Germinal centers (GCs) develop transiently within secondary lymphoid organs upon T cell–dependent antigen exposure and are the source of high-affinity antibody responses. Interactions between activated follicular helper T cells (Tfh cells) and B cells are required for the formation and function of GCs (Crotty, 2014). Intriguingly, the BCL6 transcriptional repressor protein is essential for the formation of both Tfh cells and GC B cells; BCL6-deficient mice fail to develop GCs as the result of cell-autonomous effects in each of these cell types (Cattoretti et al., 1995; Dent et al., 1997; Johnston et al., 2009; Nurieva et al., 2009; Yu et al., 2009). The requirement of BCL6 in both GC B and CD4 T cells has been puzzling because these cells have very different specialized functions and hence there were no obvious parallels pointing to similar BCL6-regulated transcriptional programs in these cell types. GC B cells proliferate rapidly and tolerate genomic damage and stress associated with somatic hypermutation. Tfh cells are a specialized subset of CD4+ T cells that migrate into B cell follicles to provide help to GC B cells via costimulatory receptors and secretion of cytokines (Crotty, 2015).

To date, few genes have been demonstrated to be directly regulated by BCL6 in Tfh cells. For example, BCL6 was shown to repress the PRDM1 locus in both Tfh and GC B cells (Tunyaplin et al., 2004; Johnston et al., 2009). BCL6 repression of PRDM1 prevents differentiation of both cell types and represents a commonality between B and T cells (Shaffer et al., 2000). Most notably, current studies have only addressed BCL6 regulation of rare single loci. Moreover, it is currently not known whether BCL6 acts predominantly as a transcriptional activator or repressor in Tfh cells. Hence, the genome-wide BCL6 transcriptional network and the BCL6 mechanisms of action in GC Tfh cells remain unknown.

Follicular helper T cells (Tfh cells) are required for T cell help to B cells, and BCL6 is the defining transcription factor of Tfh cells. However, the functions of BCL6 in Tfh cells have largely remained unclear. Here we defined the BCL6 cistrome in primary human germinal center Tfh cells to assess mechanisms of BCL6 regulation of CD4 T cells, comparing and contrasting BCL6 function in T and B cells. BCL6 primarily acts as a repressor in Tfh cells, and BCL6 binding was associated with control of Tfh cell migration and repression of alternative cell fates. Interestingly, although some BCL6-bound genes possessed BCL6 DNA–binding motifs, many BCL6-bound loci were instead characterized by the presence of DNA motifs for AP1 or STAT. AP1 complexes are key positive downstream mediators of TCR signaling and external stimuli. We show that BCL6 can directly bind AP1, and BCL6 depends on AP1 for recruitment to BCL6-binding sites with AP1 motifs, suggesting that BCL6 subverts AP1 activity. These findings reveal that BCL6 has broad and multifaceted effects on Tfh biology and provide insight into how this master regulator mediates distinct cell context–dependent phenotypes.

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whereas intergenic (17%) and intronic regions (14%) were also substantially represented (Fig. 1 B). To determine whether the BCL6-binding motif was enriched among these BCL6-binding sites, we performed an unsupervised de novo DNA motif analysis (Heinz et al., 2010). The BCL6 motif was significantly overrepresented among BCL6 peaks from GC Tfh cells \( (P = 10^{-221}) \). Moreover, the BCL6 peak summit (the region of each peak with highest enrichment of BCL6-bound DNA) strongly clustered around the BCL6 canonical DNA–binding motif, further validating this BCL6 GC Tfh ChIP-seq dataset (Fig. 1 C). To gain insight into the biological pathways targeted by BCL6 in GC Tfh cells, we identified the genes associated with BCL6-binding sites in these cells and their biological functions (Fig. 1 D). Genes encoding components of Th1 cell differentiation (Fig. 2 A), Th17 cell differentiation (Fig. 2 B), Th2 cell differentiation (Fig. 2 C), T reg cell differentiation (Fig. 2 D), and migration-associated genes (Fig. 2 E) were highly enriched for BCL6 targets in GC Tfh cells. These results suggest that BCL6 facilitates Tfh cell migration and differentiation and additionally “locks in” the GC Tfh phenotype by antagonizing alternative T cell effector programs. Although it was previously found that BCL6 could bind a few genes associated with alternative cell fates, here we show that BCL6 binds thousands of genes in GC Tfh cells, and those bound loci are highly enriched for genes involved in T cell differentiation fates. Furthermore, the whole genome BCL6 ChIP-seq analysis indicates that BCL6 has multiple redundant ways to inhibit each of the alternative effector T cell differentiation pathways. This helps explain previous observations that BCL6 could impact GATA3-associated Th2 functions and RORyt-associated Th17 functions without evidence of binding GATA3 or RORC directly (Kusam et al., 2003; Nurieva et al., 2009). Of note, we saw no evidence of BCL6 binding to the RORC gene, unlike Yu et al. (2009).

To better understand the mechanisms by which BCL6 directly regulates Tfh cells, we performed a comprehensive study of BCL6 genomic localization and transcriptional effects in primary human Tfh cells. Integration of these and other data revealed a Tfh-specific BCL6 cis-regulatory genome landscape that controls critical T cell–specific pathways, including cell migration and alternative T cell fates. Moreover, BCL6 genomic distribution exhibited distinct and characteristic features. Among these was the surprisingly prominent overlap with the major activating complex AP1, suggestive of a key counter–regulatory relation between these transcription factors in T cells. Our results reveal that BCL6 is a multifaceted regulator of the Tfh lineage, using multiple mechanisms to control Tfh cell biology.

**RESULTS**

**The GC Tfh BCL6 cistrome**

BCL6 is the central regulator of GC Tfh cell differentiation; however, the genome-wide target gene network that BCL6 regulates in these cells remains unknown. To determine the distribution of BCL6-bound cis-regulatory regions in GC Tfh cells (the BCL6 cistrome), we performed BCL6 chromatin immunoprecipitation (ChIP) sequencing (ChIP-seq) of primary GC Tfh cells (CXCR5\(^{hi}\) PD1\(^{hi}\) CD45RO\(^{+}\) CD4 T cells) freshly isolated from human tonsils (Fig. 1 A). Tonsils are a lymphoid organ rich in GCs and GC Tfh cells. Using stringent sequence abundance peak detection thresholds and the overlap of two highly correlated \( (r = 0.75) \) independent biological BCL6 ChIP-seq replicates, we identified 8,523 GC Tfh genomic loci with significant BCL6 binding. These ChIP-seq replicates were performed using chromatin from three GC Tfh isolations to minimize potential binding biases between individual tonsil donors. The BCL6-binding sites were predominantly localized to GC Tfh promoters (66%), whereas intergenic (17%) and intronic regions (14%) were also substantially represented (Fig. 1 B). To determine whether the BCL6-binding motif was enriched among these BCL6-binding sites, we performed an unsupervised de novo DNA motif analysis (Heinz et al., 2010). The BCL6 motif was significantly overrepresented among BCL6 peaks from GC Tfh cells \((P = 10^{-221})\). Moreover, the BCL6 peak summit (the region of each peak with highest enrichment of BCL6-bound DNA) strongly clustered around the BCL6 canonical DNA–binding motif, further validating this BCL6 GC Tfh ChIP-seq dataset (Fig. 1 C). To gain insight into the biological pathways targeted by BCL6 in GC Tfh cells, we identified the genes associated with BCL6-binding sites in these cells and their biological functions (Fig. 1 D). Genes encoding components of Th1 cell differentiation (Fig. 2 A), Th17 cell differentiation (Fig. 2 B), Th2 cell differentiation (Fig. 2 C), T reg cell differentiation (Fig. 2 D), and migration-associated genes (Fig. 2 E) were highly enriched for BCL6 targets in GC Tfh cells. These results suggest that BCL6 facilitates Tfh cell migration and differentiation and additionally “locks in” the GC Tfh phenotype by antagonizing alternative T cell effector programs. Although it was previously found that BCL6 could bind a few genes associated with alternative cell fates, here we show that BCL6 binds thousands of genes in GC Tfh cells, and those bound loci are highly enriched for genes involved in T cell differentiation fates. Furthermore, the whole genome BCL6 ChIP-seq analysis indicates that BCL6 has multiple redundant ways to inhibit each of the alternative effector T cell differentiation pathways. This helps explain previous observations that BCL6 could impact GATA3–associated Th2 functions and RORyt–associated Th17 functions without evidence of binding GATA3 or RORC directly (Kusam et al., 2003; Nurieva et al., 2009). Of note, we saw no evidence of BCL6 binding to the RORC gene, unlike Yu et al. (2009)
but in agreement with Nurieva et al. (2009); in contrast, we observed very robust BCL6 binding to RORA (see below), the other ROR family member that controls Th17 cell differentiation (Yang et al., 2008). A prominent BCL6 peak was also present at the IL17A/F enhancer locus (see below). Although BCL6 bound a network of T reg genes, we observed no binding to the FOXP3 locus. BCL6 ChIP-seq also revealed direct BCL6 binding to the FOXO1 promoter. FOXO1 inhibits Tfh cell differentiation (Xiao et al., 2014; Stone et al., 2015).

A defining feature of Tfh cells is their colocalization with B cells in follicles and GCs. The ChIP-seq data suggest that BCL6 regulation of T cell migration is a major mechanism by which BCL6 controls Tfh biology. Non-Tfh effector cells exit LNs in an S1PR1-dependent manner and migrate to sites of inflammation and infection. In GC Tfh cells, BCL6 bound the S1PR1 gene and a large S1PR1 proximal enhancer. BCL6 also bound the KLF2 promoter. KLF2 is a positive regulator of S1PR1 expression (Carlson et al., 2006), and repression of KLF2 is necessary for Tfh differentiation (Lee et al., 2015; Weber et al., 2015). Tfh cells localize to B cell follicles because they express CXCR5, but also because they down-regulate CCR7 and PSGL1 (SELPLG), which cause localization to the T cell zone. Both SELPLG and CCR7 are highly enriched for BCL6 binding in GC Tfh cells (see below). An additional chemotactic receptor, EBI2 (GPR183), is important for localization of B cells to follicles but specifically outside of GCs (Gatto et al., 2009; Pereira et al., 2009) and is repressed by BCL6 in GC B cells (Shaffer et al., 2000; Huang et al., 2014). EBI2 is also likely important in Tfh localization, as EBI2 expression is specifically reduced in GC Tfh cells (Rasheed et al., 2006; Ma et al., 2009; Locci et al., 2013). Combined with the recognition that BCL6 expression results in up-regulation of CXCR5, CXCR4, and SAP in human CD4 T cells (whereas up-regulation of CXCR5 in vitro on mouse CD4 T cells does not depend on BCL6 (Liu et al., 2014), but most CXCR5 expression in vivo is BCL6 dependent (Poholek et al., 2010; Choi et al., 2011)), these data indicate that BCL6 may participate in the control of most aspects of GC Tfh migration.

**BCL6 directly represses a broad network of promoters**

To understand the cis-regulatory landscape of GC Tfh cells, we next performed ChIP-seq for the epigenetic marks H3K4me1, H3K4me3, and H3K27ac, which are histone modifications that mark promoters and enhancer regions. These ChIP-seq experiments were again performed using primary human GC Tfh cells, directly ex vivo. Ranking of GC Tfh gene promoters by decreasing BCL6 binding density within ±5 kb of known transcriptional start sites (TSSs) showed that the bulk of BCL6 binding in GC Tfh cells occurs in promoters enriched

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**Figure 2.** BCL6 targets Tfh genes mainly involved in T cell signaling, differentiation, and migration pathways to drive the GC Tfh phenotype. (A–E) Graphical representation of BCL6 targeted pathway components involved in T cell receptor signaling (A), Th17 cell differentiation (B), Th2 cell differentiation (C), T reg cell signaling (D), and T cell migration (E). Proteins encoded by genes bound by BCL6 are shown in blue. Data are from two experiments, comprising primary Tfh cells from three human donors.
in H3K4me1, H3K4me3, and H3K27ac, which is a signature characteristic of actively transcribed genes (Fig. 3 A). To assess the link between BCL6 binding and nucleosome positioning at promoters, we determined the mean H3K4me1, H3K4me3, and BCL6 enrichment profile in these regions. This promoter chromatin profile indicates that BCL6, on average, occupies the nucleosome-free region just upstream of the TSS (Fig. 3 B). Enrichment of BCL6 binding to regions with active chromatin marks was unexpected, as BCL6 is known as a repressor, and it is not known to activate gene expression. Hence, its binding to genes with active chromatin marks may signify that BCL6 predominantly represses or dampens the expression of transcriptionally activated, or transcriptionally poised, genes in Tfh cells. Alternatively, BCL6 may be acting as a transcriptional activator in GC Tfh cells.

To explore these different possibilities, we performed experimental perturbations of BCL6 expression. Purified primary naive human CD4 T cells were transduced with a BCL6-expressing lentiviral vector (BCL6-LV; Fig. 3 C), and gene expression profiling was performed. Gene expression analysis at day 5 after transduction revealed that 457 genes with BCL6-bound promoters were repressed in CD4 cells upon expression of BCL6 (>1.25-fold, false discovery rate [FDR] < 0.05). There was a trend for promoters with BCL6 peaks containing BCL6 motifs to be repressed after induction of BCL6 (normalized enrichment score [NES] = −1.18).
example is loci in GC Tfh cells but was not enriched at the corresponding BCL6 bound to intergenic sites upstream of those genes repressed by BCL6 (FDR = 0.005, NES = −1.56; Fig. 3 D). These results are consistent with BCL6 acting primarily as a direct transcriptional repressor in GC Tfh cells. KLF2 expression was the most strongly repressed gene transcript overall. BCL6 target promoters with BCL6–binding sites included the genes IFNGR1, STAT4, GATA3, and RORA (Fig. 3 E), which play key roles in differentiation of Th1, Th2, and Th17 cells. BCL6 also bound a BCL6 DNA–binding motif in the promoter of GIMAP1, a regulator of T cell proliferation. We observed a significant reduction of RORA, GIMAP1, and STAT4 expression in BCL6-LV+ CD4 T cells (P < 0.001) and a moderate reduction of IFNGR1 and GATA3 (Fig. 3 F). Furthermore, quantitative PCR (qPCR) analysis of GC Tfh cells from different donors revealed reduced transcript abundance of each of these genes in GC Tfh cells compared with naive T cells (Fig. 3 G).

BCL6 represses different subsets of promoters in Tfh cells and GC B cells

GC B cells are phenotypically very different compared with Tfh cells. Given that BCL6 is a lineage-defining transcription factor of both cell types, it is logical to posit that it regulates different gene sets in each case. Interestingly, comparison of BCL6–binding sites using ChIP-seq from human primary GC B cells (Huang et al., 2013) reveals that approximately half of GC Tfh BCL6–binding sites (4,321 peaks) are shared between GC B and GC Tfh cells (Fig. 4 A). Nonetheless, a large fraction of BCL6 peaks were specific to each cell type. 49% of GC Tfh (n = 4,202) and 66% of GC B cell BCL6 peaks (n = 10,133) were unique to B and T cells, respectively (Fig. 4 A). These GC B–only and GC Tfh–only BCL6 peaks had low enrichment of BCL6 binding in the other cell type (Fig. 4, B and C). This suggests that BCL6 regulates both common and cell context–dependent functions. Notably, the vast majority of common GC Tfh B cell BCL6 peaks were localized to promoter regions (76%; TSS ± 2 kb; Fig. 4 D). The BCL6 and PRDM1 promoters were among those with robust BCL6 promoter binding in both GC Tfh and GC B cells (Fig. 4 E). A large fraction of the GC Tfh–only and GC B cell–only BCL6 peaks were in intergenic and intronic sites, suggestive of cell type–specific enhancers (Fig. 4, F and H). For example, several intergenic GC B cell–only BCL6–binding sites were present near the SYK and MSH6 loci in GC B cells but were absent in GC Tfh cells (Fig. 4 G). On the contrary, BCL6 bound to intergenic sites upstream of IL21 and PLCG1 loci in GC Tfh cells but was not enriched at the corresponding locations in GC B cells (Fig. 4 G). Another interesting example is SELPLG, a regulator of Tfh migration, for which there were different BCL6–bound loci in GC Tfh and GC B cells (Fig. 4 J). These results were validated by qChIP experiments using chromatin from GC B and Tfh cell isolations from several independent donors (Fig. 4 K).

Because most of the cell context–dependent BCL6 targets occur outside gene promoters, we examined the chromatin architecture surrounding these sites. We found that the chromatin marks H3K4me1 and H3K27ac, which are associated with enhancers, were selectively enriched in the GC B–only and GC Tfh–only BCL6 loci in B and T cells, respectively, but were largely absent in the opposite cell type (Fig. 5). On the contrary, both of these histone marks were enriched in the smaller subset of common T–B peaks both in T and in B cells. This result highlights that the role of BCL6 is associated with unique cell context–specific chromatin landscapes.

Given that BCL6 binding does not necessarily equate to transcriptional regulation, we wanted to know whether these promoters were also regulated by BCL6 in both B and Tfh cells. 457 BCL6 Tfh promoter target genes were significantly repressed by BCL6 induction in CD4 T cells, as noted earlier (Fig. 3 D). We also observed that 518 BCL6 promoter target genes were significantly up-regulated after siRNA-mediated BCL6 knockdown in GC–derived B cells (>1.5-fold, FDR < 0.05). We observed that despite the extensive overlap (70%) of BCL6 binding at B and Tfh cell promoters, only 72 genes (16%) had evidence of repression by BCL6 in both cell types. BCL6 repressed PRDM1, SIP4R, CD69, LPP, FAIM3, PTEN, CASP8, FOXO3, and CDKN1B in both GC B and Tfh cells, among other genes. The common module may be required for migration of these cells into the follicle after GC chemokine gradients or may reflect common signaling cues from the GC microenvironment. Several common components of the B and T cell receptor pathway were also targeted by BCL6 in both cells. Overall, although there are limitations to this analysis, these data indicate that many bound promoters are only repressed by BCL6 in either GC Tfh or GC B cells. BCL6 represses gene expression via recruitment of corepressors (Hatzi and Melnick, 2014), and thus these data imply a possible important role for differential expression or utilization of BCL6 corepressors in Tfh cells versus GC B cells. Alternatively, pioneering complexes and distinct chromatin nuclear topology might regulate the accessibility of specific loci to BCL6 complexes.

BCL6 targets a network of GC Tfh cell–specific enhancers

34% of BCL6–binding sites in GC Tfh were in introns or intergenic loci, indicating possible association with enhancers. Enhancers can be defined as discrete promoter–distal (upstream or downstream) genomic regions enriched in H3K4me1 but depleted of H3K27me3. The enhancer landscape (histone modifications) of human Tfh cells is distinct from non–Tfh cells (Weinstein et al., 2014). To ascertain whether nonpromoter BCL6–binding sites in GC Tfh cells corresponded to enhancers, we determined the overlap of BCL6 GC Tfh peaks with regions significantly enriched in H3K4me1 and lacking H3K27me3. From this analysis, 1,016 BCL6–binding sites mapped
Figure 4. **BCL6 binding and gene regulation in Tfh cells differs compared with GC B cells with distinct pattern mainly outside promoters.** (A) Overlap of BCL6-binding sites in Tfh (blue) and GC B cells (green). (B) Normalized BCL6 ChIP-seq read densities plotted for Tfh cells (y axis) versus B cells (x axis) for each peak corresponding to peaks common to Tfh and B cells, Tfh-only peaks, and B cell-only peaks. Density values were normalized to the total number of reads (rpm, reads per million). (C) Boxplots comparing BCL6 Tfh and GC B BCL6 ChIP-seq read densities for B cell-only peaks, Tfh-only peaks, and peaks common to Tfh and B cells. Density values were normalized to the total number of reads. (B and C) Input chromatin density was subtracted.

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**A**

BCL6 GC Tfh  BCL6 GC B
49% 4,202 4,321 10,985 28%
GC B-Tfh common peaks

**B**

(A) Overlap of BCL6-binding sites in Tfh (blue) and GC B cells (green). (B) Normalized BCL6 ChIP-seq read densities plotted for Tfh cells (y axis) versus B cells (x axis) for each peak corresponding to peaks common to Tfh and B cells, Tfh-only peaks, and B cell-only peaks. Density values were normalized to the total number of reads (rpm, reads per million). (C) Boxplots comparing BCL6 Tfh and GC B BCL6 ChIP-seq read densities for B cell-only peaks, Tfh-only peaks, and peaks common to Tfh and B cells. Density values were normalized to the total number of reads. (B and C) Input chromatin density was subtracted.

**C**

**D**

Tfh - GC B common peaks
9% 13% 76%

**E**

(GC Tfh BCL6seq) 10.7
(GC Tfh BCL6seq)

**F**

GC B only peaks
36% 35% 25%

**G**

(BCL6) 20kb

**H**

(GC Tfh only peaks)
19% 21% 57%

**I**

(SYK) 60kb

**J**

(BCL6) 12kb

**K**

(SELPLG) 20kb

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**K**

GC B BCL6 ChIP
GC Tfh BCL6 ChIP

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to GC Tfh enhancer elements. The mean pattern of BCL6 and H3K4me1 enrichment at these sites suggests that BCL6 binds between nucleosomes and directly accesses DNA containing its cognate motif localized there (Fig. 6 A). Many enhancers with critical gene regulatory functions are conserved among species. We found that the conservation index of BCL6-bound enhancers was highly significantly increased compared with random loci (P < 0.001; Fig. 6 B), suggesting that these are functionally relevant binding sites. Enhancers are known to mediate cell context-specific gene regulation. Hence, we next asked whether the enhancers occupied by BCL6 in Tfh cells are also bound by BCL6 in GC B cells. The majority of GC Tfh BCL6-bound enhancers (743/1,016; 73%) were specific to Tfh cells as they were not bound by BCL6 in GC B cells. These loci were not marked by H3K4me1 or H3K27ac in GC B cells, indicating that these loci have a gene regulatory role in Tfh cells but not in B cells (Fig. 6 C). These findings suggest that BCL6 mediates distinct functions in GC Tfh versus GC B cells by targeting cell type-specific enhancers.

Enhancers are defined as being in active or poised configuration based on the presence or absence of H3K27ac. The majority of BCL6-bound GC Tfh enhancers (76%, 757) were depleted of H3K27ac, reflecting an inactive (poised) conformation (Fig. 6 D), suggesting a putative role of BCL6 in enhancer silencing in Tfh cells. To test how BCL6 enhancer regulation may affect gene expression, we performed GSEA analysis of the gene set of the poised enhancer proximal genes, assessing ranked gene expression changes after induction of BCL6 in CD4 T cells using BCL6-LV+ or control lentivirus. We found a highly significant enrichment in repression among total genes associated with BCL6-bound poised enhancers in from each measurement. (D) Pie chart of the genome-wide distribution of BCL6 peaks common in GC B cells and GC Tfh cells based on RefSeq. Peaks occurring within ±2 kb of the TSS and TES were considered promoter and 3' UTR peaks, respectively. (E) BCL6 density tracks of BCL6 and PRDM1 loci that were commonly bound in GC Tfh and GC B cells. Read densities are shown in blue for GC Tfh BCL6 ChIP-seq and green for GC B BCL6 ChIP-seq. (F) Pie chart of the genome-wide distribution of GC B-only BCL6 peaks. (G) BCL6 density tracks of SYK and MSH6 that were bound by BCL6 in GC B cells but not in GC Tfh cells. (H) Pie chart of the genome-wide distribution of Tfh-only BCL6 peaks. (I) BCL6 density tracks the IL27 and PLCG1 loci that were bound by BCL6 in GC Tfh cells but not in GC B cells. (J) Tfh and GC B BCL6 read density on the SELPLG locus where BCL6 binds at different sites. ChIP-seq data are from two experiments, comprising primary CD4 T cells from three human donors. (K) qChIP experiments confirm BCL6 is selectively enriched at the SYK and IL27 loci in GC B and GC Tfh cells, respectively. BCL6 binds the PRDM1 promoter in both B and T cells. Nonspecific IgG antibody was used as a negative immunoprecipitation control. Data for cells from three separate donors are shown, representative of two independent experiments. Values are shown as percentage of input chromatin. The BCL6 intron 9 served as a negative control locus. Error bars indicate SEM.
Figure 6. **BCL6 targets T cell–specific enhancers associated with transcriptional repression and a poised chromatin configuration.** (A) BCL6 and H3K4me1 ChIP-seq density profiles derived from GC Tfh cells (blue). Enrichment represents the mean normalized read density in Tfh-only BCL6-bound enhancers. (B) Mean conservation score (placental mammal phastCons) of BCL6-bound enhancers relative to the BCL6 peak summit. Random regions were used as a control. (C) Heat maps representing BCL6, H3K4me1, and H3K27ac normalized read density in GC Tfh-only BCL6 enhancers in GC Tfh cells. (D) Overview of BCL6-targeted GC Tfh enhancers: "poised" enhancers (H3K27ac_{LOW}) and "active" enhancers (H3K27ac_{HIGH}). (E) Heat map of ranked gene expression changes (BCL6-LV / Ctrl-LV). (F) Enrichment score heat map of genes with BCL6-poised enhancers. (G) GC Tfh BCL6-seq heat maps for IL17A, IL17F, TBX21, and IL7R. (H) Fold repression (normalized to GADPH) of IL7A, RUNX3, TBX21, and IL7R.
GC Tfh cells (NES = −1.80, FDR < 0.001; Fig. 6 E). Enhancers containing BCL6 DNA–binding motif genes bound by BCL6 (linked to 170 genes) were equally enriched for repression in BCL6-expressing CD4 T cells (NES = −1.74, FDR = 0.001; Fig. 6 E). To understand what type of transcriptional programs BCL6 regulates though enhancer binding, we performed pathway analysis on the set of genes linked to poised BCL6 GC Tfh enhancers. We found that these genes were significantly enriched in biological pathways relevant to T cells biology, including T cell activation, Th17 biology, Th1/Th2 cell differentiation, T cell receptor signaling, protein kinase cascade, and Jak–Stat signaling (Fig. 6 F). T cell–specific gene enhancers containing a BCL6 motif directly bound by BCL6 in GC Tfh cells included TBX21, RUNX3, IL7R, and the IL17A/F enhancer (Fig. 6 G). Each of these genes is repressed in human GC Tfh cells (Fig. 6 H). The IL17A/F enhancer, in particular, is known to be a critical regulator of Th17 function (Yang et al., 2011). The reduction in H3K27 acetylation at BCL6-bound enhancers suggests that BCL6 may promote histone deacetylation at these sites to antagonize p300 histone acetyltransferase activity. This could occur via BCL6 recruitment of histone deacetylase–containing complexes such as SMRT/NCOR or NuRD (Hatz and Melnick, 2014). To further investigate this finding, we asked how the chromatin surrounding the BCL6–poised enhancers changes during T cell differentiation. Therefore, we generated and compared chromatin profiles of the enhancer histone marks H3K4me1 and H3K27ac in naive CD4 T cells (CD4+CD25−CD45RA+; Andersson et al., 2014) and GC Tfh cells (Fig. 7). Enrichment of both marks decreased in Tfh cells compared with CD4 naive T cells. This finding supports a potential role of BCL6 in decommissioning T cell enhancers during differentiation. Collectively, these data reveal that BCL6 mediates its actions in GC Tfh cells by repressing networks of gene promoters and enhancers.

**Extensive AP1 and BCL6 interactions at promoters and enhancers in GC Tfh cells**

The canonical function of BCL6 involves repression of genes by directly binding to cis-regulatory elements containing a BCL6 DNA–binding motif (Figs. 3 E and 6 G). Strikingly, the vast majority of BCL6–bound loci in GC Tfh cells, 88%, lacked a BCL6 DNA–binding motif. We considered that BCL6 may be primarily recruited to DNA by other transcription factors in GC Tfh cells. To test this hypothesis, we first performed an unbiased DNA motif discovery analysis. The BCL6 DNA motif was the top motif observed (P = 10−223), observed in 1,043 peaks, 12%; motif shown in Fig. 1 C; TTCTCTAGAAAGC), but in addition, AP1 (P = 10−112; Fig. 8 A) and STAT transcription factor motifs (P = 10−80; Fig. 8 A) were also highly ranked and highly enriched among BCL6-bound peaks in GC Tfh cells. To ascertain whether these BCL6–binding sites were bona fide STAT and AP1 targets in T cells, we cross-referenced the peaks to published ChIP-seq datasets from activated CD4 T cells. This analysis indicated that a majority of BCL6–binding peaks containing STAT– or AP1–binding motifs are indeed bound by STAT3 and AP1 family proteins (Fig. 8 B). Promoters or enhancers with consensus STAT motifs (TTCC[\{N1-3\}]GAA) within BCL6 peaks were significantly associated with repression by BCL6 in BCL6–L5+ T cells (FDR = 0.003 and FDR < 0.001; Fig. 8 C). STAT proteins mold the enhancer landscape of helper T cells (Vahedi et al., 2012), and it is known that STAT consensus motif sequences can often be observed embedded in BCL6 DNA motifs. Accumulating evidence suggests that in B cells and macrophages BCL6 may antagonize STAT signaling (Dent et al., 1997;
Therefore, a competition between certain STATs and BCL6 in Tfh cells may regulate promoter and enhancer activities, such as the IL17A/F enhancer, which is known to be regulated by competition between STAT3 and STAT5 (Yang et al., 2011).

AP1 factors are transcriptional activators that are crucial in regulating proliferation and cytokine production in T cells and have also been shown to play prominent roles in Th1, Th2, and Th17 cell differentiation (Rincón and Flavell, 1994; Wagner and Eferl, 2005; Schraml et al., 2009). Strikingly, 83% of the 889 BCL6 GC Tfh peaks containing consensus AP1 sites (TGACTCA or TGACGTCA) are occupied by AP1 in activated CD4 T cells (Fig. 8 B). To understand potential mechanistic links between AP1 transcription factors and BCL6 in GC Tfh cells, we first examined how BCL6 affected expression of genes with AP1 motif–associated BCL6 peaks. 454 genes possessed AP1 motif BCL6 promoter peaks, and 177 genes had AP1 motif BCL6 poised enhancer peaks. GSEA analysis found that genes associated with AP1/BCL6 promoters were significantly associated with repression in BCL6+ CD4 T cells (FDR = 0.035; Fig. 8 D). GSEA analysis also found that genes associated with AP1/BCL6 poised enhancers were significantly associated with repression in BCL6+ CD4 T cells (NES = -1.434, FDR = 0.019; Fig. 8 D). Similar to what we observed for the BCL6 DNA motif, both the AP1 and the STAT consensus DNA motifs occurred more frequently in peaks with the most robust amounts of BCL6 enrichment (Fig. 8 E). Notably, most Tfh BCL6 peaks with AP1 motifs did not contain BCL6 motifs within the same BCL6 peak (742/889; 84%). More specific analysis of promoters and poised enhancers also found that AP1 motif–associated BCL6-bound peaks did not overlap with genes that had BCL6 DNA motif peaks (479/617; 78%; Fig. 8 F). Thus, this results in the interesting conclusion that the BCL6 DNA–binding motif and AP1 DNA–binding motif appear to be largely independent mechanisms used to recruit BCL6 to distinct gene sets.

Collectively, these findings suggest that AP1 motifs are highly associated with repression in the presence of BCL6. AP1 factors may recruit BCL6 to genes, resulting in gene activation being converted to gene repression when BCL6 is present. This could have widespread implications for the functions of BCL6 and AP1 within the cell, given the prominent roles of AP1 in cell activation. One previous study reported BCL6 could interact with AP1 family members in B cells at PRDM1 (Vasanwala et al., 2002); however, physical association in cells (coimmunoprecipitation) was not observed, and the overall significance was unclear, with no reported follow-up. We therefore first asked whether BCL6 and AP1 factors physically interact in CD4 T cells. To this end, we used a T cell line (MCC) transduced with a BCL6-expressing retroviral vector (BCL6-RV; Fig. 9 A). AP1 immunoprecipitation showed BCL6–AP1 binding in untransduced BCL6+ RV+ cells (Fig. 9 B). Treatment of untransduced MCC cells with PMA and ionomycin induced robust endogenous BCL6 expression (Fig. 9 A). We therefore immunoprecipitated

Figure 8. BCL6-mediated repression of key Tfh target genes is linked to interaction with AP1 and recruitment to AP1 DNA–binding sites. (A) De novo motif analysis of BCL6 GC Tfh peaks using HOMER identified the AP1 and STAT DNA motifs among the most highly enriched in Tfh BCL6 peaks. P-values are indicated. (B) Human GC Tfh BCL6–binding sites identified in this study containing AP1 or STAT motifs that were homologous to sites in the mouse genome were queried for AP1 and STAT binding based on published Th17 ChIP-seq datasets. Bound versus unbound fractions are indicated. (C and D) GSEA analysis based on global gene expression changes after BCL6 lentiviral induction in CD4 T cells versus control lentivirus. Up and down indicate the relative gene up- or down-regulation after BCL6 expression. Data are from six independent replicates. The gene sets tested were as follows: (C) promoters with BCL6 peaks containing STAT motifs (left) and poised enhancers with BCL6 peaks containing STAT motifs (right); (D) promoters with BCL6 peaks containing AP1 motifs (left) and poised enhancers with BCL6 peaks containing AP1 motifs (right). FDR is based on 1,000 permutations. (E) Fraction of BCL6 peaks containing BCL6, AP1, or STAT motifs in peaks with lower, intermediate, or high BCL6 enrichment. (F) BCL6 and AP1 motif containing BCL6-bound peaks are primarily found in separate sets of gene.

Harris et al., 1999; Huang et al., 2013). Therefore, a competition between certain STATs and BCL6 in Tfh cells may regulate promoter and enhancer activities, such as the IL17A/F enhancer, which is known to be regulated by competition between STAT3 and STAT5 (Yang et al., 2011).
Figure 9. BCL6-mediated repression of key Tfh target genes is linked to interaction with AP1 and recruitment to AP1 DNA–binding sites. (A) Immunoblot of BCL6 and AP1 in MCC cells induced to express BCL6 by retroviral transduction and/or PMA/ionomycin stimulation. (B) Coimmunoprecipitation of BCL6 and AP1 in MCC cells induced to express BCL6 by retroviral transduction. (C) Coimmunoprecipitation of BCL6 and AP1 in MCC cells induced to express BCL6 by PMA/ionomycin stimulation. (A–C) TBP was used as a loading control. Nonspecific IgG served as an immunoprecipitation control.
AP1 with or without PMA and ionomycin, mimicking TCR stimulation. Endogenous BCL6 was enriched in the AP1 immunoprecipitation from stimulated cells but not in IgG control pulldown or in AP1 pulldown from untreated cells (Fig. 9 C). These results indicate that BCL6 and AP1 physically associate in CD4 T cells, in the presence or absence of TCR stimulation.

Among the AP1 motif–associated BCL6-bound genes in GC Tfh were PRDM1 and RUNX1 (promoter BCL6 peaks; Fig. 9 D) and CCR6 and CCR7 (enhancer BCL6 peaks; Fig. 9 F). Expression of each of these genes was significantly reduced in GC Tfh cells (Fig. 9 E and G; Rasheed et al., 2006; Ma et al., 2009). To determine whether AP1 is bound at predicted AP1 sites, we performed ChIP experiments for both BCL6 and AP1. Both BCL6 and AP1 were enriched at the PRDM1 promoter and the CCR6 enhancer in GC Tfh cells (Fig. 9 H). A negative control locus in Bcl6 intron 9 yielded no enrichment of either protein (Fig. 9 H). To determine whether BCL6 and AP1 are co-recruited at these loci, we performed ChIP–reChIP assays in human tonsillar Tfh cells by immunoprecipitation of AP1 (pan-Jun), followed by immunoprecipitation of BCL6. BCL6 was enriched at both the PRDM1 and CCR6 loci (Fig. 9 I), demonstrating co-recruitment of BCL6 and AP1 in vivo.

Finally, to determine whether BCL6 is dependent on AP1 for binding at these loci, we compared BCL6 binding in Bcl6-RV+ MCC cells in the presence or absence of cJun knockdown by shRNAiMIR. The majority of Bcl6 enrichment was lost at AP1 motif–associated Bcl6-bound PRDM1 and CCR6 loci after cJun knockdown (Fig. 9 J). As expected, no difference in Bcl6 enrichment was observed at the promoter region of Bcl6, which contains a Bcl6–binding motif and no AP1 motif (Fig. 9 J). Thus, overall, these results suggest that BCL6 is recruited to many genes in Tfh cells in an AP1–specific manner and may either block AP1 activity or serve as a novel AP1/BCL6 repressor complex.

DISCUSSION

In summary, in recent years BCL6 has been identified as a key regulator of Tfh cell differentiation, yet its mechanism of action in these cells remains largely undiscovered. In this work we identified the genome–wide targets of BCL6 in Tfh cells and the BCL6–regulated Tfh transcriptional program. BCL6 is primarily a repressor in Tfh cells, and it creates a wide umbrella of repression of T cell migration pathways, TCR signaling pathways, and Th1, Th17, Th2, and T reg cell differentiation pathways. Notably, we found that BCL6 is linked to repression of both Tfh promoters and enhancers. Furthermore, it was intriguing to find that BCL6 DNA–binding motifs were only present at ~10% of the bound genes. Given that observation, we then determined that, surprisingly, AP1 motifs were highly enriched within BCL6–bound loci in Tfh cells. Mechanistic experiments showed that BCL6 directly binds AP1 and AP1/BCL6 colocalizes at promoters and enhancers that are repressed by BCL6 in Tfh cells. Altogether, these findings indicate that BCL6 controls Tfh cells via multiple distinct mechanisms, including subversion of AP1. Importantly, this first head–to–head comparison of BCL6 in distinct cell lineages indicates that the BCL6 cistrome is substantially cell context dependent, explaining how this transcription factor can play essential roles in cell types with dramatically different phenotypes. Furthermore, these results provide a foundation for future studies. In particular, studies are warranted to identify the step–wise mechanisms of BCL6–mediated transcriptional repression in these cells, including the corepressor complexes involved in GC Tfh BCL6–mediated transcriptional repression.

BCL6 directly regulates Tfh promoters that are repressed by BCL6 in Tfh cells. Furthermore, these findings indicate that BCL6 controls Tfh cell differentiation and homing could help tailor better vaccination strategies (Crotty, 2014) or facilitate the design of targeted therapies for autoimmune disorders (Craft, 2012) or chronic infections (Butler et al., 2012).

MATERIALS AND METHODS

ChIP. GC Tfh cells (CXCR5hi PD1hi CD45RO- CD4+ T cells, CD19-) were isolated from human tonsils, fixed with 1% formaldehyde, lysed, and sonicated to generate fragments <500 bp. ChIP was performed by incubation of the chromatin with antibodies against BCL6 (N–3; Santa Cruz Biotechnology, Inc.), H3K4me3 (polyclonal rabbit Ab; Abcam), H4K4me1 (polyclonal rabbit Ab; Abcam), and H3K27ac (polyclonal rabbit Ab; Abcam).

E and G–J Error bars indicate SEM. *, P < 0.05; **, P < 0.001; ****, P < 0.0001.
Immunocomplexes were pulled down using protein A beads, and after increasing stringency washes, 10 ng ChiP DNA was recovered and used to generate a BCL6 ChiP-seq library according to the ChiP-seq Library preparation kit (Illumina). A negative control library was prepared in parallel using 10 ng input chromatin DNA. Libraries were quantified and validated using the 2100 Bioanalyzer (Agilent Technologies) for size, concentration, and purity. Both libraries were sequenced using HiSeq 2000 (Illumina) for 50 cycles.

**ChiP-seq data processing and peak detection.** Primary image analysis and base calling were conducted using the Illumina pipeline, and the generated reads were mapped to the human genome (UCSC hg18) using ELAND. Only sequences mapped uniquely to the genome with no more than two mismatches were accepted. Read density tracks were visualized using the UCSC browser, and ChiPseeqer algorithm (Giannopoulou and Elemento, 2011) was used for BCL6 peak calling, compared with total input control. Clonal reads (reads mapping to the same exact location) were excluded from peak calling and generation of read density tracks as amplification artifacts. Genomic regions with minimum twofold enrichment over input and negative log p-value >10 were selected. Peaks were then annotated based on the RefSeq database (hg18). Peaks localized ±2 kb of the TSS were defined as promoter peaks, peaks localized ±2 kb of the transcriptional end site (TES) were defined as 3' end peaks, and peaks >2 kb away from genes were defined as intergenic (Table S2). De novo transcription factor motif analysis was performed using HOMER (Heinz et al., 2010). Conservation analysis was performed using the ChiPseeqerCons module of ChiPseeqer. Conservation scores centered at each peak summit were computed as the mean placentum mammal conservation index (phastCons) extracted from hg18 phastCons Evansion track of the UCSC Genome Browser database. GC content, total GC, and GC distribution were calculated using the ChIPseeqerCons module of ChIPseeqer.

**Human samples.** Fresh human tonsils were obtained from Rady Children’s Hospital of San Diego. Informed consent was obtained from all donors. Tonsils were homogenized using wire mesh and passed through a cell strainer to make a single-cell suspension. Mononuclear cells were isolated using Histopaque 1077 (Sigma Aldrich). All protocols were approved by the La Jolla Institute for Allergy and Immunology, University of California, San Diego, and Rady Children's Hospital of San Diego.

**Cell sorting.** All cells were sorted using a FACS Aria (BD) as previously described (Kroenke et al., 2012). All TfH cell sorts were initially gated on CD4+CD19×, then CD45RO×, and then as CXCR5+ (non-TfH) and CXCR5× (GC-TfH). The following anti-human antibodies were used: CD45RO (clone UCHL1), CD19 (clone HIB19), PD-1 (clone J105), and CD4 (clone RPA-T4; eBioscience); and CXCR5 (clone RFP8B2).

**shRNA mirs.** transOMIC shRNAs are designed using the shERWOOD algorithm, having a proven increasing knockdown potency and specificity at low concentration. shJun shRNAs were cloned into our pLMPd vector as described previously (Chen et al., 2014). Knockdown efficiency was assessed by Western blot. shRNA selected for shJun was transOMIC #RLGM-GU36521 with guide sequence 5′-AGAAACGACCTTCTACGACGAA-3′.

**Cell culture and viral transductions.** MCC cells were maintained in D10 media (DMEM + 10% FCS supplemented with 2 mM GlutaMAX [Life Technologies] and 100 U/ml penicillin/streptomycin [Life Technologies]). Bcl6-expressing retroviral vector (Bcl6-GFP) or empty vector (GFP only) was used to produce virions from the Plat-E cell line as described previously (Johnston et al., 2009). For shRNA transductions, shJun-expressing retroviral vector (shJun-mAmetnies) or negative control vector (shCD68-mAmetnies) was used. Culture supernatants were obtained 2 d after transfections and filtered through 0.45-mm syringe filters. FCC cells were then transduced with retroviral virions two times. Transduced MCC T cells were FACs sorted based on GFP expression levels. For stimulation, cells were treated with 100 ng/ml PMA and 1 µg/ml ionomycin in D10 media for 5 h. Sorted human tonsil T cells were stimulated with anti-CD3/CD28 Dynabeads (Invitrogen) in 96-well flat-bottom plates at a starting density of 7.5 × 10⁴ cells/well. Beads were used at a concentration of 1 ml/well. RPMI 1640 medium with with 10% FCS was supplemented with 2 ng/ml recombinant human IL-7. Cells were split as necessary. Sorted naive T cells were transduced with lentiviral vectors as previously described (Kroenke et al., 2012).

**qPCR.** RNA was isolated by RNeasy spin columns (QiAGEN) and reverse transcribed into cDNA using Superscript II reverse transcription (Invitrogen). Real-time PCR reactions were set up using SybrSelect master mix (Life Technologies). Primers are listed in Table S1.

**ChiP-qPCR.** MCC cells on GC TfH cells were harvested and then cross-linked with 1% formaldehyde. Chromatin was isolated after sonication. Protein G Dynabeads (Life Technologies) were conjugated to antibodies specific to JunB, JunD, cJun, and Bcl6 (Santa Cruz Biotechnology, Inc.). Rabbit IgG was used as a control. Chromatin was immunoprecipitated using the conjugated beads, eluted, and reverse cross-linked using 0.3 M NaCl at 65°C overnight. qPCR was performed as above, and sample values were given as a percentage of input. Primers are listed in Table S1.

**ChiP-reChiP.** MCC cells were harvested and then cross-linked with 1% formaldehyde. Chromatin was isolated after sonication. Protein G Dynabeads were conjugated to antibodies specific to cJun (Abcam). Rabbit IgG was used as a control. Chromatin was immunoprecipitated using the cJun-conjugated beads and eluted, followed by immunoprecipitation with Bcl6-conjugated beads. Chromatin was then reverse cross-linked using 0.3 M NaCl at 65°C overnight. qPCR was performed as above, and sample values were calculated as a percentage of input. Primers are listed in Table S1.

**Gene expression microarrays.** Sorting of tonsil GC TfH cells (CXCR5+PD1+) cells and Tfh (CXCR5+PD1+) cells was previously described (Kroenke et al., 2012). Microarray method and data were as described previously (Locci et al., 2013). Samples from six independent donors were used. For BCL6-LV microarrays, sorted naive tonsil cells were activated with anti-CD3+CD28 antibody– coated beads and transduced with BCL6 or control lentiviral vectors as described previously (Kroenke et al., 2012). RNA was isolated at day 5 after LV infection, and microarrays were performed as previously described (Locci et al., 2013). Samples from six independent donors were used.
Accession numbers. The human GC Tfh and non-Tfh microarray data are available in the GEO database under the accession no. GSE50391 (tonsil CD4+ T cells). GC Tfh ChIP-seq data are available under accession no. GSE59933. BCL6-LV gene expression microarray has been deposited under GEO accession no. GSE66373.

Online supplemental material. Table S1 lists primers used in the study for qPCR and ChIP assays. Table S2, included as a separate Excel file, shows the peak list. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20141380/DC1.

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REFERENCES


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