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The transcription factors T-bet and GATA-3 control alternative pathways of T-cell differentiation through a shared set of target genes

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Upon detection of antigen, CD4+ T helper (Th) cells can differentiate into a number of effector types that tailor the immune response to different pathogens. Alternative Th1 and Th2 cell fates are specified by the transcription factors T-bet and GATA-3, respectively. Only a handful of target genes are known for these two factors and because of this, the mechanism through which T-bet and GATA-3 induce differentiation toward alternative cell fates is not fully understood. Here, we provide a genomic map of T-bet and GATA-3 binding in primary human T cells and identify their target genes, most of which are previously unknown. In Th1 cells, T-bet associates with genes of diverse function, including those with roles in transcriptional regulation, chemotaxis and adhesion. GATA-3 occupies genes in both Th1 and Th2 cells and, unexpectedly, shares a large proportion of targets with T-bet. Re-complementation of T-bet alters the expression of these genes in a manner that mirrors their differential expression between Th1 and Th2 lineages. These data show that the choice between Th1 and Th2 lineage commitment is the result of the opposing action of T-bet and GATA-3 at a shared set of target genes and may provide a general paradigm for the interaction of lineage-specifying transcription factors.

Genomic map | T helper differentiation | cytokines

The immune response to different pathogens is tailored by the differentiation of CD4+ T helper cell into different effector types (1, 2). Th1 cells are critical for protection against viruses and intracellular pathogens, Th2 cells for the removal of extracellular parasites, and Th17 cells function in the response to extracellular bacteria. T-cell lineage commitment also impacts upon numerous diseases processes, with defects in Th1 and Th17 responses implicated in autoimmunity and Th2 responses in the pathogenesis of allergic disease (3).

Alternative pathways of T-cell differentiation occur through the action of a number of transcription factors. Generation of Th1 cells from naïve precursors requires IL-12 and IFN-γ signaling that leads to activation of STAT4 and STAT1, respectively (1). STAT1 induces expression of T-bet (TBX21) that acts as a key regulator of Th1 cell fate determination (4). T-bet activates Th1 genetic programs whilst simultaneously suppressing Th2 and Th17 programs (5, 6). Loss of T-bet induces default commitment to Th2 and Th17 lineages and T-bet deficient mice have impaired Th1 immunity, are resistant to autoimmune disease and develop spontaneous asthma (7–10).

Generation of Th2 cells requires IL-4, which leads to STAT6 phosphorylation (11) and upregulation of GATA-3, the key regulator of Th2 development (12, 13). Deletion of GATA-3 in peripheral CD4+ T cells prevents differentiation into the Th2 lineage, causing cells to differentiate toward a Th1 phenotype in the absence of polarizing cytokines (14, 15). Conversely, overexpression of GATA-3 in Th1 cells switches their polarity to a Th2 phenotype (15). Interestingly, it has been shown that GATA-3 is also expressed at lower levels in murine Th1 cells and that this expression is necessary for the subsequent induction of a Th2 response via STAT5 (15). We have also reported an interaction between GATA-3 and tyrosine-phosphorylated T-bet in murine thymocytes (16).

Our knowledge of T-bet and GATA-3 function comes mainly from their roles at gene loci that encode IFNG and IL4/IL5/IL13, the signature Th1 and Th2 cytokines, respectively. Th1 differentiation is accompanied by acetylation of histone H3 lysine 9 (H3K9ac) at the IFNG locus and this process is dependent on T-bet (17–23). Conversely, Th2 differentiation is accompanied by hyperacetylation of the IL4/IL5/IL13 locus, dependent on GATA-3 (18, 19, 23–26). Recent murine studies also show that T-bet directly represses the expression of IL4 (23, 27) and that GATA3 directly represses IFNG (22, 28). This suggests that T-bet and GATA-3 may act to promote alternative pathways of T-cell differentiation by acting on the same target genes. However, because only a handful of genes are known to be directly targeted by T-bet and GATA-3 in primary T cells and since Th1 differentiation appears to be only partly dependent on IFN-γ (1), the mechanism by which these two factors direct alternative cell fates remains unclear.

Given the profound influence that T-cell lineage commitment has upon numerous disease processes, it is critical to understand the regulatory mechanisms that contribute to these postdevelopmental cell fate choices in humans. To determine the mechanisms of human T-cell lineage commitment, we have identified the target genes of T-bet and GATA-3 in primary human Th1 and Th2 cells. This work identifies a number of novel pathways with critical relevance to T-cell biology and reveals that master regulator transcription factors can act through a shared set of target genes to control alternative cell fates.

Results

T-bet and GATA-3 target genes in primary human T cells. To identify genes directly targeted by T-bet and GATA-3 during early human T cell differentiation, we generated Th1 and Th2 cells from primary naive human T-cells and performed chromatin immunoprecipitation coupled with microarray analysis (ChIP-Chip) (29). We first verified that our Th1 and Th2 cells were appropriately polarized. Human Th1 cells expressed relatively high amounts of T-bet and IFN-γ mRNA and low amounts of IL-4 and GATA-3 and the opposite was the case for Th2 cells (Fig. S1). Only a proportion of human Th1 cells expressed IFN-γ
Fig. 1. T-bet associates with many genes in Th1 cells. (A) Composite T-bet enrichment profile in Th1 cells for genes that show significant binding within 4 kb from their transcription start site (green), compared with the average binding profile for T-bet at the same genes in Th2 cells (blue) and an IgG control antibody ChIP in Th1 cells (black). The plot shows average fold-enrichment (normalized signal from ChIP-enriched DNA divided by the signal from whole cell extract (WCE) DNA. The start and direction of transcription of the average gene is indicated by an arrow. (B) T-bet ChIP signals at IFNG and RUNX1 in Th1 and Th2 cells. The plots show unprocessed enrichment ratios for all probes within a genomic region (ChIP vs. whole genomic DNA) for T-bet in Th1 cells (green), T-bet in Th2 cells (blue), and for an IgG control in Th1 cells (black). Chromosomal positions are from NCBI build 35 of the human genome. Genes are shown to scale below and aligned with the plots by chromosomal position (exons are represented by vertical bars, the start and direction of transcription by an arrow).

protein but almost all cells expressed T-bet, confirming the cells' Th1 phenotype (Fig. S1) and consistent with previous results (30). Th2 cells did not express T-bet but did express CRTH2 (Fig. S1). GATA-3 could be detected in both Th2 and Th1 cells, consistent with previous studies (Fig. S1) (31–33).

**T-bet associates with many genes in Th1 cells.** We performed ChIP for T-bet followed by hybridization to microarrays containing probes for 8kb surrounding the transcription start sites of 18,450 protein-coding genes. Using an error model to analyze data from replicate experiments (29) and using cells from different donors, we detected T-bet at the promoters of 832 protein-coding genes in primary human Th1 cells (Table S1 and Fig. S2). T-bet targets could be confirmed by quantitative PCR (Fig. S2), were not enriched by ChIP with an isotype-matched control antibody in Th1 cells (Fig. L4 and Fig. S2) and, as expected, did not exhibit significant T-bet binding in Th2 cells (Fig. L4 and Fig. S2), providing confirmation that these genes are specific targets of T-bet.

The average binding profile shows T-bet binding sites tend to be distributed around the transcription start site (Fig. L4). The T-bet binding profile peaks just upstream of the transcription start site with a shoulder into the gene. Examining the T-bet binding profile at individual genes, it was apparent that T-bet usually shows two or more distinct

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**Fig. 2. T-bet activates the expression of its target genes.** (A) Relative levels of expression of T-bet target genes in primary stimulated Th1 cells compared with Th2 cells (column 1) and in secondary stimulated Th1 compared with Th2 cells (column 2). Each row represents a single gene. Genes are ordered by their relative expression level in Th1 cells versus Th2 cells and this average expression ratio is shown on the left hand side. The colors represent relative gene expression levels between Th1 and Th2 cells, with shades of red indicating higher expression in Th1 cells, black equal expression levels and shades of green indicating higher expression in Th2 cells. **(B and C)** T-bet ChIP signals at NKG7 and CCL4, in human Th1 cells (green) and Th2 cells (blue), and signal from an IgG control IP in Th1 cells (black). Details as for Fig. 1B. (C and F) Real time PCR for NKG7 and CCL4 in WT (open bars) and T-bet−/− (filled bars) murine T cells. Cells were unstimulated (US), stimulated with anti-CD3 antibodies (αCD3), or stimulated with anti-CD3 antibodies and anti-CD28 antibodies (αCD3/αCD28). RNA abundance is relative β-actin RNA. (D and G) Real time PCR for NKG7 and CCL4 in CD4+ T cells from T-bet−/−× IFN-γ−/− murine CD4+ T cells transduced with empty vector (open bars) or T-bet (filled bars) expressing retrovirus. (H) Percentage of human Th1 genes whose orthologs are upregulated (at least 1.5-fold, black bars) or downregulated (white bars) in murine T-bet−/−× IFN-γ−/− CD4+ cells upon expression of exogenous T-bet. Th1 genes were defined as those expressed at least 2-fold or higher in Th1 cells than Th2 cells and divided into those bound by T-bet or those not bound by T-bet.
binding peaks, with one peak situated at, or upstream of, the transcription start site and the other in the first or second intron (Fig. 1A and B and Fig. S2).

One T-bet target we identified was IFNG, providing confidence in our dataset (Fig. 1B). T-bet also bound the TNF-α promoter, suggesting a mechanism to ensure co-ordinate regulation of Th1 cytokine production and consistent with regulation of this gene by T-bet in colonic DCs (34). We also detected binding to the promoters of a number of genes with critical functions in T-cell trafficking (CCR5, ITGAL, SELPG, and ICAM1), consistent with our data showing that T-bet controls a specific T-cell migratory program (35). We also detected binding to the promoter of RUNX1 (Fig. 1B). Overexpression of RUNX1 promotes Th1 differentiation by repressing GATA-3, and there is evidence that it may also bind to the IL-4 silencer (23, 36, 37).

To gain a better impression of the cellular functions regulated by T-bet, we used gene ontology (GO) to assign functions to our set of T-bet target genes. This revealed that the set of genes targeted by T-bet were enriched for those involved in metabolism (369 genes, $P = 6.4 \times 10^{-8}$), RNA processing (34 genes, $P = 2.4 \times 10^{-4}$), protein localization (43 genes, $P = 6.6 \times 10^{-8}$), and transcription from RNA polymerase II promoters (37 genes, $9.9 \times 10^{-4}$). Looking across all GO terms, we identified 100 T-bet target genes with roles in transcriptional regulation, including AIF4, BCL6, CREB1, and Ifi16. These functional clusters suggest previously unappreciated roles for T-bet in the biology of Th1 cells. The many transcriptional regulators targeted by T-bet likely form part of the transcriptional regulatory network that operates downstream of T-bet in Th1 cells.

**T-bet Activates the Expression of Th1 Genes.** T-bet is known to directly activate expression of IFN-γ to promote Th1 differentiation (4, 17–23).

To determine whether the human T-bet gene targets were generally more highly expressed in Th1 cells than Th2 cells, we generated expression data from primary and secondary-stimulated human naïve CD4+ T cells skewed to either Th1 or Th2 lineages (Fig. 2A). We found that although the majority of T-bet target genes showed similar expression levels between Th1 and Th2 cells, a number of genes showed increased gene expression in the former (Fig. 2B). D-16 ChIP dark blue, HG3–31 ChIP light blue. (C) Examples of genes with functions in Th1 cells that are bound by both T-bet and GATA-3. The plots show unprocessed enrichment ratios for all probes within a genomic region (ChIP vs. whole genomic DNA) for T-bet in Th1 cells (green) and GATA-3 (D16 ChIP) in Th2 cells (blue). (D) Examples of genes with functions in Th2 cells that are bound by both T-bet and GATA-3. Details as for C.

![Fig. 3. GATA-3 gene occupancy in Th2 cells. (A) Examples of GATA-3 ChIP signals in Th2 cells. The plots show unprocessed enrichment ratios for all probes within a genomic region (ChIP vs. whole genomic DNA). D-16 antibody ChIP dark blue, HG3–31 antibody ChIP light blue. (B) Composite GATA-3 enrichment profile in Th2 cells for genes that show significant binding within 4 kb from their transcription start site. D-16 ChIP dark blue, HG3–31 ChIP light blue. (C) Examples of genes with functions in Th1 cells that are bound by both T-bet and GATA-3. The plots show unprocessed enrichment ratios for all probes within a genomic region (ChIP vs. whole genomic DNA) for T-bet in Th1 cells (green) and GATA-3 (D16 ChIP) in Th2 cells (blue). (D) Examples of genes with functions in Th2 cells that are bound by both T-bet and GATA-3. Details as for C.](https://www.pnas.org/cgi/doi/10.1073/pnas.0909357106)
transduction with retroviral vectors carrying either a T-bet and a GFP transgene or a GFP transgene only, sorted for GFP expression, quantified using DNA microarrays and gene orthologs mapped between mouse and human using Homologene. We found that T-bet target genes activated in human Th1 cells were often induced upon expression of T-bet in mouse, when compared to Th1 genes that are not associated with T-bet \( P < 0.01 \) (Binomial), Fig. 2A. Genes directly activated by T-bet in this manner included IFNG, CCL4, NKG7, PRF1, TNF, IL18RAP, IL2RB, SETBP1, and PRDM1. We also identified a smaller number of genes that were bound by T-bet, including IL4, PTGER4, FOXP3a, and CD28. Therefore, T-bet directly regulates the expression of a number of genes in Th1 cells in addition to IFNG.

**GATA-3 Occupies Immune Response Genes in Th2 Cells.** GATA-3 is the lineage determining transcription factor for murine Th2 cells and seems to perform similar functions in human T cells (38). To identify GATA-3 target genes in Th2 cells, we performed ChIP with a GATA-3-specific antibody (D16) and hybridized the enriched DNA fragments to microarrays. Using our error model, we identified 344 GATA3 target genes in Th2 cells (Fig. 3A, Table S1, and Fig. S4). We also performed replicate experiments using cells from a different donor and with an alternative GATA-3 antibody (HG3–31). This antibody gave a lower enrichment of GATA-3 in both Th1 and Th2 cells (Fig. 3B, Table S2, and Fig. S5). In a pattern similar to T-bet, the average GATA-3 binding profile showed GATA-3 binding peaks occurred just upstream of the transcription start site with a second peak occurring within the gene (Fig. 3B). Examining individual gene plots, some genes showed one peak at the transcription start site (for example ITK), others two peaks (e.g., STAT6) or three peaks (e.g., IL4) (Fig. 3).

GATA-3 and T-bet both target the genes encoding IFN-\( \gamma \) and IL4/IL5/IL13 in mouse cells (22, 23). We therefore compared the sets of T-bet and GATA-3 target genes to investigate whether these factors generally target the same genes in human cells. We identified 76 genes that were associated with both T-bet in Th1 cells and GATA-3 in Th2 cells (Fig. 3C and D). These genes included those with both Th1 and Th2 functions and expression patterns (Fig. 3C and D and Fig. S5). For example, GATA-3 and T-bet co-occupied both the Th1-associated genes IFNG, CCL4, IL18RAP, and TNF as well as the Th2-associated genes IL4, PTGER4, ITK, and NFATC2. These results indicate that T-bet and GATA-3 induce alternative T-cell differentiation pathways by acting on many of the same genes.

**GATA-3 Occupies Its Target Genes in Th1 Cells.** Assuming that GATA-3 would not associate with genes in Th1 cells (as we saw with the absence of T-bet binding to promoters in Th2 cells), we performed a ChIP-Chip analysis in Th1 cells as a putative negative control. Remarkably, we instead found that GATA-3 also bound to target genes in Th1 cells (Fig. 4, Fig. S6, and Table S1). The set of GATA-3 target genes in Th1 cells encompassed most (78%) of those bound in Th2 cells (Fig. 4A) but with the addition of an extra set of genes that were only bound by GATA3.
in Th1 cells (Fig. 4B). Genes bound by GATA-3 in both Th1 and Th2 cells included Th2 genes such as IL4, STAT6, ITK, PTGER4, and CCL27 and Th1 genes such as IFNG, IL18RAP, CCL4, TNF, and NKG7 (Fig. 4 C and D and Fig. S6). Indeed, a total of 92% of Th1 lineage-specific genes occupied by GATA-3 in Th2 cells were also occupied by GATA-3 in Th1 cells. These data show that most genes targeted by GATA-3 in Th2 cells are also occupied by this transcription factor in Th1 cells.

**T-bet and GATA-3 Share Many Target Genes.** Given that we had unexpectedly found a significant number of genes bound by GATA-3 in Th1 cells and that T-bet and GATA-3 often targeted the same genes, we sought to determine whether GATA-3 and T-bet share target genes when both factors are present within the same cell. We first asked whether T-bet and GATA-3 are co-expressed in Th1 cells (Fig. 5A and Fig. S7). Using intranuclear flow cytometry, we found that T-bet and GATA-3 were co-expressed in Th1 cells and that GATA-3 was generally only expressed in Th1 cells that were positive for T-bet. As expected, in Th2 cells the expression of GATA-3 occurred in the absence of T-bet. We next compared the genes targeted by T-bet in Th1 cells with those targeted by GATA-3 in Th1 cells. We found that T-bet and GATA-3 often bound to the same genes with 39% of T-bet target genes in Th1 cells also being occupied by GATA-3 (P < 10^{-16}, hypergeometric; Fig. 5B and Table S1). Furthermore, when present at the same gene, T-bet and GATA-3 peaks were generally closely spaced, with the majority of these peaks coinciding exactly (within the ~200bp resolution of the array, Fig. 5 C and D). Genes bound by both T-bet and GATA-3 included those exhibiting differential gene expression between the two cell types (56% of Th1 genes and 37% of Th2 genes bound by T-bet were also bound by GATA-3) (Fig. 5C and Figs. S7 and S8). We also detected GATA-3 binding to these genes in tertiary stimulated Th1 cells and in a Th1 cell clone (40) using quantitative PCR (Fig. S8), indicating that the detection of GATA-3 binding in the primary stimulated Th1 cells was not due to incomplete Th1 polarization or a potentially heterogeneous cell culture. Taken together with our GATA-3 ChIP data from Th2 cells, our Th1 cell data demonstrate that GATA-3 associates with a core set of genes in both Th1 and Th2 cells and that these genes are also targeted in Th1 cells by T-bet.

The relatively invariant association of GATA-3 with lineage-specific genes in both Th1 and Th2 cells suggests that their differential expression is not due to differences in GATA-3 binding. This argues that T-bet binding in Th1 cells may be responsible for the differential expression of these genes between the two lineages. To explore this possibility, we used our human and mouse expression data to examine the effects of T-bet upon the expression of genes that were bound by both T-bet and GATA-3 in Th1 cells. Expression of T-bet in T cells from IFNγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγγgamma
action of these two transcription factors at a set of shared target genes. These data demonstrate a role for T-bet and GATA-3 in T-cell differentiation that extends beyond the regulation of IFNG and IL-4. Considering with previous results, we have shown that GATA-3 is present in both human Th1 and Th2 cells (31–33). We have also shown that almost all Th1 cells that express GATA-3 also express T-bet. The co-expression of T-bet and GATA-3 is analogous to recent data showing that FoxP3 and ROR-γt are co-expressed in murine T cells (41) and that T-bet and FoxP3 function together in a subset of regulatory T cells (42) and this suggests that this co-expression is functionally significant. We find that not only are T-bet and GATA-3 expressed in the same cells, they occupy highly overlapping sets of genes, including those that show differential expression between Th1 and Th2 cells. We have not been able to use ChiP-ReChIP to test whether T-bet and GATA-3 bind to genes simultaneously in Th1 cells due to failure of our positive control ReChIP RNA polymerase II experiments. However, the co-expression of T-bet and GATA-3 in primary human Th1 cells provides each factor with the potential to act upon their shared target genes in these cells. T-bet directly activates IFNG and represses IL-4 (Fig. 5E) (23, 27) whereas GATA-3 acts in the opposite manner to activate IL-4 and represses IFNG (22, 28). Although both factors are coexpressed in human Th1 cells, T-bet activity would appear to be dominant and these cells do not follow the bifurcation pattern that is recapitulated in murine T-cells by expression of T-bet in the absence of IFNγ. These results indicate that it may be the absence or presence of T-bet that determines T-cell lineage (16, 43) and provides a potential mechanistic rationale that explains why T cells default to the Th2 lineage in the absence of T-bet (3–5). The extension of these findings to vivo polarized human Th1 and Th2 cells is currently under active investigation, but has been limited by the cell numbers required for this type of analysis.

Human T-cell immune responses define pathological outcomes in a wide number of disease states. Identification of novel T-bet and GATA-3 target genes offers the prospect of defining rational therapeutic approaches in these conditions. Indeed, the identification of TNF-α as a direct T-bet target in T cells lead us to the definition of the transcriptional mechanism of a new model of ulcerative colitis (34), providing evidence of the utility of this approach. Other gene targets of T-bet and GATA-3 that we have identified here are also likely to play important roles in infectious diseases.

In conclusion, we have shown that Th1 and Th2 lineage-specific genes that play key roles in T-cell biology are targeted through both T-bet and GATA-3. The action of opposing master regulators through a shared set of target genes may prove to be a general mechanism in other cell types and biological systems.

Methods

Cell Lines. Human CD4 T cells from the peripheral blood of human donors were sorted for CD4+ CD25-CD45RO-HLA-DR- (purity of 98%) and activated with anti-CD3 and anti-CD28 for 48–72 h. Primary Th1 cells were generated over 10–12 days with IL-12 (10 ng/ml) and anti-IL-4 (10 ng/ml) and Th2 cells with IL-4 (10 ng/ml) and anti-IFNγ (1 ng/ml).

ChiP-Chip. We followed previously published ChiP-Chip protocols (29). Binding sites were automatically identified using an algorithm that calculates confidence values for each probe and finds sets of neighboring probes with significant P values (29).

Retrival Transcription of T-bet. T-bet and control retroviruses were produced and retrovirally described (35).

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