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NAIL: an evolutionarily conserved lncRNA essential for licensing coordinated activation of p38 and NFκB in colitis

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ABSTRACT

Objective NFκB is the key modulator in inflammatory disorders. However, the key regulators that activate, fine-tune or shut off NFκB activity in inflammatory conditions are poorly understood. In this study, we aim to investigate the roles that NFκB-specific long non-coding RNAs (lncRNAs) play in regulating inflammatory networks.

Design Using the first genetic-screen to identify NFκB-specific lncRNAs, we performed RNA-seq from the *p65*^{-/-} and *Ikkβ*^{-/-} mouse embryonic fibroblasts and report the identification of an evolutionary conserved lncRNA designated *mNAIL* (mice) or *hNAIL* (human). *hNAIL* is upregulated in human inflammatory disorders, including UC. We generated *mNAIL*^{ΔNFκB} mice, wherein deletion of two NFκB sites in the proximal promoter of *mNAIL* abolishes its induction, to study its function in colitis.

Results *NAIL* regulates inflammation via sequestering and inactivating Wip1, a known negative regulator of proinflammatory p38 kinase and NFκB subunit p65. Wip1 inactivation leads to coordinated activation of p38 and covalent modifications of NFκB, essential for its genome-wide occupancy on specific targets. *NAIL* enables an orchestrated response for p38 and NFκB coactivation that leads to differentiation of precursor cells into immature myeloid cells in bone marrow, recruitment of macrophages to inflamed area and expression of inflammatory genes in colitis.

Conclusion *NAIL* directly regulates initiation and progression of colitis and its expression is highly correlated with NFκB activity which makes it a perfect candidate to serve as a biomarker and a therapeutic target for IBD and other inflammation-associated diseases.

INTRODUCTION

Chronic inflammatory diseases such as IBD stem from dysregulated immune responses, aberrant cytokine/chemokine secretion and alterations in intestinal barrier and microbiota.¹ In IBD, intestinal microbiota releases enterotoxins that increase the permeability of intestinal mucosa, resulting in a state of intestinal inflammation caused by invasion of harmful bacteria and damaged intestinal epithelial cells.² In UC patients, NFκB is activated in epithelial cells and mucosal macrophages that enhances the secretion of a broad panel of NFκB-regulated cytokines including TNFα, IL-1 and

Significance of this study

What is already known on this subject?

- NFκB is highly activated in IBD patients and associated with disease severity.
- NFκB and p38 have been shown to be required for the expression of proinflammatory genes for full-blown inflammation.
- NFκB activity can be shut off by dephosphorylation of active p65 by the Wip1 phosphatase leading to loss of its chromatin remodelling potential and consequently reduced transcription of NFκB targets.
- A number of signalling regulators, including enzymes, adaptor proteins and non-coding RNAs such as miRNAs which regulate NFκB and p38 have been identified and characterised.
- Several lncRNAs have been proposed to be induced on TNFα stimulation and dysregulated in human inflammatory diseases.
- lncRNAs can interact with kinases and regulate their activity.

What are the new findings?

- *NAIL* is the first lncRNA to be identified by a genetic screen, and its expression is absolutely NFκB dependent. This autoregulation is unique and not documented before.
- *NAIL* forms a positive feedback loop with NFκB signalling, and its expression is enhanced in colitis.
- *NAIL* regulates the development of colitis by regulating the expression of inflammatory cytokines and recruitment of effector cells to the site of inflammation.
- *NAIL* simultaneously activates p38, a kinase and p65 a transcription factor. Such direct coordinated activation of two pathways has not been documented for any other lncRNA.
- *NAIL* deactivates a phosphatase (Wip1) essential to activate NFκB and p38 signalling for full-blown inflammation in colitis.

IL-6.³ Due to recurrent cycles of evoked inflammatory activation, IBD patients have been shown to be more susceptible to cytokine storm syndrome⁴ and cancer development. This is due to the hyperactive NFκB-dependent cytokine secretion which induces

Significance of this study

How might it impact on clinical practice in the foreseeable future?

- The critical role of NAIL in Wip1-NFκB/p38 axis in the development of colitis serves as an attractive target for the development of future therapies against inflammatory diseases such as UC or human cancers such as colitis-associated colorectal cancer

hyper proliferation of immune cells and excessive tissue damage, resulting in an accumulation of new cancer mutations.^{5,6}

There are many regulators of NFκB, including kinases, ubiquitin ligases and miRNAs. Some of these have been identified as potential therapeutic targets.⁷ However, only ~1%–2% of the human genome encodes proteins, while 75%–90% is transcribed into non-coding RNAs;⁸ it is plausible that the key regulators that activate, fine-tune or shut off NFκB activity have yet to be discovered. Long non-coding RNAs (lncRNAs) constitute the majority of this rampant transcription of the mammalian genome.^{8,9}

To identify lncRNAs that are directly and specifically regulated by NFκB, we performed RNA-seq from wild type (WT), *p65*^{-/-} and *Ikkβ*^{-/-} mouse embryonic fibroblasts (MEFs) treated with TNFα for different time points (figure 1A,B). This screen revealed a novel, evolutionarily conserved lncRNA, Gm16685 in mouse and its human homolog (loc105375914), both of which are prominently and specifically activated by the p65 subunit of NFκB. We named this unannotated lncRNA as NFκB Associated Immunoregulatory Long non-coding RNA, NAIL. *hNAIL* is upregulated in many human inflammatory disorders including UC. To model NAIL's function, we deleted the two conserved NFκB sites in the proximal mouse NAIL (*mNAIL*) promoter and generated a mouse model (*mNAIL*^{ΔNFκB}) in which NAIL gene is unresponsive to NFκB-dependent activation. *mNAIL*^{ΔNFκB} mice displayed decreased expression of NFκB targets and consequently had reduced colitis. On inflammatory stimulation, NAIL is expressed, and sequester Wip1 phosphatase and prevent Wip1 from dephosphorylating, and deactivating active p65 and other Wip1 targets such as p38 necessary for full-blown inflammatory gene expression. These results provide the first evidence that lncRNAs can directly modulate phosphatase accessibility to its substrates which has important consequences in inflammation.

RESULTS**Identification and characterisation of Gm16685 (*mNAIL*); a novel lncRNA specifically regulated by NFκB signaling**

Besides activating the NFκB pathway, TNFα receptor engagement also leads to activation of JNK, p38 and apoptotic pathways (figure 1A). Since NFκB activation is specifically impaired in *p65*^{-/-} and *Ikkβ*^{-/-} MEFs (online supplemental figure 1A,B) downstream of TNF signalling, we employed these cells to identify NFκB-driven lncRNAs. Differentially expressed lncRNAs between WT, *p65*^{-/-} and *Ikkβ*^{-/-} MEFs on TNFα stimulation were identified (figure 1B). To identify lncRNAs that were directly and specifically regulated by NFκB signalling, we selected the lncRNAs that were TNFα inducible in a time-dependent manner in WT but not in *p65*^{-/-} and *Ikkβ*^{-/-} MEFs. We selected Gm16685, a novel lncRNA expressed in the antisense direction of interleukin-7 (*IL7*), for further functional and mechanistic studies (figure 1C). Gm16685 expression was dramatically upregulated in WT MEFs stimulated with TNFα (figure 1D). However,

compared with WT MEFs, Gm16685 expression in *p65*^{-/-}, *Ikkβ*^{-/-} and *Ikkγ*^{-/-} immortalised MEFs was significantly dampened on TNFα stimulation (online supplemental figure 1C,D). Primary MEFs derived from WT, *p65*^{-/-} and *Ikkβ*^{-/-} mice also showed that activation of Gm16685 occurred specifically in an *Ikkβ*-dependent and p65-dependent manner (figure 1E). Gm16685 expression was also induced by stimulation with TLR4 ligand, lipopolysaccharides (LPS) in WT immortalised MEFs (online supplemental figure 1E). Importantly, the antisense gene *IL7* remained relatively unchanged on stimulation (online supplemental figure 1F), suggesting the specificity of induction of Gm16685 by p65 via *Ikkβ*. Taken together, these results suggest that Gm16685 expression is specifically activated by NFκB signalling, downstream of major inflammatory stimuli such as TNFα and LPS.

loc105375914 (*hNAIL*), an evolutionarily conserved human homolog of Gm16685 (*mNAIL*) is upregulated in UC

Determining which lncRNAs are evolutionarily conserved highlights the functional importance of such transcripts in fundamental biological processes.¹⁰ To identify human orthologue of Gm16685 (*mNAIL*), we converted genome coordinates of Gm16685 (chr3:7612705–7690001) in mouse (mm10) to human (hg38) genome coordinates by using UCSC liftOver with default parameters. Human lncRNA, loc105375914 (*hNAIL*) (chr8:78804578–78937681) overlapped with the converted region (chr8:78804472–78902467), hence, loc105375914 was considered as candidate orthologue of mouse Gm16685. Although lncRNAs have poor sequence conservation, their promoter regions are generally more conserved.¹¹ Hence, we performed multiple sequence alignment using promoter sequence of the mouse Gm16685 and sequences of 31 other species at the same region with reference to UCSC comparative genomic track (online supplemental figure 2A). Promoter motifs enrichment analysis identified NFκB as the most significantly enriched motif across all orthologues (online supplemental figure 2B). Most importantly, the RNA binding protein (RBP) binding motifs predicted using RBPmap server (<http://rbpmap.technion.ac.il/>) showed that >80% of RBPs predicted to bind Gm16685 are also predicted to bind the human loc105375914 lncRNA (online supplemental figure 2C) suggesting that *hNAIL* and *mNAIL* are most likely functionally conserved.

Analyses of publicly available p65 (RelA) ChIP-seq data^{12–13} from mouse dendritic cells, macrophages stimulated with LPS and human adipocytes and fibroblasts stimulated with TNFα demonstrated that p65 binds to the promoter regions of Gm16685 and loc105375914 in a stimulus-dependent manner (online supplemental figure 2D,E). Additionally, RNA Polymerase 2 (Pol2) ChIP-seq data^{14–15} from these human adipocytes and fibroblasts revealed that loc105375914 expression is driven by RNA Pol2 in conjunction with NFκB (online supplemental figure 2E). Two p65 (RelA) binding motifs located upstream of Gm16685 are highly conserved across species, suggesting that these motifs might be functionally important for regulation and subsequent transcription of Gm16685 across species. Furthermore, NAIL does not have coding potential as shown by computational analyses (online supplemental figure 2F) and 3' Flag tagging strategy (online supplemental figure 2G).

Hyperactivation of NFκB is a functional determinant in the development of inflammatory diseases such as UC that causes inflammation in the colonic mucosa.¹⁶ Since loc105375914 is transcriptionally regulated by NFκB, we analysed transcriptomes of 24 inflamed and matched non-inflamed sections from human

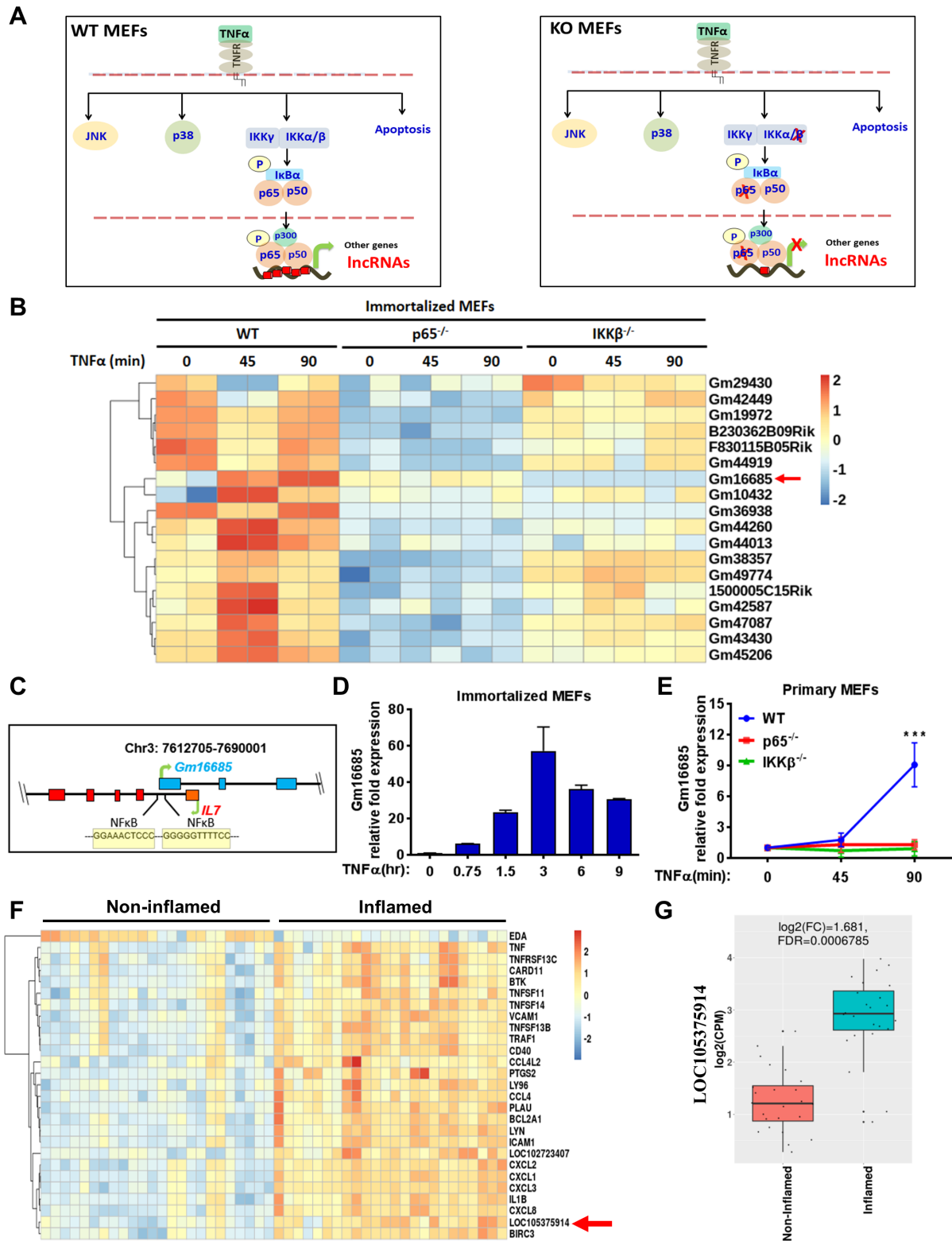


Figure 1 Identification of lncRNAs regulated by NF κ B signalling pathway. (A) NF κ B signalling pathway and screening strategy used to identify NF κ B-regulated lncRNAs. Crosses represent loss of the corresponding proteins (Ik κ β, Ik κ γ, p65) using KO MEFs. (B) Heatmap of logCPM values of differentially expressed lncRNAs in WT, *p65*^{-/-} and *Ik κ β*^{-/-} in immortalised MEFs exposed to TNF α for the indicated time points and analysed by paired-end RNA-sequencing. Gm16685 is shown with red arrow. (C) Schematic view of genetic loci of Gm16685 lncRNA (blue). Gm16685 is expressed from antisense direction to *IL7* gene (red). NF κ B binding motifs in the promoter region of Gm16685 shown as yellow highlighted sequences. (D) RT-qPCR analysis of Gm16685 transcript levels in immortalised WT MEFs exposed to TNF α for different time points as indicated. (E) RT-qPCR analysis of Gm16685 in WT (n=5), *p65*^{-/-} (n=8) and *Ik κ β*^{-/-} (n=3) independent primary MEFs treated with TNF α for indicated time courses. P values were calculated using Student's t-test method (***) p<0.001). (F) Gene expression profile of 24 inflamed and patient match non-inflamed samples. Differentially expressed genes are shown in heatmap. LOC105375914 is shown with red arrow. (G) Box plot shows upregulation of *hNAIL* (LOC105375914) expression in inflamed colon tissue of UC patients. KO, knock out; MEFs, mouse embryonic fibroblasts; WT, wild type.

UC patients. In inflamed colons, we observed dramatic upregulation of NFκB target genes including *TNFα* and *IL1β* (figure 1F) which are known to be key regulators of intestinal inflammation.¹⁷ As compared with non-inflamed controls, loc105375914 expression is significantly upregulated in the inflamed colons (figure 1G), suggesting that loc105375914 expression highly correlates with NFκB activity and may serve as a biomarker for IBDs. Collectively, these results indicate that loc105375914 is a conserved human homolog of Gm16685, and both their expressions are specifically driven by NFκB signalling in response to inflammatory and microbial triggers across species and cell types. Hereafter, we will refer to Gm16685 as *mNAIL* and loc105375914 as *hNAIL* (NFκB Associated Immunoregulatory Long non-coding RNA).

mNAIL^{ΔNFκB} mice display loss of *mNAIL* activation by NFκB and dampened inflammation

We generated *mNAIL*^{ΔNFκB} mice wherein two conserved NFκB binding motifs upstream of *mNAIL* were removed using clustered regularly interspaced short palindromic repeats (CRISPR)-Cas9 genome editing (figure 2A). *mNAIL*^{ΔNFκB} mice were born alive in normal Mendelian ratios and had no overt phenotypes (figure 2B). Deletion of the NFκB motifs was confirmed by genotyping PCR (figure 2C) and Sanger sequencing (figure 2D). Three independent *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} primary MEF clones showed that *mNAIL* expression was abolished in the *mNAIL*^{ΔNFκB} clones, whereas the expression level of *IL7* gene was unaffected on TNFα treatment (figure 2E,F). We also observed decreased expression of NFκB targets *TNFα* and *IL1β* levels on TNFα treatment in *mNAIL*^{ΔNFκB} MEFs compared with *mNAIL*^{WT} MEFs (figure 2G,H). Depletion of *mNAIL* by two independent siRNAs showed reduced NFκB target gene expression (figure 2I,J) and significant reduction in phosphorylation of p65 and p38 (figure 2K). To exclude the possibility that the observed phenotypes might stem from genome editing and its off target effects on *IL7* or it neighbouring genes, we first analysed the expression of *IL7* across different tissues in mice using NCBI fragments per kilobase of exon per million reads (FPKM) data. Since *IL7* is highly expressed in thymus, spleen and colon tissues (online supplemental figure 3A), we analysed its expression in thymus between *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} groups and observed no significant difference in RNA and protein levels (online supplemental figure 3B,C). Although we did not observe any disruption of *IL7* expression (RNA and protein level), we knocked-down *IL7* gene using siRNA to investigate the impact of *IL7* on NFκB signalling. Depletion of *IL7* did not affect NFκB activation as judged by expression of TNFα (online supplemental figure 3D–F). Taken together, these results suggest that editing the two conserved NFκB sites in the *NAIL* promoter specifically dampens inducibility of *NAIL* in response to inflammatory stimuli and that *NAIL* in turn regulates NFκB target genes in vivo.

mNAIL^{ΔNFκB} mice display reduced colitis and colon inflammation

mNAIL^{ΔNFκB} mice display reduced colitis and colon inflammation *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice were given 3% dextran sulfate sodium (DSS) to induce acute colitis (figure 3A). Initial body weights of *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice were similar, however, on DSS treatment, *mNAIL*^{ΔNFκB} mice displayed less severe body weight loss when compared with *mNAIL*^{WT} counterparts (figure 3A). On DSS treatment, colon shortening was significantly less pronounced in *mNAIL*^{ΔNFκB} mice and the disease

activity index (DAI) was lower when compared with *mNAIL*^{WT} counterparts (figure 3B–D). These results were suggestive of reduced inflammation in *mNAIL*^{ΔNFκB} mice.

DSS treatment has been shown to alter the myeloid precursor population in the bone marrow which directly correlates with the degree of inflammation in the colon.^{18–19} Especially, monocytes derived from the CD11b+Ly6C^{hi}Ly6G[–] immature myeloid cells have been shown to enter tissues under inflammatory conditions and express high levels of proinflammatory cytokines in colitis.^{20–22} Given that myeloid cells are the predominant cells that infiltrate the inflamed tissues and the precursors are made in bone marrow,²³ we analysed the myeloid populations in the bone marrow of *mNAIL*^{ΔNFκB} mice. As expected, both *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice showed increased CD11b+ population, on DSS treatment (figure 3E–G). Next, we analysed CD11b positive subpopulations CD11b+Ly6C^{hi}+Ly6G[–] and CD11b+Ly6C^{low}+Ly6G⁺ in bone marrow (figure 3F). DSS induced increase in Ly6C^{hi} activator cell population was significantly more pronounced in *mNAIL*^{WT} mice (figure 3H). Conversely, compared with *mNAIL*^{WT} mice, elevated levels of Ly6C^{low}Ly6G⁺ cells were seen in *mNAIL*^{ΔNFκB} mice (figure 3I). To further investigate if *NAIL* has any role on the stem cell population in the bone marrow, we analysed the progenitor cells in the bone marrows of *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice treated with or without DSS²⁴ (online supplemental figure 4A). Cells were initially gated as LIN[–] cells and based on the c-KIT and Sca-1 expression, the long-term and short-term hematopoietic stem cell populations (KL, KSL and K-low, S-low) were selected. KL cells were further gated based on their CD16/32 and CD34 expression to identify megakaryocyte–erythrocyte, common myeloid progenitor and granulocyte-macrophage progenitors. To identify the common lymphoid progenitors population, we further gated K-low, S-low cells based on their IL7ra expression. Representative FACS data and quantification of all samples (online supplemental figure 4A–E) are shown for *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice. No difference was observed in the stem cells and precursor cells population between *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice suggesting that *NAIL* does not alter the stem cell and progenitor populations. Instead, our data suggest that *mNAIL* is a key determinant for the differentiation of precursor cells into CD11b+Ly6C or Ly6G sub-types during inflammation.

mNAIL expression is functionally regulated in bone marrow-derived macrophages (BMDM)

To identify the major site of *mNAIL* expression in vivo, we dissected the response of *mNAIL* in different cell types and tissues including thymus, liver, spleen, lung and BMDM using the systemic LPS-induced endotoxemia model (figure 4A–E). We observed mild induction of *mNAIL* in thymus and liver (figure 4A,B) but not in spleen and lung tissues (figure 4C,D). Strong induction of *mNAIL* in BMDMs was observed (figure 4E,F). We observed dampened induction of NFκB targets *TNFα*, *CCL2* and *CXCL2* on LPS challenge in *mNAIL*^{ΔNFκB} BMDM (figure 4G–J). ChIP-qPCR results showed that the absence of *mNAIL* resulted in dramatic reduction of p65 occupancy at the promoters of these well-known NFκB target genes (figure 4K,L) suggesting that loss of *NAIL* impacts DNA binding and hence transcription by NFκB. We observed significant reduction in phospho-p65 and phospho-p38 levels in the absence of *mNAIL* in the BMDMs (figure 4M). We did not observe any change for an irrelevant protein phospho-MKK4 suggesting the specificity of *NAIL* acting on the phosphorylation of p65 and p38 (figure 4M). Crucially, myeloid lineage specific knock outs

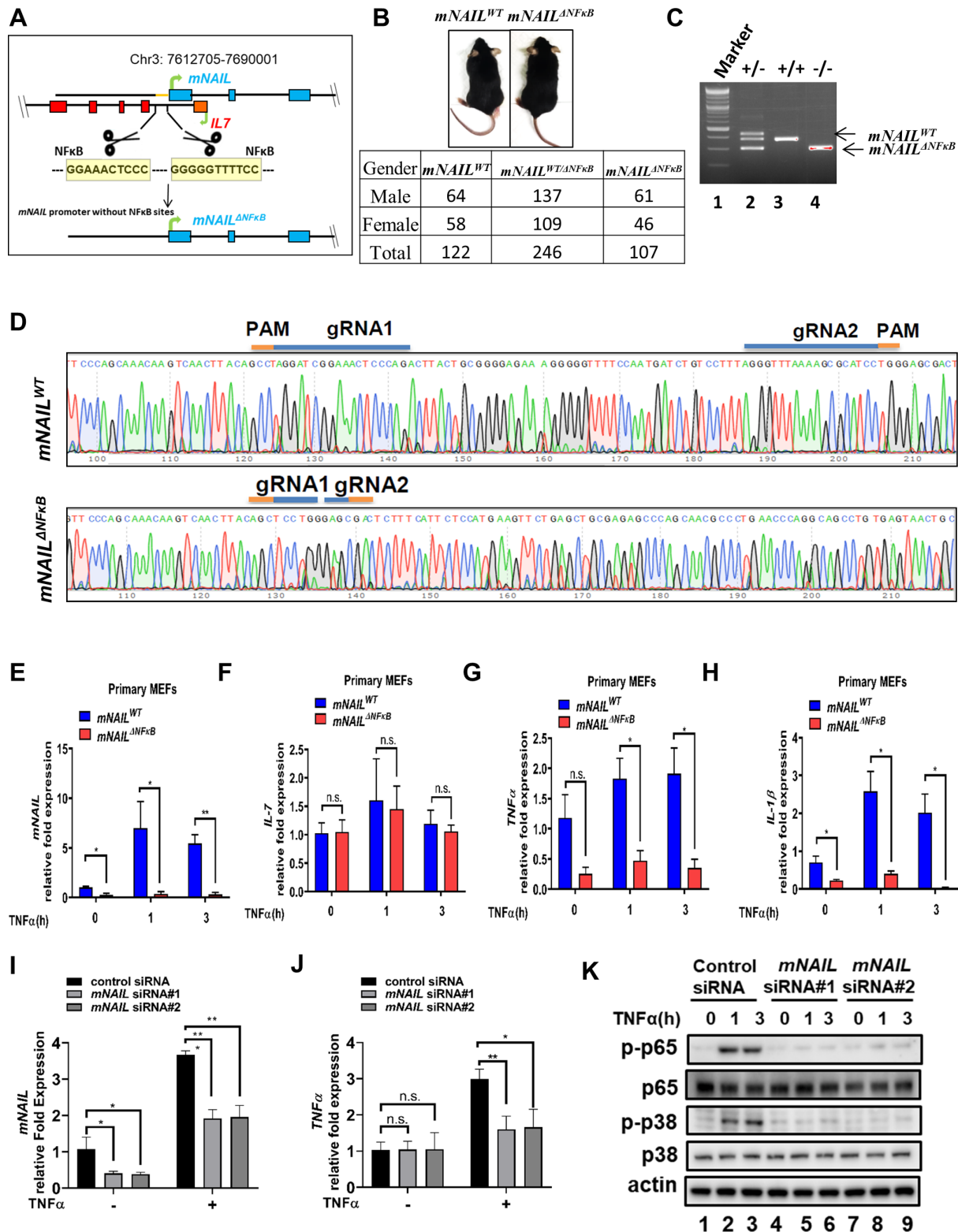


Figure 2 *mNAIL*^{ΔNFκB} mice display decreased activation of NFκB and inflammation. (A) Schematic view of *mNAIL* lncRNA promoter targeting with CRISPR-Cas9 editing. NFκB binding motifs are shown in yellow in the promoter region of *mNAIL*. (B) *mNAIL* mice were generated using standard methods (refer to 'Materials and methods' section). *mNAIL* mice were born in normal Mendelian ratio (table below) and unchallenged mice appear normal. (C) Genotyping PCR results of *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice. PCR products from *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} alleles are shown. (D) Sanger sequencing results of *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} cells. (E–H) *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} MEFs (n=3) were treated with TNFα for the indicated time points. Expression analysis was performed by RT-qPCR for (E) *mNAIL*, (F) *IL7*, (G) *TNFα* and (H) *IL1β* genes. (I–K) WT MEFs were transfected with si-Control, si-NAIL #1 and si-NAIL #2 siRNAs. After 48 hours post-transfection, cells were treated with or without TNFα and harvested for gene expression analysis. Graphs show the gene expression analysis of (I) *mNAIL* and (J) *TNFα* by RT-qPCR. Actin was used as a control. Error bars indicate mean±SD of three independent experiments. P values were calculated using Student's t-test method (*p<0.05; **p<0.01; n.s., not significant). (K) Cells lysate were analysed with western blot for the indicated proteins. MEFs, mouse embryonic fibroblasts; WT, wild type.

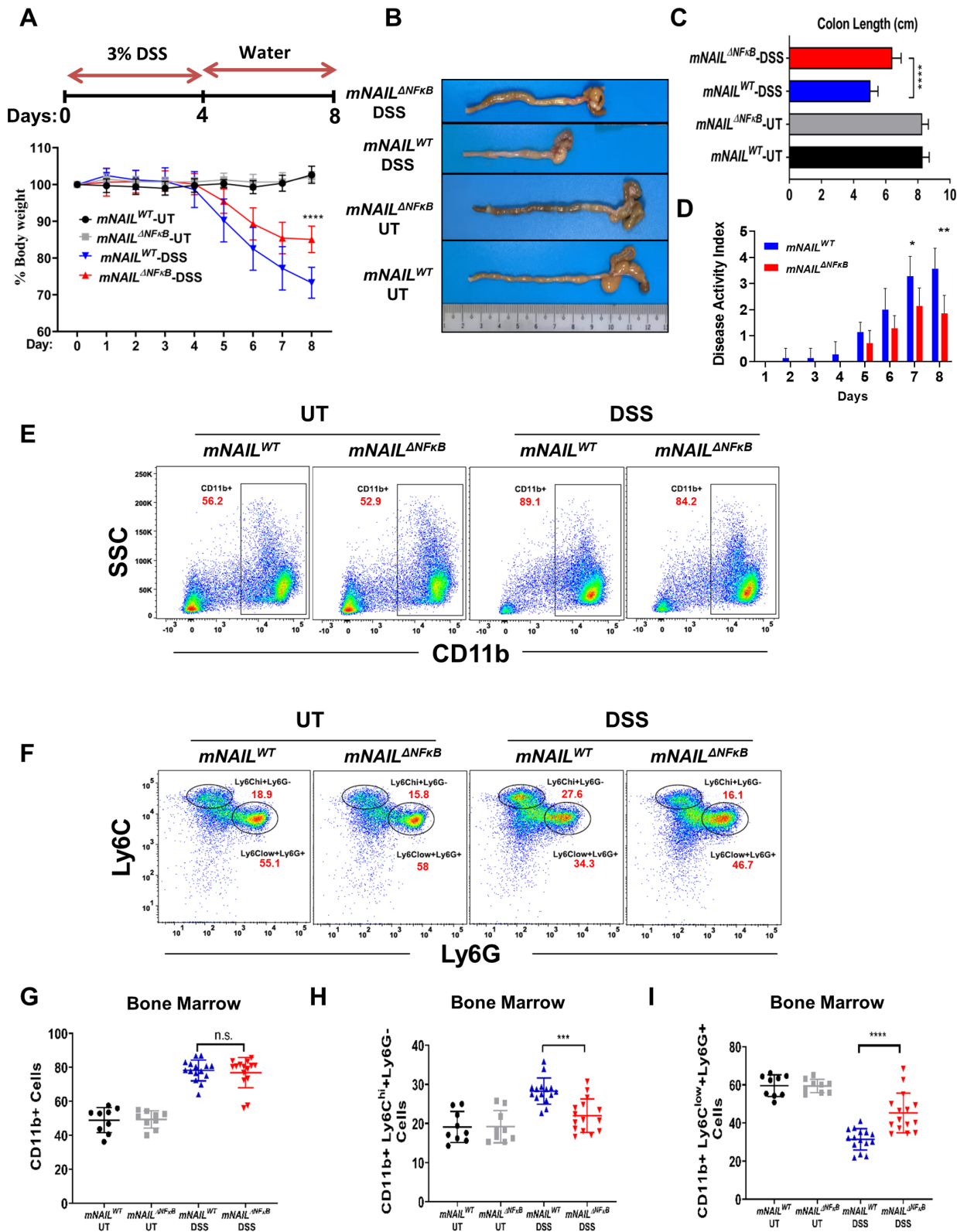


Figure 3 *mNAIL^{ΔNFκB}* mice are resistant to dextran sulfate sodium (DSS)-induced colitis. (A) Scheme of DSS-induced colitis model is shown on the top panel. *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* mice were treated as shown in top panel. Body weight measurements from mice monitored for 8 days (bottom panel). (B and C) Colon length of DSS-treated *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* mice measured at day 8. (D) Disease activity index was recorded daily. (E) Bone marrow cells were isolated from the DSS-treated *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* mice at day 8 and were stained for CD11b (PE), Ly6c (BV711) and Ly6g (FITC) cell surface markers. Cells were analysed by FACS and gated as CD11b⁺-cells. Representative FACS data were shown for *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* mice treated with or without DSS. (F) CD11b⁺-cells were further gated for Ly6c and Ly6g markers. (G–I) Quantification of CD11b⁺, CD11b⁺Ly6C^{hi}+Ly6G⁻, CD11b⁺Ly6C^{low}+Ly6G⁺-cells gated in (E) and (F). Error bars indicate mean±SD of three independent experiments (UT: n=9, DSS: n=15). P values were calculated using Student's t-test method (*p<0.05; **p<0.01; ***p<0.001; ****p<0.0001; n.s., not significant). DSS, dextran sulfate sodium; FACS, fluorescence-activated cell sorting; SSC, side scatter; side scatter; UT, untreated; WT, wild type.

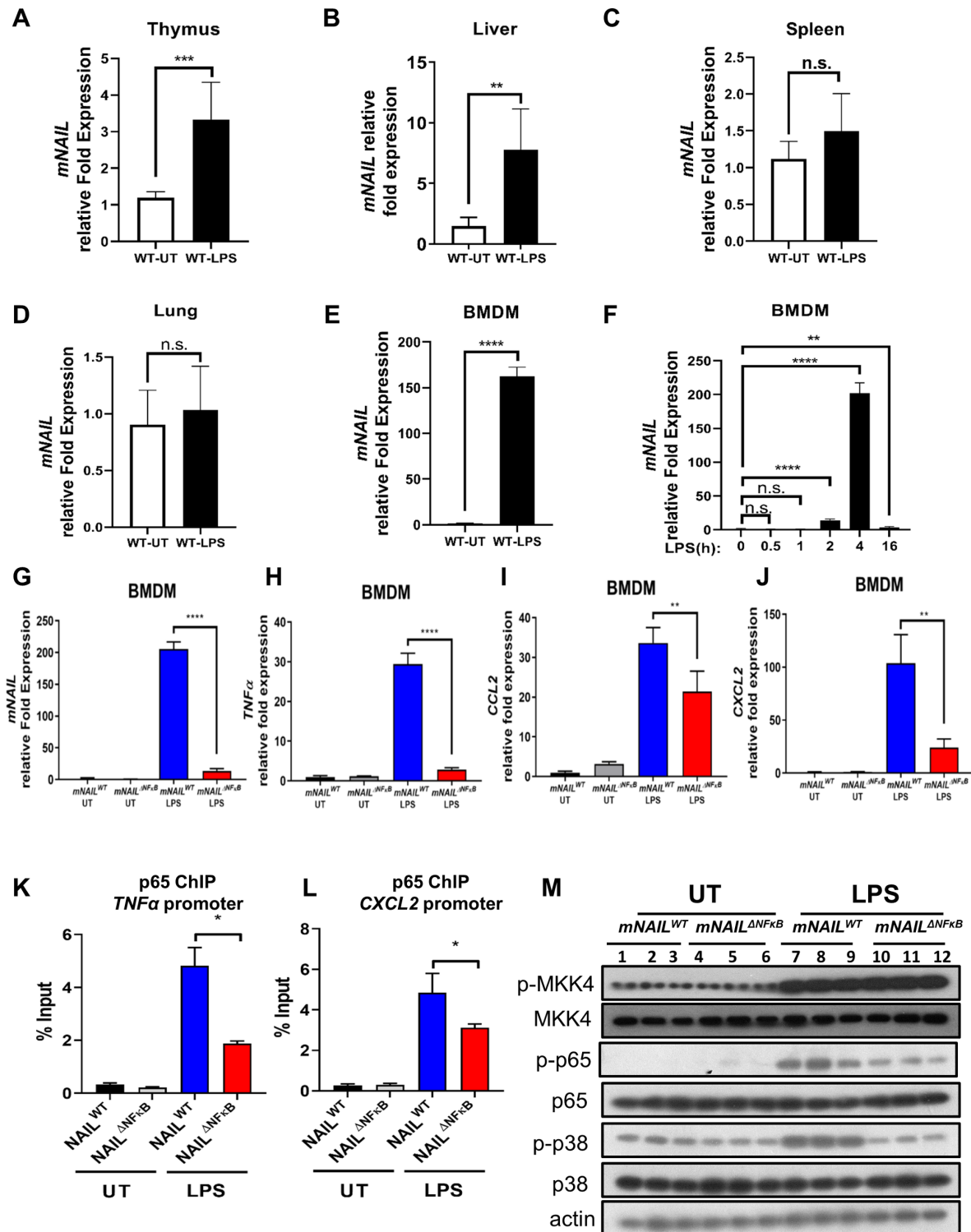


Figure 4 *mNAIL* is expressed mainly in myeloid cells and its expression amplifies activation of NFκB and inflammation. (A–E) LPS (5 mg/kg) was injected intraperitoneally into C57BL/6N WT mice (n=6). (A) Thymus, (B) liver, (C) spleen and (D) lung tissues were collected after 4 hours and expression analysis was performed by RT-qPCR for *mNAIL* gene. (E, F) Bone marrow cells isolated from WT mice (n=6) and differentiated into bone marrow-derived macrophages (BMDM) for 7 days. BMDM cells were treated with or without LPS (200 ng/mL) for 4 hours. (E) Graph shows the *mNAIL* expression analysis by qPCR. (F) Time kinetics of *mNAIL* expression is shown in LPS-stimulated BMDM cells. Data were normalised to actin. (G–J) Bone marrow cells isolated from *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice (n=6) and differentiated into BMDM for 7 days. BMDM cells were treated with or without LPS (200 ng/mL) for 4 hours. Graph shows the gene expression analysis of (G) *mNAIL*, (H) *TNFα*, (I) *CCL2* and (J) *CXCL2* by qPCR. Data were normalised to actin. (K–L) *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} BMDM cells were treated with or without LPS, and ChIP analysis was performed for p65 (n=3). Graph shows the ChIP-qPCR analysis of p65 occupancy on the (K) *TNFα* and (L) *CXCL2* promoters. Error bars indicate mean±SD of three biological replicates. P values were calculated using Student's t-test method (*p<0.05; **p<0.01; ***p<0.001; ****p<0.0001; n.s., not significant). (M) Western blot shows the total and phosphorylated p65, p38 and MKK4 in BMDM cells treated with or without LPS. Each replicate is labelled as #1, #2 and #3. UT, untreated; WT, wild type.

(KO) of p65 and p38 display reduced inflammatory responses and body weight differences in DSS model, highly resembling the observations in *mNAIL*^{ΔNFκB} mice.^{23 25 26} To determine the molecular basis of this function of *NAIL*, we performed RNA-seq using colon tissues of *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice with or without DSS treatment (figure 5A). Only a few genes were differentially expressed in the untreated (UT) group, reiterating that the deletion of two NFκB sites in the proximal promoter of *mNAIL* does not cause any global alterations in the transcriptome due to genome editing artefacts (online supplemental figure 5). However, DSS treatment caused vast changes in gene expression, with a majority of genes down-regulated in *mNAIL*^{ΔNFκB} mice (figure 5A). Gene ontology analysis revealed that the majority of differentially expressed and downregulated genes in *mNAIL*^{ΔNFκB} colons belong to inflammatory pathways regulated by NFκB (online supplemental figure 5B). To validate the RNA-seq results, we performed gene expression analysis of NFκB target genes and observed reduced expression of *TNFA*, *IL1β* and *CCL2* genes in colon tissues of *mNAIL*^{ΔNFκB} mice as compared with *mNAIL*^{WT} group (figure 5B–E). Similar to gene expression results, TNFα and IL1β protein levels were reduced in the colon tissues (figure 5F,G). We also observed reduced level of phospho-p65 and phospho-p38 in colons of DSS-treated *mNAIL*^{ΔNFκB} as compared with *mNAIL*^{WT} group (figure 5H).

Colons from DSS-treated *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice at day 8 were stained with *mNAIL*-specific fluorescence in situ hybridisation (FISH) probe, p-p65 and p-p38 antibodies and analysed using confocal microscopy. We only detected basal *mNAIL* expression and did not observe any positive staining for p-p65 and p-p38 in the UT colon sections of *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice. On DSS stimulation, infiltration of F4/80 cells increased in *mNAIL*^{WT} colons. Costaining of F4/80 and *mNAIL* showed that the infiltrated macrophages are the major source of *mNAIL* expression in DSS-treated colons (figure 6A–C-right-panel). Furthermore, only *mNAIL* expressing F4/80+ cells showed dramatic increase in p-p65 (figure 6A) and p-p38 (online supplemental figure 6A). Similar to our western blot analysis (figure 5H), we observed a dampened induction of p-p65 and p-p38 in the *mNAIL*^{ΔNFκB} mice as compared with *mNAIL*^{WT} mice treated with DSS (figure 6A–C and online supplemental figure 6B). Finally, H&E staining of the colon tissues suggested more severe pathology in *mNAIL*^{WT} mice compared with *mNAIL*^{ΔNFκB} mice (online supplemental figure 6C,D). These results, together with our previous western blot analysis, suggest that *mNAIL* expression in colon-infiltrating macrophages regulates phosphorylation of p65 and p38 and hence inflammatory programme in these immune cells.

Indeed, the reduced infiltration of immune cells in *mNAIL*^{ΔNFκB} mice, which was also observed in p38 myeloid specific KO mice,²³ can be explained by the reduction in expression of *CCL2* gene (figure 5E), a key mediator of monocyte recruitment to the inflamed tissue.²¹ In line with our finding, in colitis model of p38 but not in p65 KO mice, it has been shown that expression of chemokines such as *CCL2*, *CCL3* and *CXCL10* was reduced as compared with WT mice which resulted in reduced infiltration of immune cells.^{23 25} CCR2–*CCL2* dimerisation is very important for Ly6C^{hi} cells in bone marrow which give rise to macrophages that are later recruited to inflamed area.²⁷ Much like DSS-treated *mNAIL*^{ΔNFκB} mice, the colitis model of p38 KO mice show reduced levels of Ly6C^{hi} cells in bone marrow which results in less F4/80 infiltration to the colon.²³

Importantly, our results showed that *mNAIL* is not induced in intestinal epithelial cells and there is no difference in TNFα expression between *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice

(online supplemental figure 7A,B). These results showed that BMDMs are possibly the major source of *mNAIL* expression and phenotypes seen may stem from these cells, we performed bone marrow transplantation followed by DSS stimulation (figure 6D). We irradiated *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice and transplanted bone marrow cells isolated from *mNAIL*^{WT} or *mNAIL*^{ΔNFκB} mice (figure 6D). *mNAIL*^{WT} or *mNAIL*^{ΔNFκB} bone marrow reconstituted *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} animals were treated with DSS and body weight was measured for 8 days. As compared with *mNAIL*^{WT} bone-marrow transplanted *mNAIL*^{WT} group (blue), *mNAIL*^{WT} bone-marrow reconstituted *mNAIL*^{ΔNFκB} mice (red) showed similar degree of body weight loss, reduction in colon length and expression of *TNFA* and *IL1β* (figure 6D–G). However, when compared with *mNAIL*^{WT} bone-marrow transplanted *mNAIL*^{WT} group (blue) or *mNAIL*^{WT} bone-marrow transplanted *mNAIL*^{ΔNFκB} group (red), *mNAIL*^{ΔNFκB} bone marrow reconstituted *mNAIL*^{WT} group (green) displayed lesser reduction in body-weight and colon length and showed lower expression of *TNFA* and *IL1β* (figure 6D–G).

Taken together, these results suggest that the differences in *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice in the DSS model stem from the immune cells, particularly from the infiltrated myeloid cells. These results also suggest that *mNAIL*^{ΔNFκB} mice exhibit impaired NFκB and p38 activity leading to reduced inflammation in colon on DSS treatment. Reduced inflammatory cytokines such as *CCL2* in *mNAIL*^{ΔNFκB} mice explain the reduced precursor cell population in bone marrow of these mice.²⁸ Reduction in inflammatory cytokines expression in *mNAIL*^{ΔNFκB} mice may be attributed to reduced activation of p38 and p65, which are well-known upstream events that positively cooperate to regulate transcription of inflammatory genes in response to inflammatory stimuli.

NAIL sequesters Wip1 phosphatase away from its substrates p65 and p38

It has been previously shown that Wip1 phosphatase negatively regulates inflammatory gene expression by dephosphorylating p65²⁹ and p38.³⁰ Indeed, decreased activation of p65 and p38 was seen in the absence of *mNAIL* (figures 4M and 5H). As numerous signalling pathways can be activated by TNFα stimulation, we comprehensively analysed most immediate downstream pathways which are activated on TNF receptor ligation (online supplemental figure 7C). Based on the fact that p65 and p38 phosphorylation were specifically affected in *NAIL* expressing cells, we hypothesised that Wip1 phosphatase, which affects these two substrates,³¹ might be a major effector of *NAIL*. Indeed, enzyme sequestration is a well-known regulatory mechanism on controlling cellular events.^{32–34} To test this, we checked Wip1-p65 interaction in colons of *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice treated with and without DSS and BMDMs treated with and without LPS. We found more Wip1 is bound to p65 in *mNAIL*^{ΔNFκB} colon and BMDM cells even on stimulation and hence, p65 activation is defective (figure 7A,B and online supplemental figure 7D–F). We also found that in *mNAIL*^{ΔNFκB} MEFs and BMDM cells, si-Wip1 rescued *IL1β* and *TNFA* expression and phosphorylation levels of p65 and p38 proteins, further suggesting that Wip1 is a key determinant of *NAIL* action (online supplemental figure 8). Taken together, these results suggest that *NAIL* can exert its function in trans by sequestering Wip1 phosphatase away from p65 or other Wip1 substrates like p38. It is possible that other substrates of Wip1 may also play a role in the described phenomenon consequent to the induction of *NAIL*. Given that this induction is dependent on the p65 subunit

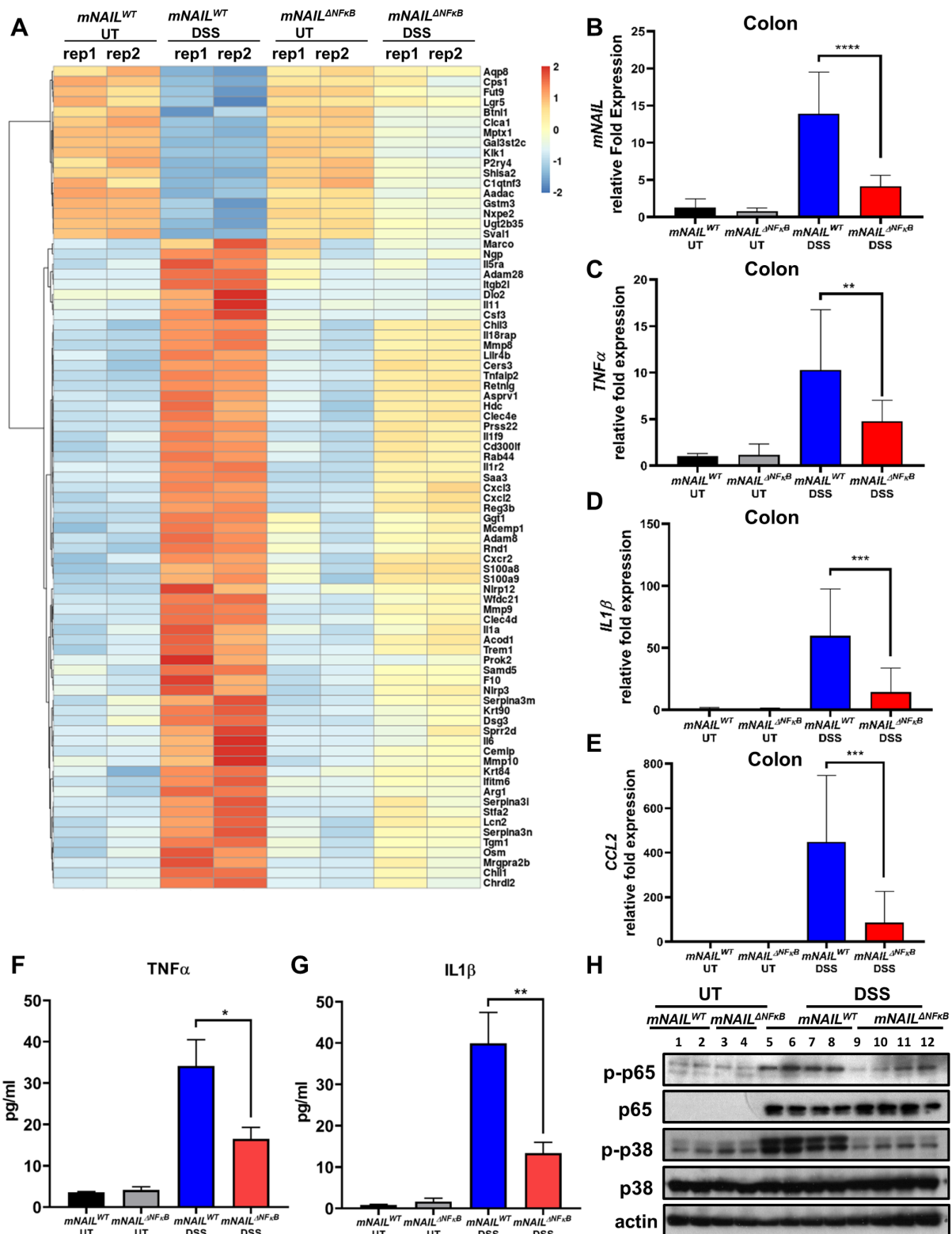


Figure 5 *mNAIL* regulates expression of inflammatory genes in colitis. (A) Heat map shows the genes that are differentially expressed between *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice treated with DSS. Genes were first filtered for differential genes on DSS treatment between *mNAIL*^{WT} untreated (UT) and DSS to identify DSS-dependent genes. Next, this gene list was used to identify the differentially expressed genes between *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice treated with DSS. Differentially expressed genes were identified based on the criteria of 2 log₂-fold difference with false discovery rate < 0.05. (B–E) Expression analysis was performed by RT-qPCR for (B) *mNAIL*, (C) *TNFα* (D) *IL1β* and (E) *CCL2* genes in the colon tissues of *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice treated with or without DSS at day 8 (n=15). (F–G) Protein levels of *TNFα* and *IL1β* cytokines were measured in colon samples by ELISA. Error bars indicate mean ± SD of three independent experiments. P values were calculated using Student's t-test method (*p < 0.01; **p < 0.001; ***p < 0.0001). (H) Total protein lysates isolated from the colon tissues of *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice treated with or without DSS at day 8 were analysed for phosphorylated and total p65 and p38. Actin was used for normalisation. DSS, dextran sulfate sodium; UT, untreated; WT, wild type.

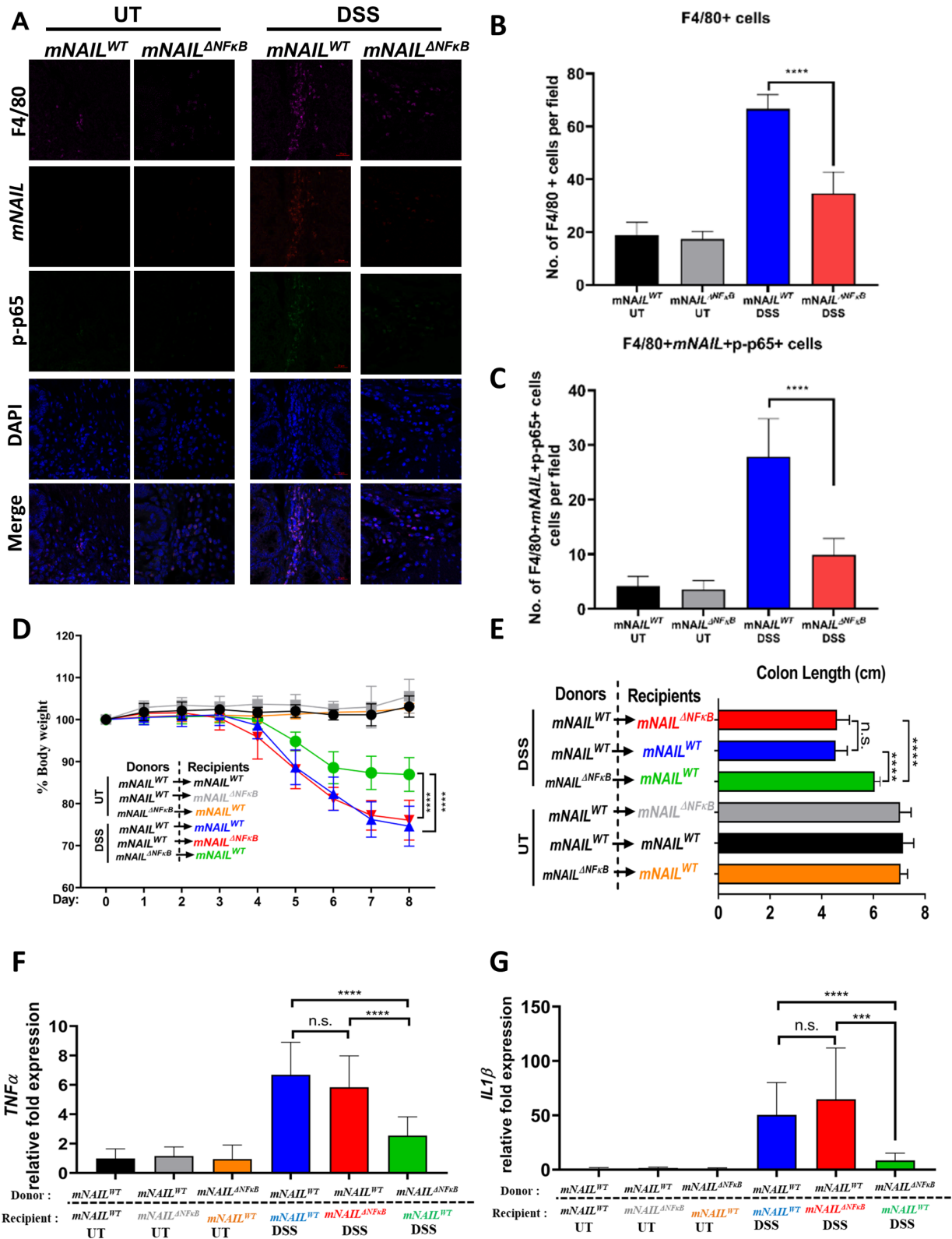
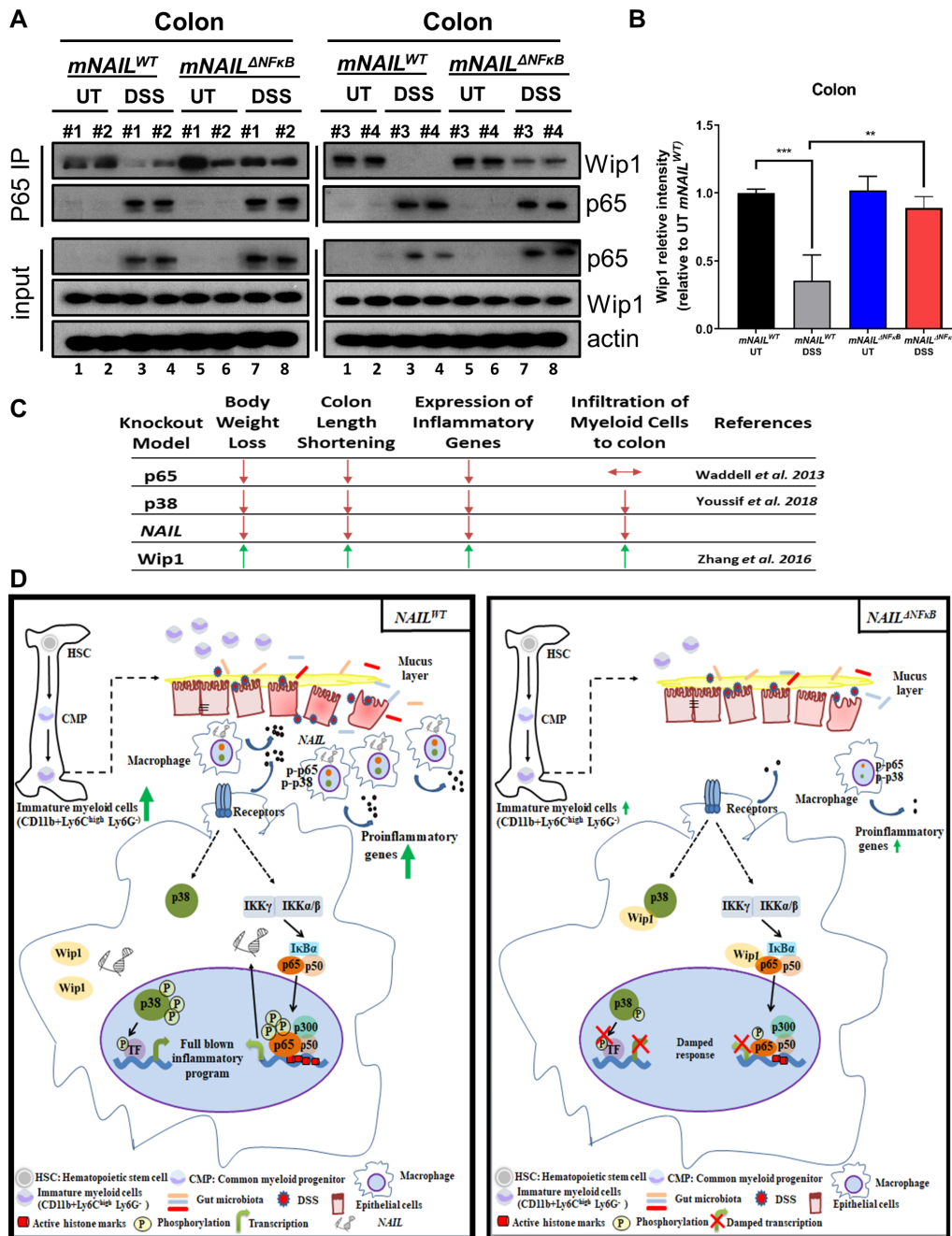


Figure 6 *mNAIL* acts via immune cells to regulate inflammation. (A) Colon sections of the DSS-treated *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice at day 8 were stained with *mNAIL* specific FISH probe, F4/80 and p-p65 antibodies. Tissue slides were analysed for the indicated molecules by confocal microscopy. (B) Graph shows quantification of the number of F4/80+ cells from A. (C) Graph shows quantification of the number of F4/80+p-p65+ cells from A. Error bars indicate mean±SD of three independent fields examined per mouse (n=3 per group). ***p<0.001; ****p<0.0001; n.s., not significant. P values were calculated by two-tailed Student's t-test method. (D) *mNAIL*^{WT} or *mNAIL*^{ΔNFκB} bone marrow reconstituted *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice were treated with DSS for 8 days. Graph shows the body weight measurements from mice monitored for 8 days (E) and the colon length at day 8. Graphs show gene expression analysis of (F) *TNFα* and (G) *IL1β* in the colon tissues of different bone marrow reconstituted groups treated with DSS for 8 days by qPCR. Data are normalised to actin. Error bars indicate mean±SD of two independent experiments (total n=12). P values were calculated using Student's t-test method (***p<0.001; ****p<0.0001; n.s., not significant). DSS, dextran sulfate sodium; UT, untreated; WT, wild type.



of NF κ B, *NAIL* could be the key factor required for full-blown inflammatory responses in an evolutionarily conserved fashion.

DISCUSSION

It is very well established that intestinal microbiota plays a key role in the development of UC. A multitude of regulatory checkpoints, regulated mainly by NF κ B and p38, collectively coordinate the intensity and duration of the inflammatory response.^{35 36} In normal conditions, mucosal immune system acts as an effective defence system against gut microbiota while maintaining the balance between proinflammatory and anti-inflammatory responses. This balance, in UC, is highly impaired and shifted towards hyperactive proinflammatory side, causing increased number of immune cells and high levels of cytokine secretion.³ While the impact of NF κ B and p38 on initiating inflammatory responses is well established, how the proinflammatory and anti-inflammatory responses are kept in balance is unclear. Hence, understanding how these signalling pathways are controlled is of great clinical value towards developing novel drug targets for therapeutic intervention in inflammatory diseases.

This study represents the first of its kind, where mouse genetic tools were used to identify novel lncRNAs that are directly and specifically regulated by NF κ B signalling. We identified *NAIL* (anti-sense direction to *IL7* gene) as the only evolutionarily conserved lncRNA across mice and humans. *NAIL* is specifically induced by inflammatory cytokines such as TNF α and TLR4 ligand LPS in a p65-dependent fashion across cell types.

CRISPR-mediated deletion of NF κ B binding motifs upstream of *NAIL* led to the loss of its expression and decreased inflammation in vivo and in vitro. *NAIL*-deficient cells showed a reduction of p38 and p65-transactivation and subsequent decline in expression of key p38 and NF κ B target genes. NF κ B and p38 have been shown to be required for the expression of proinflammatory and chemoattractant genes including *Cxcl1*, *Cxcl2*, *TNF α* and *IL1 β* ³⁰ for full-blown inflammation. Most notably, loss of *NAIL* phenocopies p38 or p65 loss in vivo in DSS model as published by other laboratories^{23 25 37} (figure 7C). Our in vitro BMDM experiments (figure 4G–M) and in vivo DSS model clearly demonstrated that infiltrated macrophage cells are the major source of *mNAIL* expression and infiltration of F4/80+ cells are dampened in *mNAIL* ^{Δ NF κ B} colons (figure 6A,B). Furthermore, only *NAIL* expressing F4/80+ cells showed dramatic increase in p-p65, suggesting that *NAIL* acts in mainly through immune cells for its mechanism of action (figure 6A–C). Indeed, bone marrow reconstitution experiment further proved this point as the DSS phenotype observed in *mNAIL* ^{Δ NF κ B} mice was rescued on reconstitution of *mNAIL*^{WT} bone marrow into irradiated *mNAIL* ^{Δ NF κ B} mice. Therefore, we can conclude that these phenotypic effects stem from the immune cells, particularly the infiltrated myeloid cells (figure 6D–G). Interestingly, unlike p38 KO mice, p65 KO mice do not show defects in the infiltration of macrophage cells to the inflamed colon, which clearly highlights that simultaneous activation of both these pathways is critical for successful mounting of inflammation.

Our findings clearly explain that *NAIL* sequesters Wip1 away from p65 and p38 simultaneously to enable an orchestrated response that leads to (a) differentiation of precursor cells into immature myeloid cells in bone marrow, (b) recruitment of macrophages to inflamed area and (c) expression of inflammatory genes. Indeed contrary to *NAIL*, p65 or p38 null mice, Wip1 null animals show enhanced inflammation³⁷ (figure 7C). As inflammatory responses and phenotypes observed in *NAIL* loss of function mice are inverse of what is seen in Wip1 KO

mice and they phenocopy the combination of the phenotypes observed in p38 and p65 KO models, it is apparent that *NAIL* is the missing link that explains the coactivation of the two essential pathways, NF κ B and p38, by Wip1 in inflammation.

In physiological conditions, regulation of inflammatory control includes multiple layers of responses. As in the example of DSS-induced colitis model, the first response is the damage in the epithelial barrier which evokes maturation and recruitment signals in bone marrow.³⁸ On sensing the damage signals and penetration of the microbiota to the colon, myeloid progenitor cells differentiate into immature myeloid cells which later on give rise to macrophages that are recruited to the inflamed areas in colon³⁹ and express inflammatory genes (figure 7D). Indeed, p38 has been shown to regulate the differentiation of the myeloid progenitor cells into immature myeloid cells and later on via regulating the required chemoattractants that lead to infiltration of the macrophages into the inflamed colon.^{23 40 41} In the next stage, on activation of NF κ B pathway, these cells express the proinflammatory cytokines for full-blown inflammatory response.⁴² As these two important pathways, p38 and NF κ B, have to be coactivated in a timely manner,⁴³ *NAIL* acts as a bridge to coordinate the activity of these two essential pathways in colitis.

In chronic diseases such as IBD, immunosuppressive drugs are used to partially block NF κ B activity.³ Finer dissection of the role of *NAIL* in chronic inflammatory diseases and cancer may present new insights to design better and more selective anti-inflammatory drugs. Such *NAIL* inhibitors could also be more efficacious since they simultaneously target both NF κ B and p38 signalling. As compared with non-inflamed controls, *NAIL* expression is significantly enhanced in the inflamed colon tissues (figure 1G) suggesting that *NAIL* expression is highly correlated with NF κ B activity and may serve as a biomarker and therapeutic target for IBDs and other diseases with inflammation as an underlying cause.

METHODS

BMDMs isolation and stimulation

BMDMs were isolated from femurs and tibias of sibling 8–10-week-old *mNAIL*^{WT} and *mNAIL* ^{Δ NF κ B} mice. Cells were plated on bacteriological plastic plates in DMEM with 30% L929 conditioned medium for 4 days. On day 4, remove non-attached cells and wash with prewarmed phosphate-buffered saline (PBS) and add fresh medium. On day 7, cells were dissociated using 5 mM PBSethylenediaminetetraacetic acid (EDTA) and cell scraper. Differentiated BMDMs were stained by CD11b and F4/80 antibodies. Differentiated BMDMs were stimulated with LPS (200 ng/mL) for the indicated time points for protein isolation, RNA isolation and chromatin immunoprecipitation.

RNA-sequencing (RNA-seq) and analysis

Total RNA was isolated by using Trizol from immortalised WT and p65 and Ikk β KO (-/-) MEFs that have been described previously.⁴⁴ Cells were either stimulated with 10 ng/mL TNF α for 45 and 90 min or vehicle control. RNA-seq library was prepared by Illumina Truseq Total RNA sequencing kit according to manufacturer instructions. Paired-end sequencing was performed using Illumina Nextseq, 2 \times 76 bp. Base quality assessment of raw sequence was performed using FastQC v0.11.8 and Trimmomatic v0.321 was used to filter low quality bases and adapter sequences. By using HISAT2 V2.1, spliced alignment was performed on the quality trimmed sequences against Mus musculus (mm10) genome index which consist of

transcriptome splicing information. Stranded alignment (first-strand) was applied and the alignment was finally coordinate sorted using Samtools V.1.8. Gene-level expression counts were quantified (strand-specific read counting) using featureCounts (subread-V.1.6.2 package) by reference to the gene annotation obtained from Ensembl (V.96) database.

Differential genes expression analysis was performed by first removal of genes with very low reads count using filterByExp() function in edgeR V.3.26.4 package. Library sizes were recalculated after the filtering. Genes with fold change ≥ 2 and a false discovery rate (FDR) < 0.05 were identified as significant differentially expressed genes (DEGs). Differentially expressed lncRNAs were extracted with reference to GENCODE lncRNA annotation of respective genomes.

RNA was isolated from the colon tissues of *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* mice treated with or without DSS. RNA-Seq library was prepared by Illumina Truseq Total RNA sequencing kit according to manufacturer instructions. Paired-end sequencing was performed using Illumina HiSeq, 2×150bp. DEGs were identified by comparing DSS-treated *mNAIL^{ΔNFκB}* versus DSS-treated *mNAIL^{WT}* and DSS-treated *mNAIL^{WT}* versus UT *mNAIL^{WT}*.

RNA sequencing data of UC patients were obtained from NCBI Sequence Read Archive (SRA), SRP125961 (GEO: GSE107593). Raw sequences were subjected to the same analysis process above and were aligned to Homo sapiens (GRCh38) genome index. DEGs were identified with the same threshold (fold change ≥ 2 , FDR < 0.05).

RNA isolation and real-time qPCR

Total RNA was extracted by Trizol and column purified using RNeasy Mini Kit for gene expression analysis. 1 μg RNA was used as a template for reverse transcription reaction using Maxima first-strand cDNA synthesis kit (Thermo Scientific). Then, real-time PCR was performed from diluted cDNAs using SsoAdvanced Universal SYBR Green Supermix (BioRad). Relative gene expression was analysed by $\Delta\Delta$ CT method by normalising experimental Ct values to β -actin. Primers are indicated in the online supplemental table 1.

Western blot analysis

Total protein was extracted using Totex buffer (20 mM Hepes at pH 7.9, 0.35 M NaCl, 20% glycerol, 1% NP-40, 1 mM MgCl₂, 0.5 mM EDTA, 0.1 mM EGTA, 50 mM NaF and 0.3 mM NaVO₃) supplemented with complete protease and phosphatase inhibitor cocktail (Roche). Immunoblotting was performed with following antibodies: anti-p-p38 (Thr180/Tyr182) 3D7 (Cell signalling; #9215S), anti-p38 (Santa Cruz; #sc-728), anti-p-p65 (Ser536) (Cell signalling; #3031L), anti-p65 (Santa Cruz; #sc-8008), anti-actin (Sigma; #A2066), anti-HSP90 α/β (F-8) (Santa Cruz; #sc-13119), p-IKK α/β (Ser176/180) (Cell signalling; #2697S), IKK α/β (H-470) (Santa Cruz; #7607) (online supplemental table 1).

ELISA analysis

Total protein was extracted from the colon tissues and TNF α and IL1 β protein levels were quantified using Quantikine ELISA kit according to manufacturer's instructions (Thermo Fisher Scientific).

Electrophoretic mobility shift assay

Electrophoretic mobility shift assay (EMSA) was performed as previously described.⁴⁵ Briefly, 20 μg of total protein was

incubated in reaction buffer (50% glycerol, 200 mM Hepes, pH 7.9, 4 mM EDTA pH 8.0, 4 mM DTT), 2 μg polydI-dC, 1% NP40 and [γ -³²P] radiolabelled DNA probe for 30 min at room temperature. Samples were separated on 6% non-denaturing polyacrylamide gel, and the gel was dried and exposed to BAS-IP Image plate (Fuji) for scanning with Typhoon FLA 7000 (GE Healthcare). Radiolabelled probes: NF- κ B: 5'-TCAACAGAGGGG ACTTTCCGAGAGGCC-3', AP-1: 5'-CGCTTGATGACTCAGC GGGAA-3'.

Flow cytometry analysis

Bone marrow cells were isolated from *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* mice treated with and without DSS for 8 days. Cells were treated with ammonium-chloride-potassium (ACK) lysis buffer (Lonza) to eliminate red blood cells. Cells ($\sim 1 \times 10^6$ cells) were incubated with CD11b (PE), Ly6C (BV711) and Ly6G (FITC) fluorescence conjugated antibodies for 20 min at room temperature. To identify the stem cell populations, cells were stained with Lineage antibody cocktail (APC), c-KIT (PerCPy5.5), Sca-1 (FITC), CD34 (PE), IL7ra (PeCy7) and CD16/32 (APCcy7) (BD Biosciences) After antibody incubation, cells were washed with PBS supplemented with 1% FBS and 2 mM EDTA and acquired using FACS LSR2 instrument (Becton Dickinson). Data were analysed using Flow Jo software.

Fluorescence in situ hybridisation

Custom Stellaris RNA FISH probes were designed against *mNAIL* by utilising the Stellaris RNA FISH Probe Designer (Biosearch Technologies, Petaluma, CA). FISH was performed using the *mNAIL* Stellaris FISH Probe set labelled with Quasar 570, according to the manufacturer's instructions available online at www.biosearchtech.com/stellarisprotocols with slight modification. Briefly, colon tissues were sectioned at a thickness of 4 μm using a microtome and mounted onto a microscope slide. Tissue sections were immersed in 100% xylene for 10 min and an additional 5 min with fresh 100% xylene. Next, slides were immersed in 100% ethanol for 20 min followed by incubation in 95% ethanol for 10 min. Subsequently, antigen retrieval by citrate buffer was performed (slides were heated in a microwave submersed in 1X citrate unmasking solution until boiling was initiated and kept for 20 min at a subboiling temperature (95°–98°C)). After allowing the slides to cool down on bench top for 20 min, slides were washed in dH₂O three times for 5 min each and blocked with blocking solution (1% bovine serum albumin (BSA), 2% fetal bovine serum (FBS) in phosphate-buffered saline (PBS)) for 1 hour at room temperature. Subsequently, blocking solution was removed and Alexa Fluor 647 antimouse F4/80 antibody (Clone:BM8, Cat# 123122, Biolegend) diluted in blocking solution and added onto slides for 1 hour at room temperature. Slides were washed with 1X PBS three times for 5 min. Tissue sections were permeabilised in 70% ethanol for at least 1 hour at room temperature. After washing slides with 1X PBS twice for 2–5 min, slides were incubated with hybridisation buffer (SMF-HB1-10, Stellaris) containing NAIL probe (125 nM) and antibodies (1:1000 dilution, phospho-NF- κ B p65 (Ser536) Rabbit mAb, Alexa Fluor 488 Conjugate, clone: 93H1, Cat#4886, CST or phospho-p38 MAPK (Thr180/Tyr182) Rabbit mAb, Alexa Fluor 488 Conjugate, clone: 3D7, #41768, CST) in a humidified chamber (150 mm tissue culture plate sealed with parafilm; a single water-saturated paper towel placed alongside the inner chamber edge) for overnight in the dark at 37°C. After incubation, slides were immersed in Wash Buffer A (SMF-WA1-60, Stellaris) in the dark at 37°C for 30 min

to allow the submerged coverglass fall off from the slides. Finally, Wash Buffer A was removed, and 4',6-diamidino-2-phenylindole (DAPI) was added to counterstain the nuclei and slides were covered with a coverglass for imaging by using ZEISS LSM800 microscopy.

Chromatin immunoprecipitation

BMDM cells derived from *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* cells were stimulated with LPS (200 ng/mL) for 4 hours. Cells were cross-linked on the plate with disuccinimidyl glutarate (DSG) (1.5 mM) (Thermo Fischer Scientific) for 30 min at room temperature with gentle shaking. After removing DSG, cells were fixed with 1% formaldehyde for 10 min and the reaction was quenched with a final concentration of 0.125 M glycine. After lysis, cell lysates were sonicated of an average size of 150–600 bp fragments and precleared with Protein A/G sepharose beads for 1 hour. After centrifugation, 2% input was stored as an input. Lysates (~4 million cells) were incubated with 4 μg p65 (Santa-Cruz, sc-372) antibody overnight at 4°C on a rotator. Immune complexes were immunoprecipitated with protein A/G sepharose beads and samples were eluted in a final volume of 60 μL after reverse crosslinking overnight. ChIP-qPCR was performed using *TNFA* and *CXCL2* promoter-specific primers. Enrichment was calculated by % input method.

Immunoprecipitation assay

Cells were harvested and lysed in IP lysis buffer [50 mM Tris-HCl pH 8.0, 150 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate, 0.1% sodium dodecyl sulfate (SDS)]. Protein concentration was measured by Bradford method. p65 was immunoprecipitated after incubating cell lysate with p65 antibody for 6 hours and an additional 2 hours with Protein G Sepharose beads (GE Healthcare). The beads were washed three times with washing buffer (10 mM Tris-HCl pH 7.5, 1 mM EDTA, 1 mM ethylene glycol-bis(β-aminoethyl ether)-N,N,N',N'-tetraacetic acid (EGTA), 150 mM NaCl, 1% Triton X-100) and immunoprecipitated proteins were eluted by boiling the beads in 2X loading sample buffer (LDS) (Invitrogen) for 10 min. Immunoblotting was performed as described above with the following antibodies: anti-Wip1 antibody (1:1000 Cell signalling; #11901), anti-p65 antibody (1:1000, Santa Cruz; sc-8008).

Generation of mice

We designed crRNAs targeting 5'-TAGGGTTTAAAAGCG-CATCC-3' and 5'-AGTCTGGGAGTTTCCGATCC-3' in the 5'UTR of Gm16685, then generated *NAIL^{ΔNFκB}* mice by introducing the CAS9/tracrRNA/crRNAs complex into the oocytes with electroporation, as described previously.⁴⁶ In brief, CARD HyperOva (0.1 mL, Kyudo, Tosu, Saga, Japan) was injected into the abdominal cavity of C57BL/6N females [Japan SLC (Hamamatsu, Shizuoka, Japan)] followed by human chorionic gonadotropin (hCG) (five units, ASKA Pharmaceutical). After 14 hours of hCG injection, the oocytes were collected from the ampulla of the oviduct and then were used for in vitro fertilisation.⁴⁷ The fertilised eggs were electroporated with the CAS9/tracrRNA/crRNA ribonucleoprotein complex, and then transplanted into the ampulla of the oviduct of pseudopregnant females. After 19 days, offspring were obtained by natural birth or Caesarean section. We genotyped the offspring by PCR with KOD-Fx neo (TOYOBO, Osaka, Japan) and a primer set (Fw: 5'-TTAGATTTGATCAGGGAGC-3', Rv: 5'-TCAGTG-GTTTTAGCCACTT-3') and direct sequencing. The PCR condition was 94°C for 3 min, denaturing at 94°C for 30 s,

annealing at 55°C for 30 s, elongation 72°C for 30 s for 40 cycles total, followed by 72°C for 2 min. Briefly, we obtained 22 pups from 89 treated zygotes and seven out of 12 analysed pups carried the 80 or 82 bp deletion. We crossed these mice to WT animals for two generations to purify 80 bp deletion mutants and also decrease the risks of off-target cleavage. Then heterozygous mutants were crossed to obtain homozygous 80 bp deletion mutant mice which was confirmed by Sanger sequencing. Subsequently, KO mouse was crossed back to WT to obtain heterozygous animals which were used as breeders to generate littermate WT, HT and KO groups for experiments.

Animal studies and ethical requirements

DSS model was conducted as previously described.⁴⁸ Briefly, *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* mice were fed with 3% DSS in drinking water for 4 days and replaced with normal water for another 4 days. Body weight was measured daily and mice were sacrificed at day 8. Colon (distal) and bone marrow tissues were harvested and used for downstream analysis including RNA, protein and flow cytometry analysis. Intestinal epithelial cells were isolated for gene expression analysis from the colon tissues as reported previously.⁴⁹ DAI was assessed by using the evaluation system: (i) body weight loss (no weight loss was scored as 0 points, weight loss of 1%–5% as one point, 5%–10% as two points, 10%–20% as three points and more than 20% as four points),⁵⁰ stool consistency (0 point was given for well-formed pellets, two points for pasty and semiformal stools that did not stick to the anus and four points for liquid stools that remained adhesive to the anus). For histological scoring, the sections were graded in a blind manner by two independent investigators with a range from 0 to 3 for the amount of inflammation, depth of inflammation and with a range from 0 to 4 for the amount of crypt damage. For bone marrow transplantation, single cell suspension of bone marrow cells was obtained from the tibia and femur of *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* 8–10 weeks old (male) donor mice. 3×10^6 bone marrow cells were injected via the tail vein into *mNAIL^{WT}* and *mNAIL^{ΔNFκB}* lethally irradiated (9.5 Grey) 8–10 weeks old recipient (male) mice. Two weeks' post-transplantation, the mice were placed on the DSS-induced colitis model.

All animal studies were conducted in accordance with the Institutional Animal Care and Use Committee at A*STAR (Singapore). All procedures were approved under the IACUC protocol ID # 181396.

Statistical analysis

Two-tailed Student's t-test was performed for statistical analysis of between two groups. PRISM software V.7 was used to plot graphs and for statistical analysis.

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Contributors SCA, BU and VT conceived the study and designed the experiments. LW and BU performed mechanistic experiments with the help of QFN. SCA, LW performed mouse DSS modelling experiments with the help of QFN, TN and MI generated *mNAIL-ΔNFκB* mice. JYHC performed RNA-seq and bioinformatics analyses. VT directed the study and wrote the paper with LW and SCA.

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Patient consent for publication Not required.

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REFERENCES

- Neurath MF. Cytokines in inflammatory bowel disease. *Nat Rev Immunol* 2014;14:329–42.
- Shen Z-H, Zhu C-X, Quan Y-S, et al. Relationship between intestinal microbiota and ulcerative colitis: mechanisms and clinical application of probiotics and fecal microbiota transplantation. *World J Gastroenterol* 2018;24:5–14.
- Atreya I, Atreya R, Neurath MF. NF-kappaB in inflammatory bowel disease. *J Intern Med* 2008;263:591–6.
- Cohn HM, Dave M, Loftus EV. Understanding the cautions and contraindications of immunomodulator and biologic therapies for use in inflammatory bowel disease. *Inflamm Bowel Dis* 2017;23:1301–15.
- Karin M, Greten FR. NF-kappaB: linking inflammation and immunity to cancer development and progression. *Nat Rev Immunol* 2005;5:749–59.
- O'Connor PM, Lapointe TK, Beck PL, et al. Mechanisms by which inflammation may increase intestinal cancer risk in inflammatory bowel disease. *Inflamm Bowel Dis* 2010;16:1411–20.
- Ruland J. Return to homeostasis: downregulation of NF-kB responses. *Nat Immunol* 2011;12:709–14.
- Derrien T, Johnson R, Bussotti G, et al. The GENCODE v7 catalog of human long noncoding RNAs: analysis of their gene structure, evolution, and expression. *Genome Res* 2012;22:1775–89.
- Harrow J, Frankish A, Gonzalez JM, et al. GENCODE: the reference human genome annotation for the encode project. *Genome Res* 2012;22:1760–74.
- Diederichs S. The four dimensions of noncoding RNA conservation. *Trends Genet* 2014;30:121–3.
- Chen J, Shishkin AA, Zhu X, et al. Evolutionary analysis across mammals reveals distinct classes of long non-coding RNAs. *Genome Biol* 2016;17:19.
- Garber M, Yosef N, Goren A, et al. A high-throughput chromatin immunoprecipitation approach reveals principles of dynamic gene regulation in mammals. *Mol Cell* 2012;47:810–22.
- Barish GD, Yu RT, Karunasiri M, et al. Bcl-6 and NF-kappaB cistromes mediate opposing regulation of the innate immune response. *Genes Dev* 2010;24:2760–5.
- Schmidt SF, Larsen BD, Loft A, et al. Acute TNF-induced repression of cell identity genes is mediated by NFkB-directed redistribution of cofactors from super-enhancers. *Genome Res* 2015;25:1281–94.
- Jin F, Li Y, Dixon JR, et al. A high-resolution map of the three-dimensional chromatin interactome in human cells. *Nature* 2013;503:290–4.
- Rogler G, Brand K, Vogl D, et al. Nuclear factor kappaB is activated in macrophages and epithelial cells of inflamed intestinal mucosa. *Gastroenterology* 1998;115:357–69.
- Schreiber S, Nikolaus S, Hampe J. Activation of nuclear factor kappa B in inflammatory bowel disease. *Gut* 1998;42:477–84.
- Zhang R, Ito S, Nishio N, et al. Dextran sulphate sodium increases splenic Gr1(+)/CD11b(+) cells which accelerate recovery from colitis following intravenous transplantation. *Clin Exp Immunol* 2011;164:417–27.
- Trottier MD, Irwin R, Li Y, et al. Enhanced production of early lineages of monocytic and granulocytic cells in mice with colitis. *Proc Natl Acad Sci U S A* 2012;109:16594–9.
- Geissmann F, Jung S, Littman DR. Blood monocytes consist of two principal subsets with distinct migratory properties. *Immunity* 2003;19:71–82.
- Bain CC, Scott CL, Uronen-Hansson H, et al. Resident and pro-inflammatory macrophages in the colon represent alternative context-dependent fates of the same Ly6Chi monocyte precursors. *Mucosal Immunol* 2013;6:498–510.
- Zigmond E, Varol C, Farache J, et al. Ly6C hi monocytes in the inflamed colon give rise to proinflammatory effector cells and migratory antigen-presenting cells. *Immunity* 2012;37:1076–90.
- Youssif C, Cubillos-Rojas M, Comalada M, et al. Myeloid p38α signaling promotes intestinal IGF-1 production and inflammation-associated tumorigenesis. *EMBO Mol Med* 2018;10.
- Mayle A, Luo M, Jeong M, et al. Flow cytometry analysis of murine hematopoietic stem cells. *Cytometry A* 2013;83:27–37.
- Waddell A, Ahrens R, Tsai Y-T, et al. Intestinal CCL11 and eosinophilic inflammation is regulated by myeloid cell-specific RelA/p65 in mice. *J Immunol* 2013;190:4773–85.
- Otsuka M, Kang YJ, Ren J, et al. Distinct effects of p38alpha deletion in myeloid lineage and gut epithelia in mouse models of inflammatory bowel disease. *Gastroenterology* 2010;138:1255–65. 65 e1–9.
- Bain CC, Mowat AM. Macrophages in intestinal homeostasis and inflammation. *Immunol Rev* 2014;260:102–17.
- Gschwandtner M, Derler R, Midwood KS. More than just attractive: how CCL2 influences myeloid cell behavior beyond chemotaxis. *Front Immunol* 2019;10:2759.
- Chew J, Biswas S, Shreeram S, et al. WIP1 phosphatase is a negative regulator of NF-kappaB signalling. *Nat Cell Biol* 2009;11:659–66.
- Kim C, Sano Y, Todorova K, et al. The kinase p38 alpha serves cell type-specific inflammatory functions in skin injury and coordinates pro- and anti-inflammatory gene expression. *Nat Immunol* 2008;9:1019–27.
- Donehower LA. Phosphatases reverse p53-mediated cell cycle checkpoints. *Proc Natl Acad Sci U S A* 2014;111:7172–3.
- Feng S, Ollivier JF, Soyer OS. Enzyme sequestration as a tuning point in controlling response dynamics of signalling networks. *PLoS Comput Biol* 2016;12:e1004918.
- Taniguchi CM, Emanuelli B, Kahn CR. Critical nodes in signalling pathways: insights into insulin action. *Nat Rev Mol Cell Biol* 2006;7:85–96.
- Blüthgen N, Bruggeman FJ, Legewie S, et al. Effects of sequestration on signal transduction cascades. *FEBS J* 2006;273:895–906.
- Schonhaler HB, Guinea-Viniegra J, Wagner EF. Targeting inflammation by modulating the Jun/AP-1 pathway. *Ann Rheum Dis* 2011;70(Suppl 1):i109–12.
- Oeckinghaus A, Hayden MS, Ghosh S. Crosstalk in NF-kB signaling pathways. *Nat Immunol* 2011;12:695–708.
- Zhang Q, Zhang C, Chang F, et al. Wip 1 inhibits intestinal inflammation in inflammatory bowel disease. *Cell Immunol* 2016;310:63–70.
- Gorjifard S, Goldszmid RS. Microbiota-myeloid cell crosstalk beyond the gut. *J Leukoc Biol* 2016;100:865–79.
- Sartor RB. Mechanisms of disease: pathogenesis of Crohn's disease and ulcerative colitis. *Nat Clin Pract Gastroenterol Hepatol* 2006;3:390–407.
- Verma A, Deb DK, Sassano A, et al. Activation of the p38 mitogen-activated protein kinase mediates the suppressive effects of type I interferons and transforming growth factor-beta on normal hematopoiesis. *J Biol Chem* 2002;277:7726–35.
- Geest CR, Buitenhuis M, Laarhoven AG, et al. p38 MAP kinase inhibits neutrophil development through phosphorylation of C/EBPalpha on serine 21. *Stem Cells* 2009;27:2271–82.
- Medzhitov R, Horg T. Transcriptional control of the inflammatory response. *Nat Rev Immunol* 2009;9:692–703.
- Bradbury CM, Markovina S, Wei SJ, et al. Indomethacin-induced radiosensitization and inhibition of ionizing radiation-induced NF-kappaB activation in HeLa cells occur via a mechanism involving p38 MAP kinase. *Cancer Res* 2001;61:7689–96.
- Tergaonkar V, Bottero V, Ikawa M, et al. IkappaB kinase-independent IkappaBalpha degradation pathway: functional NF-kappaB activity and implications for cancer therapy. *Mol Cell Biol* 2003;23:8070–83.
- Shin EM, Hay HS, Lee MH, et al. Dead-Box helicase DP103 defines metastatic potential of human breast cancers. *J Clin Invest* 2014;124:3807–24.
- Noda T, Sakurai N, Nozawa K, et al. Nine genes abundantly expressed in the epididymis are not essential for male fecundity in mice. *Andrology* 2019;7:644–53.
- Tokuhiro K, Ikawa M, Benham AM, et al. Protein disulfide isomerase homolog PDILT is required for quality control of sperm membrane protein ADAM3 and male fertility [corrected]. *Proc Natl Acad Sci U S A* 2012;109:3850–5.
- Helke K, Angel P, Lu P, et al. Ceramide synthase 6 deficiency enhances inflammation in the DSS model of colitis. *Sci Rep* 2018;8:1627.
- Macartney KK, Baumgart DC, Carding SR, et al. Primary murine small intestinal epithelial cells, maintained in long-term culture, are susceptible to rotavirus infection. *J Virol* 2000;74:5597–603.
- Ishii KJ, Koyama S, Nakagawa A, et al. Host innate immune receptors and beyond: making sense of microbial infections. *Cell Host Microbe* 2008;3:352–63.

SUPPLEMENTARY FIGURE LEGENDS

Sup. Figure 1: Validation of NFκB KO MEF cells. **A)** Western blot analysis of $p65^{-/-}$, $Ikk\beta^{-/-}$ and $Ikk\gamma^{-/-}$ immortalized MEFs with indicated antibodies. **B)** EMSA was performed using either NFκB or AP1 consensus sequence radiolabelled probes as indicated using protein extracts from WT, $p65^{-/-}$, $Ikk\beta^{-/-}$ and $Ikk\gamma^{-/-}$ immortalized MEFs treated with TNFα for the indicated time points. **C)** RNA-seq analysis of Gm16685 expression in WT, $p65^{-/-}$ and $Ikk\beta^{-/-}$ immortalized MEFs exposed to TNFα for indicated duration. **D)** RT-qPCR analysis of Gm16685 expression in WT, $p65^{-/-}$, $Ikk\beta^{-/-}$ and $Ikk\gamma^{-/-}$ immortalized MEFs exposed to TNFα for indicated duration. **E)** RT-qPCR analysis of Gm16685 expression in immortalized WT MEFs exposed to LPS (100ng/ml) for indicated time points. **F)** RT-qPCR analysis of *IL7* expression in WT, $p65^{-/-}$, $Ikk\beta^{-/-}$ and $Ikk\gamma^{-/-}$ immortalized MEFs exposed to TNFα for indicated time points. Error bars indicate mean ± SD of three independent experiments. P values were calculated using Student's t-test method (**, $p < 0.01$; ***, $p < 0.001$).

Sup. Figure 2: Conservation and analysis of Gm16685 lncRNA. **A)** Multiple sequence alignment using promoter sequence of the mouse Gm16685 and sequence of 31 other species at the same region with reference to UCSC comparative genomic track. Clustal Omega was used for the alignment. **B)** Promoter motifs enrichment analysis by using Homer-v4.1.0 package. Top 10 most enriched motifs are shown. **C)** Prediction of different RBPs that can bind to Gm16685 or loc105375914 was performed using RBPmap server (<http://rbpmap.technion.ac.il/>). Venn diagram representing numbers of overlapping predicted RBPs that can bind both Gm16685 and loc105375914 is shown. **D)** ChIP-seq analysis of p65 (RelA) binding in mouse dendritic and macrophage cells exposed to LPS for 2 hours based on

published data (GSE36099 and GSE93736 or GSE93602). **E**) CHIP-seq analysis of RelA and Pol2 signal in human adipocyte and IMR90 fibroblasts exposed to TNF α for 1 hour based on published data (GSE60462 and GSE43070). **F**) Bar plot shows the coding probability of seven NAIL (Loc105375914 for human and Gm16685 for mouse) transcript sequences (six *hNAIL* isoforms and one *mNAIL*) together with a known lncRNA sequence, TERC and actin gamma 1 (ACTG1) mRNA sequence as positive and negative control. Coding probability was calculated with CPC2 tools which indicate the closeness to the coding classification hyper-plane in the trained support vector machine (SVM) model. **G**) 293T cells were transfected with Flag, *NAIL*-Flag and PKR-Flag (Protein kinase R) vectors. Total cell lysates were resolved in SDS-PAGE gel and immunoblotted for Flag and Hsp90 proteins.

Sup. Figure 3: No difference in IL-7 expression and induction in *mNAIL*^{ANF κ B} mice. **A**) Gene expression profiles of mouse *IL7* across different mouse tissues. All the raw FPKM values were obtained from Mouse ENCODE transcriptome data in NCBI (BioProject: PRJNA66167, Publication: PMID 25409824). **B**) Gene expression analysis was performed by RT-qPCR for mouse *IL7* in the thymus tissues of *mNAIL*^{WT} and *mNAIL*^{ANF κ B} mice (n=3). **C**) Total protein lysates isolated from the thymus tissues of *mNAIL*^{WT} and *mNAIL*^{ANF κ B} mice were analysed for mouse *IL7* by western blot. Actin was used for normalization. Graph shows the quantification of *IL7* western blot band intensity. **D-F**) Wild-type MEFs were transfected with si-Control, mouse *IL7* siRNA#1 or mouse *IL7* siRNA#2. After 48h post-transfection, cells were treated with or without TNF α and harvested for gene expression analysis or western blot. Graphs show the gene expression analysis of **D**) *IL7* and **E**) *TNF α* by RT-qPCR. Actin was used as a control. Error bars indicate mean \pm SD of three independent experiments. P values were calculated using Student's t-test method (*, p<0.05; **, p<0.01; ***, p<0.001; ****,

p<0.0001; n.s., not significant). **F)** Cells lysate were analysed via western blot for the indicated proteins.

Sup. Figure 4: No difference in stem cells and precursor cells between *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice. A-E) Bone marrow cells were isolated from *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice (n=7) and treated with or without DSS at day 8. Cells were stained for Lineage marker cocktail, CD34, IL7ra, c-KIT, Sca-1 and CD16/32 cell surface markers. **A)** Cells were analysed by FACS and firstly gated as LIN- cells. LIN- cells were further gated as KL, KSL and K-low S-low cells based on the c-KIT and Sca-1 expression. KL cells were further gated into MEP, CMP and GMP according to CD16/32 and CD34 expression. K-low and S-low cells were gated for CLP population based on the IL7ra expression. Representative FACS data was shown for *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice treated with or without DSS. Quantification of **B)** GMP, **C)** CMP, **D)** MEP and **E)** CLP cells. Error bars indicate mean ± SD of 3 UT and 7 DSS treated samples. P values were calculated using Student's t-test method (n.s., not significant).

Sup. Figure 5: Genome editing process did not cause non-specific global transcriptome changes. A) RNA-seq was performed with colon tissues obtained at Day 8 from *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice treated with or without DSS. Venn diagram representing numbers of overlapping differentially expressed genes between different experimental groups is shown. **B)** GO analysis was performed from the genes that are differentially expressed between *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice treated with DSS.

Sup. Figure 6: *mNAIL*^{ΔNFκB} mice are protected from DSS induced colitis. A, B) Colon sections of the DSS treated *mNAIL*^{WT} and *mNAIL*^{ΔNFκB} mice (n=3) at day 8 were stained with *mNAIL* specific FISH probe (Quasar570), F4/80 (AF647) and p-p38 (AF488) antibodies. Tissue slides were analysed for the indicated molecules by LSM800 confocal microscopy.

Nuclei were counterstained with DAPI. Representative images for co-staining of **A)** *mNAIL*, F4/80 and p-p38 is shown. Quantification of **B)** F4/80, *mNAIL* and p-p38 positive cells is shown as bar graphs. Error bars indicate mean \pm SD of three independent fields examined per mouse (n=3 per group). *, $p < 0.05$; ****, $p < 0.0001$. p-values were calculated by two-tailed student's t-test method based on three independent experiments. **C)** Representative H & E staining and **D)** quantification of histology scores of colon sections from DSS treated *mNAIL*^{WT} and *mNAIL* ^{Δ NF κ B} mice at day 8. Error bars indicate mean \pm SD of three independent experiments (n=9 for UT, n=15 for DSS treated samples)

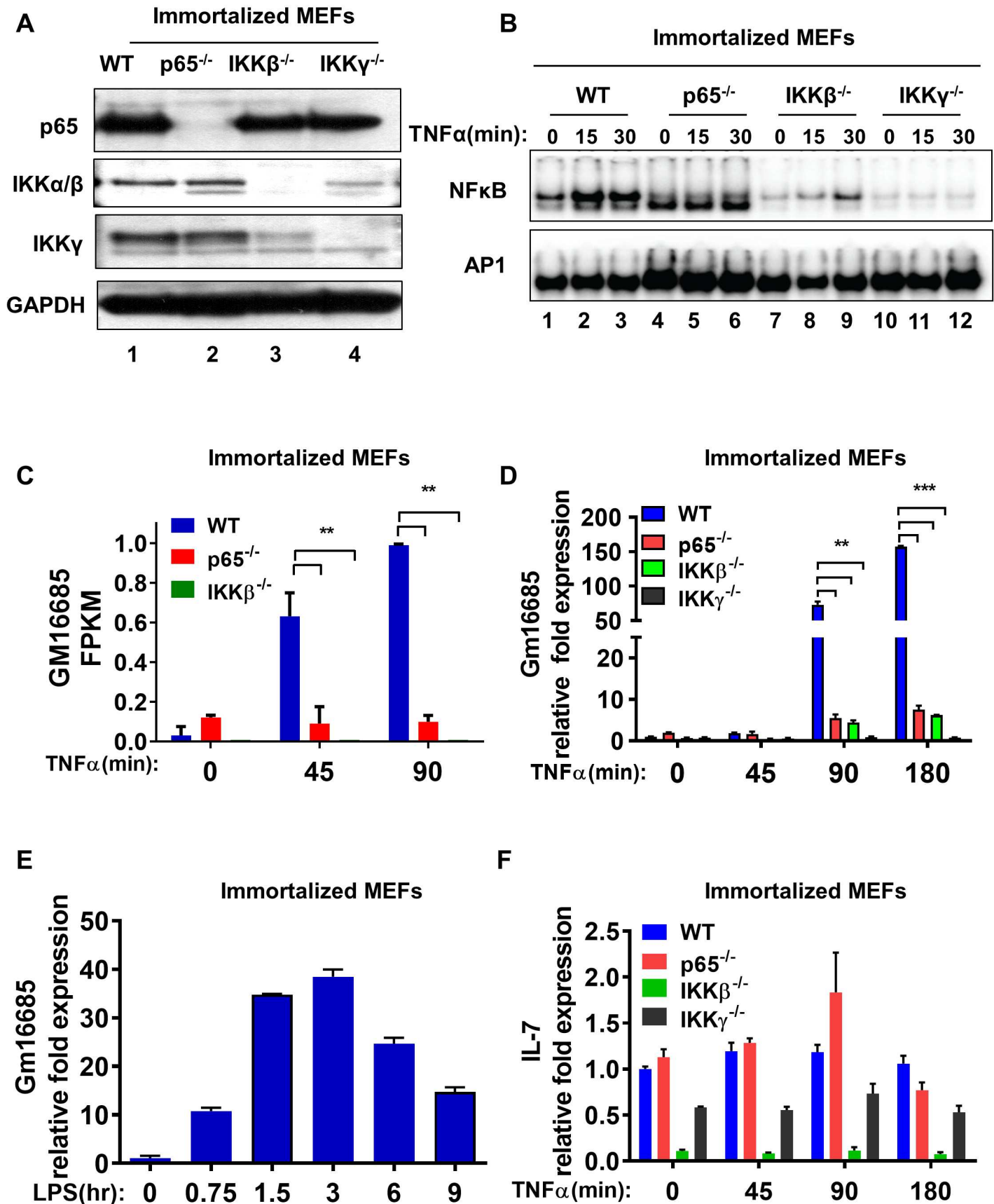
Sup. Figure 7: *NAIL* alters phosphorylation of p65 and p38 but not upstream molecules.

A-B) Graphs show the gene expression analysis of **A)** *mNAIL* and **B)** *TNF α* by RT-qPCR in intestinal epithelial cells isolated from colon tissues of *mNAIL*^{WT} and *mNAIL* ^{Δ NF κ B} mice treated with and without DSS for 8 days. Actin was used as a control. Error bars indicate mean \pm SD of 6 mice. P values were calculated using Student's t-test method (**, $p < 0.01$; ***, $p < 0.001$). n.s., not significant. **C)** *mNAIL*^{WT} MEFs were transfected with si-*Control* or si-*NAIL* and after 48h post-transfection, cells were stimulated with TNF α for indicated time points. Cell lysates were analyzed by western blot for the indicated proteins. **D-E)** Bone marrow cells isolated from *mNAIL*^{WT} and *mNAIL* ^{Δ NF κ B} mice (n=4) were differentiated into bone marrow derived macrophages (BMDM) for 7 days. BMDM cells were treated with or without LPS (200ng/ml) for 4 hours. p65 immunoprecipitation was performed. Total and co-immunoprecipitated proteins were resolved on SDS-PAGE and immunoblotted for indicated proteins. **F)** Graph shows quantification of co-precipitated Wip1 from **D-E**. Error bars indicate mean \pm SD of four replicates. P values were calculated using Student's t-test method (**, $p < 0.01$).

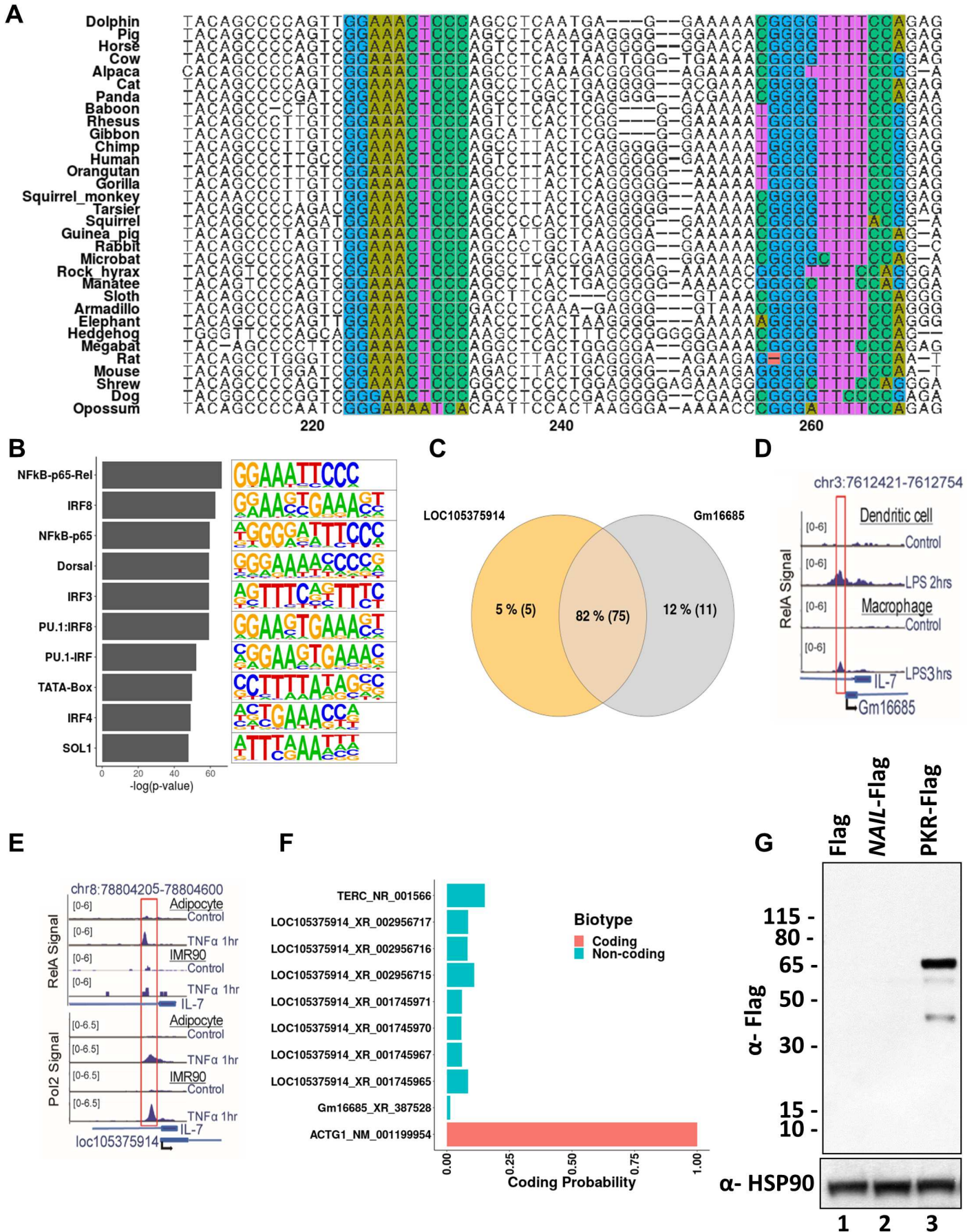
Sup. Figure 8: Wip1 is a key determinant of *NAIL* action. **A-C)** *mNAIL*^{WT} and *mNAIL* ^{Δ NF κ B} MEFs were transfected with control siRNA or Wip1 siRNA. After 48h post-transfection, cells

were treated with or without TNF α and harvested for gene expression analysis or western blot. Graphs show the gene expression analysis of **A)** *IL1 β* and **B)** *TNF α* by RT-qPCR. Actin was used as a control. **C)** siRNA transfected and TNF α stimulated MEFs cell lysates were analysed via western blot for the indicated proteins. **D-F)** *mNAIL^{WT}* and *mNAIL ^{Δ NF κ B}* BMDMs were transfected with control siRNA or Wip1 siRNA. After 48h post-transfection, cells were treated with or without LPS and harvested for gene expression analysis or western blot. Graphs show the gene expression analysis of **D)** *IL1 β* and **E)** *TNF α* by RT-qPCR. Actin was used as a control. Error bars indicate mean \pm SD of three independent experiments. P values were calculated using Student's t-test method (***, $p < 0.001$). **F)** siRNA transfected and LPS stimulated BMDM cell lysates were analysed via western blot for the indicated proteins.

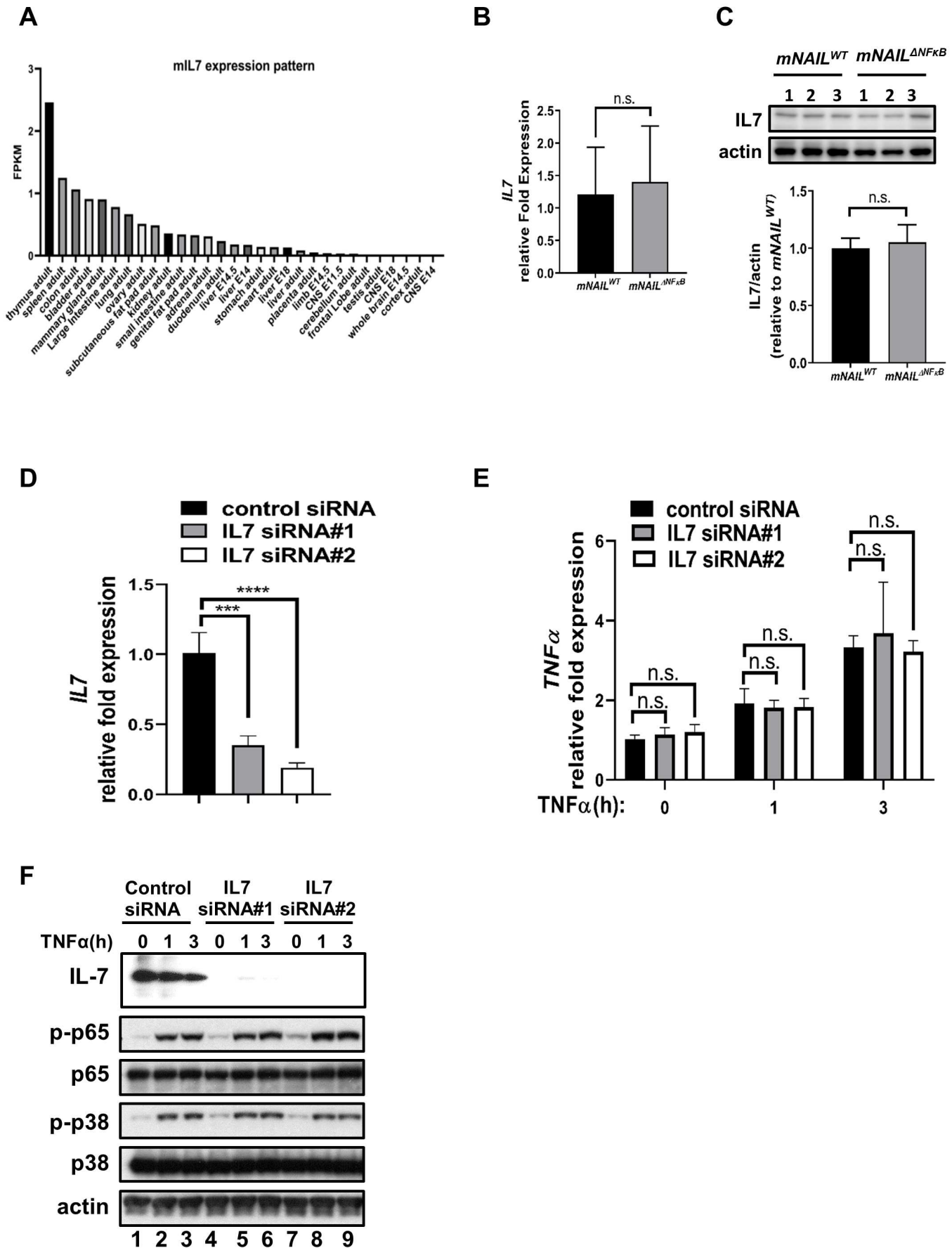
Sup-Fig.1



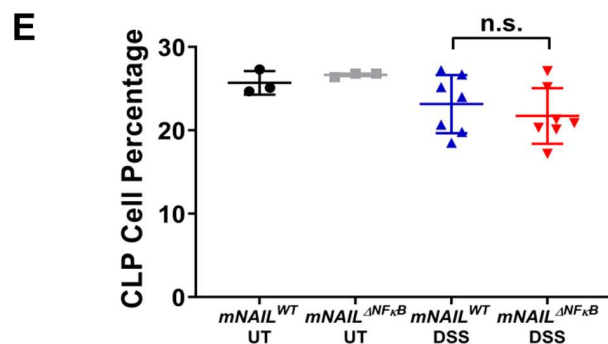
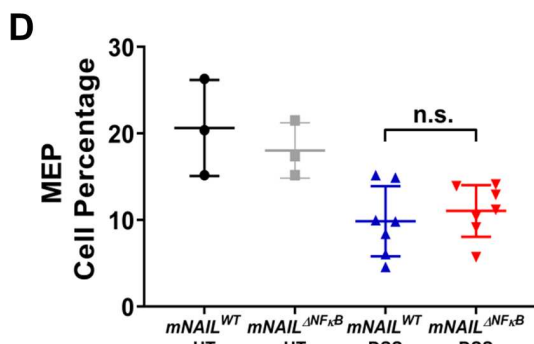
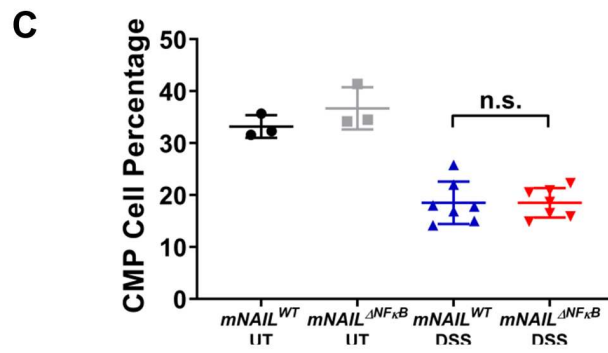
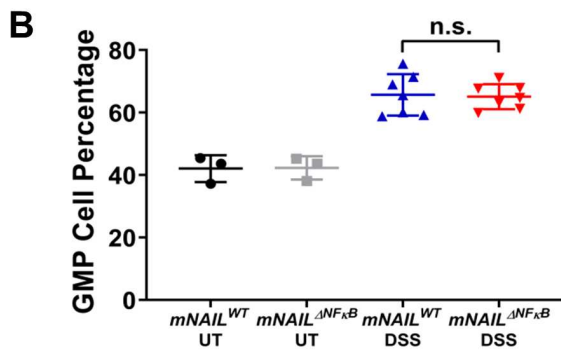
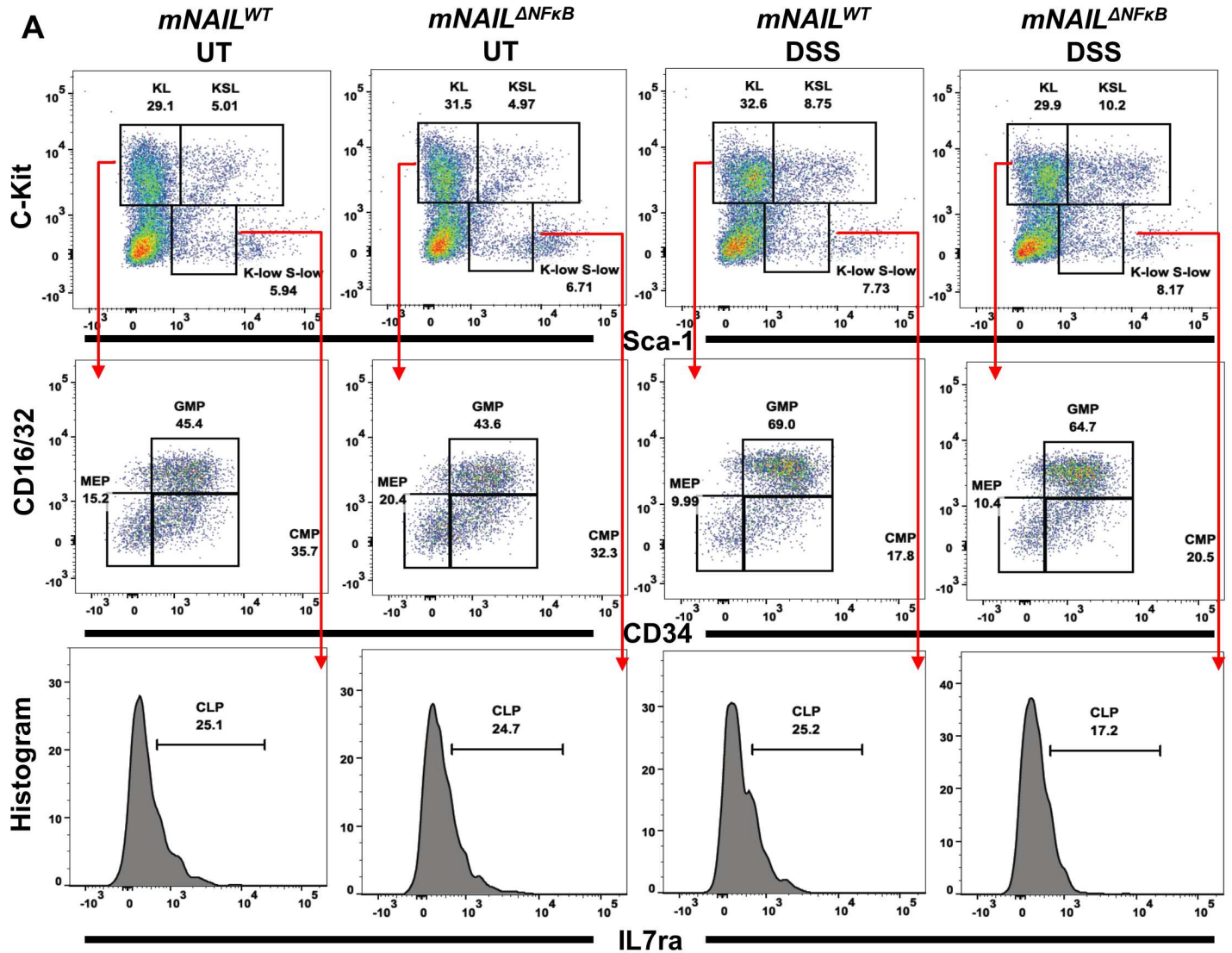
Sup-Fig.2



Sup-Fig.3

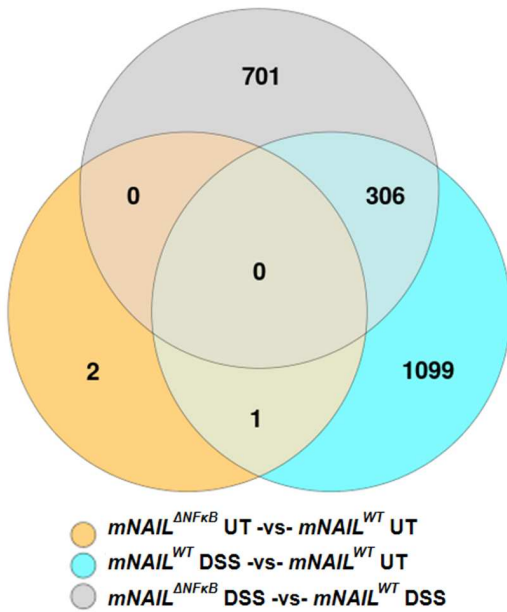


Sup-Fig. 4



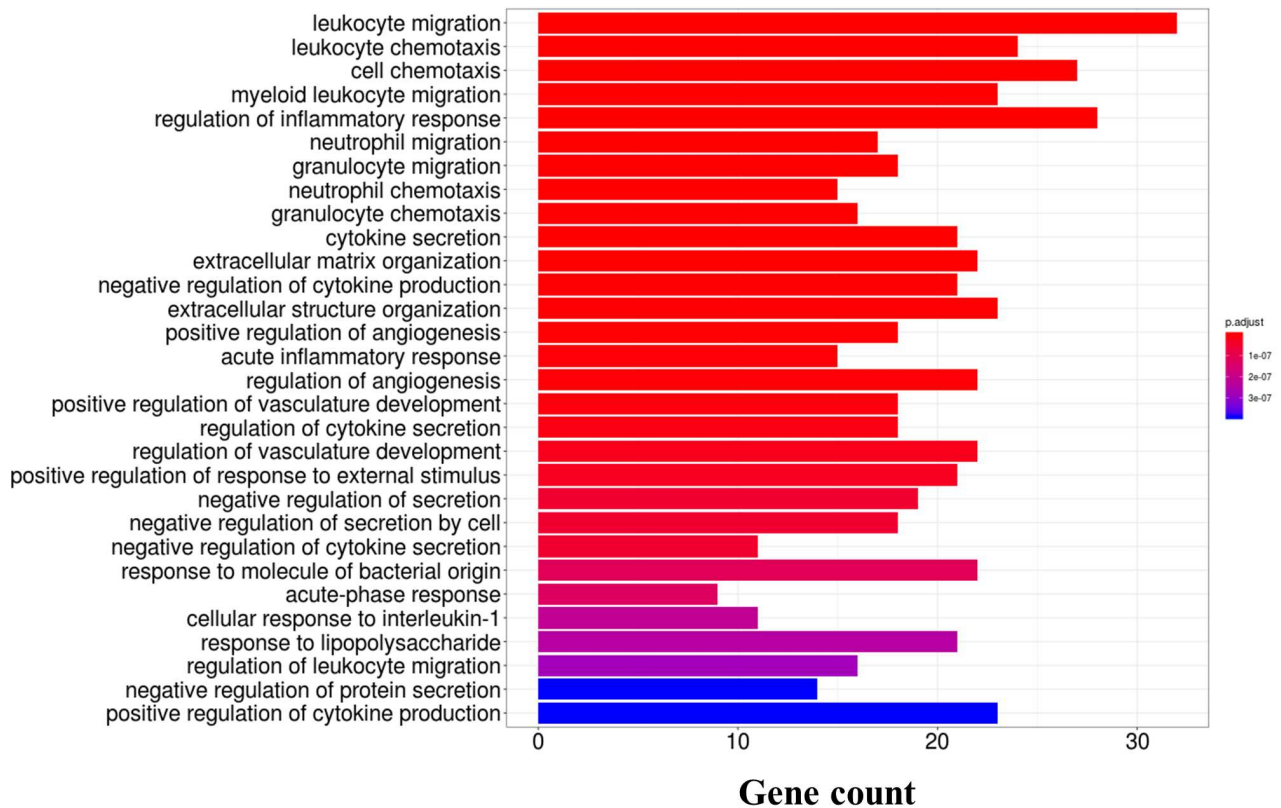
Sup-Fig. 5

A

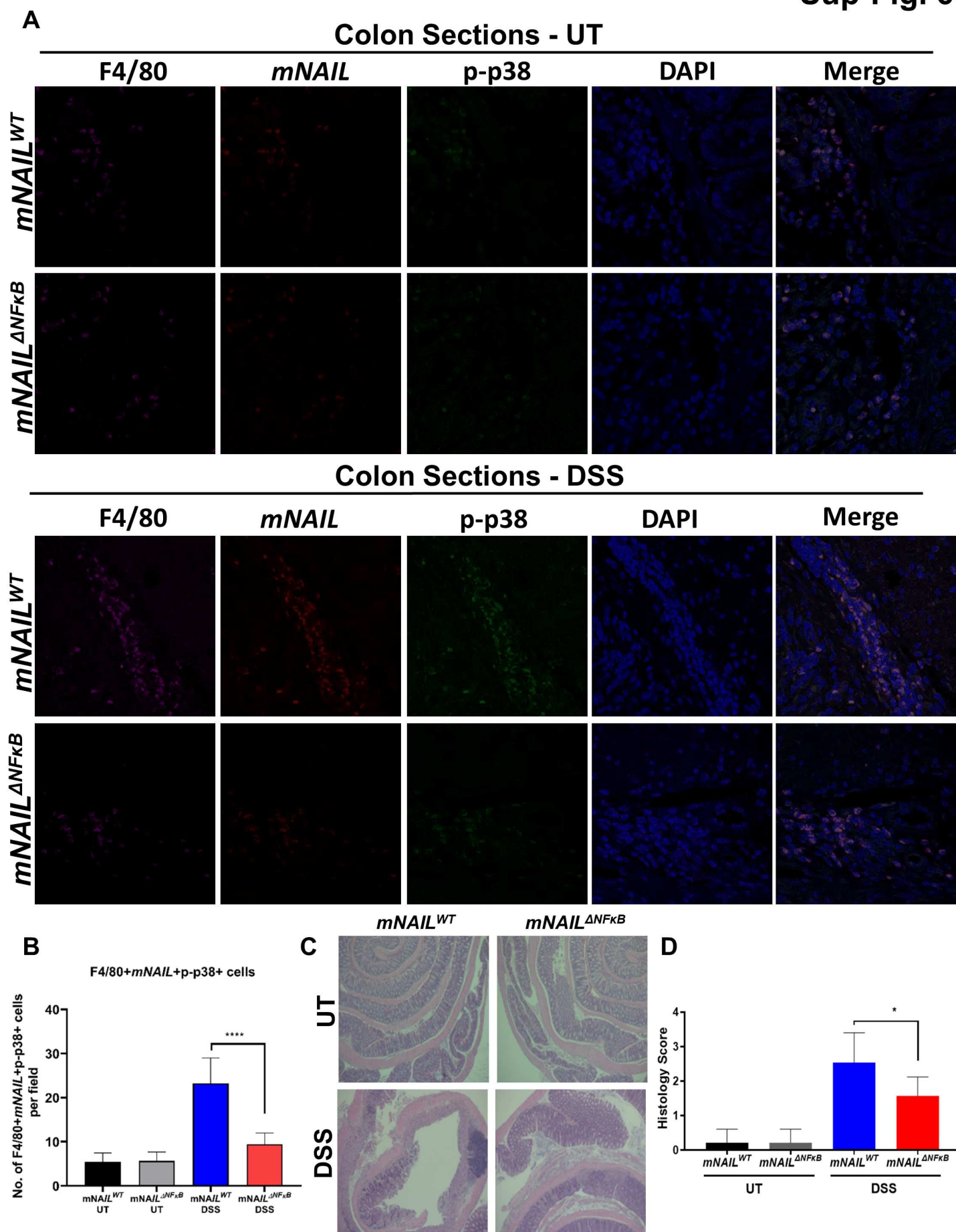


Comparisons	Total differential expressed genes
$mNAIL^{\Delta NF\kappa B}$ UT -vs- $mNAIL^{WT}$ UT	3
$mNAIL^{WT}$ DSS -vs- $mNAIL^{WT}$ UT	1406
$mNAIL^{\Delta NF\kappa B}$ DSS -vs- $mNAIL^{WT}$ DSS	1007

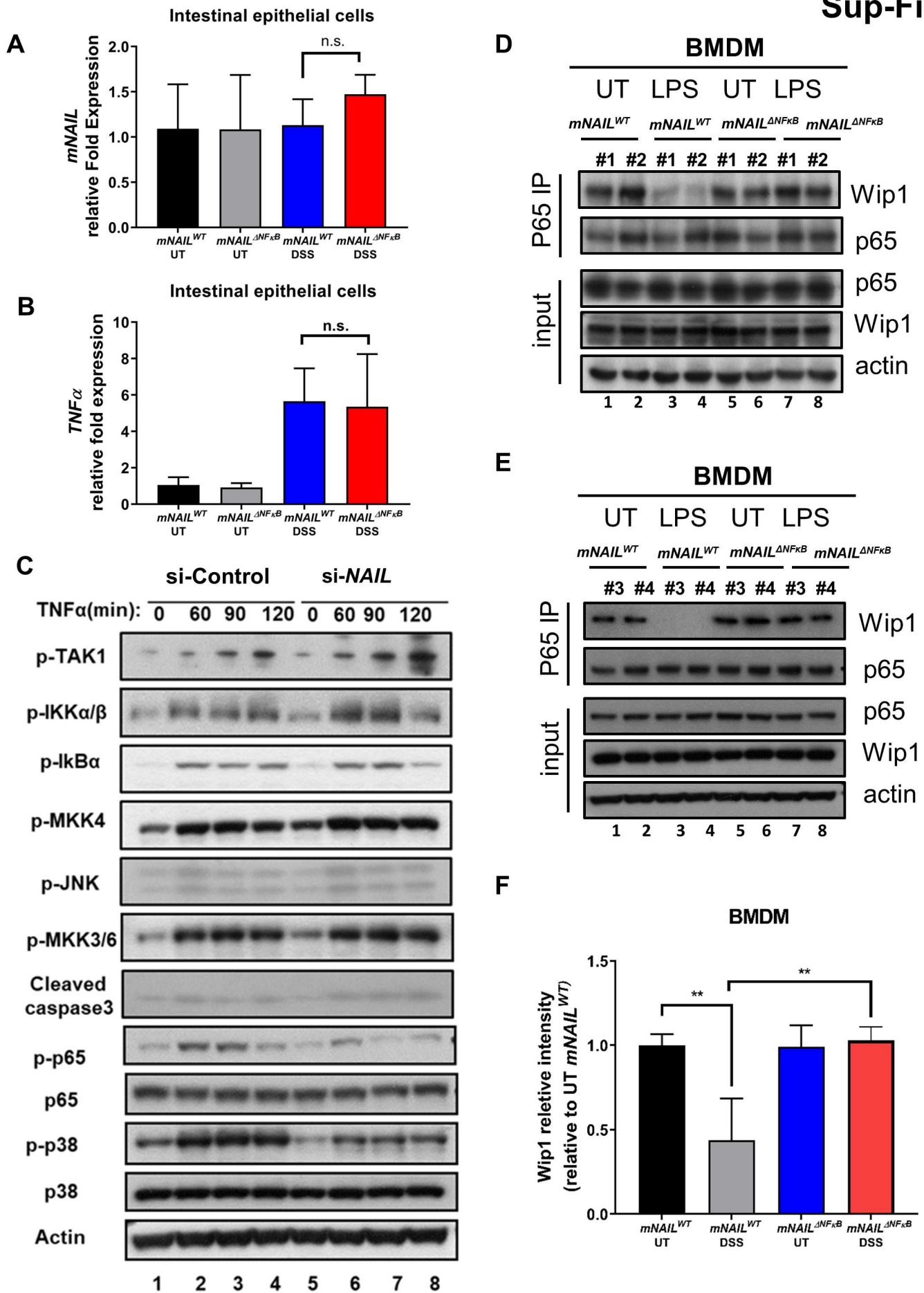
B



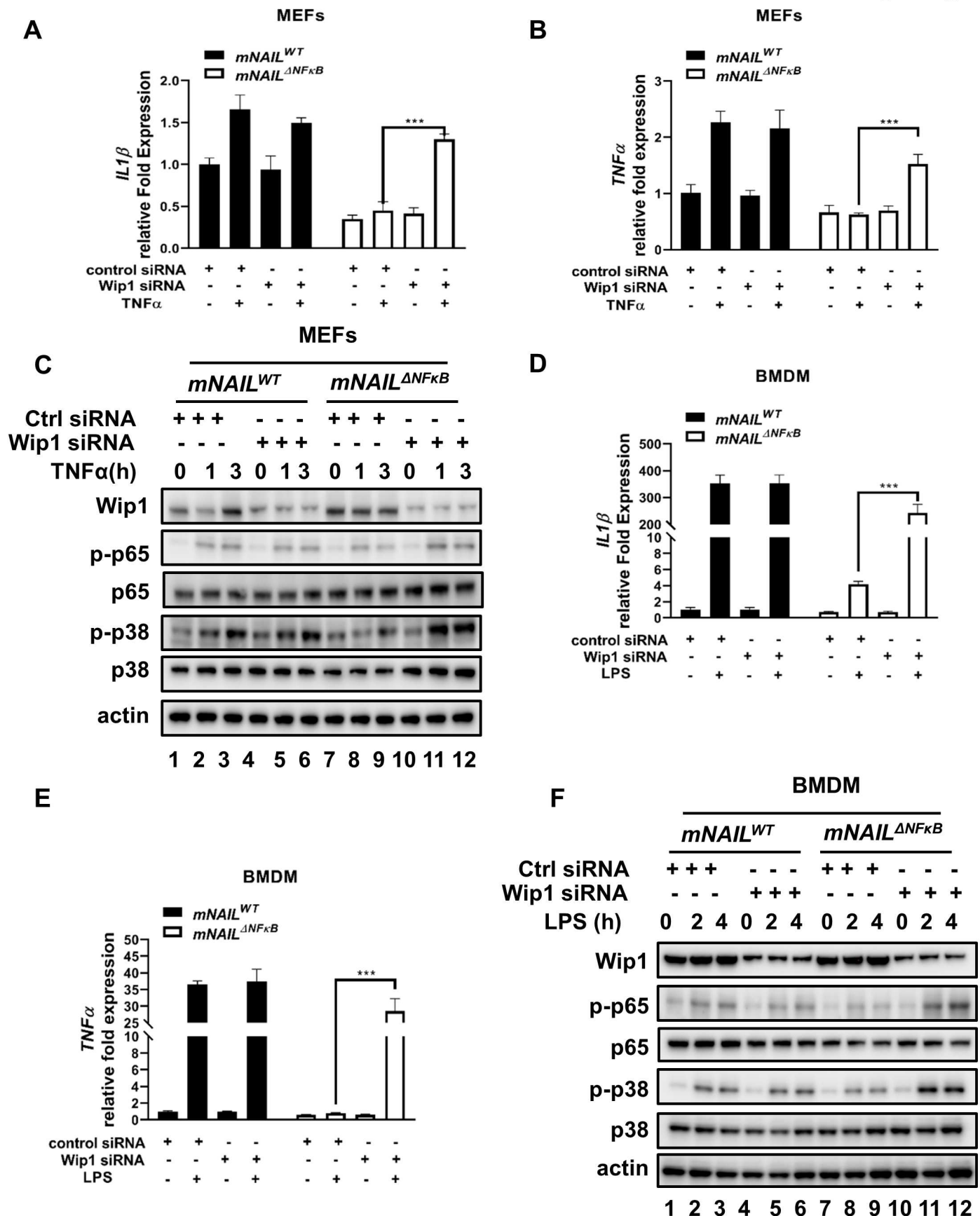
Sup-Fig. 6



Sup-Fig. 7



Sup-Fig. 8



Sup-Table 1

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
anti-p-p38 (Thr180/Tyr182) 3D7	Cell signalling	Cat #9215S
anti-p38	Cell signalling	Cat #9212
anti-p-p65 (Ser536)	Cell signalling	Cat #3031L
anti-p65	Santa Cruz	Cat #sc-8008
anti-Actin	Sigma	Cat #A2066
GAPDH (6C5)	Santa Cruz	Cat #sc-32233
p-IKK α / β (Ser176/180)	Cell signalling	Cat #2697S
IKK α / β (H-470)	Santa Cruz	Cat #sc-7607
IKK γ (FL-419)	Santa Cruz	Cat #sc-8330
anti-p65 (ChIP, IP)	Santa Cruz	Cat #sc-372
anti-Wip1 (RIP)	Cell signalling	Cat #11901
anti-Wip1(WB)	Santa Cruz	Cat # sc-376257
Dyskerin (H-300)	Santa Cruz	Cat #sc-48794
anti-IL7	R&Dsystems	Cat # AF407
P-SEK1/MKK4(T261)	Cell signalling	Cat # 9151S
anti-SEK1/MKK4	Cell signalling	Cat # 9152S
p-TAK1 (T187)	Cell signalling	Cat # 4536S
p-IkBa (S32)	Cell signalling	Cat # 2859L
p-SAPK/JNK (Thr183/Tyr185)	Cell signalling	Cat # 9251
anti-P-MKK3/MKK6 (Ser189/207)	Cell signalling	Cat # 9231
anti-Cleaved Caspase-3 (Asp175)	Cell signalling	Cat #9664L
PE-CD11b	BD	Cat #553311
FITC-Ly6G	Biolegend	Cat #127606
BV711-Ly6C	Biolegend	Cat #128037
PE-CD34	BD	Cat #551387
PeCy7-IL7RA	BD	Cat #560733
PercepCy5.5-c-kit	BD	Cat #560557
FITC-Sea1/Anti-Ly-6A/E	BD	Cat #553335
APCcy7-CD16/32	BD	Cat #560541
APC Mouse Lineage Antibody Cocktail	BD	Cat #558074

AF488-p-NF- κ B p65 (Ser536) (93H1)	CST	Cat #4886
AF488-p-p38 MAPK (Thr180/Tyr182) (3D7)	CST	Cat #41768
Chemicals, recombinants proteins, commercial kits		
Recombinant Human TNF-alpha Protein 210	R&Dsystems	Cat #210-TA
Terminal deoxynucleotidyltransferase	Promega	Cat #M1871
RQ1 RNase-Free DNase	Promega	Cat #M6101
Recombinant RNasin ribonuclease inhibitor	Promega	Cat #N2111
T7 RNA polymerase	Promega	Cat #P2075
10X Biotin RNA labelling mix	Roche	Cat #11685597910
Tak1 inhibitor 5Z-7-Oxozeaenol	Sigma	Cat #O9890-1MG
E. coli LPS	Sigma	Cat #L2654 and #L2630
Dextran sulfate sodium salt, colitis grade	MP Biomedicals	Cat # 9011-18-1
Disuccinimidyl glutarate (DSG)	Thermo Fisher	Cat #20593
X-tremeGENE 9 DNA transfection reagent	Roche	Cat #6365779001
X-tremeGENE siRNA transfection reagent	Roche	Cat #4476093001
Experimental Models: Organisms/Strains		
Mouse: wild-type: C57BL/6	Invivos	N/A
Mouse: wild type Balb/c	Invivos	N/A
Mouse: <i>mNAIL^{ANFKB}</i>	Masa's group	This study
MEFs: WT, p65 ^{-/-} , Ikk β ^{-/-} and Ikk γ ^{-/-}	Tergaonkar et al., 2003	N/A
Oligonucleotides		
RT-qPCR_Gm16685 Fw (CCCTACAGGAAGACCAACCA)	IDT	N/A
RT-qPCR_Gm16685 Rv (CTAGGCCATCCTCTGCTACG)	IDT	N/A
RT-qPCR_mTNFa Fw (ACAGAAAGCATGATCCGCGAC)	IDT	N/A
RT-qPCR_mTNFa Rv (GAAGCCCCCATCTTTGG)	IDT	N/A
RT-qPCR_mIL7 Fw (TGGAATTCCTCCACTGATCC)	IDT	N/A
RT-qPCR_mIL7 Rv (TGGTTCATTATTCGGGCAAT)	IDT	N/A
RT-qPCR_mActin Fw (CTGTATCCCCTCCATCGTG)	IDT	N/A
RT-qPCR_mActin Rv (CCTCGTCACCCACATAGGAG)	IDT	N/A

RTqPCR_mCCL2 Fw (GCCCTCCATGTATACCAGACT)	IDT	N/A
RTqPCR_mCCL2 Rv (AGACCTCTCTTGAGCTTGGT)	IDT	N/A
RT-qPCR_mIL1 β Fw (AAAGCTCTCCACCTCAATGG)	IDT	N/A
RT-qPCR_mIL1 β Rv2 (TCTTCTTGGGTATTGCTGG)	IDT	N/A
CRISPR Genotyping_Gm16685 Fw (GCATTCCCTTAGGTAGACTCC)	IDT	N/A
CRISPR Genotyping_Gm16685 Rv (GGGAGTGATTATGGGTGGTGAG)	IDT	N/A
Gm16685 gRNA #1 (CACCTAGGGTTAAAAGCGCATCC)	IDT	N/A
Gm16685 gRNA #2 (CACCAGTCTGGGAGTTTCCGATCC)	IDT	N/A
Gm16685 siRNA (GGAATAATGATAGCAACTACT)	IDT	N/A
Gm16685 siRNA (CAACCTCTCTAATCAGTCTCTTTCT)	IDT	N/A
Wip1 siRNA (CTTGTGGGGTTTCATCAAGAAGCAG)	IDT	N/A
Mouse IL7 transcript variant 4 siRNA#1 (AGCTCAGAACTTCATGGAGAATGAA)	IDT	N/A
Mouse IL7 transcript variant 4 siRNA#1 (AAAGGACAGAUCAUUGGAAAACCCC)	IDT	N/A
Mouse IL7 siRNA#1 (GACAGGAAGTATAGTAAT)	IDT	N/A
Mouse IL7 siRNA#2 (GCATATGAGAGTGTACTGA)	IDT	N/A
Plasmids		
pX458 –GFP	Addgene	Cat #48138
pX458-DsRed	Akincilar et al., 2016	N/A