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NLRX1 promotes immediate IRF1-directed antiviral responses by limiting dsRNA-activated PKR translational inhibition

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Abstract

NLRX1 is unique among nucleotide-binding domain and leucine-rich repeat (NLR) proteins in its mitochondrial localization and capacity to negatively regulate MAVS- and STING-dependent antiviral innate immunity. However, some studies suggest a positive regulatory role for NLRX1 in inducing antiviral responses. We show that NLRX1 exerts opposing regulatory effects on virus activation of the transcription factors IRF1 and IRF3, potentially explaining these contradictory results. Whereas NLRX1 suppresses MAVS-mediated IRF3 activation, NLRX1 conversely facilitates virus-induced increases in IRF1 expression, thereby enhancing control of virus infection. NLRX1 has a minimal effect on NF-κB-mediated *IRF1* transcription, and regulates IRF1 abundance post-transcriptionally by preventing translational shutdown mediated by the

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AUTHOR CONTRIBUTIONS

H.F. and S.M.L. designed the experiments; H.F., E.M.L., D.Y., D.R.M., O.G-L, and I.M performed experiments; E.W., H.G., J.M., J.K.W., N.J.M., L.M.R., J.P-Y.T. and J.A.D. provided critical reagents and intellectual input; S.M.L. supervised the study; H.F. and S.M.L. analyzed the data and wrote the manuscript; all authors reviewed the manuscript.

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The nucleotide-binding domain and leucine-rich repeat (NLR) family of proteins is recognized for its roles in inflammasome-mediated responses to both pathogen- and damage-associated molecular patterns (PAMPs and DAMPs)¹. However, several NLR proteins, including NLRX1, NLRC3, NLRC5 and NLRP12, act as negative regulators of innate immunity with the capacity to check type I interferon (IFN) responses or NF-KBinduced pro-inflammatory cytokines^{2–5}. NLRX1 is distinguished from other NLR members by its localization to mitochondria, where it interacts with the adaptor protein MAVS through its unique N-terminal X and nucleotide binding-oligomerization (NBD) domains, sequestering MAVS and suppressing virus-induced IFN responses mediated by the pathogen sensor RIG-I⁶. NLRX1 also negatively regulates lipopolysaccharide (LPS)-induced NF-κB activation, interacting with TRAF6 in unstimulated cells and being recruited to the NEMO/IKK signaling complex following LPS stimulation via its leucine-rich repeat domain³. Deletion or functionally knocking down NLRX1 results in heightened IFN responses to poly(I:C) or RNA viruses, as well as increased inflammatory responses^{3,6,7}. Acting amazingly like a Swiss Army knife, NLRX1 also interacts with the adaptor protein STING through its NBD domain, thereby inhibiting IFN responses to DNA viruses mediated through the cGAS/cGAMP signaling pathway⁸. Abundant evidence thus supports a role for NLRX1 as a checkpoint inhibitor of early innate immune responses to both DNA and RNA viruses.

However, not all studies show NLRX1 exerts negative regulatory effects on innate immune responses to viruses. Sendai virus (SeV)-induced RIG-I- and MAVS-dependent phosphorylation of the transcription factor IRF3 and IFN- β and IP10 production were reported to be unchanged in *Nlrx1^{-/-}* murine embryonic fibroblasts (MEFs), as were IL-6 and IFN- β responses to poly(I:C)⁹. Although inflammatory responses to influenza virus infection are increased in the lungs of *Nlrx1^{-/-}* mice⁷, macrophage-mediated IFN responses were impaired secondary to enhanced apoptosis¹⁰. Overexpression of NLRX1 also enhanced, rather than inhibited, NF- κ B signaling by amplifying production of reactive oxygen species (ROS) in response to several stimuli¹¹. These findings are puzzling given that the preponderance of evidence favors a negative regulatory role for NLRX1. While they may reflect different experimental conditions, they have fueled controversy about the regulatory role of NLRX1.

Here we sought to understand how NLRX1 influences innate immune responses to virus infection of human hepatocytes. Hepatocytes are targeted for infection by several medically important viruses including hepatitis A virus (HAV) and hepatitis C virus (HCV), RNA viruses that cause inflammatory diseases of the liver¹². We show that NLRX1 competes with dsRNA-activated protein kinase R (PKR) for binding to viral RNA, and that it promotes early innate immune antiviral responses by protecting NF- κ B-driven increases in expression of the transcription factor IRF1 from PKR-mediated translational suppression. Hepatocytes deficient in NLRX1 expression have reduced accumulation of IRF1 but increased IRF3 dimer formation in response to virus infection, revealing opposing actions of NLRX1 on key

signaling pathways. Our data unveil a novel and sophisticated regulation of early innate immune responses by NLRX1 that overall promotes immediate antiviral defense in hepatocytes.

RESULTS

NLRX1 is a positive immune regulator in hepatocytes

To determine how NLRX1 influences antiviral responses in hepatocytes, we used CRISPR/ Cas9 gene editing to eliminate its expression in PH5CH8 cells, which are T-antigen transformed primary human hepatocytes with functional RIG-I and TLR3 signaling^{13,14}. NLRX1 expression was detected in control cells transduced with a nontargeting ('Scramble') sgRNA, but not in either of two independent cell lines (NLRX1-T2 and NLRX1-T3) transduced with different *NLRX1*-specific sgRNAs (Fig. 1a, Supplementary Table 1). We initiated HCV replication in these cells by co-electroporating synthetic viral RNA and duplex miR-122, an essential HCV host factor lacking in PH5CH8 cells¹⁵. We demonstrated that subsequent increases in viral RNA were due to bona fide viral replication because they were blocked by a direct-acting antiviral (DAA) inhibitor (Fig. 1b, left). We also infected the cells directly with HAV¹⁶ (Fig. 1b, right). Replication of both viruses was enhanced in NLRX1-deficient cells. We similarly assessed HAV replication in primary human fetal hepatoblasts (HFHs). Partial RNAi-mediated depletion of NLRX1 boosted viral replication in cells from two donors (Fig. 1c), confirming an antiviral role for NLRX1 in human liver cells.

Although innate immune responses restrict both HAV and HCV infection in PH5CH8 cells, it is difficult to document induction of antiviral cytokines in these cells following virus challenge. We thus employed a classic agonist of RIG-I signaling, SeV, to define the impact of NLRX1 deficiency on cytokine responses. We observed significant reductions in early (3 h) *IFNB1*, *IFNL1*, *IL1B* and *IL6* mRNA responses in SeV-infected NLRX1-deficient cells (Fig. 1d). This effect was no longer evident at 8 h by which time these responses had substantially subsided (Supplementary Fig. 1a). NLRX1 deficiency also impaired *IL1B* and *IL6*, but not *IFNB1* mRNA accumulation in response to the TLR3 agonist, poly(I:C), added to medium (Fig. 1e). NLRX1 deficiency consistently reduced the amount of IL-6 protein induced in response to SeV and poly(I:C) stimulation (Fig. 1f). Likewise, siRNA-mediated depletion of NLRX1 significantly impaired IL-6 protein increases induced by SeV infection of primary HFHs (Fig. 1g). Neither HAV nor HCV replicate in murine cells, but we observed an approximate 4-fold reduction in the early (3 h) intrahepatic *Ifnb* and *II6* mRNA responses to synthetic HAV RNA administered intravenously to *Nlrx1^{-/-}* versus wild-type mice (Fig. 1h).

Supplementing PH5CH8 cells with an antioxidant (N-Acetyl-1-cysteine, NAC) did not abrogate the positive effect of NLRX1 deficiency on replication of an HCV reporter virus, nor did treating cells with inhibitors of ER stress (tauroursodeoxycholic acid, TUDCA) or autophagy (3-methyladenine, 3-MA) (Supplementary Fig. 1b). Thus, increased HCV replication in NLRX1-deficient cells is unlikely to be due to reduced ROS production^{11,17}, or the influence of NLRX1 depletion on autophagy or the response to endoplasmic reticulum (ER) stress^{18,19}.

To ascertain whether the proviral effect of NLRX1 deficiency results from a reduction in the early innate immune response, we designed a Transwell assay for soluble antiviral factors (Supplementary Fig. 1c, left). HCV replication was enhanced in indicator cells separated by a permeable membrane from NLRX1-deficient versus control PH5CH8 cells stimulated by a replication-competent HCV RNA (Supplementary Fig. 1c, right). Inhibition of JAK/STAT signaling with ruxolitinib (JAK_{1,2} inhibitor) and tofacitinib (JAK₃ inhibitor) also substantially reduced the difference in HCV replication being suppressed by cytokine responses involving JAK signaling. Depleting NLRX1 had no effect on HAV or HCV replication in RIG-I-deficient Huh-7.5 human hepatoma cells²⁰ (Supplementary Fig. 1e), or in MAVS-deficient PH5CH8 cells (Supplementary Fig. 1f). Collectively, these results define NLRX1 as a positive regulator of soluble RIG-I- and MAVS-mediated antiviral responses in human hepatocytes.

NLRX1 differentially regulates IRF3 and IRF1 responses

We carried out a series of dual luciferase promoter assays to determine if NLRX1 directly regulates cytokine gene transcription. NLRX1 deficiency had little effect on basal NF- κ B (PRDII) promoter activity, but modestly reduced its activation by SeV (Fig. 2a, left). Consistent with this, NLRX1 overexpression enhanced PRDII activation triggered by SeV (Fig. 2a, right). In contrast, NLRX1 deficiency minimally affected activation of the *IFNB1* promoter, and had no effect on an IRF3-responsive promoter (4*PRD(I/III)) (Supplementary Fig. 2a,b). We also examined the impact of NLRX1 deficiency on stability of *IL6* mRNA, as IL-6 expression is regulated in part through 3' untranslated RNA (3'UTR) sequences programmed for rapid mRNA turnover²¹. NLRX1 deficiency had no effect on *IL6* mRNA decay in cells treated with actinomycin D (Supplementary Fig. 2c), nor did NLRX1 overexpression alter luciferase expression from mRNA transcripts containing the *IL6* 3'UTR (Supplementary Fig. 2d). In aggregate, these data suggest that NLRX1 has a positive but limited effect on SeV activation of NF- κ B-responsive promoters, and no influence on *IL6* message stability.

To more directly examine the influence of NLRX1 deficiency on NF- κ B activation, we examined SeV-induced RELA signaling. NLRX1 deficiency minimally affected NFKBIA (I κ B α) or RELA phosphorylation in SeV-infected PH5CH8 cells (Supplementary Fig. 2e). Electrophoretic mobility-shift assays (EMSA) with an NF- κ B probe revealed a significant, but quantitatively small decrease in the intensity of SeV-induced band shifts in NLRX1- deficient versus control PH5CH8 cells (Fig. 2b). We conclude from these data that NLRX1 deficiency minimally suppresses SeV-induced NF- κ B signaling in PH5CH8 cells, and that this subtle impairment of NF- κ B signaling is unlikely to explain the marked reductions we observed in cytokine expression.

To further test this hypothesis, we assessed how depleting RELA influences IL-6 production upon SeV infection and the modulation of IL-6 expression by NLRX1 in PH5CH8 cells. As anticipated, RELA deficiency caused a large decrease in SeV-induced IL-6 protein expression (Fig. 2c). Importantly, however, eliminating NLRX1 expression resulted in a further reduction in IL-6 production in RELA-deficient cells (Fig. 2c). Thus, the negative

impact of NLRX1 deficiency on the IL-6 response that we observed in RELA-replete cells is preserved in RELA-deficient cells. Taken collectively, these results indicate that NLRX1 regulates IL-6 production downstream and independently of NF- κ B.

Although our data show IL-6 is strongly regulated by NF- κ B in PH5CH8 cells (Fig. 2c), CRISPR/Cas9 depletion of either of two IRF family members, IRF1 and IRF3, suppressed early (3 h) *IL1B* and *IL6* mRNA and IL-6 protein responses to SeV (Supplementary Fig. 2f–h). Because these transcription factors regulate the induction of *IFNB1* and *IFNL1* transcripts in human hepatoma cells²², we examined the impact of NLRX1 deficiency on IRF3 and IRF1 responses in PH5CH8 cells. IRF3 is constitutively expressed, and activated by phosphorylation leading to its dimerization and cytoplasmic-nuclear translocation²³. NLRX1 deficiency significantly enhanced this response, leading to an increase in SeV-induced IRF3 dimer formation (Fig. 2d). In contrast, basal IRF1 protein expression is low, but is strongly induced by SeV infection. NLRX1 deficiency substantially reduced SeV-triggered increases in IRF1 abundance (Fig. 2e). Reconstituting NLRX1 expression in NLRX1-T3 cells with recombinant lentivirus reversed both of these changes (Fig. 2f). Collectively, these data reveal that NLRX1 differentially regulates SeV-induced IRF3 and IRF1 signaling in hepatocytes, suppressing the activation of IRF3 but enhancing increases in IRF1 expression.

NLRX1 promotion of IRF1 dominates in hepatocytes

Because NLRX1 deficiency suppresses innate immune control of viral replication in hepatocytes, we reasoned that the positive regulation of IRF1 by NLRX1, rather than the negative regulation of IRF3, is likely to dominate cytokine responses in hepatocytes. To test this hypothesis, we depleted NLRX1 expression in IRF1- and IRF3-deficient PH5CH8 cells, and examined responses to SeV challenge in the resulting NLRX1-IRF3 and NLRX1-IRF1 double-deficient cells. In IRF3-deficient cells, IRF1 and IL-6 protein responses to SeV infection were suppressed and HAV replication was enhanced by depleting NLRX1 (Fig. 3a). Thus, the impact of NLRX1 depletion was similar in IRF3-deficient and wild-type PH5CH8 cells. In contrast, depleting NLRX1 in IRF1-deficient cells enhanced IRF3 dimer formation (as in IRF1-replete cells), but enhanced SeV-induced IL-6 production and also suppressed HAV replication were reversed in IRF1-deficient cells, suggesting that the NLRX1-IRF1 signaling axis is dominant in hepatocytes. Consistent with this conclusion, reduced IL-6 production in SeV-infected NLRX1-depleted HFHs (Fig. 1g) was accompanied by reductions in IRF1 protein accumulation (Fig. 3c).

Since most reports suggest NLRX1 suppresses, rather than enhances early innate immune responses in bone marrow-derived macrophages (BMDMs) and primary MEFs^{6–8}, we sought to determine whether IRF1 signaling is negatively or positively regulated by NLRX1 in these cell types. Consistent with a suppressive effect on innate immunity, we found that loss of NLRX1 enhanced SeV-mediated *Ifnb, Tnf,* and *II6* mRNA responses in murine BMDMs and MEFs (Fig. 3d). Despite this, NLRX1 deficiency reduced SeV-triggered increases in IRF1 protein abundance in both cell types (Fig. 3e). Thus, the positive regulation of IRF1 signaling by NLRX1 is not specific to hepatocytes or human cells. The

fact that cytokine responses were enhanced, while IRF1 induction was decreased in NLRX1deleted BMDMs and MEFs, indicates that IRF1 does not play a dominant role in determining cytokine responses in these cells. Thus, the ultimate impact of NLRX1 on early antiviral responses is determined by whether IRF1 or IRF3 dominates in driving the response.

NLRX1 enhances IRF1 protein synthesis in infected cells

IRF1 is the founding member of the IRF family, and its expression is induced rapidly by viral infection^{24,25}. However, the mechanisms mediating this response are poorly understood. Given its role in initiating IRF3-directed responses to SeV in hepatocytes²⁰, we reasoned that RIG-I, and its adaptor, MAVS, may mediate the IRF1 response. Consistent with this notion, depleting MAVS significantly suppressed both IRF3 and IRF1 responses to SeV infection in PH5CH8 cells (Supplementary Fig. 3a). This loss of SeV-induced IRF3 dimerization and IRF1 protein expression was not reversed by additionally depleting NLRX1 in MAVS-deficient PH5CH8 cells (Supplementary Fig. 3a).

Consistent with previous studies showing that increased *IRF1* transcription drives the IRF1 protein response to viral infection²⁵, globally inhibiting transcription with actinomycin D completely abolished the IRF1 protein response to SeV (Supplementary Fig. 3b). To better understand how NLRX1 might regulate this IRF1 response, we assessed the roles of IRF3 and NF- κ B. Importantly, IRF3 deletion had no impact on either *IRF1* transcription or IRF1 protein expression (Supplementary Fig. 3c). In contrast, deleting NF- κ B RELA significantly suppressed both *IRF1* transcription and the increase in IRF1 protein expression in response to SeV in PH5CH8 cells (Supplementary Fig. 3d). siRNA-mediated silencing of *NFKB1*, which encodes NF- κ B p50, modestly reduced the IRF1 protein response to SeV infection, and in combination with RELA deficiency severely reduced SeV-induced *IRF1* transcript abundance (Supplementary Fig. 3e,f). Collectively, these data demonstrate that the immediate (3 h) increase in IRF1 protein expression in response to SeV infection is driven by MAVS and NF- κ B signaling in PH5CH8 cells.

Importantly, although NLRX1 deficiency modestly enhanced SeV-induced *IRF3* transcript abundance (Supplementary Fig. 4a), it had no effect on either *IRF1* transcript abundance (Fig. 4a) or *IRF1* mRNA stability (Fig. 4b). These data are in agreement with our earlier observations indicating only a modest impairment of NF- κ B signaling in NLRX1-deficient cells, and collectively indicate that NLRX1 regulates IRF1 expression post-transcriptionally. To gain insight into whether this negative regulation of the IRF1 response was due to enhanced IRF1 degradation or reduced IRF1 protein synthesis, we globally inhibited protein synthesis in SeV-infected cells with cycloheximide (CHX) and monitored the subsequent decay of IRF1 protein. Despite a greater initial abundance of IRF1 in control PH5CH8 cells, the half-life (stability) of IRF1 protein was indistinguishable in NLRX1-deficient cells (Fig. 4c, Supplementary Fig. 4b). Taken collectively, these data suggest that the reduced IRF1 protein abundance in NLRX1-deficient cells results from impaired translation of *IRF1* mRNA. Consistent with this, globally blocking protein synthesis with puromycin completely abolished the impact of NLRX1 deficiency on the IRF1 response to SeV infection (Supplementary Fig. 4c).

To determine how NLRX1 might influence protein synthesis, we measured the incorporation of [³⁵S]-Met/Cys into protein. Protein synthesis was qualitatively similar in NLRX1-T3 and control cells, both before and after SeV infection (Supplementary Fig. 4d). However, whereas SeV infection induced a significant, quantitative increase in protein synthesis in control cells, it had a much reduced, nonsignificant effect in NLRX1-T3 cells (Fig. 4d). The absence of any novel protein bands incorporating [³⁵S]-Met/Cys (Supplementary Fig. 4d) indicated that this increase was due to enhanced cellular protein synthesis, not viral protein synthesis. As an independent measure of protein synthesis, we pulse-labeled NLRX1-T3 cells with a low concentration of puromycin, and used confocal microscopy to monitor its incorporation into nascent protein at a single-cell level (Fig 4e, left). SeV infection induced an increase in nascent protein synthesis in a large proportion of control cells, but not in NLRX1-T3 cells (Fig. 4e). Similar results in NLRX1-T2 cells, at both a whole cell culture and single-cell level of analysis, excluded the possibility that this reflects an off-target effect of NLRX1-T3 sgRNA (Supplementary Fig. 4e). siRNA-mediated depletion of NLRX1 expression in primary HFHs also substantially suppressed SeV-induced increases in nascent protein synthesis (Supplementary Fig. 4f).

Although NLRX1 exerts distinctly different regulatory effects on SeV-induced cytokine expression in MEFs versus human hepatocytes, depleting NLRX1expression reduced SeV-induced increases in IRF1 abundance in both cell types. Consistent with this finding, puromycin incorporation into nascent protein was also significantly reduced in SeV-infected MEFs from $NIrx1^{-/-}$ versus wild-type mice (Supplementary Fig. 4g). Thus, the negative impact of NLRX1 deficiency on protein synthesis in virus-infected cells is not limited to human hepatocytes.

To directly characterize the impact of NLRX1 deficiency on mRNA translation by ribosomes, we used sucrose density gradient fractionation to profile polysome formation following SeV infection of NLRX1-T3 versus control cells (Fig. 4f). NLRX1 deletion resulted in a significant shift in the gradient distribution of *IRF1*, *IRF3* and *ACTB* mRNAs from SeV-infected cells, with a substantially lower proportion of each of the mRNAs associated with translationally-active polysomes (fractions 7–13) (Fig. 4g). These differences were statistically significant in replicate independent experiments. Moreover, the formation of translationally competent 80S ribosomes was reduced in SeV-infected NLRX1-T3 versus control cells, as evidenced by a sharp reduction in the 80S/40S subunit ratio (Fig. 4h, Supplementary Fig. 4h), suggesting a defect in initiation of translation, puromycin labeling, and polysome profiling, indicate that mRNA translation is globally suppressed by the absence of NLRX1 expression in SeV-infected cells.

NLRX1 limits PKR-mediated global translational shutdown

The double-stranded RNA-activated kinase PKR is triggered by virus infection to phosphorylate eIF2a, thereby globally shutting down host cell translation²⁶. To ascertain whether NLRX1 influences this PKR response, we characterized PKR activation in NLRX1-deficient cells. NLRX1 deficiency enhanced SeV-induced PKR autophosphorylation, as well as phosphorylation of the PKR substrate, eIF2a (Fig. 5a, Supplementary Fig. 5a). To

determine whether NLRX1 facilitates virus-induced increases in IRF1 protein synthesis by suppressing PKR responses, we generated NLRX1-PKR double-deficient PH5CH8 cells (Fig. 5b). The negative impact of NLRX1 deletion on SeV-triggered IRF1 responses was completely abolished by PKR deficiency (Fig. 5c). PKR deficiency also fully restored the SeV-triggered immediate IL-6 protein response (Fig. 5d), and abrogated the beneficial effect conferred on HAV replication by NLRX1 deficiency (Fig. 5e). We also assessed the impact of SeV infection on nascent protein synthesis in NLRX1-PKR double-deficient cells. In contrast to the lack of an increase in [³⁵S]-Met/Cys or puromycin incorporation in NLRX1-PKR double-deficient cells, protein synthesis was increased in response to SeV infection in both NLRX1-PKR double-deficient cells and PKR-deficient cells (Fig. 5f, Supplementary Fig. 5b).

To explore the mechanism by which NLRX1 suppresses PKR responses, we investigated whether NLRX1 physically interacts with PKR in hepatocytes. However, no coimmunoprecipitation of PKR protein was detectable with either endogenous or overexpressed NLRX1 in PH5CH8 cells (Supplementary Fig. 5c,d). Since NLRX1 binds both viral RNA and poly(I:C), a common surrogate for viral dsRNA^{27,28}, we reasoned that NLRX1 might suppress PKR activation by competing for viral RNA in infected cells. To test this hypothesis, we first confirmed that NLRX1 binds genomic HAV RNA, which is known to have extensive secondary structure²⁹, in bidirectional pull down assays (Fig. 5g). Next, we designed a cell-free competition assay in which purified recombinant NLRX1 protein and biotin-tagged HAV RNA or poly(I:C) were added to lysates of NLRX1-deficient cells as a source of PKR. The addition of increasing amounts of recombinant NLRX1 protein led to a progressive reduction in PKR binding to viral RNA and poly(I:C) (Fig. 5h), indicating that NLRX1 competes with PKR for binding to viral RNA. Taken together, these results demonstrate that NLRX1 suppresses PKR activation, in part by competitively binding viral RNA, thereby preventing PKR-mediated translational shutdown that would otherwise attenuate IRF1-mediated antiviral responses (Supplementary Fig. 5e).

DISCUSSION

Although NLRX1 has been reported to be a negative regulator of innate immunity^{3,6,8,30}, others have not confirmed this^{9,31} or even suggested that NLRX1 enhances antiviral responses¹⁰. Consistent with the latter, we found NLRX1 to be an antiviral factor and a positive regulator of early innate immune responses in human hepatocytes. HAV and HCV, both common causes of viral hepatitis in humans, are unrelated, positive-strand hepatotropic RNA viruses that share many similarities in their replication cycles and in their interactions with innate immunity¹². Both produce double-stranded RNA replication intermediates that induce RIG-I and TLR3 signaling, thereby activating NF-κB and IRF family members leading to antiviral defense^{12,32}. To counter these host defenses, both HAV and HCV express proteases that target for degradation MAVS and TRIF, key adaptor proteins in the RIG-I and TLR3 signaling pathways^{33–36}. Adding to these similarities, we show here that NLRX1 acts to restrict both HAV and HCV replication.

Surprisingly, we observed that NLRX1 had clear, opposing effects on SeV-induced activation of two members of the IRF family. NLXR1 was required for maximum increases in IRF1 abundance, yet suppressed IRF3 dimerization.As key factors in inducing cytokine

transcription, IRF family members are triggered by various stimuli in all cell types. Whereas IRF3 and IRF7 are activated by virus infection in most cell types^{37,38}, other family members, including IRF1 as well as IRF5, 8 and 9, are selectively activated by different stimuli in a cell-type dependent manner³⁹⁻⁴⁴. This complexity in IRF activation provides the diversity of responses required for effective host defense. However, it also allows for a single regulatory molecule, such as NLRX1, to have different functional consequences when expressed in different cell types or pathogen contexts or, potentially, at different points of time in the response to an invading virus. The opposing regulatory activities of NLRX1 on early IRF3 and IRF1 responses may account for much of the controversy surrounding this protein^{7,8,10}. We found NLRX1 positively regulates IRF1 signaling in both BMDMs and primary MEFs. Yet, NLRX1 is known to attenuate innate immune control of viral replication in these cell types⁶⁻⁸. We infer from this that IRF1 does not play a dominant role in determining the outcome of infection in these cells, at least not under the conditions tested, whereas our data show it does in hepatocytes. Thus, whether NLRX1 functions as a pro-viral or antiviral factor, or neither^{9,31}, likely reflects which IRFs dominate in inducing antiviral responses.

Although previous studies indicate NLRX1 negatively regulates LPS/TLR4-triggered NF- κ B signaling in macrophages and HEK293T cells^{3,7}, we found that NLRX1 has a slight, positive effect on RIG-I-mediated NF- κ B signaling in hepatocytes. However, this effect is not critical for the promotion of early cytokine responses by NLRX1, as our data show that NLRX1 regulates the IL6 response downstream of the influence of NF- κ B.

Negative regulation of MAVS-mediated IRF3 responses by NLRX1 has been described previously⁶, but how NLRX1 impacts IRF1 responses has not been studied. We found that depleting NLRX1 did not alter SeV-induced *IRF1* mRNA transcription in PH5CH8 cells, but it significantly reduced the synthesis of IRF1 protein. Reductions in protein synthesis were not specific to IRF1, but rather reflected a global shutdown of protein synthesis due to enhanced PKR activation, a phenomenon that would likely have its greatest impact on proteins that turn over rapidly, such as IRF1 that has a half-life of about 60 min.

Emerging evidence is increasingly linking translational control to regulation of immediate innate immune responses. Both IRF7 and IRF8 appear to be controlled via global, eIF4Edependent regulation of translation^{45–47}. PKR restricts protein synthesis globally by phosphorylating eIF2 α , thereby blocking the production of virus⁴⁸. However, PKR activation also reduces the synthesis of effector proteins encoded by interferon-stimulated genes⁴⁹, and, as we show here, the production of IRF1 driven by an early, NF- κ B-mediated response to SeV. PKR is thus a double-edged sword, potentially weakening the cell's antiviral defenses as well as presenting a hurdle to be overcome by the virus. The negative impact of PKR activation on host defense may be particularly important when cells are infected with viruses such as HCV that can continue to translate their RNAs despite PKR activation⁴⁹. Our data suggest NLRX1 moderates the potentially negative consequences of PKR activation by competing with it for binding to viral RNA. This is distinct from how other NLR family members, including NLRP3, NLRP1 and NLRC4, physically interact with PKR to maximize inflammasome activation⁵⁰. We speculate that the opposing actions of NLRX1 on early IRF1 and IRF3 antiviral responses may underlie much of the controversy surrounding the biological function of this NLR family member. Loss of NLRX1 expression may result in contrary effects on outcome of infections by different viruses in different tissues, depending on which IRF dominates in driving the antiviral response.

METHODS

Detailed information concerning experimental design and reagents used in this study can be found in the Life Sciences Reporting Summary that accompanies this publication.

Reagents and antibodies

Puromycin and blasticidin were from Invitrogen. 3-methyladenine (3-MA), tauroursodeoxycholic acid (TUDCA), N-acetyl-L-cysteine (NAC), ruxolitinib, tofacitinib, cycloheximide and actinomycin-D were from Sigma-Aldrich. PSI-7977 (Sofosbuvir) was from Chemscene. Poly(I:C) (high molecular weight, InvivoGen) was either added to cell culture supernatants at a final concentration of 50 μ g/ml, or transfected at a concentration of 2 μ g/ml.

Antibodies used in this study included anti-NLRX1 (1:5,000 dilution; this antibody was described previously⁶); anti-IRF1 (1:1,000 dilution; clone D5E4, #8478), anti-phospho-NFκB p65 (anti-p-RELA; 1:1,000 dilution; clone 93H1, #3033), anti-NF-κB p65 (anti-RELA; 1:1,000 dilution; clone D14E12, #8242), anti-p50 (1:1,000 dilution; clone 5D10D11, #13681), anti-phospho-IkBa (1:500 dilution; clone 14D4, #2859), anti-IkBa (1:1,000 dilution; clone L35A5, #4814), anti-PKR (1:1,1000 dilution; clone D7F7, #12297) and antiphospho-eIF2a (1:1,000 dilution; clone S51, #9721) from Cell Signaling; anti-IRF3 (1:500 dilution; clone FL-425, sc-9082), anti-eIF2a (1:500 dilution; clone FL-315, sc-11386) and anti-Lamin A/C (1:1,000 dilution; clone N-18, sc-6215) from Santa Cruz; anti-phospho-PKR (1:1,000 dilution; clone E120, ab32036) and mouse IgG1 (ab18448) from Abcam; anti-Cardif (anti-MAVS; 1:2,000 dilution; clone AT107, #ALX-210-929) from Enzo Life Sciences; anti-actin (1:40,000 dilution; clone AC-74, A2228) from Sigma, and anti-GAPDH (1:50,000 dilution; AM4300) from Ambion; anti-puromycin (1: 20,000 dilution for immunoblot and 1:10,000 dilution for immunofluorescence; clone 12D10, MABE343) from EMD Millipore. Goat anti-mouse IgG (H+L) cross-adsorbed secondary antibody (1:300 dilution; A-11001) was from ThermoFisher. IRDye 680 or 800 secondary antibodies included #926-32211, #926-32212, #926-32214, #926-68020 and #926-68073 (1:12,000) from LI-COR Biosciences.

Cells

Tissues for preparation of fetal human liver cells were from random donors, and were provided by the accredited nonprofit corporation Advanced Biosciences Resources, Inc. (ABR) and obtained from fetuses between 19–21 weeks gestation during elective termination of pregnancy. Tissues were collected with written informed consent from all donors and in accordance with the US Food and Drug Administration CFR Part 1271 Good Tissue Practices regulations. Tissue processing, and hepatoblast isolation and culture were as

previously described⁵¹. The use of commercially procured fetal liver cells was reviewed by the University of North Carolina at Chapel Hill Office of Human Research Ethics, and was determined to be exempt from review by the University of North Carolina at Chapel Hill Institutional Review Board.

293FT cells and human hepatocyte-derived cell lines including Huh-7.5⁵² and PH5CH8¹³ were mycoplasma-free and cultured as previously described⁵¹. Murine bone marrow-derived macrophages (BMDMs) and mouse embryo fibroblasts (MEFs) were generated and cultured as previously described⁸.

Mice and HAV RNA inoculation

Age- and sex-matched wild-type and $NIrxI^{-/-}$ mice were previously described⁷. Animals were stratified by sex prior to randomization. Experiments were performed in a non-blinded fashion in accordance with the NIH Guide for the Care and Use of Laboratory Animals and with the approval of the Institutional Animal Care and Use Committee of the University of North Carolina at Chapel Hill. Thirteen-week-old mice were inoculated intravenously with PBS or 15 µg *in vitro*-transcribed HAV RNA in a volume of PBS equivalent to ~5% body weight. Mice were sacrificed 3 h after injection, and liver tissue collected for analysis of cytokine mRNAs by qRT-PCR.

Virus

Sendai virus (SeV, Cantell strain) was obtained from Charles River Laboratory. Cultures were exposed to SeV at a concentration of 600 HA units/ml. High-titer HAV (HM175/18f strain⁵³) stock was prepared as described previously⁵⁴. HAV infection was carried out at m.o.i = 10 for PH5CH8 cells and m.o.i = 1 for Huh-7.5 cells. Infectious molecular clones for HCV (pJFH1-QL, containing the cell culture–adaptive mutation Q221L in the NS3 helicase)⁵⁵, a reporter virus expressing *Gaussia princeps* luciferase (pJFH1-QL/GLuc)⁵¹, and HAV (pT7-18f)⁵⁶ were described previously.

Plasmids and oligonucleotides

Firefly luciferase reporter vectors including pIFNB1-Luc, pIRF3-responsive (4*PRD(I/III))-Luc and pPRDII-Luc as well as the Renilla luciferase control reporter vector, pRL-TK, were described previously⁵⁷. The *IL6*3'UTR construct was kindly provided by B. Glaunsinger (University of California, Berkeley), and was sub-cloned to psiCHECK2. Plasmids for overexpressing NLRX1 and PKR were previously described⁶. An expression vector for reconstitution of NLRX1 was generated by removal of the TurboGFP-2A region from pLOC-RFP⁵⁸, producing pLOC GFP which retains the RFP open reading frame (ORF) (empty vector, EV). The RFP ORF was then replaced by sequence encoding the NLRX1 ORF to produce pLOC-NLRX1T3 using a PCR-based strategy as previously described⁵⁹. Oligonucleotides used in this study are listed in Supplementary Table 2. All constructs were verified by DNA sequencing.

In vitro transcription and transfection

In vitro transcription and transfection of viral RNA was performed as previously described⁵¹. For biotin labeling of HAV RNA, transcription mixes were supplemented with

1× concentrated Biotin RNA Labeling Mix (Roche). siLentfect Lipid Reagent (Bio-Rad) was used for transfection of siRNAs (siRNAs: *NLRX15*'-AUCCCGACGGAAGAUGUGC-3' and control #2, both from Dharmacon; *NFKB1* from Santa Cruz). Fugen HD Reagent (Promega) was used for DNA transfections.

Lentivirus production and transduction

For sgRNA CRISPR/Cas9 lentivirus production, individual sgRNA expressing vectors and the 3rd Generation Packaging Mix vectors (ABM, Inc., Canada) were co-transfected into 293FT cells. Transduction was facilitated by the addition of 8 μ g/ml polybrene, and the resulting cells subjected to selection with puromycin (6 μ g/ml) for single-deleted cells, and puromycin (6 μ g/ml) and blasticidin (10 μ g/ml) for double-deleted cells.

RNA extraction and quantitative RT-PCR

Total RNA extraction was carried out with the RNeasy Kit (Qiagen), followed by a two-step quantitative RT-PCR analysis of indicated genes with the SuperScript® III First-Strand Synthesis System (Invitrogen) and iTaq SYBR Green Supermix (Bio-Rad). The abundance of indicated genes was normalized to that of *ACTB* (human) or *Actb* (mouse). HAV and HCV RNA abundance was quantified as previously described^{51,60}.

Nuclear protein extraction

Extraction of nuclear proteins was carried out as previously described⁶¹, with minor modification. Briefly, the washed nuclei pellet was re-suspended in 50 μ l of nuclear protein extraction buffer (20 mM Tris-HCl, pH 7.5, 400 mM NaCl, 1.5 mM MgCl₂, 0.2 mM EDTA) supplemented with complete protease inhibitor and PhoSTOP (Roche).

Protein-protein/RNA interactions

Cell lysates were prepared from 10-cm dish cultures $(1.5~2 \times 10^6$ cells) with lysis buffer (20 mM Tris-HCl, pH 7.5, 50 mM KCl, 250 mM NaCl, 10% Glycerol, 5 mM EDTA, 1× complete protease inhibitor, 1× PhoSTOP) supplemented with detergents as follows. For protein-protein interaction assays, 0.2% Triton X-100 and 0.3% NP-40 were added. The resulting lysates were used for immunoprecipitation as previously described⁶². For protein-RNA interaction, 0.5% Triton X-100 with 0.1% NP-40 was added. Recombinant NLRX1 was generated as previously described⁶³. Lysates were mixed with an equal volume of 2× incubation buffer (300 mM KCl, 40 mM HEPES, pH7.5, 10% glycerol, 200 Units/ml RNaseOUT (Thermo Fisher), 10 mM magnesium acetate, 2mM DTT). For precipitation, 500 ng biotin-labeled HAV RNA, 100 ng biotin-labeled poly(I:C) (InvivoGen) or 1 µl polyclonal anti-NLRX1 was used in each reaction. All reactions were gently rotated at 4 °C for 2 h, followed by the addition of magnetic streptavidin T1 beads (Invitrogen) for another 30 min rotation or protein G-Sepharose (GE Healthcare) for another 1 h rotation. After 5 intense washes with 1× incubation buffer, the final product was analyzed by qRT-PCR and/or immunoblotting.

EMSA, native PAGE, immunoblots and visualization

For EMSA, IRDye 700-labelled oligonucleotides containing a consensus NF-κB binding site (5'- AGTTGAGGGGACTTTCCCAGGC-3'; LI-COR Biosciences) were incubated with nuclear extracts according to the manufacturer's protocol (Odyssey Infrared EMSA kit). Electrophoresis of the resultant products was carried out with 5% TBE gel (Bio-Rad). Native PAGE for analysis of IRF3 dimerization was performed as previously described⁶⁴. Immunoblotting was carried out using standard methods. An Odyssey Imaging System (Li-COR Biosciences) was used for visualization and signal (infrared fluorescence) intensity analysis.

Polysome profiling, [³⁵S]-metabolic labeling, and puromycin incorporation assay

Control and NLRX1-depleted PH5CH8 cells (2×10^7) were infected with SeV (3×10^4 HA units) for 2 h. Cells were then harvested and polysome gradients prepared and analyzed as previously described⁶⁵. For metabolic labeling, 3×10^5 cells were infected with SeV (1.5 × 10³ HA units) for 2 h 15 min. After 15 min starvation in methionine/cysteine-free medium, newly synthesized proteins were pulse-labeled with [³⁵S]-labeled methionine and cysteine, 125 µCi [³⁵S]/ml, for 30 min as described⁶⁶. Protein synthesis was assessed in cells infected by SeV under similar conditions by monitoring puromycin incorporation into nascent protein using a method described previously⁶⁷. In brief, cells grown on glass slides were pulsed with puromycin (10 µg/ml) for 10 min, then fixed with 4% paraformaldehyde followed by staining with anti-puromycin and DAPI. Slides were imaged by laser-scanning confocal microscopy at a fixed gain in an Olympus FV1000 instrument at 40× magnification, and recorded images analyzed with ImageJ (Fiji). Individual cells were identified by DAPI staining of nuclei, and scored for intensity of the puromycin signal greater than an arbitrary threshold set with mock-infected control cells in each experiment. Alternatively, cells were lysed and puromycin-labeled proteins identified by immunoblotting, and quantified by assessing infra-red fluorescence intensity with the Odyssey Imaging System.

Luciferase reporter assays

Luciferase assays, including secreted *Gaussia* luciferase (GLuc) analysis of HCV replication and dual luciferase assays to analyze promoter activation, were carried out as described previously^{57,68}.

ELISA

IL-6 concentrations were determined with the human IL-6 ELISA Kit (eBioscience) following the manufacturer's protocol.

Statistical analysis

Unless noted otherwise, all between-group comparisons were carried out by one-way or two-way ANOVA or two-sided *t* test. Calculations were made using Prism 6.0 software (GraphPad Software, Inc.).

Data availability

Data that support the findings of this study are either shown directly in figures accompanying the main text or in supplementary figures, or are available from the corresponding author upon request.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Figure 1.

NLRX1 is a positive regulator of innate immunity and an antiviral factor in hepatocytes. (**a**) Immunoblot confirmation of NLRX1 depletion in two independent PH5CH8-derived cell lines. (**b**) qRT-PCR of HCV and HAV genomic RNAs in NLRX1-deficient cells transfected with HCV JFH1-QL RNA (left, 24 h) or infected with HAV (right, 72 h). Data are from n=2 (left) or 3 (right) technical replicates and are representative of 3 independent experiments. (**c**) qRT-PCR of HAV genomic RNA in primary human fetal hepatoblasts (HFHs) with partial NLXR1 depletion at 72 h. n=3 and 4 technical replicates for donors 1 and 2, respectively. (**d**,**e**) qRT-PCR for *IFNB1*, *IFNL1*, *IL1B* and *IL6* mRNA in NLRX1-deficient PH5CH8 cells infected with SeV (d) or stimulated with poly(I:C) added to medium for 3 h (e). Data are from 2 independent experiments with 4 technical replicates (d, *IFNB1* and

IFNL1), or are representative of 3 experiments, each with n=3 technical replicates (d, *IL1B* and *IL6*; e). (**f**,**g**) ELISA for IL-6 protein production in NLRX1-deficient PH5CH8 cells (f) and NLRX1-depleted HFHs (g). Data are representative of 3 experiments or donors, each with n=3 technical replicates. (**h**) qRT-PCR for *Ifnb* and *II6* mRNA in livers of *Nlrx1^{-/-}* mice 3 h following hydrodynamic injection of HAV RNA (n=3 per group). For all qRT-PCR analyses, host gene expression was normalized to *ACTB* (human) or *Actb* (mouse). All data are shown as mean ± S.E.M. Unless otherwise indicated, comparisons were between control and NLRX1-depleted cells by two-way ANOVA (ns, not significant; *p < 0.05; ** p< 0.01; ****p < 0.001). Comparisons in (b right,c,g,h) were performed by *t* test (*p < 0.05).

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Figure 2.

Impact of NLRX1 deficiency on NF-κB signaling and IRF1 and IRF3 activation in SeVinfected PH5CH8 cells. (a) PRDII-Luc promoter activation in cells with NLRX1 depletion (left) or overexpression (right). Data are from n=3 technical replicates and are representative of 3 independent experiments. (b) NF-κB electrophoretic mobility shift assay (EMSA) with nuclear extracts from mock- and SeV-infected NLRX1-deficient PH5CH8 cells (left). Mean infrared fluoresence intensity measurements from 4 independent EMSA experiments (right). (c) Effects of NLRX1 depletion in the absence of RELA. Immunoblots for NLRX1 and RELA in NLRX1-RELA double-deficient (RELA KO+*NLRX1*-T3) PH5CH8 cells (left). ELISA quantitation of IL-6 protein abundance in SeV-infected (3 h) control, RELA-deficient and NLRX1-RELA double-deficient cells (right). Results are representative of 2 independent experiments, each with 3 technical repeats. (d,e) Immunoblots showing SeVtriggered IRF3 dimerization (d) and IRF1 expression in NLRX1-deficient cells (e). n=3–4 independent experiments. (f) SeV-induced IRF3 dimerization and increased IRF1 protein

expression in NLRX1-reconstituted NLRX1-T3 cells. Data are representative of 2 independent experiments. All data are shown as mean \pm S.E.M. Comparisons of control versus NLRX1-depleted cells in panels (a,e) and (b-d, f) were made by two-way ANOVA and *t* test, respectively. ns = nonsignificant, *p< 0.05; **p < 0.01; ***p< 0.001; ****p< 0.0001.

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Figure 3.

NLRX1/IRF1 signaling dominates the antiviral cytokine response in hepatocytes. (**a**,**b**) ELISA measurements of SeV-induced IL-6 production and qRT-PCR quantitation of HAV replication in IRF3-deficient and IRF3-NLRX1 double-deficient PH5CH8 cells (a), or IRF1-deficient and IRF1-NLRX1 double-deficient cells (b). Panels on the left show immunoblots for NLRX1 and IRF1 (a) or IRF3 dimerization at 3 h (b). See Supplementary Fig. 2f for immunoblots of IRF1-deficient and IRF3-deficient cells. Results are representative of 2–3 independent experiments, each with 3 technical repeats. (**c**) Immunoblots of NLRX1 and IRF1 in partial NLRX1-depleted HFHs infected with SeV for 3 h. Bottom panel shows

quantification of immunoblots (infrared fluorescence intensities) from 2 independent experiments using cells from different donors. (**d**) qRT-PCR for *Ifnb*, *Il6* and *Tnf* mRNA and (**e**) immunoblots of IRF1 in SeV-infected wild-type versus $NIrx1^{-/-}$ BMDMs (top panels) and MEFs (bottom panels). Data are from n=3 technical replicates and are representative of 4 (BMDMs) and 2 (MEFs) independent experiments. All immunoblots are representative of 2–3 independent experiments or donors. Data are shown as mean ± S.E.M. ns = nonsignificant, *p<0.05; **p<0.01; ***p<0.001; ****p<0.0001 by *t* test.

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Figure 4.

NLRX1 facilitates immediate IRF1 antiviral responses by promoting global protein synthesis in SeV-infected cells. (a) qRT-PCR of SeV-induced *IRF1* mRNA responses in control and NLRX1-deficient PH5CH8 cells. (n=3). (b) *IRF1* mRNA stability in actinomycin D (Act D)-treated SeV-infected control vs. NLRX1-T3 PH5CH8 cells. (n=3). (c) IRF1 protein stability in cycloheximide (CHX)-treated cells following SeV infection assessed by quantitation of immunoblots. (n=6). See Supplementary Fig. 4b for representative immunoblots. (d) Nascent protein synthesis monitored by [³⁵S]-Met/Cys incorporation in mock- versus SeV-infected control and NLRX1-T3 cells. [³⁵S] incorporated

into TCA-precipitated protein is shown relative to that in uninfected control cells (100%). Data are from n=6 technical replicates from three experiments. See Supplementary Fig. 4d for representative immunoblots. (e) Global protein synthesis imaged by confocal microscopy in mock- versus SeV-infected control and NLRX1-T3 cells. Puromycin was detected by staining with specific antibody, and nuclei counterstained with DAPI. Scale bar, 40 µm. The percentage of cells with puromycin labeling exceeding an arbitrary threshold in 3 experiments is shown on the right. An average of 176 cells were evaluated for each condition. (f) Representative polysome profiles of IRF1, IRF3 and ACTB mRNAs in mockversus SeV-infected control and NLRX1-T3 cells. Absorbance at 254 nm (A254) is aligned with qRT-PCR quantitation of mRNA, plotted as percent of total mRNA in each fraction. (g) Proportion of IRF1, IRF3 and ACTB mRNAs associated with translationally-active polysomes (fractions 7-14) in SeV- (n=3) and mock-infected (n=2) control versus NLRX1-T3 cells. (h) 80S/40S ratios determined from the area under the curve (AUC) of 40S and 80S peaks of A_{254} traces in 2–3 experiments. Data are shown as mean \pm S.E.M. ns = nonsignificant, *p< 0.05; **p < 0.01; ***p< 0.001 by two-way ANOVA (a,d,e,g) or *t* test (h).

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Figure 5.

NLRX1 reduces PKR activation and subsequent inhibition of protein synthesis by competing for viral RNA binding. (a) Immunoblots of PKR-eIF2a signaling in NLRX1-deficient PH5CH8 cells 4.5 h after SeV infection. Quantification (infrared fluorescence intensity) of p-PKR or p-eIF2a was normalized to actin (n=4 experiments). (b) Immunoblots of PKR and NLRX1 in PKR-deficient and NLRX1-PKR double-deficient PH5CH8 cells. (c) Immunoblots of SeV-induced IRF1 expression and IRF3 dimerization in NLRX1-deficient and NLRX1-PKR double-deficient PH5CH8 cells. The image was generated from 2 separate gels in one experiment; data are representative of 2 experiments. (d,e) ELISA for SeVtriggered IL-6 protein production (d) and qRT-PCR for HAV RNA (e) in control, NLRX1deficient and NLRX1-PKR double-deficient PH5CH8 cells. Data are representative of 2-3 experiments, each with 3 technical replicates. (f) $[^{35}S]$ -Met/Cys incorporation as measurement of global protein synthesis in SeV- versus mock-infected NLRX1-PKR double-deficient PH5CH8 cells. Data are from 4 technical replicates of 2 experiments. (g) Bidirectional assessment of NLRX1-HAV RNA association. (left) qRT-PCR of HAV RNA co-immunoprecipitating with NLRX1. HAV RNA was transfected as in Fig. 1b, followed by immunoprecipitation with anti-NLRX1 at 3 h. Immunoblots of input and

immunoprecipitated (IP) NLRX1 are shown at the bottom. Data are representative of 2 experiments (n=2–3). (right) Immunoblots of NLRX1 co-precipitating with biotin-tagged HAV RNA in cell lysates. Immunoblots are representative of 3 experiments (\diamondsuit indicates endogenous NLRX1 associated with HAV RNA). "High" and "Low" indicate detection intensity. (h) Impact of NLRX1 reconstitution on PKR association with biotin-tagged HAV RNA or poly(I:C) in NLRX1-T3 cell lysates. Immunoblots are representative of 3 (HAV) or 5 [poly(I:C)] experiments. Quantification of PKR binding is summarized on the right. Data are shown as mean ± S.E.M. ns = nonsignificant, *p< 0.05; **p < 0.01; ***p< 0.001 by two-way ANOVA (a,d,f,g) and one-way ANOVA (e,h).