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DNA hypomethylation ameliorates erosive inflammatory arthritis by modulating interferon regulatory factor-8

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Epigenetic regulation plays a crucial role in the pathogenesis of autoimmune diseases such as inflammatory arthritis. DNA hypomethylating agents, such as decitabine (DAC), have been shown to dampen inflammation and restore immune homeostasis. In the present study, we demonstrate that DAC elicits potent anti-inflammatory effects and attenuates disease symptoms in several animal models of arthritis. Transcriptomic and epigenomic profiling show that DAC-mediated hypomethylation regulates a wide range of cell types in arthritis, altering the differentiation trajectories of anti-inflammatory macrophage populations, regulatory T cells, and tissue-protective synovial fibroblasts (SFs). Mechanistically, DAC-mediated demethylation of intragenic 5'-Cytosine phosphate Guanine-3' (CpG) islands of the transcription factor *Irf8* (interferon regulatory factor 8) induced its re-expression and promoted its repressor activity. As a result, DAC restored joint homeostasis by resetting the transcriptomic signature of negative regulators of inflammation in synovial macrophages (*MerTK*, *Trem2*, and *Cx3cr1*), T_{REGs} (*Foxp3*), and SFs (*Pdnp* and *Fapα*). In conclusion, we found that *Irf8* is necessary for the inhibitory effect of DAC in murine arthritis and that direct expression of *Irf8* is sufficient to significantly mitigate arthritis.

arthritis | *Irf8* | decitabine | osteolysis | methylation

Chronic inflammation and autoimmune diseases are often characterized by persistent myeloid activation and proinflammatory output occurring at the level of hematopoietic stem and primitive myeloid progenitor cells. These cells acquire enhanced *in vitro* and *in vivo* propensity to generate inflammatory myeloid cells, the key perpetrators of tissue damage in arthritis (1, 2). This hyper-myeloopoiesis (and often neutrophilia) that is prevalent in inflammatory joint diseases may be a consequence of dysregulated epigenetic programs and inflammation-mediated silencing of repressor genes. Epigenetic regulation, especially through DNA methylation that alters transcriptional accessibility, plays a key role in joint homeostasis and in pathologic states including inflammatory arthritis (IA) (3–6). Alterations in epigenetic state and transcriptional accessibility have emerged as a critical modulator in various cells in a range of inflammatory diseases, including in synovial fibroblasts (SFs), immune cells, and peripheral blood mononuclear cells (7–15).

Studies of myeloid malignancies led to the discovery of agents targeting methylation, including the cytidine analog 5-aza-2-deoxycytidine [decitabine, Dacogen® (DAC)] (16, 17), which can demethylate previously silenced tumor suppressor genes leading to their reactivation. DAC causes DNA demethylation by inactivation of DNA methyltransferase-1 (DNMT1), the enzyme responsible for methylation (18, 19). At high doses, DAC exposure leads to the formation of covalent adducts between DAC and DNMT1, causing arrest of DNA synthesis in proliferating cells and cell death (17, 20). However, at low nanomolar doses, these adducts are degraded by the proteasome, and DNA synthesis is resumed in the absence of DNMT1 (21). Consequently, proliferating cells are hypomethylated, and previously silenced genes are re-expressed (16, 17, 20). DAC is an approved therapy for the treatment of myelodysplastic syndrome (MDS) and acute myeloid leukemia (22, 23). Approximately 50% of MDS patients demonstrate a clinical response after DAC treatment, with myelosuppression as a major adverse effect (24). Notably, inactivation of certain gene repressors, such as interferon regulatory factor 8 (*Irf8*), leads to development of chronic myeloid leukemia, characterized by myeloproliferation and systemic expansion of neutrophils (25).

Work in recent years has identified a family of interferon-regulatory factors (IRFs) as key regulators of myeloid lineage determination (26–31). In particular, gene deletion studies have shown that *Irf8* plays a prominent role in myeloopoiesis (32). *Irf8* expression is negligible in early hematopoietic progenitors and increases during differentiation in common/granulocyte myeloid progenitor (CMP/GMPs) fraction (29, 33). High expression of *Irf8* was

Significance

Inflammatory arthritis (IA) is characterized by dysregulation of immune cells that leads to joint swelling, pain, and tissue breakdown. Therefore, understanding the mechanisms underlying this pathology will advance therapeutic interventions. Recent work has suggested that inflammation is governed by dysregulation of DNA methylation, an epigenetic mechanism that regulates gene expression. In this study, we show that the clinically approved hypomethylation drug decitabine (DAC) inhibits murine IA. Furthermore, using transcriptomic screening, we identify the transcription factor interferon regulatory factor 8 (*Irf8*) as a key component of this mechanism. Specifically, DAC induces expression of *Irf8*, and direct transduction of *Irf8* ameliorates murine IA. These findings position DAC and *Irf8* as potential IA therapeutic targets.

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found in monocyte-DC progenitors (MDPs) and common monocyte progenitors (cMoPs) (34, 35) limiting their proliferation, and *Irf8*^{-/-} mice display an increased number of MDPs, cMoPs, and osteoclasts and develop osteoporosis (36, 37). Most importantly, *Irf8* inhibits neutrophil development while *Irf8*^{-/-} mice exhibit severe neutrophilia (30). Mechanistically, *Irf8* inhibits neutrophilia by directly binding C/EBP α (the transcription factor that promotes neutrophil differentiation) and blocking C/EBP α chromatin binding and transactivation of target genes. Interestingly, binding of C/EBP α to target genes is facilitated by the methyl group found in the gene promoters (30, 38). Thus, hypermethylated *Irf8* loses its suppressive activity facilitating neutrophilia, inflammatory myeloid proliferation, and hyper-osteoclastogenesis (37).

In this study, using assay for whole-genome bisulfite sequencing (WGBS), unsupervised single-cell RNA sequencing (scRNA-seq), and fluorescence-activated cell sorting (FACS) analyses, we report that the demethylating drug DAC reverses inflammation-induced epigenetic, transcriptomic, and cellular changes in immune and nonimmune synovial cells. Furthermore, DAC activates cellular

suppressors, primarily *Irf8*, to attenuate inflammation and erosion in joints of murine models of IA.

Results

The Hypomethylating Agent DAC Ameliorates IA and Modifies Synovial Cell Populations. The pathology of arthritis features distal joint swelling, synovitis, and synovial tissue destruction. Intraperitoneal administration of a low dose of DAC (0.12 mg/kg) dramatically reduced clinical score and inhibited distal joint swelling when administered therapeutically after disease onset (Fig. 1 *A* and *B* and *SI Appendix*, Fig. S1 *A* and *C*) or prophylactically (*SI Appendix*, Fig. S1 *B*) in STIA (serum transfer-induced arthritis), collagen-induced arthritis (CIA), and collagen antibody-induced arthritis (CAIA) models. Given that NF- κ B activation is an indicator of inflammation in arthritis (39–41), the inflammatory index of IA was measured in RelA NF- κ B reporter mice demonstrating that DAC significantly inhibited RelA-Luciferase activity in a dose-dependent manner (*SI Appendix*, Fig. S1 *D* and *E*).

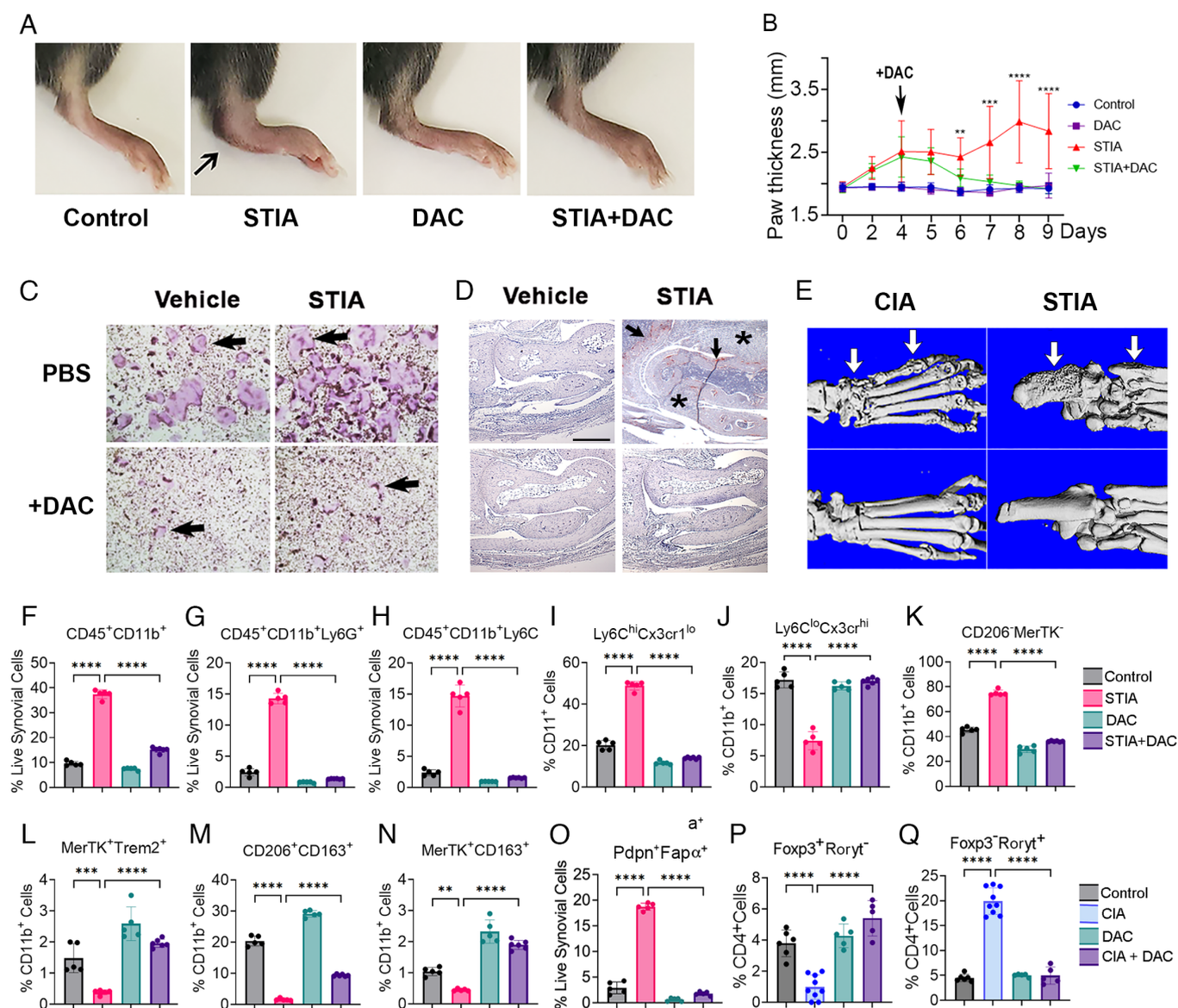


Fig. 1. Low dose of DAC inhibits IA. (*A* and *B*) Paw images and thickness measurements of therapeutically treated STIA mice. Two-way ANOVA, $N = 10$; $****P < 0.0001$ (STIA vs. STIA + DAC). (*C–E*) Ex-vivo osteoclastogenesis, IHC TRAP-staining, and 3D μ CT reconstruction of paws from CIA and STIA mice \pm DAC. Error bars represent mean \pm SD. One-way ANOVA $*P < 0.05$, $**P < 0.01$, $***P < 0.001$, and $****P < 0.0001$, unless otherwise indicated. (*F–Q*) Multichannel FACS analysis of synovial cells from IA distal joints \pm DAC (0.12 mg/kg bw).

Because IA is complicated by joint erosion with notably elevated osteoclastogenesis, these parameters were examined *in vivo* and *ex vivo*. To this end, osteoclast progenitors isolated from STIA mouse joints and bone marrow differentiated more readily and with higher propensity into multinucleated osteoclasts when cultured *ex vivo*, whereas DAC nearly completely halted osteoclastogenesis (Fig. 1C). Corroborating these findings, immunohistochemistry tartrate-resistant acid phosphatase (TRAP) staining of joint sections showed dramatic reduction of osteoclasts in DAC-treated STIA mice (Fig. 1D; arrows), and mitigated inflammation (Fig. 1D; asterisks). Further, μ CT analysis showed that DAC significantly protected against bone erosion (Fig. 1E; arrows). Buttressing the relevance of these outcomes to joint function, mouse behavioral and pain sensitivity measurements demonstrated that the joint function of DAC-treated mice is similar to control counterparts (SI Appendix, Fig. S1 F–M). Considering the potential adverse effects of this drug, DAC at the highest dose tested (250 nM) had a negligible effect on BMM and T cell cycle compared with cell cycle arrest at 1% FBS (SI Appendix, Fig. S1 N and O). Critically, there were no significant changes in mouse body weight at a dose triple the optimal dose (0.25 mg/kg) we routinely used (SI Appendix, Fig. S1P), suggesting that therapeutic dosing of DAC does not lead to systemic cytotoxic effects.

Because IA synovitis is dominated by hyper-myeloprolieration, enhanced circulating proinflammatory (MHCII⁺Ly6C⁺) macrophages (M Φ), and neutrophilia (CD11b⁺Ly6G⁺), the frequency of synovial cells in paws was quantified. FACS analysis revealed that overall CD11b⁺ population and Ly6G^{hi}, Ly6C^{hi}, and Cx3cr1^{lo} subpopulations were increased, whereas frequency of resident synovial M Φ expressing Cx3cr1, MerTK, Trem2, CD163 and the anti-inflammatory M Φ -like cell type markers CD206 and CD163 were reduced at peak arthritis (Fig. 1 F–N). Intraperitoneal administration of low doses of DAC was sufficient to reverse synovial cellular changes highlighted by reduced frequency of overall CD11b⁺ myeloid population, granulocytes/neutrophils (Ly6G⁺), inflammatory Ly6C⁺ M Φ , MerTK⁺ inflammatory M Φ , MHCII⁺/MHCII⁺ macrophage ratio, and increased (or gained) frequency of Cx3cr1^{hi}, MerTK⁺ and CD163⁺ anti-inflammatory M Φ (Fig. 1 F–N and SI Appendix, Fig. S2 A–C). Furthermore, recognizing that SFs and T cells play key roles in IA pathogenesis, the frequency and subtypes of these cells were monitored in IA and IA+DAC joints with FACS analysis using STIA, CIA, and CAIA mouse models. Whereas activated SFs were highly elevated in CIA and CAIA, DAC inhibited CIA- and CAIA-associated activation of SF resident cells underscored by a significant decrease in Pdpn⁺Fap α ⁺ inflammatory SFs in the synovial joints (Fig. 1O). Further, DAC increased T_{REG} (Foxp3⁺Roryt⁻) and inhibited T_{H17} (Foxp3⁻Roryt⁺) cells in CIA (Fig. 1 P and Q) and CAIA (SI Appendix, Fig. S2 D–F), consistent with recently published reports (42, 43).

STIA Mice Exhibited Significantly Increased Myeloid Clusters, while DAC Drug Treatment Remarkably Reduced Inflammatory Myeloid Populations in Joint Synovium of STIA Mice. To further understand how DAC modulates arthritis, we examined the transcriptional profile of synovial cells derived from control and STIA mice in the absence or presence of DAC. Employing scRNA-seq, normalized gene counts from each of the four experimental groups were evaluated using principal component analysis. We observed that samples within each group (n = 3 samples/group) clustered together, validating consistent transcriptional profile among replicates within each experimental group (SI Appendix, Fig. S3). Hierarchical clustering analysis of STIA + DAC samples demonstrated a similar gene expression profile to control samples when compared to STIA-nontreated samples (SI Appendix, Fig. S4).

This result implies that DAC treatment alters the transcriptional profile of STIA mice reflecting their nondiseased controls more closely. Next, we interrogated the effect of DAC treatment on synovial cell subpopulations. Unsupervised scRNA-seq analysis revealed that control and DAC-treated groups had 3 and 2 myeloid cell clusters, respectively. STIA mice exhibited 9 myeloid cell populations which were significantly reduced to 5 myeloid cell subsets when STIA mice received DAC drug treatment (i.e., STIA + DAC group; SI Appendix, Figs. S5–S8B). Myeloid cells were identified by conventionally used cell markers including Mrc1, Trem2, S100a8, Adgre1, Retnla, and Napsa (44). In addition, other immune cells such as B cells, T cells, and neutrophils as well as fibroblasts were also detected and annotated according to their corresponding markers (SI Appendix, Figs. S5–S8C). Unique cell populations in the different clusters in each experimental condition depicted in SI Appendix, Figs. S5–S8 are summarized in SI Appendix, Table S1. Evaluating distinct cell populations (including myeloid cells, fibroblasts, B cells, T cells, NK cells, and endothelial cells) resulting from experimental conditions (SI Appendix, Fig. S9) shows that when compared to control mice, the percentage of myeloid cells in the joint synovium was significantly increased in the STIA mice (~3.2-fold increase), and it was decreased by the DAC treatment (i.e., STIA + DAC). Remarkably, more than 80% of synovial cells in the STIA mice were myeloid cells. The percentages of the rest of synovial cell populations such as fibroblasts, B cells, T cells, and endothelial cells were decreased in the STIA mice compared with control mice (SI Appendix, Fig. S9) (Table 1).

Integrative Analysis Identified 12 Distinct Myeloid Cell Populations in the Synovium. Immune cells are pivotal constituents of the inflammatory response. To delineate how inflammation and DAC treatment modulate innate immune myeloid cell phenotypes in joint synovium, we subsetted and integrated myeloid cell clusters detected from control, STIA, and STIA + DAC mice. Unsupervised scRNA-seq analysis using canonical correlations analysis (CCA) identified 12 unique myeloid cell populations in the synovium across three experimental mouse groups: 1) proinflammatory *Wfdc21/Rdh12/Lcn2*⁺ M Φ , 2) proinflammatory *Tnfr/Csf1/Clec4n*⁺ M Φ , 3) anti-inflammatory *C1qa/Mrc1/Ppf4*⁺ M Φ , 4) anti-inflammatory *Spp1/Cd36/Arg1*⁺ M Φ , 5) *Chil3/Plac8/Ly6c2*⁺ Mo, 6) antigen-presenting *Krld1/H2-Aa/Clec10a*⁺ Mo-DCs, 7) profibrotic *Gm26917/Sparc/Col3a1*⁺ M Φ , 8) proinflammatory IFN-induced *Gm13822/lift3/Isg15*⁺ M Φ , 9) TAM-like *Nrp2/Stmn1/Atpif1*⁺ M Φ , 10) *Wdfy4/Trim35/Irf8*⁺ Mo-Dc/Neu, 11) tissue-resident *Lyz1/Mgl2/Clec4b1*⁺ M Φ , and 12) proliferating *Mpo/Ngp/Ctsg*⁺ M Φ . (Fig. 2 A–C and SI Appendix, Fig. S10). To identify which of our M Φ populations may belong to osteoclasts and/or are involved in proinflammatory signaling, we generated violin plots focused on key osteoclast genes (*Acp5*, *Ctsk*, *Tnfrsf11a*, *Fos*, *Prdm1/Blimp1*, *Nfatc1*, *Mmp9*, and *Ddstamp*) and NF- κ B pathway genes (*Rela*, *Nfkb1*, and *I11b*) (Fig. 2D). It appears that *Nrp2/Stmn1/Atpif1*⁺ M Φ had high expression of osteoclast genes, suggesting these cells may be associated with osteoclast lineages. Consistent with our previous findings, proinflammatory M Φ populations (particularly, *Wfdc21/Rdh12/Lcn2*⁺ M Φ and *Tnfr/Csf1/Clec4n*⁺ M Φ) demonstrated high expression of the genes involved in NF- κ B pathway. As expected, inflammation significantly increased myeloid cell numbers in the STIA mice compared to the myeloid cells detected in the control mice. Additionally, four distinct myeloid cell populations (i.e., *Wfdc21/Rdh12/Lcn2*⁺ M Φ , *Tnfr/Csf1/Clec4n*⁺ M Φ , IFN-induced *Gm13822/lift3/Isg15*⁺ M Φ , and *Wdfy4/Trim35/Irf8*⁺ Mo-Dc/Neu) were only detected in STIA and STIA + DAC mice but not in control mice (Fig. 2E). Next, we compared changes in the distinct macrophage populations due

Table 1. Labeled antibodies used for FACS

Reagent	Cat#	Company
Ly6G BV421	127628	BioLegend
CD101 PE-Cy7	25-1011-82	BioLegend
Cd11b BV510	101263	BioLegend
CD163 APC	17-1631-82	BioLegend
CD206 PE	141706	BioLegend
CD4 BV421	100443	Thermo
CD45 BV650	103151	BioLegend
Cell stainign buffer	420201	BioLegend
CX3CR1 FITC	149020	BioLegend
F4/80 SB600	63-4801-82	BioLegend
FAPa APC	BS-5758R	BioLegend
FCy blocker	101302	BioLegend
Fixable viability dye eFluor™ 780	65-0865-18	Thermo
foxp3 PE	12-5773-82	Thermo
Ikbz PerCP-eFluor 710	46-6801-82	Thermo
Ly6C SB780	128041	Thermo
MerTK PE-Cy7	25-5751-82	Thermo
MHCII APC	107614	Thermo
PDPN PE-Cy7	25-5381-82	Thermo
ROR AF488	53-6981-82	Thermo
Trem2 FITC	MA5-28223	Thermo
True-nuclear™ transcription factor buffer set	424401	Thermo

to experimental conditions (SI Appendix, Fig. S11). Consistent with our previous analysis, four distinct myeloid cell populations (i.e., *Wfdc21/Rdh12/Lcn2+* MΦ, *Tnf/Csf1/Clec4n+* MΦ, IFN-induced *Gm13822/lift3/Isg15+* MΦ, and *Wdfy4/Trim35/Irf8+* Mo-Dc/Neu; dashed square lines) were significantly increased in STIA mice when compared with control mice. DAC treatment showed a trend toward decreasing the percentage of *Wdfy4/Trim35/Irf8+* Mo-Dc/Neu cells in the synovium when compared to STIA treatment. Additionally, control mice had significantly higher synovial cell percentage of proliferative *Mpo/Ngp/Ctsg+* MΦ (i.e., enriched in *Top2a* and *Mki67* expression) vs. STIA and STIA mice.

DAC Treatment Significantly Restored Anti-Inflammatory Cell Populations and Decreased Inflammatory Gene Expression: Unique Signatures and Functionality of MΦ Populations.

Next, the transcriptomic profiles and functionalities of synovial macrophage populations were further explored. We observed that in STIA, proinflammatory *Wfdc21/Rdh12/Lcn2+* MΦ demonstrated high expression of *Il1a* and *Tnf* (SI Appendix, Fig. S12). GO term analysis indicated that *Wfdc21/Rdh12/Lcn2+* proinflammatory MΦ express genes enriched for immune system responses, neutrophil chemotaxis, and response to IL-1 (SI Appendix, Fig. S13A). IFN-induced *Gm13822/lift3/Isg15+* MΦ phenotype demonstrated that the cellular response to IFNβ and *Tnf/Csf1/Clec4n+* MΦ is involved in apoptotic process regulation (SI Appendix, Fig. S13A). The anti-inflammatory *Spp1/Cd36/Arg1+* MΦ had high expression in *Arg1*, *Trem2*, and *Vegfa* with moderate *Il6* expression levels, while anti-inflammatory *C1qa*

Mrc1/Ppf4+ MΦ were enriched in *Mrc1*. Tissue-resident *Lyz1/Mgl2/Clec4b1+* MΦ subset had the highest expression levels of *Cd81*, *Cd163*, and *Il10* among all MΦ populations (SI Appendix, Fig. S12). In GO term functional analysis, *C1qa/Mrc1/Ppf4+* MΦ demonstrated chemotaxis, endocytosis, regulation of ERK cascades, and cellular response to IFNγ, while *Spp1/Cd36/Arg1+* MΦ exhibited angiogenesis functionality. Tissue-resident *Lyz1/Mgl2/Clec4b1+* MΦ had a gene expression profile enriched for antigen processing and presentation functionality via MHC class II (SI Appendix, Fig. S13B and Table S1). *Chil3/Plac8/Ly6c2+* Mo exhibited genes enriched for cellular response to IL-4 and positive regulation of TNF biosynthetic process. *Wdfy4/Trim35/Irf8+* Mo-Dc/Neu showed antigen processing and presentation functionalities (SI Appendix, Fig. S13C). Collectively, these findings are consistent with FACS data and demonstrate that IA propagates proinflammatory myeloid subsets whereas DAC fortifies anti-inflammatory myeloid populations.

Identification of Differentiation Trajectories of Anti-Inflammatory MΦ Populations and Their Key Transcriptional Regulators.

As anti-inflammatory MΦ populations were greatly increased in the mice receiving DAC treatment, pseudotime analysis was conducted in the clusters containing two distinct anti-inflammatory MΦ subsets and their unique functionalities, i.e., *Spp1/Cd36/Arg1+* MΦ with functionality for angiogenesis (GO:0001525) and *C1qa/Mrc1/Ppf4+* MΦ with functionality for responding to IFNγ (GO:0071346), using *Chil3/Plac8/Ly6c2+* Mo as the root of lineage trajectories (Fig. 3A). Next, genes that are coregulated around trajectory branch points were analyzed to further obtain insights into the genetic profiles governing cell fate decisions. Trajectory branch analysis using Monocle3 identified 22 gene modules that may contribute cell fate specification of *Chil3/Plac8/Ly6c2+* Mo (SI Appendix, Fig. S14). Based on gene expression levels, modules 4 and 9 may contain genes controlling cell fate decision of *Chil3/Plac8/Ly6c2+* Mo into anti-inflammatory *Spp1/Cd36/Arg1+* MΦ, while genes in modules 6 and 8 may be responsible for *Chil3/Plac8/Ly6c2+* Mo into anti-inflammatory *C1qa/Mrc1/Ppf4+* MΦ. We observed increased expression levels of *Arg1*, *Ccl9*, *Crip1*, and *Vegfa* when *Chil3/Plac8/Ly6c2+* Mo gradually differentiated into *Spp1/Cd36/Arg1+* MΦ (SI Appendix, Fig. S15A).

To determine which transcription factors (TFs) may be regulating this differentiation trajectory, we performed the TF binding motif analysis using the RcisTarget R package and TRRUST database. The results indicated that *Sp3*, *Smarcc1*, and *JunD* are the putative key TFs driving this differentiation process with *Smarcc1* and *JunD* having higher expression levels and percentage of cells in *Spp1/Cd36/Arg1+* MΦ (Fig. 3B).

The differentiation of *Chil3/Plac8/Ly6c2+* Mo into *C1qa/Mrc1/Ppf4+* MΦ was best described by gene modules 6 and 8 (Fig. 3A and SI Appendix, Fig. S14). This trajectory showed upregulation of *Irf7* and *Stat1* at the later differentiation stage (i.e., *C1qa/Mrc1/Ppf4+* MΦ) (SI Appendix, Fig. S15B). TF binding motif analysis shows that the primary TFs regulating the differentiation route are *Stat1* and *Bach2* (Fig. 3C). Protein–protein network for modules 6 and 8 was also constructed, indicating *Ifit3*, *Ifit1*, *Ddx58*, *Usp18*, and *Isg15* are top 5 nodes with the highest degree and *Stat1-Fyn*, *Fyn-B2m*, *B2m-Tap1*, *Stat1-Ifit3*, and *Cmpk2-Dck* are top 5 edges with highest betweenness centrality (SI Appendix, Fig. S16). In this regard, *Stat1* has been shown recently to be required for efficient recruitment of the signaling complex of *Irf8* (45), and lupus GWAS studies indicated that *Irf8* directly interacts with *Ifit1* and *Stat1*, with the latter identified as disease susceptibility gene (46). Since *Irf8* was identified as a potential downstream target and epigenetically modified and activated by

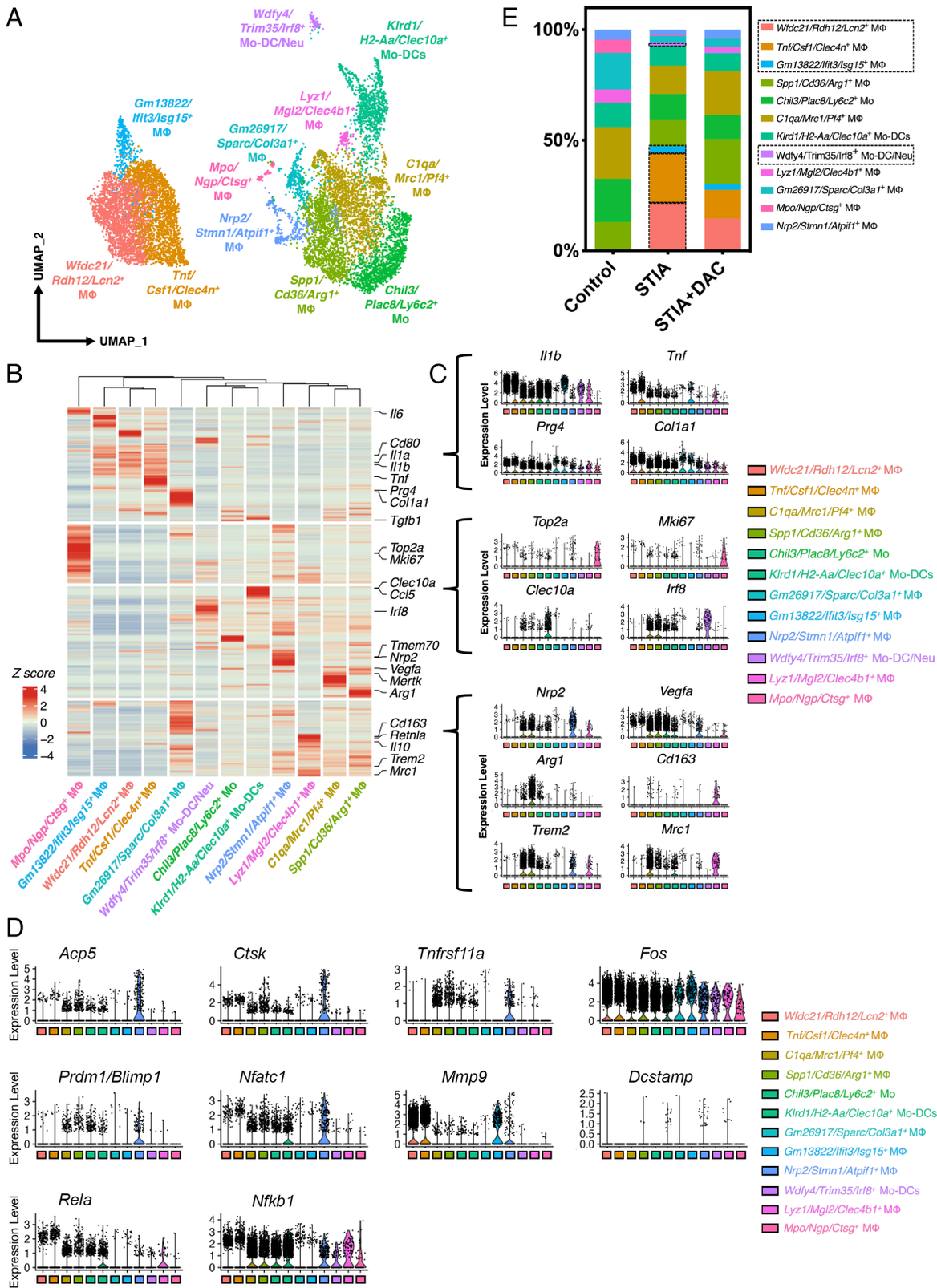


Fig. 2. (A) Twelve distinct myeloid cell populations with their specific markers were identified using CCA integration of all myeloid populations from control, STIA, and STIA + DAC mice. (B–D) Heatmap and violin plots showing unique transcriptomic profiles defining identified myeloid cell populations and preosteoclasts. (E) scRNA-seq analysis showing the changes of myeloid cell populations in the joint synovium harvested from the control, STIA, and STIA + DAC mice. Dashed rectangles indicate populations only observed in the STIA and STIA + DAC mice but not in the control group.

DAC in MΦ, DCs, and neutrophils (26, 31, 34, 47), we further investigated expression levels of *Irf8* in these two anti-inflammatory MΦ populations in control, STIA, and STIA + DAC experimental

conditions. DAC treatment significantly promoted both levels and percentage of the cells expressing *Irf7* and *Irf8* in *Spp1/Cd36/Arg1*⁺ MΦ and *C1qa/Mrc1/Pf4*⁺ MΦ populations (Fig. 3D).

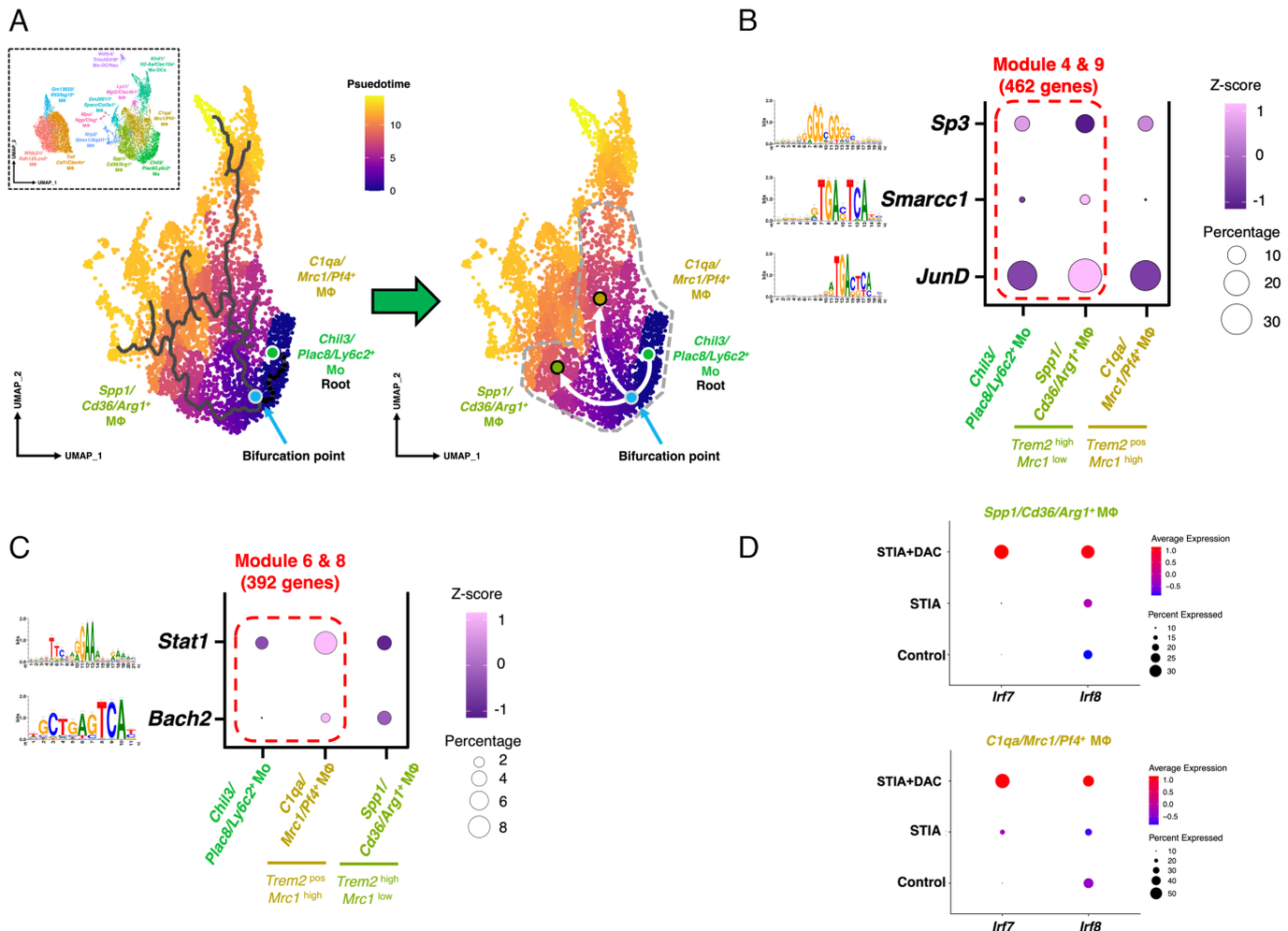


Fig. 3. (A) Pseudotime ordering indicates a lineage bifurcation of *Chil3/Plac8/Ly6c2*⁺ Mo into either anti-inflammatory *Spp1/Cd36/Arg1*⁺ MΦ or anti-inflammatory *C1qa/Mrc1/Pf4*⁺ MΦ. (B) TF binding motif analysis showing that *Sp3*, *Smarcc1*, and *JunD* are putative TFs regulating differentiation of *Chil3/Plac8/Ly6c2*⁺ Mo into *Spp1/Cd36/Arg1*⁺ MΦ. (C) *Stat1* and *Bach2* are potential regulators driving *Chil3/Plac8/Ly6c2*⁺ Mo into *C1qa/Mrc1/Pf4*⁺ MΦ lineage. (D) DAC treatment increased both levels and percentage of the cells expressing *Irf7* and *Irf8* in both *Spp1/Cd36/Arg1*⁺ MΦ and *C1qa/Mrc1/Pf4*⁺ MΦ in STIA + DAC mice.

To characterize the phenotype of these cells in vitro, synovial macrophages were collected from WT and *Irf8* null mice and subjected ex vivo to \pm DAC treatments. Flow cytometry and qPCR were carried out for markers of the two *Spp1* and *C1qa* populations. The data depicted as *SI Appendix, Fig. S17* show that cells treated with DAC express *Mrc1*, *Arg1*, *Cq1a*, *CD36*, and *Spp1* corroborating the in vivo phenotype.

DAC Caused Major Gene Methylation Changes in IA. Epigenetic regulation has emerged as a hallmark of pathologic states, including IA. To better understand potential therapeutic mechanisms, WGBS was performed on *CD11b*⁺ cells isolated from synovial tissues from control mice, STIA mice, and STIA mice after DAC treatment ($n = 2$ for WGBS per group). Approximately 18 million CpGs assayed at 10 \times coverage were identified. Comparing control to STIA mice, a significant global increase in CpG methylation was noted (Fig. 4A; red graph shift to the right), which was ameliorated by DAC treatment (Fig. 4B). Analysis of differentially methylated regions (DMRs) identified 29,808 DMRs between control and STIA mice, 75% of which were hypermethylated, and 25% were hypomethylated (Fig. 4C). In comparing STIA mice to DAC-treated STIA mice, a smaller subset of only 773 significant DMRs was identified, with 15% hypermethylated and 85%

hypomethylated (Fig. 4D). In both comparison groups, methylation changes were noted predominantly within CpG islands (Fig. 4E), and DMRs were distributed among gene bodies and intergenic regions (*SI Appendix, Fig. S15A*). Pathway analyses on DMRs highlighted peptidyl-tyrosine modification as differently methylated in control vs. STIA mice (*SI Appendix, Fig. S15B*), which aligns with the known pathologic role of tyrosine kinases in IA (48). In comparing STIA vs. DAC-treated STIA mice, pathway analysis highlighted broader pathways, including GTPase-mediated signal transduction, focal adhesions, and kinase activity (*SI Appendix, Fig. S18B*). We performed gene set enrichment analysis on differentially accessible genes, based on changes in predicted enhancer methylation at nearby loci. In the intersection of genes with both differential accessibility between STIA vs. Ctrl and those with differential expression in STIA vs. STIA + DAC, we identified 34 potentially regulated genes (*SI Appendix, Table S2*). These encompass a range of intronic and distal intergenic regions (*SI Appendix, Table S2*). Gene enrichment analysis reassuringly highlighted overrepresentation of pathways including regulation of bone resorption (by GO Biological Process), NFATC1 signaling (via Elsevier Pathway Collection), and CX3CR1⁺ cells (by MAGNET 2023) (*SI Appendix, Fig. S19*). The latter is consistent with phenotypic changes in myeloid cells described in our FACS and scRNAseq analyses.

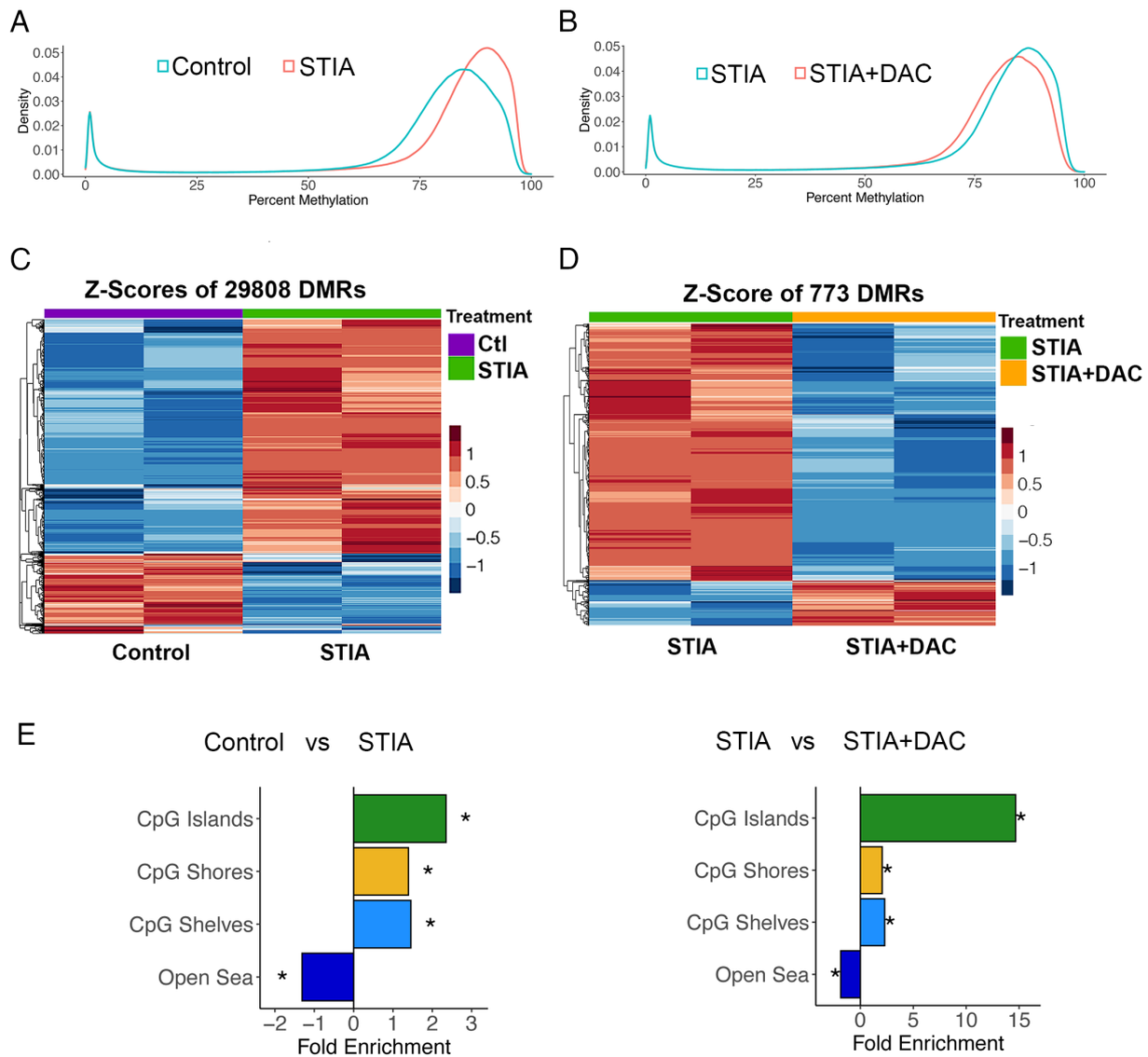


Fig. 4. IA displays significant genomic methylation and the demethylation drug DAC reverses major gene methylation changes. (A and B) Smoothed single CpG methylation density plots from control vs. STIA mice (A) and STIA vs. STIA + DAC (B) treated mice, showing overall hypermethylation of STIA mice which is largely reversed by DAC treatment. (C and D) Heatmaps showing high-level clustering of DMRs in control vs. STIA groups (C) (reflecting overall hypermethylation), and STIA vs. STIA + DAC groups (D) (reflecting hypomethylation). (E) Analysis of DMRs between control vs. STIA and STIA vs. STIA + DAC. Enrichment occurs predominantly at CpG islands, but also at other regions including the open sea.

Irf8 Is Hypomethylated by DAC and Potently Inhibits IA. Our data show that inflammation broadly reprograms gene methylation (Fig. 4) and the anti-inflammatory transcription factor *Irf8* is a potential target of DAC (Fig. 3). Therefore, we reasoned that DAC could be working through demethylating inflammation-hypermethylated suppressors, like *Irf8*, that target myelopoiesis and neutrophilia.

Combining random forest (RF) classifiers and support vector machine algorithm analyses of the top 10% DMRs from our WGBS data, *Irf8* was identified among the DAC-regulated genes in STIA, evident by DAC demethylation of *Irf8* at CpG islands (Fig. 5A). Top regulated DMRs were determined via a generalized least squares regression model including permutation testing of a pooled null distribution, as defined in DMRichR. Analysis of a specific subset of loci for methylation revealed that a region within the *Irf8* promoter is significantly hypomethylated in STIA + DAC mice (Fig. 5A). Next, we identified a 485 bp CpG island (DNA methylation/demethylation enzyme binding site, indicated as the green box in *SI Appendix*, Fig. S20A) in the murine proximal *Irf8* promoter, which is highly conserved across

species (35, 49, 50), confirming that *Irf8* is subject to epigenetic regulation.

To validate these findings, we performed the *Irf8* CpG methylation assay targeting this CpG island. Whereas inflammatory signals (LPS and TNF) and RANKL increased methylation (dark blue) of the *Irf8* promoter region in bone marrow macrophages (BMMs) by nearly 25 to 40%, DAC entirely inhibited this methylation (Fig. 5B). Even more robust was the staggering 90% *Irf8* CpG methylation observed in CD11b⁺ cells at the peak of arthritis (dark red), which was drastically prevented by DAC in vivo (Fig. 5C). Accordingly, mRNA and protein expression of *Irf8* in synovial CD11b⁺ cells isolated from ankles of control, STIA, DAC, and DAC-treated STIA mice was diminished under inflammatory conditions (e.g., STIA), whereas treatment with DAC significantly restored or elevated its expression (Fig. 5D and E and *SI Appendix*, Fig. S20B).

Confirming that *Irf8* is a direct target of DAC, the drug failed to inhibit TNF α expression and only slightly inhibited joint swelling in mice lacking *Irf8* (LysM⁻Cre mediated *Irf8* deletion).

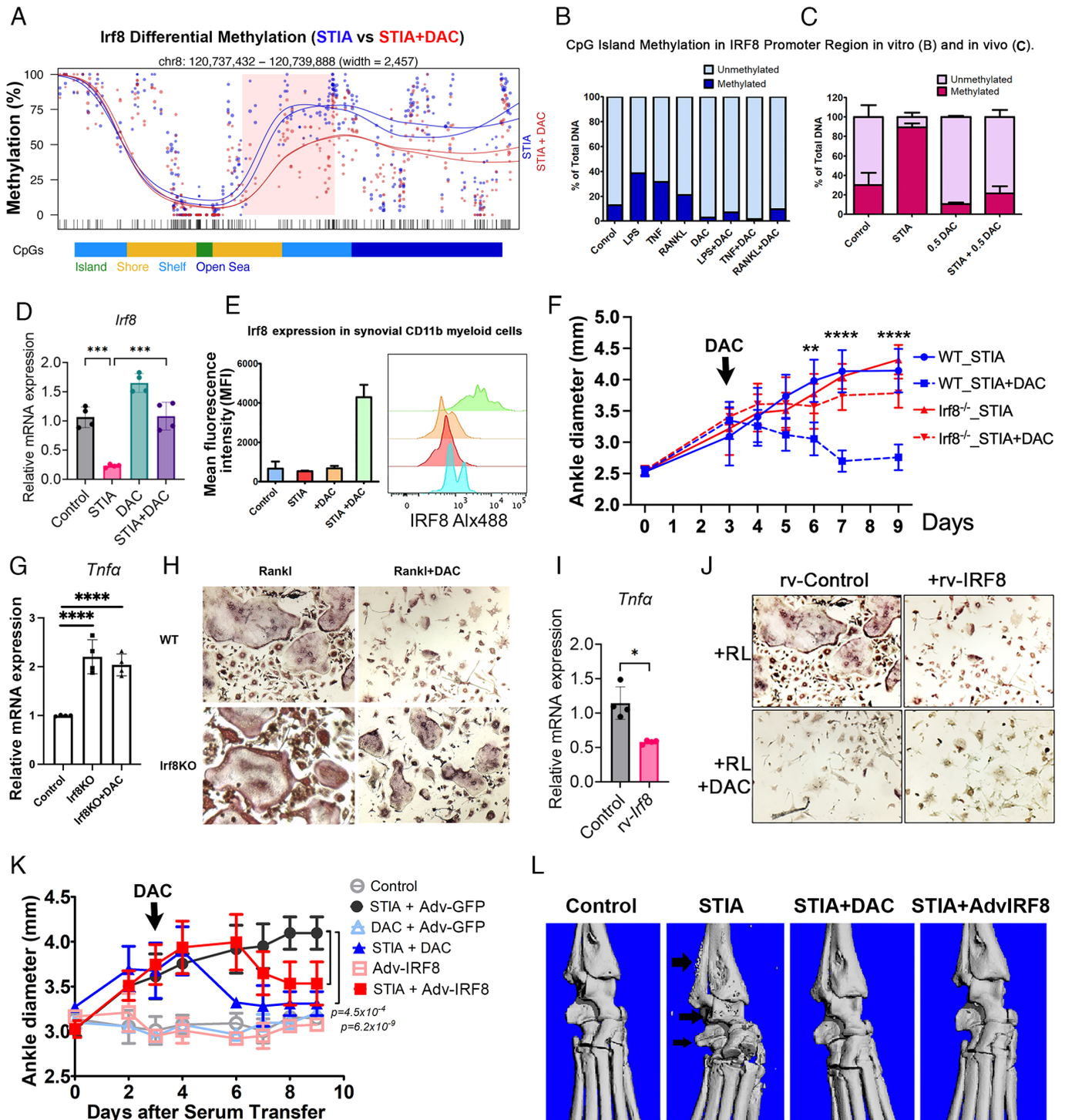


Fig. 5. *Irf8* partially mediates the DAC antiosteolytic effect. (A) The *Irf8* promoter region is significantly demethylated comparing STIA vs. STIA + DAC. (B) One million CD11b⁺ bone marrow cells were plated on a six-well tissue culture plate and stimulated with either PBS (control), 2 nM DAC, 1 ng/mL LPS, 50 ng/mL TNF, or 100 ng/mL RANKL, with or w/o 2 nM DAC for 24 h. (C) CD11b⁺ cells were isolated from paws of control, STIA ±DAC mice. Genomic DNA was subjected to methylation assay using primers specific to *Irf8* CpG island following Qiagen's Methyl II PCR Kit. (D) qPCR for *Irf8* fold change compared to control. Error bars represent mean ± SD. One-way ANOVA. ****P* < 0.001. (E) *Irf8* protein expression and protein levels (mean fluorescence intensity = MFI) quantified in CD11b⁺ cells isolated by FACS sorting from the synovium of the various treatment groups. DAC dose in vivo 0.12 mg/kg bw. (F) Ankle swelling (diameter) in WT and *Irf8* KO mice subjected to STIA ± DAC. The arrow indicates the starting time of DAC injection in the therapeutic protocol. Two-way ANOVA (Tukey's multiple comparisons test). Error bars represent mean ± SD. Two-way ANOVA (Tukey's multiple comparisons test). ***P* < 0.01; *****P* < 0.0001 WT_STIA+DAC compared with *Irf8*KO_STIA+DAC. (G and H) TNFα qPCR and osteoclastogenesis (+RANKL) from WT and *Irf8*KO cells ±DAC. Error bars represent mean ± SD. One-way ANOVA. *****P* < 0.0001. (I and J) TNFα qPCR and osteoclastogenesis from cells expressing retroviral (rv) *Irf8* ±DAC. Student's *t* test, **P* < 0.05. (K) Local injection of adenoviral (Adv) *Irf8* (10⁷ pfu/10 μL) to the joints compared with therapeutic injection of DAC (0.12 mg/kg). Error bars represent mean ± SD. Two-way ANOVA (Tukey's multiple comparisons test). *P* values are indicated in the figure. (L) μCT analysis of paws from (K).

Furthermore, DAC treatment failed to inhibit osteoclastogenesis in *Irf8* KO (knockout) cells. (Fig. 5 F–H). More importantly, forced expression of retroviral *Irf8* (*rvIrf8*) in BMMs, mimicking

DAC, potentially inhibited expression of inflammatory cytokines and osteoclasts (Fig. 5 I and J and *SI Appendix*, Fig. S20 C–E). Finally, we tested the potential therapeutic and prophylactic

efficacy of *Irf8* in vivo. Adenoviral *Irf8* particles (10^7 pfu/10 μ L) were intraarticularly injected in the knee joint space and subcutaneously near the ankle and knee joints of control and STIA mice alongside other groups treated with DAC. Adv-*Irf8* robustly inhibited joint swelling in STIA mice nearly comparable to DAC (Fig. 5K and *SI Appendix*, Fig. S20F). This protective effect was also corroborated with abolishing bone erosion as shown by μ CT (Fig. 5L). Collectively, these data show that *Irf8* is hypermethylated and inhibited under inflammatory conditions; DAC hypomethylates *Irf8* and induces its expression; virally expressed *Irf8* inhibits inflammatory cytokines, joint swelling, osteoclasts, and bone erosion; and deletion of *Irf8* not only exacerbates osteoclastogenesis and inflammation but also blunts the DAC inhibitory effects on osteoclasts and inflammatory signals. These findings strongly suggest that *Irf8* is among the principal targets of DAC in myeloid cells and is necessary to facilitate DAC actions in these cells.

Discussion

DAC Inhibits Inflammatory Characteristics of Synovial Cells and Ameliorates Murine IA. IA is characterized by hyper-myeloproliferation and neutrophilia (51) underpinned by an exuberant cytokine microenvironment. It is now apparent that cellular, metabolic, and transcriptomic alterations of hematopoietic progenitors that elicit proinflammatory phenotype are at the crux of this process (52). Likewise, the inflammatory milieu reprograms synovial FBs, CD4+ and CD8+ T cells—primarily T regulatory cells—and impedes their protective and regenerative functions (53, 54). Under inflammatory conditions, the genomes of myeloid, lymphoid, and several other cell types undergo hypermethylation that hinders homeostatic functions by dampening cellular suppressors and exacerbating disease pathology (5, 6, 55, 56). Therefore, we surmised that hypomethylating agents hold promise to reactivate methylated suppressor genes (57) and restore tissue homeostasis. Using several mouse models of arthritis, we first demonstrate that IA is accentuated by exacerbated myeloproliferation and neutrophilia. Specifically, FACS and transcriptomic analyses demonstrate increased propensity of proinflammatory (M1 type) M Φ , atypical LyC6⁺ M Φ , Ly6G⁺ neutrophils, Trem2⁻, Cx3cr1⁻ proinflammatory M Φ , Pdpn⁺Fap α ⁺ FBs, and reprogramming of Foxp3⁺ T_{REGS} into Rorygt⁺ T_H17 pathogenic cells. Furthermore, we noted a significant increase in RANK⁺ progenitors that are hypersensitive to RANKL and drive the osteolytic process. Remarkably, administration of the low dose of the hypomethylation drug DAC reversed these in vitro cellular changes and restored cellular homeostasis. Even more convincingly, administration of low doses of DAC in vivo ameliorated IA in STIA, CIA, and CAIA mouse models and therapeutically restored tissue homeostasis, highlighting the detrimental role of inflammatory macrophages, neutrophils, pathogenic T helper cells, and activated synovial FBs. These findings are supported by previous evidence wherein DAC and 5-azacitidine efficiently altered experimental IA pathogenesis (42, 58–60).

DAC Reduced Myeloid Cell Clusters in the Synovium and Modified Their Transcriptomic Signature. Significant strides have been made to identify unique synovial cell clusters to advance joint therapy (61, 62). In this regard, joint homeostasis is governed by tissue-resident M Φ (Trem2⁺ and Cx3cr1⁺), lining layer, and sublining fibroblasts (Pdpn⁺) that together form protective cellular barriers in the joint (63). We carried out scRNA-seq studies and identified 12 cell clusters reflecting the heterogeneity of synovial cells. Notably, the inflammatory phenotype of these clusters elicited by IA was largely curbed in the presence of DAC through

restoration of their homeostatic anti-inflammatory phenotype (MerTK⁺, CD206⁺, CD163⁺, and Cx3cr1⁺). These changes were corroborated by diminished or complete prevention of IA features in mice.

Our scRNA-seq analysis revealed that DAC dramatically reduced myeloid cell populations in STIA joints. In total, 12 unique myeloid populations were detected in the synovium across all treatment groups, of which four proinflammatory groups were present only in the arthritic joints, whereas DAC-treated STIA mice exhibited an increased percentage of five anti-inflammatory macrophage populations compared to arthritic mice. Of interest, expression of members of the TAM family receptor tyrosine kinase MER (MerTK) is diminished in IA, yet treatment with DAC restores its expression and its anti-inflammatory signature highlighted by elevated expression of IL-10, as has been also shown previously (63–65). Additional work is required to establish the potential anti-inflammatory role of MerTK⁺ in IA. Similar changes in Trem2 were also observed. We also noted significant changes in the expression of Cx3cr1, a recently reported marker of specialized tissue-resident M Φ that form immunological barrier at the synovial lining to protect the joint (66). IA-induced reduction of Cx3cr1 expression was significantly restored following exposure to DAC. Of special note, we identified clusters expressing *Isg15*, *Clec10a*, and *Clec4b1* with previously undescribed roles in IA pathogenesis. In this regard, we have recently unveiled that *Isg15* is an essential regulator of osteoclasts (67) and inflammatory osteolysis (39). In addition, we report a unique tissue-residing *Lyz1/Mgl2/Clec4b1*⁺ M Φ subset, which has the highest expression levels of *Cd81*, *Cd163*, and *Il10* among all M Φ populations. This cell population which is characterized by expression of galactose/GalNa-c-lectin (MGL) (68) and Fc γ R-coupled C-lectin receptor dendritic cell immune-activating receptor (Clec4b1) (69) displays immunomodulatory properties through induction of protective T cell responses to resolve inflammation (69). Additional studies are required to delineate the role of Clec4b1 and Clec10a in IA.

Collectively, these findings suggest that DAC acts systemically and at the local joint environment to rewire homeostatic immune and anti-inflammatory activities of tissue-resident and synovium-infiltrating cells. More importantly, DAC appears to restore tissue homeostasis by reversing the inflammatory phenotype of multiple cell types including myeloid, lymphoid, SFs, and potentially other cell types. Transcriptomic profiling further identified three inflammatory populations responsive to IFN γ and IL-1. On the other hand, transcriptomic signature and GO term functional analysis of other clusters revealed cellular responses including regulation of ERK signaling, angiogenesis, antigen processing, fibrosis, collagen fibril organization, and efferocytosis. These findings suggest that DAC regulates a wide array of cell functions including cell cycle, mitochondrial function, cytokine responsiveness, immune regulation, and tissue remodeling. At the transcription factor level, DAC appears to regulate specific TFs including *Stat1*, *Bach2*, *Irf7*, and *Irf8* in late-stage differentiation of Mrc1⁺ and Arg1⁺ anti-inflammatory macrophage populations, as discussed below.

Hypomethylation by DAC Restores Suppressor Gene Function. Our study demonstrates that IA dramatically hypermethylates the genome of synovial cells leading to pathogenic phenotypes, consistent with recent reports (3, 10, 15, 70, 71). We identified hypermethylation of the suppressor gene *Irf8* as a key therapeutic target. Our focus on *Irf8* as a potential modulator of DAC-induced inhibition of IA is supported by evidence pointing to its potent role as modulator of myelopoiesis and neutrophilia, and that hypermethylation of *Irf8* or lack of *Irf8* promotes neutrophilia and osteoclastogenesis (25, 30, 31, 36, 72), whereas

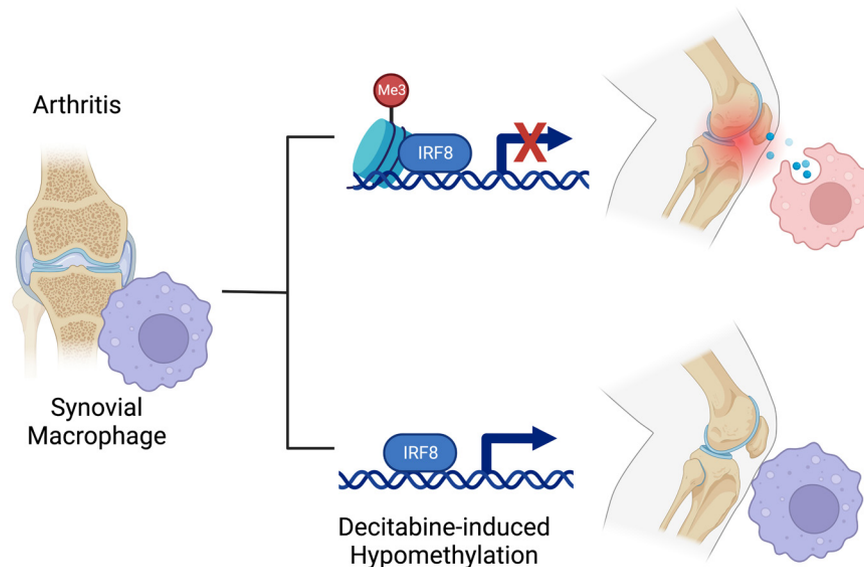


Fig. 6. Summary graphical model. IA induces hypermethylation and inhibition of the suppressor function of Irf8 resulting with joint swelling, synovitis, and tissue degradation. Administration of the hypomethylating agent DAC reactivates Irf8 leading to inhibition of joint inflammation.

hypomethylation of Irf8 increases Irf8 expression and activity, and suppresses neutrophil differentiation in C/EBP α -dependent manner (31, 37). Hence, we hypothesized that inflammation blunts expression of suppressive factor Irf8 via hypermethylating its promoter region leading to myeloid proliferation, neutrophilia, and RA propagation. We also surmised that demethylation of Irf8 by DAC may enable reprogramming of myeloid differentiation to resolve IA. Indeed, our transcriptomic studies unveiled several Irf8-expressing myeloid populations. *Wdyf4/Trim35/Irf8+ Mo-Dc/Neu* cells have the highest expression of Irf8. We also show that two anti-inflammatory macrophages (*C1aq/Mcr1/Pf4* and *Spp1/Cd36/Arg1*) have strong Irf8 expression (Fig. 2B; Irf8 violin plot). These two populations were largely increased in the STIA + DAC group (55.3% and 74%, respectively). It is noteworthy that *Spp1* and *Mcr1* have been associated with reparative and regenerative anti-inflammatory macrophages (73, 74).

Our data indicate that DAC demethylates STIA-induced hypermethylation of the genome and *Irf8* promoter regions. More importantly, DAC restores Irf8 protein expression in myeloid cells from STIA mice, inhibits osteoclastogenesis, and most notably, Irf8 mediates the DAC effect and directly inhibits IA inflammatory and erosive characteristics. These findings are consistent with the established antiosteoclast function of Irf8 (37, 72, 75, 76). It remains unclear whether DAC targets Irf8 directly or through proximal regulators. BATE, which was among the highest expressed TFs we detected, has been shown to sustain Irf8 auto-activation (35), suggesting that it is a potential target of DAC. Further, our observation that DAC regulates a unique PU.1 expressing cluster substantiates studies wherein binding of the primary myeloid transcription factor *PU.1* and the osteoclast TF *Nfatc1* with Irf8 binding sites in myeloid progenitors and differentiated OCs were described (77), with evidence that Irf8 regulates *Nfatc1* expression (72, 77, 78). Furthermore, we detected upregulation of STAT1 in *C1aq/Mcr1/Pf4* anti-inflammatory macrophages. In this regard, recent studies have shown that STAT1 is required for efficient Irf8 signaling complex assembly and function (45) and a lupus GWAS showed that Irf8 directly binds to *Ifit1* and *Stat1*, the latter identified as disease susceptibility gene (46). Thus, STAT1-Irf8 complex is potentially required

for DAC-induced differentiation of *Chil3/Plac8/Ly6c2+* monocytes into *C1aq/Mcr1/Pf4* anti-inflammatory macrophages. Our transcriptomic studies also demonstrated expression of JunD, a transcription factor that has been shown to limit expression of Irf8 (79). Consistently, recent studies have shown that JunD is hypermethylated and its expression is down-regulated under inflammatory conditions. Conversely, overexpression of JunD elicited an anti-inflammatory response (80).

Collectively, these observations suggest that the Irf8 regulatory network is far more elaborate than anticipated. Therefore, Irf8 appears as a central intersection point for several regulatory elements, and hence, it is not surprising that re-expression of Irf8 directly or through DAC-mediated hypomethylation overrides proximal suppressions and inhibits inflammation and bone erosion in IA (Fig. 6).

Materials and Methods

Note: Additional methodology is included in *SI Appendix* section.

Study Design. The objective of this study was to determine the efficacy of DAC and IRF8 in the treatment of murine IA. Mice were randomly assigned to experimental groups of the different arthritis models. The sample size was determined based on prior experience with penetrance of each model. Outliers were included in the final calculations. This study utilized a low dose of DAC and demonstrated its efficacy and nontoxic effect *in vitro* and *in vivo*. All experiments reflect a minimum of five independent biological replicates. Investigators were blinded to the experimental groups and data was assessed by at least three independent investigators.

Statistical Analysis. Male and female mice were used at equal ratios. All experiments represent biological replicates and were repeated at least three times, unless otherwise stated. Statistical analyses were performed using appropriate statistical tests using GraphPad Prism v9. All graphs were generated using Prism as well. Multiple treatments were analyzed by one-way or two-way ANOVA followed by Tukey's test multiple comparisons test. Student's *t* test was used for comparing two groups. *P* values are indicated where applicable. **P* < 0.05, ***P* < 0.01, ****P* < 0.0005, and *****P* < 0.0001. Mouse activity and pain measurements, osteoclast counts, and immunostaining data were scored by investigators blinded to the experimental conditions.

Data, Materials, and Software Availability. Transcriptomic dataset is available in NCBI GEO (81).

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