The deubiquitinase USP13 stabilizes the anti-inflammatory receptor IL-1R8/Sigirr to suppress lung inflammation

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ABSTRACT

Background: The Single immunoglobulin interleukin-1 (IL-1)-related receptor (Sigirr), also known as IL-1R8, has been shown to exhibit broad anti-inflammatory effects against inflammatory diseases including acute lung injury, while molecular regulation of IL-1R8/Sigirr protein stability has not been reported. This study is designed to determine whether stabilization of IL-1R8/Sigirr by a deubiquitinating enzyme (DUB) is sufficient to suppress inflammatory responses and lessen lung inflammation.

Methods: A molecular signature of ubiquitination and degradation of IL-1R8/Sigirr was determined using a receptor ligation chase model. The anti-inflammatory effects on USP13 were investigated. USP13 knockout mice were evaluated for stabilization of IL-1R8/Sigirr and disease phenotype in an acute lung injury model.

Findings: IL-1R8/Sigirr degradation is mediated by the ubiquitin-proteasome system, through site-specific ubiquitination. This effect was antagonized by the DUB USP13. USP13 levels correlate directly with IL-1R8/Sigirr, and both proteins were reduced in cells and tissue from mice subjected to inflammatory injury by the TLR4 agonist lipopolysaccharide (LPS). Knockdown of USP13 in cells increased IL-1R8/Sigirr polyubiquitination and reduced its stability, which enhanced LPS-induced TLR4 signaling and cytokine release. Likewise, USP13-deficient mice were highly susceptible to LPS or Pseudomonas aeruginosa models of inflammatory lung injury. IL-1R8/Sigirr overexpression in cells or by pulmonary viral transduction attenuated the inflammatory phenotype conferred by the USP13−/− genotype.

Interpretation: Stabilization of IL-1R8/Sigirr by USP13 describes a novel anti-inflammatory pathway in diseases that could provide a new strategy to modulate immune activation.

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1. Introduction

Sigirr (known as IL-1R8) belongs to the IL-1R superfamily [1]. Distinct from classic IL-1R family members, Sigirr consists of only a single extracellular Ig-like domain, a long cytoplasmic tail, and an intracellular Toll/IL-1R (TIR) domain [1]. Sigirr negatively regulates IL-1 and TLR signaling [1–7] and broadly dampens inflammatory responses during acute lung injury [8], brain inflammation [9–11], arthritis [12], colitis [2,13], and pulmonary tuberculosis [14]. Though IL-37 has been identified as a Sigirr ligand [15], the anti-inflammatory effect of Sigirr seems to be independent of ligation by IL-37 [8]. While Sigirr gene expression has been shown to be regulated by the transcriptional factor Sp1 [16], the molecular regulation of Sigirr protein stability has not been reported. Understanding the regulation of Sigirr protein turnover could provide a mechanistic basis for the modulation of Sigirr-mediated anti-inflammatory signaling.

Protein turnover is regulated by post-translational modification such as mono- or poly-ubiquitination [17]. The attachment of polyubiquitin chains to substrate proteins serves as a molecular tag for their degradation. Polyubiquitination is characterized as linear based on the type of ubiquitin chain linkage. Generally, Lysine 48 (K48)-linked ubiquitination leads to protein degradation in the proteasome [18].
2.1. Generation of USP13 deficient mice

The Usp13<sup>−/−</sup> mice were generated by the CRISPR/Cas9 system at the University of Pittsburgh. Exon 6 and Intron 18 of Usp13 (chromosome 3 between position 32,865,806 and 32,917,828) were deleted. Only the Usp13 gene is localized in the position on chromosome 3 (https://www.ncbi.nlm.nih.gov/genome/gdv/browser/?context=genomic&acc=GCF_000001635.26). In brief, Cas9 mRNA and two sgRNA were injected into the fertilized embryos, and then embryos in 2-cell stages were transferred to oviducts of pseudopregnant female mice. The RNA sequence guides are GTTGCCCGATGTGACCTGC and TCGAGTGGACTATGACACA. The potential founder F0 mice were genotyped based on genomic DNA isolated from mouse tails by PCR with the following primer sets: F52: CAGGCT CATTGTCATGACACA, and R52: ACTGACTGCTGTCACAGAA. A 481 bp or an approximately 600 bp fragment was produced from the WT allele or the null allele, respectively. Chimeric offspring were crossed with C57BL/6 to generate Usp13<sup>+/−</sup> mice. The F1 Usp13<sup>+/−</sup> mice were further crossed with C57BL/6 background for at least 6 generations before use. Usp13<sup>−/−</sup> mice identified by genotyping through PCR were intercrossed for the generation of Usp13<sup>−/−</sup> mice. Sex-matched Usp13<sup>+/−</sup> and Usp13<sup>−/−</sup> littermates at 8–10 weeks were used for animal studies.

2.2. LPS- or P. aeruginosa-induced murine models of lung injury

All mice were housed in the specific pathogen-free animal care facility at the University of Pittsburgh and the Ohio State University in accordance with institutional guidelines and guidelines of the US National Institutes of Health. All animal experiments were approved by an institutional animal care and use committee (IACUC) at the University of Pittsburgh and the Ohio State University. Lung inflammation models were performed by intratracheal administration of LPS or P. aeruginosa. Briefly, mice were given intratracheal administration of LPS (1 mg per kg body weight) or P. aeruginosa (strain PA103; 1 × 10<sup>9</sup> colony-forming units per mouse). At designated time points after LPS or PA103 challenge, the mice were anesthetized before myocardiual perforations were performed with PBS via the right ventricle until lungs were cleared of blood, and then lungs were harvested for further analyses. For BAL collection, the lungs were lavaged twice with 1 ml sterile PBS at room temperature. The cell-free supernatants were harvested for ELISA assay after centrifuging at 1000 rpm for 5 min. The cell pellets were diluted in 1 ml sterile PBS, and the cells were counted with a hemocytometer. Cytospin preparations of BAL cells were stained with hematoxylin and eosin and viewed under light microscopy for inflammatory cell differential. For lentiviral vector delivery system, cDNA encoding human Sigirr was inserted into the pLX-IRESTomato vector (Clontech, Palo Alto, CA, USA); lentiviral vectors encoding Sigirr and their controls were generated with a lentivirus packaging system (Clontech, Palo Alto, CA, USA). C57BL/6 mice were given 50 µl lentivirus vectors (2 × 10<sup>7</sup> plaque-forming units per mouse) via intratracheal administration for 5 d before intratracheal challenge with LPS or PA103 (doses described above).

2.3. H&E staining and immunohistochemistry

The left lungs from animals were inflated with 0.5 ml of 2% PFA after clearing of blood. The lung tissues were then fixed overnight, embedded in paraffin. The sections (5 µm thick) were cut and used for staining with hematoxylin and eosin to assess the degree of lung injury. Immunohistochemistry was performed as described below. In brief, sections were deparaffinized and rehydrated through graded alcohol. Antigen retrieval was performed by high-pressure heating with citrate buffer (Thermo Scientific, Fremont, CA, USA), then tissues were incubated with different antibodies at 4 °C overnight and HRP-polymer secondary antibodies (Santa Cruz Biotechnology, Santa Cruz, CA, USA) for 15 min and then incubated and developed using DAB solution (Santa Cruz Biotechnology, Santa Cruz, CA, USA). The antibody specific for USP13 or IL-1β was used for staining. Images were captured by EVOS inverted microscope.

2.4. Cells and reagents

Mouse lung epithelial cells (MLE12, SV40-immortalized mouse alveolar cell line with epithelial characteristics), RAW 264.7 cells (mouse macrophage-like virus-induced leukemia cell line), and BEAS-2B (human lung epithelial cell line) were obtained from American Type Culture Collection (ATCC, Manassas, VA). Human macrophage-like cells (monocyte-derived macrophages differentiated in vitro with M-CSF and IL-4) were purchased from StemCell Technologies (Vancouver, BC, Canada). Human TLR4 stable transfected HEK293 (HEK293/TLR4) cells were from InvivoGen (San Diego, CA, USA). Recombinant human IL-37 (a truncated form, Lys27-Asp192) was from R&D Systems (Minneapolis, MN, USA). Cycloheximide, leupeptin, and LPS were from Sigma Aldrich (St Louis, MO, USA). MG-132 was from EMD Chemicals (Gibbstown, NJ, USA). Immobilized protein A/G beads and control IgG,
anti-Sigirr (for mouse, rat, and human origin), anti-phospho-JNK1, anti-JNK1, anti-NF-κB p65, anti-GAPDH, anti-IL-1β, and anti-Lamin A/C and anti-TLR4 were from Santa Cruz Biotechnology (Santa Cruz, CA, USA). Anti-USP13 and anti-IκB were purchased from ProteinTech (Chicago, IL, USA). Anti-USP13 was purchased from Bethyl Laboratories (Montgomery, TX, USA). Antibodies against phospho-p38, p38, phospho-IκB, HA tag, K48 ubiquitin, K63 ubiquitin, and ubiquitin were from Cell Signaling (Beverly, MA, USA). Anti-V5, mammalian expression plasmid pcDNA3.1/His-V5-topo and Sigirr reverse, TCACACCGACCTTCAC, as well as hSigirr forward, TGTCGCAAGGCTGTGTACTT, were inserted into pCDNA3.1/V5-His-Topo vector (Invitrogen, Carlsbad, CA, USA). The sequences of specific primer pairs are described above: Sigirr forward, CACCATCGCAAGGCTGTGTACTT; Sigirr reverse, AACCTGGGCAACAGCAGCTATCTCAGC. Plasmid transfection in MLE12 cells was performed with Lonza electroporation according to the manufacturer’s protocol. Lipofect™ reagent was used for transfection of plasmids into RAW 264.7 cells and HEK293/TLR4 cells according to manufacturer instructions (Life Technologies, Gaithersburg, Maryland, USA). Cellular transfection of siRNA in BEAS-2B cells was done with GeneMute siRNA transfection reagent system.

2.5. Plasmid and siRNA transfection

The cDNA encoding human Usp13 or human Sigirr and its truncations were inserted into pCDNA3.1/V5-His-Topo vector (Invitrogen, Carlsbad, CA, USA). The sequences of specific primer pairs are described below: Sigirr forward, CACCATCGCAAGGCTGTGTACTT; Sigirr reverse, AACATATCATCCTTGGACACCA, Sigirr (1-160) reverse, GTTATATCCACCTCCCAT, Sigirr (1-290) reverse, GGGCTCTCCAGACGCAAGG, Sigirr (1-374) reverse, CACCAGTGGTAGTCGGTGGAG, Usp13 forward, CACCATGTACGAGCCGCGCCGGCTC, Usp13 reverse, GCTTGGATATCTCGGGTAA. Usp13C345A mutant plasmid was carried out using QuikChange Lightning Site-Directed Mutagenesis Kit (Agilent Technologies, La Jolla, CA). The sequences of specific primer sets are described below: Usp13C345A forward, GAGACAGCTAGATGCTGTTGACCA, and Usp13C345A reverse, AACCTGGGCAACAGCAGCTATCTCAGC. Plasmid transfection in MLE12 cells was performed with Lonza electroporation according to the manufacturer’s protocol. Lipofect™ reagent was used for transfection of plasmids into RAW 264.7 cells and HEK293/TLR4 cells according to manufacturer instructions (Life Technologies, Gaithersburg, Maryland, USA). Cellular transfection of siRNA in BEAS-2B cells was done with GeneMute siRNA transfection reagent system.

2.6. qRT-PCR analysis

RNA isolation and qRT-PCR analysis were performed as previously described [24]. The expression of Usp13 and Sigirr was performed using iQ SYBR Green Supermix and the iCycler real-time PCR detection system (Bio-Rad, Hercules, CA, USA). GAPDH was used as an internal control. The sequences of specific primer pairs are described below: hUsp13 forward, TGGTCGAAAGGTGTGATCAG, hUsp13 reverse, CACCGGCTTCAGCAAAATCTG; hSigirr forward, GAAGCTCTAGATGGCTGTTGACCA, and hSigirr reverse, TCTGGGGTGCTCTCTCAAGC; hGAPDH forward, TGGAGGTCACACGATCATCCTCC, and hGAPDH reverse, GCCCTCTGAGACATCATCCTCC. Usp13 forward, AGTGGAGGTCACACGATCATCCTCC, and Usp13 reverse, AGTGGAGGTCACACGATCATCCTCC; mSigirr forward, CAGCTGCTGGTGTTCGACT, and mSigirr reverse, GTCTGCCAAGAGGTGAAGGA; mGAPDH forward, ACCCTTAAAGAGGATGCTGCC, and mGAPDH reverse, TCACACCGAATCCTGAC. 2.7. Immunostaining

MLE12 cells were grown in glass bottom dishes until 80–90% confluence. After treatment or transfection, cells were fixed with 3.7% formaldehyde for 20 min, blocked with 1% BSA in TBST for 30 min, then immunostained with HA tag or V5 tag antibody for 1 h followed by washes with PBS for three times, and incubated with the fluorescent probe-conjugated secondary antibodies. Images were captured by a Nikon ECLIPSE TE 300 inverted microscope.

2.8. Western blotting analysis

The proteins for cells or lung tissues were extracted using cell lysis buffer following standard protocols as described before [25]. The cell lysates were then sonicated on ice for 12 s, followed by centrifugation at 4 °C at 5000 rpm for 10 min to remove insoluble fragments. Protein concentrations were determined with a Bio-Rad Protein Assay Kit (Bio-Rad Laboratories, Inc., Hercules, CA, USA). Equal amounts of samples were subjected to SDS-PAGE gel. Blots were washed with 25 mM Tris HCl (pH 7.4), 137 mM NaCl, and 0.1% Tween 20 (TBST) and incubated with a primary antibody for 2 h or overnight. The membranes were then washed three times at 10 min intervals with TBST prior to the addition of a secondary antibody for 1 h. Blots were developed with an Enhanced Chemiluminescence Detection Kit (ThermoFisher Scientific, Waltham, MA) according to the manufacturer’s instructions. Western signalizing was detected by Imagej software (Image J, National Institutes of Health, Bethesda, MD, USA; http://imagej.nih.gov/). Comparisons between several groups were analyzed using one-way or two-way ANOVA. Data are expressed as mean ± SEM of triplicate samples from at least three independent experiments.
experiments and values that were p < 0.05 were considered statistically significant.

2.13. Enzyme-linked immunosorbent assay (ELISA)

Levels of TNF-α, IL-1β, IL-6, and IL-8 proteins in the BAL and cell supernatant were measured with commercial ELISA kits (eBioscience, San Diego, CA, USA) according to the manufacturers’ instructions.

3. Results

3.1. Sigirr is degraded in the proteasome

Most cell surface receptors are post-translationally modified, internalized, and degraded upon ligation as a primary control for receptor abundance and intracellular signaling [26–28]. IL-37 has been identified as Sigirr’s only ligand [1]. We tested if IL-37 modulates Sigirr abundance. Treatment of MLE12 mouse lung epithelial cells or human macrophage-like cells with IL-37 decreased total Sigirr protein levels but not IL-18R1 as assessed by immunoblotting (Fig. 1a, b). Flow cytometry analysis with an anti-Sigirr antibody revealed that IL-37 reduced its abundance on the cell surface over time (Supplemental Fig. S1a). IL-37 also significantly reduced overexpressed Sigirr-V5 in a dose-dependent manner (Supplemental Fig. S1b). Notably, Sigirr mRNA levels were not reduced by IL-37 in 2 h, while at 4 h, Sigirr mRNA level was significantly increased (Supplemental Fig. S1c), possibly as a compensatory response to reduced protein levels. These results indicate that Sigirr internalization and degradation by IL-37 is a suitable molecular model for investigating the molecular mechanisms of Sigirr degradation. To examine if the proteasome or lysosome pathways were involved in the degradation of Sigirr, we incubated MLE12 cells with inhibitors of the proteasome (MG-132) or lysosome (leupeptin) activity prior to IL-37 treatment. MG-132, but not leupeptin, prevented Sigirr degradation (Fig. 1c, Supplemental Fig. S1d). After IL-37 treatment, the transfected Sigirr-V5 protein product did not localize in the lysosomes in MLE12 cells (Supplemental Fig. S2a), suggesting that Sigirr degradation does not occur in the lysosome. In addition, the half-life of Sigirr (~4 h) in the presence of the protein synthesis inhibitor cycloheximide (CHX) was prolonged by MG-132, but not by leupeptin (Supplemental Fig. S2b), further implying that proteasomal degradation is the major pathway of cellular Sigirr elimination.

Polyubiquitination can be characterized by Lys (K)48-, K63-, and other linkages of ubiquitin chains based on which lysine residue of ubiquitin is bound to a growing ubiquitin chain [29]. Next, we examined Sigirr ubiquitination with a linkage-specific polyubiquitin pulldown and revealed that the addition of K48- and not K63-linked ubiquitin chain is the predominant mode of Sigirr modification (Fig. 1d). Ubiquitin chains are usually added to lysine residues on substrate proteins [30]. To identify putative ubiquitin-acceptor sites within Sigirr, we replaced several K candidate residues of human Sigirr with arginine (R). These K-to-R mutations identified the K163 residue of Sigirr as a putative ubiquitin acceptor residue for IL-37-mediated modification, as SigirrK163R was not degraded and displayed a reduction in ubiquitination compared to WT or other Sigirr K-to-R mutants (Fig. 1e, f). K163 seems to be localized in a surface-exposed loop in the intracellular domain of Sigirr (Fig. 1g). Notably, this ubiquitin site

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*Fig. 1.* Sigirr degradation is mediated by the ubiquitin-proteasome system. (a, b) MLE12 (a) or human macrophage-like cells (b) were treated with IL-37 (100 ng/ml) for the indicated times with immunoblots to examine the protein levels of endogenous Sigirr, IL-18R1, and β-actin. (c) MLE12 cells were pretreated with MG-132 (20 μM) or leupeptin (100 μM) for 2 h and then treated with IL-37 (100 ng/ml) for the indicated times. (d) MLE12 cells were treated with IL-37 for the indicated times. *In vitro ubiquitination assay* was performed with a modified immunoprecipitation (IP). Smear bands indicate Ubi(n). (e) MLE12 cells were transfected with plasmids encoding Sigirr-V5, K196R-V5 mutant, or K163R-V5 mutant for 48 h and then treated with IL-37 for the indicated times. (f) MLE12 cells were transfected with plasmids encoding Sigirr-V5 and its mutant K163R-V5 for 48 h and then treated with IL-37 for 1 h. Denatured cell lysates were subjected to IP with a V5 antibody, followed by immunoblotting with an antibody against ubiquitin (Ubi). Smear bands indicate Ubi(n). (g) Predicated Sigirr protein structure analysis by Swiss-Model reveals that K163 is localized on the protein surface. (h) The scheme shows that IL-37 induces Sigirr degradation in the ubiquitin-proteasome system. Sigirr is polyubiquitinated by a K48-linked chain on the K163 residue. (For all panels, all data are mean ± SEM. n = 3 *p < 0.05; **p < 0.01 by one-way (a, b) or two-way (c-e) ANOVA with Post hoc Tukey’s test compared to 0 h or control. Representative blots from three independent experiments are shown. Bands densitometry were analyzed with ImageJ.)
mutation of Sigirr (SigirrK163R) had no effect on ligand-induced Sigirr internalization (Supplemental Fig. S3a, b), suggesting that ubiquitination at K163 is not necessary for Sigirr internalization. These data suggest that Sigirr can be ubiquitinated by K48-linked ubiquitin chains on the K163 residue in response to IL-37 ligation, resulting in its proteasomal degradation (Fig. 1h).

3.2. USP13 deubiquitinates and stabilizes Sigirr

Since DUBs remove ubiquitin chains, they can oppose degradation signals to stabilize substrate proteins. We performed a screen to identify a DUB responsible for stabilizing Sigirr (Supplemental Fig. S4). Only ectopic expression of V5-tagged USP13 (USP13-V5) significantly diminished degradation of endogenous or exogenous Sigirr (Fig. 2a, b). Three separate short hairpin RNAs (shRNAs) targeting mouse USP13 mRNA effectively reduced Sigirr protein levels by ~70–90% (Fig. 2c), and likewise, human USP13 siRNAs also decreased Sigirr protein by ~80% in BEAS-2B human epithelial cells (Fig. 2d). Knockdown of USP13 did not affect Sigirr mRNA transcript levels (Fig. 2e), indicating that USP13 stabilizes Sigirr protein without altering its gene expression. Overexpression of USP13-V5 attenuated Sigirr ubiquitination in MLE12 cells (Fig. 2f), while knockdown of USP13 with shRNA (Fig. 2g) enhanced Sigirr ubiquitination, suggesting that USP13 deubiquitinates Sigirr to stabilize the protein.

3.3. USP13 is associated with Sigirr

Next, we evaluated USP13-Sigirr association using co-immunofluorescence staining. MLE12 cells were co-transfected with HA-tagged Sigirr (Sigirr-HA) and USP13-V5. Sigirr-HA and USP13-V5 were highly co-localized on the inside of the cell membrane with some puncta also observed in the cytoplasm (Fig. 3a). Co-immunoprecipitation (Co-IP) also revealed USP13’s association with Sigirr (Fig. 3b, c). The association between DUB and a transmembrane receptor protein can be regulated by ligation [28]. As shown in Fig. 3c, the complex of USP13 and Sigirr is disrupted after IL-37 treatment, indicating that IL-37 increases Sigirr ubiquitination through the reduction of its binding to USP13. To further identify the USP13-binding region on Sigirr, we incubated purified USP13-HA together with a series of C-terminal deletion mutants of Sigirr-V5 (Fig. 3d). The Co-IP assays revealed that the 290-374aa region in the C-terminal of Sigirr is required for its association with USP13 (Fig. 3e).
3.4. USP13 dampens LPS-induced inflammatory responses via the stabilization of Sigirr

To investigate the biological function of USP13 in inflammatory responses, we first confirmed the anti-inflammatory activity of Sigirr. Overexpression of Sigirr attenuated LPS-induced phosphorylation of JNK, ERK, and IL-6 gene expression (Supplemental Fig. S5a, b). We then modulated USP13 and observed downstream inflammatory signals in a cellular model. LPS induces NF-κB and MAPK signaling to mediate inflammatory signaling and cytokine release. Overexpression of USP13 attenuated LPS-induced phosphorylation and degradation of IκB, phosphorylation of p38 (Fig. 4a), as well as LPS-induced IL-6 release in RAW 264.7 cells (Fig. 4b). Downregulation of USP13 with shRNA or siRNA promoted LPS-induced phosphorylation and degradation of IκB, activation of p38 (Fig. 4c, d), NF-κB p65 subunit nuclear translocation (Fig. 4e), and IL-8 release (Fig. 4f). To determine if the inhibitory effect of USP13 on LPS signaling is dependent on its activity, we overexpressed a catalytically inactive mutant of USP13 (C345A) in 293TLR4 cells. As shown in Supplemental Fig. S5c, USP13C345A did not attenuate LPS-induced phosphorylation of p38 but rather increased p38 phosphorylation, suggesting that the enzyme activity of USP13 is required for inhibiting LPS signaling. These data indicate that USP13 exhibits anti-inflammatory activity in LPS-exposed cells.

3.5. USP13 is reduced in response to endotoxin exposure in vitro and in vivo

Following LPS exposure, USP13 and Sigirr, but not TLR4, were both depleted over time in RAW 264.7 cells and MLE12 cells (Fig. 5a, b). Importantly, we noticed that there were faint bands close to the molecular weight of USP13 in lung tissue lysates from USP13−/− mice. It is possible that the faint bands are from non-specific detections or an undiscovered USP13 splicing mutant. USP13 deficient mice exhibit no significant phenotype differences compared to wild type mice in the murine models of acute lung injury, USP13 and Sigirr levels were likewise decreased in mice lung homogenates after intratracheal challenge with either LPS or P. aeruginosa (strain PA103) (Fig. 5c, d). Immunohistochemical staining from these experiments revealed reductions in USP13 in the lung epithelium, which correlated with increased inflammatory injury indicated by H&E staining and IL-1β abundance (Fig. 5e). These data suggest that USP13 is downregulated during acute inflammatory injury. We evaluated possible molecular mechanisms of LPS-induced USP13 downregulation and observed that USP13 mRNA levels were reduced by LPS (Fig. 5f), suggesting that LPS stimulation diminishes USP13 transcription. We then tested several downstream pathways, and pharmacologic blockade of NF-κB, JNK, and p38 pathways did not affect the LPS-mediated reduction of USP13 (Fig. 5g). Further studies will be needed to determine if and how LPS might accelerate USP13 mRNA degradation or cause epigenetic modification to the USP13 promoter or promoter-related histone subunits (e.g., methylation or acetylation).

3.6. Usp13 deficient mice exacerbate LPS or Pseudomonas aeruginosa-induced lung inflammation

To investigate the relevance of USP13 in the inflammatory response, we used CRISPR/Cas9 to generate Usp13 deficient mice (Supplemental Fig. S6a, b). We noticed that there were faint bands close to the molecular weight of USP13 in lung tissue lysates from Usp13−/− mice. It is possible that the faint bands are from non-specific detections or an undiscovered Usp13 splicing mutant. Usp13 deficient mice exhibit no significant phenotype differences compared to wild type mice in the murine models of acute lung injury. USP13 and Sigirr levels were likewise decreased in mice lung homogenates after intratracheal challenge with either LPS or P. aeruginosa (strain PA103) (Fig. 5c, d). Immunohistochemical staining from these experiments revealed reductions in USP13 in the lung epithelium, which correlated with increased inflammatory injury indicated by H&E staining and IL-1β abundance (Fig. 5e). These data suggest that USP13 is downregulated during acute inflammatory injury. We evaluated possible molecular mechanisms of LPS-induced USP13 downregulation and observed that USP13 mRNA levels were reduced by LPS (Fig. 5f), suggesting that LPS stimulation diminishes USP13 transcription. We then tested several downstream pathways, and pharmacologic blockade of NF-κB, JNK, and p38 pathways did not affect the LPS-mediated reduction of USP13 (Fig. 5g). Further studies will be needed to determine if and how LPS might accelerate USP13 mRNA degradation or cause epigenetic modification to the USP13 promoter or promoter-related histone subunits (e.g., methylation or acetylation).

Fig. 3. USP13 is associated with Sigirr. (a) Sigirr-HA and USP13-V5 co-overexpressed MLE12 cells were double-immunostained with V5 and HA antibodies. Scale bar, 5 μm. Yellow (shown by arrows) indicates co-localization. (b) Overexpressed USP13-V5 in MLE12 cells were co-immunoprecipitated with a Sigirr antibody and then analyzed by immunoblotting with a V5 antibody. Input cell lysates were immunoblotted with V5 and β-actin antibodies. (c) MLE12 cells were treated with or without IL-37 for 1 h. Cell lysates were assessed by IP with an antibody against Sigirr and immunoblot analysis of precipitates with antibodies against USP13 and Sigirr. Input lysates were analyzed with antibodies against USP13 and β-actin. (d) Schematic representation of V5-tagged full-length Sigirr (1–410 aa) and two C-terminal deletion mutants. Ig: immunoglobulin; TIR: the Toll/IL-1R domain. (e) Control vector (V), USP13-HA, and V5-tagged Sigirr or its C-terminal deletion mutants were synthesized with a TnT translation system. USP13-HA was incubated with Sigirr-V5 or its mutants, followed by IP with anti-V5 beads and immunoblotted with an antibody to HA. Input lysates were analyzed by antibodies to HA or V5 tag. The arrows indicate overexpressed Sigirr-V5 and mutants. A star (*) indicates a non-specific band. For all panels, Representative blots or images from two (e) or three (all others) independent experiments are shown.
normal condition. Consistent with our cellular USP13 knockdown studies (Fig. 2c, d), Sigirr levels in lungs from Usp13−/− mice were significantly reduced (Supplemental Fig. S6b).

Based on the in vitro data, we hypothesized that Usp13−/− mice would display enhanced inflammation, and we tested this with intratracheal administration of LPS or *Pseudomonas aeruginosa* (strain PA103) to Usp13−/− or Usp13+/− (WT) mice. Compared to WT mice, Usp13−/− mice displayed a significantly greater cellular infiltration, particularly with neutrophils, 6 h after LPS or 4 h after *P. aeruginosa* challenge (Fig. 6a–d). Usp13−/− mice also displayed increased inflammatory injury shown by H&E staining (Fig. 6e, f), and enhanced TNF-α, IL-1β, and IL-6 (Fig. 6g–i) release in BAL fluid. These data demonstrate that the deletion of USP13 promotes lung inflammation in murine models of lung injury. Taken together, this study indicates that the reduction of USP13 in the setting of inflammation may partly contribute to the pathogenesis of lung injury.

### 3.7 Sigirr reverses LPS-induced acute lung inflammation in Usp13−/− mice

To investigate if the enhanced inflammation observed in Usp13−/− mice is due to the dysregulation of Sigirr, we constructed a lentiviral vector encoding a fusion of Sigirr (LentiSigirr) and inoculated WT or Usp13−/− mice with this construct or an empty control lentiviral vector (LentiCont) by intratracheal administration 5 d prior to intratracheal LPS injection. Expression of Sigirr by LentiSigirr infection in lung tissues was confirmed by Western blotting (Supplemental Fig. S7). The LentiSigirr transduction effectively dampened the enhanced LPS-induced lung inflammation phenotype observed in Usp13−/− animals, with reduced cellular infiltration and neutrophils (Fig. 7a–c), TNF-α (Fig. 7d), and IL-1β (Fig. 7e). These data indicate that the augmentation of pro-inflammatory responses caused by USP13 depletion is in part due to the destabilization of Sigirr.
Fig. 5. USP13 is reduced in response to endotoxin exposure. (a, b) RAW 264.7 (a) or MLE12 cells (b) were treated with LPS (0.1 μg/ml for RAW 264.7 cells; 5 μg/ml for MLE12 cells) for 0, 6, and 24 h. Cell lysates were analyzed by immunoblotting with indicated antibodies. (c, d) C57BL/6J mice were challenged with an intratracheal injection of LPS (1 mg/kg body weight; 24 h) or P. aeruginosa (strain PA-103) for 4 h (d). Lung tissue lysates were analyzed by immunoblotting with antibodies against USP13, Sigirr, and β-actin. (e) Lung tissues were stained by hematoxylin and eosin (H&E) and antibodies against IL-1β or USP13. Representative images are shown (n = 4–6 per group). Bars, 100 μm. (f) RAW 264.7 cells were treated with LPS (0.1 μg/ml) for 0, 4, and 6 h. qRT-PCR was performed to analyze the relative mRNA expression of USP13. (g) RAW 264.7 cells were treated with an NF-κB inhibitor (10 μM), JNK inhibitor (10 μM), or p38/SAP2 inhibitor (10 μM) for 1 h prior to LPS (0.1 μg/ml, 30 min) treatment. Cell lysates were analyzed by immunoblotting with antibodies against USP13 and β-actin. For all panels, all data are mean ± SEM. n = 3 *p < 0.05; **p < 0.01 by two-way ANOVA with Post hoc Tukey’s test compared to 0 h (a, b). Representative blots from three independent experiments are shown. Bands densitometry were analyzed with ImageJ.

Fig. 6. Usp13−/− mice increase lung inflammation after LPS or PA103 challenge. WT and Usp13−/− mice were subjected to intratracheal injection of LPS (1 mg/kg body weight) for 6 h (a, c, e, g, h) or P. aeruginosa (strain PA103; 1 × 10⁴ colony-forming units/mouse) for 4 h (b, d, f, i, j). (a, b) Total cell counts in BAL were measured by a hemocytometer. (c, d) Neutrophil influx into alveolar spaces was examined by cytospin. (e, f) Lung sections were stained with hematoxylin and eosin. Representative images of the staining are shown (n = 8–10 per group). Bars, 100 μm. (g–j) TNF-α, IL-1β, and IL-6 levels in BAL were measured by ELISA and overall. For all panels, the p-value was calculated by two-way ANOVA with post hoc Tukey’s test *p < 0.05, **p < 0.01.
post-translational modification (such as phosphorylation) may be necessary for Sigirr internalization. Receptor internalization and degradation are uncoupled in other systems we have studied [28], and ubiquitination as well as phosphorylation are known molecular signals for receptor internalization [32]. Other Sigirr modifications that impact internalization, signaling, and stability will be areas of interest for future studies. Sigirr can also be constitutively degraded without ligand stimulation. In this study, we show that Sigirr is degraded in IL-37-dependent and -independent manners. Notably, we used a truncated form of IL-37 that Sigirr abundance is reduced after LPS treatment in human and mouse models. Though LPS has been shown to increase IL-37 degradation [19], our finding indicates that USP13 regulates Sigirr stability through the association and deubiquitination of Sigirr in both IL-37-dependent and -independent pathways. Notably, we used a truncated form of IL-37 in this study. We will confirm the effects of other IL-37 isoforms on Sigirr degradation in future works.

Protein ubiquitination can be reversed by DUBs, which cleave ubiquitin or ubiquitin-like proteins from substrate proteins to prevent their degradation [19]. Our finding indicates that USP13 regulates Sigirr stability through the association and deubiquitination of Sigirr in both IL-37-dependent and -independent pathways. In this study, we show that Sigirr abundance is reduced after LPS treatment in human and mouse cell and animal models. Though LPS has been shown to increase IL-37 mRNA stability in human cells [33], IL-37 is not present in mice. The decrease in Sigirr level we observe in mouse models may therefore be attributable to USP13 reduction rather than IL-37 signaling. USP13 targets multiple ubiquitinated proteins to modulate tumorigenesis through deubiquitination and regulation of PTEN [34], Myc [35], Beclin-1 [36], microphthalmia-associated transcription factor [37], and Siah2 [38]. USP13 also has been shown to regulate viral responses through the deubiquitination of STAT1 [39] and STING [40]. Our data indicate that USP13 exerts anti-inflammatory activity by stabilizing Sigirr. The inhibitory effect of USP13 on LPS signaling is dependent on its activity. Usp13 deficient mice display increased severity of lung injury. While destabilization of other USP13 targets involved in inflammation such as STAT1, STING, or PTEN may also contribute to the pro-inflammatory effects we observe with Usp13 knockout, none of these substrates is known to suppress inflammation as Sigirr does. Though most immune cells express Sigirr, it has been shown that M1 differentiated macrophages express more surface and total Sigirr compared with M2 macrophages and monocytes [41]. The molecular regulation of Sigirr degradation in different cell types needs more in-depth investigations. The expression of USP13 in M1 and M2 differentiated macrophages and monocytes needs to be studied in future works.

In summary, we propose a working model with USP13 as a positive regulator of Sigirr stability, and thus USP13 activity maintains anti-inflammatory IL-37-Sigirr activity with decreases in the abundance of both proteins during inflammation. USP13 deubiquitinase activity reverses Sigirr polyubiquitination and stabilizes the Sigirr protein to preserve anti-inflammatory signaling (Fig. 7g). Based on this information, stabilization of Sigirr could be achieved through enhancing USP13 activity, and this may be a practical strategy to block pulmonary inflammation.

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Declarations of interests

The authors declare no conflict of financial interests.
Author contributions
Y.Z. and J.Z. designed the study. LL, Y.Z. performed the experiments, analyzed the data, and wrote the manuscript. J.W., S.L., A.M.J., J.Z., R.K.M., and N.W. performed experiments, analyzed the data. N.W. and R.K.M. edited the manuscript.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ebiom.2019.06.011.

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