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Study of Human RIG-I Polymorphisms Identifies Two Variants with an Opposite Impact on the Antiviral Immune Response

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Abstract

Background: RIG-I is a pivotal receptor that detects numerous RNA and DNA viruses. Thus, its defectiveness may strongly impair the host antiviral immunity. Remarkably, very little information is available on RIG-I single-nucleotide polymorphisms (SNPs) presenting a functional impact on the host response.

Methodology/Principal Findings: Here, we studied all non-synonymous SNPs of RIG-I using biochemical and structural modeling approaches. We identified two important variants: (i) a frameshift mutation (P229fs) that generates a truncated, constitutively active receptor and (ii) a serine to isoleucine mutation (S183I), which drastically inhibits antiviral signaling and exerts a down-regulatory effect, due to unintended stable complexes of RIG-I with itself and with MAVS, a key downstream adapter protein.

Conclusions/Significance: Hence, this study characterized P229fs and S183I SNPs as major functional RIG-I variants and potential genetic determinants of viral susceptibility. This work also demonstrated that serine 183 is a residue that critically regulates RIG-I-induced antiviral signaling.

Introduction

Among all viral components that trigger the antiviral screen of the host, nucleic acids have been viewed as the most important [1]. In mammals, there are at least two receptor systems in place to detect such viral motifs and to further mount a type I interferon (IFN)-dependent antiviral immune response. The endosomal TLR3, 7, 8, 9 and 13 interact with extracellular viral nucleic acids while the cytosolic helicases RIG-I and MDA-5 sense intracellular double-stranded (ds)RNA and/or 5’ triphosphate single-stranded RNA, two common byproducts of viral infection and replication [2,3,4].

Current knowledge posits RIG-I as a particularly critical surveillance molecule that detects numerous viruses such as the human pathogens influenza and hepatitis C (HCV) viruses [6,7]. RIG-I interacts with its ligands by means of its central ATP-binding helicase domain as well as its carboxyterminal regulatory domain (RD; see the schematic representation in Fig. 1A). For its amino-terminal tandem Caspase Recruitment Domains (CARDs), RIG-I homocomplexes relay a signal by binding MAVS (also known as IPS-1, CARDIF or VISA), an adapter protein that mediates CARD-dependent interactions with RIG-I. This signaling complex further activates the transcription factors NF-kB and interferon regulatory factor (IRF)-3 to ultimately upregulate the expression of pro-inflammatory and antiviral mediators and the subsequent induction of adaptive immune responses [2,3,4].

Importantly, the receptor function of RIG-I is non-redundant, as confirmed by knock-out studies [8]. Moreover, the Huh7.5 hepatocytic cell line is especially permissive to HCV as the result of an elaborate viral evasion strategy as well as to a defective RIG-I protein bearing a single mutation [9,10]. In that regard, unequivocal evidence shows that genetic mutations may be important determinants of increased susceptibility to viral diseases [11,12]. Among them, single-nucleotide polymorphisms (SNPs) are DNA sequence variations that occur when a single nucleotide is altered. There are more than 4 million SNPs in the human genome, 200,000 of which occur in coding regions, underlying the extent of genetic variability...
and its potential positive or negative effects on the host antimicrobial defense [13,14]. Interestingly, studies aiming to characterize RIG-I polymorphisms are scarce. Here, we characterized functional effects of two RIG-I SNPs that might help us to understand the basis of individual variations between normal and abnormal innate immune responses to viral pathogens as well as to better appreciate the molecular mechanism by which RIG-I is triggered by non-self RNA.

**Results**

**Genetic variability profile of human RIG-I**

Information collected on 04/2009 from NCBI SNP database indicates that at least 342 SNPs are present in the human RIG-I gene. Among them, 14 are situated within coding sequences but only 7 result in amino acids substitutions, i.e. R7C, S144F, S183I, T260P, I406T, D580E, F789L (Fig. 1A). An additional SNP corresponds to a thymidine insertion at nucleotide position 845 of RIG-I mRNA (accession number NM_014314), which results in a frameshift (fs) and in a truncated RIG-I protein. This mutant is herein defined as P229fs as it includes the first 229 residues (instead of 925 residues in the WT RIG-I protein) followed by 4 unintended residues [i.e. FRSV; Fig. 1B] and thus, does not contain the helicase and the RD domains. As illustrated in Fig. 1A, RIG-I SNPs map to the different domains of the protein. Next, we found by sequence alignments, that the S183I, T260P, I406T, D580E, F789L mutations did not alter RIG-I function.

**Missense SNPs differentially affect RIG-I-mediated innate immune signaling**

Elucidating the functional role of non-synonymous SNPs in RIG-I may enhance our understanding of viral pathogenesis and host defense mechanisms as well as to contribute to a more detailed knowledge in structure-function relationship of RIG-I. To this effect, plasmids containing the eight SNPs were generated by site-directed PCR mutagenesis. We first observed that R7C, S144F, S183I, I406T, D580E, F789L mutations did not alter expression and/or stability of RIG-I protein using western-blot (Fig. 2A) and flow cytometry (Fig. 2B) analyses. Also, the molecular weight of all RIG-I proteins was similar, with the exception of P229fs RIG-I which resulted in a truncated protein with a size comparable to the 2CARD module (Fig. 2A).

To determine whether non-synonymous SNPs can alter RIG-I-induced antiviral and/or pro-inflammatory signaling pathways, we used a functional cell-based assay to evaluate RIG-I-dependent promoter activities, at a level above that induced by the WT RIG-I and lower in cells expressing the R7C, S144F, S183I, I406T, D580E, F789L mutants. To this end, we reported activities of RIG-I-dependent promoter, respectively. We first observed that R7C, S144F, S183I, I406T, D580E, F789L mutations did not alter induction of IFN-β promoters or an NF-κB-dependent promoter, respectively. We first checked the level of constitutive activation of the RIG-I constructs in absence of any signal-eliciting stimulus in HEK 293T or BEAS-2B cells (Fig. 2C) shows a moderate but highly significant constitutive IFN-β expression and NF-κB activity - but no NF-kB activity (not shown) - in WT RIG-I-transfected cells (n = 3, p < 0.001, when compared to control vector-expressing cells), in agreement with the fact that RIG-I is especially prominent in signaling pathways leading to type I IFNs [2,3,4,15].

Interestingly, IFN-β expression and IRF-3 activity in cells expressing T260P, I406T or F789L mutants was similar to that induced by WT RIG-I and lower in cells expressing the R7C, S144F, D580E RIG-I (n = 3, p ≤ 0.0002). With regard to P229fs RIG-I, we observed a salient constitutive IFN-β and NF-kB reporter activities, at a level well above that induced by the full-length form of WT RIG-I (n = 3, p < 0.0001; Fig. 2D). In addition, P229fs SNP induces the expression of endogenous inflammatory and antiviral chemokines such as IL-8 and RANTES, respectively (Fig. 2E), at a level comparable to that of 
(grey bars) and/or HEK 293T cells (black bars) were co-transfected with a three is shown. // in (vector-transfected cells (activity of triplicate samples minus basal activity measured in empty expressed as the mean 6

SNPs RIG-I (filled bars) (luciferase-reporter plasmid and a vector encoding WT (empty bars) or

Stimulated HEK 293T cells as shown in panels (C–E) BEAS-2B (grey bars) and/or HEK 293T cells (black bars) were co-transfected with a β-galactosidase reporter plasmid and either a NF-κB- or IFN-β-luciferase-reporter plasmid and a vector encoding WT (empty bars) or SNPs RIG-I (filled bars) (C–E) or WT 2CARD (E) or a control plasmid. Data were collected 42 h (C) or 24 h (D–E) post-transfection and are expressed as the mean ± SD of RLU normalized to β-galactosidase activity of triplicate samples minus basal activity measured in empty vector-transfected cells (C–D). One representative experiment out of three is shown. // in (C) means that this condition was not tested. (E) Stimulated HEK 293T cells as shown in panels (D) were assessed for IL-8 and RANTES release by ELISA. Data are mean ± SD of triplicate samples and are representative of three independent experiments.

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trigged by the 2CARD module. This finding is particularly important as it suggests that individuals carrying such mutation may constitutively produce exaggerated amounts of immune mediators.

By contrast, no constitutive IFN-β expression was triggered by the S183I RIG-I mutant (n = 3, p<0.0001; Fig. 2C). More importantly, S183I SNP uniquely inhibited IRF-3 (not shown), IFN-β and NF-κB reporter activities elicited by the viral mimetic poly(I:C), in agreement with previous studies that have shown that poly(I:C) is a potent RIG-I stimulus ([16,17,18,19]; Fig. 3A; n = 3

p<0.0001). To confirm the pathophysiological relevance of the above findings, we sought to address the responsiveness of the mutant proteins to viral infection. We clearly demonstrated that S183I mutation had a deleterious effect on RIG-I antiviral activity as it drastically reduced IFN-β and NF-κB-mediated responses triggered by intact, replicative Sendai or influenza A viruses (Fig. 3B and 3C). Noteworthy, while R7C SNP slightly inhibited RIG-I signaling triggered by Sendai virus stimulation, D300E inhibited RIG-I signaling in response to dsRNA and IAV, but not to Sendai virus infection (n = 3, p<0.003). Nevertheless, as S183I SNP uniquely resulted in the strongest inhibition of RIG-I-dependent signaling induced by all stimuli, we decided to focus the rest of our study on this specific mutation. Thus, the clear loss-of-function effect of S183I RIG-I SNP was confirmed by measuring the secretion of IL-8 (n = 3 p<0.0001; Fig. 3D) and RANTES (not illustrated) in the supernatants of stimulated HEK 293T cells. This result well extends Shigemoto et al.'s findings using RIG-I-deficient murine embryonic fibroblasts [20]. Finally, specificity controls are

Impact of RIG-I Polymorphisms

![Figure 2](image2.png)

**Figure 2.** RIG-I-mediated constitutive innate immune signaling, but not expression level, is differentially affected by RIG-I SNPs. Expression of wild-type (WT) and non-synonymous SNPs RIG-I as assessed by western-blot using BEAS-2B cells (A) and flow cytometry (B) using an anti-Flag antibody and BEAS-2B and/or HEK 293T cells 42 h post-transfection. RIG-I SNP proteins are expressed at the same level as WT RIG-I with the exception of P229fs in BEAS-2B cells. (C–E) BEAS-2B cells (+) were co-transfected with a β-galactosidase reporter plasmid and either a NF-κB- or IFN-β-luciferase-reporter plasmid and a vector encoding WT (empty bars) or SNPs RIG-I (filled bars) (C–E) or WT 2CARD (E) or a control plasmid. Data were collected 24 h (C) or 48 h (D–E) post-transfection and are expressed as the mean ± SD of RLU normalized to β-galactosidase activity of triplicate samples minus basal activity measured in empty vector-transfected cells (C–D). One representative experiment out of three is shown. // in (C) means that this condition was not tested. (E) Stimulated HEK 293T cells (+) were assessed for IL-8 and RANTES release by ELISA. Data are mean ± SD of triplicate samples and are representative of three independent experiments.

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![Figure 3](image3.png)

**Figure 3.** SNPs differentially modulate RIG-I-mediated innate immune signaling in response to a viral dsRNA mimetic as well as to influenza A and Sendai viruses. HEK 293T cells (A–B) and BEAS-2B cells (C) were co-transfected with WT (open bars) or non-synonymous SNP RIG-I (black bars) expression vectors and IFN-β- or NF-κB-dependent luciferase reporter plasmids. 24 h later, cells were challenged for 18 h by poly(I:C) (p(I:C), 1 μg/well) (A) or infected with Sendai virus (SeV, 2 HAU/well) (B) or influenza A virus (IAV, MOI = 1) (C). Data are expressed as in Fig. 2c and are representative of three independent experiments. // in (B) means that this condition was not tested. (D) The stimulated or infected cells as shown in panels (A–B) were subsequently assessed for IL-8 release by ELISA. Data are mean ± SD of triplicate samples and are representative of three independent experiments. IL-8 was undetectable in supernatants of non-stimulated transfected cells. (E) S183I SNP does not alter RIG-I-independent signaling. BEAS-2B and HEK 293T cells were co-transfected with WT or S183I RIG-I vectors or empty vector (control) and a NF-κB-dependent luciferase reporter plasmid. 24 h later cells were stimulated with PMA (100 nM) or TNF-α (20 ng/ml).

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provided to make sure the alteration of cell signaling by S183I variant is specific to the RIG-I-dependent pathway. Thus, NF-κB signaling in HEK 293T or BEAS2B cells triggered by two non-viral stimuli (i.e., the cytokine TNFα and the potent PKC signaling activator FMA) was not down-modulated by S183I RIG-I, in comparison with WT RIG-I (Fig. 3E). Altogether, these data stressed the critical role of S183 residue in mediating RIG-I-induced innate immune signaling.

**RIG-I 2CARD module carrying the S183I SNP is unable to trigger signal transduction**

Next, we investigated the mechanism by which S183I SNP results in inhibition of RIG-I antiviral immune response. First, we found that this was neither due to an alteration at a very early step of RIG-I signaling, i.e., the ligand-binding capacity (Fig. 4A) nor to a RIG-I cellular mislocalization (not illustrated). In a very recent report, Fujita’s laboratory also demonstrated that the inhibitory phenotype of S183I RIG-I was neither due to a failure of ubiquitination [20]; a post-translational process essential for RIG-I activity [21]. Next, we took advantage of the fact that the isolated tandem WT 2CARD elicits a vigorous and spontaneous induction of downstream signaling [7] to examine whether S183I mutation could also inhibit this constitutive cell response. As shown in Fig. 4B, contrary to WT 2CARD, the S183I 2CARD could not induce IFN-β and NF-κB activities in HEK 293T (n = 3, p<0.0001) and BEAS-2B cells (not shown). This loss-of-function effect was confirmed by measuring the secretion of endogenous mediators in supernatants of HEK 293T cells (n = 3, p<0.0001; Fig. 4C).

**RIG-I isoleucine 183 residue closes off RIG-I homodimers and RIG-I/MAVS complexes**

CARD domains mediate homotypic or heterotypic interactions

![Figure 4. Analysis of the loss-of-function mechanism of S183I SNP: evidence for an inhibition of the constitutive signal transduction triggered by 2CARD RIG-I.](image)

**Figure 4. Analysis of the loss-of-function mechanism of S183I SNP: evidence for an inhibition of the constitutive signal transduction triggered by 2CARD RIG-I.** (A) S183I does not affect dsRNA binding activity of RIG-I as assessed by a pull-down of Flag-tagged WT and S183I RIG-I proteins using poly(I:C)-coated agarose beads, 42 h after transfection of HEK 293T cells. (B, C) S183I strongly inhibits RIG-I 2CARD-induced IFN-β-dependent antiviral and NF-κB-dependent pro-inflammatory signaling as demonstrated by luciferase reporter assays (B) or by measuring RANTES and IL-8 release by ELISA in HEK 293T cells (C) co-transfected for 42 h with WT or S183I 2CARD or empty expression vector and luciferase reporter plasmids. Data are mean ± SD of triplicate samples and are representative of three independent experiments.

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S183I SNP exerts a down-modulatory effect

Phenotypes of several heritable disorders are linked to missense mutations in single alleles. In some cases, the mutant protein exhibits a regulatory effect whereby heterozygous co-expression of mutant and WT gene has a deleterious consequence, relatively to the case in which two WT alleles are expressed [25,26]. Such a down-regulatory effect usually involves homomeric or heteromeric proteins. In regard to the ability of S183I SNP to impair antiviral
signaling through an increase of RIG-I homocomplexes and RIG-I/MAVS heterocomplexes, it might be speculated that in a heterozygous host, the mutant protein would interfere with the function of the normal protein being produced from the WT allele. To test this hypothesis, we titrated WT RIG-I with increasing amounts of S183I RIG-I in mock treated-HEK 293T and in HEK 293T cells activated by the viral mimetic poly(I:C) or infected by Sendai virus (Fig. 6, panels A–C). As a single example, IFN-β response was reduced by 50% in cells co-transfected with an equimolar concentration of WT RIG-I and S183I RIG-I expressing vectors and further activated by these stimuli (n = 3, p < 0.0001). We also observed that this S183I 2CARD mutant reduced IFN-β activity of WT 2CARD by 50% when transfected at a 1:1 ratio and up to 70% at a fourfold excess of transfected plasmid DNA (n = 3, p < 0.0001, Fig. 6D). Interestingly, the negative impact of S183I SNP was less potent when considering...
RIG-I-mediated NF-κB activity triggered by poly(I:C) or Sendai virus, consistently with the primary role of RIG-I in type I-IFN-inducing antiviral signaling pathways [2,3,4,27].

**Discussion**

The efforts conducted by international consortiums – such as the HapMap Project and Perlegen – to identify and characterize the levels of polymorphic variation in humans has yielded an ever-growing list of SNPs [28]. These include variation located in genes involved in innate immunity, which may account for individual differences in the response to pathogens. For instance, mutations in TLR2, TLR4, TLR5 and IRAK4 have all been associated with increased risk to develop infectious diseases [13,29,30]. In regard to genes encoding CARD-containing proteins, mutations in the peptidoglycan receptors NOD1 and NOD2 have been associated to several inflammatory disorders, including Crohn’s disease, Blau syndrome and asthma [23]. A non-synonymous SNP in MDA-5 was also reported to show an association with type I diabetes [31].

Remarkably, no human disease has yet been linked to RIG-I. Nonetheless, the defective response of a human hepatoma cell line, found permissive to HCV replication, was due to a single mutation (T55I) [9,21,32]. Here, we characterized functional effects of RIG-I SNPs that might help us to understand the basis of individual variations between normal and abnormal innate immune responses to viral pathogens as well as to better appreciate the molecular mechanism by which RIG-I is triggered by non-self RNA.

Among the eight RIG-I SNPs reported in NCBI SNP database, we characterized two distinct functional SNPs which strongly alter RIG-I-mediated signaling. First, we identified P229fs as a SNP which results in a truncated constitutively active RIG-I. This finding is particularly important as it suggests that individuals carrying such mutation may constitutively produce exaggerated amounts of antiviral and pro-inflammatory mediators. Conversely, in agreement with a very recent study from T. Fujita’s laboratory [20], we characterized the loss-of-function S183I SNP. Interestingly, this natural mutation allowed us to further demonstrate the importance of S183 in the transient complex formation that is required for proper RIG-I signaling. Thus, our findings strongly support the hypothesis that regulation of RIG-I/RIG-I and RIG-I/MAVS association/dissociation constitutes a major checkpoint of this antiviral signaling pathway.

CARDs-containing proteins are members of a large group of the ‘death domain superfamily’, which also include the DD (death domain) subfamily and the DED (death effector domain) subfamily. ‘death domain superfamily’, which also include the DD (death domain) subfamily and the DED (death effector domain) subfamily.

**Figure 6. Down-regulatory effect of S183I RIG-I SNP.** (A–C) S183I exerts a down-modulatory effect on full-length WT RIG-I-mediated responses as revealed by IFN-β and NF-κB-dependent reporter assays with HEK 293T cells transfected at different ratio with WT RIG-I and/or S183I RIG-I vectors. An empty vector was used to maintain the total plasmid quantity constant. Data represent the mean ± SD of percentage of inhibition by S183I RIG-I of WT RIG-I-dependent constitutive responses (A, “mock”), or after challenge by poly(I:C) (p(I:C); C) of triplicate samples. (D) S183I exerts a down-regulatory effect on WT 2CARD-induced antiviral, but not on pro-inflammatory, responses. HEK 293T cells were transfected as in (A–C) except that expression of full-length WT or S183I RIG-I was replaced by the corresponding 2CARD modules. Data represent the mean ± SD of percentage of inhibition by S183I 2CARD of WT 2CARD-dependent constitutive responses measured in triplicate samples. (A–D) are representative of three independent experiments. doi:10.1371/journal.pone.0007582.g006

Collectively, on the basis of the data presented here, we consider that scirr 183 residue plays a central role in the molecular ordering that leads to RIG-I-mediated NF-κB and IRF-3 activation pathways. Nevertheless, one can wonder how S183I SNP inhibits RIG-I-induced signaling pathways despite its enhancing effect on RIG-I complexes formation. Based on our biochemical assays and structural modeling showing that this mutation does affect hydrophobicity and flexibility of the CARD2 domain of RIG-I but does not influence its ligand binding activity, we hypothesize that S183I rather induces an abortive conformation of RIG-I, rendering it incapable of downstream signaling. Concerning the inhibitory effect of S183I on RIG-I/MAVS-dependent signal transduction, a recent study clearly supports the concept that MAVS association with RIG-I is not per se sufficient for inducing immune gene expression [37].

Thus, a splicing form of MAVS called MAVS 1a, which shares little sequence similarity with WT MAVS but still contains CARD domain as well as a TRAF-binding motif, can interact strongly with RIG-I but cannot trigger cell signaling. Therefore, like S183I, expression of MAVS 1a interferes with the formation of...
productive RIG-I/MAVS signaling complexes, which likely contributes to its inhibitory outcome.

Elucidating the functional role of RIG-I SNPs may enhance our understanding of the pathogenesis of viral infections, to ultimately decrease morbidity and mortality through improved risk assessment and early administration of prophylactic therapies [13,29]. Clinical studies assessing $S_{183}^{I}$ SNP frequency in control healthy individuals and patients infected by viruses will certainly clarify the contribution of RIG-I variation to the pathogenesis of viral diseases. Likewise, investigating the clinical relevance of the potent immunostimulatory $P_{256}^{A}$ SNP may be particularly interesting in patients with autoimmune diseases where cytokines play a pivotal pathogenic role. Among them, evidence linking IFN-$
\alpha$ with autoimmune diseases such as lupus and insulin-dependent diabetes mellitus in humans are the most convincing [30]. Meanwhile, our study demonstrates that serine 183 is a pivotal residue involved in communication between CARD modules of RIG-I themselves as well as with MAVS and emphasizes the complexity of molecular events that governs RIG-I-induced antiviral signaling.

Materials and Methods

Viruses and reagents

Influenza A/Scotland/20/74 (H3N2) virus was previously described as previously described [39]. Sendai virus (Cantell strain, ATCC VR-907 Parainfluenza 1) was a kind gift of E. Meurs (Institut Pasteur, Paris, France). The viral dsRNA mimetic polynucleosome: polycytidylic acid (poly(LC)) and phosphor 12-myristate 13-acetate, (PMA) were from Sigma. Human recombinant TFN-$\alpha$ was purchased from Peprotech.

Phylogenetic analysis of RIG-I SNPs

RIG-I SNPs were as described in NCBI’s SNP database (cf. http://www.ncbi.nlm.nih.gov/SNP/snp_ref.cgi?locusId = 23586). RIG-I sequences from human to platypus were aligned using EMBL ClustalW software and manually arranged.

Plasmids construction and site-directed mutagenesis

The pEFBOS(+)-Flag-RIG-I (amino acids 2–925) or 2CARD (amino acids 2–229) vectors were previously described [17] and pcDNA3-Flag-MAVS and pcI-V5-WT 2CARD plasmids were a kind gift of Dr. Z. Chen and Dr. E. Meurs, respectively. SNPs containing plasmids were made using the QuickChange II XL Site-Directed Mutagenesis kit (Stratagene). Sequences of oligonucleotides used for mutagenesis are indicated in Table S1. An Site-Directed Mutagenesis kit (Stratagene). Sequences of oligonucleotides used for mutagenesis are indicated in Table S1. An in vitro recombination-based cloning (Gateway system; Invitrogen) was used to generate biotin-tagged WT or $S_{183}^{I}$ RIG-I, as previously described [40]. Briefly, biotin-tagged RIG-I (WT or $S_{183}^{I}$) expression vectors were generated by PCR using pEFBOS(+) Flag-RIG-I as a template and the following forward (5’-ggggacaactttgtagaantaacagccagt ACCACCGGACGCAGGCA-GCA-3’) and reverse primers (5’-ggggacaccttctgattagggtgattattgtaTTGGGACA TTTCCTGTGGATGATAATTG-G-3’), as well as the Gateway technology for a final cloning in pcDNA6/BioEase-DEST plasmid (Invitrogen), according to manufacturer’s instructions.

All constructs were entirely sequenced to confirm that no unintended mutations were generated during PCR reaction.

Cell culture, transfection, ELISA and luciferase assays

Detailed protocols were described before [39]. Data are expressed as the mean ($\times 10^{-5}$) of relative luciferase units (RLU) normalized with $\beta$-galactosidase activity minus basal activity measured in empty vector-transfected cells.

Immunoblot and protein-protein interactions analysis

EK 293T cells were transiently co-transfected with 1 $\mu$g of Flag-tagged MAVS or V5-tagged 2CARD vectors (for tandem 2CARD/MAVS interaction analysis) or 3 $\mu$g of Flag-tagged 2CARD and biotin-conjugated RIG-I vectors (for tandem 2CARD/RIG-I interaction). After cell disruption and a pre-clearing step, pull-down of biotin-tagged RIG-I was performed using streptavidin sepharose beads (GE Healthcare). For co-immunoprecipitation assay, cell lysates were incubated with a monoclonal anti-Flag M2 antibody, followed by the addition of protein G sepharose beads. More detailed protocols can be provided upon request. After centrifugation and protein denaturation, samples were analyzed by immunoblot as described in reference [39].

dsRNA binding assay

Assay of dsRNA binding activity of RIG-I (WT or $S_{183}^{I}$) was previously described [41]. Briefly, HEK 293T cells were seeded in 100 mm tissue culture dishes and transiently transfected with 0 $\mu$g of control plasmid or vector encoding Flag-tagged RIG-I (WT or $S_{183}^{I}$). 48 h post-transfection, cells were disrupted in 1.5 ml of RIPA lysis buffer and 400 $\mu$g of cell lysates were incubated with poly(LC)- or control poly(C)-coated agarose beads (Sigma) in RIPA lysis buffer supplemented with proteases inhibitors cocktail and 50 U/ml of RNAse inhibitor (Promega) for 1 h at 4°C. Agarose beads were then collected by centrifugation and washed three times with lysis buffer before resuspension in 30 $\mu$l sample denaturing buffer.

Flow cytometry and fluorescence microscopy analysis.

To evaluate RIG-I (WT or SNP) protein expression levels and subcellular localization, BEAS-2B and HEK 293T cells were transfected and processed as previously described [42], using the following antibodies: anti-Flag antibody (2 $\mu$g/ml) and Alexa flour conjugated secondary antibody (4 $\mu$g/ml, A11001, Molecular probes).

Computational modeling and structural analysis of RIG-I CARD #2

Homology modeling and molecular dynamics of the human RIG-I CARD #2 domain were carried out as previously described by Kuharenko et al. [24] based on several CARD domain structures: 1cww [43], 2vqg [34], 3crd [44], 1dgn [45] and 2b1w [46]. The sequence identity for 2vqg and RIG-I CARD domains is between 21–26% (depending on the alignment algorithm used). The method of comparative/homology modeling was therefore applied [47]. Structure analysis was carried using the following software (referenced in [24]) SwissPBD Viewer and PyMol (www. pymol.org) for visualization; HotPatch [48] for hydrophobicity and PDB2PQR [49], PropKa [50] and APBS [51] for charged surface calculation. Further details are available on request.

Statistical analysis

Statistical differences were tested using a one-way ANOVA followed by a Fisher test, with a threshold of $p<0.05$.

Supporting Information

Table S1 Plasmids containing SNPs were made by site-directed mutagenesis using the QuickChange II XL Site-Directed Mutagenesis kit (Stratagene), 125 ng of specific forward and reverse primers and 25 ng of RIG-I WT vector as a template in 50 $\mu$l reaction volume. After an initial denaturation step at 95°C for 1 min, mutagenesis was performed by 18 cycles of amplification (1 min at 95°C, 50 s at 60°C and 9 min 30 s at 68°C), followed by
Figure S1 Alignment of amino acid sequence of CARD#2 domain of RIG-I with other CARD structures. Alignment of CARD domain sequences from different CARD proteins which were used for RIG-I CARD#2 modeling. Helix 1 colored in red, helix 2 in orange, helix 3 in yellow, helix 4 in green, helix 5 in blue and helix 6 in brown. For Apa1, MAVS, RAIDD, ICEBERG and NOD1 CARDs, helix boundaries were determined directly from the respective PDB files 1cwv, 2vgq, 3crd, 1dgn and 2b1w, based on a 3D alignment of these structures. Found at: doi:10.1371/journal.pone.0007582.s001 (0.03 MB DOC)

Figure S2 Intermediate frames of WT and S183I CARD#2 structures during molecular dynamic simulation. Eleven frames from the molecular dynamics simulation of RIG-I CARD#2 WT (A) or S183I mutant (B), showing one frame per picosecond. First frame (0 ps) corresponds to the initial raw model formation. Helices from 1 to 6 are rainbow-colored, helix 6 which harbors S183 is colored red. Found at: doi:10.1371/journal.pone.0007582.s002 (0.91 MB DOC)

Figure S3 Comparison of the stability of WT and S183I CARD#2 structures over the course of molecular dynamic simulation. Root mean square deviation (RMSD) calculated from 10 ps molecular dynamics simulation mapped onto the initial raw models of WT RIG-I CARD#2 (A) or S183I mutant (B) structures. More stable elements are colored green, more flexible regions in red. Found at: doi:10.1371/journal.pone.0007582.s004 (1.37 MB DOC)

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Author Contributions

Conceived and designed the experiments: JP MST. Performed the experiments: JP AB AK BS ANW. Analyzed the data: JP AB AK GC FT POV ANW MC MST. Contributed reagents/materials/analysis tools: GC MBA LQM AH JD POV. Wrote the paper: JP AK ANW MST.
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