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CD4+ Teff cell heterogeneity:

The perspective from single-cell transcriptomics

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Abstract

Single-cell transcriptomics (scRNAseq) holds the promise to generate definitive atlases of cell types. We review scRNAseq studies of conventional CD4⁺ $\alpha\beta$ T cells performed in a variety of challenged contexts (infection, tumor, allergy) that aimed to parse the complexity and representativity of previously defined CD4⁺ T cell types, lineages and cosmologies. With a few years' experience, the field has realized the difficulties and pitfalls of scRNAseq. With the very high-dimensionality of scRNAseq data, subset definitions based on low-dimensionality marker combinations tend to fade or blur: cell types prove more complex than expected; transcripts of key defining transcripts (cytokines, chemokines) are distributed as broad and partially overlapping continua; boundaries with innate lymphocytes are blurred. Tissue location and activation, either cytokine- or TCR-driven, determine Teff heterogeneity in sometimes unexpected ways. Emerging techniques for lineage and trajectory tracing, and RNA-protein connections, will further help define the space of differentiated CD4⁺ T cell heterogeneity.

Introduction

CD4⁺ T cells initiate and orchestrate immune responses to a wide array of pathogens by detecting non-self antigens, help humoral and cellular responses, orchestrate tissue repair, and have even more esoteric effects on the homeostasis of organs systems outside the classic confines of Immunology. This range of functions and response abilities has long raised the question of their heterogeneity: are diverse functions all mediated by one multifunctional cell, or do distinct subsets possess specific effector functions, like helping B cell responses or provoking delayed-type hypersensitivity. Single-cell transcriptomics holds the promise to reopen this long-standing question from a radically different perspective and to provide a genome-scale parsing of true identities in the CD4⁺ T cell subsets.

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Declaration of interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Almost half a century ago, it was shown that different T cell clones can have different functions, with a division of labor between cells responsible for cell-mediated immunity and for help to antibody responses [1,2]. Mosmann and co-workers [3] were the first to classify different CD4⁺ T cell clones by the cytokines they produce, a fundamental notion that is followed to this day, and spread to classifications of other immunocytes such as innate lymphoid cells (ILCs), gamma delta T cells or non-conventional T cells. They analyzed independent T cell clones in culture and noted that their cytokine secretion profiles could distinguish them in relation to their functional ability [3]: Th1 cells responsible for cellular immunity against intracellular pathogens, Th2 cells that orchestrated the antibody response by B cells [4]. It was soon realized that naïve T cells could also be induced to adopt these biased cytokine-producing phenotypes by culture with cocktails of inducing cytokines [5].

The CD4⁺ T effector (Teff) cosmology has since expanded beyond the Th1/Th2 dichotomy, which was quickly found to be too constraining, with identification of Th17, Tfh, Th9, or Th22 (and further "sub-subsets" thereof, like Th17*, Tfh2, etc), and with the reallocation of several functions (i.e. Tfh are now seen as the main B cell helpers). These distinctions remain primarily based on cytokine but now bolstered by "master" transcription factors which are thought to anchor these programs[6]. Moreover, sub-subsets such as pathogenic and non-pathogenic or homeostatic and inflammatory Th17s have been identified[7–9]. Conventional gene expression profiling of *in vitro* polarized T cell cultures were used to identify differentially expressed signatures[10–14], or differential epigenetic marks and chromatin states[15,16], but usually from cells derived in biasing tissue culture, rather than from *ex-vivo* cell populations. Importantly, it has proven impossible to identify cell surface markers that reliably and consistently identified Th-x subsets (some chemokine receptors have been proposed, but proved not to have unequivocal expression).

In hindsight, was there an obligate reason why cytokine expression had to be the organizing principle for CD4⁺ T subset distinction? Cytokines are certainly essential to T cell function, but other functionally important molecules might have served. Indeed, an orthogonal frame of reference for Teff heterogeneity was developed, based primarily on organismal location and the expression of chemokine receptors and adhesion molecules that drive it[17,18], distinctions that have direct relevance to the rapid detection of pathogens or inflammation states in the tissue where they occur. As for CD8⁺ T cells, CD4⁺ T cells have been distinguished as T_N , T_{CM} , T_{EM} , T_{RM} (but also less frequently used categories like T_{EMRA} , T_{PM} , etc[19], a classification orthogonal to the Th-x nomenclature and that also relies on very few (low dimensionality) flags.

Against this background, several groups have begun to apply single-cell (sc) RNAseq to probe T cell populations more deeply in the context of various challenges, to chart the relationships between subsets, to search for unrecognized heterogeneity within subsets, or perchance to discover entirely new planets in the CD4⁺ system. We will focus here on effector CD4 T cells but occasionally refer to regulatory T cells and CD8⁺ T cells for comparisons or parallels.

Caveats and pitfalls

Several good recent reviews have discussed the revolutionary potential of single-cell RNAseq in parsing the complexity of cell-types and -states[20–22]. Overcoming the averaging that occurs during gene expression profiling of sorted cells, scRNAseq has the potential to reveal hidden heterogeneity in cell populations, characterize distinct cell subsets, and map the trajectories of cell differentiation. So, much exciting promise.

The field having a few years' experience, it is important to avoid rose-colored glasses and be conscious of the caveats and pitfalls of scRNAseq. Not so much the question of its sensitivity, as one still hears occasionally: low abundance transcripts are indeed difficult to analyze robustly in individual cells, owing to the high probability of technical dropouts as well as biological noise in the low expression range. But the scRNAseq process (in its different variants) is actually remarkably faithful: aggregation of cells into microclusters, and/or imputation of missing values[23–26], yields profiles akin to conventional RNAseq, showing that rare transcripts are correctly detected.

A key obstacle is that scRNAseq data are simply difficult to grasp and handle: defining a cell-type based on a multi-dimensional transcriptome is more difficult, conceptually and experimentally, than from a handful of flow cytometry markers. Computational pipelines have been developed to process, verify, and analyze scRNAseq data and are well codified by now (e.g. Seurat[27]). But the follow-up, the interpretation of the results is far less codified and can be misleading. For instance, 2-dimensional projections (tSNE, UMAP) are commonly used to summarize and make human-readable multidimensional complexity. But those maps are by no means to be taken as gold-standard representations of the reality (play with https://distill.pub/2016/misread-tsne/ for sobering examples). Similarly, partition clustering tools produce groupings (e.g. the colored outputs of common processing pipelines) that may, or may not, have anything to do with biological reality. Clustering tools accurately distinguish distant cells, neurons from hepatocytes, B from T cells, but are far less trustworthy when trying to parse closely related cells - one should remember that a clustering algorithm will *always* return clusters (conditioned by its set parameters), even when there aren't any biologically compelling or rational ones [28]. Inferring precursorproduct relationships between cells using the early "trajectory" tools can be hazardous (fine to map intermediates in a transition sequence that is known beforehand, haphazard if trying to infer an unknown relationship)[29]. Annotation of cell types (regions in a 2D projection, or clusters) is surprisingly challenging; quick and superficial annotation based on recognizing a handful of favorite transcripts is too often used, but rigorous annotation based on defining signatures is much more challenging[30]. Finally, the common experience is that distinction between cells that one would imagine should be readily distinguished (e.g. NK and T cells) are often not straightforward: T cells and ILCs differ fundamentally by the rearrangement and expression of TCR genes, but much less when the rest of the functional transcriptome is taken into account. Whether a cell's identity should be defined by one or two key functional genes that deeply condition the cell's function, or by the fingerprint of its entire transcriptome, is a question that the field is still grappling with.

Did scRNAseq reveal novel CD4+ T cell-types: "Tis"

Discovering new cell-types is a long-established tradition in Immunology, and a novel CD4+ T cell would be a valuable trophy. In other domains, scRNAseq has allowed the discovery of rare but important cell types, primarily epithelial: CFTR-expressing ionocytes in the lung surface, tuft-like cells in the thymic epithelium[31–33]. For CD4⁺ T cells, there may be one such revelation: in several scRNAseq studies, a minor but very distinct population of CD4⁺ T cells showed an inordinately high expression of Interferon-stimulated Genes (ISGs), with no other notable characteristic. These cells (hereafter "Tis") (for T interferon signature) were observed in a range of immunologic challenges: in airways after allergic sensitization with house dust mites or Alternaria extract[34,35], in kidney infiltrates of lupus nephritis[36], or in colon lamina propria of Salmonella- or Citrobacter- infected mice[37]. With the exception of the lupus kidney, these are not contexts where Type-1 interferon would be expected. Tis were rare or absent in lymphoid tissues at baseline, however, indicating that they are induced or summoned during active immune defenses. The origin of their high ISG expression is mysterious. Their Interferon (IFN) signaling pathway may be unusually sensitive; Tibbitt et al. [34] showed that the Type-I IFN receptor is not over-expressed in Tis, but other signaling molecules might be involved (e.g. MyD88). Alternatively, Tis cells may reside in microcompartments that contain high levels of IFN, a reaction that can occur in any tissue. Interestingly, though, a similar population was also seen observed in thymic CD4+ singlepositive, suggesting that Tis may differentiate as an independent branch in the thymus [38]. Finally, the interferon-activated transcriptional module may have been activated in Tis but never switched off. Tis function is totally unknown (and may prove difficult to explore, given their characteristics and rarity), but one might speculate that constitutively high ISG expression may have pathological consequences (e.g. participating in lupus pathogenesis?).

Discrete subsets or a continuum of states?

Several studies took the unbiased approach of analyzing the entire pool of CD4⁺ T cells in a given location, in the context of a tumor or some immune challenge[34,36,37,39–41]. A strong effect of location was commonly observed, an anticipated conclusion based on much conventional profiling.

Ciucci et al.[41] performed scRNAseq on splenic activated (CD44^{hi}) T cells, 7 days after LCMV infection, polyclonally or after tetramer selection of cells reacting to the immunodominant GP66 peptide. Although polyclonal splenocytes appear to split into clusters according to conventional Th1/Tfh/Tmem phenotypes, the GP66-specific population proved more continuous, with few or no well-defined and reproducible clusters (perhaps an illustration of issues with default clustering and coloring), and with overlapping representations of transcripts classically used to identify T subsets (*Tbx21, Bcl6, Ccr7*). In a study of activated CD4⁺ T cells from bronchoalveolar lavage after house dust mite re-challenge[34], clustering only poorly resolved cell subsets (other than Tregs and Tis). Cells producing IL13 and IL5 were predominantly found in one corner of the tSNE plot, but the expression of *Gata3* (highly represented) or *Tbx21* (less common) was spread among the clusters. Similarly, in a study of colon lamina propria T cells after challenge with a panel of bacteria or parasites known to induce Th1, 17 or 2 biased responses, Kiner et al.[37] found

that Teff could not be parsed into discrete Th cell clusters, even with pliable artificial intelligence tools, cytokine and other signature transcripts being distributed along biased but continuous gradients. Variance in expression was tied to that infecting pathogen, more than to the cytokines produced or to any classic Th signature. Thus, in these several conditions where pathogenic challenges might have been expected to elicit distinct Th-x subsets, more complex populations of responding CD4⁺ T cells were observed [34,37,41]; cell distributions were biased in accordance with the challenge, but with far "messier" representations of key transcripts, and no distinct cell-types.

Several studies undertook single-cell analyses in tumors in order to study tumor heterogeneity as well as the heterogeneity of the immune response to tumors. Zheng et al. [39] profiled single T cells from hepatocellular carcinoma patients and found several clusters amongst CD4⁺ T cells, including naive CD4s, Tregs, Teffs, exhausted T cells, as well as cytotoxic CD4⁺ T cells. As expected from conventional profiling, they observed a strong tissue effect that differentiated the transcriptomes of peripheral blood T cells and tumorinfiltrating T cells. Azizi et al.[40] profiled total CD45⁺ cells from breast tumor patients and described 21 CD4⁺ clusters, which were split into 9 naive, 7 central memory, 15 effector memory, and 5 Treg clusters (over-clustered?). Tumor-specific signatures were not found in discrete cell state clusters, but rather along "gradual trends of variation", or a single continuous trajectory.

Other scRNAseq studies drilled down into subsets of T cells, identified by knockin reporters driven by cytokine-encoding genes and/or marker combinations, aiming to uncover further heterogeneity within CD4⁺ T cell subtypes. In a pioneering study, Gaublomme et al.[42] used scRNAseq on IL17-producing cells derived in vitro and from the central nervous system of EAE-bearing mice. This allowed for the unbiased characterization of cell-states within Th17 cells, which had been previously described to vary phenotypically, with regard to their exposure to IL23 and on their pathogenicity[8]. Indeed, variation within Th17s was observed, in vivo and in vitro, but as a phenotypic continuum rather than discrete subsubsets. The variation could be decomposed into several components: overall activation; a module containing classic Th1 transcripts including Ifng, and one linked to the IL23 signature or pathogenicity. These results were paralleled by IL17-producing CD8⁺ T cells in the skin, after challenge with S. epidermis [43] where a significant proportion of the cells could also co-express type 2 cytokines. Two other studies similarly revealed heterogeneity within IL10-producing CD4⁺ T cells in anti-CD3 treated or LCMV infected mice[44,45]. Xin et al. showed a dominant population of IL10-producing cells with several characteristics related to Th1 cells (*Tbx21, Id2*) but also an outlier cluster with classic Tfh transcripts (*II21*, Cxcr5, Bcl6). Building on the scRNAseq data, elegantly constructed mixed bone-marrow chimeras were then used to show the functional relevance of these IL10-producing Tfh in the humoral response to LCMV. These studies, focused on cells making an individual cytokine, illustrated that cytokine production does not result from a single and discrete cell population.

Gowthaman et al.[35] took apart the Tfh subset (strenuously purified as CD44^{hi}PD1^{hi}CXCR5^{hi}) elicited in an intranasal model of allergic sensitization. What might have been expected to be a well-defined group of cells actually contained a number of distinct cell-types: some Tis cells (confirming that the Tis phenotype cuts across marker-

identified subsets), groups of cells superficially annotated as IL4-producing Tfh2, and a small but distinct group of cells with dual production of IL4 and IL13 (follow-up experiments showed that IL13+ cells were indeed essential for allergic sensitization and anaphylactic IgE production; other allergy models may use other phenotypes for Type 2 cytokine production, however). In a similar vein, Patil et al.[46] sorted classic fractions of effector/memory cells (Tcm, Tem, Temra) from human donors originating from different continents, also based on low dimensionality flow cytometry criteria (CD45RA and CCR7). The Temra pool proved to contain several different clusters, although it was not clear how much of the clustering was driven by inter-donor differences vs true cell-states. Some of these clusters were hypothesized to represent effector or precursor pools, based on the differential representation of some homeostatic or precursor transcripts, although these relationships remain to be substantiated experimentally. Thus, both these studies illustrate that previously defined cell-types that rest on limited marker combinations actually hide far more extensive complexity when analyzed in the multidimensional transcript space.

Taken together, these results imply that Teff states are non-discrete, vary between tissues of residence, and are strongly influenced by cytokine- or TCR-induced activating signals. The variation in gene expression between T cells distributes as broad continua, rather than as discrete entities. Conversely, populations that were thought to be well-defined turned out to be more complex than anticipated. We have proposed elsewhere[37] a "Cloud Model" for Teff cell heterogeneity, wherein cytokines or other key functional or regulatory molecules are expressed along partially overlapping gradients, rather than in discrete Th-x entities. This concept is different from the "plasticity" that has been often evoked over the last decade, in that plasticity implies defined states or lineages between which T cells could switch. The Cloud Model raises conceptual difficulties. It is easier to think of interactions between well-defined and consistent cell types than between proteoform cell populations, in which the expression of any one effector or regulatory molecule is loosely distributed. It raises practical difficulties for experimentation: for instance, it was comfortable to sort CCR6+ cells to test their activity and properties, under the assumption that they represented a coherent functional cell entity. But how does one sort a segment of a cloud?

Do variations in TCR clonotypes drive CD4+ T heterogeneity?

T cells see the world through their TCR. Other factors (inductive influences in specific tissue locations, microbe-derived ligands, cytokines, etc) can sway T cell phenotypes, but a key question is how much of the phenotypic choice rests in the characteristics of signals caused by TCR engagement (affinity, avidity, duration). ScRNAseq, and in particular its recent advances[40,47,48], allows the determination of TCR sequences (both chains) while evaluating the cells' transcriptomes. Several scRNAseq studies have addressed these questions, in localized settings (tumors, infection, immunization) where clonotype expansion allows the comparison of several cells that express the same TCR (noting that this focus on expanded clones may introduce an experimental bias).

The overall conclusion is that the TCR does influence phenotypes, but to varying degrees according to location. Very demonstrative were the results of Azizi et al. in breast cancer infiltrates, where cells expressing the same $\alpha\beta$ TCR clonotype tended to occupy confined

regions of the tSNE projection, although to variable extent for different TCRs[40,49]. A multiple regression procedure estimated that 30–50% of the T cell diversity could be explained by TCR clonotype. Similarly, among ~3,000 CD4⁺ T cells infiltrating basal cell carcinoma tumors, Yost et al. found no clonotypic overlap between cell regions annotated as Treg, Tfh and Th17, contrary to some CD8 subsets that transitioned from memory or effector to activated phenotypes[50]. On the other hand, Pratama et al.[51] reported some sharing of TCR clonotypes between Treg and Tconv pools in the colonic lamina propria of ex-germ-free mice colonized with a single microbe. Several of the highly represented clones evidenced by Tu et al.[48] after immunization with HPV-E7 were also spread throughout the phenotypic space. Overall, these studies show that the TCR is not the only driver of T cell heterogeneity but definitely an important factor.

Mechanistically, TCR signals may control the transcriptional programs directly or indirectly. Early TCR signaling is almost always the most variable gene module in single-cell surveys of T cell populations [42,44,45,49,52,53]. Zemmour et al.[49] tested the connection between signaling intensity and phenotypes in Tregs, using Nr4a1-gfp reporter mice in which the reporter intensity was directly related to the strength of TCR signals[54]. Cells belonging to different windows of GFP intensity were analyzed by scRNAseq. GFP levels proved highly related to the position of the cells in the multidimensional phenotypic space. These results suggest that TCR signals are intrinsically related to phenotypes. On the other hand, the same range of phenotypes was found in Treg cells of the adipose tissue, whether they were polyclonal or expressed a single transgenic TCR [53], showing that extrinsic cues can dominate in some contexts.

What lies ahead?

Single-cell transcriptomics is a recent and very fast-moving technology, and novel approaches to study T cells at a single-cell level are continuously emerging. "CITE-Seq" uses DNA-tagged antibodies that recognize molecules on the cell surface, and the tags are then quantitated in the same microfluidic process as mRNAs[47,55,56]. This allows the simultaneous quantitation of surface proteins and mRNAs and can be scaled to tens or even hundreds of cell surface proteins. CITE-seq promises to be invaluable for T cell biology, connecting complex structures in the transcriptome with the surface markers that can be used to select cells for functional experiments. Along the same lines, tagging reagents that enable input multiplexing ("hash-tags") can be used for creative experimental designs, combining controls with experimental variants, time- or dose-shifted points, to enable direct comparisons and alleviate dreaded batch effects [37,47,55]. Single-cell analysis of chromatin accessibility is also coming of age[57], with new versions that associate chromatin and transcriptome data on the same cells. T cell differentiation and heterogeneity can be probed at the more "primordial" chromatin level, free of confounders from variation in RNA stability or transcriptional bursting. Finally, spatially-resolved transcriptomics (e.g. [58,59]) will shed light on the locations within organs where T cells of particular phenotypes live (building here on an already extensive knowledge from multi-color imaging), and on transcriptional changes happening in T cells in situ. One should acknowledge that spatial transcriptomics may be of more value to study immobile cells (neurons) than for T cells that are highly mobile, constantly probing other cells in search of excitement. Finally, new

experimental supports for cell and lineage tracing (e.g.[60–63]), and more performant computational tools[29] should build on early explorations [64] and allow a robust analysis of differentiation trajectories, what paths and fluctuations are followed by a T cell during activation, holding a key for understanding the molecular underpinnings of T cell differentiation and immune or autoimmune responses

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- Single-cell transcriptomics can reveal unrecognized heterogeneity and CD4 T subsets.
- An unexpected CD4+ T population with high levels of IFN-I responsive transcripts.
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- The end of Thx: continuous, rather than discrete, distribution of key cytokines.
- Phenotypes are driven by TCR signals, but not solely.