc, Id3, Sox13 and Rorc identified mature γδ thymocytes (Fig. 2d). This analysis revealed that multiple distinct populations correspond to each of the canonical DN stages and suggests that DN thymocytes are much more heterogenous than previously thought.

We also tested the effect of cell cycle gene expression on the clustering. Cell cycling scores were calculated from the integrated data and regressed using Seurat’s built-in regression model (Supplementary Fig. 3a, b). There was
a slight variation in the number of output clusters, but we did not observe any biological variability that could be
accounted for by cell cycle status, with the genetic profiles nearly identical between output with cell cycle genes
left in or regressed out (Supplementary Fig. 3c). Therefore, exclusion of cell cycle genes was not necessary for
downstream analyses.

**Trajectory analysis suggests that αβ and γδ T cell development occurs in parallel**

To understand how the single cell transcriptomic map of DN and γδ thymocytes aligned with the established
understanding of early T cell development, we reconstructed a developmental pathway *de novo* from the identified
clusters by ordering the cells based on tracking gene expression in pseudotime analysis.

Pseudotime inference of the integrated data with Slingshot (Street et al., 2018) ordered the cells in the expected
sequence from DN1 through to DN4, corresponding to αβ development (Germain, 2002). There were also two
branch points predicted, one corresponding to the γδ lineage branching from DN1 cells and another branch point
at DN2 and ending with a subpopulation of DN3 cells (Fig. 3a).

We also performed pseudotime inference using Monocle 2 (Qiu et al., 2017). This assembled the cells along an
asymmetric trajectory that divided into six states (Fig. 3b), which were then analyzed for signature gene
expression (Supplementary Fig. 4) to assign the cells to the conventional DN development stages (Fig. 3c, d).
State 2 comprising DN1 and some DN2a cells, was identified as the starting point. State 3 corresponded to the
main αβ pathway, with DN4 as the endpoint. States 5 and 6 terminated with DN3 and corresponded to β-selection
failure (Fig. 3b, d). State 1 corresponded to the γδ branch, and like Slingshot, Monocle 2 inferred that γδ cells
develop directly from DN1. Interestingly, while *Rag* expression was highest in DN3 cells, which is expected,
expression could already be detected in some DN1 cells (Fig. 3e, f). Thus, trajectory analyses suggest that the γδ
lineage decision may be much earlier than the current paradigm of branching at DN2b›DN3 (Supplementary Fig.
1).
Fig. 3. Trajectory analyses predict αβ and γδ T cell development occurs in parallel from the DN1 stage. (a) Pseudotime analysis of the integrated 10X dataset with Slingshot characterizing the course of development from DN1 to DN4 (αβ lineage) and γδ T cells, visualized by UMAP. Cells are color-coded by developmental stage. (b) Pseudotime analysis of total DN and γδ thymocytes (first 10X run) with Monocle 2. The six states identified in the asymmetric trajectory are indicated. (c) Color-coding of the Monocle 2 analysis by thymocyte development stage based on expression of
marker genes by the individual clusters. (d) Shown are the proportions of DN developmental stage or TCRγδ+ cells that contributed to each state of the Monocle 2 analysis. (e) Expression of Rag1 and Rag2 overlaid onto the Monocle 2 analysis. (f) Expression of Rag1 and Rag2 overlaid onto the Slingshot analysis.

**Extensive heterogeneity within DN1 and DN2 thymocytes**

The trajectory analyses, as well as the OP9-DL1 kinetic studies, predicted that γδ T cell development might be occurring in parallel with αβ development, rather than branching off at the later DN2b>DN3 point. To better delineate the earlier stages of the developmental model, we analyzed the second 10X run that was performed specifically on DN1 and DN2 thymocytes. Unsupervised clustering of the 8,851 cells that pass quality control checks identified 26 distinct subpopulations (Fig. 4a). Analysis for marker genes identified eight DN1, five DN2a and 10 DN2b subpopulations (Fig. 4a). There were also three small clusters of non-T cells consisting of doublets and B cells that, for simplicity, were excluded from downstream analysis. DN1s expressed high levels of progenitor markers, including Hhex, Il7r and Cd44 but low levels of T-commitment markers, such as Il2ra, Tcf7, Notch1, Cd24a and Myb (Supplementary Fig. 5a, b). Late T-lineage commitment genes, including Rag1, Rag2, Notch3 and Ptcra, were used to separate DN2a and DN2b subpopulations (Supplementary Fig. 5a, b). While heterogeneity in DN1 thymocytes was reported previously (Porritt et al., 2004), the degree of heterogeneity among both DN1 and DN2 thymocytes was far more extensive than expected, which could be clearly seen by the differential expression of many marker genes (Supplementary Fig. 5b).

**Identification of novel cell surface markers for delineating DN1 and DN2 subpopulations**

In order to study these novel subpopulations of DN1 and DN2 thymocytes, we needed to identify useful cell surface markers that could be used for flow cytometry. Differential expression analysis was performed to select features (p<0.05 and 2-fold difference) that were cross-referenced to GO terms for cell surface proteins (Supplementary Fig. 6a, b). We then tested commercial antibodies against these candidates. While not all
antibodies produced a staining pattern consistent with mRNA expression levels, we were able to identify a minimal panel to delineate both DN1 and DN2 thymocytes.
Fig. 4. Identification of cell surface markers for delineating DN1 and DN2 the subpopulations inferred from scRNA-seq. (a) t-SNE visualization of 8,851 DN1 and DN2 thymocytes from the second 10X run clustered with Seurat. This yielded 26 distinct clusters (left plot), which were then annotated as DN1a, DN1b, DN1c, DN1d, DN1e, DN2a or DN2b (right plot). (b) Genes found to be differentially expressed between DN1 subpopulations were annotated against Gene Ontology (GO) terms for cell surface markers (Figure S6). Shown is a heatmap showing the average expression of the genes encoding cell surface markers selected to delineate the DN1 subpopulations. (c) Flow cytometric validation of the selected markers used to identify DN1 subpopulations. Total DN1 cells (CD4^−CD8^−CD44^+CD25^−Lin^−) were first divided into five populations based on c-Kit and CD24. The c-Kit^−CD24^+ cells (total DN1e) was then divided into three populations based on Ly6a and Bst2. Finally, the Ly6a^−Bst2^− cells were divided into two populations based on Klrk1. Lin = B220, CD11b, CD11c, TCRβ and TCRγδ. (d) Heatmap showing the average expression of genes encoding cell surface proteins selected to delineate the DN2 subpopulations. (e) Flow cytometric validation of the selected markers used to identify DN2 subpopulations. Total DN2 cells (CD4^−CD8^−CD44^+CD25^+Lin^−) were first divided into two populations based on c-Kit and Thy1. The c-Kit^+Thy1^int cells (total DN2a) were divided into three populations based on Ly6d and CD53. Then, the c-Kit^+Thy1^intLy6d^intCD53^− cells were divided into two populations based on CD53 and CD3e. The c-Kit^−Thy1^+ cells (total DN2b) were divided into three populations based on Ly6d and CD53. Then, the c-Kit^+Thy1^intLy6d^intCD53^int/+ cells were divided into five populations based on CD53 and CD3e.

The combination of five markers encoded by the genes *Cd24a, Kit, Bst2, Ly6a* and *Klrk1* distinguished the eight DN1 subpopulations (Fig. 4b). CD24 and c-Kit was previously shown to separate DN1 thymocytes into five subpopulations, known as DN1a, DN1b, DN1c, DN1d and DN1e(Porritt et al., 2004). DN1a to DN1d each corresponded to a single cluster (Fig. 4a). Bst2, Ly6a and Klrk1 were then used to delineate DN1e-1 to DN1e-4 (Fig. 4c).
The pair of DN2a clusters 7/15 could not be separated by cell surface markers as well as the pairs of DN2b clusters 4/23, 9/13 and 8/19 due to very similar gene expression profiles. Thus, these cluster pairs were combined. The resulting 11 DN2 clusters were annotated as DN2a-1 to 4 and DN2b-1 to 7 based on the expression of genes encoded by *Thy1, Kit, Cd53, Ly6d* and *Cd3e* (Fig. 4d, e).

**Distinct DN1 subpopulations are primed to develop into either αβ or γδ lineage**

The heterogeneity of the DN1 thymocytes could potentially represent different progenitors of αβ and γδ lineages, which would explain the early branch point inferred by pseudotime trajectory analyses (Fig. 3a-c). To test this, we sorted the DN1 and DN2 subpopulations using our antibody panels and accessed their αβ versus γδ lineage outcomes in OP9-DL1 cultures after 14d and 20d (Fig. 5a-f).

At 14d, we observed strong αβ lineage restriction in DN1a, DN1b and DN1c cultures, whereas DN1d and DN1e cultures only generated TCRγδ+ cells (Fig. 5a-c and Supplementary Table 2). At 20d, there was significant increase in the percentage and number of αβ cells produced from DN1a and DN1b, while DN1c had given rise to some γδ cells as well as αβ cells (Fig. 5a-c and Supplementary Table 3). DN1d and DN1e subpopulations continued to exhibit γδ restriction, with DN1e-1 and DN1e-2 producing the highest percentage and number of γδ T cell among DN1 subpopulations (Fig. 5a-c and Supplementary Table 2).

At 14d, all DN2b subpopulations and DN2a-1 produced a high percentage and number of αβ cells and few TCRγδ+ cells, while the rest of DN2a subpopulations produced mostly TCRγδ+ cells and few αβ lineage cells (Fig. 5d, e, Supplementary Fig. 7 and Supplementary Table 4). By day 20, all DN2 subpopulations had produced αβ lineage cells, while TCRγδ+ cells had been lost from the DN2a cultures (Fig. 5d, e, Supplementary Fig. 7 and Supplementary Table 5). There was also a dramatic reduction in cell numbers in the DN2b cultures, which may be a result of the cells undergoing cell death after reaching the DP stage (Fig. 5e).
Fig. 5. Restricted αβ or γδ potential of DN1 subpopulations. (a) Sorted DN1 subpopulations were cultured on OP9-DL1 cells to assess their lineage potential. The cultures were analyzed at 14d and 20d of culture by flow cytometry. αβ lineage cells were identified as CD8α+ while γδ lineage cells were identified as TCRγδ+. Representative flow cytometric plots are shown. (b, c) Pooled data analyzing αβ versus γδ differentiation from sorted DN1 subpopulations. The means ± S.E.M. of four
to nine replicates performed over four independent experiments are shown. See Table S2 and S3 for
$P$-value calculations. (d, e) Pooled data analyzing $\alpha\beta$ versus $\gamma\delta$ differentiation from sorted DN2
subpopulations. The means $\pm$ S.E.M. of four to nine replicates performed over four independent
experiments are shown. See Supplementary Table 4 and 5 for $P$-value calculations. (f) Repopulation
of dGuo-depleted fetal FTOCs with select DN1 subpopulations. The lobes were then analyzed by
flow cytometry for CD8$\alpha$ versus TCR$\gamma\delta$ expression after 14d. Shown is a representative from three
independent experiments.

We also tested the differentiation of select DN1 subpopulations in fetal thymic organ cultures (FTOCs). Like in
the OP9-DL1 co-cultures, DN1b cells only produced $\alpha\beta$ cells, DN1c cells primary produced $\alpha\beta$ cells and some
$\gamma\delta$ cells, while DN1d and DN1e-4 cells were entirely restricted to the $\gamma\delta$ lineage (Fig. 5f). Thus, both OP9-DL1
and FTOC systems support the model that the decision to develop into $\alpha\beta$ versus $\gamma\delta$ lineages is largely hardwired
from the DN1 stage rather than a decision made later in T cell development. However, there remains some
flexibility because small numbers of TCR$\gamma\delta^+$ cells can also differentiate from DN1c and DN2a subpopulations.

**Gene expression analysis of DN1 subpopulations suggest potential relationships with distinct mature $\gamma\delta$
subsets**

In our initial clustering analyses of the DN and $\gamma\delta$ thymocyte scRNA-seq data, $\gamma\delta$ thymocytes were found to
segregate into two distinct but closely related clusters (Fig. 2c). We therefore wanted to determine how these
related to the different DN1 subpopulations. Firstly, differential expression analysis revealed substantial
differences between two $\gamma\delta$ clusters, with 93 genes expressed at significantly higher levels by the $\gamma\delta$-1 cluster
while 117 genes were expressed at significantly higher levels by $\gamma\delta$-2 cluster (Fig. 6a). Genes that were highly
expressed by the $\gamma\delta$-1 cluster included *Gzma, Blk, Maf, Sox13, Etv5, Gata3, Ccr9, Rorc, Sox4, Tcf12, Lgals9,*
*Cmak4* and *Bcl11b*, which are all associated with IL-17A-producing $\gamma\delta$ T cell subset (Sagar et al., 2020). This
suggests that the $\gamma\delta$-1 cluster is likely to be $\gamma\delta$ thymocytes that go on to mature into the IL-17-producing subset,
while the $\gamma\delta$-2 cluster are those that are likely to mature into the IFN$\gamma$ producing $\gamma\delta$ T cell subset. This is consistent with a previous study that suggested the eventual effector function of $\gamma\delta$ T cells, may already be acquired in the thymus (Sumaria et al., 2017). The unique transcriptional profiles of $\gamma\delta$-1 and $\gamma\delta$-2 thymocytes appears to be evidence of this.

**Fig. 6.** The gene expression profiles of the different DN1 subpopulations correlate with distinct $\gamma\delta$ effector subsets. (a) Heatmap showing genes differentially expressed ($P$-value <0.05 and 2-fold difference) between the $\gamma\delta$-1 and $\gamma\delta$-2 thymocyte subpopulations identified in the cluster analysis (clusters 11 and 19) of DN and $\gamma\delta$ thymocyte scRNA-seq data from Figure 2c. Each column is an individual cell in the dataset while each row is a differentially expressed gene. Genes associated with either IL-17A (left side) or IFN$\gamma$ production (right side) are indicated. (b) Analysis of DN1 subpopulations for expression of the 210 genes differentially expressed between the two $\gamma\delta$ thymocyte populations. Transcription factors associated with either IL-17A or IFN$\gamma$ are indicated.
Next, to investigate the relationship between the mature $\gamma\delta$ thymocyte subpopulations and the DN1 subpopulations that differentiate into TCR$\gamma\delta^+$ cells, the DN1c, d and e subpopulations were analyzed for the expression of the 210 differentially genes that distinguished the two mature $\gamma\delta$ thymocyte populations. This revealed a similar transcriptional profile between DN1d and $\gamma\delta$-1 thymocytes, while DN1c and all the DN1e subpopulations overlapped significantly with $\gamma\delta$-2 thymocytes (Fig. 6b). However, there were also clearly differences between the DN1e subpopulations.

We then focused on transcription factors because they are regulators of gene expression and could potentially play key roles in the hardwiring $\gamma\delta$ effector outcomes in DN1 thymocytes. $Sox13$ was highly expressed by DN1d cells (Fig. 6b), which was previously shown to be important for the differentiation of a subset of DN1 thymocytes into T$\gamma\delta$17 cells (Spidale et al., 2018). Interestingly, DN1e subpopulations also express transcription factors associated with IL-17-producing $\gamma\delta$ T cells. Notably, DN1e-1 and DN1e-2 cells expressed high levels of $Maf$, while $Gata3$ was highly expressed by both DN1e-1 and DN1d cells (Fig. 6b). We thus predict that $\gamma\delta$ effector outcomes could be pre-wired from the DN1 stage, with DN1d going on to develop into IL-17-producing cells and DN1e potentially producing both $\gamma\delta$ effector subsets.

**Hardwiring of $\gamma\delta$ subset outcomes in DN1 subpopulations.**

To determine if different DN1 subpopulations give rise to different $\gamma\delta$ T cell subsets, the TCR$\gamma\delta^+$ cells that developed from the different DN1 subpopulations in OP9-DL1 co-cultures were analyzed for effector phenotype by staining for intracellular IL-17A and IFN$\gamma$ expression and for expression of specific $V\gamma$ chains (Fig. 7a). Cytokine production is highly associated with specific $V\gamma$ chain usage, with $V\gamma1^+$ cells enriched for IFN$\gamma$ and $V\gamma2^+$ cells enriched for IL-17A production (Narayan et al., 2012; O'Brien & Born, 2010).

DN1c thymocytes generated both $V\gamma1^+$ and $V\gamma2^+$ cells, with only a low percentage of IFN$\gamma$ expression by the $V\gamma1^+$ cells (Fig. 7b-d and Supplementary Table 6). DN1d primarily produced $V\gamma2^+$ cells that secreted IL-17A.
Similarly, DN1e-1 and DN1e-2 were restricted to a Vγ2+ IL-17A+γδ. On the other hand, DN1e-3 and DN1e-4 exhibited plasticity and could generate both IL-17A and IFNγ producing cells (Fig. 7b-d and Supplementary Table 6). This strongly suggests that not only is the αβ versus γδ lineage decision largely determined by the DN1 stage in T cell development, but even γδ effector outcomes are already hardwired.
Fig. 7. Restriction of γδ effector subset outcomes in different DN1 subpopulations. (a) Gating strategy for analyzing the phenotype of γδ T cells generated from DN1 subpopulations after co-culturing with OP9-DL1 cells for 14 days. TCRγδ+ (CD4−CD8−TCRβ+) were first divided based on Vγ1 versus Vγ2 expression. The three subpopulations, including Vγ1−Vγ2− double negative (DN) cells were then analyzed for intracellular IL-17A and IFNγ expression. (b) Shown are the flow cytometric plots from a representative of three independent experiments for cytokine expression by γδ T cells derived from DN1 subpopulations. The top row shows the Vγ1 versus Vγ2 expression. Gated Vγ1+, Vγ2+ and double negative (DN) cells were then analyzed for IL-17A versus IFNγ expression in the bottom three rows. (c) Pooled data analyzing the percentage of Vγ1 versus Vγ2 cells differentiated from sorted DN1 subpopulations. The means ± S.E.M of four to six replicates performed over three independent experiments is shown. See Supplementary Table 6 for P-value calculations. (d) Pooled data analyzing percentage of Vγ1+IFNγ+ (left) and Vγ2+IL-17A+ (right) cells out of total TCRγδ+ cells. The means ± S.E.M is shown (*P<0.05, **P<0.01).

Discussion

A precise understanding of the earliest events in T cell development has until now been complicated by two factors. First, it has been unclear if the earliest thymocytes constitute a truly bipotent homogenous population that can differentiate into both αβ and γδ lineages. Second, previous studies that analyzed whole populations by flow cytometry largely lacked the resolution to reveal the asynchronous development of rare subpopulations. Leveraging the power of scRNA-seq for rare cell type identification, our study has revealed the extensive heterogeneity within the early stages of T cell development. Significantly, we show that the αβ versus γδ lineage decision is already hardwired within DN1 stage. This earliest stage of T cell development in the thymus, instead of being an uncommitted T cell progenitor population, in fact consists of distinct subpopulations that primed to develop into αβ T cells or γδ T cells. This is well before the expression of the lineage-defining TCRs.
The concept that the early stages of T cell development may consist of distinct subpopulation has previously been considered. It was shown that DN1 thymocytes can be subdivided into five subpopulations (DN1a-e) based on CD24 and c-Kit expression (Porritt et al., 2004), while neonatal IL-17-producing cells were shown to develop from a subset of DN1d thymocytes that is determined by a cell intrinsic program prior to TCR signaling (Spidale et al., 2018). Despite these findings, the cellular heterogeneity of DN1e thymocytes and their lineage specification remained unclear. We have now defined eight DN1 subpopulations, including four novel DN1e subpopulations and showed that DN1a and DN1b are restricted to αβ lineage whereas DN1d and DN1e subpopulations are restricted to γδ lineage. Therefore, lineage commitment is already determined by the DN1 stage, which is well before the apparent lineage bifurcation at the DN2b>3a stage.

Unlike αβ T cells, γδ T cells are thought to acquire their effector potential in the thymus rather than upon antigen exposure in secondary lymphoid organs (Ribot et al., 2009). Indeed, a previous study showed that some IL-17-producing γδ T cells develop from a distinct subset of DN1 thymocytes (Spidale et al., 2018). Although none of the DN1 subpopulations expressed any definitive markers of specific mature T cell developments, we observed substantial transcriptomic overlap between the IL-17-primed subpopulations with mature IL-17-expressing γδ thymocytes and between the IFNγ-primed subpopulations with mature IFNγ-expressing γδ thymocytes. Notably, we observed expression of many important transcription factors. Thus, critical components of the IL-17 or IFNγ γδ T cell transcriptional programs are already in place within distinct DN1 subpopulations.

We showed that DN1d thymocytes express many of transcription factors that are expressed by mature IL-17-producing γδ thymocytes but not IFNγ-producing thymocytes like Bcl11b, Etv5, Sox13, Rcf7, Rorc and Maf. Sox13 has previously be shown to be an important lineage determining factor for the neonatal IL-17A producing cells (Spidale et al., 2018). It is thought to act in concert with other transcription factors like Bcl11b, Tcf7, Rorc and Maf to specify the IL-17A-effector program in γδ T cells (Malhotra et al., 2013; Spidale et al., 2018).
We also showed that DN1e-1 and DN1e-2 are also restricted to Vγ2^+ IL-17A- producing γδ T cell fate, while only DN1e-3 and DN1e-4 cell display the plasticity to also develop into Vγ1^+ IFNγ^+ cells. This plasticity is likely to involve the differential expression of transcription factors that contribute to distinct effector fates. Although all four DN1e subpopulations expressed Stat1, which regulates IFNγ stimulated genes (Qiao et al., 2013), DN1e-1 and DN1e-2 also expressed Maf. cMaf is known to positively regulate IL-17A- producing γδ T cell development (Zuberbuehler et al., 2019), whereas a lack of cMaf expression by γδ T cells correlates with increased IFNγ expression (Gabrysova et al., 2018; Iwata et al., 2017; Zuberbuehler et al., 2019). Furthermore DN1e-1 cells were found to express Gata3, an important regulator of IFNγ expression (Yagi et al., 2010). The interplay between transcription factors may thus be key to the lineage decisions of DN1 subpopulations. Further analysis of chromatin states and epigenetic mechanisms associated with these transcription factors will likely to be valuable to reveal the intricacies of the regulatory cascades driving the different γδ effector outcomes.

While it is clear that the majority of DN1 thymocytes exhibited a restricted lineage outcome, the fact that DN1e-3 and DN1e-4 can differentiate into both IL-17A and IFNγ-producing cells, and that DN1c and DN2a still displayed some γδ potential in addition to their dominant αβ outcome, means that we cannot entirely rule out that at least some decisions could be made at the DN2b>3 stage. While largely hardwired, there remains some lineage flexibility and there must be other pathways that can contribute to the αβ versus γδ lineage decision. TCR signaling may have a role in these remaining undecided precursors. Moreover, appropriate TCR signaling may still be required for maturation of early progenitors even if they are already hardwired to develop into either αβ or γδ cells.

Current models for the αβ versus γδ lineage decision frequently center on the role of TCR signals at the DN2b>3 stage. The evidence for a role of differing TCR signals comes from studies that have employed genetic mouse models to manipulate TCR signaling at the DN3 stage. This revealed that strong signals activate the ERK-EGR-ID3 pathway to drive γδ T cell differentiation, whereas a weak signal promotes adoption of the αβ fate (Hayes et
TCR\(\gamma\delta\) signaling is also thought to play a role in subsequent \(\gamma\delta\) T cell effector differentiation (Munoz-Ruiz et al., 2017). Weak TCR signaling appears to promote an IL-17A effector phenotype, while strong signaling promotes IFN\(\gamma\) phenotype (Sumaria et al., 2017). However, it is still unclear whether TCR signal strength is dependent on instructive extrinsic signals or simply a result of stochastic selection of TCR chains. The instructive model proposes that TCR\(\gamma\delta\) signal competes with pre-TCR signal and the lineage decision is determined by cell specific interactions that activate key transcription factors, which in turn instructs gene expression program (Carpenter & Bosselut, 2010). In contrast, the stochastic model suggests that there is random rearrangement of TCR genes, where thymocytes that successfully assemble a \(\beta\) chain progress along the \(\alpha\beta\) lineage, while those that assemble specific \(V_\gamma\) chains differentiate into a specific subset of \(\gamma\delta\) T cells following appropriate TCR signaling. It is also possible that there may a component of both instruction and random selection to these decisions. Further analysis of whether TCR signaling occurs in a deterministic manner or in a random manner will be important to provide mechanistic insight into lineage commitment and subsequent \(\gamma\delta\) effector differentiation of those DN1c and DN2a. Moreover, as mentioned, appropriate TCR signaling may still be important for maturation of the \(\gamma\delta\)-restricted DN1 subpopulations, even if lineage outcome has already been determined.

Our novel finding of highly heterogeneous DN1 and DN2 subpopulations highlights the concept that progenitor populations may actually not be multi/bipotent cells at all that are. Progenitors may in fact be a grouping of lineage-primed cells at a similarly early stage in differentiation. On the surface, they may lack lineage defining markers and they are similar enough to appear to constitute a population, but individually, these cells have unique transcriptional programs that already specify a lineage-fate. Such heterogeneity in progenitor populations have been recently describe in the neural cortex (Johnson et al., 2015), prostate epithelium (Crowley et al., 2020) and in the heart (Churko et al., 2018). Within the hematopoietic system, lymphoid-primed multipotent progenitors in the bone marrow also appear to exhibit substantial restriction to specific lineage fates (Naik et al., 2013). More specifically, myeloid progenitors can be divided into subpopulations that are primed towards neutrophil, basophil, eosinophil, monocyte, dendritic cell, erythrocyte or megakaryocyte fate (Paul et al., 2016).
In conclusion, we have made the novel discovery that the $\alpha\beta$ versus $\gamma\delta$ T lineage decision, as well as the effector outcomes of $\gamma\delta$ T cells, are already hardwired in DN1 thymocytes. Such hardwiring in apparently uncommitted progenitor and prior to the apparent lineage decision, at DN2b>3 in the case of T cell development, could potentially be a common feature of many cellular developmental pathways. It will thus be of interest to perform a similar analysis of other immune developmental pathways, including other lymphocytes, in the future.

**Materials and Methods**

**Mice and thymocyte preparations**

Thymuses was harvested from C57BL/6 mice at around 6 weeks of age. All experiments were approved by the St Vincent’s Hospital Animal Ethics Committee and performed under the Australian code for the care and use of animals for scientific purposes.

Total thymocytes were obtained by crushing the thymus through a metal sieve to generate a single cell suspension. The cells were washed with PBS and filtered through a 70 $\mu$m sieve to remove any clumps. CD4$^+$ and CD8$^+$ expressing thymocytes were depleted using anti-CD4 and CD8 magnetic-activated cell-sorting (MACS) beads (Miltenyi Biotec), according to the manufacturer’s instructions. The depleted thymocyte preparation was stained with surface antibodies for sorting on a FACSaria (BD Biosciences).

**Flow cytometry**

For the analysis of cell surface phenotype, the cells were simply stained with antibodies. For the analysis of intracellular cytokine expression, cells were first restimulated *in vitro* with 50ng/mL phorbol 12-myristate 13-acetate (PMA) + 2 $\mu$g/mL ionomycin (both Sigma-Aldrich) in the presence of Monensin (BD Biosciences) at 37°C for 2.5h before staining with cell surface antibodies. The cells were then fixed with the Intracellular Fixation and Permeabilization buffer set (eBioscience) and stained with antibodies against cytokines. All antibodies were purchased from eBioscience or BD Biosciences. Flow cytometry data were then acquired on a BD LSR Fortessa.
III (BD Biosciences) and analyzed with FlowJo software v10.7.0 (Treestar). When only analyzing cell surface phenotype, dead cells were excluded by DAPI. For intercellular cytokine analyses the cells were not stained with DAPI but were gated on live cells determined by size.

**OP9-DL1 co-cultures**

Thymocyte subpopulations for interest were purified by MACS depletion and cell sorting then plated onto OP9-DL1 monolayers (Holmes & Zuniga-Pflucker, 2009). The OP9-DL1 cells were inactivate with Mitomycin C (Stem Cell) immediately prior to use. 5×10^2 to 3×10^3 sorted thymocytes were seeded per well of a 96-well plate in aMEM (Life Technologies) supplemented with 20% FCS (Bovogen Biologicals), penicillin/streptomycin/gentamycin (Gibco), 2ng/mL murine IL-7 (Peprotech) and 5ng/mL human FLT3L (Peprotech) and were incubated at 37°C and 5% CO₂. The media was refreshed every 2d and freshly inactivated OP9-DL1 cells were added every 4d. The cells were analyzed by flow cytometry after 9-26d of culture.

**Fetal thymic organ culture (FTOC)**

Fetal thymic lobes were isolated from embryos at gestational 14d following timed pregnancies of C57BL/6 female mice. They were cultured for 5-6d on 0.8mm isopore membranes (Millipore) atop surgical gelfoam sponge (Ferrosan Medical Devices) soaked in RPMI-1640 (Sigma) supplemented with 10% FCS (Bovogen Biologicals), 20mM HEPES (Sigma-Aldrich), 50mM 2-mercaptoethanol (Sigma) and 1.35mM 2’-deoxygyanosine (dGuo, Sigma) to deplete endogenous thymocytes. The depleted thymic lobes were then transferred onto new sponges soaked in fresh media with supplements but without dGuo for 2d before repopulation with thymocyte progenitors. To repopulate thymic lobes, they were placed in 20mL hanging drop cultures on Terasaki plates (Sigma-Aldrich) containing 5×10^2 to 2×10^3 sorted thymocytes for 24h before returning to fresh sponges. The media was refreshed every 3-4d. Single cell suspensions of the thymic lobes were generated by passing through a 70mm sieve for analysis by flow cytometry.

**Barcode transduction**
Sorted total DN1 thymocytes were pre-cultured on OP9-DL1 for 24h. The cells were then transduced with barcode lentivirus library (Naik et al., 2013) in StemSpan medium (Stem Cell Technologies) by centrifugation at 900 ×g for 1.5h at room temp. A viral titer pre-determined to give 10% transduction efficiency was used to ensure that the cells are not transduced with multiple barcodes. 2ng/mL murine IL-7 and 5ng/mL human FLT3L was then added to each well and the cells were returned to the incubator. The following day, fresh αMEM with supplements was added. αβ (CD90.2+ CD8α+ TCRγδ-) and γδ (CD90.2+ CD8α- TCRγδ+) lineage cells were sorted after 14d and 20d OP9-DL1 co-culture.

Barcode amplification and sequencing

Barcode library construction was performed as described previously (Naik et al., 2013). The cells were lysed in 0.5mg/ml Proteinase K (Invitrogen) in Direct PCR Lysis Buffer (Viagen) at 55°C for 2h. The Proteinase K was then inactivated at 85°C for 30min and 95°C for 5min. The lysate was split into 2 wells for technical replicate PCRs. A first round of PCR was performed using 1× Standard-Taq magnesium free reaction buffer pack (NEB) with 2 mM MgCl2 (NEB), 0.2mM dNTPs (made in house), 0.5μM TopLiB forward primer (TGCTGCCGTCAACTAGAACA) and 0.5μM of BotLiB reverse primer (GATCTCGAATCAGGCGCTTA) for 32 cycles (1 cycle at 95°C for 5min, 30 cycles at 95°C for 15sec, 57.2°C for 15sec and 72°C for 15sec followed by 1 cycle at 72°C for 10min). A second round of PCR was then performed to add different Illumina index to each sample by amplifying the first round PCR product with a sample specific Illumina forward index primer and a common Illumina reverse index primer for 32 cycles (1 cycle at 95°C for 5min, 30 cycles at 95°C for 15sec, 57.2°C for 15sec and 72°C for 15sec followed by 1 cycle at 72°C for 10min). An aliquot of the PCR product was run on a 2% agarose gel to check for barcode amplification, then the samples were pooled and the DNA was cleaned using NucleoMag SPRI beads (Machery-Nagel). A 75-cycle sequencing run was performed on a NextSeq instrument (Illumina).

Cellular barcode data processing and analysis
The data was demultiplexed and aligned to the reference barcode library using *processAmplicons* function from edgeR package (Robinson, McCarthy, & Smyth, 2010). The barcode counts were then processed in the following steps: 1) barcodes with 0 read counts in all sample/cell types were excluded from the analysis; 2) Barcodes that were detected in the water control were also removed from the analysis; 3) The read count between technical replicates of the same sample was averaged and the total read count in each sample was then normalized to 100%.

t-distributed stochastic neighbor embedding (t-SNE) was used to cluster the normalized barcode profiles and visualize any lineage biases of individual barcodes. DBSCAN (version 1.1-8) was then used to classify barcodes based on their corresponding t-SNE coordinates. Heatmaps were plotted to visualize lineage output of barcodes that were identified by transforming the normalized reads value for each barcode using logarithmic transformation.

Two-tailed Pearson’s correlation analysis with 95% confidence interval was calculated to determine the correlation between each lineage outcomes using Prism v9 (GraphPad).

**Single-cell RNA sequencing**

Sorted thymocytes were counted and checked for viability, then loaded onto the 10X Chromium platform (10X Genomics) for scRNA-seq library construction using the Single Cell V2 or V3.1 Reagent Kit according to the manufacturer’s instructions. Libraries were sequenced using 150-cycle/150bp-reads NextSeq500 (Illumina) or 300-cycle/150bp-reads Novaseq (Illumina). Sequencing files were demultiplexed and aligned to *Mus musculus* transcriptome reference (mm10), and count matrix was extracted using CellRanger Single Cell software v2.1.1 or v4.0.0 (10X Genomics) (Zheng et al., 2017) (Supplementary Table 1).

The Illumina sequencing output was pre-processed with Seurat (v2.3 or v3.2.2) on R (v3.6.3 and 4.1.0). Cells with <500 genes, >5000 genes or >7% mitochondria gene expression were filtered out as low-quality cells. Following normalization and removal of confounders, highly variable genes were identified and selected using VST selection method (Zwiener, Frisch, & Binder, 2014). Unsupervised linear principal component analysis (PCA) was performed on these highly variable genes to grouped them into 20 principal components. Cell
clustering was implemented using the number of components that retain >90% of variance of gene expression in the data (stated in Supplementary Table 1).

DoubletFinder (v2.0.3) was applied to remove likely sequencing doublets before downstream analyses (McGinnis, Murrow, & Gartner, 2019). The expected number of doublets was calculated as 0.75% of every 1,000 targeted cells of recovery. The remaining cells were re-clustered and were visualized with t-SNE or uniform manifold approximation and projection (UMAP) dimensional reduction with a resolution of 2.0 (Table S1). Differential expression between subclusters was carried out using FindAllMarkers function, with default parameters; differentially expressed genes with adjusted p-value <0.05 and fold change >0.5 or <-0.5 (log2FC) were considered unless otherwise stated. The heatmaps was generated using function DoHeatmap. Hierarchical clustering heatmaps were generated with the R package ComplexHeatmap (v2.8.0) using Euclidean distance measures (Gu, Eils, & Schlesner, 2016).

Cell cycle genes specific to either the G1, S, G2M stages was used to perform cell cycle scoring and assign cells to their respective stage of the cell cycle (Macosko et al., 2015). Cell cycle genes were regressed using Seurat’s built-in regression model.

**Merging of multiple scRNA-seq datasets**

To compare cell types and proportions across three independent sequencing runs, the datasets were integrated as described at [https://satijalab.org/seurat/archive/v3.0/integration.html](https://satijalab.org/seurat/archive/v3.0/integration.html) (Stuart et al., 2019). The Seurat package (v.3.2.2) was used to assemble multiple distinct scRNA-seq datasets into an integrated dataset and cell cycling scores were calculated. To remove technical variability, the datasets were pre-processed and normalized using SCTransform (Hafemeister & Satija, 2019). To correct for experimental batch effect, integration anchors were identified between the experiments then merged using canonical correlation analysis. Linear dimensional reduction was applied and principal components that retain >90% of variance of gene expression in the data were
Unsupervised clustering was implemented on the integrated data and clustered the cells using a resolution of 2.0.

**Pseudotime trajectory construction**

Filtered 10X data was imported into Monocle 2 by generating a cell dataset from the raw counts slot of the Seurat object. Cells were ordered into a branch pseudotime trajectory according to the procedure recommended in the Monocle 2 documentation (Qiu et al., 2017). The highly variable genes identified by Seurat were chosen as ordering genes to recover pseudospatial trajectories using the setOrderingFilter, reduceDimension and orderCells functions in Monocle 2 using default parameters. Differential expression between pseudotime states were determined using the Seurat function FindAllMarkers.

**Slingshot trajectory construction**

Cluster information from merged 10X dataset were used as input for Slingshot package (v2.0.0) to infer cell trajectory (Street et al., 2018). Default parameters was applied and DN1b was set as the starting point for the minimum spanning tree. The previously generated UMAP clustering from the integrated data was set as the cellular embedding on which Slingshot performed trajectory inference computation.

**GO term enrichment analysis**

The biomaRt package (v3.13) was loaded from the Bioconductor using R studio (Amezquita et al., 2020). The Gene Ontology (GO) terms Cell surface markers (0009986) and Transcription factors (0003700, 0008134, 0140110) were obtained from ‘mmusculus_gene_ensembl’ and applied to the input (differentially expressed) gene lists. Gene ontology enrichment analysis for GO Biological Process was conducted using ShinyGo (v0.66) (Ge, Jung, & Yao, 2020).

**Statistical analysis**
Statistical testing was performed with one-way analyses of variance (ANOVA) using Prism v9 (GraphPad). \( P \) values are shown as \(* < 0.05, ** < 0.01, *** < 0.001, \) and **** \( < 0.0001 \) where each statistical significance was found, and all data are represented as means \( \pm \) S.E.M.

**Data availability**

All scRNA-seq datasets have been deposited in the NCBI Gene Expression Omnibus repository under accession number GSE188913.

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**Competing interests**

The authors declare that they have no competing interests.

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