Epigenetic regulation of NfatC1 transcription and osteoclastogenesis by nicotinamide phosphoribosyl transferase in the pathogenesis of arthritis

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Abstract

Nicotinamide phosphoribosyltransferase (NAMPT) functions in NAD synthesis, apoptosis, and inflammation. Dysregulation of NAMPT has been associated with several inflammatory diseases, including rheumatoid arthritis (RA). The purpose of this study was to investigate NAMPT’s role in arthritis using mouse and cellular models. Collagen-induced arthritis (CIA) in DBA/1J Nampt+/− mice was evaluated by ELISA, micro-CT, and RNA-sequencing (RNA-seq). In vitro Nampt loss-of-function and gain-of-function studies on osteoclastogenesis were examined by TRAP staining, nascent RNA capture, luciferase reporter assays, and ChIP-PCR. Nampt-deficient mice presented with suppressed inflammatory bone destruction and disease progression in a CIA mouse model. Nampt expression was required for the epigenetic regulation of the Nfatc1 promoter and osteoclastogenesis. Finally, RNA-seq identified 690 differentially expressed genes in whole ankle joints which associated (P < 0.05) with Nampt expression and CIA. Selected target was validated by RT-PCR or functional characterization. We have provided evidence that NAMPT functions as a genetic risk factor and a potential therapeutic target to RA.

Introduction

Rheumatoid arthritis (RA) is characterized by synovial inflammation and bone erosion1,2. Unfortunately, current therapies for arthritis are inadequate and there remains a need for additional therapeutic targets.

Nicotinamide phosphoribosyltransferase (NAMPT) is an essential gene3 which functions in NAD synthesis, apoptosis, and inflammation4. NAMPT is expressed in nearly all organs, tissues, and cells examined4. Because of its pleiotropic functions, dysregulated NAMPT expression has been implicated in the pathogenesis of several diseases, including arthritis, though the role of NAMPT in these disorders remains to be elucidated4. Using a collagen-induced arthritis (CIA) mouse model, Busso et al. demonstrated that FK866, a NAMPT inhibitor, effectively reduced the severity and progression of arthritis5. The progression of CIA was also slowed by selective siRNA knockdown of NAMPT in Ly6Chigh monocytes6. However, no study has been conducted to systematically evaluate the molecular mechanisms of Nampt in arthritis in well-established Nampt knockdown (Nampt−/−) and Nampt overexpression (NamptOE) mice to substantiate that Nampt is a genetic risk factor and potential therapeutic target in RA.

This study investigated the molecular mechanisms of Nampt in arthritis through integrative approaches of CIA mouse models, in vitro cellular experimentation, and...
transcriptional profiling. We validated Nampt’s involvement in arthritis using a CIA mouse model\(^7,8\) in wild-type and Nampt heterozygous knockdown DBA/1J mice, and we investigated further the pathways underlying Nampt’s mechanism in arthritis through loss-of-function and gain-of-function studies experiments in mouse bone marrow-derived macrophages (BMM) and RNA sequencing (RNA-seq) of CIA mouse tissue. Selected targets from RNA-seq discovery were experimentally validated. Our results support the hypothesis that Nampt is a genetic risk factor and a potential therapeutic target in RA.

**Results**

**Decreased inflammation and suppressed bone erosion in collagen-induced arthritis in DBA/1J Nampt\(^{-/-}\) mice**

To explore the molecular mechanisms of altered Nampt expression in arthritis, we characterized CIA in Nampt\(^{-/-}\) and Nampt\(^{+/+}\) mice. The progression of observable inflammation was less severe in Nampt\(^{-/-}\) mice compared to Nampt\(^{+/+}\) mice with a significant difference in the median arthritic index from day 28 to 70 post-immunization (Fig. 1a). The incidence of arthritis (index score > 1) was decreased slightly in Nampt\(^{-/-}\) mice (85%, 11/13) compared to Nampt\(^{+/+}\) (100%, 11/11). At day 70 post-immunization, the mice were euthanized for tissue isolation and sample analysis.

To elucidate potential mechanisms by which the induction of arthritis is suppressed in Nampt\(^{+/+}\) mice, we measured the serum levels of the arthritogenic anti-mouse CIA auto-antibody, which plays a crucial role in the initiation of CIA\(^7\). The serum levels of the CIA antibody were decreased significantly in Nampt\(^{-/-}\) mice compared with Nampt\(^{+/+}\) mice (Fig. 1b). The decreased immune response corresponded with lower levels of circulating Nampt in the heterozygous mice compared to wild-type mice (Fig. 1b).

To evaluate the effect of CIA on focal bone loss, we analyzed the hind paw and talus by micro-computed-tomographic (micro-CT) imaging. Micro-CT revealed visible differences between the hind paw and talus of Nampt\(^{+/+}\) and Nampt\(^{-/-}\) mice with CIA (Fig. 1c). To quantify bone loss, we contoured the talus to determine bone volume (BV). The average total BV of the talus were significantly lower in both CIA Nampt\(^{+/+}\) and Nampt\(^{-/-}\) mice compared with their non-immunized controls (Fig. 1d). Correction of the BV by the total volume of the talus (BV/TV) revealed bone loss was significantly milder in CIA Nampt\(^{+/+}\) compared with CIA Nampt\(^{-/-}\) mice (Fig. 1d).

**Attenuated osteoclastogenesis in Nampt-deficient primary BMM and RAW 264.7 cells**

Our finding that Nampt\(^{-/-}\) mice were protected against bone erosion in CIA led us to hypothesize that Nampt plays a critical role in osteoclast differentiation and that decreased Nampt expression attenuates osteoclast formation. To test this potential mechanism, we examined Rankl-dependent osteoclast differentiation in BMM isolated from Nampt\(^{+/+}\) and Nampt\(^{-/-}\) mice (Fig. 2). TRAP activity, a histochemical marker of osteoclastogenesis\(^9\), was detected by cell staining (Fig. 2a). M-CSF-dependent Nampt\(^{+/+}\) BMM were able to produce TRAP\(^+\) cells following a 3-day stimulation with M-CSF and Rankl, while stimulated Nampt\(^{-/-}\) BMM produced significantly fewer TRAP\(^+\) cells. The relative number of differentiated osteoclasts from Nampt\(^{+/+}\) BMM was significantly lower compared with Nampt\(^{+/+}\) BMM (Fig. 2a). Decreased differentiation of Nampt-deficient macrophages into TRAP\(^+\) cells correlated with lower expression of Acp5, the gene encoding TRAP protein. mRNA levels for key osteoclast markers, including Nfatc1, Dc-stamp, and Cathepsin K, were also lower in osteoclasts derived from Nampt\(^{+/+}\) BMM compared with Nampt\(^{+/+}\) controls, thus validating the attenuation of osteoclastogenesis (Fig. 2b). Western blot analyses verified that Nampt expression was decreased in Nampt\(^{+/+}\) BMM relative to Nampt\(^{+/+}\) BMM (Fig. 2c).

To investigate the mechanism by which Nampt-deficiency attenuated macrophage differentiation into TRAP\(^+\) cells, we examined the expression of Nfatc1, an essential transcriptional regulator of osteoclast differentiation\(^10\). Silencing of Nampt expression by siRNA in RAW 264.7 cells inhibited expression of Nfatc1 protein and mRNA (Fig. 3a). To determine the regulatory mechanism of Nfatc1 expression, we tested mRNA stability in response to decreased Nampt levels. Although there was a decrease in mRNA stability, it was not sufficient to account entirely for the loss in protein levels (data not shown). Therefore, we investigated Nfatc1 transcription using luciferase reporter and nascent RNA capture assays. The relative luciferase activity decreased significantly in RAW 264.7 cells co-transfected with the Nfatc1 promoter luciferase reporter and Nampt siRNA relative to the Nfatc1 promoter luciferase reporter and scrambled siRNA control cells (Fig. 3b). Nascent RNA capture in RAW 264.7 cells validated further that synthesis of Nfatc1 mRNA required Nampt expression (Fig. 3b).

Epigenetic re-modeling of the Nfatc1 promoter plays a critical role in Nfatc1 expression\(^11\). Therefore, we performed ChIP-PCR analyses in Nampt-deficient RAW 264.7 cells to characterize the transcriptional regulation of the Nfatc1 promoter. The interaction of acetylated histones, which represents open chromatin, with the Nfatc1 P1 promoter decreased following Nampt knockout (Fig. 3c). Conversely, the presence of methylated histones, which represents closed chromatin, increased at the Nfatc1 P1 and P2 promoter regions following Nampt
siRNA knockdown compared with scrambled siRNA transfected controls (Fig. 3c). These observations corresponded with decreased histone acetyltransferase (HAT) activity in RAW 264.7 cells subjected to Nampt knockdown (Fig. 3d). The epigenetic remodeling was consistent with the decreased transcriptional activity observed by luciferase reporter and nascent RNA capture assays (Fig. 3b).

To determine if Nampt enzymatic activity was required for the Nampt–NfatC1–osteoclastogenesis pathway, we treated RAW 264.7 and Nampt+/+ BMM with the enzymatic inhibitors, FK866 and MC4 during Rankl-induced differentiation. Formation of TRAP+ cells was decreased significantly in RAW 264.7 cells treated with FK866 and MC4 relative to the DMSO-vehicle control cells (Fig. 4a, b). Differentiation of Nampt+/+ BMM also decreased significantly in response to FK866 and MC4 relative to the control cells (Fig. 4c, d). In both RAW 264.7 cells and Nampt+/+ BMM, MC4 was as effective as MTX in blocking the formation of TRAP+ cells. The combination of MC4 and MTX in RAW 264.7 and Nampt+/+ BMM significantly blocked osteoclast differentiation compared with MC4 or MTX treatment alone (Fig. 4b, d). These findings strongly support the requirement of Nampt enzymatic activity to promote osteoclastogenesis.

Transcriptomic profiling of whole ankle joints in CIA mice identifies genes and pathways associated with osteoclastogenesis

To investigate the molecular mechanisms of Nampt in the pathogenesis of arthritis, we sequenced RNA isolated from whole ankle joints of Nampt+/+ and Nampt−/− mice with and without CIA (Fig. 5). We initially determined the DEG in two comparison groups. There were 1613 DEG in CIA Nampt+/+ ankle compared with non-CIA control Nampt+/+ ankle with and without CIA (Fig. 5a). We initially determined the DEG in two comparison groups. There were 1613 DEG in CIA Nampt+/+ ankle compared with non-CIA control Nampt+/+ ankle with and without CIA (Fig. 5a). We initially determined the DEG in two comparison groups. There were 1613 DEG in CIA Nampt+/+ ankle compared with non-CIA control Nampt+/+ ankle with and without CIA (Fig. 5a). We initially determined the DEG in two comparison groups. There were 1613 DEG in CIA Nampt+/+ ankle compared with non-CIA control Nampt+/+ ankle with and without CIA (Fig. 5a). We initially determined the DEG in two comparison groups. There were 1613 DEG in CIA Nampt+/+ ankle compared with non-CIA control Nampt+/+ ankle with and without CIA (Fig. 5a). We initially determined the DEG in two comparison groups. There were 1613 DEG in CIA Nampt+/+ ankle compared with non-CIA control Nampt+/+ ankle with and without CIA (Fig. 5a). We initially determined the DEG in two comparison groups. There were 1613 DEG in CIA Nampt+/+ ankle compared with non-CIA control Nampt+/+ ankle with and without CIA (Fig. 5a).
Cxcl5, Tnn, Fam229a, and Atp6v0d2, while the top five down-regulated genes were Lep, Polr3g, Serpina3c, Pnpla3, and Nnat. Although the majority of the 690 DEG in the CIA Nampt<sup>+/−</sup> ankle were protein-coding genes, there were 10 non-coding RNA genes that were differentially expressed (Supplemental Table 1).

The finding that two osteoclast-specific genes, Oscar and Atp6v0d2, were among the top 5 up-regulated genes...
supported our observation that decreased Nampt expression limited osteoclastogenesis in both RAW 264.7 and BMM. Additionally, the osteoclast-specific genes \textit{Mmp9}, \textit{Acp5}, \textit{Ctsk}, and \textit{Dcstamp} were differentially expressed with increased expression in CIA \textit{Nampt}\textsuperscript{+/+} mice (Supplemental Table 2). Calcium influx plays a critical role in the activation of \textit{Nfatc1}\textsuperscript{14}. We identified 19 significantly upregulated calcium metabolism genes...
within the 690 DEG (Supplemental Table 3). Our previous finding that knockdown of NAMPT expression significantly attenuated calcium influx into human pulmonary artery endothelial cells\(^{15}\), supports our hypothesis that decreased Nampt expression may inhibit osteoclast differentiation by inhibiting calcium influx via mediating expression of calcium metabolism genes.

To gain further insight into the biological functions associated with Nampt mediated pathogenesis of CIA, we submitted the 690 DEG for pathway analysis. These include several pathways associated with osteoclastogenesis, including the Inhibition of Matrix Metalloproteases and the Role of Osteoblasts, Osteoclasts and Chondrocytes in Rheumatoid Arthritis (Fig. 5c, Supplemental Table 4).

IPA analysis predicted target molecules in the dataset of 690 DEG that are either activated or inhibited by well-characterized upstream regulators. TNF\(\alpha\) (5.575 activation \(z\)-score; 7.73E\(-46\) P-value of overlap), TGF\(\beta\)1 (5.436; 9.4E\(-32\)), and LPS (5.985; 4.87E\(-32\)) are the top 3 activators, while the top 3 inhibitory regulators are the drugs dexamethasone (\(-3.074; 2.12E\(-34\)) and rosiglitazone (\(-4.791; 7.56E\(-24\)), and the kinase inhibitor PD98059 (\(-3.847; 1.33E\(-23\)) (Supplemental Table 5). Upstream regulator analysis also identified factors, such as cytokines (Tnfsf11, Csfr1), transcriptional regulators (Fos, Nfatc1), and signaling proteins (Nfkb, Erk1/2, P38 Mapk, Src, P13k, Akt) that are known to promote osteoclast differentiation and bone resorption (Supplemental Table 6).

We next performed functional enrichment analyses for GO terms and KEGG to predict potential biological processes and pathways involved in the Nampt associated pathogenesis of CIA. The top biological processes linked with the 690 DEG include GO:0007155—cell adhesion, GO:0022610—biological adhesion, GO:001501—skeletal system development, GO:0001503—ossification and GO:0060348—bone development (Fig. 5d, Supplemental Table 6).
Table 7). The top pathways identified by KEGG analysis include mmu04512:ECM—receptor interaction and mmu04142:Lysosome (Supplemental Table 8).

Validation of RNA-seq result by RT-PCR assay of the selected targets as well as functional validation of GM26870, a long non-coding RNA

To validate RNA-seq results, we performed RT-PCR assay of 3 selected targets: Nampt, Chemokine (C–C motif) ligand 12 (Ccl12), and Vascular cell adhesion protein 1 (Vcam1). In the control group, Nampt expression levels in Nampt+/− mice were 40% of Nampt+/+ mice, while in the CIA group, Nampt expression levels were lower than those in the control group. The trend was similar to the RNA-seq result (Fig. 6a). Ccl2 expression in Nampt+/− mice was 40% of Nampt+/+ mice in the control group, while in the CIA group, Ccl2 expression levels were lower than the control group with Ccl2 expression in Nampt+/− mice approximately 50% of Nampt+/+ mice. The trend was also similar to the RNA-seq result (Fig. 6b). Vcam1 expression in Nampt+/− mice was about 20% higher than Nampt+/+ mice in the control group while in the CIA group, Vcam1 expression levels were all higher than in those in the control group. The trend was also similar to the RNA-seq result (Fig. 6c). Although we only assayed a limited number of targets, the RT-PCR analyses confirmed the RNA-seq results.

To validate the RNA-seq results functionally and to initiate signal transduction analyses of Nampt mediated pathways in CIA Nampt+/− mice, we focused on GM26870, a differentially expressed long non-coding RNA (lncRNA), to examine whether it may be a component underlying the
protective role of Nampt knockdown in arthritis. GM26870 (ENSMUSG00000097312), located on the reverse strand of Chromosome 9: 3,000,282–3,038,313, encodes 3 transcripts (splice variants) (Fig. 7a), which all have a length around 1100 bp. Among them, GM26879-201 is 1170 bp. We employed GM26870-201 dsiRNA synthesized by IDT (Fig. 7b) to knockdown GM26870-201 in mouse primary BMM to examine its effect on osteoclast formation. The knockdown of GM26870-201, successfully measured by RT-PCR (data not shown), blocked osteoclast formation (Fig. 7c) in mouse primary BMM.

Overexpression of Nampt promotes osteoclast formation

Since knockdown of Nampt expression inhibits osteoclast formation, we hypothesized that overexpression of Nampt would promote osteoclast formation. To test this hypothesis, we isolated BMM from Nampt+/+ and NamptOE DBA/1J male mice and initiated cell differentiation into TRAP+ osteoclasts (Fig. 8a). As presented in Fig. 8b, the number of osteoclasts is as high as 450/well from NamptOE mice cells while only about 200/well in Nampt+/+ mice cells. Osteoclast density is above 0.25 in NamptOE mice cells vs. 0.1 in Nampt+/+ mice cells. Overexpression of Nampt in NamptOE mouse cells was confirmed by western blotting (Fig. 8c). Nfatc1 was also upregulated in NamptOE mouse cells. Semi-quantitatively, Nampt and Nfatc1 expressions were 150% and 130% higher in NamptOE cells than those in Nampt+/+ cells, respectively. These results indicate that the overexpression of Nampt promotes osteoclast formation via upregulating Nfatc1.

Discussion

In this study, we discovered that heterozygous knockdown of Nampt suppressed inflammatory bone destruction and disease progression in a CIA mouse model. We characterized one potential mechanism by which Nampt affects arthritis through its transcriptional regulation of the osteoclastogenesis essential transcription factor Nfatc1. Nampt expression was required for the epigenetic regulation of the Nfatc1 promoter and osteoclastogenesis. Finally, we performed transcriptome analysis of whole ankle joints isolated from Nampt+/+ and NamptOE mice which demonstrated the enrichment of osteoclastogenesis genes and pathways and provided insight into the roles of Nampt in the pathogenesis of CIA. Selected targets were validated by RT-PCR for functional characterization. Our findings support our hypothesis that NAMPT is a genetic risk factor and potential therapeutic target for RA.

The role of Nfatc1 in osteoclastogenesis has been well characterized. However, the epigenetic regulation of Nfatc1 transcription by Nampt is a novel finding. Knockdown of Nampt expression inhibits Rankl expression (Fig. 2) while overexpression of Nampt upregulates Rankl expression (Fig. 7). Rankl is an upstream regulator of Nfatc1. RANKL belongs to the tumor necrosis factor superfamaly and plays a critical role in osteoclast differentiation and bone destruction in RA16. Knockdown of Nampt expression inhibits Nfatc1 expression as well as Acp5, Dc-stamp, Cathepsin K mRNA expression which are known to be involved in osteoclastogenesis (Fig. 2). Nfatc1 is required for sufficient osteoclast differentiation. It plays the role of a master transcription regulator of osteoclast differentiation17. The dysregulation of the NAMPT–RANKL–NFATC1–osteoclastogenesis axis may play a major role in bone erosion associated with chronic arthritis, which is underlying the role of NAMPT in the pathogenesis of RA. Two lines of evidence, luciferase reporter assays and RNA capture experiments, support that Nampt regulates the transcription of Nfatc1 (Fig. 3). Our ChIP-PCR analyses in Nampt-deficient RAW 264.7 cells found that the interaction of acetylated histones with the Nfatc1 P1 promoter was decreased (Fig. 3c) while that of methylated histones was increased at the Nfatc1 P1 and
P2 promoter regions. These observations corresponded with decreased HAT activity in RAW 264.7 cells subjected to Nampt knockdown (Fig. 3d). Histone acetylation is correlated with gene expression activation. The presence of methyl moieties inhibits gene expression. Knockdown of Nampt expression inhibits histone acetylation while enhancing histone methylation of the Nfatc1 gene promoter and hence its expression. Our study provides the first evidence that Nampt knockdown inhibits osteoclast formation via epigenetic inhibition of Nfatc1 gene expression. The epigenetic remodeling was consistent with the decreased transcriptional activity observed by luciferase reporter and nascent RNA capture assays (Fig. 3b). Our finding that the formation of Rankl-stimulated TRAP+ cells was blocked by treatment with the small molecule Nampt inhibitors, FK866 or MC4, provides evidence for the requirement of NAD+ in osteoclastogenesis. In addition, the increased inhibitory effect of an MC4 + MTX combination relative to individual treatments of each drug, support our earlier work which demonstrated that the inhibition of NAMPT potentiated the effectiveness of MTX.

A few RNA-seq studies have been applied to rheumatic diseases. In the present study, RNA-seq analysis of mouse whole ankle joints identified 690 genes that are associated with Nampt's role in CIA pathogenesis. They revealed not only “usual suspects” in the pathogenesis such as upregulation of inflammatory activator (TNFα, TGFβ1, and LPS), transcriptional regulators (Fos, Nfatc1), and signaling proteins (Nfkβ, Erk1/2, P38 Mapk, Src, PI3k, Akt)
that are known to promote osteoclast differentiation and bone resorption but also provided a number of new targets and novel insight into the role of Nampt in the pathogenesis of arthritis. First, we found that two osteoclast-specific genes, Oscar and Atp6v0d2, were among the top 5 up-regulated genes in CIA Nampt+/− mice vs. CIA Nampt+/+ mice. OSCAR–collagen interaction stimulates RANK-dependent osteoclastogenesis. OSCAR can play a proinflammatory role in monocyte-derived cells and contribute crucially on multiple levels to RA pathogenesis. The RANK/c-Fos/ATP6V0D2 signaling pathway is an important pathway in the osteoclastogenesis. Second, functional enrichment analyses for GO terms and KEGG to predict potential biological processes and pathways involved in the Nampt associated pathogenesis of CIA has identified many new targets in biological processes: GO:0001501—skeletal system development, GO:001503—ossification, and GO:0060348—bone development (Fig. 5d, Supplemental Table 7). Third, a number of non-coding RNAs was differentially expressed (Supplemental Table 7) and they are novel targets which may underlie Nampt’s role in the pathogenesis of RA. This short discussion just scratches the surface of our RNA-seq data. It should be mentioned that we validated RNA results by RT-PCR of selected targets: Nampt, Ccl2, and Vcan1 (Fig. 6), support the validity of our RNA-seq results. Thus our RNA-seq data provide a rich resource for us and others to further experimentation to characterize new targets in the pathogenesis of RA in the future. We also functionally validated one of the differentially expressed lncRNA, GM26870, and found that knockdown of GM26870 inhibited osteoclast formation (Fig. 8). It may in part be among the Nampt mediated pathways. lncRNAs can function as modular scaffolds to specify higher-order organization in RNP complexes and in chromatin states. It forms extensive networks of ribonucleoprotein (RNP) complexes with numerous chromatin regulators. It is increasingly recognized that lncRNAs play critical roles in multiple biological processes across all kingdoms of life. GM26870’s biological role is thus far unknown. Our study here provides the first glimpse into GM26870’s function or pathological role.

In conclusion, we demonstrated that decreased Nampt expression attenuates inflammatory bone loss in a Nampt+/− CIA mouse model. In vitro assays revealed impaired osteoclastogenesis in Nampt-deficient RAW 264.7 and BMM which corresponded with epigenetic suppression of Nfatc1 transcription and may provide a potential mechanism by which the Nampt–NfatC1–osteoclastogenesis pathway promotes arthritis. RNA-seq analysis further supported these observations and uncovered new insights into the pathways associated with arthritis. In total, our findings suggest that NAMPT is a genetic risk factor and potential therapeutic target for RA.
Materials and methods

Antibodies and chemicals

RPMI 1640 and DMEM were purchased from Life Technologies. Lipopolysaccharide-\textit{Escherichia coli} 055: B5 was obtained from Sigma Aldrich (#L56529; St. Louis, MO). TRACP and ALP double staining kit (#MK300) was purchased from Clontech (Mountain View, CA). Anti-NFATc1 antibody (#ab151239) was from Abcam (Cambridge, MA). Phospho-MAPK family antibody sampler kit (#9910), pNF-kB p105 (#4806), pNF-kBp65 (#3033), Acetyl-Histone H3 (Lys9) (#9649), Tri-Methyl-Histone H3 (Lys27) (#9733) antibodies, Simple ChIP Enzymatic Chromatin IP kit (#9003), and cell lysis buffer (#9803) were purchased from Cell Signaling Technology (Beverly, MA). Recombinant Mouse M-CSF (#576406) and purified anti-NFATc1 antibody (#649601) were purchased from Biolegend (San Diego, CA). GAPDH (#sc25778), iRISC lab as described previously33.

Cell culture

The murine macrophage cell line RAW 264.7 (TIB-71, ATCC®) was maintained in RPMI 1640 media containing 10% FBS and antibiotics. All cells were grown at 37 °C, 5% CO₂.

Animal studies

Mice were maintained in an Association for Assessment and Accreditation of Laboratory Animal Care accredited institution in a temperature-controlled, pathogen-free facility with a 12 h light/dark cycle. The mice were gang housed with free access to food (Irradiated Teklad Global 18% Rodent Diet, Envigo, cat# 2918.15) and water. DBA/1J mice (The Jackson Laboratory) were crossed with C57BL/6 Nampt+/− mice33. Offspring were backcrossed to DBA/1J for >10 generations to develop congenic DBA/1J Nampt+/−. To induce arthritis, 10-week-old, male mice were immunized with bovine collagen type II (CII, 100 μg, Chondrex) in Complete Freund’s Adjuvant (CFA, 100 μg \textit{Mycobacterium tuberculosis}, Chondrex) by intradermal injection at the base of the tail. A collagen booster (100 μg) in Incomplete Freund’s Adjuvant (IFA, Chondrex) was administered at day 218. Mice were evaluated for the onset of inflammation and scored (scale 0–4 per paw) as described by Brand et al.8. Mice were scored twice per week for 10 weeks. At the end of the 10-week period, the animals were euthanized and tissue was isolated for analysis. Circulating serum levels of anti-mouse CIA antibody (Chondrex) and Nampt (AdipoGen®) were determined by ELISA. The left hind limb was scanned with a vivaCT40 (Scanco) in the Skeletal Imaging and In vitro-In vivo Mechanical Core in the University of Missouri Kansas City, School of Dentistry, as described previously32. The right hind limb was collected for RNA isolation and gene expression analyses.

Overexpression (Nampt<sup>OE</sup>) mice were generated in our lab as described previously33.

Isolation of bone marrow-derived macrophage

Isolation of BMM was performed as previously described34.

In vitro osteoclastogenesis

Osteoclasts were generated from bone marrow as described previously35.

siRNA mediated Nampt knockdown

RAW 264.7 cells were 4D-nucleofected with Nampt and scrambled siRNA (50 nM/2 × 10⁶ cells) in SF solution (Amaxa). Cells were seeded at 5 × 10⁶ nucleofected cells/well/24-well plate for TRAP staining and 1 × 10⁵ cells/well/6-well plate for protein isolation.

Protein extraction and western blot analysis

Protein extraction and western blot analysis were carried according to our previous procedure33.

ChIP assay

SimpleChIP® Enzymatic Chromatin IP assays (Cell Signaling Technology) with Acetyl-Histone H3 (Lys9) and Tri-Methyl-Histone H3 (Lys27) antibodies were utilized according to the manufacturer’s instruction. Immunoprecipitated DNA was reverse cross-linked, purified and analyzed by PCR (primers: NFATc1-617-F 5′-ACGAAACGGGAAGGAAAG-3′, NFATc1-426-R 5′-GGAAGCCTGCGATTTTACAT-3′, NFATc1-125-F 5′-GGGACGCGTTCGCTTATCCCAT-3′, NFATc1-617-F 5′-ACGAAACGGGAAGGAAAG-3′). svv

Histone acetyltransferase (HAT) assay

HAT enzyme activity was quantified in Nampt-deficient and scrambled control cells using an EpiQuik HAT assay (EpiGentek) according to the manufacturer’s instruction.
Luciferase reporter assays

Luciferase reporter assays were performed as described previously.34

RNA isolation, quantitative RT-PCR, and nascent RNA capture

RNA isolation, quantitative RT-PCR, and nascent RNA capture were performed as we previously described.36

RNA-seq

RNA was isolated from flash frozen whole ankle joints collected after the 10-week CIA observation period. cDNA sequencing libraries were prepared with an Illumina TruSeq Stranded Total RNA Sample Prep Kit and subjected to 2 × 101 paired-end sequencing as described previously3. Mapping of RNA-seq reads and transcript assembly and abundance estimation were conducted using Tuxedo Suite pipeline (TopHat v1.3.0/Bowtie assembly and abundance estimation were conducted using Tuxedo Suite pipeline (TopHat v1.3.0/Bowtie v0.12.7/Cufflinks v1.0.3) and reported in Fragments Per Kilobase of exon per Million fragments mapped (FPKM). To identify genes which were differentially expressed, fold changes for each gene were calculated by dividing the average FPKM for the case by the average FPKM for the control. We determined fold changes for two comparison groups: (1) CIA Nampt+/+/ mice compared with non-CIA control Nampt+/+ mice, and (2) CIA Nampt+/+ mice compared with CIA Nampt+/− mice.

Functional pathway analysis

Kyoto Encyclopedia of Genes and Genomes (KEGG) analyses to identify cellular pathways and biological processes associated with differentially expressed genes (DEG) were performed with DAVID37. Ingenuity Pathway Analysis (IPA) (Ingenuity Systems) predicted functional and canonical pathways.

Statistics

Statistical analyses were performed with Sigma Stat (v4.0, Systat Software, Inc.). Results were expressed as mean ± SD. P < 0.05 was considered statistically significant.

Data availability

The RNA-seq data have been deposited to Gene Expression Omnibus (http://www.ncbi.nlm.nih.gov/geo; accession number GSE121793).

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Author contributions

D.P.H., L.G. and S.Q.Y conceptualized and designed the research. X.L., S.I., D.P.H. and S.Q.Y. wrote the manuscript. X.L., S.I., M.X., N.N.N and D.P.H. acquired data. X.L., S.I., D.P.H., L.G. and S.Q.Y. analyzed data. M.W.L. and Y.U. provided samples, reagents, and other project assistance. All authors critically revised the manuscript. D.P.H., L.G. and S.Q.Y. provided overall project guidance and supervision.

Conflict of interest

The authors declare that they have no conflict of interest.

Study approval

Animal studies were approved by the University of Missouri Kansas City’s Institutional Animal Care and Use Committee.

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