Covalent modification of histones is a fundamental mechanism of regulated gene expression in eukaryotes, and interpretation of histone modifications is an essential feature of epigenetic control. Bromodomains are specialized binding modules that interact with acetylated histones, linking chromatin recognition to gene transcription. Because of their ability to function in a domain-specific fashion, selective disruption of bromodomain:acetylated histone interactions with chemical probes serves as a powerful means for understanding biological processes regulated by these chromatin adaptors. Here we describe the discovery and characterization of potent and selective small molecule inhibitors for the bromodomains of CREBBP/EP300 that engage their target in cellular assays. We use these tools to demonstrate a critical role for CREBBP/EP300 bromodomains in regulatory T cell biology. Because regulatory T cell recruitment to tumors is a major mechanism of immune evasion by cancer cells, our data highlight the importance of CREBBP/EP300 bromodomain inhibition as a novel, small molecule-based approach for cancer immunotherapy.

Regulatory T cells (Tregs) are essential for the maintenance of immunological self-tolerance, imposing a crucial check on the incidence of autoimmunity (1, 2). Tregs can also accumulate within tumors and are recruited from circulation by chemotactic mechanisms. Increased Treg loads are associated with poor prognosis in several cancer types, such as breast, ovarian, and gastric cancers (3–8). Intratumoral Tregs dampen effector T cell responses to tumor antigens, engendering an immunosuppressive microenvironment, and, as such, represent a major impediment to immune system clearance of tumor cells; specific inhibition of this subset therefore represents a powerful approach for bolstering anti-tumor immune responses, known as cancer immunotherapy (9–12).

Chromatin modifications govern diverse aspects of cellular function (13, 14). Lysine acetylation in histones and non-histone proteins (such as transcription factors) has been identified as a commonly utilized post-translational modification, allowing for widespread, dynamic, and reversible control of gene expression (15). Histone acetylation impacts gene transcription both by direct relaxation of chromatin architecture through alteration of DNA-histone interactions and by providing docking sites for recruitment and assembly of transcriptional complexes (16). Although histone acetyltransferase (HAT) enzymes deposit acetyl marks on histones, a group of conserved proteins, bromodomains (BRDs), has evolved to bind acetyl lysine motifs, forging an essential link between chromatin modifications and transcriptional control (16–18).

The highly homologous, multifunctional mammalian proteins, CREBBP and EP300 (cAMP response element binding-protein binding protein and E1A binding protein 300, respectively; hereafter referred to as CBP/EP300) are especially important in this regard. CBP and EP300 each encode a single protein binding protein and E1A binding protein 300, respectively; hereafter referred to as CBP/EP300) are especially important in this regard. CBP and EP300 each encode a single

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importance of the bromodomain in transactivation functions of CBP/EP300 (23). The conserved BRD fold provides a deep hydrophobic pocket suited to the binding of inhibitors, whereas diversity in surface and loop residues across BRD proteins allows for selective targeting by pharmacologic agents (23).

Here we describe the discovery and characterization of inhibitors of CBP/EP300 bromodomains. These compounds are potent, are highly selective over other bromodomains, and demonstrate target inhibition in cellular assays. A more detailed description of the optimization of this series is reported elsewhere (24). Employing these chemical probes to query bromodomain function, we find that inhibition of CBP/EP300 bromodomains in ex vivo differentiated human Tregs is sufficient to reduce FOXP3 expression, as well as mediators of Treg suppressive function, such as LAG-3, CTLA-4 and TIM-3. Our data reveal the importance of CBP/EP300 bromodomains in maintaining pro-tolerance programs in Tregs and point toward manipulation of CBP/EP300 bromodomain function as an approach to counter immunosuppression in cancer.

**Experimental Procedures**

**Chemical Synthesis**—CPI098 (4-methyl-1,3,4,5-tetrahydro-2H-benzo[bi][1,4]diazepin-2-one) is commercially available from Sigma-Aldrich and was used without further purification. Detailed descriptions of the chemical synthesis of CPI703, CPI644, CPI644(−), and CPI571, along with descriptions of biochemical assays and crystallography are included in the supplemental text.

**Thermal Shift Assay**—All assays were carried out in 384-well plates. In a conical tube, CBP (4 mM) was combined with Sypro Orange (Life Technologies) to a final dye concentration of 5× in 50 mM Tris, 1 mM DTT, pH 8.5. The tube was centrifuged briefly to remove precipitate, and the protein:dye solution was then added to a black Optiplate™ plate (Greiner) and spun briefly (1 min, 900 × g), and then 23 μl of solution was transferred to either DMSO controls or fragments plated from 100 mM DMSO stocks into clear-bottomed Fluotrac200™ plates. In a conical tube, CBP (4 mM) was combined with Sypro Orange (Life Technologies) to a final dye concentration of 5× in 50 mM HEPES, pH 7.5, 150 mM NaCl, 0.1 mg/ml BSA). Streptavidin acceptor beads were added to the protein solution and compounds were equilibrated in the calorimeter cell (15 μM, 200-μl volume, 25.0 °C, stirrer speed of 1000 rpm, 5 μcal/s reference power), and to this was added CBP (150 μM) in the same buffer as a series of sixteen 2.49-μl injections (4.98 s each) with an equilibration time of 120 s between each injection. The resulting data were fitted using the instrument’s Origin 7.0 software.

**AlphaLISA**—Inhibitory activity of compounds was determined by competition with the binding of purified His-FLAG-tagged bromodomains to H4-TetraAc-biotin peptide (New England Peptide) using AlphaLISA technology (PerkinElmer Life Sciences). Compound at varying concentrations were dispensed into 384-well Proxiplates (PerkinElmer Life Sciences) using Echo technology (Labcyte). For the assays, 0.5 μM His-FLAG-tagged CBP bromodomain (amino acids 1082–1197) was incubated with 0.003 μM H4-TetraAc-biotin for 20 min at room temperature in 1× reaction buffer (50 mM HEPES, pH 7.5, 1 mM tris(2-carboxyethyl)phosphine, 0.069 mM Brij-35, 150 mM NaCl, and 0.1 mg/ml BSA). Streptavidin acceptor beads and nickel donor beads (PerkinElmer Life Sciences) were added to 15 μg/ml with a Combi multidrop dispenser. Plates were sealed and incubated at 90 min in the dark at room temperature, and plates were read on an Envision plate reader (PerkinElmer Life Sciences) according to the manufacturer’s instructions. Other bromodomains were assessed by AlphaLISA in a similar manner.

**NanoBRET Cellular Assays**—NanoBRET was carried out using the NanoBRET™ protein-protein interaction system (Promega) according to the manufacturer’s instructions. Briefly, HEK293 cells were transiently co-transfected with a vector for histone H3.3-HaloTag and either a NanoLuc tagged CBP bromodomain, EP300 bromodomain, or full-length BRD4 expression construct. Transfected cells were plated in 96-well plates in the presence or absence of ligand and then treated with dose titrations of indicated compounds. The readings were performed on an Envision plate reader (PerkinElmer Life Sciences), and BRET readings were calculated by dividing the acceptor emission value (600 nm) by the donor emission value (460/50 nm).

**Mesoscale Cellular Assay**—Mesoscale was carried out according to the manufacturer’s instructions (Mesoscale Discovery). CBP/EP300 double knock-out mouse embryonic fibroblasts (25) or HCT116 cells (compound treatment, 16 h) were used. Coat Standard MSD plates (catalog no. L15XA-3) with 30
µl/well capture antibody anti-histone (Millipore catalog no. MAB3422) at 4 µg/ml final concentration was used. The anti-H3K18ac (catalog no. 9675) and anti-histone H3 (catalog no. 4499) antibodies were obtained from Cell Signaling and used at 0.125 µg/ml. Data capture was performed using a MSD Sector Imager 2400 (Meso Scale Discovery).

Acetyltransferase Assay—EP300 acetyltransferase reactions were carried out in 50 µM Tris, pH 8, 100 mM NaCl, 1 mM DTT, 0.1 mM EDTA, 0.069 μM Brij-35, and 0.1 mg/ml BSA. A solution of EP300 (Active Motif) and 1.9 H3-labeled (PerkinElmer Life Sciences)/unlabeled (Sigma-Aldrich) acetyl-CoA was added to each well of a black 384-well plate (Greiner) containing 200 nl of compound (final concentration, 0.8% DMSO). Compounds were preincubitated with enzyme and acetyl-CoA for 30 min. Reactions were initiated by the addition of 200 µM Tris, pH 8, 200 µM EDTA, 160 µM anacarcid acid (Sigma Aldrich), transferred to a 384-well streptavidin FlashPlate (PerkinElmer Life Sciences), and counted by TopCount (PerkinElmer Life Sciences).

Bromo Dot Cellular Assays—U2OS-C413 cells express the bromodomain of CBP in the context of the BRD9 protein. Specifically, amino acid residues 114–253 of BRD9 (UniProt Q9H8M2) have been replaced by amino acids 1064–1205 of CBP (UniProt Q92793). This chimeric protein is expressed in fusion with a C-terminally located ZsGreen (ZsG) fluorescent tag in a tetracycline-inducible manner. For the bromo dot assay, 5000 cells/well were seeded in 384-well imaging plates in the presence of 2 µg/ml doxycycline for 16 h. Compounds diluted in fresh medium were added to the cells for 2 h at 37 °C. The cells were fixed with 4% paraformaldehyde for PBS in 15 min at room temperature. Images of cells were acquired using ImageXpress Micro (Molecular Devices) and processed with the Transfluor Module of MetaXpress software. Foci per nucleus values were obtained by four adjacent images in each well, and there were two technical replicates for each compound concentration. Dose-response curves were generated by plotting these numbers for each compound concentration, and EC₅₀ values were calculated by a four-parameter non-linear regression model in GraphPad Prism. The average values from DMSO-treated wells were used to constrain the bottom of the curve.

Human T Cell Cultures—Leukopak samples were procured from the Biological Specialty Corporation (Colmar, PA), and peripheral blood mononuclear cells were isolated by Ficoll (GE Biosciences) density gradient centrifugation. Naïve CD4⁺ CD45RA⁺ T cells were isolated from peripheral blood mononuclear cells to a purity >98% using Miltenyi naïve human T cell isolation kits (130-094-131; Miltenyi Biotech). Isolated cells were cultured at 10⁶ cells/ml under Treg-polarizing conditions, using human T-activator CD3/CD28 Dynabeads® (11132D; Invitrogen), human TGFβ at 10 ng/ml and human IL-2 at 10 units/ml (100-B and 202-IL, respectively; R&D Biosystems). For unpolarized T₁₀ cultures, isolated cells were cultured with Dynabeads alone, without the addition of exogenous cytokines.

FACS Antibodies—Tregs were stained for CTLA-4 (46-1529-42; eBioscience), CD25 (12-0259-42; eBioscience), LAG-3 (17-2239-42; eBioscience), PD-1 (47-2799-42; eBioscience), TIM3 (25-3109-42; eBioscience), and FOXP3 (12-4777-42; eBioscience). For FOXP3 intracellular staining, cells were fixed and permeabilized using the FoxP3/transcription factor staining kit (00-5523-00; eBioscience). In addition to staining for CTLA-4, LAG-3, PD-1, and TIM-3, CD8 T cells were also fixed and permeabilized using the FoxP3/transcription factor staining kit and stained for Perforin (11-9994-42; eBioscience) and Granzyme B (561142; BD Biosciences). The cells and markers were quantified on the BD FACSCanto™ II flow cytometry analyzer (BD Bioscience), and data were analyzed using FlowJo Software, gating on live cells for all FACS plots shown.

Lentiviral shRNA Knockdown of CBP and EP300—Naïve human T cells were cultured under Treg-inducing conditions as described above and at ~16 h post-activation were infected with lentivirus harboring shRNAs specific for either EP300 or CBP (three independent hairpins per protein were cloned into pLKO.1-based lentiviral vectors (25) (see supplemental Table S2). Lentiviral supernatants were added to T cells in the presence of 8 μg/ml Sequa-brene (S2667-1VL; Sigma) followed by spin infection at 2100 rpm, 90 °C. Transduced cells were selected by the addition of 1 μg/ml puromycin after 24 h; infection rates were monitored by measuring GFP fluorescence.

Global Expression Profiling—Naïve CD4⁺ T cells were treated with 4 µM CPI703 or DMSO under Treg polarizing conditions (described above) for 4 days. Total RNA was prepared by using a Qiagen RNeasy Plus mini kit (with columns for elimination of genomic DNA) according to the manufacturer’s protocol. Samples were processed and hybridized on Affymetrix exon arrays, and the data were acquired at ALMAC Diagnostics. CEL files were analyzed with the Affymetrix Expression Console software.

Treg Suppression Assay—Human Tregs were differentiated in vitro (as described above) for 4 days. Additional IL-2 (at 10 units/ml) was added to the cultures on day 2, and FOXP3 expression was checked on day 4 (~80% FOXP3 positive). The cells were taken off Dynabead stimulation, washed, and counted. Naïve T cells were labeled with carboxyfluorescein succinimidyl ester (C34554; Life Technologies) using the manufacturer’s protocol. Co-cultures of naïve T cells and Tregs were set up at a 1:1 ratio. Human T-Activator CD3/CD28 Dynabeads® were added at a 1:4 ratio of beads to cells.

Deep ChIP Sequencing (ChIP-seq) and RNA Sequencing (RNA-seq)—Naïve human CD4⁺ T cells were treated with 4 µM CPI703 or DMSO under Treg polarizing conditions (described above) for 4 days. For ChIP-seq, the cells were pelleted, fixed in 1% formaldehyde for 10 min, lysed, and sonicated. Chromatin samples were precleared with protein A Dynabeads (Life Technologies) and incubated overnight at 4 °C with anti-H3K18 Ac (9675; Cell Signaling), anti-H3K27 Ac (ab4729; Abcam), and anti-H3K4 Me3 (ab8580; Abcam). Chromatin-antibody complexes were precipitated using protein A Dynabeads followed by washes in radioimmune precipitation assay buffer and Tris/EDTA. Samples were digested with RNase A and treated with proteinase K and 10% SDS, followed by cross-link reversal at 65 °C. DNA was purified using MinElute PCR purification kits (Qiagen).
DNA libraries for ChIP-seq were prepared using Ovation Ultralow DR multiplex system kits (0330-32; NuGEN) followed by Illumina sequencing at the MIT BioMicro Center. For RNA-seq, RNA was isolated using Qiagen RNeasy Plus mini kits and sequenced at Ocean Ridge Biosciences (Palm Beach Gardens, FL).

**Luminex Cytokine Assays**—Cytokines were quantified from 72-h cell supernatants using Luminex multiplex assays (HTH17MAG-14K-12; Millipore), as per the manufacturer’s protocol.

**Cell Viability**—Cell viability was assessed using CellTiterGlo®, which detects any change in the number of viable cells based on quantitation of ATP (G7572; Promega). Absolute live cell numbers were determined by trypan blue staining followed by analysis using the Countess automated cell counter (Invitrogen).

**Real Time RT-PCR**—RNA was purified from cells using an RNeasy Plus mini kit (Qiagen) according to the manufacturer’s protocol. First strand cDNA was synthesized using SuperScript III reverse transcriptase (Invitrogen). Quantitative real time PCR was performed using FastStart Universal Probe master mix (Roche) and TaqMan probes (FOXP3, LAG3, HAVCR2 (TIM3), PDCD1 (PD1), PRF1, GRZMB, and EOMES) on the Stratagene MxPro3005p. Glucose-6-phosphate dehydrogenase was used as housekeeping gene (05-046-246-001; Roche).

**Western Blotting**—Cell pellets were resuspended in radioimmunoprecipitation assay buffer containing complete protease inhibitor mixture (04693116001; Roche), incubated on ice for 15 min, and spun at 12,000 × g. Supernatants thus obtained were boiled in loading buffer for 10 min at 95 °C and loaded on 4–20% Tris-glycine mini gels (EC6028BOX; Novex) followed by overnight transfer at 35 V onto PVDF membranes. The membranes were blocked with 5% milk in TBS, 0.1% Tween. Rabbit anti-human EP300 (sc-584; Santa Cruz) and rabbit anti-human CBP (sc-369; Santa Cruz) were used to detect EP300 and CBP, respectively, with goat anti-rabbit:HRP as secondary antibody. Blots were developed using the Pierce ECL Western blotting substrate.

**Global Expression Profiling Data Analysis**—The gene level RNA algorithm was applied, with the core set of probe sets and the annotation file Huex-1_0-st_v2_na31_hg19 provided by Affymetrix. The data were exported using the function “Export Probe Set Results (pivot table) with Annotations to TXT.” The RMA algorithm provides data in log space. The replicate profiles were averaged, and each probe set was displayed on scatter plot as a point. Gene set enrichment analysis (GSEA) was done using the GseaPreranked tool in GSEA with the following parameters: Score_scheme, Weighted; Make_sets, TRUE; Mode, Max_probe; Gene set collection, C7; Include_only_symbols, TRUE; Set_min, 15; Nperm, 1000; Rand_seed, timestamp. The gene expression data are available at the Gene Expression Omnibus with accession no. GSE66596.

**Immunoprecipitation of FOXP3 and Acetyl-FOXP3 Detection**—Human Tregs were differentiated in the presence of 4 μM CPI703 for 36 h or 4 days. Whole cell extracts were prepared using complete lysis buffer (40010; Active Motif) and FOXP3 was immunoprecipitated using anti-FoXP3 (14-4777-82; eBioscience) directly coupled to NHS-activated magnetic beads (88826; Thermo Fisher; per manufacturer’s protocol), followed by immunoblotting with a pan-acetyl Lys antibody (Ac-K2–100; 9814S; Cell Signaling) or anti-Foxp3. MG132 was added at 20 μM to reduce proteasome-induced FOXP3 degradation ~6 h before preparation of whole cell extracts. Band densities were quantified using ImageJ software.

**Results**

**Identification of Selective Small Molecule Inhibitors of CBP/EP300 Bromodomains**—To identify small molecule ligands for CBP/EP300 bromodomains, we screened a library of ~1200 fragments by differential scanning fluorimetry. Compounds that shifted the melting temperature of the CBP bromodomain by at least 1 °C and demonstrated a dose-response in the melting temperature shift when titrated were considered hits. Compounds were further validated by 13N heteronuclear single quantum coherence NMR, ITC, and a peptide competition AlphaLISA assay (data not shown). These studies identified 4-methyl-1,3,4,5-tetrahydro-2H-benzo[b][1,4]diazepin-2-one (CPI098) as a 14 μM inhibitor of CBP by AlphaLISA that displayed promising selectivity over other bromodomains tested, including 13-fold selectivity over the BRD4 bromodomain 1 (BD1) (Fig. 1A).

The binding mode of the series was established with co-crystal structures of the CPI098 fragment (1.65 Å resolution) and a more advanced derivative CPI703 (1.86 Å resolution) bound to the CBP bromodomain. The lactam carbonyl of the scaffold formed key H-bonding interactions with Asn1168 and, through water, with Tyr1125 (Fig. 2A, left panel). The lactam NH provided a further interaction with Asn1168, helping position the 4-methyl substituent toward the water pocket, in the space normally occupied by the acetyl methyl group of an acetyl lysine ligand (Protein Data Bank code 3P1C). Interestingly, only the R enantiomer of CPI098 was observed in the electron density, despite use of a racemic mixture for crystallization. Assaying individual enantiomers confirmed that this enantiomer was substantially more potent (~50-fold).

Analysis of interactions observed in the complex structure supported structure-activity relationships among compounds with modifications to the CPI098 core. Exploring substitutions around the benzene ring, we determined that a variety of substituents at C6 increased potency (a detailed structure-activity relationship account for this series will be described elsewhere). Of note were two molecules, CPI703 and CPI644, which had IC50 values of 0.47 and 0.18 μM, respectively, in the AlphaLISA assay (Figs. 1B and C, and 2B). A co-crystal structure of CPI703 bound to CBP revealed favorable interactions between the C6 tert-butyl-pyrazole of the compound and the LFP shelf of the bromodomain (Fig. 2A, right panel), accounting for the significant boost in potency. Importantly, these compounds retained selectivity over the BET family and other bromodomain-containing proteins (Fig. 1, B and C). We chose CPI571 (Fig. 2B) as a negative control compound, because it lacked activity on CBP but otherwise demonstrated physicochemical properties very similar to CPI703 and CPI644. The S-enantiomer, CPI644 (−), was found to be significantly less potent than CPI644, with an IC50 value of 6.0 μM in the peptide AlphaLISA assay. The potencies of these compounds were corroborated by ITC, using the isolated bromodomain of CBP (Figs. 1D and 2 and B and C, and data not shown). Extensive profiling by differential scanning fluoro-
rimetry using 19 different isolated bromodomains confirmed the selectivity of CPI703 and CPI644 for the CBP bromodomain (Table 1).

Recent work has shown that a subset of small molecule kinase inhibitors show additional binding and functional inhibition of BET bromodomains (26). To test whether benzodiazepinone
TABLE 1

Differential scanning fluorimetry binding data for CPI703, CPI571, CPI644, and CPI644(−) on isolated bromodomains

| Protein     | CPI703 | CPI571 | CPI644 | CPI644(−) |
|-------------|--------|--------|--------|-----------|
| ASH1L       | 0.00 ± 0.73 | −0.35 ± 0.40 | −0.64 ± 0.25 | −0.23 ± 0.88 |
| BAZ1A       | −0.12 ± 0.20 | 0.00 ± 0.00 | −0.12 ± 0.20 | −0.23 ± 0.20 |
| BAZ2B       | −0.12 ± 0.40 | 0.23 ± 0.53 | 0.12 ± 0.00 | −0.23 ± 0.35 |
| BRPF        | 0.46 ± 0.40 | 0.46 ± 0.53 | 0.58 ± 0.20 | 0.46 ± 0.72 |
| BRD1        | 0.23 ± 0.00 | 0.46 ± 0.20 | 0.46 ± 0.20 | 0.23 ± 0.00 |
| BRD4        | 0.70 ± 0.35 | 0.00 ± 0.00 | 1.39 ± 0.70 | 0.12 ± 0.20 |
| BRD8        | 0.52 ± 0.25 | 0.46 ± 0.20 | 0.87 ± 0.25 | 0.35 ± 0.35 |
| BRD9        | 0.93 ± 0.53 | 0.46 ± 0.53 | 0.70 ± 0.35 | 0.23 ± 0.20 |
| BRG1        | 0.00 ± 0.00 | 0.35 ± 0.35 | 0.12 ± 0.20 | 0.12 ± 0.40 |
| BRPF1       | 0.23 ± 0.00 | −0.12 ± 0.00 | −0.12 ± 0.35 | −0.46 ± 0.35 |
| BRPF3       | 0.35 ± 0.35 | 0.00 ± 0.00 | 0.46 ± 0.20 | 0.35 ± 0.60 |
| BRPF4       | 0.35 ± 0.10 | 0.58 ± 0.00 | 0.70 ± 1.22 | 1.28 ± 0.35 |
| CEC2        | 0.06 ± 0.25 | 0.06 ± 0.25 | 0.12 ± 0.20 | 0.93 ± 0.00 |
| GCN5        | 0.00 ± 0.20 | 0.00 ± 0.20 | −0.12 ± 0.20 | −0.12 ± 0.20 |
| MLL         | 0.00 ± 0.20 | 0.58 ± 0.35 | 0.70 ± 0.53 | 0.46 ± 0.20 |
| PCAF        | 0.81 ± 0.20 | 0.35 ± 0.35 | −0.23 ± 0.53 | 0.00 ± 1.04 |
| TRAF1D1−D2D | 0.35 ± 0.00 | 0.46 ± 0.20 | 0.23 ± 0.20 | 0.35 ± 0.00 |
| TRIM24      | 0.23 ± 0.20 | 0.23 ± 0.20 | 0.23 ± 0.20 | 0.00 ± 0.00 |
| WDR9        | 0.23 ± 0.20 | 0.12 ± 0.00 | 0.12 ± 0.00 | 0.12 ± 0.00 |

CBP/EP300 bromodomain inhibitors show similar polypharmacology, we profiled the in vitro activity of CPI644 against a broad panel of serine/threonine and tyrosine protein kinases. At a concentration of 1 μM (well above the IC50 for bromodomain binding), CPI644 showed negligible activity against all of the kinases profiled (supplemental Table S1), suggesting that the phenotypic effects of CBP/EP300 bromodomain inhibitors (see below) are not the result of direct kinase inhibition.

To characterize CBP-bromodomain inhibition in cells, we utilized several orthogonal assays. First, we utilized NanoBRET technology (Promega) to directly monitor the interaction between the isolated CBP bromodomain and histone H3.3. CPI703 and CPI644 inhibited CBP bromodomain binding in a dose-dependent manner with cellular EC50 values of 2.1 and 0.53 μM, respectively (Fig. 2D and Fig. 3A). Inhibition of EP300 bromodomain in cells occurred at similar concentrations (Fig. 3A). Next, we established a complementary assay to monitor the interaction between chromatin and a fluorescent fusion protein containing the CBP bromodomain. This chimeric protein was expressed as a fusion of the CBP bromodomain with a ZsG fluorescent tag (in the context of BRD9 protein i.e. the bromodomain of BRD9 was replaced with that of CBP) (27). Chromatin release was measured by conversion of the largely homogeneous nuclear localization of the ZsG-CBP protein to a more pronounced dot-like pattern (Figs. 2E and 3B). Dot formation occurred in a compound-specific and dose-dependent manner, resulting from displacement of the fluorescently tagged protein from chromatin and its subsequent aggregation (Figs. 2E and 3B). The number and intensity of these foci were measured by high content imaging, and analysis afforded a quantitative determination of target engagement by inhibitor (see “Experimental Procedures”). The number of dots increased upon titration of the inhibitor, yielding a sigmoidal curve when plotted as a function of inhibitor concentration. Cellular EC50 values calculated using this assay (Fig. 2E; IC50 = 2.2 μM; CPI644 = 0.33 μM) correlated extremely well with the potency measured by NanoBRET.

CBP and EP300 modulate gene expression in part through the acetylation of H3K18 and H3K27 (28), and the bromodomain has been linked to its transcriptional activity (29). We therefore sought to determine whether pharmacologic inhibi-
Treg Modulation by CBP/EP300 Inhibition

FIGURE 3. Cellular characterization of CBP/EP300 probe molecules. A, cellular potency of CPI644 using NanoBRET. A dose-response of CPI644 using the isolated bromodomain of CBP (top panel) or EP300 (bottom panel) and histone H3.3 with calculated EC_{50} values of 0.53 and 0.36 µM are shown. B, cellular potency of CPI644 determined by the CBP-bromo or BRD4 (full-length protein) dot assays. Representative images show ZsG-CBP and ZsG-BRD4 protein aggregates from 5.0 µM (CPI644 and CPI644−) and 0.3 µM (CPI203) treatment (top panel). The dose-dependent increase in the number and intensity of these nuclear foci was used to determine the EC_{50} of CPI644 in the CBP (0.33 µM) or BRD4 (12.8 µM) dot assays (bottom panel). C, GFP (control) or Cre (knock-out) cells (25) were utilized to probe H3K18ac by Mesoscale (left panel). The relative change in H3K18ac signal was plotted setting the GFP control cells to 100%. The right panel shows a Western blot using the indicated antibodies in the GFP (control) or Cre (knock-out) cells (25). D, cellular potency of CPI703 or CPI644 using BRD4 and H3.3 NanoBRET. No appreciable inhibition was observed.

Tregs, as demonstrated by GFP expression (supplemental Fig. S1A). All three independent hairpins (for either gene) brought about at least a 50% reduction in the level of the corresponding endogenous protein (supplemental Fig. S1B), and consequently, completely suppressed Treg differentiation (Fig. 4D). To determine whether CBP/EP300 bromodomains could regulate FOXP3 expression in established Tregs, we added CPI703 after naïve human T cells were differentiated into Tregs for 4 days in vitro. This regimen also resulted in a dose-dependent inhibition in the percentage of FOXP3^{+} cells (Fig. 4E), suggesting an important contribution of CBP/EP300 bromodomains in maintenance of the Treg lineage. Indeed, in vitro differentiated Tregs exhibited reduced suppressive capability when co-cultured with carboxyfluorescein succinimidyl ester-labeled naïve CD4^{+} T cells in the presence of CPI703, as evidenced by a ~3-fold increase in naïve T cell proliferation in co-cultures relative to the inactive compound (Fig. 4F, upper panels); this points toward down-modulation of Treg function as a result of CBP/EP300 bromodomain inhibition. Proliferation of naïve T cells alone was minimally affected in the presence of the inhibitor (Fig. 4F, lower panels).

CBP/EP300 Bromodomain Inhibition Alters the Human Treg Transcriptional Signature—To delineate molecular mechanisms underlying the phenotypic and functional impact of CBP/EP300 bromodomain inhibition, we performed full-genome transcriptional profiling, comparing samples from Treg cultures treated either with 4 µM CPI703 or DMSO (4 µM was chosen to ensure robust target coverage, ~1C_{75}). As an additional control we cultured naïve T cells in the absence of differentiating cytokines (hereafter referred to as T_{0}). Double-pairwise comparisons ([CPI703-treated Treg versus DMSO-treated Treg] versus [DMSO-treated Treg versus DMSO-treated T_{0}] (Figs. 5, A and B) revealed a bias toward preferential inhibition by CPI703 of genes that are up-regulated in Tregs compared with those up-regulated in T_{0} cells. Scatter plot analysis of Treg cells treated with CPI703 compared with DMSO controls (Fig. 5A) showed more genes down-regulated than up-regulated in compound-treated Tregs, suggesting that bromodomain function is necessary for full CBP/EP300-mediated transcriptional activation. FOXP3 was among the genes significantly down-regulated, as expected (Fig. 5, A and C), serving as an internal control to validate the analysis. Of the genes up-regulated in Tregs (at least 1.5-fold over T_{0}), 69 genes were inhibited 1.5-fold or more at the transcript level upon CPI703 treatment (Fig. 5C and supplemental Table S3). Another important co-inhibitory receptor, TIM-3 (encoded by HAVCR2), was significantly reduced in Tregs upon compound treatment (~70% down; p < 0.005), although not strongly up-regulated in Tregs compared with T_{0} cells (data not shown). GSEA identified the Treg versus Tconv curated gene set (GSE15659_TREG_VS_TCONV_DN) as significantly coincident with the signature elicited by CPI703 treatment (Fig. 5D), confirming the impact of CPI703 on Treg transcription. These transcriptional data for key mediators of Treg function were also recapitulated...
by flow cytometric analysis. Indeed, levels of LAG-3, CTLA-4, and, to a lesser extent, PD-1, were reduced in a dose-dependent manner by CPI703 and CPI644 (Fig. 5, E and F), with no significant impact on cell viability. Dose-dependent reduction in transcription of FOXP3, LAG3, CTLA4, and PDCD1 (PD-1) was also corroborated by quantitative RT-PCR (data not shown). Thus, we conclude that CBP/EP300 bromodomain inhibition results in the suppression of a network of genes that transcriptionally define Treg cells and their biological functions.

CPI703 Modulates Acetylation in Human Tregs—Mechanistically, CBP/EP300 bromodomain inhibition may facilitate reprogramming of Treg cells by altering the histone acetyl landscape. To address this, we combined global RNA-seq with chromatin immunoprecipitation followed by ChIP-seq to investigate molecular changes in chromatin induced by CBP/EP300 bromodomain inhibition and their association with transcriptional changes. In keeping with the role of CBP/EP300 as transcriptional activators, we found a strong positive correlation between the activation mark H3K4me3 and CBP/EP300-mediated H3K18ac (Fig. 6A) and H3K27ac (not shown) across the genome of human Treg cells. RNA-seq analysis showed that CPI703-mediated CBP/EP300 bromodomain inhibition resulted in the down-regulation of 482 genes by 1.5-fold or more (Fig. 6B). Importantly, CBP/EP300 bromodomain inhibition resulted in diminished H3K18 and H3K27 acetylation (Fig. 6C), similar to previous findings in a different cellular context (Fig. 2F). Consistent with these effects of CPI703 on transcription and CBP/EP300-mediated histone acetylation, we observed a significant concomitant reduction in global H3K4me3 (Fig. 6C). Correlation analysis indicated a strong statistical association between reduced H3K18ac (and H3K27ac) with genes that were down-regulated in their expression by 1.5-fold or more (Fig. 6C). From these data we conclude that optimal deposition of acetyl marks on chromatin by CBP/EP300 necessitates full activity of their bromodomains. H3K18ac, H3K27ac, and H3K4me3 marks were localized primarily at the transcription start sites (TSS; Fig. 6D), in agreement with their role in transcriptional activation. Individual analysis of target loci confirmed this observation, with H3K4me3 being almost exclusively restricted to the TSS, whereas H3K18ac and H3K27ac were highly enriched at the TSS with modest extension toward the gene bodies (Fig. 6, D and E, and supplemental Fig. S2). Furthermore, integrated ChIP-seq signal analysis using the SICER interval analysis algorithm demonstrated that CPI703
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[Graphs and figures are not transcribed, but the text describes the findings and methods related to Treg modulation by CBP/EP300 inhibition.]

Treatment reduces H3K18ac, and to a lesser extent H3K27ac, and that this reduction takes place almost exclusively at the TSS regions (Fig. 6E and supplemental Fig. S2), with a minimal impact on histones located 3′ of the TSS. Reduction of H3K18ac is evident around the TSS of exemplar loci such as LAG3, FOXP3, and RASGRP4 (Fig. 6E) (see also HAVCR2 and IRF7; supplemental Fig. S2), matching a corresponding decrease in H3K4me3 and reduction in exon RNA accumulation. We did not, however, observe a global expulsion of EP300 from chromatin as a consequence of bromodomain inhibition, either in Tregs or in other cellular contexts (data not shown).

These results are consistent with recent publications exploring the function of CBP/EP300 bromodomain inhibition in cancer cell line contexts (32, 33). The most prevalent functional effect of CBP/EP300 bromodomain inhibition was reduced acetylation of H3K18 and H3K27 (and other proteins; see below). Together, our ChIP-seq and RNA-seq studies indicate that CPI703 inhibits optimal acetylation of histones at the transcriptional start sites of target loci, suggesting a novel mechanistic basis for the role of CBP/EP300 bromodomains in human Treg cells.

Additionally, stabilization of FOXP3 protein by EP300-mediated acetylation has been previously reported (34). Therefore we asked whether CBP/EP300 bromodomains might also play a significant role in this process, similar to their effects on histone acetylation. We immunoprecipitated FOXP3 early during Treg differentiation (36 h), followed by immunoblotting with a pan-acetyl antibody. Because FOXP3 is known to be degraded via the proteasome pathway (34), we used the proteasomal inhibitor, MG132, to minimize changes in acetylated FOXP3 levels caused by mere changes in total FOXP3 protein. CPI703 treatment led to a clear reduction in the ratio of acetylated FOXP3 to total FOXP3 (Fig. 7A), indicating a role for CBP/EP300 bromodomains in modulating FOXP3 acetylation. Importantly, CPI703 does not directly inhibit EP300 acetyltransferase activity (Fig. 7B). CPI703 treatment over a longer time period (4 days) led to a marked reduction in the ratio of acetylated FOXP3 to total FOXP3 protein. CPI703 treatment led to a clear reduction in FOXP3 protein levels, as observed previously by FACS (Figs. 4 and 7C), potentially reflecting an increased susceptibility to proteasomal degradation caused by reduction in protein acetylation. Thus, in addition to a transcriptional effect, these data suggest another level of FOXP3 regulation by CBP/EP300 bromodomains via direct FOXP3 acetylation. Taken together, our results highlight at least two separate, but related, mechanisms underlying the novel role of CBP/EP300 bromodomains in human Treg differentiation and function.
Modulation of T Helper Subsets by CBP/EP300 Bromodomain Inhibition—Given the importance of CBP/EP300 bromodomain function within the Treg lineage, we sought to ascertain the specificity of our observed effects vis-à-vis the effect of CPI703 on differentiation of naïve T cells into other TH cell lineages, namely TH1, TH2, and TH17. TH1 and TH2 cell differentiation were largely unperturbed by CPI703, because the percentage of cells producing their canonical cytokines, IFNγ and IL-4, respectively, were unchanged (except for a modest reduction of IFNγ+ T H1 cells at 15 and 20 μM concentrations; Fig. 8A, top panels). However, CBP/EP300 bromodomain inhibition significantly affected T H17 differentiation (Fig. 8, B and C). Both transcript and protein for the characteristic T H17 cytokines, IL-17A, IL-17F, and IL-21, were suppressed by CPI703 with no effect on TNFα production or cell viability (Fig. 8, B and C).

Discussion

Chemical probes allow for temporal, titratable, and domain-specific manipulation of molecular targets in human cells. In the current report, we introduce a series of potent and selective CBP/EP300 bromodomain inhibitors with low micromolar cellular activity using numerous cell systems and contexts. Using these inhibitors, we discover novel effects of CBP/EP300 bromodomain modulation on the biology of human Treg cells. Recent reports have shown that mice with EP300 or CBP-deficient Tregs exhibit enhanced effector responses, reduced peripheral Treg generation, and Treg apoptosis, resulting in rapid allograft rejection and decreased tumor growth. CBP/EP300 HAT inhibition phenocopies the consequences of genetic ablation of either protein in mice (21, 22). Here we show that lentivirus-mediated knockdown of CBP or EP300 in human Tregs severely and non-redundantly inhibited FOXP3 up-regulation with a decrease in cell viability, likely because of a loss of global CBP/EP300-dependent acetylation. In contrast, specific inhibition of CBP/EP300 bromodomains by treatment of differentiating Tregs with CPI703 or CPI644, led to a marked reduction in FOXP3 expression both at protein and transcript levels, with no appreciable effect on cell viability. CBP/EP300 are known to interact with over 300 protein partners and are among the best connected protein “hubs” in the intracellular transcriptional network (19, 35). Given their ubiquitous distribution and the importance of their...
HAT domains in maintaining basal acetylation levels, we propose that bromodomain inhibition represents a more nuanced and less cytotoxic means for querying the importance of context-specific CBP/EP300 scaffolding function in primary cells and a viable option for therapeutic intervention.

Our results point to the requirement for CBP/EP300 bromodomains in the differentiation of human Tregs from naïve T cells. Transcriptional profiling of inhibitor-treated Tregs reveals significant down-regulation of genes encoding crucial Treg factors (FOXP3), and receptors such as LAG-3, CTLA-4, and TIM-3, which participate in cognate Treg responder cell interactions (36–38). CPI703 treatment also reduced transcript levels of other co-inhibitory receptors such as B- and T-lymphocyte attenuator (BTLA) and CD160 (39, 40), as well as the chemokine receptor CCR4, known to be expressed on the highly suppressive effector Treg subset which comprises the bulk of tumor-infiltrating lymphocytes in cancer patients (4). In accordance with this effect on the Treg transcriptome, CPI703 dampened Treg-mediated suppression of naïve T cell proliferation in Treg + responder T cell co-culture assays, underscoring important functional effects of the altered transcriptional signature elicited by CBP/EP300 bromodomain inhibition. As chromatin regulators, the important role of bromodomain function in the optimal acetylation of histones at target loci was perhaps not unexpected. The robust association between reduced acetylation, namely H3K18ac, and gene transcription upon treatment with CPI703 suggests a mechanistic framework to understand how these chromatin reader domains contribute to gene expression. Furthermore, molecular discrimination between H3K18 and H3K27 acetylation might provide additional layers of nuanced transcriptional control (at least in human Tregs), because we observed a generally stronger effect of CPI703 on H3K18ac than on H3K27ac. It is worth noting that we did not observe complete elimination of H3K18ac, which could be the consequence of partial inhibition of CBP/EP300 HAT activity, or a limitation of the reagents used, as double knock-out of CBP/EP300 in mouse embryonic fibroblasts appeared to reduce H3K18ac by \( \sim 60\% \).

Additionally, CBP/EP300 bromodomains could impact optimal acetylation of chromatin associated transcription factors, including FOXP3. EP300-mediated acetylation of FOXP3 has been shown to contribute to its stability (34), and it is conceivable that bromodomain-dependent binding to histones maximizes the probability of CBP/EP300-mediated FOXP3 acetylation in cells. Indeed our data on the regulation of FOXP3 acetylation by CPI703 lends support to this hypothesis. In sum, we propose that the precise positioning of CBP/EP300 at TSS-bound transcriptional complexes is critically aided by their bromodomains, allowing optimal
H3K18 acetylation (and potentially of transcription factors, such as FOXP3), to promote gene activation at these sites. In agreement with a recent report (41), we found a significant disruption of TH17 cytokine production by CBP/EP300 bromodomain inhibition. In addition to the well understood role for TH17 cells and associated cytokines in autoimmune and inflammatory processes, their role in tumor immunology has garnered increased interest, especially their pro-tumorigenic effects (42, 43). In that context, there could be a potential synergistic effect of CBP/EP300 bromodomain inhibition within the tumor microenvironment, by a “double hit” to Tregs and TH17 cells, thereby reversing both suppressive and pro-tumorigenic pathways. The lack of effect on IFNγ/TH1 cells suggests that at least this anti-tumor effector subset might be refractory to CBP/EP300 inhibition, thus leaving a major cytotoxic pathway intact for tumor attack.

Collectively, our results suggest that CBP/EP300 bromodomains are capable of fine-tuning transcriptional programs across different biological pathways that converge on specific T cell states. Potent, selective, and cell-active chemical probes that disrupt CBP/EP300 bromodomain interactions with chromatin therefore constitute crucial starting points for the development of pharmacologic agents. By impairing Treg differentiation and function, such agents may reverse dampened effector responses in the context of cancer and repotentiate anti-tumor immunity, thus complementing more established cancer immunotherapies, particularly those aimed at enhancing tumor eradication by reversal of cytotoxic CD8+ T cell exhaustion through antibody-mediated checkpoint inhibition (44–46).

**Author Contributions**—S. G., M. C., A. S., P. J. D., D. M., G. H., D. D.-N., E. C., and T. R. designed, executed, and analyzed biological experiments. A. T., A. C., F. A. R., V. T., and T. C. designed and supervised small molecule inhibitors. A. T., F. A. R., J.-C. H., J. E. A., A. G. C., S. R. M., S. F. B., B. K. A, and R. J. S. designed and supervised drug discovery studies. H.-R. H., P. S., J. A. M., and A. R. C. designed and executed target engagement cell-based assays. H. J., F. P., and J. W. S. designed, executed, and analyzed crystallography experiments. E. P., H.-R. H., L. E., V. G. W., R. J. S., and R. T. C. designed, executed, and analyzed biochemistry experiments. R. J. S. and R. T. C. supervised biochemistry studies. C. H. and B. M. B. conducted and analyzed bioinformatics analysis of gene transcription and ChIP-seq studies. S. G., J. L. G., and J. M. L. supervised the immunology section of the work. R. J. S. conceived the drug discovery section and wrote the chemical probe section of the manuscript. J. M. L. conceived the biology section, supervised the work, and wrote the immunology section of the manuscript. All authors discussed the results and commented on the manuscript at all stages.

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