The Type I Interferon-IRF7 Axis Mediates Transcriptional Expression of Usp25 Gene*

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Viral infection or lipopolysaccharide (LPS) treatment induces expression of a large array of genes, the products of which play a critical role in host antipathogen immunity and inflammation. We have previously reported that the expression of ubiquitin-specific protease 25 (USP25) is significantly up-regulated after viral infection or LPS treatment, and this is essential for innate immune signaling. However, the mechanism behind this phenomenon is unclear. In this study, we found that viral infection-induced up-regulation of USP25 is diminished in cells lacking interferon regulatory factor 7 (IRF7) or interferon receptor 1 (IFNAR1) but not p65. Sendai virus- or type I interferon-induced up-regulation of USP25 requires de novo protein synthesis of IRF7. Furthermore, IRF7 directly binds to the two conserved IRF binding sites on the USP25 promoter to drive transcription of USP25, and mutation of these two sites abolished Sendai virus-induced IRF7-mediated activation of the USP25 promoter. Our study has uncovered a previously unknown mechanism by which viral infection or LPS induces up-regulation of USP25.

Host pattern recognition receptors recognize pathogen-associated molecular patterns and initiate a series of signaling cascades that lead to activation of transcription factors including NF-κB and interferon regulatory factor 3 (IRF3) (1–3). It has been well documented that activation of NF-κB (p65/p50 heterodimer) is dependent on inhibitors of κB kinase (IKK) complex (IKKα/β/γ)-mediated phosphorylation and degradation of IκBα, whereas activation of IRF3 requires phosphorylation by TBK1 or IKKe (4–9). The activated NF-κB and IRF3 enter into nucleus, bind to the conserved κB or IRF binding sites of promoters, and recruit co-activators to activate the transcription of target genes.

Viral nucleic acid and lipopolysaccharide (LPS) of Gram-negative bacteria are two common pathogen-associated molecular patterns that trigger signaling cascades to activate NF-κB and IRF3 and induce the production of type I interferons (IFNs) (3, 10). Type I IFNs further induce the expression of hundreds of downstream genes in an autocrine or paracrine manner, and the products of these genes including interferon-induced GTP-binding protein (Mex), 2′-5′-oligoadenylate synthase (OAS), double-stranded RNA-activated protein kinase (PKR), ISG56, and ISG15 orchestrate inhibition of pathogen replication and spread and promote apoptosis and clearance of the infected cells (11). In addition to the direct effect on innate immune cells for antipathogen responses, type I IFNs also regulate adaptive immunity including T cell activation and differentiation and antitumor immunity (12, 13).

The type I IFN family is composed of 13 functional IFNA genes in humans (14 in mice), a single IFNB gene, and others. The IFNα family shares 80% sequence homology among them, whereas the homology between various IFNα and IFNβ is 30% (14, 15). However, all the type I IFNs bind to the same receptors, IFNAR1 and IFNAR2, with affinities varying from picomolar to micromolar orders to recruit tyrosine kinase 2 (TYK2) and Janus kinase 1 (JAK1) for signal transduction, respectively. TYK2 and JAK1 are cross-phosphorylated and activated to phosphorylate several conserved tyrosine residues on IFNAR1 and IFNAR2, which provides docking sites for the downstream effector proteins including STAT1 (16, 17). It has been shown that STAT2 interacts with IFNAR2 constitutively, whereas STAT1 is recruited to IFNAR2-IFNAR1 receptor complex in both STAT2-dependent and -independent manners (18–20). TYK2 and JAK1 further phosphorylate Tyr-701 of STAT1 and Tyr-690 of STAT2, which form the ISGF3 transcription factor complex together with IRF9 to bind to the IFN-stimulated response elements on the promoters of and activate the transcription of ISGs (16). Type I IFN treatment also results in Tyr(P)-STAT1 homodimers that are responsible for the regulation of IFNγ-activated sequence elements (21, 22). In addition to phosphorylation of STAT1 at Tyr-701, the phosphorylation of STAT1 at Ser-708 by IKKe accounts for transcriptionsal activation of about 30% of the ISGF3 target genes (23). Thus, it is conceivable that type I IFN-triggered transcription of ISGs is
regulated at multiple steps ranging from the ligand subtypes to the modifications of transcription factors.

IFR7 is strongly induced by type I IFN-mediated signaling in a manner that is dependent on the TYK2-mediated phosphorylation of Tyr-701 of STAT1 but independent of IKKe-mediated phosphorylation of Ser-708 of STAT1 (23, 24). Although IFR3 and IFR7 share a similar structure to bind the conserved IFR binding sites and are activated by TBK1- or IKKe-mediated phosphorylation, studies with Ifr7−/− and Irf7−/− mice or cells suggest that IFR3 is required for early induction of IFNβ and IFNα, whereas IFR7 is a master transcription factor essential for later induction of IFNα subsets (25–30). Whether and how IFR3 and IFR7 differentially regulate transcription of other genes are of great interest.

We have previously observed that LPS or viral infection substantially up-regulates the expression of Usp25 gene (31, 32). In this study, we found that virus- or LPS-induced expression of Usp25 was significantly abolished in cells lacking IFR7 or IFNAR1. Importantly, type I IFN-triggered signaling indirectly induces up-regulation of Usp25 by inducing expression of IFR7. Furthermore, we have identified two conserved IFR7 binding sites on the promoter of Usp25 gene, and mutation of these two sites impaired SeV-induced or IFR7-mediated activation of the Usp25 promoter. Our study has uncovered the type I IFN-IRF7 axis-mediated expression of Usp25 gene.

**Experimental Procedures**

*Mice—Ifnar11−/− mice were purchased from The Jackson Laboratory and maintained and crossed to obtain Ifnar11+/− and Ifnar1−/− littermates in the specific pathogen-free facility of Wuhan University. Age- and sex-matched Ifnar11+/− and Ifnar1−/− littermates were used for all experiments. All animal experiments were in accordance with protocols approved by the Institutional Animal Care and Use Committee of Wuhan University.

Cells—Ifnar11+/− and Ifnar1−/− MEFs were isolated from E14.5 embryos. Bone marrow from Ifnar11+/− and Ifnar1−/− mice was isolated and differentiated into bone marrow–derived dendritic cells (BMDCs) with GM-CSF (20 ng/ml). MLFs were isolated as described previously (31). Ifr7−/− Irf7−/− MEFs were kindly provided by Dr. Pinghui Feng (University of Southern California). P65+/− and P65−/− MEFs were kind gifts from Dr. Tom Maniatis (Columbia University). Tbk1−/− MEFs were described previously (33, 34) and provided by Dr. Wen-Chen Yeh (University of Toronto). The cells were cultured in DMEM containing 10% FBS, 1% streptomycin-penicillin, and 10 μM mercaptoethanol.

Constructs, Antibodies, and Reagents—Mouse IFR3 or IFR7 was cloned into phage-6tag vector via standard molecular methods. USP25 promoter (~5000 to −1) was cloned into the pGL3-Basic vector (Promega). Site-directed mutagenesis was performed with a kit (Life Technologies) and sequenced for confirmation. FLAG-tagged RelB, p52, and p50 were kindly provided by Dr. Jin Jin (Zhejiang University). IFNα, IFNβ, anti-IFNβ, anti-IFNα (PBL Assay Science), actinomycin D (Sigma), IMD0354, ZM449829, amlexanox, and p38 mitogen-activated protein kinase inhibitor (Abcam) were purchased from the indicated manufacturers. Mouse anti-FLAG (KM8002), mouse anti-β-actin (KM9001), and HRP-conjugated goat anti-mouse or -rabbit IgG (Thermo Scientific, PA1-86717 and SA1-9510) were from the indicated manufacturers. Rabbit anti-USP25 was described previously (35) and kindly provided by Gemma Marfany (University de Barcelona, Barcelona, Spain).

Real Time Quantitative PCR—Cells treated with various stimuli were harvested in TRIzol (Invitrogen), and first strand cDNA was synthesized with a reverse transcription kit (Biotool). Gene expression was examined with a Bio-Rad CFX Connect system with a SYBR Green One Step Real-Time PCR kit (Biotool). Data were normalized to the expression of β-actin. Real time quantitative PCR primers were described previously (31) and are as follows: Ifr7: forward, CGG AAA GAA GTG TTG CGG TT; reverse, TTT TCC TGG GAG TGA GGC AG; Ifnan: forward, AGA GGG CGT TTT ATC TTG CG; reverse, TGG AGC CCA GCA TTT TCT CT; and Ifnan: forward, TCA AAG GAC TCA TCT GCT GC; reverse, GGT TCC TGC ACC CCC ACC TG.

Viral Infection—Cells were seeded into 24-well plates (2 × 10^3 cells/well) or 6-well plates (10^6–10^7 cells/well). Twenty-four hours later, cells were treated with LPS or infected with SeV or HSV-1 for the indicated time points. The cells were collected for quantitative PCR (qPCR) or immunoblotting assays.

Virus-mediated Gene Transfer—For lentivirus-mediated gene transfer, phage-6tag-IFR3, phage-6tag-IRF7, phage-6tag-rTBK1, phage-6tag-rIKKe, or the empty vector was cotransfected with the packaging vectors pSPAX2 and pMD2G into HEK293T cells. Eight hours after transfection, the medium was changed with fresh full medium (10% FBS, 1% streptomycin-penicillin, and 10 μM β-mercaptoethanol). Forty hours later, the supernatants were harvested to infect Ifr7−/− Irf7−/− MEFs, or wild-type MEFs followed by puromycin (1 μg/ml) selection for 2 weeks.

siRNA—siRNA targeting mouse IFR3 or IRF7 was synthesized and transfected into cells by Lipofectamine 2000 (Life Technologies) according to the manufacturer’s protocol. The sequences of siRNA are as follows: IRF3-siRNA1, 5′-GGA AAG AAG UGU UGC GGU UTT-3′; IRF3-siRNA2, 5′-GGC UAU UGU UUC UGA UCC UTT-3′; IRF3-siRNA3, 5′-GGU UGC UCC AUC AUC UCU UTT-3′; IRF7-siRNA1, 5′-CUU CGG CCA AGA CAA UUC ATT-3′; IRF7-siRNA2, 5′-CU GGA UGC GAC CAU CAU GUTT-3′; IRF7-siRNA3, 5′-CUA CGU CCA UCC GAG AAC UTT-3′; siTBK1, 5′-CCC ACA ACA CGA UUG CCA UTT-3′; siIKKe, 5′-CCC ACA ACA CGA UUG CCA UTT-3′, and control siRNA, 5′-GAU GAC GGG AAC UAG AAC ATT-3′.

Reporter Gene Assays—HEK293 cells (4 × 10^4 cells/well) cultured in 24-well plates were transfected with the reporter plasmid (100 ng) and an internal control vector, phRL-TK- Renilla luciferase (Promega) (2.5 ng). The pGL3-Basic vector served as a negative control, and empty vector was used to equalize the total amount of DNA. Twenty-four hours after transfection, cells were lysed in passive lysis buffer, and the firefly and Renilla luciferase activities were determined using a Dual-Luciferase reporter assay kit (Promega). The firefly luciferase activity was normalized by Renilla luciferase activity and expressed as the
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-fold stimulation relative to the activity in vector-transfected cells.

**Chromatin Immunoprecipitation Assays**—Briefly, $5 \times 10^6$ cells were fixed with 1% formaldehyde and quenched by glycine. The cells were washed three times with PBS and then harvested in chromatin immunoprecipitation (ChIP) lysis buffer (50 mM Tris-HCl, pH 8.0, 1% SDS, 5 mM EDTA) followed by sonication until the sizes of DNA were 400–600 bp. The
lysate was centrifuged at 4 °C for 15 min, and ChIP dilution buffer (20 mM Tris·HCl, pH 8.0, 150 mM NaCl, 2 mM EDTA, 1% Triton X-100) was added to the supernatant (4:1 volume). The resulted lysate was then incubated with anti-FLAG at 4 °C overnight. The protein G beads were added into the lysate on the next morning and incubated at 4 °C for 3 h. DNA was eluted using ChIP elution buffer (0.1 m NaHCO₃, 1% SDS, 30 μg/ml proteinase K) through incubation at 65 °C overnight, and DNA was purified with a DNA purification kit (TIANGEN). The purified DNA was assayed by quantitative PCR with an CFX Connect system with a SYBR Green One Step Real-Time PCR kit.

**Statistical Analysis**—Differences between experimental and control groups were determined by Prism software with two-way analysis of variance and Bonferroni test. p values less than 0.05 were considered statistically significant.

**Results**

**IRF7 Plays an Essential Role of LPS- or Virus-induced Expression of Usp25**—In our previous studies, we have observed that the expression of *Usp25* is up-regulated by LPS treatment or viral infection in various types of cells (31, 32). Interestingly, treatment with actinomycin D, a compound that inhibits transcription, almost abolished the up-regulation of *Usp25* and the increase of USP25 protein by LPS or SeV or HSV-1 infection in BMDCs (Fig. 1A). Sequence analysis of the promoter of mouse *Usp25* gene identified two IRF binding sites (−1399 to −1375 and −4648 to −4624), two ISGF3 binding sites (−3154 to −3130 and −3345 to −3321), and at least five NF-κB binding sites (Fig. 1B). However, p65 deficiency did not affect SeV-induced expression of *Usp25* but did inhibit SeV-induced expression of *Ifnb* and *Il6* (Fig. 1C). We reconstituted either IRF3 or IRF7 into IRF3−/−IRF7−/− MEFs and examined SeV-induced expression of *Usp25*. We found that reconstitution of IRF7 into IRF3−/−IRF7−/− MEFs promoted SeV-induced up-regulation of *Usp25* more robustly than did reconstitution of IRF3 (Fig. 1D). In addition, SeV-induced up-regulation of *Usp25* was substantially inhibited by knockdown of IRF7 and to a lesser extent by knockdown of IRF3 in MEFS (Fig. 1E). These data suggest that IRF7 and IRF3 (to a lesser extent) but not p65 are essential transcription factors for virus-induced up-regulation of *Usp25*.

**LPS- and Virus-induced Up-regulation of Usp25 Depends on Type I IFN-triggered Signaling**—We next examined the effects of various kinase inhibitors on virus- or LPS-induced up-regulation of *Usp25* in BMDCs or MLFs. Consistent with the notion that IRF7 and IRF3 are essential for transcriptional up-regulation of *Usp25*, inhibition of the upstream kinases TBK1 and IKKe by amlexanox but not IMD0354 (inhibitor for IKKβ) or the p38 kinase inhibitor impaired LPS- or virus-induced expression of *Usp25* (Fig. 2A). Interestingly, we also found that ZM449828 (a JAK1 inhibitor) strongly inhibited up-regulation of *Usp25* induced by LPS or viral infection, indicating that JAK1-mediated signaling is critical for the induction of *Usp25*.

Because JAK1 is critical for type I IFN-triggered signaling, we reasoned that LPS or virus up-regulates the expression of *Usp25* through type I IFN-triggered signaling. To test this hypothesis, we treated BMDCs with anti-IFNα, anti-IFNβ, or both followed by LPS stimulation or viral infection. As shown in Fig. 2B, blocking IFNα, IFNβ, or both strongly inhibited LPS- or virus-triggered induction of *Usp25* and *Ifi7*. Furthermore, LPS- or virus-induced up-regulation of *Usp25* was substantially diminished in *Iṣfα1−/−* MEFS and almost completely abolished in *Ifi7−/−* BMDCs compared with the wild-type controls (Fig. 2C). These data together suggest that LPS- or virus-induced expression mainly depends on type I IFN-triggered signaling.

**Type I IFN-induced Expression of Usp25 Is Dependent on TBK1/IKKe and de Novo Synthesized IRF7**—Considering that *Usp25* gene promoter contains two ISGF3 binding sites and that type I IFN-triggered signaling is critical for the up-regulation of *Usp25*, we hypothesized that type I IFNs activate transcription of *Usp25* through ISGF3. However, treatment with cycloheximide, a compound that inhibits mRNA translation, impaired IFNα- or IFNβ-induced expression of *Usp25*. In contrast, IFNα- or IFNβ-induced expression of *Ifi7*, a direct target of ISGF3, was not affected by cycloheximide treatment (Fig. 3A), indicating that virus-triggered type I IFN-mediated up-regulation of *Usp25* requires *de novo* protein synthesis. In addition, IFNα-triggered up-regulation of *Usp25* was impaired by knockdown of IRF7 but not IRF3 in MEFS and restored by reconstitution of IRF7 but not IRF3 into IRF3−/−IRF7−/− MEFS (Fig. 3, B and C), indicating that the *de novo* synthesized IRF7 is required for type I IFN-induced up-regulation of *Usp25*.

It has been recognized that TBK1- or IKKe-mediated phosphorylation of IRF7 is critical for its transcriptional activity. Consistent with this notion, we found that inhibition of TBK1 or IKKe impaired type I IFN-triggered up-regulation of *Usp25* (Fig. 4A). To further characterize the role of TBK1 and IKKe in type I IFN-triggered induction of *Usp25*, we reconstituted empty vector (Vec), TBK1, or TBK1(K38A) into *Tbk1−/−* MEFS and examined IFNα-triggered up-regulation of *Usp25*. As shown in Fig. 4B, reconstitution of TBK1 or TBK1(K38A) substantially promoted or inhibited IFNα-induced up-regulation of *Usp25* compared with reconstitution of the empty vector, respectively, indicating that TBK1(K38A) functions as a domi-
nant negative mutant regulating type I IFN-induced up-regulation of \textit{Usp25}. In addition, knockdown of IKKe in \textit{Tk1} \textsuperscript{−/−} MEFs reconstituted with the empty vector but not in those reconstituted with TBK1 significantly impaired IFNα-induced expression of \textit{Usp25} (Fig. 4C). We further transfected siRNA-resistant TBK1 (rTBK1) or IKKe (rIKKe) into wild-type

![Graphs](image-url)
MEFs followed by simultaneous knockdown of endogenous TBK1 and IKKε. Interestingly, IFNα-induced up-regulation of Usp25 was not affected in MEFs transfected with either rTBK1 or rIKKε (Fig. 4D). Furthermore, simultaneous knockdown of TBK1 and IKKε by siRNA significantly inhibited IFNα-induced expression of Usp25 in BMDCs (Fig. 4E). Taken together, these data suggest that TBK1 and IKKε function redundantly to regulate type I IFN-induced expression of Usp25.
To further confirm that IRF7 drives transcription of Usp25, we cloned the upstream 5000 bp starting from the transcription start site of Usp25 into the pGL3-Basic luciferase vector (USP25 promoter), made constructs with various mutations in the IRF or NF-κB binding sites, and performed luciferase reporter assays (Fig. 5A). Interestingly, IRF7 or SeV potently activated the luciferase activity of USP25 promoter, which was substantially impaired by mutations.
FIGURE 5. IRF7 binds to USP25 promoter. A, a schematic model of USP25 promoter and its mutations. B, HEK293 cells were transfected with the indicated luciferase (Luc.) reporter (100 ng) together with Vec, IRF7 (0.1 μg), IRF3 (0.5 μg) (left graph), p65, p50, or p52-RelB (0.2 μg) (middle graph). Twenty hours later, luciferase reporter assays were performed. HEK293 cells were transfected with the indicated luciferase reporter (100 ng). Twenty hours later, cells were infected with SeV for 8 h followed by luciferase reporter assays (right graph). C, HEK293 cells were transfected with the indicated luciferase reporter (100 ng) together with Vec, IRF7 (0.02–0.1 μg), or IRF3 (0.1–0.5 μg) (left graph). Twenty hours later, luciferase reporter assays were performed. Immunoblotting analysis was performed to examine the expression of transfected plasmids (right panels). D, Irf3−/−Irf7−/− MEFs were reconstituted with the empty vector (Irf3−/−Irf7−/− + Vector), IRF3 (Irf3+/−Irf7−/− + FLAG-IRF3), or IRF7 (Irf3−/−Irf7+/− + FLAG-IRF7) through lentivirus-mediated gene transfer. Cells were left untreated or infected with SeV for 12 h followed by ChIP analysis. TSS, transcription start site. Data shown are representatives of four (B) or three independent experiments (C and D). *p < 0.05; **p < 0.01; ***p < 0.001. Error bars represent S.D. Rel., relative.
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**TABLE 1**

qPCR primers of USP25 promoter and Ifnb promoter for ChIP analysis

| Primer name | Primer sequence |
|-------------|-----------------|
| USP25 IB1 (−1384 to −1319) | AAGTTACAGCGCTGAGCTCT |
| USP25 IB2 (−4706 to −4615) | TTGAGAGATCAGGAGGCTG |
| USP25 NS (−1018 to −907) | CTGTTTTTCGGCCGGATG |
| Ifnb promoter | GAGAGATCCGAGGAGGCTT |

**Discussion**

We have previously shown that LPS and viral infection strongly induce up-regulation of USP25. In this study, we further confirmed that LPS and viral infection activated transcription of **USP25** through type I IFN-triggered signaling. In addition, type I IFN-triggered signaling induced expression and protein synthesis of IRF7, which was activated by TBK1 and IKKε and bound to the USP25 promoter to activate transcription of **USP25** gene (Fig. 6).

Sequence analysis of USP25 promoter indicated that multiple NF-κB binding sites exist in the USP25 promoter. In our study, p65 deficiency did not affect virus-induced expression of **USP25**. Mutation of the five NF-κB binding sites did not affect basal, SeV-triggered, or IRF7/3-mediated activation of USP25 promoter. In addition, overexpression of p65, p50, or p52-RelB complex did not activate USP25 promoter, indicating that the NF-κB sites on USP25 promoter are dispensable for virus- or LPS-induced up-regulation of **USP25**. However, whether the NF-κB sites are involved in the induction of **USP25** by other stimuli is unknown. The USP25 promoter also contains two potential ISGF3 binding sites. We found that cycloheximide treatment impaired type I IFN-triggered induction of **USP25**, indicating that ISGF3 does not directly regulate transcription of **USP25** but instead activates de novo synthesis of other transcription factor(s) to mediate transcription of **USP25**.

IRF3 and IRF7 are two structurally related transcription factors that bind to the conserved IRF binding site (5'-GAAANNNGAAA-3') on the promoters and are essential for induction of hundreds of genes involved in innate immunity and inflammation. IRF3 exhibits more restricted DNA binding site specificity compared with IRF7. Mutation of a single nucleotide in either of the two GAAA core sequences impairs IRF3 binding and transcription activity, whereas the G and the third A in the GAAA core sequence are variable for IRF7 binding activity (36). According to this standard, IB1 “gaacataaa” and IB2 “gaatgaga” in **USP25** promoter are preferentially recognized and bound by IRF7 but not IRF3. Consistent with this observation, we found that (i) IRF7 but not IRF3 was sufficient to activate the IB1- or IB2-driven reporters and required for virus-triggered type I IFN-mediated up-regulation of **USP25**, (ii) IRF7 activated USP25 promoter more potently than did IRF3 in luciferase reporter assays, and (iii) IRF7 bound to the USP25 promoter more potently than did IRF3. However, it should be noted that IRF3 activated USP25 promoter (~2–4-fold) when transfected at a high dosage and partially rescued USP25 induction in IRf3−/−/IRf7−/− MEFs after viral infection. In addition, we observed that IFNAR1 deficiency in MEFs partially inhibited virus-triggered up-regulation of **USP25**, whereas IRFAR1 deficiency in BMDCs completely abolished up-regulation of **USP25** after viral infection, indicating that virus-induced expression of **USP25** might be differentially regulated by IRF3 and IRF7 in distinct types of cells. Taken together, it is likely that IRF3 is responsible for minimal expression of **USP25** in MEFs after viral infection, whereas the de novo synthesized IRF7 induced by type I IFNs is a master transcription factor for **USP25** expression in MEFs and BMDCs.

Unlike IRF3, which is constitutively expressed and resides in the cytosol, IRF7 is expressed at a low level and strongly induced by type I IFN-triggered signaling. Both IRF3 and IRF7 undergo TBK1- or IKKε-mediated phosphorylation, dimerization, and nuclear translocation after LPS treatment or viral infection. We found that treatment with TBK1 and IKKε inhibitor severely abolished type I IFN-triggered induction of **USP25**. Furthermore, reconstitution of TBK1 (K38A) into Tbk1−/− MEFs inhibited...
IFNα-induced up-regulation of Usp25, and knockdown of IKKε in Tbk1−/− + Vec MEFs but not in Tbk1−/− + TBK1 MEFs substantially impaired IFNα-induced up-regulation of Usp25, indicating that TBK1 and IKKε function redundantly for USP25 induction downstream of type I IFN stimulation. Further investigations are required to fully address how TBK1 and IKKε are involved in type I IFN-triggered signaling. Nonetheless, our data have clearly demonstrated that the type I IFN-IRF7 axis critically regulates viral infection- or LPS-induced transcription of Usp25 and contribute to our understanding of positive feedback regulation of cellular antiviral responses.

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