

RNase H2 catalytic core Aicardi-Goutières syndrome-related mutant invokes cGAS–STING innate immune-sensing pathway in mice

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The neuroinflammatory autoimmune disease Aicardi-Goutières syndrome (AGS) develops from mutations in genes encoding several nucleotide-processing proteins, including RNase H2. Defective RNase H2 may induce accumulation of self-nucleic acid species that trigger chronic type I interferon and inflammatory responses, leading to AGS pathology. We created a knock-in mouse model with an RNase H2 AGS mutation in a highly conserved residue of the catalytic subunit, *Rnaseh2a*^{G37S/G37S} (G37S), to understand disease pathology. G37S homozygotes are perinatal lethal, in contrast to the early embryonic lethality previously reported for *Rnaseh2b*- or *Rnaseh2c*-null mice. Importantly, we found that the G37S mutation led to increased expression of interferon-stimulated genes dependent on the cGAS–STING signaling pathway. Ablation of STING in the G37S mice results in partial rescue of the perinatal lethality, with viable mice exhibiting white spotting on their ventral surface. We believe that the G37S knock-in mouse provides an excellent animal model for studying RNASEH2-associated autoimmune diseases.

Aicardi-Goutières syndrome (AGS) is a rare neuroinflammatory disorder (Crow et al., 2015). In most cases, it is present at birth with symptoms indistinguishable from those associated with congenital viral infection, such as elevated levels of type I IFN in the serum and cerebrospinal fluid. AGS arises from mutations in seven different genes: *RNASEH2A*, *RNASEH2B*, *RNASEH2C*, *TREX1*, *SAMHD1*, *ADAR1*, and *IFIH1*, all of which are nucleic acid-transacting enzymes. AGS is believed to result from activation of the innate immune pathway by nucleic acids accumulating in the cytosol when an AGS-associated gene is defective. AGS shares some disease hallmarks with systemic lupus erythematosus (Günther et al., 2015). Over 50% of AGS patients carry biallelic mutations in the genes encoding the three subunits of the heterotrimeric RNase H2 complex (RNase H2A, RNase H2B, and RNase H2C; Crow et al., 2015).

RNase H2 provides the main RNase H activity in humans (Cerritelli and Crouch, 2009) and is essential for removing ribonucleotides incorporated in genomic DNA during replication, as well as for resolving R-loops formed during transcription (Nick McElhinny et al., 2010; Reijns

et al., 2012; Chon et al., 2013). The crystal structures of human and mouse RNase H2 revealed the interactions of the subunits and the positions of the more than 50 known AGS-related mutations in the three subunits (Figiel et al., 2011; Reijns et al., 2011; Reijns and Jackson, 2014). Some mutations are located near the catalytic center and affect catalysis, whereas others affect stability or alter protein interactions. The most common mutations reported in AGS patients are found in the B subunit (Crow et al., 2015) and are associated with less severe disease phenotype than mutations in the catalytic A subunit. A mutation in a highly conserved glycine (G37S) near the catalytic center on the RNase H2A subunit causes a severe early onset presentation of AGS, likely as a result of a substantial loss of RNase H activity (Crow et al., 2006). Many in vitro studies showed that Serine substitution for Glycine 37 in RNases H2 of eukaryotes reduces RNase activity (Crow et al., 2006; Rohman et al., 2008; Chon et al., 2009; Coffin et al., 2011). Mouse models using deletions of *RNASE2B* and *RNASEH2C* exhibit significant DNA damage, resulting in embryonic lethality at E9.5 (Hiller et al., 2012; Reijns et al., 2012). These mice have elucidated important information on the role of RNase H2 in genome stability, but because of their early death, have not yielded insight into the innate immune pathways responsible for disease mani-

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Abbreviations used: AGS, Aicardi-Goutières syndrome; CBX, carbenoxolone; ISG, IFN-stimulated gene; VSV, vesicular stomatitis virus.

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festation. Likewise, neither do mice with residual levels of RNase H2B (R2B KOF; Hiller et al., 2012), or the viable and asymptomatic *Rnaseh2b*^{A174T/A174T} mouse model (Reijns and Jackson, 2014). Therefore, it remains unclear how *RNASEH2* mutations lead to the development of AGS.

RESULTS AND DISCUSSION

G37S homozygous mice are perinatal lethal, and G37S embryos show increased expression in IFN-stimulated genes

We generated *Rnaseh2a*-G37S knock-in mice to mimic the exact mutation present in AGS patients. *Rnaseh2a*^{G37S/+} mice were viable, with no evident deleterious phenotype. Breeding of *Rnaseh2a*^{G37S/+} mice generate still-born pups that were homozygotes for the mutation (Fig. 1 A). No viable G37S homozygotes were observed (dead at or within hours after birth). The G37S homozygote embryos were smaller from an early embryogenesis period of E10.5, present at expected Mendelian ratio, and were ~20% smaller in size compared with their uterine mates (Fig. 1 A). Magnetic resonance imaging and micro-computed tomography scans did not reveal any additional phenotypic abnormalities in *Rnaseh2a*^{G37S/G37S} embryos in the brain (Fig. 1 B) or elsewhere (not depicted). We also did not observe any inflammation in histology staining of tissues of *Rnaseh2a*^{G37S/G37S} embryos, including the brain (unpublished data). The lack of neuroinflammation could be caused by early death of the animal, or by different disease presentation in mouse versus human. Other AGS mouse models such as *Trex1*^{-/-} or *Samhd1*^{-/-} also lack evidence of neuroinflammation (Gall et al., 2012; Rehwinkel et al., 2013).

We next examined where we can detect a molecular signature of immune activation in *Rnaseh2a*^{G37S/G37S} embryos, as it would be expected from its association with AGS. We performed RNA-seq analysis comparing gene expression profiles of WT, *Rnaseh2a*^{G37S/+}, and *Rnaseh2a*^{G37S/G37S} primary MEFs isolated from E13.5 embryos. We found that 388 genes were up-regulated twofold or more in *Rnaseh2a*^{G37S/G37S} MEFs compared with WT; of those, the most enriched gene network was “immune response” (DAVID GO term analysis; Fig. 1, C and D). Many of the highly up-regulated genes in *Rnaseh2a*^{G37S/G37S} cells were IFN-stimulated genes (ISGs), such as *Ifit44*, *Usp18*, *Ifit1*, *Rsad2*, *Isg15*, *Irf7*, and *Cxcl10* (Fig. 1 D). We validated increased expression of *Ifit1*, *Ifit3*, *Rsad2*, and *Cxcl10* by quantitative RT-PCR (Figs. 1 E and 2). As ISGs provide defense mechanisms against viral infection, we infected WT, *Rnaseh2a*^{G37S/+}, and *Rnaseh2a*^{G37S/G37S} primary MEFs with vesicular stomatitis virus (VSV)-PeGFP, to assess their functionality. We found that *Rnaseh2a*^{G37S/G37S} MEFs were highly refractory to VSV infection, as measured by reduced GFP fluorescence at 24 h or by reduced VSV G and M RNA from 6 to 24 h after infection (Fig. 1 F). Collectively, our data showed that the homozygous G37S mutation in mice invokes innate immune activation of ISGs, similar to that of AGS patients (Crow et al., 2015).

Immune activation in *Rnaseh2a*^{G37S/G37S} primary MEFs requires the cGAS-STING innate immune pathway

We next aimed to determine the signaling pathways responsible for the innate immune activation in *Rnaseh2a*^{G37S/G37S} cells. Many of the up-regulated ISGs we observed are direct targets of the transcription factor IRF3 (Diamond and Farzan, 2013; Lazear et al., 2013), which is activated by phosphorylation by TBK1. We treated *Rnaseh2a*^{G37S/G37S} cells with a TBK1 inhibitor BX795, and observed reduced expression of activated genes, *Cxcl10*, *Ifit1*, and *Rsad2* in *Rnaseh2a*^{G37S/G37S} cells (Fig. 2 A), suggesting the involvement of a cytosolic immune-sensing pathway. Using shRNA directed against *Mavs* or *Sting* (adaptor proteins for cytosolic RNA or DNA sensing, respectively) in *Rnaseh2a*^{G37S/G37S} cells, we found that shRNA against *Sting* restored the low expression levels of *Cxcl10* mRNA to WT levels. In contrast, *Mavs* knockdown significantly reduced poly(I:C)-induced IFN response, but had no effect on the G37S-induced ISG expression (Fig. 2, B and C), suggesting that the G37S mutation leads to activation of a DNA- but not RNA-sensing pathway.

We next used shRNA to knock down components of the cytosolic DNA-sensing pathway in *Rnaseh2a*^{G37S/G37S} cells, and then examined expression of ISGs. shSting and shTbk1 effectively restored mRNA levels to those seen in WT cells (Fig. 2 D). A prominent driver of cytosolic DNA detection, cGAS, responds to microbial or self-DNA and long, homopolymeric RNA/DNA hybrids (Ablasser et al., 2014; Mankan et al., 2014; Gao et al., 2015; Gray et al., 2015). shRNA-mediated knockdown of cGAS in *Rnaseh2a*^{G37S/G37S} cells also returned ISG expression to low WT levels (Fig. 2 D). As the production of cGAMP by cGAS leads to activation of STING-mediated signaling, we next examined the presence of cGAMP in *Rnaseh2a*^{G37S/G37S} cells using the gap junction transfer cGAMP bioassay (Ablasser et al., 2013). cGAMP can be transferred from producing cells to neighboring cells through gap junctions, thereby enabling a co-culture-based transactivation assay for cGAMP detection (Ablasser et al., 2013). We co-cultured *Rnaseh2a*^{G37S/G37S} MEFs (producing cells) with human fibroblasts (target cells), and then used human-specific primers to probe the immune activation status of the human cells. *Rnaseh2a*^{G37S/G37S} MEFs induced strong immune activation of the human ISGs tested, whereas WT MEFs had no effect (Fig. 2 E). Importantly, we also found that inhibiting gap junctions with carbenoxolone (CBX), a non-toxic gap junction inhibitor (Fig. 2 E), or trans-well separation completely abolished *Rnaseh2a*^{G37S/G37S} MEFs transactivation activity (Fig. 2 F). These data suggest that *Rnaseh2a*^{G37S/G37S} primary MEFs produce cGAMP that activates the STING pathway.

Sting^{-/-} partially rescues perinatal lethality of G37S mice

In the AGS mouse model of *Trex1* gene deletion, viability of *Trex1*^{-/-} mice is extended dramatically when the response to secreted type I IFN is ablated by deleting the IFN receptor, *Infar1*, or by eliminating the adaptive immune response through *Rag2* gene deletion (Stetson et al., 2008). We thus bred G37S to *Infar1*^{-/-} or *Rag2*^{-/-} backgrounds and found that nei-

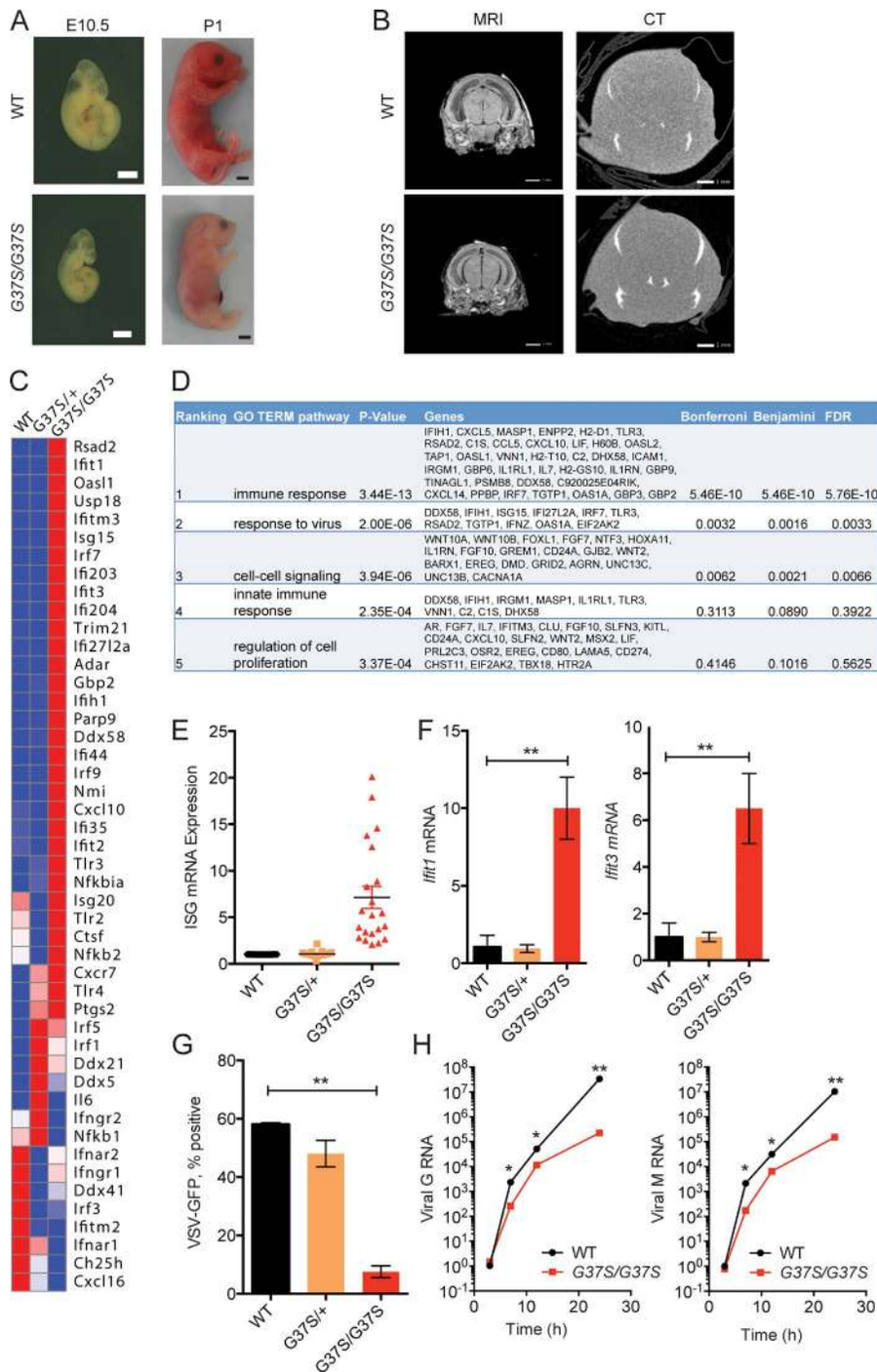


Figure 1. Primary cells from *Rnaseh2a*^{G37S/G37S} embryos show increased expression in ISGs. (A) Images of WT and *Rnaseh2a*^{G37S/G37S} embryos at indicated stage (bottom). Bars, 2 mm. (B) Transcranial images of ex vivo mouse E18.5 embryos WT and *Rnaseh2a*^{G37S/G37S} in MRI gradient echo image and micro computed tomography scans. No appreciable calcium, other than in the bones of the cranium (top arrow) and jaw (bottom arrow), which are still under formation. Bars, 1 mm. (C) A heat map of immune gene expression in WT, *Rnaseh2a*^{G37S/+} and *Rnaseh2a*^{G37S/G37S} primary MEFs. Data from RNA-seq (Table S1). (D) Gene ontology analysis of 388 genes that are increased by twofold or more in *Rnaseh2a*^{G37S/G37S} compared with WT MEFs. Top five enriched pathways are shown. (E) Expression of ISGs in WT, *Rnaseh2a*^{G37S/+}, and *Rnaseh2a*^{G37S/G37S} primary MEFs. Each dot represents a different ISG. Data from RNA-seq. (F) Quantitative RT-PCR analysis of *Ifi1* and *Ifi3* mRNA (ISGs) in WT, *Rnaseh2a*^{G37S/+}, and *Rnaseh2a*^{G37S/G37S} primary MEFs. (G and H) VSV-GFP replication in WT, *Rnaseh2a*^{G37S/+}, and *Rnaseh2a*^{G37S/G37S} primary MEFs. FACS analysis measures VSV-GFP signal at 24 h after infection (G). Quantitative RT-PCR analysis of VSV G and M RNA measure viral RNA replication at indicated time after infection (H). *, $P < 0.05$; **, $P < 0.01$. Mice were compared with littermate controls. Data are representative of at least three independent experiments. Error bars represent the SEM. Unpaired Student's *t* test (F–H).

ther of these genetic knockouts was able to rescue the perinatal lethal phenotype (Table 1). These findings are consistent with lack of inflammation in the *Rnaseh2a*^{G37S/G37S} embryos, further suggesting cell-intrinsic defects may be responsible. Partial rescue of the embryonic development defect seen in *Rnaseh2b*-null mice was obtained by deleting the *p53*^{-/-} gene (Reijns et al., 2012). However, we found *Rnaseh2a*^{G37S/G37S} *p53*^{-/-} mice also exhibited perinatal lethality indistinguishable

from *Rnaseh2a*^{G37S/G37S} alone. We also did not observe elevated expression of p53 pathway genes from our RNA-seq analysis (unpublished data).

We next bred *Rnaseh2a*^{G37S/G37S} onto *Mavs*^{-/-} or *Sting*^{-/-} background. *Rnaseh2a*^{G37S/G37S} remains perinatal lethal on *Mavs*^{-/-} background (Table 1), consistent with RNA-sensing pathways being uninvolved (Fig. 2). Remarkably, we obtained viable pups of *Rnaseh2a*^{G37S/G37S}

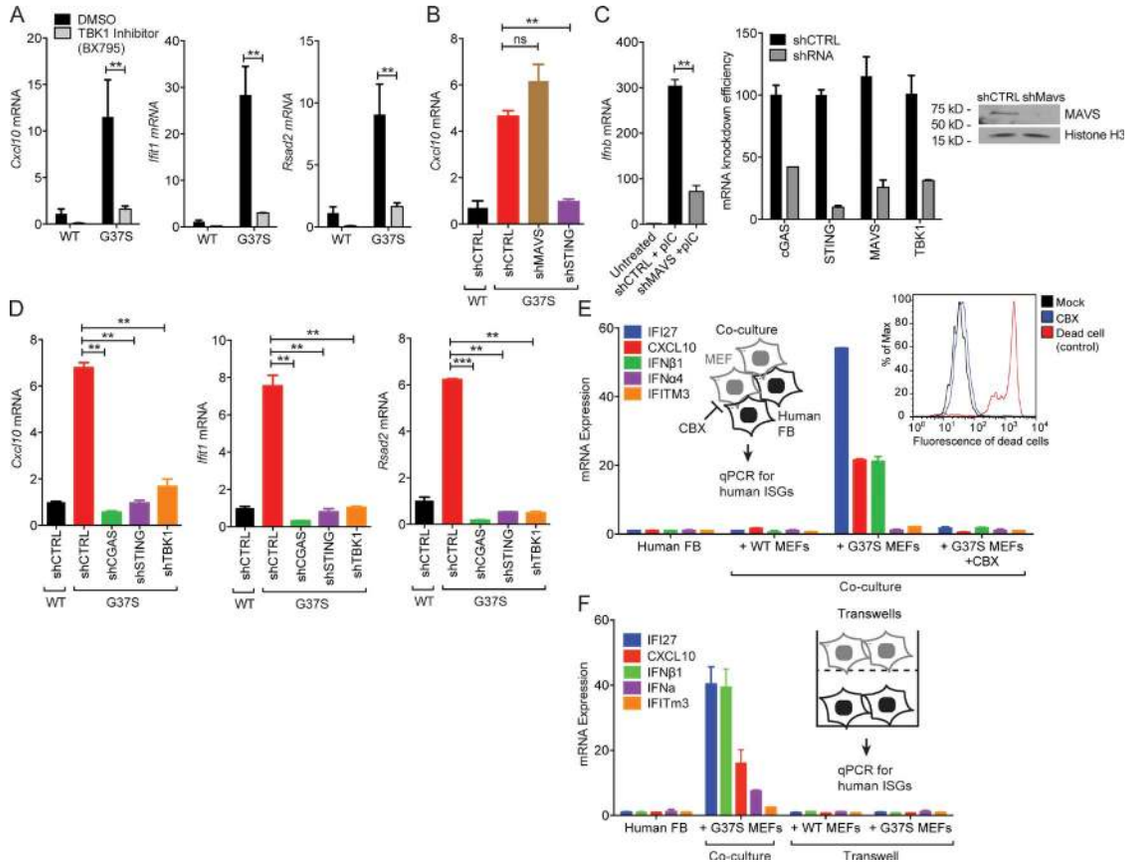


Figure 2. Immune activation in *Rnaseh2a*^{G37S/G37S} primary MEFs requires the cGAS–STING innate immune pathway. (A) Quantitative RT-PCR analysis of *Cxcl10*, *Ifi1* and *Rsad2* mRNA (all ISGs) in WT and *Rnaseh2a*^{G37S/G37S} (G37S, same below) MEFs treated with DMSO or TBK1 inhibitor BX795 (10 μM) for 6 h. (B) Quantitative RT-PCR analysis of *Cxcl10* mRNA in WT and G37S MEFs treated with shRNA against indicated genes involved in cytosolic nucleic acid-sensing. (C) shMAVS knockdown reduces poly(I:C)-induced IFN response. Knockdown efficiency is shown on the right. (D) Quantitative RT-PCR analysis of *Cxcl10*, *Ifi1*, and *Rsad2* mRNA in WT and G37S MEFs treated with shRNA against indicated genes involved in DNA-sensing pathway. (E) Quantitative RT-PCR analysis of a panel of human ISGs and IFN genes in human fibroblasts (BJ-1 cells) co-cultured with WT or G37S MEFs for 18 h, with or without CBX treatment (inhibits gap junction). Left inset shows a schematic diagram of the gap junction assay. Right inset shows FACS analysis of cell death in mock- and CBX-treated cells. (F) Quantitative RT-PCR analysis of human ISGs in human fibroblasts in a trans-well assay co-cultured with WT or G37S MEFs for 18 h. Mice were compared with littermate controls. **, P < 0.01; ***, P < 0.001. ns, not significant. Data are representative of at least three independent experiments. Error bars represent the SEM. Unpaired Student's t test (A–D).

Sting^{-/-} genotype, albeit at 6% of expected frequency (or 2% of weaned pups from heterozygous crosses on the *Sting*^{-/-} background; Table 1). Most of the *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} still exhibited perinatal lethality, similar to *Rnaseh2a*^{G37S/G37S} *Sting*^{+/-} or *Sting*^{+/+} (Table 1). Mice were either perinatal lethal or survived after birth. The failure to rescue any le-

thality in *G37S*^{-/-} *Ifnar1*^{-/-} could reflect the limited number of offspring examined. We compared ISG expression in primary E14.5 MEFs, and found that *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} completely returned ISG expression to the low level in *Rnaseh2a*^{+/+} *Sting*^{-/-}, whereas *Mavs*^{-/-} had no effect (Fig. 3 A). We bred *Rnaseh2a*^{G37S/G37S} onto *cGAS*^{-/-} back-

Table 1. Genetic crosses of G37S mice

Mice	<i>p53</i> ^{-/-}		<i>INFAR</i> ^{-/-}		<i>Rag2</i> ^{-/-}		<i>Mavs</i> ^{-/-}		<i>Sting</i> ^{-/-}	
	Neonates (no. embryos)	Weaned pups (no. mice)	Neonates (no. embryos)	Weaned pups (no. mice)	Neonates (no. embryos)	Weaned pups (no. mice)	Neonates (no. embryos)	Weaned pups (no. mice)	Neonates (no. embryos)	Weaned pups (no. mice)
WT	23% (7)	33% (17)	24% (8)	44% (67)	26% (10)	32% (30)	22% (4)	41% (52)	23% (7)	32% (95)
<i>G37S</i> ⁺	58% (18)	67% (34)	52% (17)	56% (87)	53% (20)	68% (65)	56% (10)	59% (74)	60% (18)	66% (196)
<i>G37S</i> / <i>G37S</i>	19% (6)	0% (0)	24% (8)	0% (0)	21% (8)	0% (0)	22% (4)	0% (0)	17% (5)	2% (6)

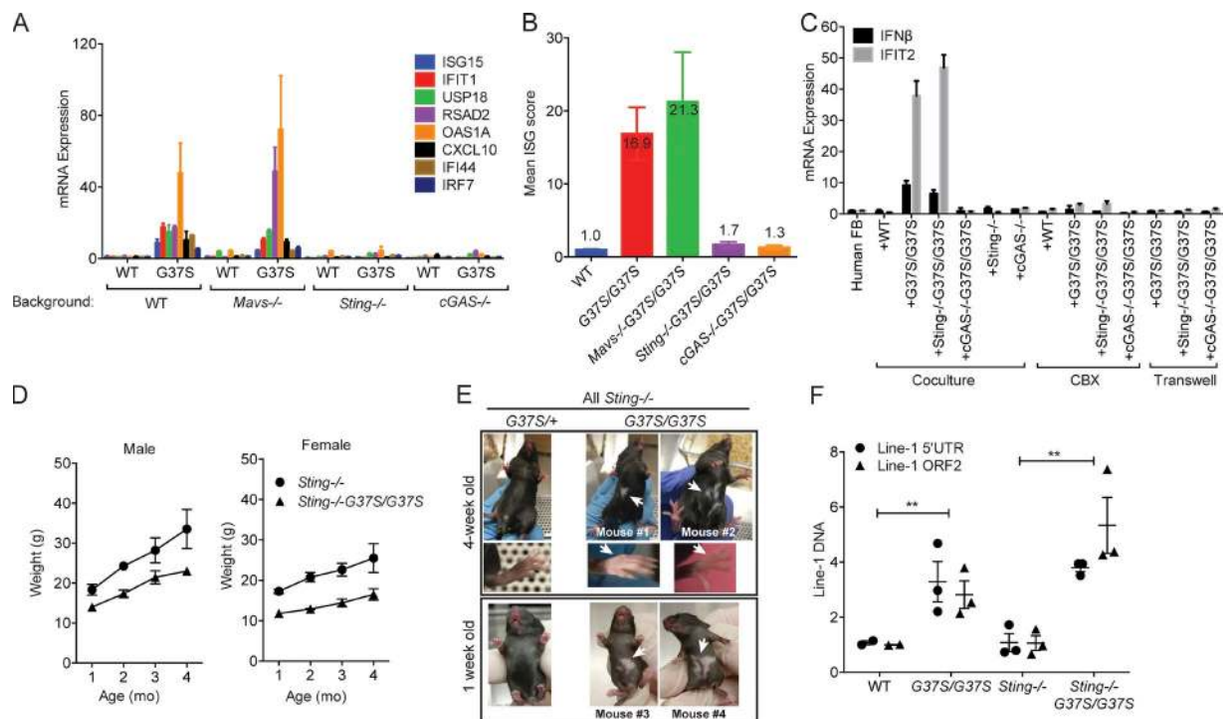


Figure 3. *Sting*^{-/-} partially rescues perinatal lethality of G37S mice. (A) Quantitative RT-PCR analysis of a panel of mouse ISGs in WT or G37S embryos on *Mavs*^{-/-} or *Sting*^{-/-} or *cGAS*^{-/-} background. Total RNA was isolated from primary MEFs (E13.5) of indicated genotype. (B) Mean ISG score of indicated genotypes. Data from A. (C) Gap junction cGAMP bioassay. As in Fig. 2 E, MEFs of indicated genotype were co-cultured with human BJ-1 cells for 18 h, with or without CBX or direct contact (indicated on the bottom). Quantitative RT-PCR analysis of IFN-β and IFIT1 indicates cGAMP activity in MEFs. (D) Mouse body weights. *n* = 4. (E) White-spotting phenotype in *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} viable adults. (F) Quantitative PCR analysis of mouse Line-1 5' UTR and ORF2 DNA in WT or G37S embryos (isolated from E13.5 or E15.5; *n* = 3). Each dot represents an individual embryo. Mice were compared with littermate controls and with age-matched knock-out mice **, *P* < 0.01. Data are representative of at least two independent experiments (A–C), or pooled data from multiple animals (D–F). Error bars represent the SEM. Unpaired Student's *t* test (F).

ground and found that *Rnaseh2a*^{G37S/G37S} *cGAS*^{-/-} embryos also restored ISG expression to WT levels (Fig. 3 A). Moreover, the mean ISG score is highly elevated in *Rnaseh2a*^{G37S/G37S} and *Rnaseh2a*^{G37S/G37S} *Mavs*^{-/-} MEFs, whereas both *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} and *Rnaseh2a*^{G37S/G37S} *cGAS*^{-/-} MEFs show similar baseline values as in WT (Fig. 3 B). We also measured cGAMP in both rescued MEFs using the gap junction bioassay. Both *Rnaseh2a*^{G37S/G37S} and *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} produce cGAMP, whereas *Rnaseh2a*^{G37S/G37S} *cGAS*^{-/-} MEFs did not (Fig. 3 C). These data further demonstrate that the cGAS–cGAMP–STING pathway is mediating the immune activation in G37S mice. Because only a small fraction of the progeny with *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} genotype is viable, our data also suggest that innate immune activation through the cGAS–cGAMP–STING pathway only partially contributed to the lethality of G37S mice.

White-spotting phenotype and increased LINE-1 expression

We also observed several interesting phenotypes in the viable *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} mice. These mice are ~70% in body size and weight compared with WT or heterozygous controls (Fig. 3 D). We have observed both male and female progeny

for *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-}, and all have so far failed to produce offspring, whereas littermate controls are fertile. These rescued *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} mice are grossly healthy, with the oldest animal approaching 1 yr of age. Intriguingly, all of the viable *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} mice presented a ventral white spotting phenotype, as well as white hind- and forepaws that are not observed in WT or heterozygous littermates (Fig. 3 E). This phenotype was consistent from birth and remained throughout the lifespan of the mice. Histopathology analysis did not find any abnormalities or inflammation in internal organs, including the brain, of *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} mice (not depicted). Skin histology from the white patches of *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} mice is structurally normal, although lack of melanin in hair shafts is evident (unpublished data). As endogenous retroelements have been implicated in the pathogenesis of AGS (Volkman and Stetson, 2014) and the RNA/DNA hybrids or DNA of murine endogenous retroviruses can function as a ligand of the cGAS–STING pathway (Mankan et al., 2014), we measured LINE-1 element in WT, *Rnaseh2a*^{G37S/G37S}, and *Rnaseh2a*^{G37S/G37S} *Sting*^{-/-} E13.5 and E15.5 embryos. Indeed, we found that LINE-1 DNA level from cytosolic extract is increased in both *Rnaseh2a*^{G37S/G37S}

and *Rnaseh2a*^{G37S/G37S}*Sting*^{-/-} embryos to similar levels compared with littermate WT embryos, suggesting that it is independent of immune activation (Fig. 3 F). However, we failed to observe elevation of L1 ORF1 protein by Western blot (not depicted). It remains unclear whether the increase in LINE-1 DNA is a result of an increase in LINE element activity, or because of defects in genomic structures where LINE elements are enriched as was recently suggested (Lim et al., 2015). Together, our data suggest that the G37S mutation causes white-spotting phenotype in *Rnaseh2a*^{G37S/G37S}*Sting*^{-/-} mice, likely resulting from defects in melanocyte development or migration from progenitors at the neural crest. G37S mutation also causes increased level of LINE-1 DNA, which may contribute to the activation of the cGAS–STING pathway.

In summary, the G37S mouse represents the first RNase H2 mouse model with a clear immune activation phenotype, making it uniquely useful for understanding the associated human disease. Embryonic development until birth allowed the expression of innate immune signaling proteins or immune ligands, which uncovered the active cGAS–STING innate immune pathway in G37S mice. RNA/DNA hybrids or rNMPs in DNA could be a direct source of nucleic acids activating the DNA-sensing pathway. Alternatively, specific nucleic acids, such as LINE-1–derived nucleic acids, could elicit the innate immune response. The perinatal lethality of the G37S mice is likely caused in large part by a yet-to-be-identified biochemical defect associated with the mutation. *Sting*^{-/-} only partially rescued the lethality, despite complete suppression of ISG expression, and *p53*^{-/-} failed to rescue the lethality. Similarly, another AGS mouse model, the *Adar1*^{-/-} mouse, exhibits early embryonic lethality, and it can be partially rescued to birth by *Mavs*^{-/-} and to adult by *Ifih1*^{-/-} (Mannion et al., 2014; Liddicoat et al., 2015). These findings indicate that *Rnaseh2* and *Adar1* genes associated with AGS have important functions that are critical for embryonic development in mice, beyond that of prohibiting formation of aberrant nucleic acids that activate innate immunity (Pestal et al., 2015). Further biochemical analysis comparing G37S and other existing *Rnaseh2* knockout mouse models are necessary to elucidate the differences in biochemical defects and to shed light on the possible source of nucleic acids that trigger the cGAS–STING pathway in the G37S mice. Recently studies also showed that *cGAS*^{-/-} can rescue the inflammation and mortality of *Trex1*^{-/-} mice (another AGS mouse model; Gao et al., 2015; Gray et al., 2015). Therefore, our genetic and immunological analysis of the G37S mice further establish a critical role of the cGAS–STING pathway in G37S-induced immune activation, and suggest that therapeutic intervention of this pathway may be beneficial for treating AGS patients.

MATERIALS AND METHODS

Mice, cells, and viruses. G37S mice were generated by introducing the human disease associated point mutation into the conserved residue of mouse *Rnaseh2a* gene. *Infar1*^{-/-} and *Rag2*^{-/-} mice were obtained from Taconic Biosciences.

p53^{-/-} mice were obtained from The Jackson Laboratory. *Mavs*^{-/-} and *cGAS*^{-/-} mice were obtained from Z. Chen (University of Texas Southwestern Medical Center, Dallas, TX) and *Sting*^{-/-} mice were obtained from G. Barber (University of Miami, Miami, FL). Primary MEFs were isolated from embryos of indicated embryonic dates. These cells were maintained in DMEM with 20% (vol/vol) heat-inactivated FCS, 2 mM L-glutamine, 10 mM HEPES, and 1 mM sodium pyruvate (complete DMEM) with the addition of 100 U/ml penicillin and 100 mg/ml streptomycin and were cultured at 37°C with 5% CO₂. VSV-PeGFP is a gift from A. Pattnaik (University of Nebraska, Lincoln, NE; Das et al., 2014). Cells were plated overnight and, the next day, infected overnight with VSV-GFP at a multiplicity of infection of 1. Cells were washed with PBS before standard fixation with 4% paraformaldehyde in PBS (Affymetrix). Percentage of infectivity was assessed with FACSCalibur (BD). For viral RNA measurement, total RNA was extracted at various time points after infection, and VSV G and M RNA were measured with specific primers (Hasan et al., 2013). Experiments performed in BSL2 conditions were approved by the Environmental Health and Safety Committee at University of Texas Southwestern Medical Center. Experiments involving mouse materials were approved by the Institutional Animal Care and Use Committees of the University of Texas Southwestern Medical Center and the National Institute for Child Health and Human Development (Bethesda, MD).

RNA isolation and quantitative RT-PCR. Total RNA was isolated with TRI reagent according to the manufacturer's protocol (Sigma-Aldrich), and cDNA was synthesized with iScript cDNA synthesis kit (Bio-Rad Laboratories). iTaq Universal SYBR Green Supermix (Bio-Rad Laboratories) and an ABI-7500 Fast Real-Time PCR system (Applied Biosystems) were used for quantitative RT-PCR analysis (primer sequences listed in Table S2). *Hprt* and *Gapdh* were used as housekeeping genes for data normalization. RNA-seq was performed as previously described (Hasan et al., 2013).

cGAMP activity bioassay. cGAMP activity in MEFs was measured by a co-culture bioassay as previously described (Ablasser et al., 2014). In brief, 2×10^5 human fibroblasts/ml were plated overnight. After attachment, 4×10^5 primary murine fibroblasts/ml were plated onto the human cells, with or without 200 μM CBX (Sigma-Aldrich) treatment or with separation by 0.4-μm polycarbonate Trans-well inserts (Corning) for 18 h. Subsequent analysis was performed using a human-specific PrimePCR Array plate (Bio-Rad Laboratories).

shRNA knockdown and TBK inhibitor. shRNA oligos were synthesized (Sigma-Aldrich) and cloned into a pLKO.1-TRC cloning vector following Addgene's protocol. Lentiviral particles were packaged in HEK-293 T cells and filtered with Amicon Ultra-15 centrifugal filters (EMD Millipore). shRNA-harboring lentiviruses were allowed to

infect cells overnight with polybrene (10 µg/ml) treatment, and cells expressing the shRNA were selected with a puromycin concentration (Life Technologies) for several days. Puromycin selection was removed several days before subsequent analysis. shRNA oligo sequences are listed in Table S1. For TBK1 inhibitor experiments, cells were treated with DMSO (% vol/vol) or 10 µM BX795 (InvivoGen) for 6 h. Cells were then washed with PBS and grown in DMEM with 20% FCS alone for several hours before subsequent analysis.

Line-1 retroelements. E13.5 or E15.5 embryos were homogenized into single-cell suspensions and cytosolic lysates obtained using a cytosolic extraction buffer as previously described (Yang et al., 2007). In brief, PBS-washed cell pellets were lysed in 10 mM Hepes, pH 7.9, 10 mM KCl, 1.5 mM MgCl₂, 0.34 M sucrose, 10% glycerol, and 0.1% Triton X-100 for 5 min on ice. Cell lysates were treated with Proteinase K at 55°C for 1 h and with RNase I (Life Technologies) before phenol/chloroform extraction and ethanol precipitation overnight. Isolated DNA was resuspended in nuclease free H₂O and directly subjected to quantitative PCR analysis using Line-1 primers (Table S1).

Statistical methods. Data are presented as the mean ± SEM. Prism 6 (GraphPad) was used for statistical analysis. Statistical tests performed are indicated in figure legends. *, P < 0.05; **, P < 0.01; ***, P < 0.001; and ****, P < 0.0001.

Online supplemental material. Table S1 lists RNA-seq data. Table S2 lists oligonucleotides used in this study. Online supplemental material is available at <http://www.jem.org/cgi/content/full/jem.20151464/DC1>.

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Author contributions: V. Pokatayev performed immunology related experiments, including crosses between G37S mice and *Mavs*^{-/-} or *Sting*^{-/-} or *cGAS*^{-/-} mice. S.M. Ceritelli was responsible for production of *Rnaseh2a*^{G37S} ES cells. N. Hasin, H. Chon, and K. Sakhuja generated and characterized the initial phenotypes of the G37S mouse, including crosses between G37S mice and *Infar*^{-/-}, *Rag2*^{-/-}, or *P53*^{-/-} mice. J.M. Ward performed pathology observations, H. Douglas Morris performed MRI, CT and analyzed the images. N. Yan and R.J. Crouch jointly supervised the study. N. Yan, R.J. Crouch, V. Pokatayev, N. Hasin, and S.M. Ceritelli wrote the paper.

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