Type I IFN is siloed in endosomes

Jennie B. Altmana, Justin Tafta, Tim Wedekingb, Conor N. Gruberb, Michael Holtmannspötterb,c, Jacob Piehlerbd,e, and Dusan Bogunovicaf,g

aDepartment of Microbiology, Icahn School of Medicine at Mount Sinai, New York, NY, USA 10029; bDivision of Biophysics, Department of Biology, Osnabrück University, 49076 Osnabrück, Germany; cIntegrated Bioimaging Facility, Osnabrück University, 49076 Osnabrück, Germany; dCenter for Cellular Nanoanalytics, Osnabrück University, 49076 Osnabrück, Germany; eDepartment of Pediatrics, Icahn School of Medicine at Mount Sinai, New York, NY, USA 10029; fPrecision Immunology Institute, Icahn School of Medicine at Mount Sinai, New York, NY, USA 10029; and gMindich Child Heath and Development Institute, Icahn School of Medicine at Mount Sinai, New York, NY 10029

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Type I IFN (IFN-I) is thought to be rapidly internalized and degraded following binding to its receptor and initiation of signaling. However, many studies report the persistent effects mediated by IFN-I for days or even weeks, both ex vivo and in vivo. These long-lasting effects are attributed to downstream signaling molecules or induced effectors having a long half-life, particularly in specific cell types. Here, we describe a mechanism explaining the long-term effects of IFN-I. Following receptor binding, IFN-I is siloed into endosomal compartments. These intracellular “IFN silos” persist for days and can be visualized by fluorescence and electron microscopy. However, they are largely dormant functionally, due to IFN-I–induced negative regulators. By contrast, in individuals lacking these negative regulators, such as ISG15 or USP18, this siloed IFN-I can continue to signal from within the endosome. This mechanism may underlie the long-term effects of IFN-I therapy and may contribute to the pathophysiology of type I interferonopathies.

Type I interferon | endosome | cytokine retention

Type I IFN (IFN-I) is a potent antiviral and inflammatory cytokine, with a relatively short half-life (1). It disappears from the plasma several hours after intramuscular administration (2). Even pegylated IFN-I has a half-life of just 2 d in humans (3). Nevertheless, IFN-I has long-term functional effects. For example, expression of OAS1, an IFN-stimulated gene (ISG), remains close to peak levels for 1 wk in humans given pegylated IFN-I (4). Similarly, following influenza infection and resolution in mice, bone marrow leukocytes express antiviral genes and are resistant to influenza infection despite lack of detection of IFN-I in the bone marrow or serum on enzyme-linked immunosorbent assay (5). Likewise, individuals lacking ISG15 or USP18, both key negative regulators of IFN-I, have high levels of ISGs in peripheral blood mononuclear cells (PBMCs), as expected, but circulating IFN-I is only detectable in half of ISG15-deficient individuals (6), highlighting the potency of IFN-I. Here, we explore the cellular mechanisms governing this human phenotype.

Results/Discussion

Only half of ISG15-deficient patients have detectable circulating IFN-I, but all display high levels of ISGs in their PBMCs (6), a feature common in all type I interferonopathies (7). We investigated this phenomenon by quantifying ISG messenger RNA (mRNA) after 12 h of priming with IFN-I and 36 h of rest, in hTert-immortalized fibroblasts. At the end of this 48-h period, ISG mRNA levels were significantly higher in ISG15- and USP18-deficient cells than control cells (Fig. 1D). These high ISG mRNA levels persisted up to 5 d after IFN-I priming (mostly, therefore, in absence of cytokine) in ISG15-deficient cells, but returned to prepriming values in control cells after 24 h (8). Given the persistence of ISGs for several days after removal of IFN-I, we investigated whether ISGs accumulated due to a lack of down-regulation during the initial IFN-I exposure or due to active transcription after the cytokine was withdrawn. RT-qPCR on nascent RNA transcripts isolated 36 h to 60 h after the removal of IFN-I revealed active transcription of ISGs in ISG15- and USP18-deficient cells, but not control cells (Fig. 1B). Thus, transcription occurs long after the elimination of IFN-I in ISG15- and USP18-deficient cells.

We investigated molecules acting farther upstream and detected pSTAT1 and pSTAT2 (mediators of IFN-I proximal signaling) (9) in ISG15- and USP18-deficient cells, but not control cells (Fig. 1C), consistent with transcriptional initiation. Immunofluorescence (IF) staining for total STAT2 at this time point confirmed its nuclear localization (Fig. 1E). We ensured no infinitesimally small amount of residual soluble cytokine was causing this late transcription, by incubating cells with an anti–IFN-I antibody (Fig. 1F and G), and confirmed IFN-I was not being transcribed in the cells (Fig. 1H). We did detect minute amounts of IFN-I in the supernatant, but it was completely blocked by the IFN-I antibody (Fig. 1I). Together, these results demonstrate that IFN-I–mediated signaling can occur in the absence of soluble cytokine.

These results imply either that there is “cytokineless” signaling or that cytokine is present within the cells. We tested these hypotheses by performing volumetric fluorescence imaging after treatment with fluorescently labeled IFN-I. HeLa cells were stimulated with IFN-I for 17 h, washed, rested for 25 h, then imaged by lattice light sheet microscopy (LLSM). We detected IFN-I colocalized with Rab5, an early endosomal marker, demonstrating that IFN-I was being retained in what we call “IFN silos” (Fig. 2A), previously unappreciated sites of IFN-I storage. We detected these silos in all genetic backgrounds (Fig. 2B). While IFN-I receptor (IFNAR) is known to undergo endocytosis (10), this is documentation that IFN-I was present endosomally after signaling cessation. We demonstrate that IFNAR is required for the formation of active IFN-I silos (Fig. 2C). Using single molecule array (SiMoA) (Fig. 2D) and electron microscopy (EM), we confirmed IFN-I intracellular retention (Fig. 2E). Finally, while an endocytosis inhibitor minimally affected proximal signaling, it completely reversed the phenotype in cells lacking negative regulation (Fig. 2F).

The data presented here help resolve the paradox of the long-lasting effects of a cytokine after it is no longer detectable. This phenomenon is explained by the retention of IFN-I in Rab5+ endosomes after signaling, with the continuation of signaling largely prevented by USP18 in wild-type cells. Thus, signaling is abolished not by elimination of the cytokine but by the action of negative regulators. The relationship between the turnover of USP18 and the

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The authors declare no competing interest.

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1To whom correspondence may be addressed. Email: dusan.bogunovic@msm.edu.
Cells were hTert-immortalized dermal fibroblasts from control, ISG15-, or USP18-deficient patients. RT-qPCR was performed as previously described (8). For nascent RNA capture, ethynyl uridine (EU) was incubated for 24 h. The mRNA containing EU was isolated for RT-qPCR (12). Western blotting was performed as previously described (8). For confocal microscopy, cells were stained for DAPI and STAT2 (sc-476; 1:100). Leica SPS DMI microscope and Cell Profiler were used for analyses. For IFN-I blocking, anti-human IFNα antibody (PBL 31110-1) was used at 0.2 µg/mL, with sheep serum (Millipore S3772, 1:3 for equivalent concentration) as vehicle. Site-specific labeling of IFN was performed as previously described (13, 14). For fluorescence microscopy, cells were calcium phosphate transfected with pSEMS Rab5 mNeonGreen (pSEMS-26m Covalys), and treated with Cerdulatinib (Cerd) or vehicle (dimethyl sulfoxide [DMSO]) for 4 h following A, and Western blotting was performed. (F) IFN-I primed-rested lysates were run on a SiMoA device as a positive control, and RSAD2 mRNA was quantified. SEM represented, unpaired Student t tests: *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001. ns = not significant.

**Materials and Methods**

Cells were hTert-immortalized dermal fibroblasts from control, ISG15-, and USP18-deficient patients. RT-qPCR was performed as previously described (8). For nascent RNA capture, ethynyl uridine (EU) was incubated for 24 h. The mRNA containing EU was isolated for RT-qPCR (12). Western blotting was performed as previously described (8). For confocal microscopy, cells were stained for DAPI and STAT2 (sc-476; 1:100). Leica SPS DMI microscope and Cell Profiler were used for analyses. For IFN-I blocking, anti-human IFNα antibody (PBL 31110-1) was used at 0.2 µg/mL, with sheep serum (Millipore S3772, 1:3 for equivalent concentration) as vehicle. Site-specific labeling of IFN was performed as previously described (13, 14). For fluorescence microscopy, cells were calcium phosphate transfected with pSEMS Rab5 mNeonGreen (pSEMS-26m Covalys), and treated with Cerdulatinib (Cerd) or vehicle (dimethyl sulfoxide [DMSO]) for 4 h following A, and Western blotting was performed. (F) IFN-I primed-rested lysates were run on a SiMoA device as a positive control, and RSAD2 mRNA was quantified. SEM represented, unpaired Student t tests: *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001. ns = not significant.

**Data Availability.** All supporting data are included in the manuscript.
Fig. 2. IFN-I is retained intracellularly. (A) HeLa cells expressing Rab5-mNeonGreen (green) incubated with 5 nM DyLight IFNα2 (magenta) for 17 h, washed, rested for 24 h, and visualized by LLSM (Top). Untreated cells show background autofluorescence (Bottom). Z projections are of five sections (0.52-μm total thickness). Box plot shows correlation coefficients for colocalization of IFNα2 and Rab5 (n = 10 cells). (B) Control, ISG15−, and USP18-deficient cells were given DyLight IFNα2 for 17 h and imaged, or rested for 24 h and IFN-I vesicles quantified. (C) EM of Control or IFNAR2-deficient cells given 0.5 nM of IFN-I biotin for 15 min. Yellow arrows denote positive immunoreactivity. (D) Prime-rested cells evaluated by SiMoA, P = 0.0093. (E) EM of control cells given 0.5 nM of IFN-I biotin for 12 h, washed and rested in media containing 0.2 μg/ml anti-IFNα or vehicle (sheep serum) for 36 h. Representative example of (1) negative control without IFN-I biotin; (2) IFN-I biotin 12 h, 36 h rest; (3) IFN-I biotin 12 h, anti-IFNα during 36 h rest; and (4) IFN-I biotin 12 h, sheep serum during 36 h rest. Asterisk denotes unlabeled endosome; arrow denotes positive immunoreactivity within endosome; n = 50 cells per condition. (F) Cells were primed-rested in DMSO or 100 μM PitStop2, and qPCR was performed. Cells given IFN-I with DMSO or PitStop2 for 6 h served as controls. SEM represented, unpaired Student t tests: *P < 0.05, ****P < 0.0001.

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