DNA Methylation Patterns in CD4⁺ T Cells of Naïve and Influenza A Virus-Infected Mice Developmentally Exposed to an Aryl Hydrocarbon Receptor Ligand

Catherine G. Burke,¹,³ Jason R. Myers,² Christina M. Post,³ Lisbeth A. Boulé,¹,³ and B. Paige Lawrence¹,²,³

¹Department of Microbiology and Immunology, University of Rochester School of Medicine and Dentistry, Rochester, New York, USA
²Genomics Research Center, University of Rochester School of Medicine and Dentistry, Rochester, New York, USA
³Department of Environmental Medicine, University of Rochester School of Medicine and Dentistry, Rochester, New York, USA

BACKGROUND: Early life environmental exposures can have lasting effects on the function of the immune system and contribute to disease later in life. Epidemiological studies have linked early life exposure to xenobiotics that bind the aryl hydrocarbon receptor (AhR) with dysregulated immune responses later in life. Among the immune cells influenced by developmental activation of the AhR are CD4⁺ T cells. Yet, the underlying affected cellular pathways via which activating the AhR early in life causes the responses of CD4⁺ T cells to remain affected into adulthood remain unclear.

OBJECTIVE: Our goal was to identify cellular mechanisms that drive impaired CD4⁺ T-cell responses later in life following maternal exposure to an exogenous AhR ligand.

METHODS: C57BL/6 mice were vertically exposed to the prototype AhR ligand, 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD), throughout gestation and early postnatal life. The transcriptome and DNA methylation patterns were evaluated in CD4⁺ T cells isolated from naïve and influenza A virus (IAV)-infected adult mice that were developmentally exposed to TCDD or vehicle control. We then assessed the influence of DNA methylation-altering drug therapies on the response of CD4⁺ T cells from developmentally exposed mice to infection.

RESULTS: Gene and protein expression showed that developmental AhR activation reduced CD4⁺ T-cell expansion and effector functions during IAV infection later in life. Furthermore, whole-genome bisulfite sequencing analyses revealed that developmental AhR activation durably programmed DNA methylation patterns across the CD4⁺ T-cell genome. Treatment of developmentally exposed offspring with DNA methylation-altering drugs alleviated some, but not all, of the impaired CD4⁺ T-cell responses.

DISCUSSION: Taken together, these results indicate that skewed DNA methylation is one of the mechanisms by which early life exposures can durably change the function of T cells in mice. Furthermore, treatment with DNA methylation-altering drugs after the exposure restored some aspects of CD4⁺ T-cell functional responsiveness. https://doi.org/10.1289/EHP7699

Introduction

An individual’s long-term health trajectory is shaped by the development of complex tissues and organ systems during early life. Development begins in the womb and, for some organ systems, such as the immune system, continues for a period of time after birth (Robin et al. 2003; Zhu and Emerson 2002). Appropriate development relies on integrated signaling networks that regulate cell fate, differentiation, and functional properties (Basson 2012; Partridge et al. 2010; Sanz-Ezquerro et al. 2017). Because this early life window is so critical, it is vulnerable to environmental influences that can durably change health and disease across the life span (Renz et al. 2017). In fact, early life exposures to environmental insults have been associated with increased incidences of obesity, cardiovascular disease, and cancers later in life (Boekelheide et al. 2012; Cao et al. 2016), and with altered function of the immune system (Boule and Lawrence 2015). This idea, which is generally referred to as the developmental origins of health and disease, has been recognized for nearly 40 y; yet we still have much to learn about the mechanisms driving these long-term changes (Boekelheide et al. 2012; de Boo and Harding 2006; Hoffman et al. 2017).

One of the barriers to understanding how early life exposures cause long-lasting consequences to health is that the cellular and molecular mechanisms underlying these durable changes remain poorly characterized. Several epidemiological studies have reported an association between early life exposures to pollutants that bind a ligand-activated transcription factor known as the aryl hydrocarbon receptor (AhR) and impaired immune responses later in life. Associations include increased incidence of respiratory tract and ear infections, increased wheezing, and decreased responses to vaccinations (Dallaire et al. 2004; Glynn et al. 2008; Heilmann et al. 2010; Hochstenbach et al. 2012; Jusko et al. 2014, 2016; Stølevik et al. 2011, 2013). An important immune cell type that is involved in all of these responses is the CD4⁺ T cell. CD4⁺ T cells help fight infections, bolster immunological memory, and aid in responses to vaccination (Crotty 2014; MacLeod et al. 2009; Sant and McMichael 2012; Swain et al. 2012; Zhu and Paul 2008). Given this central role of CD4⁺ T cells, and that all immune cells, including CD4⁺ T cells, express the AhR (Esser and Rannug 2015), we investigated how early life AhR activation modulates CD4⁺ T-cell responses later in life.

Maternal exposure of mice to the prototype AhR ligand, 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD), causes impaired CD4⁺ T-cell responses in adult offspring (Boule et al. 2014, 2015a, 2015b; Burke et al. 2019). With specific regard to CD4⁺ T cells, developmental AhR activation reduces conventional helper T-cell responses and increases regulatory T cell responses during influenza A virus (IAV) infection at adulthood (Boule et al. 2014). Adoptive transfer studies revealed that reduced CD4⁺ T-cell responses were due to factors intrinsic to CD4⁺ T cells (Boule et al. 2014). Furthermore, these changes in CD4⁺ T-cell responses in developmentally exposed offspring were durable and observed up to 1 y after birth (Boule et al. 2015a). Given that the half-life of TCDD in mice is 7–10 d (Gasiewicz et al. 1983), differences in T-cell responses were maintained long after the exogenous AhR stimulus was cleared from the system and when AhR target genes were no longer induced (Winans et al. 2015). The fact that alterations persist well after exposure suggests that...
the long-term responsive capacity of CD4+ T cells is programmed during development and that this programming is sensitive to environmental cues. One way that exposures can cause durable modifications is by altering the epigenetic landscape. To date, whether early life activation of the AhR by exposure to an exogenous ligand changes epigenetic modifications within CD4+ T cells later in life remains unknown.

One type of epigenetic mechanism that influences cell differentiation and function is DNA methylation (Suárez-Álvarez et al. 2013; Tirado-Magallanes et al. 2017). There are several reports that early life activation of the AhR changes DNA methylation patterns in nonimmune organs such as the testes (Ding et al. 2018; Manikkam et al. 2012), ovaries (Zhang et al. 2019), placenta (Ding et al. 2018), mammary tissues (Papoutsis et al. 2015), muscle (Sommer et al. 2013), and liver (Sommer et al. 2013). Taken together, this information led us to hypothesize that one of the mechanisms by which early life AhR activation impairs CD4+ T-cell responses later in life is via remodeling of DNA methylation patterns in CD4+ T cells. To test this, mice were developmentally exposed to a vehicle control or to TCDD, which specifically activates the AhR (Boule et al. 2014; Winans et al. 2015). Unbiased whole-genome approaches were used to define how developmental AhR activation impacts DNA methylation and gene expression patterns in CD4+ T cells isolated from adult animals. We also examined the causal relationship between changes in the level of DNA methylation and altered CD4+ T-cell responses using pharmacological interventions that elevate or dampen DNA methylation in offspring.

Methods

Animals and Developmental Exposure

C57BL/6 (Ahp/h) mice (5–6 wk of age) were purchased from the Jackson Laboratory. For each experiment, between 20 and 40 nulliparous females were housed with C57BL/6 males. At the initiation of experimental breeding, there were 2 or 3 females with each male. As vaginal plugs were identified, females were removed and singly housed thereafter. In the morning, females were checked daily for the presence of a vaginal plug, which was labeled as gestational day (GD) 0. Anisole was used to dissolve the TCDD (99% purity; Cambridge Isotope Laboratories), which was then diluted in peanut oil. The vehicle control was peanut oil containing 0.01% anisole. Impregnated female mice were alternately assigned to treatment groups and treated with the peanut oil vehicle control (vehicle) or 1 μg TCDD/kg body weight by gavage between 1100 hours and 1300 hours on GD0, -7, and -14 and 2 d after parturition (PND2). Cages were clearly labeled to denote dam treatment and to ensure accurate dosing across pregnancy and postpartum. All members of the research team were aware of which animals were in which treatment group. Offspring were weaned at PND21, and no culling of litters was performed. Littermates were housed in same-sex groups. Offspring were assigned a number, and at the time the mice were sacrificed, only the person in charge of the experiment knew which number correlated with a specific mouse. Mice were housed at the University of Rochester in microisolator cages in a specific-pathogen free facility that is maintained in a 12-h light: dark cycle, at 22°C±2°C and 30–50% relative humidity. Mice were provided food (LabDiet 5010) and reverse osmosis purified water ad libitum. All experiments were performed using female offspring. All animal treatments had prior approval of the Institutional Animal Care and Use Committee of the University of Rochester, and followed all guidelines and regulations. The University has accreditation through the Association for Assessment and Accreditation of Laboratory Animal Care. Animals were treated humanely and with due consideration to alleviation of any distress and discomfort. All U.S. Public Health Service Policy on Human Care and Use of Laboratory Animals guidelines for the handling of vertebrate animals were followed (https://olaw.nih.gov/policies-laws/phs-policy.htm).

S-Adenosyl Methionine and Zebularine Treatment

For some studies, at weaning, developmentally exposed offspring were randomly placed into three treatment groups. One group of mice (control) remained on drinking water provided by the vivarium. The second group (SAM) received drinking water containing 0.5 mg/mL S-adenosyl methionine (SAM; Sigma-Aldrich). The third group (Zeb) received drinking water containing 0.2 mg/mL Zebularine [Zeb; NCS309132, provided by the Developmental Therapeutics Program, National Cancer Institute, National Institutes of Health (NIH)]. These in vivo doses of SAM (Y Li et al. 2012) and Zeb (Yoo et al. 2008), administered via drinking water, have been previously shown to alter DNA methylation levels in mice. Mice were provided water ad libitum. To conserve drugs, mice within the same developmental exposure group were co-housed at up to five mice per cage. Water consumption was measured every other day, and body weight was recorded once a week (Figure S1). Mice were infected with IAV at 6–8 wk of age, and body weight was monitored daily throughout infection (Figure S1). All mice were maintained on treated water for the remainder of the experiment.

Influenza Virus Infection

IAV (strains HKx31, H3N2 or PR8, and H1N1) were initially obtained from M. Coppola (Argonex, Charlottesville, VA). Virus stocks were separately propagated in embryonated specific-pathogen free chicken eggs (Charles River), and clear allantoic fluid was aliquoted and snap frozen in liquid nitrogen, as described previously (Warren et al. 2000). Virus titers were determined by hemagglutination of erythrocytes (Barrett and Inglis 1985). Briefly, serial dilutions of allantoic fluid (1:2 to 1:8,192) were prepared in phosphate-buffered saline (PBS), and 100 μL was added to U-bottom 96-well microtiter plates. To each well, 100 μL of 1% (vol/vol) chicken erythrocytes (Cat. No. 31152; Colorado Serum Co.), diluted in PBS, was added, and the plate was incubated for 45 min at room temperature. PBS alone was included as a negative control, and undiluted IAV from a previously titered frozen stock was included as a positive control. To infect the mice, adult offspring (6–10 wk of age) were anesthetized with 0.225–0.250 mL of avertin (2% 2,2,2-tribromoethanol; Sigma-Aldrich), administered by intraperitoneal injection. Infections were performed in the morning, between 0700 hours and 0900 hours. For primary infection, mice were infected intranasally with 120 hemagglutination units (HAU) of HKx31, which is a sublethal inoculum in immunocompetent mice (Warren et al. 2000; Boule et al. 2014, 2015b; Winans et al. 2015). The primary response was examined 9 d later, which is the peak of the T-cell response to acute primary IAV infection (Boule et al. 2014; Brown et al. 2006). To examine immunological memory, it is important to allow time for full resolution of the primary response and establishment of anamnestic cells. Mice were challenged with a heterosubtypic infection (0.1 HAU PR8) on Day 79 (i.e., 70 d after Day 9 post primary infection with 3 HAU HKx31). This dose of PR8 causes significant mortality during primary infection (Giannandrea et al. 2012). In all experiments, infection and tissue collection were initiated in the morning. All work with infectious agents was conducted with prior approval of the Institutional Biosafety Committee of the University of Rochester.

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following guidelines of the NIH and the Centers for Disease Control and Prevention (https://www.cdc.gov/labs/BMBL.html).

**Tissue Collection**
At the termination of each experiment, mice were sacrificed using a lethal dose of anesthetic (Avertin >0.5 mL) and either cervical dislocation or exanguination. Tissue collection was initiated in the morning between 0730 hours and 0900 hours. The mediastinal lymph nodes (MLNs) were harvested, and single-cell suspensions prepared by pressing MLNs from a single mouse between the frosted ends of two microscope slides. Cells were then resuspended in cold Hanks’ balanced salt solution (HBSS) containing 2.5% fetal bovine serum (FBS) (Hyclone). Lysis solution (0.15 M ammonium chloride, 10 mM sodium hydrogen carbonate, 1 mM ethylenediaminetetraacetic acid) was added to the cells for 5 min at room temperature to lyse red blood cells. Lysis was quenched using cold HBSS/FBS. Debris was removed by passing cells through a 70-μm filter. Cells were resuspended in cold HBSS/FBS for enumeration using a TC20 cell counter (BioRad). Cells were immediately used for analysis by flow cytometry or subjected to further treatment to enrich and purify a specific cell type and extract cellular materials. When peripheral lymph nodes were also collected, they were processed in the same manner.

**Flow Cytometry**
Prior to staining with fluorochrome-conjugated antibodies, single-cell suspensions containing 2×10^6 lymph node cells were incubated for 10 min at 4°C with anti-mouse CD16/32 monoclonal antibody (mAb; clone 93) to block nonspecific staining. Cells were then incubated for 20 min at 4°C with previously determined optimal concentrations of fluorochrome-conjugated antibodies against cell-surface antigens (Table 1). To identify CD4+ T cells that recognize IAV, prior to extracellular antibody staining, cells were incubated for 1 h at 37°C with high histocompatibility class II tetramers containing an immunodominant peptide epitope of Hx31 [nucleoprotein (NP), I-A^B/NP311-325; provided by the National Institute of Allergy and Infectious Diseases, NIH Tetramer Core Facility; http://tetramer.yerkes.emory.edu/]. To detect effector proteins, cells were incubated in the presence of recombinant mouse interleukin-2 (IL-2; R&D Systems) and brefeldin A (Sigma-Aldrich) for 1 h at 37°C prior to CD16/32 mAb block or extracellular antibody staining. Anti-CD107a [LAMP-1 (lysosomal-associated membrane protein 1)] was added, and the cells were incubated for an additional 4 h at 37°C prior to extracellular staining (Post et al. 2019). The cells were then fixed with a solution containing 2% formaldehyde in PBS for 20 min and resuspended in staining buffer containing 1% saponin before being incubated at room temperature with an antibody against perforin (Table 1). To detect Ki67, cyclin-dependent kinase inhibitor 2A (p16^INK4A; p16), and T-box transcription factor (Tbet), cells were fixed/permeabilized using either the FoxP3 Staining Kit (eBioscience) or the BD Transcription Factor Buffer Set (BD Biosciences) following extracellular staining per the manufacturer’s instructions. Cells were then stained with intracellular antibodies to Ki67, p16, or Tbet for 20 min at 4°C. Prior to use, the antibody against p16 was directly conjugated to CF680 using the Mix-n-Stain Antibody Labeling Kit, following the manufacturer’s instructions (Sigma-Aldrich). Table 1 lists all antibodies used. Fluorescence minus one controls were used to determine nonspecific fluorescence staining and to define gating parameters. Data were obtained using an LSRII flow cytometer (BD Biosciences) to collect approximately 500,000 events per sample. Data were analyzed using the FlowJo software program (version 10; Tree Star).

**CD4+ T Cell Purification**
Pregnant mice were administered TCDD (1 μg/kg body weight) or peanut oil vehicle control on GD0, -7, and -14 and PND2. CD4+ T cells were isolated from lymph nodes of adult female offspring (8–10 wk of age). Peripheral lymph nodes (cervical, axillary, brachial, inguinal, and sacral) were obtained from immunologically naïve mice. MLNs were harvested from mice 9 d after infection with IAV (HKx31). For naïve offspring, peripheral lymph nodes from a single mouse were combined, and there were three replicates per treatment group, with each offspring in a replicate from a unique dam. For IAV-infected offspring, MLNs were used. Mice were sacrificed 9 d after infection, and there were three replicates per treatment group, with 4–6 offspring in each unique pool of cells. All of the mice were sacrificed on the same day. Lymph nodes were processed to single-cell suspensions, and CD4+ T cells were isolated using immunomagnetic negative selection to >90% purity (Figure S2) using the MojoSort Mouse CD4+ T cell kit (BioLegend), using reagents provided by the vendor and following the vendor’s instructions (https://www.biolegend.com/protocols/mojosort-isolation-kits-protocol-1/4599/), with the following change: the MagCelllect Magnet (R&D Systems) was used.

**RNA-Sequencing**
RNA was isolated from purified CD4+ T cells using the RNaseasy Mini Kit (Qiagen). Total RNA concentration was determined with a Nanodrop 1000 spectrophotometer (NanoDrop), and RNA quality measured using an Agilent Bioanalyzer 2100 (Agilent). RNA (1 ng) was preamplified with the SMARTer Ultra Low RNA Kit for Illumina Sequencing (Clontech), following the manufacturer’s recommendations. The quantity and quality of cDNA was determined using the Qubit Fluorometer (Life Technologies), and Agilent Bioanalyzer 2100 (Agilent), complementary DNA (cDNA) (150 pg) was used to generate Illumina-compatible sequencing libraries with the NexterXXT library preparation kit (Illumina), following the manufacturer’s protocols. Libraries were hybridized to the Illumina single-end flow cell and amplified using the cBot (Illumina) at a concentration of 10 pM per lane. Single-end reads of 100 nucleotides (nt) were generated for each sample using Illumina’s HiSeq2500 v4. Raw reads from the Illumina HiSeq2500 sequencer were demultiplexed using bc2fastq (version 2.19.0; Illumina). Quality filtering and adapter removal were performed using Trimomatic-0.36 (Bolger et al. 2014) with the following parameters: “TRAILING:13 LEADING:13 ILLUMINACLIP:adapters.fasta:2:30:10 SLIDINGWINDOW:4:20 MINLEN:15.” Processed/cleaned reads were then mapped to the Mus musculus reference sequence (GRCm38.p5 + gencode M12) with STAR-2.5.2b (Dobin et al. 2013) with the following parameters: “--twopassMode Basic –runMode alignReads –genomeDir $[GENOME]–readFilesIn $[SAMPLE] –outsAMType BAM SortedByCoordinate –outSAMstrandField intronMotif –outFilterIntronMotifs RemoveNoncanonical.” The subread-L.0p3 (Liao et al. 2014) package (featureCounts) was used to derive gene counts given the following parameters: “-s 0 -t exon -g gene_name.” Differential expression analysis and data normalization was performed using DESeq2-1.14.1 (Love et al. 2014) with an adjusted p-value threshold of 0.05 within an R (version 3.3.2; R Development Core Team) environment. Genes were considered differentially expressed (DEGs) when they had an adjusted p < 0.05. Pathway analysis was performed on DEGs using Ingenuity Pathways Analysis (Ingenuity Systems; http://www.ingenuity.com).
CD3e PE hamster anti-mouse 145-2C11 BD Biosciences 553064 0.13 μg
CD3e PE-CF594 hamster anti-mouse 145-2C11 BD Biosciences 562286 0.13 μg
CD3e PE-Cy5 hamster anti-mouse 145-2C11 BD Biosciences 552774 0.25 μg
CD4 BV650 rat anti-mouse GK1.5 BD Biosciences 563232 0.06 μg
CD4 FITC rat anti-mouse H129.19 BD Biosciences 553651 0.16 μg
CD4 PerCP-Cy5.5 rat anti-mouse RM4-5 Invitrogen/E Bioscience 45-0042-82 0.06 μg
CD9 APC-R700 rat anti-mouse 1D3 BD Biosciences 565473 0.06 μg
CD9 BB515 rat anti-mouse 1D3 BD Biosciences 564531 0.06 μg
CD19 BV421 rat anti-mouse 1D3 BD Biosciences 562701 0.06 μg
CD19 BV711 rat anti-mouse 1D3 BD Biosciences 563157 0.06 μg
CD19 PE rat anti-mouse 1D3 BD Biosciences 565487 0.06 μg
CD44 BB515 rat anti-mouse IM7 BD Biosciences 563970 0.25 μg
CD44 BV711 rat anti-mouse IM7 BD Biosciences 563971 0.25 μg
CD45R APC rat anti-human, mouse RA3-6B2 Invitrogen/E Bioscience 56-0452-82 0.25 μg
CD45R AF700 rat anti-mouse RA3-6B2 BD Biosciences 557957 0.03 μg
CD62L BV605 rat anti-mouse MEL-14 BD Biosciences 563252 0.06 μg
CD62L PE-Cy7 rat anti-mouse MEL-14 Invitrogen/E Bioscience 25-0621-82 0.06 μg
CD107a (LAMP-1) PE rat anti-mouse 1D4B BD Biosciences 558661 0.80 μg
CD152 (CTLA-4) APC-R700 hamster anti-mouse UC10-4F10-11 BD Biosciences 565778 0.13 μg
CD185 (CXCR5) biotin rat anti-mouse 2G8 BD Biosciences 551961 1.25 μg
CD279 (PD-1) BV421 hamster anti-mouse J43 BD Biosciences 562584 0.50 μg
CD279 (PD-1) FITC hamster anti-mouse J43 Invitrogen/E Bioscience 11-9905-82 0.23 μg
CD366 (TIM3) PE rat anti-mouse 25-0508 R&D Systems FAB1529P 3.75 μL/10^6 cells
KLRG1 PerCP-Cy5.5 hamster anti-mouse 2FI BD Biosciences 563595 0.06 μg
MHCII tetramer [I-A(b)] containing influenza A virus nucleoprotein (NP) peptide 311–325], APC Streptavidin PE NA BD Biosciences 550401 0.15 μg
Tbet PE-Cy7 mouse anti-human, mouse, rhesus monkey 4B10 Invitrogen/E Bioscience 25-5825-82 0.50 μg
Perforin FITC rat anti-mouse eBioGM1D BD Biosciences 11-9392-82 1.00 μg
Sigma Mix Non-Stain CF680 Antibody labeling kit NA Sigma-Aldrich MX805620 NA
CDKN2A (p16INK4a) unconjugated mouse 2D9A12 Abcam ab54210 2 μL
CD19 purified rat anti-mouse 1D3 BD Biosciences 550284 0.25 μL
Fixable Viability Dye eFluor506 NA Invitrogen/E Bioscience 65-0866-14 1:500 dilution
BD Horizon Brilliant Stain Buffer NA BD Biosciences 563794 NA
Transcription Factor Buffer Set NA BD Biosciences 562574 NA

Whole-Genome Bisulphite Sequencing
DNA was isolated from purified CD4^+ T cells using the DNaseq blood and tissue kit (Qiagen), and processed for whole-genome bisulphite sequencing (WGBS). Genomic DNA was quantified using the Qubit fluorometer (Life Technologies). DNA integrity was determined using the TapeStation with genomics DNA tape reagents (Agilent). Methylation libraries were generated with Illumina’s TruSeq DNA Methylion Kit. Briefly, bisulphite conversion was carried out on 100 ng of genomic DNA using Zymo EZ DNA Methylation Gold per the manufacturer’s instructions (Zymo Research). Conversion efficiency was determined using a BioAnalyzer 2100 (Agilent). Single-stranded cDNA was generated and tagged from bisulphite converted DNA using random hexamers containing a 5’ tag followed by terminal tagging to mark the 3’ end of the DNA. Illumina-specific adaptors were added during polymerase chain reaction amplification. DNA libraries were purified with AmpureXP beads and quantified with the BioAnalyzer 2100 and Qubit fluorometer. The amplified libraries were hybridized to the Illumina pair end flow cell and amplified using the cBot (Illumina) at a concentration of 8 pM per lane. Pair end reads of 126 nt were generated on the Illumina HiSeq2500 for each sample. Raw reads generated from the Illumina HiSeq2500 sequencer were demultiplexed using bcftools. Sequence reads were cleaned and adapter trimmed using trim-galore (version 0.4.3; FelixKrueger; https://github.com/FelixKrueger/TrimGalore) with the following parameters: “-paired –clip_r1 8 –clip_r2 8 –three_prime_clip_r1 8 –three_prime_clip_r2 8”. Alignment of each sample individually to the bisulphite converted mouse reference genome (GRCm38.p5) was accomplished with Bismark (version 0.18.1; Krueger and Andrews 2011) and the following parameters: “--bowtie2 --maxins 1000”. The accessory programs “filter_non_conversion” and “deduplicate_bismark” were applied to the resulting BAM files. The methylKit (version 1.4.0; Akalin et al. 2012) R package within an R (version 3.4.1) environment was used to extract methylation information in CpG context without overlap, filter loci with fewer than five reads and quality score lower than 10 out, normalize coverage across samples, and produce quality control plots. The genome was divided into 500-bp windows/tiles to assess differentially methylated regions (DMRs). Regions were united across all of the samples using the “min.per.group = 1L” option (window covered in at least one sample in every group). DMRs were defined as a 500-bp window with a q < 0.05 and a change in DNA methylation of at least 25%. The DMRs were annotated with the mm10 annotations using HOMER (version 4.9.1; Heinz et al. 2010).

Statistical Analysis
For these studies, the dam was the statistical unit. Hence, at each point in time relative to infection and in each developmental

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exposure group, the offspring were from a different treated dam. Except for RNAseq and WGBS data, all data were analyzed using JMP software (SAS Institute Inc.). Differences between means of vehicle or TCDD-treated groups at a single point in time were evaluated using a Student’s t-test. A two-way analysis of variance, followed by a Tukey honestly significant difference post hoc test, was used to compare differences between multiple independent variables. The specific statistical test used is indicated in the legend of each figure. Differences were considered statistically significant when p-values were ≤0.05. Error bars on all graphs and tables represent the standard error of the mean.

Data Availability
All WGBS and RNA-sequencing (RNA-seq) data have been deposited to the NIH Gene Expression Omnibus under accession number GSE143893 (WGBS) and GSE143894 (RNA-seq).

Results

Gene Expression Patterns in CD4+ T Cells of Developmentally Exposed Mice

To characterize how triggering the AhR during development modifies CD4+ T-cell responses later in life, the transcriptomes of CD4+ T cells isolated from immunologically naïve mice and IAV-infected mice were examined using high-throughput RNA-seq (Figure 1A). In the absence of infection, gene expression profiles in CD4+ T cells from mice that were developmentally exposed to vehicle control or TCDD were similar (Figure 1B). In fact, there were no DEGs in CD4+ T cells from immunologically naïve adult offspring from the two treated groups of dams. However, this does not mean there was an absence of gene expression. It indicates that in the absence of infection, there were not any genes whose expression was significantly different due to maternal exposure to the AhR agonist TCDD. In contrast, following IAV infection, developmental exposure resulted in the differential expression of 163 genes (Figure S3). Thus, triggering the AhR during development, by maternal treatment with the ligand TCDD, induced differences that were not revealed until after the offspring’s immune system was challenged to respond.

To further examine how developmental exposure shapes CD4+ T-cell responses to infection, we compared DEGs in cells from naïve and IAV-infected mice within each exposure group. In CD4+ T cells from IAV-infected mice that were not exposed to TCDD during development, there were 2,903 DEGs vs. immunologically naïve offspring of vehicle-treated dams (Figure 1C). In CD4+ T cells from the offspring of TCDD-treated dams, there were 2,395 DEGs in infected vs. naïve mice (Figure 1D). In both developmental exposure groups, roughly 60% of the DEGs represented genes with higher expression (Figure 1C,D). Among the up-regulated genes, 1,070 DEGs were the same, regardless of maternal exposure group, whereas 354 genes with higher expression were unique to offspring of the TCDD exposed dams (Figure 1E). Similarly, more of the down-regulated DEGs were shared between CD4+ T cells from vehicle and TCDD developmentally exposed offspring. Specifically, 587 infection-associated DEGs were common in vehicle and TCDD developmentally exposed offspring. Additionally, 384 genes were uniquely down-regulated as a result of developmental exposure (Figure 1F). Thus, when responding to antigen challenge, developmental exposure to TCDD altered the transcriptional profile of CD4+ T cells, with differences that included enhanced and diminished gene expression.

Additional examination of the pathways influenced by IAV infection indicated several similarities and differences between CD4+ T cells from the offspring of dams that were or were not exposed to an exogenous AhR agonist. For example, in both exposure groups, T-cell activation and T helper differentiation were among the major affected pathways (Figure 1G,H). In contrast, developmental AhR activation resulted in decreased activity of cell cycle-related pathways (e.g., cell cycle G2/M). When ranking the top 20 pathways enriched in infected vs. naïve comparisons, 14 were shared between vehicle and TCDD exposure groups (Figure S4). Therefore, IAV infection affected gene expression patterns in many similar pathways, regardless of developmental exposure. Despite this overlap, affected pathways in T cells from the offspring of control dams were at a lower rank in cells from mice developmentally exposed to TCDD (Figure S4). Thus, even among pathways shared by vehicle and TCDD groups, the manner in which they were affected was not the same.

Influence of Developmental AhR Activation on Senescence and Exhaustion

One of the most strongly enhanced pathways in both groups of developmentally exposed offspring was T-cell exhaustion (Figure 1G,H; Table S1). To determine whether developmental exposure to TCDD enhanced metrics of CD4+ T-cell exhaustion, the expression of cytotoxic T-lymphocyte-associated protein 4 (CTLA4), programmed cell death protein 1 (PD1), and T cell immunoglobulin and mucin domain-3 (TIM3) was measured because increased expression of these molecules is associated with T-cell exhaustion (Akbar and Henson 2011). Flow cytometry analysis revealed that the number of CD4+ T cells that expressed PD1 was significantly lower in mice that were exposed to TCDD during development (Figure 2A). Furthermore, the percentage of PD1+CD4+ T cells was lower following developmental exposure to TCDD (Figure 2B). However, PD1 is also a cell-surface receptor expressed by T follicular helper (Tfh) cells. Developmental AhR activation reduced the percentage and the number of Tfh cells during IAV infection in mice (Boule et al. 2014; Burke et al. 2019). Therefore, the observation that there were fewer PD1+CD4+ T cells may be due to a reduction in the number of Tfh cells, rather than a reduction in T-cell exhaustion. Similar to PD1, the number of CD4+ T cells that expressed the inhibitory receptors CTLA4 (Figure 2C) and TIM3 (Figure 2E) was lower in mice developmentally exposed to TCDD. Yet, unlike PD1, the percentage of CD4+ T cells that expressed CTLA4 (Figure 2D) and TIM3 (Figure 2F) were similar in mice regardless of developmental exposure group.

In addition to T-cell exhaustion, pathways associated with cellular senescence (Franceschini et al. 2000) were influenced by developmental exposure (Figure 1H). These include proliferation (e.g., mitotic roles for polo-like kinase, chromosomal replication, cell cycle: G2/M, and cell cycle control), and effector functions [e.g., leukocyte extravasation, nuclear factor kappa-light-chain-enhancer of activated B cells (NF-kB) activation by viruses, IL-6 signaling, and CD40 signaling], which were dampened or missing from the top pathways in CD4+ T cells from the offspring developmentally exposed to TCDD during IAV infection. Two common markers associated with cellular senescence are killer cell lectin-like receptor subfamily G member 1 (KLRG1) and p16 (Henson and Akbar 2009; Liu et al. 2009; Suen et al. 2016). In mice that had been developmentally exposed to TCDD, there was a lower number (Figure 3A) and percentage (Figure 3B) of CD4+ T cells that expressed KLRG1. Although the number was not different (Figure 3C), developmental AhR activation increased the percentage of cells that express p16 during IAV infection (Figure 3D). Effector functions of CD4+ T cells during IAV infection include cytokine production and degranulation (Brown et al. 2004, 2016). Up-regulation of surface CD107a was
Figure 1. Gene expression profiles in CD4⁺ T cells of naïve or influenza A virus (IAV)-infected mice developmentally exposed to vehicle or TCDD. (A) Pregnant mice were administered vehicle or TCDD (1 µg/kg body weight) on GD0, 7, and 14 and 2 d after parturition. CD4⁺ T cells from lymph nodes of naïve or IAV-infected offspring (8–10 wk of age) were purified, RNA was isolated, and RNA-sequencing (RNA-seq) was performed. For immunologically naïve offspring, peripheral lymph nodes from a single mouse were combined, and there were a total of three replicates per treatment group, with 1 offspring from each unique pool of cells for a total of 30 total mice used. (B) All of the mice (naïve and infected) were sacrificed on the same day with CD4⁺ T-cell purification, RNA isolation, and RNA-seq performed in parallel. The number of differentially expressed genes (DEGs) was assessed. (C,D) Volcano plots of DEGs. On each plot, the direction of change indicates increased (right; red) or decreased (left; green) gene expression with infection compared with CD4⁺ T cells from immunologically naïve (C) vehicle or (D) TCDD developmentally exposed mice. (E) Venn diagram of the number of DEGs that were up-regulated in both vehicle and TCDD groups, or uniquely up-regulated in only one group. (F) Venn diagram of the number of DEGs that were down-regulated in both vehicle and TCDD groups, or uniquely down-regulated in only one group. (G) Venn diagram of the number of DEGs that were down-regulated by both vehicle and TCDD exposure, or uniquely down-regulated in only one group. (H) Ingenuity Pathway Analysis was used to identify cellular pathways affected by DEGs identified in (C) and (D). T-cell relevant pathways were ranked according to z-score, in descending order of highest to lowest z-score. Graphs depict pathways from (G) IAV-infected vs. naïve vehicle offspring or (H) IAV-infected vs. naïve TCDD offspring. The x-axes denote –log (p-value) for each pathway. Pathways were plotted according to z-score, with highest z-scores at the top (red) to lowest at the bottom (green). Gray bars were used when a z-score was not predicted. Numerical details are provided in Table S1. Graphic art in (A) was manually created or adapted from Servier Medical Art templates (https://smart.servier.com) and licensed under the Creative Commons Attribution 3.0 Unported License agreement (https://creativecommons.org/licenses/by/3.0/). Note: ATM, ataxia telangiectasia mutated protein; BW, body weight; GD, gestational day; IL, interleukin; iCOS, inducible T-cell costimulator; iCOSL, inducible T-cell costimulator ligand; iNOS, inducible nitric oxide synthase; NF-jB, nuclear factor kappa-light-chain-enhancer of activated B cells; PND, postnatal day; TCDD, 2,3,7,8-tetrachlorodibenzo-p-dioxin; Th, T helper; TNFR, tumor necrosis factor receptor.
Persistence of Impaired CD4+ T-cell Responses

A robust primary CD4+ T-cell response to IAV infection influences anamnestic responses to subsequent infections (Sun and Braciale 2013; Wilkinson et al. 2012). Given that developmental activation of the AhR-damped CD4+ T-cell responses to primary IAV infection, we hypothesized CD4+ T-cell responses would also be reduced during a subsequent IAV infection. A key metric of CD4+ T-cell responsiveness is the clonal expansion of virus-specific cells (Chapman et al. 2005; Sant and McMichael 2012). Thus, the number of CD4+ T cells that recognize an immunodominant epitope of IAV NP was evaluated in developmentally exposed offspring before infection, during primary IAV infection, and then prior to and after challenge with a second heterosubtypic IAV (Figure 4A). In naïve mice, the number of virus-specific CD4+ T cells was similar in offspring that were developmentally exposed to vehicle or TCDD (Figure 4B). In response to primary infection (HKx31; H3N2), the number of NP-specific CD4+ T cells increased in the offspring of vehicle control-treated dams. However, 9 d after infection, the number of virus-specific CD4+ T cells in infected offspring of TCDD-treated dams was significantly lower than in the vehicle offspring (Figure 4C). After mice recovered from primary IAV infection, the number of virus-specific CD4+ T cells was no longer significantly different between the two groups (Figure 4D). However, when they were inoculated with a heterosubtypic IAV (PR8; H1N1), expansion of virus-specific CD4+ T cells was significantly lower in mice that had been developmentally exposed to TCDD.

Influence of Developmental Exposure and Immune Challenge on DNA Methylation Patterns in CD4+ T Cells

Because changes to CD4+ T-cell responses and gene expression patterns were long lasting, we assessed epigenetics as an underlying mechanism. Specifically, DNA was purified from the same CD4+ T cells that were used for RNA-seq in Figure 1. DNA was subjected to WGBS to evaluate DNA methylation patterns across the genome of these cells. In the absence of infection, CD4+ T cells from vehicle- and TCDD-exposed offspring had similar levels of global DNA methylation (Figure 5A). Compared with CD4+ T cells from naïve animals, CD4+ T cells from animals infected with IAV had significantly lower global DNA methylation levels, and this was more pronounced in the offspring of the dams given TCDD. Specifically, in the offspring of control dams, infection was associated with 1.8% lower overall DNA methylation ($p = 0.0002$; Table S4). In CD4+ T cells from mice that were exposed developmentally to TCDD, the overall DNA methylation level was lower by 2.6% ($p < 0.0001$; Table S4). Thus, CD4+ T cells from infected offspring that were developmentally exposed to TCDD had significantly lower levels of global DNA methylation compared with IAV-infected offspring of vehicle dams ($p = 0.0118$). Thus, developmental exposure impacted DNA methylation levels in CD4+ T cells, and, at a genome-wide level, differences became apparent after an immune challenge.

To better understand the extent to which developmental AhR activation impacted DNA methylation, DNA methylation patterns were characterized within discrete regions across the genome. DMRs were defined as having a significant methylation change of $>25\%$ within 500 bp regions. Triggering the AhR during development caused about 22,000 DMRs in naïve T cells, and 18,752 DMRs in CD4+ T cells isolated from IAV-infected mice (Figure 5B). In naïve animals, the number of DMRs that were hypo- or hypermethylated were roughly similar, which is in accordance with similar global levels of DNA methylation (Figure 5B). Moreover, DMRs spanned all chromosomes prior to and after infection, in both exposure groups (Figure 5C–F). Following infection, a greater proportion (57%) of the DMRs in the TCDD group represented hypomethylated DNA, which is consistent with the overall reduction in global DNA methylation (Figure 5B). However, infection with IAV caused 20% more DMRs (29,563) in CD4+ T cells from vehicle-treated dams. However, these DMRs were not preferentially a hypomethylated feature of DNA, as the number of DMRs was significantly different between the two groups (Figure 5B). Following infection, a greater proportion (57%) of the DMRs in the TCDD group represented hypomethylated DNA, which is consistent with the overall reduction in global DNA methylation (Figure 5B). However, infection with IAV caused 20% more DMRs (29,563) in CD4+ T cells from vehicle-treated dams. However, these DMRs were not preferentially a hypomethylated feature of DNA, as the number of DMRs was significantly different between the two groups (Figure 5B).

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the CD4+ T-cell genome were annotated. This analysis showed that expression and function, only 2% of DMRs were found in promoter regions. These results show that both an immune challenge and developmental exposure influenced DNA methylation patterns across all genomic features in CD4+ T cells.

**Integrating Gene Expression and DNA Methylation**

One of the best-studied ways in which DNA methylation affects cellular function is by regulating gene expression. Therefore, we integrated transcriptomic and genome-wide DNA methylation analyses. Gene expression fold changes did not correlate with DMRs across all genomic regions, or within specific regions associated with each DEG (Figure S5). Others have reported a similar lack of correlation between promoter DNA methylation and gene expression (Akemann et al. 2019). Yet, a subset of the DMRs that we observed had a strong correlation with gene expression changes (blue dots; Figure S5). This was observed regardless of maternal exposure group. For instance, compared

CD4+ T cells but, instead, changed DNA methylation patterns overall such that DMRs throughout the entire genome of CD4+ T cells included regions with greater and lower DNA methylation.

DNA methylation at different genomic features influences cellular functions, although precisely how is not fully understood (Moore et al. 2013; Schmidl et al. 2018; Tirado-Magallanes et al. 2017). To identify specific genomic features that were preferentially influenced by developmental exposure, the DMRs across the CD4+ T-cell genome were annotated. This analysis showed that although the number of DMRs differed between all four comparison groups (Figure 5B), they spanned similar genomic features (Figure 5G–J). In all groups, over 60% of all DMRs were found in intergenic regions. In addition, DMRs were found in promoters, gene bodies (introns and exons), transcription termination sites (TTS), and untranslated regions (UTRs). Although promoter methylation generally has a strong influence on gene expression and function, only 2% of DMRs were found in promoter regions. These results show that both an immune challenge and developmental exposure influenced DNA methylation patterns across all genomic features in CD4+ T cells.

Figure 3. Evaluation of senescence-associated characteristics of CD4+ T cells from the offspring of vehicle and TCDD-treated dams during IAV infection. Adult (6–8 wk of age) developmentally exposed mice were infected with IAV. Mediastinal lymph nodes (MLNs) were analyzed 9 d after infection. (A,C,E,G) The number of CD4+ T cells from vehicle (7 mice) or TCDD (8 mice for KLRG1 and p16; 3 mice for CD107a and perforin) developmentally exposed offspring that express (A) KLRG1, (C) p16, (E) CD107a, or (G) perforin are shown as bar graphs. (B,D,F,H) Scatter plots show the percentage of (B) p16+, (D) KLRG1+, (F) CD107a+, or (H) perforin+ CD4+ T cells from infected mice that were developmentally exposed to vehicle (left) or TCDD (right). The numerical information is provided in Table S2. All offspring within a group were from separate dams. Values are mean ± SEM. *p ≤ 0.05, using Student’s t-test.
with CD4+ T cells from naïve mice, cells from IAV-infected offspring of vehicle dams had DMRs associated with 8,922 genes. Of these genes, 766 were also differentially expressed (Figure 6A). Although this accounts for only a quarter of all DEGs, these genes were involved in cellular pathways important for CD4+ T-cell functions during IAV infection, including NF-κB signaling, T-cell receptor signaling, and Th cell activation pathways (Figure 6B). Performing a similar comparison in CD4+ T cells from developmentally exposed offspring revealed DMRs associated with 11,559 genes, of which 707 were differentially expressed (Figure 6C). Among the DEGs with DMRs, some were similar to the vehicle group, such as those involved in NF-κB signaling and Th cell activation pathways (Figure 6D). Unlike vehicle offspring CD4+ T cells, the TCR signaling pathway was not among the top pathways influenced by DNA methylation and gene changes in offspring that had developmental AhR activation. Furthermore, CD4+ T cells from mice that were developmentally exposed to TCDD had reduced activities of co-stimulatory pathways, and Tec kinase signaling was increased.

**Modifying CD4+ T-cell Responses with DNA Methylation-Altering Drugs**

Given that important T-cell function pathways were influenced by developmentally induced changes in DNA methylation and gene expression, we sought to determine whether influencing DNA methylation modifies the impaired CD4+ T-cell responses caused by developmental exposure. To do this, we used pharmacological interventions that influence DNA methylation levels (Chen et al. 2012; TWH Li et al. 2012; Mato et al. 2013; Yoo et al. 2008). To assess the contribution of DNA hypomethylation we used SAM, which increases global DNA methylation levels by increasing the supply of methyl group donors (TWH Li et al. 2012; Mato et al. 2013). We posited that if developmental exposure affected T-cell function by reducing DNA methylation, then SAM treatment would abrogate this effect. At weaning, offspring that were developmentally exposed to vehicle or TCDD were randomly assigned to remain on untreated water (control group) or to receive water that contained SAM (Figure 7A). At maturity, mice were infected with IAV and then maintained on treated water throughout infection. SAM treatment did not influence the amount of water consumed by the mice during the treatment period (Figure S1A). Similarly, age-related weight gain was not influenced by treatment with SAM (Figure S1B). Treatment with SAM for ~4 wk did not impair the overall growth or health of the mice, consistent with prior reports that used DNA methylation-altering drugs in older mice for shorter periods of time (Chiang et al. 1996; TWH Li et al. 2012; Mato et al. 2013; Yoo et al. 2008). To the best of our knowledge, long-term SAM treatment has not been used in combination with IAV infection. Thus, we also determined whether drug treatment affected morbidity or mortality upon IAV infection. Upon infection, all mice lost weight (Figure S1), but there was no statistically significant difference between the groups. Furthermore, within each developmentally exposure group, SAM treatment did not influence morbidity and none of the offspring died as a result of infection (Figure S1). This suggests that the overall health of the mice prior to and during IAV infection was not adversely affected by SAM treatment.

The expansion of CD4+ T cells during IAV infection was not evaluated in developmentally exposed mice that were not treated with SAM from weaning through maturity. Consistent with previous observations, developmental exposure to TCDD resulted in significantly lower total CD4+ (Figure 7B) and virus-specific CD4+ T cells (Figure 7C) in IAV-infected mice by half and two-thirds, respectively. In contrast, after treatment with SAM, the number of CD4+ (Figure 7B) and virus-specific CD4+ T cells (Figure 7C) were not significantly different in infected vehicle and TCDD offspring. Although it did not restore the number to the same level observed in infected offspring of vehicle-treated
Figure 5. DNA methylation patterns in CD4+ T cells from naïve IAV-infected mice that were developmentally exposed to TCDD or vehicle. Mice were developmentally exposed to vehicle or TCDD, as described in Figure 1. Whole-genome bisulfite sequencing was performed using DNA from purified CD4+ T cells from naïve or IAV-infected adult mice (8–10 wk of age). For naïve offspring, peripheral lymph nodes from 1 mouse were pooled, and there was a total of three replicates per treatment group, with 1 offspring from unique dams used per replicate, for a total of 6 mice. For IAV-infected offspring mediastinal lymph nodes were used. Mice were sacrificed 9 d after infection, and there were three replicates per treatment group, with 4–6 offspring in each unique pool of cells for a total of 30 total mice used. All of the mice were sacrificed at the same time. (A) Box plots show global DNA methylation levels. The horizontal line denotes the mean, and boxes depict SEM. Differences between groups were evaluated using a two-way ANOVA with Tukey’s HSD post hoc test, and p-values are reported in Table S4. Groups with the same letter were not significantly different from each other. (B) The number of differentially methylated regions (DMRs) that were hypo- (blue; left side) or hyper- (red; right side) methylated are shown. The length of the bar is proportionate to the number of DMRs, and the number of DMRs is denoted on the bar segments. (C–F) Modified Manhattan plots show the distribution of hypomethylated (blue dots; left cluster) and hypermethylated (red dots; right cluster) DMRs across chromosomes in CD4+ T cells. (C,D) DMRs in CD4+ T cells from naïve (C) or IAV-infected TCDD (D) mice compared with vehicle controls. (E,F) DMRs in CD4+ T cells from vehicle (E) or TCDD (F) mice during IAV infection compared with naïve controls. The direction of hyper- and hypomethylation in (B) top two comparisons, (C) and (D) refer to the effect of developmental exposure to TCDD, or refers to the effect of infection in (B) bottom two comparisons, (E) and (F). Global methylation levels were analyzed by ANOVA with a Tukey’s HSD post hoc test. (G–J) DMRs were mapped to the genome. The total numbers of DMRs are shown. The percent of DMRs in intergenic, intron, TSS, TTS, 3'UTR, exon, noncoding, and 5'UTR regions are shown. The following comparisons were made: (G) DMRs from CD4+ T cells from naïve TCDD vs. naïve vehicle mice, (H) DMRs from CD4+ T Cells from IAV-infected TCDD vs. IAV-infected vehicle mice, (I) DMRs from CD4+ T cells from vehicle IAV-infected vs. vehicle naïve mice, and (J) DMRs from CD4+ T cells from TCDD IAV-infected vs. TCDD naïve mice. Note: ANOVA, analysis of variance; HSD, honestly significant difference; IAV, influenza A virus; SEM, standard error of the mean; TCDD, 2,3,7,8-tetrachlorodibenzo-p-dioxin; TSS, transcription start site; TTS, transcription termination site; UTR, untranslated region.
dams, treatment with SAM negated the down-modulatory effects of developmental AhR activation on these metrics of CD4+ T-cell expansion upon infection.

DNA methylation also influences CD4+ T-cell activation and differentiation (Durek et al. 2016; Schmidl et al. 2018). During IAV infection, developmental AhR activation resulted in a lower percentage of activated (CD44hiCD62L−) CD4+ T cells (Figure 7D) and correspondingly higher percentage of naïve (CD44loCD62L+) CD4+ T cells (Figure 7E). A similar difference (e.g., lower numbers of activated T cells and higher numbers of
naïve T cells) was observed in mice that were developmentally exposed to TCDD and treated with SAM (Figure 7D,E). Developmental exposure to TCDD resulted in a lower percentage of CD4+ T cells that differentiated into T helper-1 (Th1) cells (Figure 7F) and Tfh cells (Figure 7G). Although SAM was able to restore the lower Th1 response in IAV-infected offspring of TCDD-treated dams (Figure 7F), SAM was unable to restore the Tfh response in the same mice (Figure 7G). Thus, increasing DNA methylation levels alleviated deficits in clonal expansion and some, but not all, aspects of T helper differentiation following developmental exposure of mice to TCDD.

To define the effects of developmentally associated DNA hypermethylation, similar experiments were conducted by treating offspring with Zeb (Figure 8A), which lowers global DNA
methylation by blocking DNA methyltransferases (Chen et al. 2012; Yoo et al. 2008). Similar to SAM, water consumption and body weight were not affected by Zeb treatment (Figure S1A,B). Likewise, 4-wk treatment with Zeb did not adversely affect host resistance to infection (Figure S1C–E). However, unlike treatment with SAM, Zeb did not ameliorate the effects of developmental exposure on CD4⁺ T-cell clonal expansion. That is, the numbers of CD4⁺ (Figure 8B) and virus-specific CD4⁺ T cells (Figure 8C) were still significantly lower in IAV-infected Zeb-treated offspring of TCDD-treated dams. In contrast, treatment with Zeb restored the skewed percentage of activated and naïve CD4⁺ T cells (Figure 8D,E) in the offspring of TCDD-

**Figure 8.** Effects of Zebularine (Zeb) on CD4⁺ T cells from developmentally exposed mice after infection with IAV. (A) Mice were developmentally exposed to vehicle or TCDD. At weaning (21 d of age), mice were randomly selected be given normal vivarium water to maintain DNA methylation levels or water containing Zeb (0.2 mg/mL) to decrease DNA methylation levels. At 6–8 wk of age, mice developmentally exposed to vehicle (white bar) and TCDD (orange bar) were infected with IAV. Mediastinal lymph nodes (MLNs) were harvested 9 d after infection, and cells stained for flow cytometry. (B) The number of CD4⁺ T cells or (C) virus-specific [I-A⁺NP31–325⁺] CD4⁺ T cells were quantified from MLNs of mice on control (black outline) or Zeb (green outline) water. The percentage of (D) activated (CD44hiCD62L⁻) or (E) naïve (CD44loCD62L⁺) CD4⁺ T cells from control or Zeb mice. The percentage of (F) Th1 (Tbet⁺CD4⁺) or (G) Tfh (CXCR5⁺PD1⁺CD44hiCD4⁺) isolated from control or Zeb mice. For each group, 5–9 developmentally exposed offspring were used: control water vehicle (8), control water TCDD (5), Zeb water vehicle (9), and Zeb water TCDD (6). Due to a limited number of offspring, individual offspring were defined as the statistical unit, rather than the dam. All data shown are mean ± SEM. *, p ≤ 0.05, using a two-way ANOVA with Tukey’s HSD and Student’s t-test. #, p ≤ 0.05, using Student’s t-test. The numerical data in the graphs and p-values are provided in Table S7. Note: ANOVA, analysis of variance; HSD, honestly significant difference; H₂O, water; IAV, influenza A virus; SEM, standard error of the mean; T, TCDD; Tbet, T-box transcription factor; TCDD, 2,3,7,8-tetrachlorodibenzo-p-dioxin; Tfh, T follicular helper; Th, T helper; V, vehicle.
treated dams. Likewise, Zeb treatment abrogated the diminished Th1 cells (Figure 8F) and Tfh cells (Figure 8G) in TCDD developmentally exposed mice such that the frequency was similar to that of the vehicle-treated group.

Discussion

There is growing recognition that human health is influenced not only by contemporaneous exposures, but also by the environment experienced earlier in life, including in the womb (Boekelheide et al. 2012; Cao et al. 2016; Dallaire et al. 2004; Glynn et al. 2008; Heilmann et al. 2010; Hochstenbach et al. 2012; Jusko et al. 2014, 2016; Stolevnik et al. 2011; 2013). Yet, how exposures during development shape the immune system’s capacity to respond later in life are not well understood. The results of the present study show that early life exposure to the AhR agonist TCDD not only impacted the primary immune response to IAV, but also led to long-lasting consequences that also affected CD4+ T-cell responses to a second immune challenge with a similar pathogen. This illustrates the durable effect that early life exposure to an AhR agonist, such as TCDD, can have on the functional capacity of CD4+ T cells. Early life exposure of mice to environmentally derived AhR ligands affects CD4+ T cells in an enduring manner, but the cellular mechanisms remain poorly defined (Boule et al. 2014, 2015a, 2015b; Burke et al. 2019). Our results have elucidated one of the cellular mechanisms—developmental signaling through the AhR programmed DNA methylation and gene expression profiles of CD4+ T cells—and these alterations contributed to their impaired response to infection. Yet, this was not due to AhR targeting a single-cellular function or pathway. For instance, prior to infection, differences in gene expression were difficult to discern and differences in DNA methylation were modest. However, after infection, there were pronounced differences in both gene expression and DNA methylation patterns. This indicates that the AhR likely influences multiple pathways in T cells, which collectively regulate the responsive capacity of CD4+ T cells in a durable manner. This also suggests that some effects of early life exposures are cryptic and are only revealed upon another challenge, such as the call to arms triggered by infection.

A key finding from this present study was that developmental AhR activation changed DNA methylation patterns in CD4+ T cells prior to and during an immune challenge. This is consistent with a previous report that developmental AhR activation caused genome-wide changes in DNA methylation patterns in CD8+ T cells from mice (Winans et al. 2015). Similar to CD8+ T cells, CD4+ T cells from TCDD developmentally exposed mice had lower levels of global DNA methylation during IAV infection compared with cells from infected vehicle control offspring (Winans et al. 2015). This suggests that a common mechanism driving durable changes to cellular function after developmental exposure to an AhR agonist may be dysregulation of DNA methylation patterns in T cells. Further support for this idea was recently described within a human population. After early life exposure to AhR ligands, changes in DNA methylation patterns in human whole blood were observed, and they persisted into adulthood (Su et al. 2019). Moreover, methylation of 11 specific genes correlated with maternal exposure levels (Su et al. 2019). Seven of these 11 genes were differentially methylated in CD4+ T cells from naïve mice that were developmentally exposed to TCDD, and 5 of the 7 genes remained differentially methylated during IAV infection in mice (Table S8). Thus, in mouse and human studies, exposure to AhR-binding chemicals durably changes DNA methylation patterns in immune cells. These findings indicate that changes in DNA methylation may be a candidate biomarker for early life exposures.

Another key finding was that activation of the AhR during development modulated aspects of functional states associated with impaired responsive capacity: cellular senescence and exhaustion. These processes are characterized by poorer cellular proliferation and dysregulated effector functions, although depressed effector function is more typical of exhausted T cells (Crespo et al. 2013). Consistent with previously published reports that developmental AhR activation reduced interferon-gamma (IFNγ) production by CD4+ T cells during IAV infection (Boule et al. 2014; Burke et al. 2019), the results presented here further demonstrate that developmental AhR activation reduced other metrics of T-cell effector function, such as CD107a and perforin levels. Cellular exhaustion and senescence are commonly observed during chronic infection and aging, respectively (Akbar and Henson 2011). However, neither should occur when nonaged animals are infected with a pathogen that does not persist or form latent reservoirs, such as IAV. Thus, observing enhancement of markers related to these processes in nonaged animals and in the absence of chronic immune challenge is indicative of dysfunctional T-cell responses (Akbar and Henson 2011; Kasakovski et al. 2018). Interestingly, there was a reduction in the number and percentage of CD4+ T cells that express the exhaustion marker PD1. Using this classical marker of CD8+ T-cell exhaustion proved an inconclusive sole marker to identify CD4+ T-cell exhaustion because PD1 is also expressed by an important CD4+ T-cell helper subset: Tfh cells. Developmental AhR activation reduced the frequency of mouse Tfh cells during an immune response to IAV (Boule et al. 2014; Burke et al. 2019). Thus, it is more likely that the lower percentage of PD1+ cells observed in the present study reflects the impaired ability of CD4+ T cells to differentiate into Tfh cells, rather than diminished T-cell exhaustion. A previous study showed that CD4+ T cells from mice exposed to TCDD during development proliferated less in vitro after immune challenge, and changes in cellular metabolism may play a role in their reduced function (Burke et al. 2019). The work reported here extends this, and suggests that altered cellular metabolism may contribute to inappropriate T-cell exhaustion or premature senescence (Schurich and Henson 2014). In addition, analysis of DEGs in CD8+ T cells from IAV-infected mice that were developmentally exposed to TCDD indicated that early life AhR activation enhanced aspects of T-cell exhaustion in CD8+ T cells as well (Winans et al. 2015). Given that exhausted cells have an impaired ability to fight infections and protect the host, this could be one of the reasons that T-cell responses are reduced following developmental AhR activation.

Despite changes in gene expression in CD4+ T cells from infected mice, early life exposure to TCDD was not associated with significant differences in gene expression in CD4+ T cells from naïve mice. There are multiple theories that could account for this. A simplistic explanation is that this reflects the overall lack of overt changes to immune system development in mice developmentally exposed to this dose of AhR agonist (Boule et al. 2014; Vorderstrasse et al. 2004). Yet, a more likely explanation is the transcriptionally quiescent state of resting CD4+ T cells in the absence of an immune challenge (Feng et al. 2010, 2011; Zhang et al. 2018). The dearth of DEGs in naïve CD4+ T cells is similar to observations in CD8+ T cells from uninfected offspring of TCDD and control treated dams, in which there were only a handful of DEGs (Winans et al. 2015). Thus, developmentally induced changes in the responsive capacity of T cells are cryptic or latent in naïve cells. Yet, they are revealed once the transcription machinery is stimulated, and the cells respond to an immune challenge. Pervasive differences in DNA methylation were present across all genomic features of CD4+ T cells from naïve mice, which suggests that exposures during early life shape the way in
which CD4+ T cells are poised to respond to immune challenges encountered later in life and that DNA methylation patterns likely contribute to this poised or programmed state.

Another novel finding reported here was that developmentally induced differences in CD4+ T-cell functions were modifiable later in life by DNA methylation-altering drugs. This supports the idea that changes in DNA methylation patterns are part of the causative mechanism driving altered CD4+ T-cell responses that result from developmental AhR activation. Yet, altered CD4+ T-cell responsive capacity was not solely due to skewing of DNA methylation in a single direction, given that increasing or decreasing overall DNA methylation affected different aspects of CD4+ T-cell responses to IAV in offspring of TCDD-treated dams. Specifically, postnatal Zeb treatment restored diminished CD4+ T-cell differentiation in developmentally exposed mice. This is consistent with evidence that DNA de-methylation of specific genes plays a role in naive CD4+ T-cell differentiation into helper subsets (Komori et al. 2013; Y Li et al. 2012). Zeb reduces DNA methylation by blocking DNA methyltransferases, whereas SAM increases DNA methylation by providing an additional supply of donor methyl groups (Chiang et al. 1996; Zhou et al. 2002). As a consequence, SAM’s methyl group can be transferred to both DNA and histones (Chiang et al. 1996; Detich et al. 2003; Mews and Berger 2016). Thus, perhaps it is not surprising that SAM treatment elicited different outcomes than Zeb treatment. Although Zeb treatment did not affect the metrics of clonal expansion, SAM treatment restored the reduced clonal expansion of CD4+ T cells in mice that had been developmentally exposed to TCDD. This suggests that increasing DNA methylation benefits T-cell proliferation. This idea is supported by evidence from cancer studies, where aberrant DNA hypermethylation results in the silencing of tumor suppressor genes, increased proliferation, and tumor progression (Dahl et al. 2007; Gonzalez-Zulueta et al. 1995; Kane et al. 1997; Lin et al. 2017; Pilozzi et al. 2004). Although not examined in the context of immune cells, it is reasonable to hypothesize that SAM treatment restored impaired CD4+ T-cell proliferation through a similar mechanism. In addition, unlike Zeb, SAM was able to restore only the depressed Th1 cell response, and not the Th2 cell response. This subset-specific effect suggests that differentiation into Th1 and Th2 cells is differentially regulated by DNA methylation. Together, these observations indicate that the mechanisms underlying changes to CD4+ T-cell responses in developmentally exposed animals are complex and that DNA methylation levels influence cellular responses in discrete ways.

Given that enhancing or reducing overall DNA methylation restored some, but not all, of the impaired CD4+ T-cell responses in the adult offspring of TCDD-treated dams, developmental programming is not solely controlled by DNA methylation. Transfer of naive CD4+ T cells from developmentally exposed mice into recipients that have never been exposed revealed that they maintained the impaired responsive capacity (Boule et al. 2014). When considered together, these findings suggest that differences established by early life exposure to an AhR binding pollutant, such as TCDD, can be carried within the CD4+ T-cell lineage and that additional regulatory mechanisms contribute to their altered functional capacity. Similar to DNA methylation marks, histone modifications can be inherited by daughter cells throughout the lifetime of an organism, contributing to long-lasting influences on gene expression patterns (Henikoff and Greally 2016; O’Kane and Hyland 2019; Zheng et al. 2016). Although not studied in the context of developmental exposure or in T cells, the AhR is capable of interacting with histone deacetylases (HDACs), including HDAC1, HDAC2, and HDAC8 (Chang et al. 2014; Gomez-Duran et al. 2008; Schnekenburger et al. 2007; Wang et al. 2017). Although these studies were performed in tumor cell lines, they suggest that AhR activation influences histone modifications. Further support that early life exposures may elicit durable changes to histone modifications stems from investigations of developmental exposure to bisphenol A, which causes long-term histone modifications in nonimmune organs (Doherty et al. 2010; Greathouse et al. 2012; Strakovský et al. 2015). Together, these observations support the idea that in addition to DNA methylation, other epigenetic mechanisms likely contribute to the ways in which early life AhR activation influences CD4+ T cells in a durable manner.

The results presented in this study have broad-spanning implications because CD4+ T cells play integral roles in numerous immune responses, ranging from fighting infections and destroying tumor cells to driving allergic and autoimmune diseases (Knochelmann et al. 2018; MacLeod et al. 2009; Zhu and Paul 2008). We identified DNA methylation changes as one of the mechanisms underlying durable differences in CD4+ T-cell responsive capacity in mice resulting from early life AhR activation. Given that changes to DNA methylation patterns can be passed on to future generations, our results suggest that a person’s health may be programmed not only by the environment they experience, but also by the environmental exposures experienced by their parents. The work described here expands our understanding of how early life exposures can cause long-lasting changes to the immune system. Moreover, it brings us one step closer to devising optimally targeted therapeutics by uncovering DNA methylation as a mechanism of early life exposure-induced immune dysfunction.

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