| 1 2 | ETV7 represses a subset of interferon-stimulated genes that restrict influenza viruses |
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| 11 12 13 14 15 16 17 18 19 | |
| 20 21 | Keywords: |
| 22 23 24 25 26 | CRISPRa, screening, type I interferon, interferon-stimulated genes, gene regulation |
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42 Abstract

43 The type I interferon (IFN) response is an important component of the innate immune 44 response to viral infection. Precise control of interferon responses is critical, as insufficient 45 levels of interferon-stimulated genes (ISGs) can lead to a failure to restrict viral spread, 46 while excessive ISG activation results in interferon-related pathologies. While both 47 positive and negative regulatory factors can control the magnitude and duration of IFN 48 signaling, it is also appreciated that a number of ISGs regulate aspects of the interferon 49 response themselves. However, the mechanisms underlying complex ISG regulatory 50 networks remain incompletely defined. In this study, we performed a CRISPR activation 51 screen to identify new regulators of type I IFN responses. We identified ETS variant 52 transcription factor 7 (ETV7), a strongly induced ISG, as a protein that acts as a negative 53 regulator of the type I IFN response; however, ETV7 did not uniformly suppress ISG 54 transcription. Instead, ETV7 specifically targeted a subset of ISGs for regulation based 55 on their promoter sequences. We further showed the subset of ETV7-modulated ISGs is 56 particularly important for control of influenza viruses. Together, our data demonstrate that 57 ETV7 is a component of the complex ISG regulatory network by controlling the expression 58 of a subset of ISGs with a potential role in directing the interferon response against 59 specific viruses.

60 Significance

61 Interferons (IFNs) were first described in 1957 and are now known to be critical for 62 restriction of viruses. Still, our understanding of the complex web of interactions that 63 underlie IFN responses remains incomplete. In particular, negative regulation of interferon 64 responses has received disproportionately less study. In this work, we performed a 65 genome-wide overexpression screen for factors capable of suppressing IFN response 66 signaling. We identified a DNA binding transcription factor (ETV7) that, after induction by interferon, acts to suppress a subset of IFN-stimulated genes required for control of 67 68 influenza viruses. Our work highlights the importance of understanding negative IFN 69 signaling not only with respect to the magnitude and duration of the response, but also 70 the specificity of its antiviral effects.

71 Introduction

72 The type I interferon (IFN) response is a transient innate immune defense system that, 73 upon activation by viral infection, induces the transcription of hundreds of interferon-74 stimulated genes (ISGs) (1). Many ISGs have characterized antiviral roles that restrict viral replication by either interfering with viral processes directly or regulating the cellular 75 76 pathways required for viral replication (2). However, because replication mechanisms and 77 points of interaction with the cell differ between viruses, individual ISGs have varying 78 potencies against different viruses (3–5). As a result, unique combinations of ISGs are 79 thought to mediate successful antiviral responses against distinct viruses (1, 6).

80

81 The activation of the type I IFN signaling pathway in response to viral infection is well 82 understood (7, 8). Extracellular IFN, which is released after a cell recognizes virus-derived 83 nucleic acid, is bound by its cognate plasma membrane-localized receptor (IFNARs). 84 Downstream effectors (JAK proteins) are phosphorylated to then activate interferon-85 stimulated gene factor 3 (ISGF3) complex formation. Finally, the ISGF3 complex of 86 STAT1, STAT2, and IRF9 translocates to the nucleus (7). There, ISGF3 binds the 87 interferon sensitive response element (ISRE), with the consensus DNA motif 88 GAAANNGAAA, to activate transcription of ISGs (9).

89

As infection is cleared and virally derived innate immune activators become scarce, interferon production is reduced and the interferon-stimulated gene response is downregulated. To facilitate this return to cell homeostasis, negative regulators are induced and act at multiple levels in the signaling pathway (10). For example, PKD2 is an

94 ISG that recruits ubiquitin to the IFN receptor, IFNAR1, resulting in its degradation (11).
95 SOCS1 and SOCS3 are upregulated during, and act to limit, the IFN response through
96 direct interactions with JAK proteins, while SOCS1 also ubiquitinates other pathway
97 components (12). USP18 is induced by IFN to help return the cell to homeostasis by
98 removing the ubiquitin-like ISG15 from target proteins (13). Thus, negative regulators of
99 IFN responses are an important group of IFN-stimulated genes that control the duration
100 of ISG induction and activity.

101

102 In addition to activating or suppressing IFN responses, there are a number of interferon-103 induced regulators that enhance, limit, or fine-tune antiviral activity (14). Many ISGs 104 themselves participate in innate immune signaling to amplify IFN, and other pro-immune, 105 responses. For example, IFN signaling increases the levels of STAT1/2 and IRF9, thus 106 forming a positive feedback loop that enhances further ISG expression (15). Activators 107 also add complexity by inducing non-canonical IFN response pathways or specific groups 108 of ISGs. Interferon responsive factors (IRFs) 1 and 7 are ISGs and transcription factors 109 that activate subsets of ISGs (16, 17). Recent work has shown ELF1 (E74-like ETS 110 transcription factor) is induced by IFN, resulting in the expression of a group of genes not 111 otherwise activated by the IFN response (18). These differential ISG profiles are thought 112 to allow the cell to fine-tune its antiviral activity for an effective and appropriate response. 113 While interferon-induced positive regulators of the IFN response are known to shape the 114 complexity of ISG activation, reports of analogous roles for negative regulators remain 115 conspicuously absent.

116

117 To address this gap in knowledge and identify genes able to shape the IFN response 118 through negative regulation, we performed a CRISPR activation (CRISPRa) screen that 119 selected for factors sufficient to prevent expression of an ISRE-containing IFN response 120 reporter. We identified ETV7 (ETS variant transcription factor 7) as a negative regulator 121 of the type I IFN response with a role in controlling the expression of a subset of ISGs. 122 We further showed the ETV7-modulated ISGs are important for control of influenza 123 viruses. Together, these data demonstrate ETV7 is a suppressive component of the 124 complex ISG regulatory network that could be targeted to enhance specific antiviral 125 responses against influenza viruses (1, 19).

126

127 **Results**

<u>A CRISPR activation screen identifies potential negative regulators of the type I IFN</u> response.

130 In order to identify negative regulators of the type I IFN response, we developed a type I 131 IFN response reporter (IFNrsp) that included seven copies of the consensus interferon 132 sensitive response element (ISRE) ahead of a minimal CMV promoter controlling 133 expression of sfGFP (Fig. 1A). To make our reporter temporally specific, sfGFP was 134 fused to a mouse ornithine decarboxylase (MODC) protein degradation domain to 135 decrease its half-life (20). We stably introduced this construct into the A549 lung epithelial 136 cell line along with a dCAS9-VP64 fusion protein and a MS2-p65-HSF1 activator complex 137 required for the SAM CRISPR activation system (21). After clonal selection, 99.8% of the 138 A549-SAM-IFNrsp cells expressed GFP in response to type I IFN treatment (Fig. 1B and 139 **C**). To perform the screen, we took the A549-SAM-IFNrsp cell line and transduced 2×10^8

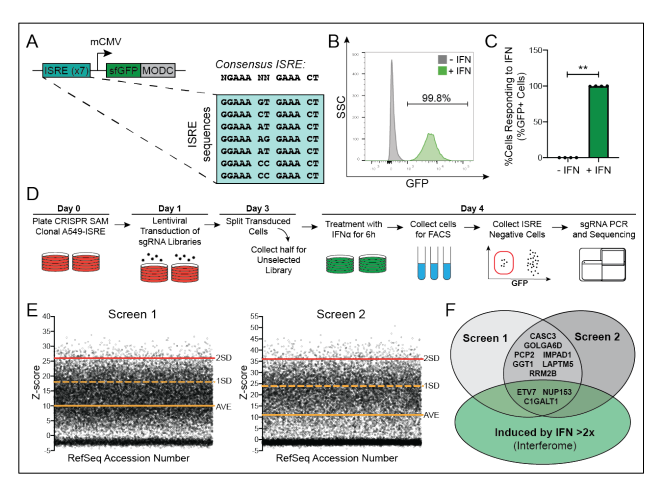


Fig. 1. A CRISPR activation screen to identify negative regulators of the type I interferon response. A) Diagram of the IFN response reporter (IFNrsp) used to identify cells responding to IFN. ISRE = interferon sensitive response element, MODC = protein degradation domain. B) Flow cytometry histogram and C) bar graph of A549-SAM-IFNrsp cells before and after IFN- α treatment (1000 U/mL, 6 h) (data shown as mean ± SD, n=4). Data shown are representative of two independent experiments. P-values calculated using unpaired, two-tailed Student's t-tests, *p<0.05, **p<0.001. D) Diagram of CRISPRa screen workflow to identify negative regulators of the type I IFN response. E) Results of the two independent CRISPRa screens. Z-score values from the replicate screens with a cutoff of 2 standard deviations from the mean were used to identify top "hits". F) Venn diagram indicating overlapping hits from the replicate screens and genes upregulated by interferon at least two-fold, according to the Interferome database (23).

- 140 cells at a multiplicity of infection (MOI) of 0.5 with a lentivirus library containing sgRNAs
- designed to activate every putative ORF in the human genome (21) (Fig. 1D). After 48
- 142 hours, half of the cells were collected to determine the transduction efficiency and the
- remaining cells were re-plated for IFN stimulation. At 72 hours post-sgRNA introduction,

144 the cells were treated with 4,000 U/mL IFN- α for 6 hours and collected for fluorescence-145 activated cell sorting. During sorting, we eliminated reporter-positive cells and collected 146 only cells that were nonresponsive to IFN, because this population should theoretically 147 be overexpressing a negative regulator of the IFN response. We performed two 148 independent biological replicates of the screen and sequenced the sgRNA-containing 149 amplicons derived from our input DNA, unselected transduced cells, and cells that were 150 nonresponsive to type I IFN. Raw sequencing data was aligned and mapped and 151 subsequently analyzed using the MAGeCK pipeline (22) to generate z-score values for 152 each gene. Genes were defined as "hits" if their z-scores exceeded two standard 153 deviations from the mean, resulting in an overlap of 10 genes between the two screen 154 replicates (Fig. 1E, Supplementary Data 1 and 2, and Supplementary Table 1). We 155 were seeking to identify regulators of the IFN response that are regulated by IFN 156 themselves; therefore, we selected hits for validation previously reported to have at least 157 a two-fold induction after IFN stimulation in the Interferome database (23). This analysis 158 identified three hits (C1GALT1, ETV7, and NUP153) as potential negative regulators of 159 the type I IFN response (Fig. 1F and Supplementary Table 1).

160

161 Overexpression of ETV7 is sufficient to negatively regulate the type I IFN response.

To validate our three hits, and to avoid potential false positive results as the result of offtarget effects of CRISPRa, we cloned the three ORFs and validated overexpression of the genes in 293T cells (**Supplementary Figure 1**). Co-transfection of the overexpression plasmids and IFNrsp plasmid, followed by stimulation with IFN- α , resulted in significantly fewer GFP-expressing cells compared to a control mCherry-expressing

167 plasmid (Fig. 2A and B). To verify this repressive activity was specific to the IFN response 168 and the hits were not general inhibitors of transcription or translation, we transfected the 169 overexpression plasmids along with a constitutively active GFP-expressing plasmid (Fig. 170 **2C**). We included a positive control (EIF2AK1/HRI), which is known to shut off translation 171 when overexpressed (24). C1GALT1 overexpression significantly downregulated GFP 172 expression, indicating the repressive activity of C1GALT1 is not completely specific to the 173 IFN response. While the overexpression of either ETV7 or NUP153 specifically affected 174 the IFNrsp plasmid, NUP153 has previously been shown to control the distribution of 175 STAT1 in the cell (25). We therefore chose ETV7 for further characterization because: 1) 176 ETV7 had not been previously reported to play a role in the IFN response, and 2) it had 177 the strongest inhibitory phenotype against the IFNrsp reporter.

178

179 After confirming overexpression of ETV7 at the protein level (Fig. 2D), we verified the 180 inhibitory effects of ETV7 were not restricted to the reporter plasmid. We collected mRNA 181 and protein from IFN- α stimulated ETV7 overexpression cells to quantify effects on the 182 expression of endogenous ISGs. ETV7 overexpression significantly repressed the 183 induction of three prototypical ISGs (IFIT1, MX1, and ISG15) at the RNA level (Fig. 2E). 184 The reduction of ISG expression during ETV7 overexpression was also demonstrated at 185 the protein level for IFIT1 (Fig. 2F). These experiments show that overexpression of ETV7 186 is sufficient to repress ISG induction by type I IFN.

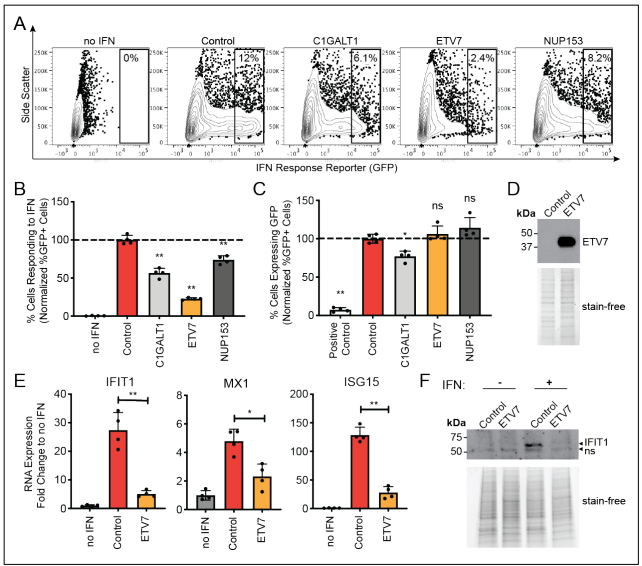


Fig. 2. ETV7 overexpression suppresses ISG expression. A) Flow cytometry plots of 293T cells transfected with the IFNrsp reporter and overexpression plasmids for the indicated screen hits then treated with IFN-a (100 U/mL, 6 h). B) Quantification of A showing normalized percentage of cells expressing GFP compared to the mCherryexpressing control (data shown as mean ± SD, n=4). C) Normalized percentage of cells expressing GFP from a constitutively expressing plasmid in cells overexpressing the indicated genes (positive control = EIF2AK1/HRI, shuts off translation) compared to control (data shown as mean ± SD, n=4). D) Western blot showing ETV7 protein levels in 293T cells transfected with the ETV7 overexpression plasmid. Stain-free gel imaging was used to confirm equal loading. E) Endogenous ISG mRNA expression levels measured using RT-gPCR after IFN-α treatment (100 U/mL, 9 h) (data shown as mean ± SD, n=4). F) Western blot comparing IFIT1 protein levels in control and ETV7 overexpressing cells after IFN-α treatment (500 U/mL, 18 h). ns = nonspecific band. Stainfree gel imaging was used to confirm equal loading. For all panels: Data shown are representative of two independent experiments. P-values calculated using unpaired, twotailed Student's t-tests (*p<0.05, **p<0.001) compared to IFN-stimulated, mCherryexpressing control samples.

189 ETV7 acts as a transcription factor to repress the type I IFN response.

| 190 | ETV7 is known to be a repressive transcription factor (26, 27), although a role in |
|-----|---|
| 191 | repressing type I IFN responses has never been reported. To determine whether ETV7 |
| 192 | acts as a transcription factor in this context, we generated a previously validated mutant |
| 193 | of ETV7, called ETV7(KALK), which is unable to bind DNA (Fig. 3A and B) (28). |
| 194 | Overexpression of ETV7(KALK) and stimulation with IFN- α had no measurable effect on |
| 195 | expression of the IFNrsp reporter, in contrast to WT ETV7 overexpression (Fig. 3C). |
| | |

196

197 ETV7 has been reported to bind the canonical ETS family DNA motif, GGAA (29), known 198 as an "ETS" site (Fig. 3D). Since consensus ISREs can either contain or lack a GGAA 199 motif (Supplementary Table 2), we hypothesized ETV7 could act on specific ISGs based 200 on the presence of ETS sites in their promoters. The original IFN response reporter design 201 contained multiple ETS sites (Fig. 3E), which potentially explains why it is negatively 202 impacted by ETV7. To test the requirements of ETS sites for ETV7 repressive activity, we 203 generated an IFN response reporter containing seven consensus ISREs from canonical 204 ISGs that all lack ETS sites (ISRE -ETS) (Fig. 3E). We transfected the two reporter 205 plasmids (ISRE +ETS and ISRE -ETS) independently into 293T cells and stimulated with 206 IFN- α . As expected, both reporter plasmids responded to IFN treatment, but the 207 repressive activity of ETV7 was restricted to the reporter plasmid containing ETS motifs 208 (Fig. 3F). These experiments together demonstrate that ETV7's repressive activity 209 requires both its ability to bind DNA and the presence of ETS sites in ISG promoters.

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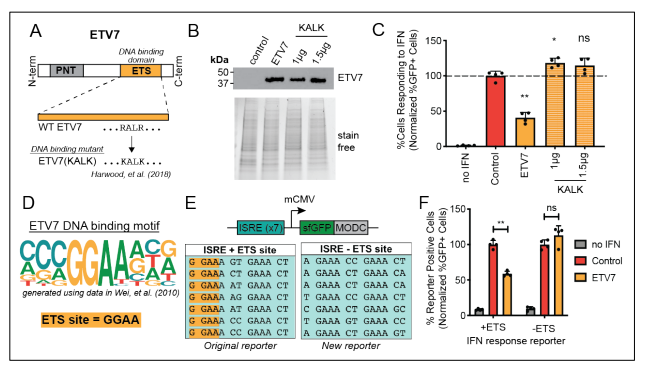


Fig. 3. ETV7 acts as a transcription factor to negatively regulate the type I IFN response. A) Diagram showing the ETV7 protein domains and amino acid changes made to generate the DNA binding mutant, ETV7(KALK). B) Western blot showing ETV7 protein levels in 293T cells transfected with WT (1µg) or DNA binding mutant (KALK) ETV7 expression plasmids. Stain-free gel imaging was used to confirm equal loading. C) Normalized percentage of 293T cells expressing GFP from the IFNrsp reporter with overexpression of WT or DNA binding mutant (KALK) ETV7 after IFN-α treatment (100 U/mL, 6 h) compared to control (data shown as mean ± SD, n=4, statistical analysis relative to IFN-stimulated, mCherry-expressing control samples). D) ETV7's DNA binding position weighted matrix (PWM) generated using enoLOGOS (69) with data from Wei et al. (70) and the conserved ETS family binding site, GGAA, highlighted in yellow. E) Diagrams of the IFNrsp reporters containing (+ETS) and not containing (-ETS) potential ETV7 binding sites (ETS site, highlighted in yellow). F) Normalized percentage of 293T cells expressing GFP from IFNrsp reporters either containing or not containing ETS sites after overexpression of ETV7 and IFN-α treatment (100 U/mL, 6 h) compared to mCherryexpressing control (data shown as mean \pm SD, n=4). For all panels: Data shown are representative of two independent experiments. P-values calculated using unpaired, twotailed Student's t-tests (*p<0.05, **p<0.001).

211 ETV7 differentially regulates genes based on their ISRE sequence.

- 212 Our data suggested ETV7 likely does not affect all ISG promoters. To perform an
- 213 unbiased examination of ETV7's repressive activity against ISGs with a variety of
- 214 potential regulatory sites, we performed RNA sequencing in cells with or without ETV7

215 overexpression and IFN stimulation (Supplementary Data 3). We then generated a 216 dendrogram using the 2,000 most differentially expressed genes after IFN treatment to 217 compare the four conditions: overexpression of control protein (mCherry) or ETV7, and 218 with or without IFN- α treatment. When comparing the impact of IFN treatment on control 219 and ETV7-overexpressing cells, we observed a larger divergence in the transcriptional 220 profile of control cells compared to ETV7-overexpressing cells after IFN treatment (Fig. 221 **4A**). This difference demonstrates that ETV7 generally "dampens" the transcriptional 222 impact of IFN treatment. Using a heat map to observe patterns in genes that increased 223 at least two-fold upon IFN treatment, we found some genes are more suppressed during 224 ETV7 overexpression than others (Fig. 4B). We divided these genes into three groups 225 (from I = most affected to III = least affected) depending on their response to ETV7 226 overexpression and we examined their promoters to identify motifs associated with ETS 227 transcription factors and IFN regulation. Unexpectedly, comparing the number of ETS 228 binding sites (GGAA) across these three groups revealed no significant difference 229 between the differentially affected groups (Fig. 4C). However, it is known that ETS sites 230 sometimes occur as a part of combined motif related to ISREs, known as ETS-IRF 231 combined elements (EICEs) with the consensus sequence GGAANN(N)GAAA (30, 31). 232 We therefore tested the hypothesis that ETV7 negatively regulates ISGs with EICE sites. 233 The number of EICE sites was significantly different between the most and least ETV7-234 affected groups (Fig. 4D), indicating ETV7 impacts the expression of specific ISGs by 235 targeting an extended DNA binding motif.

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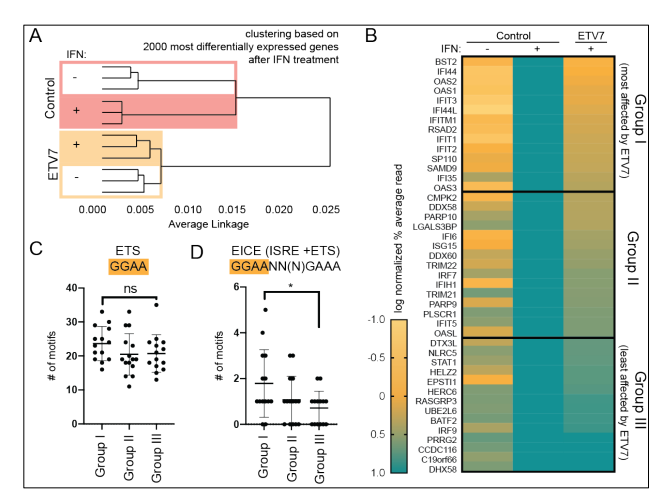


Fig. 4. ETV7 differentially regulates ISGs during the type I IFN response based on ISRE-related regulatory elements. A) Dendrogram of genes most differentially expressed in cells overexpressing either a control protein (mCherry) or ETV7 before and after IFN-α treatment (100 U/mL, 9 h) as measured using RNA sequencing. Three independent, biological replicates per condition. Red box highlights control samples, yellow box highlights ETV7-expressing samples, shading indicates IFN-stimulated samples. The box width indicates the linkage distance between samples before and after IFN, indicating control cells' transcriptional profile is more diverged after IFN treatment compared to ETV7-expressing cells. B) Heat map displaying RNA levels of genes upregulated at least two-fold following IFN-α treatment (100 U/mL, 9 h) in control cells. Expression was normalized to control cells after IFN treatment (averaged across replicates). Yellow = downregulated, blue = upregulated. C.D. Motif counts in promoter regions (-1000 bp, +500 bp) for the genes most and least affected by ETV7 overexpression in the RNA sequencing results. ETS sites (GGAA) highlighted in yellow. P-values calculated using unpaired, two-tailed Mann-Whitney U tests (*p<0.05, **p<0.001).

238

240 ETV7 is required to negatively regulate specific ISGs.

241 Our experiments to this point used an overexpression system to demonstrate that ETV7 242 is sufficient to suppress ISG expression. However, this approach leads to constitutive 243 ETV7 expression at high levels relative to the physiological magnitude and IFN-induced 244 expression of ETV7 (Fig. 5A). To determine the importance of ETV7 induction during the 245 IFN response, we performed a series of loss of function experiments that we expected 246 would have the reciprocal effect on IFN responses (32). We transduced A549-IFNrsp 247 reporter cells (the original reporter with ISRE +ETS sites) with Cas9 and one of two 248 different sgRNAs targeting ETV7 (ETV7 KO1, ETV7 KO2), selected for edited cells, and 249 then stimulated with IFN- α . Both guides resulted in significantly more IFN-induced sfGFP 250 expression compared to a control sgRNA (Fig. 5B). We next generated clonal ETV7 251 knockout A549 lung epithelial cell lines and sequenced the resulting DNA lesions to 252 confirm ETV7 knockout (Supplementary Figure 2). Since ETV7 is normally only 253 expressed after IFN stimulation, we treated with IFN- α and verified a reduction in ETV7 254 expression at the RNA level (**Fig. 5C**). We then selected five ISGs for RT-qPCR analysis. 255 Three (IFI44L, RSAD2/Viperin, IFIT3) were from the group most affected by ETV7 (Group 256 I) that contained multiple EICEs in their promoters. Two (IFIT5, IRF9) were chosen from 257 the less affected groups (Groups II and III). The EICE-rich genes (Group I) showed 258 significantly higher levels of RNA expression in the ETV7 knockout cells (Fig. 5D), while 259 genes with few EICEs were not significantly impacted by the loss of ETV7 (Fig. 5E). Thus, 260 the physiological level of ETV7 induction after IFN stimulation affects the expression of a 261 subset of ISGs.

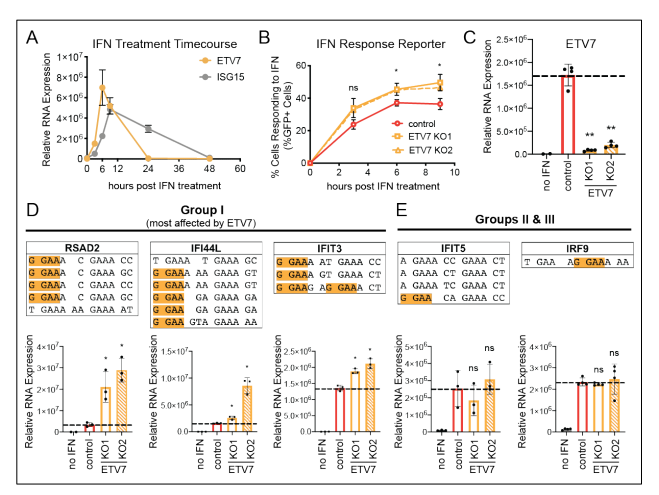


Fig. 5. ETV7 loss enhances expression of specific ISGs. A) ISG15 and ETV7 mRNA levels in A549 lung epithelial cells after IFN-α treatment (1000 U/mL, 6 h) (data shown as mean ± SD, n=4). B) Percentage of A549 cells expressing GFP from IFNrsp reporter after knockout of ETV7 and IFN-α treatment (1000 U/mL, 6 h) (data shown as mean ± SD, n=3, statistical analysis represents p-values for both of the two ETV7 KO sgRNAs compared to a non-targeting control). C) mRNA levels of ETV7 in non-targeting control and ETV7 KO A549 cells after IFN-α treatment (1000 U/mL, 6 h) (data shown as mean ± SD, n=4). D,E) Representative genes were chosen from the groups D) most affected by ETV7 (Group I) and E) least affected (Groups II and III) in the RNA sequencing analysis (**Fig. 4B**). Each gene's potential ISRE sequences (ETS sites highlighted in yellow) are shown, along with its mRNA levels in control and ETV7 KO cells after IFN-α treatment (1000 U/mL, 6 h) (data shown are representative of two independent experiments. P-values calculated using unpaired, two-tailed Student's t-tests (*p<0.05, **p<0.001) compared to IFN-stimulated, non-targeting sgRNA control samples unless otherwise indicated.

- 263 Loss of ETV7 restricts influenza viral replication.
- 264 For a successful antiviral response, individual ISGs are thought to work together to restrict
- 265 multiple parts of the virus replication cycle (1). To determine whether the effects of ETV7

266 suppression of ISG expression were relevant in the context of a viral infection, we wanted 267 to identify a virus restricted by the genes regulated by ETV7 (i.e. Group I genes) (33). 268 Considering the Group I genes with well recognized antiviral functions (IFITM1, IFIT1-3, 269 OAS1-3, BST2, RSAD2), we found each had been reported to play important roles in the 270 restriction of influenza viruses (34). IFITM1 has been shown to prevent viral entry (35), 271 OAS proteins activate RNase L to degrade viral RNA (36), IFITs bind viral RNA and 272 promote antiviral signaling (37), and BST2/Tetherin and RSAD2/Viperin restrict viral 273 budding and egress (38, 39).

274

275 To determine whether ETV7 regulation affected influenza virus infection, we first infected 276 our ETV7 knockout A549 cells with a laboratory-adapted H1N1 influenza A virus (IAV), 277 A/Puerto Rico/8/1934 (PR8). Using a hemagglutination (HA) assay to measure the 278 number of viral particles released over time, we observed reduced virus production in our 279 ETV7 KO cells compared to control cells (Fig. 6A). This was the anticipated outcome 280 because loss of a negative regulator (i.e. ETV7) is expected to enhance expression of 281 antiviral ISGs. We also measured infectious viral titers and found a significant reduction 282 in our ETV7 KO cells compared to control cells (Fig. 6B). Using a fluorescent reporter 283 strain of PR8 (PR8-mNeon) (40), we next visualized infection and spread. As expected, 284 we observed fewer cells expressing mNeon in ETV7 KO cells using both microscopy (Fig. 285 6C) and flow cytometry readouts (Fig. 6D). Next, we tested whether ETV7's impact on 286 influenza virus infection and spread would extend to a more contemporary H1N1 IAV 287 strain, A/California/07/2009 (Cal/09), as well as an unrelated Victoria lineage influenza B 288 virus strain, B/Malaysia/2506/2004 (Mal/04) (41). Using these fluorescent reporter

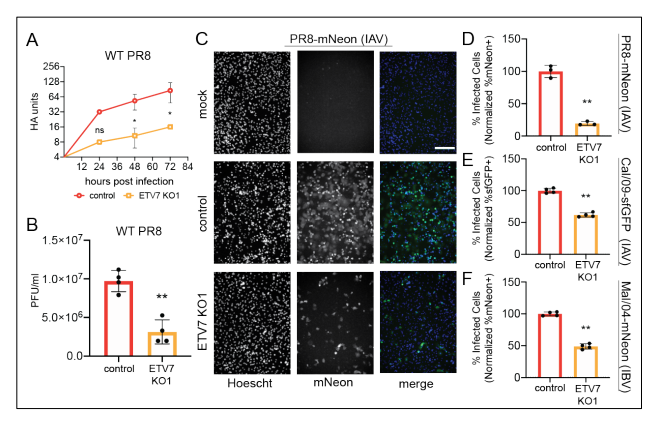


Fig. 6. Loss of ETV7 limits replication of multiple influenza viruses. A) Hemagglutination (HA) assay of virus collected at indicated time points from non-targeting control and ETV7 KO A549 cells after infection with WT PR8 virus (MOI=0.05, multicycle infection) (data shown as mean ± SD, n=3). B) Titer of virus collected from control and ETV7 KO A549 cells after infection with WT PR8 virus (18 h, MOI=0.05, multicycle infection) (data shown as mean ± SD, n=4). C) Control or ETV7 KO A549 cells after mock or PR8-mNeon reporter virus infection (24 h, MOI=0.1, multicycle infection). Green = mNeon, blue = nuclei. Scale bar, 200 µm. D) Flow cytometry quantification of control or ETV7 KO A549 cells after infection with PR8-mNeon reporter virus (24 h, MOI=0.01, multicycle infection) (data shown as mean ± SD, n=3). E,F) Normalized percentage of infected (reporter+) cells in ETV7 KO A549 cells compared to control cells after infection with E) Cal/09-sfGFP or F) Mal/04-mNeon reporter viruses (24 h, multicycle infection) (data shown as mean ± SD, n=4). For all panels: Data shown are representative of two independent experiments. P-values calculated using unpaired, two-tailed Student's ttests (*p<0.05, **p<0.001) compared to influenza infected, non-targeting sgRNA control samples.

- viruses, we observed significant decreases in the number of Cal/09- and Mal/04-infected
- cells when comparing ETV7 KO cells to control cells (**Fig. 6E and F**). These experiments
- 291 demonstrate that loss of ETV7 leads to decreased viral replication across multiple,
- 292 unrelated influenza viruses.

293 Discussion

294 In this study, we performed a CRISPR activation screen to identify negative regulators of 295 the type I IFN response. Specifically, we were interested in negative regulators that 296 contribute to the types of differentiated ISG profiles IFN-induced activators are reported 297 to produce. From this screen, we identified ETV7 as a negative IFN regulator and showed 298 it acts as a transcription factor to repress subsets of ISGs dependent on a motif related 299 to the ISRE, the EICE. We also showed ETV7's regulatory activity impacts the replication 300 and spread of multiple strains of influenza viruses. These findings demonstrate the 301 importance of ETV7 in fine-tuning the IFN response through specificity and transcriptional 302 repression to regulate particular, antiviral ISG targets.

303

304 ETV7 is a member of the ETS family of transcription factors. This family performs diverse 305 functions despite recognizing the same core DNA sequence, GGAA, by acting on 306 extended motifs requiring binding partners (42, 43). In our work we identified the EICE, a 307 recognized ETS transcription factor-associated motif, as a regulatory element related to 308 ISREs that ETV7 uses to discriminate genes for regulation. The EICE has previously been 309 reported to require an IRF binding partner to direct ETS transcription factor activity (30, 310 31); therefore, it is likely ETV7 has an IRF binding partner. If ETV7 does require a binding 311 partner, this protein's induction and distribution likely contribute to the timing, gene 312 targets, and activity of ETV7 during the IFN response. It is known that IRFs can be basally 313 expressed (IRF2, IRF3) or IFN-induced (IRF1, IRF7) (44) and the availability of a binding 314 partner could dramatically affect the timing and magnitude of effects on EICE-controlled 315 ISGs. Future work will define if ETV7 has specific binding partners and how those

interactions may contribute to the nonuniform, repressive activity of ETV7 during the typeI IFN response reported in this study.

318

IFN-induced regulators control the magnitude and duration of IFN responses in addition 319 to the temporal regulation of specific waves of ISGs (45). These coordinated waves of 320 321 ISG induction can peak early or late during the IFN response and are thought to 322 correspond to specific stages of virus replication or immune processes (1, 6). We 323 compared the induction of ETV7 and ISG15 and observed ETV7 is both upregulated and 324 downregulated at earlier time points than this prototypical ISG (Fig. 5A). We expanded 325 our analysis to published datasets of human gene expression during respiratory infections and concluded that ETV7 is generally induced earlier than many ISGs (46). Although not 326 327 the focus of our study, ETV7's early and short induction pattern suggests it may be a key 328 regulator of the first stages of IFN-mediated gene induction. We favor a model wherein 329 early ETV7 expression is responsible for reducing the accumulation, or delaying the 330 expression, of ISGs controlled by EICE motifs (**Supplementary Fig. 3**).

331

ETV7 is induced during infections across many vertebrate species (47, 48), indicating a potential conserved, relevant role in the immune response; however, ETV7 has been lost in mice and closely related rodents (49). Since mice and rodents have an intact interferon response pathway, a natural question is: how are the activities of ETV7 being accounted for in these animals? While we have no clear answer from the data in this study, it is wellrecognized that IFN responses contains many redundancies (33). Accordingly, we believe other ETS family members, potentially the closely related ETV6 (which is also induced by

339 IFNs), may perform the role of ETV7 in mice (50). Future studies will be required to test 340 the hypothesis that mice induce an ETV7-related alternative during the type I IFN 341 response.

342

343 Another important question is why ETV7's IFN-induced activity has been maintained 344 throughout evolution. In this report, we provide evidence that ETV7's activity reduces a 345 cell's ability to restrict influenza virus infection; this seems counterintuitive to ETV7 346 benefitting the host. We hypothesize that regulators like ETV7 are important to prevent 347 excessive inflammatory signaling. It is appreciated that negative regulators of the IFN 348 response are required to prevent extreme and prolonged immune responses, which are 349 associated with poor disease outcomes after infection (51-53). ETV7 potentially 350 contributes to the cumulative activities of negative IFN regulators to limit IFN responses 351 during pathogen clearance. Additionally, it stands to reason that different individual ISGs 352 have differing toxic effects on the cell. It is tempting to speculate that ETV7 suppresses 353 ISGs whose accumulation is particularly harmful to cell viability and host recovery after 354 infection.

355

Additionally, the relevance of controlled IFN responses goes beyond infectious disease; patients with dysfunctional USP18, a negative regulator of the IFN response, develop a type I interferonopathy that results in a severe pseudo-TORCH syndrome (54). Mouse knockouts for other negative regulators of the IFN response (SOCS1, SOCS3, USP18) also develop non-pathogen associated, chronic inflammatory diseases (55–58). ETV7's lack of murine homolog eliminates an easily generated animal-knockout model to

experimentally show ETV7's relevance as a general innate immune repressor. However, genome wide association studies (GWAS) have linked ETV7 to autoimmune diseases including rheumatoid arthritis and multiple sclerosis (59, 60); both of these autoimmune diseases have evidence of enhanced ISG expression (61, 62). Thus, although the specific contributions of ETV7 activity to IFN regulation are currently undefined, its potential role is not limited to viral infections.

368

369 In conclusion, we identified ETV7 as a negative regulator of the type I IFN response. 370 Previously, ETV7 was appreciated to be an ISG; however, a specific function during the 371 IFN response was unknown. We determined that ETV7 acts as a transcription factor to target specific ISGs for repression, potentially contributing to the complex ISG 372 373 transcriptional landscape. Additionally, many of the ETV7-modulated ISGs restrict 374 influenza viruses (34) and we showed that loss of ETV7 limits influenza virus spread. 375 Further work is required to understand the complexity of IFN regulation, while therapeutic 376 targeting of factors like ETV7 could lead to the development of a new class of host-377 directed antivirals that tailor ISG responses to specific viruses.

378 **Experimental Procedures**

379 Cloning

380 To generate reporters sensitive to IFN, we designed gBlocks (IDT) containing ISREs to 381 be cloned into the pTRIP vector ahead of a minimal CMV promoter controlling expression 382 of sfGFP. To clone and express the open reading frames (ORFs) of our screen hits, we 383 designed primers for cloning into the pLEX-MCS vector using Gibson Assembly (NEB). 384 To amplify ETV7 and NUP153, we used cDNA templates from Transomic Technologies. 385 To amplify C1GALT1 and EIF2AK1, we used RNA from IFN-stimulated A549 cells. The 386 DNA binding mutant, ETV7(KALK) (28), was also generated using a gBlock. Non-387 targeting and ETV7-targeting CRISPR KO sgRNAs were cloned by annealing oligos 388 encoding the desired sqRNA sequence and ligating them directly into the lentiCRISPRv2 389 vector (Addgene). DNA was transformed into NEB 5-alpha high efficiency competent 390 cells. Insert size was verified with PCR and purified plasmids were sequenced using 391 Sanger sequencing.

392

393 **Cells**

All cells were obtained from ATCC and grown at 37°C in 5% CO₂. A549 and 293T cells were grown in Dulbecco's Modified Eagle Medium (DMEM) supplemented with 5% fetal bovine serum, GlutaMAX, and penicillin-streptomycin. Madin-Darby canine kidney (MDCK) cells were grown in minimal essential media (MEM) supplemented with 5% fetal bovine serum, HEPES, NaHCO₃, GlutaMAX, and penicillin-streptomycin. The A549 CRISPR-SAM cells were previously validated (63) and transduced with the IFNrsp

- 400 reporter three times before being clonally selected. The A549 CRISPR KO cells were 401 transduced and then selected using puromycin (10 μ g/mL).
- 402

403 Flow Cytometry

- 404 Cells were trypsinized and analyzed on a Fortessa X-20 (BD) machine with standard laser
- 405 and filter combinations. Data was visualized and processed with FlowJo software.
- 406

407 CRISPR Activation Screen

408 The sgRNA library was packaged into lentivirus as previously described (63). After 409 packaging and titering the lentivirus, 2x10⁸ A549-CRISPR-SAM-IFNresp cells were 410 seeded onto 15 cm plates (10 plates total). The next day they were transduced with the 411 packaged sqRNA library (MOI=0.5). After 48 h, the transduced cells were split and half 412 were collected as a transduction control, while the remaining half were plated back onto 413 15 cm plates. The next day, cells were treated with IFN- α (4x10³ U/mL) for 6 h. Cells were 414 then collected and sorted on a Beckman Coulter Astrios cell sorter. Specifically, gates 415 were set to sort GFP-negative cells as the population of interest, as well as GFP-positive 416 cells as a control population of cells still capable of signaling. This screen was performed 417 in duplicate. Genomic DNA was extracted from sorted cells using the Zymo Quick gDNA 418 micro prep kit. PCR was subsequently performed using barcoded primers as previously 419 described using the NEB Next High Fidelity 2x PCR master mix (63). PCR bands were 420 gel purified using the Thermo GeneJET gel extraction kit. Samples were then sequenced 421 via next-generation Illumina MiSeg using paired-end 150 bp reads.

422

423 Screen Analysis

Raw MiSeq read files were aligned to the CRISPR SAM sgRNA library and raw reads for each sgRNA were counted using the MAGeCK pipeline (22). sgRNA enrichment was determined using the generated count files and the MAGeCK-MLE analysis pipeline. Genes were sorted based on z-score and determined to be significantly enriched if their z-score was at least two standard deviations above the average z-score of the entire sorted population.

430

431 Western Blotting

Cells were trypsinized and 1×10^6 cells were pelleted at 800 x g for 5 min. Equal amounts 432 433 of cellular material were loaded into 4-20% acrylamide gels (Bio-Rad) and imaged using 434 a ChemiDoc Imaging System (Bio-Rad). Protein was transferred to a nitrocellulose membrane at 60V for 60 min. PBS with 5% (w/v) non-fat dried milk and 0.1% Tween-20 435 436 were used to block for 1 h at 4°C. Primary antibodies were then incubated with the 437 membrane overnight at 4°C. Antibodies used were rabbit anti-ETV7 (Sigma, HPA029033) 438 and rabbit anti-IFIT1 (Cell Signaling, D2X9Z). Membranes were washed five times in PBS 439 with 0.1% Tween-20 and then an anti-rabbit-HRP secondary antibody (Thermo, A16104) 440 was added for 1 h. The membrane was then washed five times and Clarity or Clarity Max 441 ECL substrate (Bio-Rad) was added before being exposed to film and developed.

442

443 **RT-qPCR**

444 Total RNA was collected using Monarch Total RNA Miniprep Kits (NEB). One-step RT-445 qPCR was performed with commercial TaqMan assays from Thermo for ETV7

446 (Hs00903229 m1), C1GALT1 (Hs00863329 g1), NUP153 (Hs01018919 m1), ISG15 447 (Hs00196051 m1), MX1 (Hs00895608 m1), IFIT1 (Hs00356631 g1), RSAD2 448 (Hs00895608 m1), IFI44L (Hs00915292 m1), IFIT3 (Hs01922752 s1), IFIT5 449 (Hs00202721 m1), and IRF9 (Hs00196051 m1) using the EXPRESS One-Step 450 Superscript qRT-PCR Kit on an Applied Biosystems StepOnePlus instrument. RNA was 451 normalized using an endogenous 18S rRNA primer/probe set (Applied Biosystems).

452

453 **RNA sequencing**

454 293T cells were transfected with ETV7- or control-expressing plasmids and selected 455 using puromycin (20 μ g/mL) for 24 h before treatment with IFN- α (100 U/mL). Total RNA 456 was collected at 9 h post-IFN treatment using Monarch Total RNA Miniprep Kits (NEB). 457 RNA was prepped for RNA sequencing submission using the NEBNext Poly(A) mRNA 458 Magnetic Isolation Module (NEB), NEBNext Ultra II RNA Library Prep Kit for Illumina 459 (NEB), and NEBNext Multiplex Oligos for Illumina (NEB). Samples were analyzed on one 460 lane of an Illumina HiSeq 4000 using 50 bp single strand reads. Mapping of the raw reads 461 to the human hg19 reference genome was accomplished using a custom application on 462 the Illumina BaseSpace Sequence Hub (64). After data normalization, average read 463 values were compared across samples. For comparisons in which some samples had 464 zero reads detected for a specific gene, one read was added to all values in the sample 465 to complete analyses that required non-zero values. Dendrograms were generated by 466 identifying the 2,000 most differentially expressed genes in the control samples with and 467 without IFN treatment using a Student's t-test and plotted using Heatmapper (65). The 468 heat map shows genes upregulated 2-fold (after normalization) with IFN treatment in the

469 control samples. Values shown are log normalized to the control samples with IFN470 treatment.

471

472 Viruses

PR8-mNeon was generated via insertion of the mNeon fluorescent gene (66) into 473 474 segment 4 of the virus (40). Mal/04-mNeon was generated by inserting the mNeon 475 fluorescent gene (66) into segment 4 of the Mal/04 genome (41). Cal/09-sfGFP was 476 generated via insertion of the sfGFP gene (67) into segment 8 of the virus using the same 477 scheme previously used to insert Cre recombinase (68). For influenza virus infections, cells were either mock- or virus-infected for 1 h and then cultured in OptiMEM 478 479 supplemented with bovine serum albumin (BSA), penicillin-streptomycin, and 0.2 µg/mL 480 TPCK-treated trypsin protease (Sigma). PR8 WT, PR8-mNeon, Cal/09-sfGFP, and were 481 incubated at 37°C, Mal/04-mNeon was incubated ay 33°C.

482

483 Viral Growth Assays

484 Hemagglutination (HA) assays to measure the amount of viral particles were performed 485 by diluting influenza infected cell supernatants collected at the indicated time points in 486 cold PBS. An equal amount of chicken blood diluted 1:40 in PBS was mixed with serially 487 diluted virus and incubated at 4°C for 2-3 h before scoring. Infectious viral titers were 488 determined using standard plaque assay procedures on MDCK cells. Infected cell 489 supernatants were collected at 18 h, serially diluted, and used to infect confluent 6-well 490 plates for 1 h before removing the virus and adding the agar overlay. Cells were then 491 incubated at 37°C for 48 h before being fixed in 4% PFA overnight. The 4% PFA was then

492 aspirated, and the agar layer was removed before washing cells with PBS. Serum from 493 WT PR8 infected mice was diluted 1:2,000 in antibody dilution buffer (5% (w/v) non-fat 494 dried milk and 0.05% Tween-20 in PBS) and incubated on cells at 4°C overnight. Cells 495 were then washed twice with PBS and incubated for 1 h with anti-mouse IgG horseradish 496 peroxidase (HRP)-conjugated sheep antibody (GE Healthcare) diluted 1:4,000 in 497 antibody dilution buffer. Assays were then washed twice with PBS and exposed to 0.5 mL 498 of TrueBlue peroxidase substrate (KPL) for 20 min. Plates were then washed with water 499 and dried before plagues were counted.

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501 Data Availability

All next generation sequencing data are available at NCBI GEO under accession numberGSE140718.

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682 Acknowledgements

683 We would like to acknowledge Brook Heaton for generating the A549-CRISPR-SAM-684 IFNrsp cell line. We acknowledge assistance from Mike Cook and the Duke Cancer 685 Institute Flow Cytometry Core. We thank Robert Lefkowitz and his laboratory for assistance with and use of the ChemiDoc Imaging System. We also thank Ephraim Tsalik 686 687 and Micah McClain for helpful discussions and Ben Chambers, Stacy Webb, and other 688 members of the Heaton lab for critical reading of the manuscript. H.M.F. and A.T.H. were 689 supported by NIH training grant T32-CA009111. N.S.H. is supported by R01-HL142985, 690 R01-AI137031, and funding from the Defense Advanced Research Projects Agency's 691 (DARPA) PReemptive Expression of Protective Alleles and Response Elements (PREPARE) program (Cooperative agreement #HR00111920008). The views, opinions 692 693 and/or findings expressed are those of the author and should not be interpreted as 694 representing the official views or policies of the U.S. Government.

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696 Author Contributions

A.T.H. and N.S.H. designed the screen. A.T.H performed the screen experiments and initial analysis. H.M.F. and N.S.H. designed the screen validation studies. H.M.F. performed the screen hit determination analysis, flow cytometry, RNA and protein quantification experiments, sequencing and promoter analysis, and virus infection experiments. A.T.H. generated the Cal/09-sfGFP virus. H.M.F. and N.S.H. wrote the manuscript.

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