Characterisation of innate lymphoid cell populations at different sites in mice with defective T cell immunity [version 2; referees: 1 approved, 1 approved with reservations]

Emma E. Dutton¹, Ana Camelo², Matthew Sleeman²,³, Ronald Herbst⁴, Gianluca Carlesso⁴, Gabrielle T. Belz⁵,⁶, David R. Withers ¹

¹Institute of Immunology and Immunotherapy, College of Medical and Dental Sciences, University of Birmingham, Birmingham, UK
²MedImmune, Aaron Klug building, Granta Park, Cambridge, UK
³Immunology & Inflammation Group, Regeneron, Tarrytown, NY, USA
⁴Department of Cancer Biology, MedImmune, Gaithersburg, MD, USA
⁵Department of Medical Biology, University of Melbourne, Parkville, Victoria, Australia
⁶The Walter and Eliza Hall Institute of Medical Research, Parkville, Victoria, Australia

Abstract

Background: Innate lymphoid cells (ILCs) have now been identified within most tissues of the body and current evidence indicates that this family of cells play a fundamental role in maintaining tissue homeostasis. However, few studies have compared the ILC populations between several tissues.

Methods: We sought to generate a comprehensive characterisation of the ILC populations in different tissues of C57BL/6 WT and genetically modified mice targeting costimulatory pathways, using transcription factor expression to define specific groups.

Results: Consistent with studies individually describing the ILC composition in different tissues, our analysis revealed different ILC groups dominate the ILC population in different tissues. Additionally, we observed a population of IL-7Ra⁺Id2⁺ cells lacking expression of lineage markers but also lacking expression of GATA-3, RORgt or T-bet. This population was most evident in ear skin where it outnumbered the defined ILC groups, however, further experiments demonstrated that detection of these cells was influenced by how the tissue was digested, raising concerns as to its real nature. Since both ILC2 and ILC3 express ICOS, we then investigated the requirement for ICOS:ICOSL interactions in the homeostasis of ILC populations at these sites. Surprisingly, no significant differences were detected in the number of ILC1, ILC2 or ILC3 between WT and ICOSL⁻/⁻ mice in any tissue, indicating that this pathway is not required for ILC homeostasis at these sites. These data were compared with CD80⁺CD86⁻/⁻ mice given evidence of CD28 expression by some ILC and ILC crosstalk with activated T cells. Notably, the absence of CD28 ligands resulted in a significant increase in ILC2 and ILC3 numbers in the intestine.

Conclusions: Together, these data provide new insight into ILC composition in different tissues in both WT and genetically modified mice where key costimulatory pathways are genetically deleted, providing a useful resource for further research into ILC biology.
Corresponding author: David R. Withers (d.withers@bham.ac.uk)

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Competing interests: AC, GC and RH are full time employees of MedImmune LLC. MS is a full time employee of Regeneron.

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### Introduction

In recent years, innate lymphoid cells (ILCs) have received significant attention and current evidence indicates they play a key role in the maintenance of tissue homeostasis and the initiation of inflammatory responses. Whilst initial studies of ILCs focused on the intestine, it is now evident that these cells are located in most, if not all tissues in the body, including mucosal barrier sites, primary and secondary lymphoid tissue, skin and organs such as the liver, heart and bladder. Formal nomenclature of the ILC groups brought together key phenotypic and functional data and provided a clear framework in which to further investigate this family of immune cells. Three distinct groups of ILCs, characterised on the basis of transcription factor expression and associated cytokine production have been described and whilst plasticity may blur our understanding of how different populations developed within a tissue, transcription factor expression has provided a robust means of unequivocally identifying ILC populations, superior to reliance on surface markers that show tissue-specific variations in expression.

Although ILCs appear ubiquitous in the body, within given tissues it is evident that distinct ILC groups dominate. For example, whilst all ILCs are present in the intestinal tract, the majority of these cells belong to the RORγt-expressing group 3 population (ILC3). In contrast, the vast majority of ILCs in lung or adipose tissue express high levels of GATA-3 characteristic of the group 2 ILCs (ILC2). Furthermore, developmentally distinct subsets of ILC within a specific group are evident, particularly amongst ILC3s and these also exhibit tissue tropism. Our previous analysis of ILC populations within secondary lymphoid tissues relied on transcription factor expression rather than surface markers and this approach clearly identified three distinct ILC groups that accounted for the total IL-7Rα-Lineage- (Lin) population of cells. Thus our aim was to extend this approach to compare ILC populations within a range of lymphoid and non-lymphoid tissues, reasoning that this would provide a useful resource from which to better develop understanding of ILC populations. Several studies have also indicated that ILC-T cell cross talk shapes the outcome of CD4 T cell responses and that both cell types require common γ chain cytokines for their homeostasis. Furthermore, ILCs express a number of costimulatory molecules whose expression is shared with T cells, although the role these molecules play in ILC function requires further investigation. Thus, the secondary aim of this work was to characterise ILCs in mice in which specific costimulatory molecules were absent.

### Methods

#### Mice

All mice used were bred and maintained in accordance with Home Office guidelines at the University of Birmingham, with C57BL/6 background, including CD80−/−, ICOS−/− (provided by Medimmune), ICOSL−/−, Id2-eGFP and WT. All mice were used at 6 weeks of age or older (up to 12 weeks of age). Within experiments, mice were age and sex matched, but not randomised. Mice were housed within IVC cages at up to 4 mice per cage of the same sex dependent upon the composition of the litter. Mice were checked daily by trained staff and every effort was made to ameliorate any suffering of the animals. Mice were checked daily by trained staff. Mice were culled by cervical dislocation.

#### Cell isolation

**Secondary lymphoid tissue digestion protocol.** Lymph nodes (LN) and spleen were cleaned and teased in Roswell Park Memorial Institute (RPMI) 1640 Medium (life technologies). For digestion, collagenase dispase (final concentration, 0.25 mg/ml) (Roche Life Sciences) and DNase (0.025 mg/ml) (Roche Diagnostics) were added followed by incubation at 37°C for 20 minutes. Digestion was stopped through the addition of Ethylenediaminetetraacetic acid (EDTA) (0.01 M) (Sigma-Aldrich) and the tissues were crushed through a 70 μm filter. Samples were centrifuged (5 minutes, 1,500 rpm, 4°C) and supernatant removed. LNs were resuspended in appropriate amount of DPBS supplemented with 2% FBS and 0.5% EDTA, and spleen samples resuspended in 5 ml of Gey’s red blood cell lysis buffer (70% H2O, 20% solution A, 5% solution B, 5% solution C) (Table 1) and incubated for 5 minutes on ice. Spleens were diluted with RPMI, centrifuged and re-suspended in the appropriate amount of DPBS supplemented with 2% FBS and 0.5% EDTA (staining buffer).

**Lung digestion protocol.** Prior to the removal of the lungs they were perfused with DPBS (Life Technologies). Briefly, the right atrium of the heart was pierced and the left ventricle injected with 10 ml of DPBS, inflating the lungs and flushing out the blood. The lung was cleaned and teased apart in RPMI-1640 supplemented with 1% Penicillin Streptomycin solution, 1% L-Glutamine and 10% Fetal Bovine Serum (culture medium). Liberase TM/DNase solution (for one lung; 500 μL Liberase TM (42.4 μg/ml) and DNAse (0.025 mg/ml) (Roche Diagnostics) were added followed by incubation at 37°C for 20 minutes. Digestion was stopped through the addition of Ethylenediaminetetraacetic acid (EDTA) (0.01 M) (Sigma-Aldrich) and the tissues were crushed through a 70 μm filter. Samples were centrifuged (5 minutes, 1,500 rpm, 4°C) and supernatant removed. LNs were resuspended in appropriate amount of DPBS supplemented with 2% FBS and 0.5% EDTA, and spleen samples resuspended in 5 ml of Gey’s red blood cell lysis buffer (70% H2O, 20% solution A, 5% solution B, 5% solution C) (Table 1) and incubated for 5 minutes on ice. Spleens were diluted with RPMI, centrifuged and re-suspended in the appropriate amount of DPBS supplemented with 2% FBS and 0.5% EDTA (staining buffer).

#### Table 1. Geys Red Cell Lysis Buffer.

<table>
<thead>
<tr>
<th>Solution</th>
<th>Additive</th>
<th>Quantity</th>
</tr>
</thead>
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<tr>
<td>A</td>
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</tr>
<tr>
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<td>KCl</td>
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</tr>
<tr>
<td></td>
<td>Na2HPO4</td>
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<td></td>
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<td></td>
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<tr>
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<td>MgCl2</td>
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<td></td>
<td>CaCl2</td>
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</tr>
<tr>
<td>C</td>
<td>NaHCO3</td>
<td>22.5g</td>
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</table>
(Roche Life Sciences), 10 μL DNase (10 mg/ml) and 3.5 ml of culture media was added per sample for digestion. The tissue was incubated at 37°C and shaken for 45 minutes before being crushed through a 70 μm filter and washed with culture media. Samples were then centrifuged, supernatant removed, resuspended in Gey’s solution and incubated on ice for 2 minutes, before being diluted in culture media and then filtered, washed and re-suspended in an appropriate amount of staining buffer.

Small intestine digestion protocol. The small intestine (SI) was dissected from below the stomach and above the caecum then placed in a petri dish containing Hank’s Balanced Salt Solution (HBSS) (Sigma-Aldrich) 2% FBS. The fat and Peyer’s patches were removed before the SI was cut longitudinally and the contents washed out. The tissue was cut into small pieces, placed in HBSS and shaken vigorously, before being filtered through nitex mesh. The SI underwent a series of incubations in various digestion media; it was placed in a specific digestion media, shaken vigorously for 20 seconds, incubated and shaken at 37°C for 20 min (HBSS/EDTA wash) or 15 min (collagenase digestion). This is followed by a washing process where the SI was resuspended in HBSS and vigorously shaken for a further 20 seconds before being filtered again. This process was carried out with the following digestion media: HBSS 2mM EDTA (20 minutes) twice and pre-warmed culture media containing 1 mg/mL collagenase VIII (Sigma-Aldrich) (15 minutes). The SI was filtered through 100 μm and 70 μm cell strainers before being centrifuged and re-suspended in an appropriate amount of staining buffer.

Ear skin digestion protocol. Ears were cut into small sections and incubated in a shaker at 37°C for 30 minutes in 1 ml of Liberase TM/DNase digestion solution (3 ml per ear; 3 ml DMEM, 75μL Liberase TM (0.28 Wunsch units/mL) and 75 μL DNase (200 mg/mL)). The tissue was filtered through a 70 μm filter and washed with DMEM, and the supernatant collected. The ear was then removed from the strainer and this process repeated twice more. After the third incubation, the ear was crushed through the filter and washed with DMEM. Samples were centrifuged and suspended in an appropriate amount of staining buffer.

Flow cytometry
Samples were stained using the antibodies in Table 2. A minimum of 1% cell suspensions were stained. To identify in ILCs dead cells were excluded using a LIVE/DEAD cell viability assay in APC-Cy7. A lineage cocktail containing monoclonal Abs to B220, CD11c, CD11b, CD3 and CD5 was used except where stated and ILC were identified as IL-7Rα-Lin-. Surface staining was conducted with antibodies diluted in staining buffer at 4°C for 30 minutes. Cells were fixed and intracellularly stained using the Foxp3 Staining Buffer Set (eBioscience, catalogue number 00-5523-00) according to manufacturers instructions. Samples were run on the BD LSRSFortessa™ X-20 (BD Biosciences) and data collected using BD FACSDiva Software (BD biosciences).

Statistical analysis
Data collected were analysed using Flow Jo software vX.0.7 (Treestar) and GraphPad Prism 6. Pairs of samples were compared using a two-tailed Mann-Whitney T test; multiple samples were compared using Kruskal-Wallis one-way ANOVA with post hoc Dunn’s test: * p ≤ 0.05, ** p ≤0.01, *** p ≤ 0.005. The bar represents the median and where appropriate the median has been shown. Where a p-value is not indicated, no statistical difference was observed.

Results
Characterisation of ILC populations in different lymphoid and non-lymphoid tissues
Over the last few years, enormous progress has been made identifying ILC populations within different murine tissues. A range of surface markers were initially used to describe these cells, however, tissue specific differences in expression of these molecules is evident and has created some discrepancies in the populations described. We sought to assess all ILCs in a collection of non-lymphoid tissues such as the intestine, lung and peripheral skin and lymphoid (different lymph nodes) to better compare these populations at different sites. All ILC populations express IL-7Rα and we have previously found this a robust initial means of identifying ILC amongst cells lacking expression of markers associated with other lineages, with subsequent characterisation of ILC groups based largely on transcription factor expression\textsuperscript{13}. Thus, cells were prepared from SI lamina propria (SI LP), lung, ear skin, mesenteric LNs (mLN, a pool of 5 LN per sample) and auricular LNs (auLN, a pool of 2 LN per sample). The mLNs were chosen to contrast with the SI LP and the auLN chosen to compare with the ear skin. ILCs were identified as IL-7Rα^Lin^-cells with cells expressing intracellular CD3ε (iCD3ε) also excluded to minimise potential T cell contamination of the ILC gate (Figure 1A). Consistent with other studies, enumeration of ILCs within these different tissue identified that this family of cells was most numerous in the mucosal barrier sites (intestine and lung) as either a proportion of hematopoietic cells (Figure 1B) or as total cell numbers per tissue (Figure 1C). While a substantial (~10%) of the live hematopoietic cells within ear skin were located within the IL-7Rα^Lin^-gate, the number of CD45^-cells isolated from ear tissue was very low (Figure 1B, C). A distinct ILC population was evident in the auLN and mLNs, but numerically these were small populations and vastly out-numbered by naïve lymphocytes\textsuperscript{1}. Analysis of GATA-3, RORγ and T-bet expression amongst ILC clearly identified the three described ILC groups. In the SI LP and mLN this accounted for the vast majority of the IL-7Rα^Lin^-cells (Figure 1A). Graphical representation of the mean proportion of each subset in each tissue from multiple mice illustrated the substantial differences in the proportion of ILC subtypes in the various tissues (Figure 1D). In all the tissues analysed, but most evident within the ear skin, we detected a population of IL-7Rα^Lin^-cells lacking clear expression of GATA-3, RORγ and T-bet. We termed these cells ‘Triple Negative’ (TrN) cells in the context of this study. An obvious concern was that these cells were wrongly identified as ILC, due a failure to completely discriminate them on the basis of the lineage-defining antibodies used. Given the substantial proportion of these TrN cells identified in the ear skin, we repeated the analysis of ILC populations in ear skin using an extended cocktail of Abs against different immune cell lineages to potentially further restrict the cells termed IL-7Rα^Lin^- (Supplementary Figure 1). Whilst this revised lineage discrimination modestly reduced the proportion of TrN cells, they remained a substantial population (almost 50%) of the IL-7Rα^Lin^-cells, arguing that these cells could not be accounted for solely through contamination with other immune cell populations. To provide evidence that these
Table 2. Antibodies used for flow cytometry.

<table>
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<th>Conjugate (dilution)</th>
<th>Clone</th>
<th>Concentration (mg/ml)</th>
<th>Manufacture</th>
<th>Host</th>
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...cells displayed further characteristics of ILCs, we assessed their expression of Id2 using Id2-eGFP reporter mice. The vast majority of IL-7RαLin cells in the ear skin or draining auLN were eGFP compared with WT controls (Supplementary Figure 2), demonstrating that most of the TrN population expressed Id2, although levels of eGFP appeared lower than in splenic NK cells. However, a further concern was that the manner in which these cells were isolated was affecting the ILC that could be identified. The protocol for the digestion of ear skin tissue uses the same enzymes used in the digestion of lung tissue, but at a higher concentration and for longer. To test whether the ear digestion protocol could affect the ILC populations identifiable by flow cytometry, we digested lung tissue with the normal lung digest protocol or the protocol used to digest the ear. Strikingly, the ear digestion protocol resulted in changes in the proportions of different ILC populations in the lung tissue, reducing the GATA-3+ population approximately 2-fold and doubling the proportion of TrN cells (Supplementary Figure 3). We also assessed whether the ear digestion protocol was affecting any of the lineage markers detected by the cocktail of Abs used to identify ILC. Whilst some markers were clearly affected by the prolonged ear digestion, these were not ones expressed by IL-7RαLin cells, minimising the impact on ILC identification (Supplementary Figure 4). Thus, while the TrN cells described here share several key features of ILCs, there are clear concerns that this population is either an artefact of the protocol used to isolate cells or at the very least likely exaggerated because of the digestion employed.
Figure 1. Identification of ILC groups in different tissues using transcription factor expression. To characterise ILC groups in lymphoid and non-lymphoid tissues, cells were isolated from the ear, auLN, SI LP, lung and mLN and the ILC groups identified on the basis of expression of the transcription factors GATA-3, RORγt and T-bet. A) Representative flow cytometry plots showing the gating approach used to identify the populations in the different tissues. ILCs were identified as live CD45+IL-7Rα-Lin (CD3, CD5, B220, CD11b, CD11c)-iCD3- cells. Coloured gates used to show specific groups of cells with ILC1 (T-bet+, blue), ILC2 (GATA-3+, red), ILC3 (RORγt+, green) and ‘Triple Negative’ cells lacking expression of the GATA-3, RORγt and T-bet (black). B) ILCs as a percentage of the hematopoietic population in each tissue. C) The total numbers of ILCs isolated from each tissue. D) Pie charts showing the mean proportion of each ILC group using the group-specific colours used in part ‘A’. Cells falling outside of the transcription factor gates were termed ‘ungated’ and included to allow analysis of the entire ILC compartment. Each data point represents cells isolated from 2x ears, 2x auLN, 1x SI LP, 1 x lung and 1 pool of mLNs isolated from one mouse. The data shown for ear (n=6), auLN (n=10), SI LP (n = 10), lung (n = 13) and mLN (n=26) are pooled from a minimum of 3 independent experiments for each tissue. Values on flow cytometry plots represent percentages, bars on scatter plots represents the median, which is also shown numerically for clarity.
It is now evident that ILC3s can be further split into developmentally distinct subsets based upon expression of CCR6 and NKp46, with CCR6^+NKp46^- LTi-like ILC3, CCR6^+NKp46+ ILC3 and a CCR6^+NKp46^- population that contain cells able to differentiate into the CCR6^+NKp46+ and CCR6^+NKp46^- subsets\(^6,12\). Thus, CCR6 and NKp46 expression amongst the ROR\(\gamma\)^+ ILC3 in the different tissues was analysed and consistent with other studies, all three populations were evident in the SI LP (Figure 2A, B) and all populations were most numerous in this tissue compared with other sites (Figure 2C). In the different LNs, the majority of the ILC3 were CCR6^+NKp46-, again consistent with previous work\(^11\). Within the ear skin and lung, the number and proportion of ILC3 was very low, but this small population contained both CCR6^+NKp46+ and CCR6^+NKp46- subsets. Thus, whilst most CCR6^+NKp46+ ILC3 were present in the intestine, these were detected at low number in the other tissues. The NKp46^-CCR6+ ILC3 appeared to be more restricted to the intestine than the other subsets of ILC3, in regard to the tissues assessed in this study.

To investigate the extent to which the tissue of origin affected their expression of reported surface markers, we further characterised the ILC2 in ear skin, lung, SI LP and mLN. Whilst the most notable difference in phenotype was between ILC2 isolated from ear skin compared with other tissues, these data must be viewed with caution given the prolonged enzymatic digestion

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**Figure 2. Identification of ILC3 subsets in different tissues using CCR6 and NKp46 expression.** To identify the different subtypes of ILC3 in the tissues, ROR\(\gamma\)-expressing ILC3s were assessed for expression of CCR6 and NKp46. **A** Representative flow cytometry plots showing the gating approach used to identify the subsets in each tissue, having first gated on ROR\(\gamma\)-expressing ILC3 as shown in Figure 1. **B** The percentage of each ILC3 subset in the different tissues. **C** The total numbers of each ILC3 subset isolated from each tissue. Each data point represents cells isolated from 2x ears, 2x auLNs, 1x SI LP, 1x lung and 1 pool of mLNs isolated from one mouse. The data shown for ear (\(n=7\)), auLN (\(n=10\)), SI LP (\(n=30\)), lung (\(n=13\)) and mLN (\(n=26\)) are pooled from a minimum of 3 independent experiments for each tissue. Values on flow cytometry plots represent percentages, bars on scatter plots represents the median, which is also shown numerically for clarity.
required to isolate cells from skin (Figure 3A, B). ILC2 isolated from either the SI LP or lung expressed CD25, ICOS, KLRG-1 as previously reported, however, ST2 expression was very low in the intestine (Figure 3A, B). ILC2 have also been reported to express major histocompatibility complex class II (MHCII), facilitating interactions with CD4 T cells\(^1\) and whilst ILC2 isolated from tissues had little detectable MHCII expression, amongst those isolated from the mLN, there appeared to be a clear MHCII\(^+\) subset (Figure 3A, B).

**Role of ICOS:ICOSL interactions in ILC homeostasis**

Given the relatively recent identification of many ILC populations, it is unsurprising that our understanding of the role of many molecules expressed by ILC is limited. Interestingly, the initial characterisation of ILC2, alongside early descriptions of LTi-like cells, have identified ILC expression of costimulatory molecules with known roles in CD4 T cell responses\(^1\). In our hands, ILC3 expression of ICOS in the SI LP was restricted to the CCR6\(^-\) fraction, with a homogenous level of expression comparable to that observed on ILC2 (Figure 4A). Signalling through ICOS has been reported to increase ILC2 proliferation in both the lung and intestine, whilst ICOS\(^-\) mice are described as having fewer ILC2 in the lung\(^19,20\). Employing our broad characterisation of tissue ILC populations, we sought to assess the importance of signals through ICOS in ILC homeostasis. Thus, ILCs were isolated from the SI LP, lung and mLN of WT and ICOSL\(^-\) mice and the total number of ILC as well as the proportion and total number of specific ILC groups assessed (Figure 4B–D). In contrast to earlier reports defining ILC populations with cell surface markers alone, no differences in the total number of the ILC2 or ILC3 were observed in any tissue from WT and ICOSL\(^-\) mice, indicating that ICOSL signals are not essential for ILC homeostasis in these tissues. Similarly, no change in the total number of ILC2 or ILC3 was evident when ILCs were isolated from the SI LP of ICOS\(^-\) mice compared with WT ILCs, although modest differences in the number of ILC3 in mLN were observed (Supplementary Figure 5). Although only CCR6\(^-\) ILC3 in the SI LP uniformly express ICOS, no differences were detected in the total number of different ILC3 subsets in ICOSL\(^-\) SI LP (Figure 4E) or mLN (Figure 4F), arguing that CCR6 ILC3 populations are not ICOSL dependent.

ILC populations in mice lacking activated T cells

Our analysis of ILC populations in mice deficient in ICOS or ICOSL indicated no clear effect on ILC populations despite some of the cells expressing ICOS expression. Whilst there is little clear evidence that ILC express the CD28 ligands CD80 and CD86, there is evidence to support ILC expression of CD28\(^25\). Furthermore, ILC cross talk with activated T cells has been suggested by several studies\(^7,14,15\), arguing that disruption of T cell activation may impact upon ILC homeostasis. In addition, the cytokines that support ILC homeostasis, IL-7 and IL-15, are similarly required for T cell populations, and competition for these molecules has been proposed as a mechanism limiting ILC and T cell numbers\(^16,26\). Thus we considered that it would be informative to extend our analysis of ILC populations to mice deficient in both CD80 and CD86, where signals through CD28 are deficient and thus T cell activation is blocked (Figure 5 A–C). No differences were observed within lung ILC

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**Figure 3. Characterisation of ILC2 surface phenotype in different tissues.** To further characterise the ILC2 population identified in the different tissues, expression of CD25, ICOS, KLRG-1, MHCII and ST2 was assessed. A) The percentage of each GATA-3\(^+\) ILC expressing the stated surface marker isolated from ear skin, SI LP, lung or mLN. B) The median fluorescence intensity (MFI) of the stated surface marker total numbers of each ILC2 subset isolated from ear skin, SI LP, lung or mLN. Each data point represents cells isolated from 2x ears, 1x SI LP, 1 x lung and 1 pool of mLNs isolated from one mouse. The data shown for ear (n=5), SI LP (n = 8), lung (n = 6) and mLN (n=6) are pooled from 3 independent experiments for each tissue, bars on scatter plots represents the median. Statistical significance was tested using Kruskal-Wallis one-way ANOVA with post hoc Dunn’s test: *p≤0.05, **p≤0.01, ***p≤0.001.
Figure 4. Normal numbers of ILC1, ILC2 and ILC3 in ICOSL−/− mice. To investigate the role of ICOS:ICOSL interactions in ILC homeostasis, cells were isolated from the SI LP, lung and mLNs of WT and ICOSL−/− mice and ILC groups enumerated. ILCs were identified as live CD45−IL-7Rα+Lin−iCD3− cells, transcription factor expression was used to identify ILC1 (T-bet+, blue), ILC2 (GATA-3+, red), ILC3 (RORγt+, green) and ‘Triple Negative’ cells lacking expression of the GATA-3, RORγt and T-bet (black).

A) Representative flow cytometry plots showing ICOS expression by GATA-3+ ILC2 and RORγt+ ILC3 isolated from the SI LP of WT mice.

B)–D) The total number of ILCs (left hand graph), the percentage of the ILCs within each group (centre graph) and the total number of each ILC group isolated from the SI LP (B), lung (C) and mLN (D). E)–F) The percentage (left hand graph) and total number (right hand graph) of each ILC3 subset isolated from the SI LP (E) and mLN (F) of WT and ICOSL−/− mice. Each data point represents cells isolated from 1 x SI LP, 1 x lung and 1 pool of mLNs isolated from one mouse. The data shown for SI LP WT (n=10–13), SI LP ICOSL−/− (n=8–11), lung WT (n=6–13), lung ICOSL−/− (n=4–11), mLN WT (n=5–7) and mLN ICOSL−/− (n=5–7) are pooled from a minimum of 3 independent experiments for each tissue. Values on flow cytometry plots represent percentages, bars on scatter plots represents the median, which is also shown numerically for clarity. Statistical significance was tested using an unpaired, non-parametric, Mann-Whitney two tailed T test: *p≤0.05, **p≤0.01, ***p≤0.001.
Figure 5. Mice deficient in CD80 and CD86 have increased ILC2 and ILC3 populations in the small intestine. To investigate the effect of blocking T cell activation on ILC populations, cells were prepared from the SI LP, lung and mLNs of WT and CD80−/−CD86−/− mice and ILC populations enumerated. ILCs were identified as live CD45+IL-7RαLin (CD3, CD5, B220, CD11b, CD11c)-iCD3- cells, transcription factor expression was used to identify ILC1 (T-bet+, blue), ILC2 (GATA-3+, red), ILC3 (RORγt+, green) and ‘Triple Negative’ cells lacking expression of the GATA-3, RORγt and T-bet (black). A)–C) The total number of ILCs (left hand graph), the percentage of the ILCs within each group (centre graph) and the total number of each ILC group isolated from the SI LP (A), lung (B) and mLNs (C). D)–E) The percentage (left hand graph) and total number (right hand graph) of each ILC3 subset isolated from the SI LP (D) and mLNs (E) of WT and CD80−/−CD86−/− mice. Each data point represents cells isolated from 1 x SI LP, 1 x lung and 1 pool of mLNs isolated from one mouse. The data shown for SI LP WT (n=7), SI LP CD80−/−CD86−/− (n=5), lung WT (n=5), lung CD80−/−CD86−/− (n=5), mLN WT (n=7) and mLN CD80−/−CD86−/− (n=7) are pooled from a minimum of 3 independent experiments for each tissue. Values on flow cytometry plots represent percentages, bars on scatter plots represents the median, which is also shown numerically. Statistical significance was tested using an unpaired, non-parametric, Mann-Whitney two tailed T test: *p≤0.05, **p≤0.01, ***p≤0.001.
populations of WT and CD80\(^{-/}\) mice, however the ILC3 population in the SI LP was significantly enhanced (Figure 5A), with expanded numbers of all ILC3 subsets (Figure 5D). In addition, the ILC3 populations in the mLN were reduced in the absence of CD80 and CD86 signals. (Figure 5C, E). These data suggest that ILC populations in some tissues are enhanced in the absence of T cell activation or perhaps that T cell activation can restrain ILC numbers.

Discussion

In this study, we have used a flow cytometric approach based upon identification of transcription factor expression to characterise the ILC populations present within different murine tissues. This approach was taken to avoid confusion resulting from tissue specific variations in surface molecule expression and to provide a simple overview of the composition of ILC populations within mice from the same colony. Whilst our data is consistent with existing descriptions of ILCs in tissues such as the intestine, lung and mLN\(^{3,13}\), a surprising observation that emerged was the identification of IL-7R\(^{α}\)Lin cells lacking clear expression of the transcription factors used to define the ILC groups. The proportion of IL-7R\(^{α}\)Lin cells with this phenotype differed markedly between tissues and was most noticeable within ear tissue which, to our knowledge has not been assessed with in this manner previously. The robust expression of the different transcription factors in samples analysed alongside the ear skin argues against an inability to detect expression of these transcription factors. Furthermore, we were unable to diminish this population through extending the cocktail of lineage defining Abs used in the study. Additionally, we provide evidence that these cells express Id2, a defining feature of ILCs. However, when we tested whether the manner in which we digest tissue can affect the proportion of different ILC populations, it is was evident that for lung tissue at least, using the same enzymatic mix for longer and at a higher concentration changed the ILC populations that were detectable. While a novel regulatory ILC population lacking expression of ROR\(γ\)T, GATA-3 and T-bet was very recently described\(^{3}\), the weight of evidence presented here indicates that the ROR\(γ\)T GATA-3 T-bet population we observe in the skin is at least partially influenced by the prolonged digestion and high enzyme concentration used to isolate the cells and likely includes ILCs that are losing transcription factor expression as a result of how they are isolated from the tissue. Whilst ILC progenitor populations within different tissues remain to be fully characterised, these observations highlight how the manner in which different tissues are digested can impact on our ability to properly identify different ILC populations.

The tissue specific composition of ILC groups and also ILC3 subsets are consistent with many studies of ILCs, but are striking when total ILC groups in multiple tissues are viewed together. Having characterised these populations in WT mice, we sought to compare this cellular distribution with mice in which costimulatory molecules were deficient, focusing initially on ICOS, since in addition to its described expression by ILC2\(^{1}\), we detected ICOS protein expression on the surface of ILC3 in the SI LP, but only those lacking CCR6 expression. Surprisingly, we observed no clear defect in the total number of ILC populations in the different tissues analysed, in contrast to other studies\(^{19,20}\). Whilst this argues that this pathway is not essential for normal ILC homoeostasis, it is worth pointing out that this is analogous to the role of ICOS in T cell development, where ICOS is required for differentiation of effector T cells rather than normal naïve T cell development\(^{26}\). We decided to characterise ILC populations in CD80\(^{-/-}\)CD86\(^{-/-}\) mice since there is evidence to support ILC expression of CD28 and a block in T cell activation might reveal further insight into the effects of ILC:T cell cross talk\(^{26-31}\). It was noticeable that in the intestine, the ILC3 population were enhanced, which might reflect greater availability of cytokines such as IL-7 and IL-15, although the absence of T cell responses to the commensal microbial population is obviously a further potential factor. Alternatively, T cell populations may constrain ILCs and the inability form activated T cell populations and regulatory T cells may permit ILC expansion in some tissues.

Conclusions

In this study we have tried to broadly, but definitively assess ILC populations in a range of tissues. While using this analysis in mice is a fairly blunt tool for understanding ILC biology, it does provide some initial experimental observations as to ILC requirements \textit{in vivo}. ILCs express an array of different costimulatory receptors and ligands, yet we know very little about how many of these molecules contribute to ILC function. As we look to understand the roles of ILCs within an intact immune system, this is an important and growing area of research, particularly given the wealth of therapeutics that target these pathways.

Data availability

Data are stored on the online depository, Open Science Framework: http://doi.org/10.17605/OSF.IO/WGZA\(^{M}\). Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

Competing interests

AC, GC and RH are full time employees of MedImmune LLC. MS is a full time employee of Regeneron.

Grant information

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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.
Supplementary material
Supplementary File 1: Supplementary Figure 1–Supplementary Figure 5.
Click here to access the data.

Supplementary Figure 1. IL-7Rα’Lineage’ cells in ear skin.
To better characterise the IL-7Rα’Lineage’ population identified in ear skin, cells were isolated from the tissue and stained with an extended panel of lineage specific markers (comprising B220, CD11b, CD11c, CD5, CD3, CD19, Ter119, CD123, Gr1, F4/80, FcεR1 and CD49b) and the population assessed by flow cytometry. A) Representative flow cytometry plots showing live CD45εIL-7RαεLinεCD3ε cells, with coloured gates identifying ILC1 (T-bet, blue), ILC2 (GATA-3, red), ILC3 (RORεγεεε, green) and ‘Triple Negative’ cells lacking expression of the GATA-3, RORεγεεε and T-bet, black). B) Pie chart showing the mean proportion of each ILC group using the group-specific colours used in part ‘A’.
Cells falling outside of the transcription factor gates were termed ‘ungated’ and included to allow analysis of the entire ILC compartment. Data shown for 2x ears (n=8), values on flow cytometry plots represent percentages.

Supplementary Figure 2. IL-7Rα’Lineage’ cells in ear skin express Id2.
To provide further evidence that the IL-7Rα’Lineage’ lacking Gata-3, RORεγεεε and T-bet resembled an ILC population, the IL-7Rα’Lineage’-population in the ear skin of Id2-eGFP reporter mice was assessed. A) Representative flow cytometry plots showing gating on live CD45εIL-7RαεLinεiCD3ε cells prepared from ear skin and auLN of Id2-eGFP mice versus WT controls (to aid gating on eGFP expression, shown in red). B) The proportion of eGFP+ ILC (live CD45εIL-7RαεLinεiCD3ε cells) isolated from ear skin and auLN of Id2-eGFP mice. Data shown for 2x ears (n=5) and 2x auLN (n=5) pooled from two independent experiments.

Values on flow cytometry plots represent percentages, bars on scatter plots represent the median.

Supplementary Figure 3. ILC transcription factor expression is reduced under the ear digestion protocol.
To further evaluate the effect of the ear digestion protocol (EDP) on identifying ILCs within the ear, cells were isolated from the ear, using normal EDP, the lung using normal lung digestion protocol (LDP) and compared to cells isolated from the lung using EDP. ILCs were identified as live CD45εIL-7RαεLinε (CD3, CD5, CD4, CD49b, CD49d, FcεR1, CD4 and CD8 on live CD45ε cells) isolated from the ear using EDP, lung using LDP and lung using LDP. A) The percentage (left hand graph) and total number (right hand graph) of live CD45ε cells within the lung under EDP (n=5) or LDP (n=5). B) The percentage (left hand graph) and total number (right hand graph) of ILC within the lung using EDP or LDP. F) The percentage (left hand graph) and total number (right hand graph) of ILC subsets within the lung using EDP or LDP. Data are from one experiment, values of flow cytometry plots represent percentages, bars on scatter plots represent the median, which is also shown numerically for clarity.

Supplementary Figure 4. Surface markers used to identify ILC are not affected by the ear digestion protocol.
To evaluate whether cell surface markers identified by Abs in the lineage cocktail were affected by the EDP, cells were isolated from the ear, using normal EDP, the lung using normal lung digestion protocol (LDP) and compared to cells isolated from the lung using EDP. A) Representative flow cytometry plots showing expression of the cell surface markers CD3ε, CD5, CD49b, F4/80, FcεR1, CD4 and CD8 on live CD45ε cells isolated from the ear using EDP, lung using LDP and lung using LDP. B) Percentage of live CD45ε cells expressing the cell surface markers CD3ε, CD5, CD49b, F4/80, FcεR1, CD4 and CD8 within the lung using EDP (n=3) and lung using LDP (n=3). Data are from one experiment, values of flow cytometry plots represent percentages, bars on scatter plots represent the median.

Supplementary Figure 5. Numbers of ILC2 and ILC3 in ICOSε mice.
To further assess the role of ICOS:ICOSL interactions in ILC homeostasis, cells were isolated from the SI LP and mLNs of WT and ICOSε mice and ILC groups enumerated. ILCs were identified as live CD45εIL-7RαεLinε (CD3, CD5, B220, CD11b, CD11c, CD19, Ter119, Gr1, CD123, F4/80, CD49b, FcεR1 and CD49b)ε iCD3ε ε cells, transcription factor expression was used to identify ILC1 (T-bet, blue), ILC2 (GATA-3, red), ILC3 (RORεγεεε, green) and ‘Triple Negative’ cells lacking expression of the GATA-3, RORεγεεε and T-bet (black). A-B) The total number of ILCs (left hand graph), the percentage of the ILCs within each group (centre graph) and the total number of each ILC group isolated from the SI LP (A), and mLN (B). Each data point represents cells isolated from 1x SI LP and 1 pool of mLNs isolated from one mouse. The data shown for SI LP WT and ICOSε (n = 10) and mLN WT and ICOSε (n = 10) are pooled from 3 independent experiments. Bars on scatter plots represent the median, which is also shown numerically. Statistical significance was tested using an unpaired, non-parametric, Mann-Whitney two tailed T test: *p≤0.05, **p≤0.01, ***p≤0.001.
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Nick Powell
School of Immunology & Microbial Sciences, King's College London, London, UK

The paper addresses 4 issues.
1) A descriptive comparison of the relative and absolute ILC subset composition in some tissues of adult B6 mice in the steady state.
2) Identifies that different tissue digestion protocols employed to isolate immune cell fractions may be linked to loss of particular lineage defining markers in ILCs (and other immune cells)
3) An analysis of the impact of ICOSL deficiency on the composition of ILCs in various tissues
4) An analysis of the impact of CD80/86 deficiency on the composition of ILCs in various tissues

General comments
As well as providing a useful “atlas” of ILC subset distribution in different tissues in the steady state, the study also highlights the presence of a population of cells, that are deemed ILCs (defined by absence of major lineage markers and IL7+), that lack expression of any lineage defining transcription factors (RORgt, GATA3, T-bet). The authors term these cells triple negative (TrN) ILCs.

The paper also nicely demonstrates a potentially important technical issue in the ILC field, in that tissue digestion protocols (most importantly dose and duration of enzyme exposure) seems to significantly impact ILC composition using conventional gating strategies. Although it is easy to envisage how high dose enzymes might shave off some surface antigens, it is more conceptually challenging to understand how tissue processing impacts intracellular transcription factor expression. Another possibility is that enzyme treatment leads to loss of lineage defining antigens, resulting in lineage positive cells erroneously appearing in the lineage negative gate (and hence potentially contributing to the variable ILC and TrN ILC pool).

Notwithstanding that a separate technical paper about isolation protocols and ILC marker stability might be needed, I would be intrigued to see how the different digestion protocols impact the expression of all ILC markers in tissue where these cells can be isolated without enzymatic digestion (e.g. splenic ILCs can be visualized by flow cytometry by mechanical dissociation without the need for enzymatic digestion), allowing systematic analysis of ILC marker changes without digestion vs all other digestion protocols.

Specific comments
The manuscript could be improved if the following issues were addressed.
1) Regulatory ILCs should now be considered within the extended ILC family, and given that the presence of "lineage undefined" ILCs in different tissues is a major finding of the paper, there would be merit in more in depth phenotyping of the TF negative population. Unsupervised transcriptomics (e.g. RNA-seq) would provide an unbiased assessment of the cellular identity these cells. However, given the potential relevance of ILCregs, an alternative approach would be to consider performing qPCR to evaluate ILCreg molecules (e.g. Il10, Sox4, etc) in the TF negative population.

2) I would question the value of the ID2 reporter experiments. ID2 is expressed by a wide variety of lymphocytes (and certainly not just by ILCs), and therefore, I'm not sure it helps convince me that the unknown TF negative cells are ILCs, or even narrow down the search (beyond lymphocytes).

3) I would be intrigued to see how the composition of ILC subsets differs in the different digestion protocols using TF reporter mice (rather than intracellular TF staining), as these data raise some important considerations for the field, and strategies to mitigate against this problem are now needed. This is probably be beyond the remit of the current paper, but would be a useful "technical report" to help shape future experiments in the field.

4) I would like to see representative flow plots of MHCII, CD25 etc (Figure 3) as a supplemental figure.

5) In this study there was no impact on the composition of tissue ILC populations in mice with germ line genetic deletion of ICOSL. This is in variance to other published reports that have shown significant reductions in lung ILC2 in mice with disruption of the ICOS/ICOSL axis (albeit with ICOS deficient animals). Although in previous published work this tends to be most apparent in IL33 treated animals - it is also clearly apparent in the steady state (60-70% reduction in lung ILC2 in Icos-/- animals). The manuscript would benefit from a more in depth discussion reconciling this discrepancy.

6) The interpretation of the Cd80-/- x Cd86-/- experiments is probably overstated. I would be cautious of attributing the ILC phenotype observed (expansion of intestinal ILC2 and ILC3) to unactivated T-cells, especially as ILCs may express CD28 (as well as checkpoint molecules, PD-1, which modulate CD28 signalling). Further experiments are needed to conclusively demonstrate whether the observed shift in ILC composition in Cd80-/- x Cd86-/- animals is due to a direct effect of loss of CD80/86 interaction with ILCs, or as the authors imply, by loss of T-cell activation. The authors could address this by "relaxing" the interpretation of their findings and by entertaining some alternative possibilities (as described above). This would potentially include changing the title of the manuscript.

In short, the m/s brings important new information to the table. A few minor (mostly editorial) changes would improve the article.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Partly

Competing Interests: No competing interests were disclosed.

Referee Expertise: Mucosal Immunology, IBD

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Emma Dutton, University of Birmingham, UK, UK

We thank the reviewer for the constructive criticism and have responded to the specific comments below:

Question 1: Regulatory ILCs should now be considered within the extended ILC family and given that the presence of "lineage undefined" ILCs in different tissues is a major finding of the paper, there would be merit in more in depth phenotyping of the TF negative population. Unsupervised transcriptomics (e.g. RNA-seq) would provide an unbiased assessment of the cellular identity these cells. However, given the potential relevance of ILCregs, an alternative approach would be to consider performing qPCR to evaluate ILCreg molecules (e.g. Il10, Sox4, etc) in the TF negative population.

Answer 1: We thank the reviewer for this suggestion. Whilst we have considered a more in depth assessment of the TrN population which could be described as resembling a putative regILC cell, the fact that the detection of these cells was very dependent upon the digestion protocol performed greatly reduced our enthusiasm to study these cells further. We are concerned that the ‘novel’ population described here is artefactual and reflects a loss of transcription factor expression under harsh isolation conditions. The fact that changing the digestion of the lung tissue greatly affects the detection of these cells (alongside the conventional ILC groups) really argues that these cells are unlikely to be a real biologically relevant population. If regILC really do exist then we would agree that further more in depth experiments from other groups would substantiate this potentially new field.

Question 2: I would question the value of the ID2 reporter experiments. ID2 is expressed by a wide variety of lymphocytes (and certainly not just by ILCs), and therefore, I'm not sure it helps convince me that the unknown TF negative cells are ILCs, or even narrow down the search (beyond lymphocytes).

Answer 2: As mentioned in response to a similar comment from Reviewer 1, we agree that Id2 is not a definitive ILC marker, but having gated out lineage expressing cells, detection of Id2 expression by the majority of the IL-7Rα+ Lin- population was supportive of the cells being ILC-like. The aim of this experiment was to ask whether the TrN population, which was >50% of the IL-7Rα+ Lin- population in the ear, expressed Id2 and thus might be considered to be ILC-like in this regard. This was the only point we were trying to make with the Id2-reporters and we have tried to amend the text to make this clearer.
Question 3: I would be intrigued to see how the composition of ILC subsets differs in the different digestion protocols using TF reporter mice (rather than intracellular TF staining), as these data raise some important considerations for the field, and strategies to mitigate against this problem are now needed. This is probably be beyond the remit of the current paper, but would be a useful "technical report" to help shape future experiments in the field.

Answer 3: We agree this would be an interesting experiment to perform but we do not have access to the relevant mouse models.

Question 4: I would like to see representative flow plots of MHCII, CD25 etc (Figure 3) as a supplementary figure.

Answer 4: We agree with the reviewer that this should have been shown and these data have now been added to the revised supplementary figures (Supplementary Figure 7).

Question 5: In this study there was no impact on the composition of tissue ILC populations in mice with germ line genetic deletion of ICOSL. This is in variance to other published reports that have shown significant reductions in lung ILC2 in mice with disruption of the ICOS/ICOSL axis (albeit with ICOS deficient animals). Although in previous published work this tends to be most apparent in IL33 treated animals - it is also clearly apparent in the steady state (60-70% reduction in lung ILC2 in Icos<sup>-/-</sup> animals). The manuscript would benefit from a more in depth discussion reconciling this discrepancy.

Answer 5: We thank the reviewer for this comment and have tried to discuss this point further. We were surprised that in our hands we failed to observe differences in ILC2 numbers in either ICOS<sup>-/-</sup> nor ICOSL<sup>-/-</sup> mice versus controls, given the other reports. Having repeated the experiments a number of times we felt that whilst there was no reproducible difference in our hands this was clearly distinct from the findings of others for reasons unclear. It seems likely that ILC numbers will vary between different animal facilities and this may be influencing our findings. We have inserted the following sentence:

‘Whilst these data conflict with previous reports on the requirement for ICOS-ICOSL in ILC2 homeostasis, this may reflect differences arising from the animal facility in which the studies were conducted.’

Question 6: The interpretation of the Cd80<sup>-/-</sup> x Cd86<sup>-/-</sup> experiments is probably overstated. I would be cautious of attributing the ILC phenotype observed (expansion of intestinal ILC2 and ILC3) to unactivated T-cells, especially as ILCs may express CD28 (as well as checkpoint molecules, PD-1, which modulate CD28 signalling). Further experiments are needed to conclusively demonstrate whether the observed shift in ILC composition in Cd80<sup>-/-</sup> x Cd86<sup>-/-</sup> animals is due to a direct effect of loss of CD80/86 interaction with ILCs, or as the authors imply, by loss of T-cell activation. The authors could address this by "relaxing" the interpretation of their findings and by entertaining some alternative possibilities (as described above). This would potentially include changing the title of the manuscript.

Answer 6: We agree with the reviewer and have modified our discussion here and included the following text:

‘Having only assessed ILC populations in total knockout mice, it is also impossible to really separate ILC-intrinsic effects arising from expression of CD28 or microenvironmental effects resulting from changes in T cell activation. To really understand these pathways, novel conditional knockout mice targeting these costimulatory pathways in ILC versus T cells need to be developed.’

**Competing Interests:** No competing interests were disclosed.
Amy E. Saunders
Manchester Collaborative Centre for Inflammation Research, University of Manchester, Manchester, UK

This article provides a detailed comparison of ILC subsets in different tissues including ear skin, lamina propria, lung and various lymph nodes, including analysis of NKp46 and CCR6 expressing subsets of ILC3.

This analysis identified a subset of ILCs that do not express any of the lineage defining transcription factors (TrN cells), which are particularly abundant in skin and lung. The authors have made efforts to show that these cells are ILCs, and their analysis of different digestion protocols suggests that these cells may have lost expression of transcription factors due to the tissue digest procedure. This is an important consideration for other research examining ILCs in these tissues.

The authors also examine the effects of the costimulatory molecules ICOSL and CD80/86 on the subsets of ILCs present in tissues. They show conclusively that the absence of ICOSL does not have major effects on the ILCs present, however the absence of CD80/86 causes an increase in ILC2 and ILC3 populations in the small intestine. It is unclear if the later finding is due to the lack of CD80/86-CD28 interactions on ILCs themselves, or if it is due to the absence of activated T cells.

To improve the clarity of the article I suggest the following changes:

1. The materials and methods could be written more clearly in places. In particular, there is a reference to spleen digestions however, there doesn’t appear to be any data from spleen included in the manuscript. It is unclear if collagenase, or dispase, or both were used to digest lymph nodes and the centrifugation performed is described in rpm, but it is more useful for the reader to quote the force used. For lung and ear digests the final concentrations of enzymes should be clearly stated, as it is currently unclear if the concentrations given are stock concentrations or the final concentrations used in the digests.

I also suggest that the following additions could be made to enhance the article:

1. The data shown here suggest that prolonged tissue digestion, or a high concentration of enzyme used to digest the tissue, leads to a higher proportion of the ILCs being TrN cells, which is likely due to a reduction in transcription factor expression. It would be interesting to ascertain if this is due to the activity of the enzymes directly affecting the transcription factor expression, or if it is due to prolonged incubation at 37 degrees. and this additional information may aid protocol optimization.

2. The authors show that the majority of the CD127 positive, lineage negative cells are Id2 positive, which suggests that these cells are ILCs. However, despite Id2 being required for ILC differentiation, it is not a specific marker for ILCs as other immune cells have been shown to express this transcriptional regulator, and these data are therefore not conclusive, but are suggestive. An approach to more definitively analyse the TrN population would be to perform RNAseq analysis and compare the findings to known immune cell subsets, however this may be outside of the scope of this paper.

3. It would be interesting to digest the lung with the ear digest protocol and then analyse the cell surface markers examined in Figure 3 (CD25, ICOS, KLRG-1, MHCII and ST2), to ascertain if the
digest protocol used affects the expression of these markers, particularly given the large differences seen in expression of these markers between lung and ear skin.

4. The expression of markers such as CD25, KLRG1 and ST2 are shown in this report to be expressed at different levels on ILC2s in skin versus ILC2s in lung and gut. The potential reasons behind this differential expression could be addressed more fully in the discussion.

5. The data showing increased populations of ILC2 and ILC3 in CD80/86 deficient mouse gut, but lower numbers of ILC3s in mesenteric lymph nodes is intriguing, but it is unclear if this is caused by the lack of CD80/86 signals acting directly on ILCs, or if it is due to a lack of activated T cells in the absence of CD80/86. It would therefore be interesting to analyse ILCs in other transgenic mice where T cell activation is impaired to determine if ILC populations are affected by the presence of activated CD4 or CD8 T cells. Given that the ILC3 population is increased in the lamina propria and decreased in the mesenteric lymph node, it would also be interesting to determine if the increase in these ILC populations in the lamina propria is due to increased turnover, or increased trafficking of these cells or their precursors.

Overall this article makes important contributions to the field by describing the ILC subsets in different tissues, highlighting the TrN ILC cell population and demonstrating that the costimulatory molecule ICOSL does not affect ILC numbers, whereas CD80/86 does affect ILC subsets in the gut and mesenteric lymph nodes.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Referee Expertise:** Skin immunology including ILCs present in skin.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

We thank the reviewer for the constructive criticism and have responded to the specific comments below:
Comment 1: The materials and methods could be written more clearly in places. In particular, there is a reference to spleen digestions however, there doesn’t appear to be any data from spleen included in the manuscript. It is unclear if collagenase, or dispase, or both were used to digest lymph nodes and the centrifugation performed is described in rpm, but it is more useful for the reader to quote the force used. For lung and ear digestes the final concentrations of enzymes should be clearly stated, as it is currently unclear if the concentrations given are stock concentrations or the final concentrations used in the digests.

Answer 1: We thank the reviewer for spotting this and we have amended the text in the methods in an effort to more clearly describe what was done. We have omitted the description of spleen cell preparation as this was not used in the data shown.

Comment 2: The data shown here suggest that prolonged tissue digestion, or a high concentration of enzyme used to digest the tissue, leads to a higher proportion of the ILCs being TrN cells, which is likely due to a reduction in transcription factor expression. It would be interesting to ascertain if this is due to the activity of the enzymes directly affecting the transcription factor expression, or if it is due to prolonged incubation at 37 degrees. and this additional information may aid protocol optimization.

Answer 2: We agree with the reviewer that the prolonged tissue digestion seems to reduce expression of the transcription factors. We have provided further data in the revised supplementary figures showing expression of the transcription factors and different surface markers with different digestions of lung tissue. These data indicate that loss of T-bet and Gata-3 expression occur with the prolonged digestion protocol used to isolate cells from the ear. Based on expression of the transcription factors after ex vivo stimulation in other experiments (e.g. PMA/IONO), we think that the loss of expression is not simply a result of culture at 37 degrees and reflects the presence of high concentrations of enzyme. It may be that the harsh conditions in the culture result in some of the cells down regulating these transcription factors rather than a direct effect of the enzymes (which is conceptually easier to consider where surface markers are expressed). The key point that we felt we needed to make is that ILC appearing to lack the three ‘signature’ transcription factors could be enhanced in number through different digestions demonstrating an artificial element to their detection and questioning their relevance in vivo.

Question 3: The authors show that the majority of the CD127 positive, lineage negative cells are Id2 positive, which suggests that these cells are ILCs. However, despite Id2 being required for ILC differentiation, it is not a specific marker for ILCs as other immune cells have been shown to express this transcriptional regulator, and these data are therefore not conclusive, but are suggestive. An approach to more definitively analyse the TrN population would be to perform RNAsseq analysis and compare the findings to known immune cell subsets, however this may be outside of the scope of this paper.

Answer 3: We agree that Id2 is not a specific marker for ILC and is widely expressed in activated T cells. BUT, in this data we had first gated on IL-7Rα+ Lineage- cells and then asked whether Id2 was expressed within this population in the ear where many ‘TrN’ cells were detected. This data provided further evidence that the ‘TrN’ cells might be ILC given other known populations of Id2-expressing cells were gated out by surface markers. We considered this data supportive but not definitive.

Question 4: It would be interesting to digest the lung with the ear digest protocol and then analyse the cell surface markers examined in Figure 3 (CD25, ICOS, KLRG-1, MHCII and ST2), to ascertain if the digest protocol used affects the expression of these markers, particularly given the large differences seen in expression of these markers between lung and ear skin.

Answer 4: We thank the reviewer for this suggestion and have performed these experiments and included the data within the revised supplementary figures (Supplementary Figure 5 and 6). These new data indicate that expression of both CD25 and KLRG-1 are reduced on cells from the lung
when digested with the ear protocol. This then in turn indicates that the ILC2 phenotype reported in the ear is influenced by the digestion protocol and we have tried to make this point more clear in the text. We did try other means of digesting the ear tissue, but the yield was very poor. We have added this text to the discussion:

‘We demonstrate that by increasing the digestion of lung tissue, the frequency of GATA-3\(^+\) ILC2 isolated from the tissue is reduced and the proportion of ILC lacking expression of GATA-3, T-bet and ROR\(\gamma\)t is increased. Thus we conclude that whilst ILC lacking expression of the three group defining transcription factors may exist, care must be taken in interpreting data from digested tissue and this population can certainly be ‘contaminated’ by ex vivo changes to known ILC populations during their isolation and analysis. Whilst ILC progenitor populations within different tissues remain to be fully characterised, these observations highlight how the manner in which different tissues are digested can impact on our ability to properly identify different ILC populations. 

Whilst we have tried to phenotype ILC populations isolated from different tissues, our data again highlights the potential caveats to studying phenotypic markers following substantial digestion of tissue. Whilst we could detect a robust ILC2 population in ear skin following the digestion protocol used, we are left unable to comment on the expression of CD25 and KLRG-1 at the protein level due to clear loss of these markers during cellular isolation. We do provide evidence that ILCs can still be clearly distinguished since expression of key lineage markers were not lost. ’

Question 5: The expression of markers such as CD25, KLRG1 and ST2 are shown in this report to be expressed at different levels on ILC2s in skin versus ILC2s in lung and gut. The potential reasons behind this differential expression could be addressed more fully in the discussion.

Answer 5: As outlined above, the experiment suggested by the reviewer provides evidence that the low CD25 and KLRG-1 expression is due to the digestion of the ear and we have made this clearer in the text. We did not find evidence that ST2 expression is lost during the digestion, indicating that ILC2 in the ear skin do indeed lack ST2 expression.

Question 6: The data showing increased populations of ILC2 and ILC3 in CD80/86 deficient mouse gut, but lower numbers of ILC3s in mesenteric lymph nodes is intriguing, but it is unclear if this is caused by the lack of CD80/86 signals acting directly on ILCs, or if it is due to a lack of activated T cells in the absence of CD80/86. It would therefore be interesting to analyse ILCs in other transgenic mice where T cell activation is impaired to determine if ILC populations are affected by the presence of activated CD4 or CD8 T cells. Given that the ILC3 population is increased in the lamina propria and decreased in the mesenteric lymph node, it would also be interesting to determine if the increase in these ILC populations in the lamina propria is due to increased turnover, or increased trafficking of these cells or their precursors.

Answer 6: We agree with the reviewer that further analysis of ILC in different mice lacking costimulatory molecules might be interesting as well as exploring the mechanisms behind such changes. We think this is beyond the scope of this current study and perhaps conditional knockout mice may be more informative should the models become available.

**Competing Interests:** No competing interests were disclosed.