

Severe viral respiratory infections in children with *IFIH1* loss-of-function mutations

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Viral respiratory infections are usually mild and self-limiting; still they exceptionally result in life-threatening infections in previously healthy children. To investigate a potential genetic cause, we recruited 120 previously healthy children requiring support in intensive care because of a severe illness caused by a respiratory virus. Using exome and transcriptome sequencing, we identified and characterized three rare loss-of-function variants in *IFIH1*, which encodes an RIG-I-like receptor involved in the sensing of viral RNA. Functional testing of the variants *IFIH1* alleles demonstrated that the resulting proteins are unable to induce IFN- β , are intrinsically less stable than wild-type IFIH1, and lack ATPase activity. In vitro assays showed that IFIH1 effectively restricts replication of human respiratory syncytial virus and rhinoviruses. We conclude that *IFIH1* deficiency causes a primary immunodeficiency manifested in extreme susceptibility to common respiratory RNA viruses.

respiratory syncytial virus | rhinovirus | *IFIH1* | RIG-I-like receptor family | severe pediatric infectious disease

Viral respiratory tract infections are the most common childhood infections worldwide, with close to 100% of children being infected during the first years of life. Whereas the vast majority of viral respiratory infections are mild and self-limiting, more severe disease leads to the hospitalization of about 3% of individuals in each birth cohort (1). In-hospital mortality rates are limited to <1% with intensive care support; still these infections account for 21% of childhood mortality worldwide (2, 3). The main viral pathogens causing lower respiratory tract infections are human respiratory syncytial virus (RSV), enteroviruses [including human rhinoviruses (HRV)], adenoviruses, human metapneumovirus, coronavirus, influenza, and parainfluenza viruses, with RSV being responsible for the majority of the hospitalized pediatric cases (4, 5).

A number of risk factors including socioeconomic and environmental influences, preterm birth, chronic diseases, and immunosuppression are associated with more severe clinical presentation (6). However, \sim 1 out of 1,000 children without any known risk factor will require intensive care support due to life-threatening manifestations of common viral respiratory infections. In the absence of established differences in pathogen virulence, we hypothesized that human genetic variation contributes to unusual susceptibility to severe disease due to common viruses. Supporting evidence is provided by a recent study, which showed that rare variants in *IRF7* resulted in life-threatening influenza in an otherwise healthy child (7).

We combined exome sequencing, transcriptomic analysis, and in vitro functional testing to identify and characterize potentially causal genetic variants in a prospective cohort of previously healthy children requiring intensive care support for common respiratory viral infections. We report the identification of a pathogen-restricted immunodeficiency due to loss-of-function variants in *IFIH1*, which result in defective innate recognition of RNA viruses, preventing the activation of an efficient antiviral IFN response.

Results

Study Participants. We enrolled 120 previously healthy children admitted to pediatric intensive care units (PICUs) with respiratory failure due to a common viral respiratory infection. The most common clinical presentation was bronchiolitis (n = 105, 88%) and the median age was 78 d (interquartile range, IQR: 37–769). RSV was the most common pathogen, identified in 67 (56%) of the cases, followed by HRV in 31 (26%) of the cases (Table 1).

Exome Sequencing and Analysis. DNA samples were sequenced to a mean coverage of 70×, with 96% of exonic bases achieving at least 10× coverage and 78% achieving at least 30× coverage. The final set of variants included 2,793 stop-gained single-nucleotide variants (SNVs), 297 splice-site SNVs, and 951 frame-shift indels. Among these putative loss-of-function variants (LoFs), we searched for variants that were homozygous in at least one study participant, and with a higher minor allele frequency in our cohort than in the genome Aggregation Database (gnomAD) (8) and in an in-house

Significance

Life-threatening susceptibility to common respiratory infections in previously healthy children can be indicative of pathogenspecific primary immunodeficiencies due to rare deleterious variants in key genes and pathways of the immune system. These findings have implications for prevention and treatment of susceptible children.

Author contributions: A.T., D.G., and J.F. designed research; S. Asgari, L.J.S., S. Anchisi, C.H., G.M.-O., K.M.P.-B., D. Longchamp, M.S., S.C., L.K., and T.R. performed research; L.J.S., K.M.P.-B., D. Longchamp, M.S., T.K., D. Long, and A.S. contributed new reagents/analytic tools; S. Asgari, S. Anchisi, C.H., I.B., T.J., G.M.-O., S.C., C.T., and P.J.M. analyzed data; and S. Asgari, L.J.S., S. Anchisi, and J.F. wrote the paper.

The authors declare no conflict of interest

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Table 1. Baseline characteristics of the 120 study participant	Table 1.	Baseline	characteristics	of the	120 study	participants
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Parameter	Variable	Median (IQR) or N (%)
Age, d		78 (37–269)
Weight, kg		5.9 (4.4–9.8)
Country of recruitment	Australia	100 (83%)
	Switzerland	20 (17%)
Ethnicity	Caucasian	90 (78%)
	African	4 (4%)
	Asian	4 (4%)
	Australian aboriginal	6 (5%)
	Pacific Islander	11 (10%)
Sex	Male	70 (58%)
	Female	50 (42%)
Clinical phenotype	Bronchiolitis	105 (88%)
	Pneumonia	8 (7%)
	Laryngotracheobronchitis	5 (4%)
	Reactive airway disease	2 (2%)
Virus identified in	RSV	67 (56%)
respiratory sample	Entero/rhinovirus	31 (26%)
	Adenovirus	17 (14%)
	Human bocavirus	9 (8%)
	Influenza	2 (2%)
	Parainfluenza	6 (5%)
	Human metapneumovirus	3 (2%)
	CoV-HKU1	3 (2%)
	CoV-NL63	1 (1%)
Respiratory support	HFNC	99 (83%)
	CPAP or BiPAP	30 (25%)
	Invasive ventilation	31 (26%)
	HFOV	6 (5%)
Length of PICU stay, d		2.7 (1.6–5.0)
Expected mortality	Pediatric index of mortality 2 (%)	0.02 (0.01–0.7)
Observed mortality	Pediatric index of mortality 2 (%)	No fatal case

CoV-HKU1, coronavirus HkU1; CoV-NL63, coronavirus NL63.

collection of 485 exomes. Six putative LoFs fitting these criteria were identified (*SI Appendix*, Table S3). This set included a variant in the IFN induced with helicase C domain 1 (*IFIH1*) gene, which encodes an RIG-I-like cytoplasmic sensor of long double-stranded RNA (dsRNA) and plays a major role in innate immune recognition of RNA viruses (9).

LOF Variants in IFIH1. In total, eight study participants carried putative LoF variants in *IFIH1* (Table 2). Four study participants carried a rare [gnomAD minor allele frequency (MAF) = 0.64%]

splicing variant, rs35732034 (Fig. 1A): one in homozygous and three in heterozygous form. We used RNA sequencing to characterize the transcriptomic impact of this variant. We observed that the minor allele T causes skipping of exon 14 (IFIH1- Δ 14) (Fig. 1B), which results in a frame shift and an early stop codon in exon 15. The resulting protein lacks the final 153 amino acids of wild-type IFIH1, including the C-terminal regulatory domain (CTD), which is essential for binding to viral dsRNA (10) (Fig. 1C). Western blot analyses of peripheral blood mononuclear cells from the homozygous patient and her heterozygous parents demonstrated that the IFIH1- Δ 14 protein is expressed upon in vitro RSV infection (SI Appendix, Fig. S1). We identified two additional rare LoF variants in IFIH1, present in heterozygous form in a total of four study participants (Fig. 1A): the splicing variant rs35337543 (n = 3, gnomAD MAF = 0.67%) and the stopgained variant rs35744605 (n = 1, gnomAD MAF = 0.32%). RNA sequencing showed that the minor allele G at rs35337543 causes skipping of exon 8 (IFIH1- Δ 8) (Fig. 1*B*), which removes 39 amino acids at the end of the helicase 1 domain and in the linker part between helicase 1 and helicase 2, but does not result in a frame shift. rs35744605 is a stop-gained SNV in exon 10 that leads to the loss of 399 amino acids from the C-terminal end of IFIH1 (IFIH1- Δ CTD) (Fig. 1*C*).

Description of Study Participants Carrying IFIH1 LoF Variants. A 16-moold girl was homozygous for rs35732034. She presented with respiratory failure due to RSV infection requiring invasive ventilation. The disease course was complicated by a pulmonary superinfection with Staphylococcus aureus. She had a full recovery and did not develop any other severe infection up to the age of 3. Her phenotype and history was otherwise unremarkable. In particular, she did not develop any complication after live vaccine administration. Full blood count, Ig levels and IgG subclasses, and lymphocyte subclasses were within normal limits. Three infants requiring noninvasive respiratory support for bronchiolitis were heterozygous for rs35732034. One of these had recurrent severe viral lower respiratory tract infections leading to repeated PICU admissions during childhood. Three children were heterozygous for rs35337543 and required noninvasive ventilatory support for RSV bronchiolitis. One infant was heterozygous for rs35744605 and required invasive ventilatory support for HRV-positive bronchiolitis. Parental DNA was available for three of the eight children. Targeted genotyping confirmed that the relevant IFIH1 variant was present in heterozygous form in one of the parents for two heterozygous individuals, and in both parents for the homozygous patient. Parental medical history was unremarkable in all cases.

Functional Characterization of IFIH1 Variants. To functionally characterize the identified variants, we first measured the ability of wild-type (IFIH1-wt) and mutant IFIH1 isoforms to induce IFN β (IFN β) in vitro. We transfected plasmids carrying IFIH1-wt,

Table 2.	Characteristics of the	eight study particip	ants carrying an IFI	11 loss-of-function variant

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Patient ID	Zygosity	Sex	Age at admission, d	Ethnicity	Virus	Ventilation required	Clinical presentation	Parental allele	Variant ID	Nucleotide change	Amino acid change	gnomAD AC (#hom), MAF
PRI_022	hom	F	493	White	RSV	I	Bronchiolitis pneumonia	het in both parents	rs35732034	2:163124596 C/T	p.lle872Ter	1673 (7), 0.006
PRI_050	het	М	34	Aboriginal	HRV	NI	Recurrent bronchiolitis	NA				
PRI_061	het	М	41	White	RSV	NI	Bronchiolitis	het in father				
PRI_122	het	F	48	White	RSV	NI	Bronchiolitis	het in mother				
PRI-063	het	М	22	White	RSV	NI	Bronchiolitis	NA	rs35337543	2:163136505 C/G	p.Leu509_Glu547del	1847 (8), 0.006
PRI-065	het	М	38	White	RSV	NI	Bronchiolitis	NA				
PRI-116	het	М	479	White	RSV	NI	Bronchiolitis	NA				
PRI-080	het	Μ	343	White	HRV	I	Bronchiolitis pneumonia	NA	rs35744605	2:163134090 C/A	p.Glu627Ter	887 (1), 0.003

NA, not available; het, heterozygous; hom, homozygous; I, invasive; NI, noninvasive; AC, allele count.

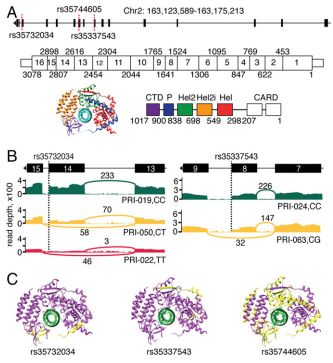


Fig. 1. LoF variants identified in *IFIH1*. Related to *SI Appendix*, Fig. S1. (A) Schematic representation of IFIH1 DNA, mRNA, and protein. The identified variants are indicated with dashed red lines. Exon boundaries are marked with nucleotide coordinates. Protein domain boundaries are marked with amino acid coordinates. (*B*) Alternative splicing of IFIH1 associated with rs35732034 and rs35337543 genotypes, as seen in RNA sequencing data. The T allele at rs35732034 leads to skipping of exon 14. The G allele at rs35337543 leads to skipping of exon 14. The G allele at rs35337543 leads to skipping of exon 8. The Sashimi plots illustrate the genotype-dependent abundance of splice junctions. The number of observed reads spanning the respective splice junctions is indicated on the Bezier curves, which connect exons. (C) Schematic 3D representation of IFIH1 (Protein Data Bank ID code: 4GL2, image produced using UCSC Chimera). The parts of the protein that are predicted to be missing due to rs35732034, rs35744605 and rs35337543 variants are indicated in yellow. Hel, helicase domain; P, pincer.

IFIH1- Δ 8, IFIH1- Δ 14, and IFIH1- Δ CTD into 293T cells. Overexpression of IFIH1-wt, but not of any of the mutant IFIH1 isoforms, led to IFNβ induction. Cotransfection of IFIH1-wt with each of the mutant IFIH1 isoforms showed interference with IFIH1-wtinduced IFN β production (P < 0.05, Fig. 2A). We then tested the ATPase activity of recombinant IFIH1-wt and mutant IFIH1 isoforms. IFIH1-wt was able to hydrolyze ATP and showed a typical dsRNA-dependent increase in enzymatic activity, whereas the mutant isoforms had no detectable ATPase activity, even upon stimulation with polyinosinic:polycytidylic acid (polyI:C), a synthetic analog of dsRNA (Fig. 2B). Furthermore, all mutant isoforms decreased IFIH1-wt ATPase activity in a dose-dependent manner, an interference that was specific to mutant IFIH1 isoforms, as demonstrated by the absence of any effect of BSA on IFIH1-wt ATPase activity (P < 0.05, Fig. 2C and SI Appendix, Fig. S2 A and B). Finally, we checked the stability of the various IFIH1 protein isoforms by performing pulse-chase experiments in transfected 293T cells. The three mutant isoforms were less stable than IFIH1-wt (Fig. 2D) and had a negative impact on the stability of the wild-type isoform when cotransfected (Fig. 2E). Jointly, these experiments demonstrate that the three putative LoF variants identified in our study population lead to severe disruption of IFIH1 signaling function, enzymatic activity, and protein stability in vitro. In addition, the observations that the mutant IFIH1 isoforms interfere with the wild-type protein in terms of IFN β induction, and enzymatic activity and protein stability suggest a dominant negative role for heterozygous LoF variants in *IFIH1*.

Role of IFIH1 in RSV and HRV Replication. Viral testing of respiratory samples showed that six of the patients harboring *IFIH1* LoF alleles were infected with RSV and two with HRV. To study the effect of IFIH1 on RSV and HRV replication, we used Huh7.5 cells, which lack endogenous expression of IFIH1 and express a mutated, inactive form of RIG-I, and thus are completely unreactive to the RNA pathogen-associated molecular patterns that normally activate these pathways (11). The cells were transduced with an IFIH1-wt-expressing lentiviral vector, which made them highly responsive to polyI:C stimulation (*SI Appendix*, Fig. S3*A*) without causing any nonspecific or constitutive activation of the IFN system (*SI Appendix*, Fig. S3*B*). We observed a much higher level of viral replication in native than in IFIH1-wt-transduced Huh7.5 cells upon infection with HRV-B14, HRV-

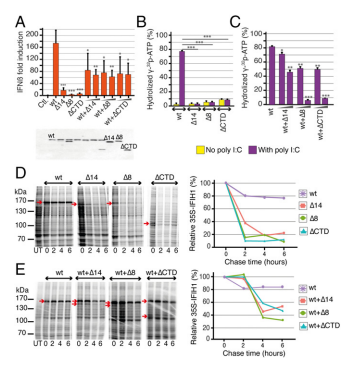


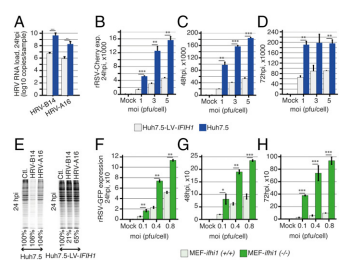
Fig. 2. Functional characterization of the IFIH1 variants. Related to SI Appendix, Fig. S2. (A) Transfection with IFIH1-wt plasmid (20 ng) results in strong IFN_B induction in 293T cells, whereas transfection of plasmids harboring any of the IFIH1 variants cannot induce IFN β (240 ng, n = 4). Cotransfection experiments demonstrate an interference of the mutant plasmids with IFIH1-wt on IFN β induction (20-ng wt plasmid, 20- and 120-ng mutant plasmids, n = 4). Expression levels of the IFIH1 isoforms are shown under the plot in the Western blot gel. *P < 0.05, **P < 0.01, ***P < 0.001. (B) RNA-induced ATPase activity of purified IFIH1-wt protein and alternate IFIH1 isoforms; IFIH1-wt can hydrolyze ATP in the presence of polyl;C, whereas IFIH1- Δ 14, IFIH1- Δ 8, and IFIH1-ACTD lack ATPase activity, with (purple) or without (yellow) polyI:C stimulation. (C) ATPase activity of IFIH1-wt is reduced upon coincubation with the alternate isoforms in a dose-dependent manner (300-ng wt protein, 300- or 600-ng of each alternate isoform, 10-ng polyl:C, n = 2). *P < 0.05, **P < 0.01, ***P < 0.001. (D and E) Protein stability followed by pulse chase in 293T cells expressing IFIH1-wt, IFIH1-∆14, IFIH1-∆8, or IFIH1-∆CTD. Each protein is marked on the gel by an arrow, and relative amounts of proteins are shown in the graphs. IFIH1-wt is more stable than the alternate IFIH1 isoforms, and the stability of IFIH1-wt is reduced upon coexpression with any of the alternate isoforms. Molecular mass markers are shown on the left of each gel (kilodaltons) and the bands corresponding to IFIH1-wt, IFIH1- Δ 14, IFIH1- Δ 8, or IFIH1- Δ CTD are indicated using a red arrow (2- μ g plasmid expressing wt protein, 2 μ g of each mutant plasmid, n = 1). Data are represented as mean ± SD; polyI:C, polyinosinic:polycytidylic acid.

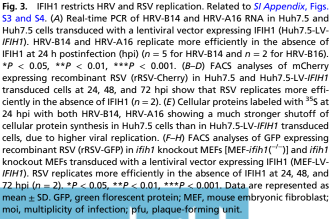
A16, and RSV (Fig. 3 *A–D*). Furthermore, RSV replication level was higher in cells transduced with the mutant forms of IFIH1 than in IFIH1-wt-transduced cells (P < 0.05, *SI Appendix*, Fig. S4 *A* and *B*). The role of IFIH1 in HRV restriction was further demonstrated by ³⁵S labeling of infected cells, which showed a stronger shutoff of cellular protein synthesis in native than in IFIH1-transduced Huh7.5 cells, due to higher replication of the virus in the absence of IFIH1 (Fig. 3*E*). We also measured RSV replication in mouse embryonic fibroblasts (MEFs), *ifih1*(^{+/+}), and in IFIH1-knockout MEFs, *ifih1*(^{-/-}), and obtained similar results (Fig. 3 *F–H*). Together, these results affirm the central role of IFIH1 in innate immune recognition of RSV and HRV (12, 13). Therefore, LoF variants in *IFIH1* can be reasonably expected to increase susceptibility to these viruses.

Discussion

We hypothesized that extreme susceptibility to common viral respiratory infection in previously healthy children—a rare, potentially lethal phenotype—could reflect an underlying primary immunodeficiency. Using an unbiased exomewide approach in a prospective cohort of carefully selected individuals requiring intensive care support, we identified a rare monogenic defect predisposing to severe clinical presentations of RSV and HRV infections.

Three deleterious variants were observed in *IFIH1*, which encodes a cytoplasmic receptor critical for viral RNA sensing. It has been shown previously that IFIH1, alone or in combination with RIG-I, recognizes and limits the replication of many RNA viruses including: positive single-stranded RNA (ssRNA) viruses





like picornaviruses (14–16), negative ssRNA viruses like paramyxoviruses (17–19), and dsRNA viruses like reoviruses (20).

IFIH1 recognizes viral RNA via interaction of its CTD and helicase domains with long dsRNA molecules. This is an ATPdependent reaction that leads to polymerization of IFIH1 molecules into a filament and assembly of IFIH1 caspase activation recruitment domains (CARDs) (21, 22). This in turn initiates a signaling cascade that results in type 1 IFN production and activation of antiviral genes (23). Our transfection and transduction analyses show that this process is disrupted in the presence of any of the *IFIH1* rare variants found in our study population. Our exome and RNA sequencing data predict that the loss of IFIH1 function is due to loss of the CTD (rs35732034 and rs35744605) or to partial loss of the helicase domain (rs35337543).

We observed interference between IFIH1-wt and the three mutant proteins in terms of stability, ATPase activity, and capacity to induce IFN β production, suggesting a dominant negative effect, which provides a rationale for the unusual susceptibility to respiratory viruses observed in heterozygous individuals. The exact interfering mechanism is not known but could involve physical interaction between IFIH1-wt and mutant proteins, preventing the formation of normal, multimeric IFIH1 filaments.

While the revised version of this paper was under review, an independent study showed an association between IFIH1 deficiency and life-threatening infections with HRV and other respiratory viruses in a child carrying another homozygous missense *IFIH1* variant (24). This observation further supports a causal role for IFIH1 deficiency in extreme susceptibility to common respiratory viruses.

The three IFIH1 variants described in this study have allele frequencies of 0.32-0.67% in gnomAD. The cumulative frequency of all putative LoF alleles is 1.89% in the same database, which is significantly less than the 3.75% cumulative frequency observed in our study population (P = 0.037, Fisher's exact test). Nevertheless, the presence of alleles of potentially devastating consequences at such frequency in the general population is intriguing, as they are expected to be removed by purifying selection. Two nonexclusive mechanisms can explain this observation: balancing selection and incomplete penetrance. We here show that some IFIH1 alleles increase susceptibility to viral respiratory infections, but the same LoF variants are known to be protective against type 1 diabetes and other autoimmune diseases (25–30), strongly suggesting a role for balancing selection in their maintenance. In a comparable example, rare nonsynonymous variants in TYK2, a known primary immunodeficiency gene, were shown to be protective against rheumatoid arthritis (31). Incomplete penetrance, on the other hand, could be due to modulating effects of environmental or genetic factors, like compensatory mutations, or to functional redundancy in innate immune response to RNA viruses (32, 33). This hypothesis is in line with several recent publications (34-37), which suggest that incomplete penetrance and genetic heterogeneity are likely to be the rule rather than the exception in severe clinical presentations of infectious diseases.

On top of their associations with autoimmunity, more common *IFIH1* variants have also been associated with hepatitis C virus clearance (38). Additionally, rare gain-of-function mutations in *IFIH1* dramatically up-regulate type I IFN production, resulting in Aicardi–Goutières syndrome or Singleton–Merten syndrome (39–41). At the functional level, Gorman et al. recently studied the effects on viral sensing and autoimmune pathogenesis of rs1990760, a missense *IFIH1* variant that is associated with multiple autoimmune diseases (30). They showed that the allele providing better defense against viral infection also bolsters autoimmune responses against self-RNA (42). Together, these results underscore the pivotal role of innate immune recognition and activation in the intricate balance between host defense, inflammation, and autoimmunity.

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Our study demonstrates the power of using an unbiased, exome sequencing approach to variant discovery in prospective cohorts of extreme infectious disease phenotypes. Nevertheless, LoF variants in *IFIH1* were only found in a minority (n = 8, 6.2%) of the 120 children enrolled in our study, suggesting that other genetic or nongenetic risk factors remain to be discovered. Larger sample sizes will be required to delineate the relevance of other rare potentially causal alleles. Whole genome sequencing will also be needed to obtain a more complete coverage of exonic regions (43), and to explore noncoding and large-scale structural variation.

RSV and rhinovirus infections are the two most common viral respiratory infections in children. The elucidation of the human genetic basis of extreme susceptibility to these viruses provides insight into pathogenesis and innate immune response. An immediate practical implication is the possibility to develop diagnostic assays to identify susceptible individuals who could benefit from specific preventive and interventional measures. By highlighting the genes and pathways that play an essential role in host–pathogen interaction, genetic discovery in individuals with extreme phenotypes also provides the opportunity to design new therapeutic strategies that could be useful for the vast majority of patients with milder clinical presentation.

Materials and Methods

Between December 2010 and October 2013, we prospectively enrolled previously healthy children below 4 y of age suffering from severe lower respiratory tract infection and requiring invasive or noninvasive respiratory support in five specialized PICUs from Australia and Switzerland. The study was approved by the respective institutional Human Research Ethics Committees. Written informed consent was obtained from parents or legal guardians.

Exclusion criteria were the presence of any significant underlying disease or comorbidity, including prematurity, congenital cardiac disease, chronic lung disease, sickle cell disease, hepatic, renal, or neurologic chronic

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conditions, solid and hematological malignancies, and known primary immunodeficiency. Respiratory support was defined as noninvasive ventilation including high-flow nasal cannulae (HFNC) and continuous or bilevel positive airway pressure (CPAP and BiPAP), or invasive ventilation including conventional and high-frequency oscillation ventilation (HFOV). The following demographic and clinical information was collected: age, gender, weight, ethnicity, type of ventilation, length of ventilation in days, clinical outcome, microbiological diagnostic procedures and results including rapid antigenic test for RSV and influenza, respiratory virus PCR panel, and viral cultures. For each study participant, we obtained a nasopharyngeal aspirate or endotracheal tube aspirate, 1 mL EDTA blood in vacutainer tubes, and 2.5 mL blood in PAXgene blood RNA tubes. Samples were immediately frozen at -70 °C until shipment, and then analyzed in batch.

We generated high-coverage exome sequencing data for all study participants (*SI Appendix*, Table S1). We then used a combination of three variant calling methods (GATK, Platypus, and SAMtools) and only kept SNVs and small insertion and deletions (*SI Appendix*, Table S2). Assuming that causal genetic variants are likely to be highly deleterious, we focused on rare gene knockout events, defined as homozygous, putative LoF variants (stopgained and splice-site SNVs, frame-shift indels) with an MAF of <1% in the ExAC (*SI Appendix*, Table S3). We performed RNA sequencing to assess the transcriptomic impact of candidate DNA variants, and characterized the infecting viruses using multiplex PCR assays. We then used in vitro functional testing to demonstrate the biological relevance of candidate variants and tested in vivo expression of the affected protein (*SI Appendix*).

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