The TH1 cell lineage-determining transcription factor T-bet suppresses TH2 gene expression by redistributing GATA3 away from TH2 genes

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ABSTRACT

Lineage-determining transcription factors (LD-TFs) drive the differentiation of progenitor cells into a specific lineage. In CD4+ T cells, T-bet dictates differentiation of the TH1 lineage, whereas GATA3 drives differentiation of the alternative TH2 lineage. However, LD-TFs, including T-bet and GATA3, are frequently co-expressed but how this affects LD-TF function is not known. By expressing T-bet and GATA3 separately or together in mouse T cells, we show that T-bet sequesters GATA3 at its target sites, thereby removing GATA3 from TH2 genes. This redistribution of GATA3 is independent of GATA3 DNA binding activity and is instead mediated by the T-bet DNA binding domain, which interacts with the GATA3 DNA binding domain and changes GATA3’s sequence binding preference. This mechanism allows T-bet to drive the TH1 gene expression program in the presence of GATA3. We propose that redistribution of one LD-TF by another may be a common mechanism that could explain how specific cell fate choices can be made even in the presence of other transcription factors driving alternative differentiation pathways.

INTRODUCTION

Cell fate choice is determined by lineage-determining transcription factors (LD-TF). LD-TFs often antagonise each other’s expression and function in order to drive mutually exclusive gene expression programmes. However, differentiating cells frequently express more than one LD-TF but the effect that this has on the function of the factors and on cell fate choice is often unclear.

Mutually antagonistic LD-TFs are frequently co-expressed in CD4+ helper T cells (1,4–6). Naive CD4+ helper T cells respond to antigen and cytokine cues by differentiating into a number of specialised effector cell lineages (7–10). This functional segregation allows the development of tailored immune responses against specific types of pathogens. The classical example of mutually exclusive effector programs is the differentiation of CD4+ T cells into the TH1 or TH2 lineage in response to intracellular viral and bacterial infections and extracellular parasites, respectively (11,12). Precise control of effector cell lineage determination is critical for an effective immune response and inappropriate T cell differentiation may result in inefficient pathogen clearance or immune pathology (6,13–15).

TH1 and TH2 cell differentiation are driven by the LD-TFs T-bet (TBX21) and GATA3, respectively. Robust differentiation of each lineage is attained through transcription factor and cytokine-mediated positive feedback loops and by cross-inhibition of the alternative cell fate. Specifically, T-bet directly activates the gene encoding the TH1 subtype-defining cytokine interferon gamma (IFNγ) and represses genes encoding the TH2 cytokines IL-4, IL-5 and IL-13, while GATA3 displays the opposite functionality (16–27). Furthermore, both IFNγ and IL-4 function in an autocrine manner to further stabilise the corresponding lineage. In addition, both LD-TFs repress the other’s expression (28) and can bind and auto-activate their own genes (29,30).

These positive and negative feedback loops theoretically generate a bistable switch between TH1 and TH2 cell states.
However, T-bet and GATA3 are co-expressed in in vitro differentiated primary human Th1 cells (29,31,32) and in human CCR5+ T_h memory cells (33). Furthermore, although in vitro polarised murine T_h2 cells stably maintain a T_h2 phenotype, transfer into a mouse model of lymphocytic choriomeningitis virus infection leads to co-expression of GATA3 and T-bet and development of T_h1 properties (T_h2+1 cells) (34). CD4+ T cells co-expressing T-bet and GATA3 can also be generated in vitro by activating naïve cells in the presence of IFNγ, IL-12 and IL-4 and such T_h2/1 hybrid cells have been observed in vivo, during infections with intestinal helminths such as *Heligmosomoides polygyrus*, *Schistosoma mansoni* and *Strongyloides ratti* in mice and *Strongyloides stercoralis* in humans (35,36). T_h2/1 hybrid cells generated from naïve T cells were refractory to cell lineage reprogramming during exposure to T_h1 and T_h2 polarizing cytokines in vitro and remained detectable in a helminth infection model for months after pathogen clearance (35). In addition, experimentally boosting the numbers of T_h2/1 hybrid cells in *H. polygyrus* infected mice increased IFNγ production and parasite fecundity (37). Together, these previous findings demonstrate that T-bet/Sco-expressing cells do not represent uncommitted intermediates, are stable long-term, and can actively modify the nature of an immune response.

In addition to T-bet and GATA3, other pairs of LD-TFs have been found to be co-expressed in CD4+ T cells. T-bet can be co-expressed with Bel6 or Rorγt (38,39), while the regulatory T cell (T_REG) LD-TF FoxP3 can be co-expressed with Bel6, T-bet, GATA3 or Rorγt (4,40–46). However, the effect of LD-TF co-expression on the function of the proteins and the impact on the resultant cell phenotype are not well understood. In human T_h1 cells, GATA3 exhibits a distinct binding profile characterised by loss from T_h2 genes and gain at T_h1 genes (29) suggesting that T-bet may alter GATA3’s DNA binding profile when the two factors are co-expressed. To address this hypothesis, we have expressed T-bet and GATA3 alone or together in T cells and measured the effect on the function of the two factors. We show that T-bet interacts with GATA3 and redistributes it from T_h2 genes to T-bet binding sites at T_h1 genes, switching T cells from a T_h2 to a T_h1 expression program. We propose that direct sequestration of one LD-TF by another may be a common mechanism through which lineage-determination can proceed in cells co-expressing antagonistic LD-TFs.

**MATERIALS AND METHODS**

**Cell culture**

EL4-GFP, EL4-T-bet, EL4-GFP + Plum, EL4-T-bet + Plum, EL4-GFP + GATA3 and EL4-T-bet + Plumb cells were described previously (29,57). EL4 cells expressing mPlum or HA-GATA3 were transduced with FLAG-Tbx21 Y525F-T2A-BSD-ires-EGFP or FLAG-Tbx21 R163/R164A-T2A-BSD-ires-EGFP retrovirus and selected with blasticidin as described (29). EL4-GFP + Plum cells were transduced with retrovirus encoding a doxycycline (dox)-inducible FLAG-T-bet construct and selected with G418. Cells were treated with between 0 and 10 μg/ml dox for 48 hours prior to harvesting. HEK293T cells and EL4 cells were cultured in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum, 4 mM L-alanyl-L-glutamine, 25 mM D-glucose, 1 mM sodium pyruvate, 10 mM HEPES, 100 U/ml penicillin and 100 μg/ml streptomycin. EL4 cells were stimulated with phorbol 12-myristate 13-acetate (50 ng/ml) and ionomycin (1 μM) for 4 hours prior to formaldehyde crosslinking.

**ChIP**

Cells were crosslinked by the addition of one-tenth volume of fresh 11% formaldehyde solution for 20 minutes at room temperature before the reaction was quenched by addition of glycine, as described (57). Cells were rinsed twice with 1x PBS and flash frozen in liquid nitrogen. Cells were lysed with non-ionic detergent, the nuclei washed and then lysed with ionic detergent. Cells were sonicated on ice to solubilize and shear crosslinked DNA (27W for 10 × 30 second pulses using a Misonix Sonicator 3000). The resulting whole cell lysate was cleared by centrifugation and then incubated overnight at 4°C with 50 μl of Protein G magnetic Dynabeads that had been pre-incubated with 5 μg of purified antibody (anti-FLAG, M2 Sigma; anti-HA, 3F10 Roche; anti-H3K27ac, ab4729 Abcam). Beads were washed 6 times with RIPa buffer and once with TE containing 50 mM NaCl. Bound complexes were eluted from the beads by heating at 65°C with rocking for 2 hrs and crosslinks then reversed in IP and input DNA by incubation at 65°C for 6 hrs. IP and input DNA were then purified by treatment with RNase A, proteinase K and isolated with KAPA Pure beads.

For ChIP in transiently transfected HEK293T cells, 4.5 × 10^6 cells were plated onto a 15 cm plate. The cells were transfected 24 hours later with 70 μg of wild-type (WT) HA-GATA3 or HA-GATA3 C320G expression plasmids using polyethyleneimine. The cells were washed with PBS 36 hrs after transfection and crosslinked as above. The remaining steps were carried out as outlined above for EL4 cells except that the cells were sonicated using a Bioruptor Pico (Diagenode) (3 × 30 sec pulses with 30 sec gaps in between).

Specific DNA sequences were quantified in triplicate in ChIP-enriched and input DNA by quantitative PCR (ThermoFisher QuantStudio) using QuantiTect SYBR Green PCR master mix (Qiagen) and enrichment calculated relative to input. Standard curves were performed for all primer pairs to ensure linear amplification. The following primers were used to quantify ChIP enrichment:

**Ilf4 CNS2 forward:** 5’-ATC ACG TCG TCT TAC CCA AAC A-3’,

**Ilf4 CNS2 reverse:** 5’-TGT GGG AGA GCG TCT GAT CTG T-3’,

**Ifng TSS forward:** 5’-CCT GTG CTG TGC TGT G TG G-3’,

**Ifng TSS reverse:** 5’-ACT CCT TGG GCT CTC TGA CG-3’,

**Ifng CNS-6 forward:** 5’-GAC GAG GTC TGC A AC CCT TGA AGC TGT GGG TAC-3’,

**Ifng CNS-6 reverse:** 5’-TGA CTC GAG AGA TTG CCG TCT GGT CTT GGC GT-3’.
ChIP-seq

Libraries were constructed from ChIP DNA by standard Illumina protocols, except that DNA in the range 150–350 bp was gel-purified after PCR-amplification. The libraries were quantified using a Qubit fluorometer and Agilent bioanalyzer, pooled and subjected to 50 bp single-end sequencing with a HiSeq 2500 sequencer (except for samples from the dox-induced T-bet experiment, which were subjected to 75 bp single-end sequencing with a NextSeq 550 sequencer). Raw reads were filtered for base quality using FastQC with default parameters and adapters sequences were removed with Trim Galore! Reads passing quality filtering criteria were aligned to GRCh38 using BWA with the default settings (58). Significantly enriched regions were identified with MACS v2 using a significance threshold of \( p < 0.01 \) (59). The significance of changes in T-bet and GATA3 binding to the diffbind R package based on two ChIP replicates (60). The different patterns of GATA3 binding were defined by the following criteria: gain sites: T-bet + GATA3 > GATA3 + GFP at FDR < 0.05 and called by MACS2 in both T-bet + GATA3 replicates; loss sites: T-bet + GATA3 < GATA3 + GFP at FDR < 0.05 and called by MACS2 in both GATA3 + GFP replicates; invariant sites: T-bet + GATA3 vs GATA3 + GFP binding ratio \( < = 1.25 \), differentially bound between sample groups at FDR > 0.05 and called by MACS2 in both replicates of both sample groups. All GATA3 sites: called by MACS2 in both replicates of GATA3 + GFP sample. Binding sites that overlapped ENCODE blacklist version 2 regions (https://github.com/Boyle-Lab/Blacklist/tree/master/lists) were removed. Genes whose transcriptional start sites was closest to the center of GATA3 gain, invariant and loss sites were determined with the ChIPpeakAnno R package (61) using the GRCh38 reference transcript assembly. We downloaded previously published T-bet ChIP-seq data from mouse T\(_{H1}\) cells (activated with \( \alpha \)CD3/CD28 for 3 days followed by 4 days of culture with IL-2 and IL-12: Rep1: GSM998272 (62) and Rep2: GSM836124 (63)) and GATA3 ChIP-seq data from mouse T\(_{H2}\) cells (activated with \( \alpha \)CD3/CD28 and polarized with IL-4 in the presence of \( \alpha \)IFN-\( \gamma \) and \( \alpha \)IL-12 for 8 days: GSM293180 (64)). The data were processed with the same pipeline used for the EL4 data. Only binding sites called by MACS in both T\(_{H1}\) cell T-bet ChIP-seq datasets were considered.

Average binding profiles (in reads/million) and heatmaps across sets of T-bet and GATA3 binding sites were generated with ngsplot (65). To control for differences in ChIP efficiency when comparing average profiles between ChIPs of the same factor in different cells, average reads/million across a set of sites were then normalized by the average signal across all sites.

Locations in the human genome (hg19) orthologous to GATA3 binding sites in EL4 cells were identified using liftOver and average GATA3 binding profiles in human T\(_{H1}\) cells (GSM776558 (29)) and T\(_{H2}\) cells (GSM776559 (29)) plotted across these sites using ngsplot.

RNA-seq

EL4 cells were stimulated with phorbol 12-myristate 13-acetate (50 ng/ml) and ionomycin (1 \( \mu \)M) for 4 hours and total RNA was then purified with TRIsure (Bioline). Polyadenylated RNA was isolated with Oligotex (Qiagen) and libraries were prepared using the NEBNext Multiplex Small RNA Library Prep kit and then sequenced on an Illumina HiSeq 2500 using 100 bp paired-end reads. Raw reads were filtered for base quality using FastQC with default parameters and adapters sequences were removed with Trim Galore! Transcript-level read abundances were created with kallisto (66) using the Gencode M20 transcript models and combined with gene-level abundances using tximport (67). Expression estimates were then modelled using the DESeq2 package (68). Ranked gene lists were generated by ordering the expression ratios by -log10 \( p \)-value divided by the rank of the fold-change and enrichment for selected gene sets was computed using the fgsea R package (69). T\(_{H1}\) and T\(_{H2}\) specific gene sets were previously defined (70).

ATAC-seq

2 \( \times \) 10\(^6\) EL4-GFP or EL4-T-bet cells were stimulated with 50 ng/ml PMA and 1 \( \mu \)g/ml ionomycin at 37°C for 4 hrs or left unstimulated. For each ATAC-seq reaction, 50,000 cells were washed in 50 \( \mu \)l cold PBS (500g, 5 min, 4°C) and resuspended in 50 \( \mu \)l cold lysis buffer I (10 mM Tris-Cl pH 7.4, 10 mM NaCl, 3 mM MgCl\(_2\), 0.5% IGEPA L-CA-630) and spun (500g, 5 min, 4°C). The supernatant was removed, and lysed cells resuspended in 50 \( \mu \)l transposase reaction mix, containing 10 mM Tris pH8.0, 5 mM MgCl\(_2\), 10% dimethylformamide and 2.5 \( \mu \)l Tn5 transposase + transposase reagent (TDE1 Illumina). Reactions were incubated at 37°C for 45 min, with shaking at 500 rpm. Following transposition, 150 \( \mu \)l reverse-crosslinking buffer (50 mM Tris-Cl, 1 mM EDTA, 1% SDS, 0.2 M NaCl) was added to reactions, and incubated at 65°C overnight at 1000 rpm before DNA purification, using the MinElute PCR purification kit (Qiagen).

Transposed DNA was amplified by PCR for 5 cycles, using KAPA HiFi HotStart ReadyMix, 1.25 \( \mu \)M Ad1 primer and 1.25 \( \mu \)M Ad2.x indexed primers, as described (71). 5 \( \mu \)l of each reaction was taken for qPCR as described, using KAPA HiFi HotStart ReadyMix and 50X KAPA Low ROX for signal normalisation, to determine the additional number of PCR cycles for library amplification. PCR reactions were purified using the MinElute gel extraction kit (Qiagen) according to the manufacturer’s instruction, and fragments between 100–1000 bp in size were selected for sequencing. DNA fragmentation of the library was measured using the Bioanalyzer High Sensitivity DNA assay (Agilent), and library concentration quantified by qPCR using the Illumina Library Quantification kit (KAPA). DNA from multiple libraries was pooled at equimolar concentrations to a final concentration of 4 nM. Pooled libraries were sequenced on a HiSeq 3000 (Illumina) using 75 bp paired-end reads. Raw reads were filtered for base quality using FastQC with default parameters and adapters sequences were removed with Trim Galore! Reads passing quality filtering criteria were aligned to GRCm38 using BWA-MEM with default settings (58).

FRET

HEK293T cells were plated into 24-well plates at 5.5 \( \times \) 10\(^4\) cells/well and transfected 24 hours later with 730 ng of
each vector encoding mTagBFP and mNeonGreen tagged proteins using polyethyleneimine. Fluorescence signals were recorded 32 hours after transfection with a FACSymphony flow cytometer (BD Biosciences). The FRET signal was recorded in the 405 nm laser line using a 530/30 bandpass filter. DFRET efficiency was calculated as described (72).

FLIM-FRET

HEK293T cells were plated into 8-well glass coverslip bottom slides (Ibidi) at 2.7 × 10^4 cells/well. The cells in each well were transfected 24 hours later with 760 ng of each vector as above. The cells were washed with PBS 36 hrs after transfection, fixed with 2% formaldehyde for 15 min, washed twice with PBS and left covered in PBS for image acquisition.

Time-domain fluorescence lifetime images were acquired via time correlated single photon counting (TCSPC) on an Open FLIM microscope (73) at a resolution of 256 by 256 pixels, with 256 time bins and 100 frames accumulated over 300 seconds, via excitation and emission filters suitable for the detection of mTagBFP fluorescence (Excitation filter: Semrock FF01-40/40-25; Beam Splitter: Edmund 48NT-392 30R/70T; Emission filter: Semrock FF01-470/22–25). For each sample a ‘donor’ control image and a ‘donor with acceptor’ image was acquired. FLIM analysis was performed with TRI2 software (v3.0.1.15) using monoeXponential Levenberg-Marquardt fitting (74). The FRET efficiency (FRETeff) for each cell sample region of interest was calculated according to the equation \( \text{FRETeff} = 1 - (tD/tD) \), where TD is the average lifetime of mTagBFP in the absence of a FRET acceptor from the donor image and tDA is the average lifetime of mTagBFP in the presence of the acceptor mNeonGreen (mNG), from the ‘donor with acceptor’ image.

Co-immunoprecipitation and immunoblot analysis

HEK293T cells were transiently transfected with expression vectors for mNeonGreen-tagged or HA-tagged proteins using polyethyleneimine. Cells were lysed in lysis buffer (50 mM Tris pH 7.5, 300 mM NaCl, 0.5% IGEPLA CA-630, 1 mM EGTA, 2 mM MgCl₂) supplemented with 250 U/ml benzonase (Santa Cruz), 1 mM phenylmethylsulfonyl fluoride, 1 mM DTT and 1X complete EDTA-free protease inhibitor (Roche). The cell lysates were sonicated for 10 sec using a Bioruptor Pico (Diagenode) and then incubated on ice for 1 hour with rotation. Insoluble material was removed by centrifugation (17,000 g for 10 min), diluted in an equal volume of lysis buffer without NaCl and incubated with 10 μl of mNeonGreen-Trap magnetic agarose beads (ChromoTek) on an over-head shaker for 1 hr at 4°C. Beads were washed 2x with 200 μl lysis buffer containing 150 mM NaCl and 4x with lysis buffer containing 500 mM NaCl, resuspended in 1x Laemmli buffer and incubated at 95°C for 9 min. Proteins were resolved on a 12% Tris-glycine gel and transferred to a nitrocellulose membrane. Membranes were probed with anti-β-Tubulin (ab6046, Abcam), anti-HA (3F10, Roche), anti-FLAG (M2, Sigma), anti-GATA3 (HG3-31 (Santa Cruz), anti-T-bet (4B10, eBioscience), anti-β-actin (4967S, Cell Signaling Technologies), and anti-mNeonGreen (#53061, Cell Signaling Technologies).

Motif analysis

De novo and known DNA binding motifs enriched in the sets of gain, loss and invariant GATA3 binding sites were identified with the findMotifsGenome.pl program from the HOMER motif discovery library (75) using the following parameters: mm10 – size 200 – mask. The occurrences of T-box and GATA motifs were quantified using the FIMO motif scanning tool from the MEME suite using a p-value < 0.0004 cutoff (76).

Oligo-Flow

Oligo-Flow was performed as described (77). We designed DNA oligonucleotides that incorporated the synthetic T-box motif and a control oligo corresponding to the DNA sequence ~7 kb upstream of Il5 that lacks a T-box motif: T-box sense: 5′-TTA GCT AGT CGG CGC TAT AGT TTT CAC ACC TAC GTA ACT GCT GCT AGC TAG TAC-3′ T-box antisense: 5′-GTA CTA GCT AGC AGC AGT GTA GGT GTG AAA ACT ATA GCG CCG ACT AGC TAA-3′ Il5 sense: 5′-TAC TGA TCT GTA GCA CAT TAA AGG AGA TAG AGG GCT TAG GGC ACA GGG GGT AAA-3′ Il5 antisense: 5′-TTT ACC CCC TGT GCC CTA AGC CCT GTA TCT CCT TTA ATG TGC TAC AGA TGA TAA-3′

The biotinylated sense oligos and complementary unmodified antisense oligos were annealed in annealing buffer (10 mM Tris-HCl pH 8.0, 50 mM NaCl, 10 mM MgCl₂ and 1 mM DTT) and incubated with 12 μl streptavidin-coated paramagnetic microspheres (7.5 – 8.5 μm COMPEL, Bangs Laboratories) for 1 hr at RT with rotation. Oligo/bead complexes were washed once in protein binding buffer (10 mM Tris pH 8.0, 100 mM KCl, 10% glycerol, 1 mM EDTA, 1 mM DTT and 160 μg / ml BSA) and resuspended in 462 μl protein binding buffer supplemented with 20 μg / ml poly(dI-dC) (Sigma), 1 mM phenylmethylsulfonyl fluoride and 1X complete EDTA-free protease inhibitor (Roche).

HEK293T cells were transiently transfected with expression vectors for mNeonGreen-tagged WT or mutant T-bet DBD using polyethyleneimine. Cells from one 10 cm plate were lysed in 100 μl lysis buffer (20 mM Tris pH 7.5, 150 mM NaCl, 0.5% IGEPLA CA-630, 0.5 mM EDTA, 2 mM MgCl₂, 1 mM phenylmethylsulfonyl fluoride, 1 mM DTT and 1X complete EDTA-free protease inhibitor (Roche)). Insoluble material was removed by centrifugation (17,000 xg for 10 min) and 20 μl of cleared lysate was incubated with oligo/bead complexes on an over-head shaker for 2 hrs at 4°C. Oligo/bead/protein complexes were washed twice with 200 μl protein binding buffer and resuspended in 600 μl protein binding buffer. Fluorescence was recorded on an LSRFortessa X-20 flow cytometer (BD Biosciences). The background signal from the Il5 sequence oligo was subtracted from the signal from the T-box motif oligo and the signals for the mutant proteins were then normalised to the signal of the WT T-bet sample.
Expression constructs

FLAG-T-bet was inserted into BamHI and EcoRI sites of TiTMPV-Neo (Addgene plasmid #27993) to generate TRE-T-bet. The cDNA clones of the Tbx21 Y525F and R1634/R164A mutants were kindly provided by Eun Sook Hwang and Amy Weinmann, respectively, and cloned into pMY to generate FLAG-Tbx21 Y525F-T2A-BSD-IRESC-EGFP and FLAG-Tbx21 R1634/R164A-T2A-BSD-IRESC-EGFP, respectively. Other T-bet and GATA3 mutations were generated via standard two-step overlap PCR mutagenesis using primers containing modified codon sequences. GATA3 R352A was cloned into pMY-PAC-FT2A-HA-GATA3 R352A-IRES-mPlum, as described (29).

RESULTS

T-bet redistributes GATA3 away from its canonical binding sites

We first sought to determine if co-expression of T-bet and GATA3 had an effect on the genome binding profiles of either factor. To address this question, we expressed FLAG-T-bet and HA-GATA3, either individually or together, in the C57BL/6 mouse T lymphoblast cell line EL4 (78) (Supplementary Figure S1A). Expression of T-bet resulted in production of IFNγ by EL4 cells, as observed previously (17), but expression of GATA3 did not lead to production of IL-4 (data not shown). The cells were stimulated with PMA and ionomycin and T-bet and GATA3 binding sites identified across the genome by chromatin immunoprecipitation sequencing (ChiP-seq).

In order to validate the EL4 cell model, we first compared the sites occupied by T-bet and GATA3 when expressed in EL4 cells individually with the sites occupied by these factors in primary TH1 cells (62,63) or primary TH2 cells (64), respectively (Supplementary Figure S1B-D). This showed that the binding profiles of T-bet and GATA3 were similar between EL4 cells and primary T cells, but that some sites exhibited greater occupancy in one cell type or the other. Analysis of transcription factor binding motifs at these sites revealed enrichment of similar motifs, including strong enrichment of T-box motifs at sites uniquely bound by T-bet in EL4 cells (Supplementary Figure S1E). Similarly, sites uniquely bound by GATA3 in EL4 cells exhibited strong enrichment for GATA motifs (Supplementary Figure S1F).

Thus, we conclude that T-bet and GATA3 bind similar sets of sites with similar motifs in EL4 cells compared with primary T cells.

We next determined whether co-expression of T-bet and GATA3 had any effect on T-bet genome occupancy. Comparison of all T-bet binding sites between cells expressing T-bet alone and cells co-expressing T-bet and GATA3 did not reveal any significant changes in T-bet binding in the presence of GATA3 (Figure 1A). In contrast, we observed dramatic changes in GATA3 genome occupancy when T-bet was present (Figure 1A and B). Specifically, 7% (988) of GATA3 binding sites were lost in the presence of T-bet and 13% (1,760) of GATA3 binding sites were gained in the presence of T-bet (Figure 1A). In addition, around half (6,622) of GATA3 binding sites were invariant (significantly bound by GATA3 in the presence and absence of T-bet). Thus, when both factors were co-expressed, GATA3 had no effect on T-bet localisation while T-bet induced a major redistribution of GATA3 to a different set of sites.

We have previously noted that GATA3 adopts an alternative binding profile in primary human TH1 cells versus TH2 cells (29). We have also demonstrated that a large proportion of transcription factor binding sites are shared between human and mouse CD4+ T cells (70). Thus, to determine if the alternative binding profile adopted by GATA3 in TH1 cells could be related to the redistribution of GATA3 by T-bet in EL4 cells, we identified locations in the human genome orthologous to murine GATA3 binding sites using liftOver and plotted GATA3 occupancy at these locations in human TH1 and TH2 cells (Supplementary Figure S2A). We found that sites at which GATA3 was gained in the pres-
Figure 1. T-bet redistributes GATA3 from its canonical binding sites to T-bet binding sites. (A) Top: Changes in T-bet occupancy in cells co-expressing T-bet and GATA3 compared with cells expressing T-bet alone (T-bet + Plum). Each point represents a site bound by T-bet in one of the two cell types. Bottom: Changes in GATA3 occupancy in cells co-expressing T-bet and GATA3 compared with cells expressing GATA3 alone (GFP + GATA3). Each point represents a site bound by GATA3 in one of the two cell types. For both transcription factors, sites with significantly higher occupancy (FDR < 0.05; gain sites), significantly lower occupancy (FDR < 0.05; loss sites), or no change in occupancy (FDR > 0.05, -1.25 < log2 FC < 1.25; invariant sites) in the presence of the other factor are coloured as indicated (n = 2 biological replicates). (B) Heatmaps showing GATA3 occupancy (reads/million) at GATA3 gain, invariant and loss sites defined in A in the presence or absence of T-bet (representative of two biological replicates). (C) Heatmap showing T-bet occupancy at GATA3 gain, invariant and loss sites defined in A in the presence or absence of T-bet (representative of two biological replicates). (D) Binding profiles (reads/million) for T-bet and GATA3 at genes associated with GATA3 gain (top), invariant (center) and loss (bottom) sites in EL4 cells expressing each factor alone or co-expressing the two factors (representative of two biological replicates). T-bet occupancy in mouse T_{H1} cells and GATA3 occupancy in mouse T_{H2} cells are shown below for comparison.
ence of T-bet exhibited higher levels of GATA3 occupancy in T\(_{H1}\) cells compared to T\(_{H2}\) cells. Reciprocally, sites at which GATA3 was lost in the presence of T-bet exhibited lower levels of GATA3 occupancy in T\(_{H1}\) cells, while sites at which GATA3 was invariant in the presence or absence of T-bet showed equal GATA3 occupancy in T\(_{H1}\) and T\(_{H2}\) cells. Thus, these results suggest that the phenomenology of T-bet-mediated GATA3 redistribution occurs in both EL4 cells and primary T cells.

We considered two scenarios by which T-bet could bring about redistribution of GATA3. Firstly, T-bet could compete with GATA3 for its binding sites, evicting it and forcing GATA3 to redistribute to alternative locations. Alternatively, T-bet could induce GATA3 to localise to its own binding sites, thereby depleting GATA3 from its canonical binding sites. To distinguish between these possibilities, we measured T-bet binding at sites at which GATA3 was lost, gained or invariant in the presence of T-bet (Figure 1C). This showed that T-bet was absent from sites at which GATA3 was lost, which rules out an eviction mechanism. In contrast, T-bet was bound with GATA3 at sites to which GATA3 was invariant or gained in the presence of T-bet. This suggests that T-bet recruits GATA3 to its own binding sites. These binding patterns could also be observed at individual genes (Figure 1D and Supplementary Figure S2B). For example, when expressed alone, GATA3 occupied sites at Ccr3 but these disappeared in the presence of T-bet. In contrast, GATA3 did not bind to Ilfng when expressed alone, but co-occupied the gene with T-bet when the two factors were co-expressed. Thus, when both factors are present, T-bet induces redistribution of GATA3 from its canonical binding sites to T-bet binding sites.

**The extent of GATA3 redistribution is dependent on the relative levels of T-bet and GATA3**

We considered whether the relative levels of T-bet and GATA3 might affect the ability of T-bet to redistribute GATA3. To test this, we generated an EL4 cell line that expressed higher levels of HA-GATA3 and a doxycycline (dox)-inducible FLAG-T-bet construct through which the T-bet expression level could be controlled by titration of dox (Figure 2A). These cells contained higher levels of GATA3 and lower levels of T-bet, which were more comparable to primary T\(_{H1}\) cells (compare Figure 2A with Supplementary Figure S1A). To determine the effect varying the level of T-bet expression on GATA3 occupancy, we treated the cells with 0.4, 2 or 10 mg/ml dox for 48 hours and then performed ChIP-seq for HA-GATA3 and FLAG-T-bet. We also compared these data with GATA3 occupancy in the absence of T-bet in our previously generated GATA3 + GFP cell line. We found that regardless of the level of T-bet expression, T-bet recruited GATA3 to T-bet-dependent gain sites and the greater the level of T-bet expression, the greater the level of GATA3 recruitment to these positions (Figure 2B). GATA3 recruitment to T-bet target sites in proportion to the level of T-bet occupancy was also apparent at individual genes (Figure 2C) and was also confirmed at Ilfng by ChIP-qPCR (Figure 2D). We also found that the greater the expression of T-bet, the greater the depletion of GATA3 from loss sites (Figure 2B), although the lower level of T-bet expression in these cells was not sufficient to completely remove GATA3 from these sites. These results demonstrate that the greater the amount of T-bet relative to GATA3, the greater the degree of GATA3 redistribution.

There was also evidence of T-bet recruitment to GATA3 loss sites when present at its lowest level in the 0.4 mg/ml dox-treated sample (Figure 2B), suggesting that GATA3 may be able to recruit T-bet to GATA3 binding sites when GATA3 is in excess. However, the level of T-bet recruitment to GATA3 binding sites was low compared to the level of GATA3 recruitment to T-bet-dependent sites. These data suggest that T-bet and GATA3 are both able to recruit the other factor to their binding sites, but that the effect of T-bet on GATA3 is dominant, consistent with the previously observed redistribution of GATA3 to T-bet binding sites in human T\(_{H1}\) cells (29).

**T-bet-mediated GATA3 redistribution is associated with silencing of the T\(_{H2}\) gene expression program**

We next sought to determine whether the redistribution of GATA3 in the presence of T-bet was associated with changes in gene expression. To address this, we performed RNA-seq in the T-bet-expressing, GATA3-expressing, and T-bet and GATA3 co-expressing cells that we used for the initial transcription factor binding studies. Using gene set enrichment analysis (GSEA) we found, as expected, that expression of T-bet alone upregulated T\(_{H1}\)-specific genes and downregulated T\(_{H2}\)-specific genes (Supplementary Figure S3A). Correspondingly, expression of GATA3 alone had the opposite effect, upregulating T\(_{H2}\) genes and downregulating T\(_{H1}\) genes (Supplementary Figure S3B). Thus, in isolation each LD-TF can establish its canonical lineage-specific gene expression program in EL4 cells.

We then determined the effect of T-bet on gene expression when co-expressed with GATA3. Comparing the gene expression changes induced by T-bet alone with those induced by T-bet and GATA3, we found that T-bet established essentially the same expression profile in the presence and absence of GATA3 (\(r = 0.83, p < 2.2e-16\); Figure 3A). In contrast, there was no correlation between the expression profile of T-bet / GATA3 co-expressing cells and cells containing GATA3 alone (\(r = 0.26, p < 2.2e-16\), demonstrating that T-bet prevents GATA3 from establishing its expression programme (Figure 3B). As for cells containing T-bet alone, cells expressing both T-bet and GATA3 upregulated T\(_{H1}\)-specific genes in the absence of significant upregulation of T\(_{H2}\) genes (Figure 3C and D). Thus, when both factors are expressed together, T-bet function dominates, with cells adopting a T\(_{H1}\) expression profile, characterised by induction of T\(_{H1}\) genes and repression of T\(_{H2}\) genes compared to cells expressing GATA3 alone.

We next asked how these changes in gene expression were related to GATA3 redistribution. We tested whether genes associated with sites at which GATA3 was lost, gained or invariant in the presence of T-bet were enriched among the sets of genes up- or down-regulated in cells co-expressing T-bet and GATA3 versus cells expressing GATA3 alone. This analysis revealed that genes downregulated in the presence of T-bet were enriched for sites at which GATA3 was lost
Figure 2. The extent of GATA3 redistribution is dependent on the relative levels of T-bet and GATA3. (A) Immunoblot with αFLAG, αHA, αGATA3 and αACTB in EL4 cells constitutively expressing FLAG-T-bet and HA-GATA3 (used in Figure 1) and EL4 cells that constitutively express HA-GATA3 and a dox-inducible FLAG-T-bet construct (TRE-T-bet) after treatment with 0–10 μg/ml dox for 48 hours. The positions of endogenous GATA3 and HA-GATA3 in the GATA3 blot are labelled. Representative of two biological replicates. (B) Heatmaps showing T-bet and GATA3 occupancy (reads/million, according to the scales to the right) at GATA3 gain, invariant and loss sites defined in Figure 1A after treatment of TRE-T-bet + GATA3 EL4 cells with 0.4, 2 or 10 μg/ml dox for 48 hours (one replicate). T-bet and GATA3 occupancy in GATA3 + GFP cells (used in Figure 1, not treated with dox) are shown at the same sites for comparison. (C) Binding profiles (reads/million) for T-bet and GATA3 at genes associated with GATA3 gain sites in cells described in B (one replicate). (D) Enrichment of Ifng promoter DNA relative to input DNA by ChIP for FLAG-T-bet (white) or HA-GATA3 (grey) in cells shown in A, measured by qPCR (mean and SD, n = 3 technical replicates of one biological replicate).
and that upregulated genes were enriched for sites at which GATA3 was gained (Figure 3E). In contrast, GATA3 invariant sites were not significantly enriched among either upregulated or downregulated genes (Figure 3E).

We also asked whether the recruitment of GATA3 to T-bet binding sites had any effect on the expression of the associated genes. Comparing gene expression between cells co-expressing T-bet and GATA3 with cells expressing T-bet alone, we found no enrichment of sites to which GATA3 was gained at the genes that were up- or down-regulated in the presence of GATA3 (Figure 3F), suggesting that GATA3 recruitment had no effect on the expression of T-bet target genes. Thus, we conclude that T-bet-mediated GATA3 redistribution results in suppression of the TH2 gene expression program.
gene expression program without impacting T\(_H\)1 gene expression.

**GATA3 removal depletes H3K27ac from its binding sites**

Our results show that T-bet removes GATA3 from a subset of its binding sites and that this is associated with gene repression. We sought to determine whether this repressive effect of T-bet was reflected in changes to chromatin at GATA3 binding sites by profiling the active histone modification H3K27ac by ChIP-seq in the presence or absence of T-bet and GATA3. We found increased levels of H3K27ac at sites to which GATA3 was recruited in the presence of T-bet (Figure 4A), consistent with the positive effect of T-bet and GATA3. We found increased levels of H3K27ac by ChIP-seq in the presence or absence of T-bet and GATA3. We found increased levels of H3K27ac at these GATA3 gain sites (Figure 4A), consistent with the lack of effect of GATA3 on the expression of these genes (Figure 3F). Reciprocally, we found decreased levels of H3K27ac at sites from which GATA3 was lost in the presence T-bet (Figure 4A), consistent with the reduced expression of these genes in the presence of T-bet. We conclude that T-bet-mediated GATA3 redistribution is associated with changes in chromatin modification at GATA3 binding sites that is consistent with silencing of the T\(_H\)2 gene expression program.

**GATA3 redistribution to new sites requires T-bet DNA binding activity**

We considered that T-bet may recruit GATA3 to new binding sites by opening chromatin and making the sites accessible. To test this, we performed ATAC-seq in unstimulated and PMA + ionomycin stimulated EL4 cells in the presence and absence of T-bet. We found that GATA3 loss sites and GATA3 invariant sites tended to be accessible in the absence of T-bet and that this accessibility increased upon cell stimulation (Figure 4B). In the presence of T-bet, chromatin accessibility was reduced at GATA3 loss and GATA3 invariant sites, consistent with the repressive effect of T-bet on the genes associated with these sites. In contrast, GATA3 gain sites tended to be inaccessible in the absence of T-bet and exhibited increased chromatin accessibility when T-bet was present independently of the cellular activation status. Thus, this ATAC-seq analysis indicates that T-bet may recruit GATA3 to gain sites by opening chromatin and thereby allowing GATA3 to access otherwise inaccessible DNA sequence motifs.

To determine whether GATA3 could be directly binding to DNA at gain sites, we identified T-box and GATA binding motifs significantly enriched at their binding sites in cells expressing T-bet or GATA3 alone, respectively, and quantified the occurrences of these motifs at GATA3 gain, loss and invariant sites (Figure 4C). We found that a GATA motif was present at 32% of loss sites but at only 14% of gain sites. We found the opposite pattern for the T-box motif, with 43% of gain sites containing the motif compared with only 8% of loss sites (Figure 4C). These data suggest that it is T-bet DNA binding motifs rather than GATA DNA binding motifs that are important for GATA3 redistribution.

To determine if additional factors may have a role in changes in GATA3 occupancy in the presence of T-bet, we systematically identified motifs enriched at GATA3 gain, loss and invariant sites (Supplementary Figure S4A). In comparison to GATA3 invariant sites, we found that T-bet-dependent GATA3 gain sites exhibited strong enrichment for T-box motifs. No other motif type exhibited this relative enrichment at GATA3 gain sites, consistent with a model in which T-bet directly recruits GATA3 to these sites. Comparison of GATA3 invariant sites to GATA3 loss sites showed no differences in motif enrichment, consistent with the model that GATA3 loss is caused by sequestration by T-bet rather than changes in binding of a third factor to these sites (Supplementary Figure S4B). We also looked for enrichment of potential composite T-bet and GATA3 DNA binding motifs at GATA3 gain sites through which T-bet and GATA3 may bind together but could not find any evidence of such motifs (Supplemental Figure S4C).

The low frequency of GATA motifs coupled with the high frequency of T-box motifs at T-bet-dependent GATA3 binding sites indicated a requirement for T-bet DNA binding for GATA3 recruitment to these sites. To test this, we measured GATA3 recruitment in the presence of the T-bet DNA-binding mutant R163A + R164A (79). ChIP for HA-GATA3 showed that mutation of the T-bet DNA binding domain (DBD) abrogated its ability to recruit GATA3 to Ifng (p = 0.0002, Student’s t-test, Figure 4D). Thus, we conclude that recruitment of GATA3 to new sites requires T-bet DNA binding.

**Recruitment of GATA3 to T-bet binding sites does not require its canonical DNA binding activity**

Given that GATA3 was recruited to sites enriched for the T-box motif in a manner that required T-bet DNA binding, we considered the possibility that GATA3 was bound to these sites via interaction with T-bet. In support of such a model, two independent studies have reported that T-bet and GATA3 interact in co-IP experiments (23,80). T-bet has been reported to interact with GATA3 via its C-terminus, with mutation of Y525 to phenylalanine disrupting this interaction (23). However, we found that T-bet Y525F still interacted with GATA3 (Supplementary Figure S5A) and, consistently, could still recruit GATA3 to Ifng (Supplementary Figure S5B).

The T-box family member TBX5 and the GATA family member GATA4 interact in cardiomyocytes (81). Mutations in the TBX5 and GATA4 DBDs disrupt the interaction between the proteins suggesting that interactions between T-box and GATA factors may be mediated by their DBDs (81). To test whether T-bet and GATA3 interacted via their DBDs, we co-expressed mTagBFP-tagged GATA3 DBD and mNeonGreen-tagged T-bet DBD in HEK293T cells and measured Förster resonance energy transfer (FRET) between the fluorescent proteins by flow cytometry. We identified a strong FRET signal in the presence of fluorescent T-bet and GATA3 that was absent if either protein was replaced with the fluorophore alone (Figure 5A). Thus, these results support a model in which T-bet recruits GATA3 to its binding sites through direct interaction between their DBDs.
**Figure 4.** T-bet recruits GATA3 to sites of open chromatin in a manner dependent on T-bet DNA binding. (A) Heatmaps showing H3K27ac enrichment at GATA3 gain, invariant and loss sites in cells expressing GFP and Plum, T-bet and Plum, GFP and GATA3 or GATA3 and T-bet (one biological replicate). (B) Heatmaps showing chromatin accessibility (measured by ATAC-seq) at GATA3 gain, invariant and loss sites in unstimulated (US) and stimulated (Stim.) cells expressing GFP or T-bet (representative of two biological replicates). (C) Left: T-box and GATA DNA binding motifs enriched at T-bet binding sites and GATA3 binding sites, respectively, in cells expressing either factor alone. Right: Proportion of GATA3 gain, invariant and loss sites that contain the T-box or GATA3 DNA binding motifs. (D) Enrichment of Ifng promoter DNA relative to input DNA by ChIP for FLAG-T-bet (white) or HA-GATA3 (grey) in EL4 cells expressing GFP and GATA3, WT T-bet and GATA3, T-bet DNA binding mutant R163A/R164A and Plum or T-bet R163A/R164A and GATA3 measured by qPCR (mean and SD, n = 3 technical replicates of one biological replicate, one-tailed Student’s t-test).

**Figure 5.** Recruitment of GATA3 to T-bet binding sites does not require its canonical DNA binding activity. (A) Flow cytometric measurement of FRET between mTagBFP-GATA3 DBD and mNG-NLS, mTagBFP and mNG-T-bet DBD or mTagBFP-GATA3 DBD and mNG-T-bet DBD. mTag-BFP was excited with the violet laser and fluorescence emission from mNG measured in the same laser line with a 530/30 bandpass filter. The proportion of FRET-positive cells are indicated for each combination of constructs. Representative of two biological replicates. (B) Enrichment of Il4 CNS2 and Ifng promoter DNA relative to input DNA by ChIP for anti-FLAG (white) or anti-HA (grey) in EL4 cells expressing WT GATA3 or GATA3 R352A DNA binding mutant in the absence (GFP) or presence of T-bet (mean and SD, n = 3 technical replicates of one biological replicate, one-tailed Student’s t-test).
Our findings thus far are compatible with two functionally different models of GATA3 binding to gain sites; either T-bet-mediated chromatin opening allows GATA3 to bind to its canonical GATA motif at these sites or that T-bet binds GATA3 and brings it to T-bet binding sites without the requirement for GATA3 DNA binding function. To distinguish between these models, we measured the recruitment of a previously identified GATA3 DNA binding mutant (R352A) to T-bet-dependent binding sites in the absence and presence of T-bet. We found that while the R352A mutation reduced GATA3 binding to a canonical GATA3-containing site at II4 CNS2, it had no effect on T-bet mediated GATA3 recruitment to Ifng (Figure 5B). Thus, together with our T-bet DNA binding mutant ChiP data (Figure 4D), these results demonstrate that T-bet DNA binding is required for T-bet-mediated GATA3 recruitment while GATA3 DNA binding activity is dispensable. This suggests that GATA3 does not directly bind DNA at T-bet-dependent binding sites and is instead associated with these sites through interaction with T-bet.

DNA binding and GATA3 interaction are tightly linked aspects of T-bet function

Given that the T-bet DBD is sufficient for interaction with GATA3, we considered how T-bet’s GATA3 binding activity was related to its DNA binding activity. The two residues in TBX5 (G80R and R237W/Q) previously shown to be required for interaction with GATA4 [81] are also required for TBX5 DNA binding [81,83,84], indicating the two activities may be tightly linked. To understand the relationship between T-bet’s GATA3 and DNA binding activities in more detail, we utilised a previously performed in silico docking of the crystal structures of the TBX5 DBD and the GATA3 DBD [85]. This analysis identified twelve amino acids in TBX5 that were located at the GATA3 DBD interaction surface. Using sequence and structural homology search against a murine T-bet DBD structure [86], we mapped ten of these amino acids to positions within T-bet, while we were unable to find equivalent residues for two others (TBX5 A130 and M131) (Figure 6A). Notably, the ten residues that could be mapped to murine T-bet all cluster together in an area close to the DNA below the dimerization interface (in the crystal structure T-bet forms a dimer that bridges two DNA molecules [86]).

Based on the hypothesis that these ten residues would contribute to the interaction of T-bet with GATA3, we generated point mutations to test their effect on GATA3 interaction. We excluded three amino acids because they were either known to contact DNA phosphates (S261) or were located next to amino acids that do so (L262 and A303) [86]. Furthermore, we included two additional residues in this analysis: I257, because of its location within the same region predicted to contact GATA3, and R324, equivalent to R237 in TBX5, because of its importance for both DNA binding and interaction with GATA4 [81,83,84]. To determine the contribution of these residues to the interaction of T-bet with GATA3, we generated mNeonGreen (mNG)-tagged mutant T-bet proteins by changing the WT residues to amino acids of altered electrical charge, polarity, hydrophilicity or size. Additionally, we included the known T-bet DNA-binding mutant R163A/R164A [79] to assess the importance of T-bet DNA binding activity for its interaction with GATA3. Flow-cytometric FRET measurements of HEK293T cells transfected with mTagBFP-tagged GATA3-DBD and mNG-tagged WT or mutant T-bet DBD revealed a ~50% reduction in the interaction of T-bet with GATA3 while I257E completely abrogated interaction with GATA3 (Figure 6B and Supplementary Figure S6A). The inability of the I257E mutant to interact with GATA3 was confirmed by fluorescence lifetime imaging microscopy (FLIM)-FRET (Supplementary Figure S6B). In addition, the R163A/R164A and R324W mutations also completely abrogated interaction with GATA3, indicating that residues required for DNA binding are also involved in the interaction with GATA3.

To confirm that the reduction in FRET activity reflected reduced interaction between T-bet and GATA3, we also performed co-IP experiments with mNG-T-bet DBD and HA-tagged GATA3 in cell lysates that were treated with benzonase to degrade DNA that could potentially mediate indirect interactions between the proteins (Figure 6C). These experiments confirmed that the interaction of the DNA-binding mutants R163/4A and R324W with GATA3 were diminished compared to the WT T-bet DBD, with T-bet Q160A, I257E and Q305W also exhibiting reduced GATA3 interaction. These results demonstrate that residues in the T-bet DNA binding domain are required for interaction with GATA3. Thus, we next determined the effect of these mutations on T-bet DNA binding activity using our previously developed in vitro OligoFlow assay [77]. Biotinylated DNA oligos incorporating a synthetic T-box motif were conjugated to streptavidin-coated polystyrene beads, incubated with cell lysates from HEK293T cells transfected with mNG-tagged T-bet and fluorescence intensity measured by flow cytometry (Figure 6D). This revealed that all the mutants that displayed reduced interaction with GATA3 also showed a corresponding reduction in DNA binding activity. We conclude that T-bet residues required for interaction with GATA3 are also required for DNA binding, indicating that T-bet’s GATA3 binding and DNA binding activities are tightly linked.

We next asked whether GATA3’s T-bet binding and DNA binding activities were also tightly linked in the same way. We selected four GATA4 mutants [C273G, G296S and K299A + H301L [81,87,88]] that were previously shown to impair both DNA binding and interaction with TBX5 and identified the equivalent residues in GATA3 (C320G, G342S, K346A + H348L) by aligning the primary amino acid sequence of the two proteins. Co-IP of mNG-tagged T-bet and HA-GATA3 in HEK293T cells showed that G342S and K346A + H348L maintained interaction with T-bet but that C320G showed a strongly reduced interaction (Figure 6E). We then assessed the effect of the C320G mutation on GATA3 DNA binding by ChiP. We found that occupancy of HA-GATA3 C320G at the II4 locus in HEK293T cells was strongly reduced compared to that of WT HA-GATA3 (Supplementary Figure S6C). These data indicate that GATA3 interacts with T-bet through a subset of the residues that mediate its interaction with DNA and, taken together with the results from the T-bet mutants, this is con-
Figure 6. T-bet residues required for interaction with GATA3 are also required for DNA binding. (A) Structural alignment of the mouse T-bet DBD (PDB ID 5T1J; grey) and DNA-bound human TBX5 DBD (PDB ID 2×6U; orange) crystal structures. TBX5 residues in close proximity to GATA3 according to molecular docking of the TBX5 and GATA3 crystal structures (81) are highlighted in red and the equivalent T-bet residues are in green. (B) FRET efficiency (DFRET; relative to mNG-T-bet and mTagBFP) measuring interaction between WT or mutant mNG-T-bet DBD and mTagBFP-GATA3 DBD in HEK293T cells measured by flow cytometry (representative of two biological replicates). (C) Immunoblots of HA-GATA3, mNG and β-tubulin in input samples and mNG immunoprecipitates from HEK293T cells expressing full-length (FL) GATA3-HA with NLS-mNG or WT or mutant mNG-T-bet DBD (representative of two biological replicates). (D) Binding of mNG-tagged WT and mutant T-bet DBD to biotinylated T-box motif-containing DNA oligos measured by flow cytometry. For each protein, the mean fluorescence intensity (MFI) from non-specific binding to DNA oligos lacking T-box motifs was subtracted. Representative of two biological replicates. (E) Immunoblots of HA-GATA3, mNG and β-tubulin in input samples and mNG immunoprecipitates from HEK293T cells expressing WT or mutant HA-GATA3 with NLS-mNG or mNG-T-bet. Representative of two biological replicates.

consistent with a model in which the interaction between T-bet and GATA3 is mediated by their DNA binding surfaces.

DISCUSSION

LD-TFs that drive mutually exclusive cell differentiation programs are frequently co-expressed in cells but the impact of this on transcription factor function and cell state are unclear. Using co-expression of the T_H1 and T_H2 cell LD-TFs T-bet and GATA3 as a model, we have found that T-bet directly interacts with GATA3, redistributing it from its canonical sites at T_H2 genes to a new set of binding sites at T_H1 genes, driving T_H1 gene expression while silencing the T_H2 gene expression program (Figure 7).

The mechanism of T-bet-mediated GATA3 redistribution that we have defined here likely has important consequences for the regulation of CD4+ T cell differentiation. GATA3 adopts an alternative binding profile in primary human T_H1 cells, which is characterised by depletion of GATA3 from T_H2 genes and gain at T-bet binding sites that lack GATA motifs at T_H1 genes (29). Thus, despite the differences between the cell types, the GATA3 redistribution mechanism that we describe in EL4 cells also appears to operate in primary T cells. However, directly testing the importance of T-bet-mediated GATA3 redistribution to T_H1 and T_H2 differentiation in vivo is complicated by our discovery that the same residues are required for T-bet’s GATA3 interaction and DNA binding functions, which therefore cannot be separated.

The question arises as to the advantage of the GATA3 redistribution mechanism over other possible means by which the T_H2 gene expression program could be repressed. Al-
though T-bet is a transactivator, T-bet-mediated GATA3 redistribution allows T-bet to mediate both activation of TH1 genes and the repression of TH2 genes. The ability of T-bet to repress GATA3 function in this way likely relates to the addition role of GATA3 in T cell maturation, the outcome of which is that GATA3 is already present in cells at the time that T-bet becomes expressed. Repressing TH2 genes by redistributing GATA3 rather than silencing GATA3 expression completely may also provide a degree of plasticity in which a reduction in T-bet expression levels could release GATA3 from sequestration and allow re-activation of TH2 genes. Consistent with this possibility, we found that reducing the level of T-bet relative to GATA3 reduces the extent to which GATA3 is redistributed.

We have demonstrated that the redistribution of GATA3 to T-bet binding sites requires T-bet DNA binding function but not GATA3 DNA binding function. This suggests that GATA3 is tethered to T-bet binding sites by T-bet rather than directly contacting the DNA itself. However, although T-box motifs were the only motifs that exhibited stronger enriched at T-bet-dependent GATA3 binding sites compared with invariant GATA3 binding sites, we cannot rule out the possibility that an additional T-bet-dependent factor is also required for GATA3 redistribution. Why T-bet only results in the removal of GATA3 from some of its binding sites (‘loss’ sites) and not others (‘invariant sites’) is unclear, but the reason may be that invariant sites are co-occupied by T-bet whereas loss sites are not.

Our data further demonstrate that the interaction of T-bet with GATA3 requires the same residues that T-bet uses to bind DNA. This is consistent with previous findings that TBX5 and GATA4 mutants which lack the capacity to interact with the other factor also display deficient DNA binding activities (81,83,84,87–89). These data demonstrate that the DNA binding domains of transcription factors do not only govern the DNA binding specificity of the proteins but can also mediate interaction with other transcription factors. This dual function of the T-bet DBD raises the question of how the protein can interact with GATA3 and simultaneously recruit it to DNA. The answer may lie in the ability of T-bet to dimerise (86), with one monomer potentially interacting with DNA and the second monomer interacting with GATA3 (Figure 7).

Transcription factor redistribution through the action of another factor is likely to be a common mechanism in T cell differentiation and beyond. In addition to interacting with GATA3, T-bet directly interacts with the TFIH LD-TF BCL6 and targets this factor to a subset of T-bet DNA binding elements in committed TH1 cells (38). However, unlike its sequestration of GATA3, T-bet utilises BCL6 to repress expression of genes to which it is recruited. In addition to the altered GATA3 binding profile we have observed in human TH1 cells (29), GATA3 exhibits alternative binding profiles at different stages of thymocyte development (90,91), suggesting it can be redistributed by other factors in addition to T-bet. Furthermore, interactions also occur between other T-box and GATA family members, notably TBX5 and GATA4 during heart development (81,92), suggesting that the mechanism we describe for T-bet and GATA3 may be common to co-expressed T-box and GATA factors. Interestingly, in induced pluripotent stem (iPS) cells expressing a GATA4 G296S mutant that disrupts interaction with TBX5, TBX5 is lost from super-enhancers of cardiac genes and instead becomes mislocalized in the genome, resulting in disruption of the cardiac gene program (88). Thus, GATA4 may redistribute TBX5 during cardiomyocyte differentiation. The weak binding of T-bet to GATA3 binding sites that we observed when T-bet was expressed at low levels suggests that there may also be physiological contexts in which GATA3 can alter T-bet’s DNA binding profile.

Redistribution of one transcription factor from its canonical binding sites to a different set of sites by another transcription factor may contribute to the widely observed deviation between DNA sequence motifs and transcription factor occupancy and to the differences in the patterns of transcription factor occupancy between cell types or development stages. The redistribution of FLI1 from GATA sites by RUNX1 during embryonic stem cell differentiation (93), the redistribution of SATB1, GATA3 and RUNX1 by PU.1 (94) and RUNX1 by BCL11B (95,96) during thymocyte development provide additional examples of how redistribution of one transcription factor by another can drive gene expression changes that alter cell state.

In summary, we have demonstrated that T-bet directly redistributes GATA3 from its canonical binding sites at TH2 genes to T-bet binding sites at TH1 genes and that this is associated with suppression of the TH2 gene expression program. We propose that this mechanism allows T-bet to suppress GATA3 function in cells in which the two LD-TFs are co-expressed and that similar mechanisms may underlie the interplay between other pairs of co-expressed LD-TFs during cell differentiation and development.

**DATA AVAILABILITY**

All raw and processed ChIP-seq, RNA-seq and ATAC-seq data reported in this study have been deposited in the Na-
nitional Center for Biotechnology Information (NCBI) Gene Expression Omnibus under accession number GSE171410. Accession numbers of previously published T-bet and GATA3 ChIP-seq data sets are: GSM836124, GSE62486 and GSE109109 (57,63,64). Raw flow cytometry data files have been deposited at the FlowRepository under repository ID FR-FCM-Z3VC.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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