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A systematic analysis of recombination activity and genotypephenotype correlation in human recombination-activating gene 1 deficiency

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Disclosure of potential conflict of interest:

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The rest of the authors declare that they have no relevant conflicts of interest.

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Abstract

Background—The recombination-activating gene (RAG) 1/2 proteins play a critical role in the development of T and B cells by initiating the VDJ recombination process that leads to generation of a broad T-cell receptor (TCR) and B-cell receptor repertoire. Pathogenic mutations in the *RAG1/2* genes result in various forms of primary immunodeficiency, ranging from T⁻B⁻ severe combined immune deficiency to delayed-onset disease with granuloma formation, autoimmunity, or both. It is not clear what contributes to such heterogeneity of phenotypes.

Objective—We sought to investigate the molecular basis for phenotypic diversity presented in patients with various *RAG1* mutations.

Methods—We have developed a flow cytometry–based assay that allows analysis of RAG recombination activity based on green fluorescent protein expression and have assessed the induction of the *Ighc* locus rearrangements in mouse $Rag1^{-/-}$ pro-B cells reconstituted with wild-type or mutant human RAG1 (hRAG1) using deep sequencing technology.

Results—Here we demonstrate correlation between defective recombination activity of hRAG1 mutant proteins and severity of the clinical and immunologic phenotype and provide insights on the molecular mechanisms accounting for such phenotypic diversity.

Conclusions—Using a sensitive assay to measure the *RAG1* activity level of 79 mutations in a physiologic setting, we demonstrate correlation between recombination activity of RAG1 mutants and the severity of clinical presentation and show that RAG1 mutants can induce specific abnormalities of the VDJ recombination process.

Keywords

Recombination-activating gene 1; V(D)J recombination; severe combined immune deficiency; Omenn syndrome; autoimmunity; genotype-phenotype correlation; immune repertoire

The recombination-activating gene (RAG) 1 and RAG2 proteins initiate the VDJ recombination process by generating DNA double-strand breaks at the recombination signal

sequences (RSSs) that flank the variable (V), diversity (D), and joining (J) gene segments of the immunoglobulin and T-cell receptor (TCR) genes. These DNA double-strand breaks are then joined by the ubiquitous mechanism of nonhomologous end-joining machinery. As a result of VDJ recombination, a polyclonal pool of functional T and B lymphocytes is generated, expressing a diverse repertoire of productive TCR and B-cell receptor rearrangements.

Null mutations of *RAG1* and *RAG2* genes result in the T⁻B⁻ severe combined immune deficiency (SCID) phenotype.³ However, hypomorphic *RAG* mutations have been associated with a spectrum of clinical and immunologic phenotypes that include Omenn syndrome (OS),^{4–10} with erythroderma, lymphadenopathy, eosinophilia, increased serum IgE levels, and the presence of autologous, oligoclonal, and activated T lymphocytes; leaky/atypical SCID,¹⁰ with varying numbers of T and B cells but without the typical features of OS; SCID with expansion of $\gamma\delta$ T lymphocytes ($\gamma\delta$ -T),^{11,12} which us often associated with cytomegalovirus infection; delayed-onset combined immune deficiency with granuloma and/or autoimmunity (CID-G/A)^{13–15}; and in a single case of idiopathic CD4⁺ T cell lymphopenia (ICL), presenting with extensive chickenpox and recurrent pneumonia.¹⁶

Attempts to correlate the phenotypic diversity of RAG-related disorders in human subjects with functional activity of the mutant proteins were largely based on a transient transfection assay in nonlymphoid adherent cells. ¹⁷ In this assay cells are cotransfected with plasmids encoding for wild-type (or mutant) human RAG1 (hRAG1) and hRAG2 and a third plasmid containing a suitable recombination substrate that would allow expression of an antibiotic resistance gene upon recognition and cleavage by hRAG1 and hRAG2 and nonhomologous end joining–mediated ligation. However, with this assay, the recombination activity of RAG proteins is analyzed on an extrachromosomal substrate (ie, a nonphysiologic setting), and functional impairment of mutants that specifically affect nuclear translocation of the hRAG proteins might be missed. Furthermore, it has been shown that stability and posttranslational modifications of the RAG proteins differ significantly in lymphoid versus nonlymphoid cells. ¹⁸

Recently, Abelson murine leukemia virus (A-MuLV)— transformed pro-B cells containing an inverted green fluorescent protein (GFP) cassette flanked by RSS (pMX-INV) have been used to measure VDJ recombination activity on an intrachromosomal substrate by using flow cytometry with GFP expression as a read-out. ¹⁹ On the basis of this platform, we have analyzed the expression and recombination activity of 79 naturally occurring hRAG1 mutant proteins and thereby performed the largest comprehensive analysis of genotype-proteotype-phenotype correlation for hRAG1 deficiency. Our results provide novel insights into the molecular mechanisms underlying phenotypic diversity in patients with this disease.

METHODS

Patient selection and assignment to phenotypic subgroups

Deidentified patients' clinical, immunologic, and molecular data were provided by an international network of physicians in Europe, the Middle East, South America, and the United States, according to protocols approved by the local institutional review boards. On

the basis of phenotype, each patient was assigned to one of the following subgroups: T^B^- SCID, OS, $\gamma\delta$ -T, atypical/leaky SCID-G/A, and ICL.

Determination of recombinase activity level of wild-type and mutant RAG1

Analysis of recombination activity of wild-type and mutant RAG1 constructs was performed, as previously described. ¹⁴ For detailed methods, see the Methods section in this article's Online Repository at www.jacionline.org.

Immune repertoire analysis—Analysis of rearrangements at the endogenous Ighc locus was performed by using as a template the mRNA extracted from A-MuLV pro-B cells transduced with retroviral vectors encoding for wild-type or mutant hRAG1. A set of nested primers specific for various V_H and C_H elements of the Ighc locus were used for the first amplification cycles, followed by amplification with communal primers according to the manufacturer's protocol for MBCR (iRepertoire, Huntsville, Ala). By using this protocol, ampliconrescued multiplex PCR allows semiquantitative amplification of the immune repertoire. 20 Purified PCR products were sequenced with the GS Junior 454 platform (Roche, Mannheim, Germany).

Raw sequences were filtered for PCR errors, and a tree map and 2-dimensional map were generated from the total complementarity-determining region (CDR)–H3 sequences to analyze V_H - J_H pairing and the relative distribution of distinct rearrangements (iRepertoire). The filtered sequences, excluding duplicates, were further analyzed for V_H , D_H , and J_H gene usage; composition; CDR-H3 length; reading frame (RF) determination; and the diversity index of Shannon entropy by using IMGT HighV-QUEST output files²¹ and IgAT software.²² Rarefaction curves were generated by using the VDJ statistics file from IgAT analysis and the PAST program.²³ To assess reproducibility, 2 runs on independent amplification libraries were performed for R699Q, and 2 to 3 repeats of the same amplification library were run for the R314W and M435V mutants.

Statistical analysis—Results of recombination activity of hRAG1 proteins were analyzed with the Mann-Whitney U test. The χ^2 test was used to assess skewing in the use of V_H , D_H , and J_H genes by various hRAG1 mutant proteins compared with wild-type hRAG1. The unpaired 2-tailed t test was used to analyze the average CDR-H3 length, the average index of hydrophobicity according to a normalized Kyte-Doolittle scale, differential use of 3 RFs, and the average Shannon entropy of the CDR-H3 loop. Nonparametric Spearman correlation was used to determine correlation between the recombination activity level and protein expression of various hRAG1 mutants by using the Spearman correlation coefficient (rs).

RESULTS

Recombination activity and protein expression of hRAG1 mutants

We have compiled a list of 79 distinct, naturally occurring h*RAG1* genetic variants, including 3 known polymorphisms (G99S, H249R, and K820R) and 76 distinct mutations identified in 68 patients (among which were 34 novel cases, Table I).^{4–9,11–16,24–28} To investigate correlation between the expression and function of mutant hRAG1 and the

patients' clinical and immunologic phenotypes, we used A-MuLV- transformed Rag1^{-/-} tg.E μ -bcl2 pro-B cells (hereafter referred to as $Rag l^{-/-}$ Abl pro-B cells), containing a single integration of the pMX-INV GFP cassette flanked by RSSs (Fig 1, A, and see Fig E1 in this article's Online Repository at www.jacionline.org), ²¹ which were further transduced with retroviral vectors encoding wild-type or mutant hRAG1 or wild-type mouse Rag1 (mRag1) and hCD2 as a reporter. To control for expression of wild-type and mutant RAG1 proteins, $Rag 1^{-/-}$ Abl pro-B cells expressing either wild-type or mutant hRAG1 were equally enriched to contain greater than 90% cells expressing the hCD2 reporter. Comparable levels of GFP expression were observed on induction of wild-type hRAG1 and mRag1 expression with STI-571 (imatinib; Fig 1, B). This treatment arrests cells in the G0/G1 phase of the cell cycle while RAG proteins are expressed.²⁹ We then measured the VDJ recombination activity of 79 different naturally occurring hRAG1 variants by normalizing GFP expression to that observed in the presence of wild-type hRAG1 (Fig 1, C, and see Table E1 in this article's Online Repository at www.jacionline.org). Wild-type levels of VDJ recombination activity were observed for 3 known hRAG1 polymorphisms (G99S, H249R, and K820R) and for N106K, R449K, R474C, H612R, A868V, and M1006V mutants, suggesting that the latter represent neutral variants and not disease-causing mutations.

We investigated whether the specific location of RAG1 mutations might impinge on recombination activity. Fig 2, A, shows the location of the mutations analyzed in this study along the length of the hRAG1 protein; different colors were used to indicate the diverse clinical and immunologic phenotypes. For mutations associated with more than 1 phenotype, the color code corresponds to the least severe phenotype under the assumption that the mutation is potentially permissive for some V(D)J recombination activity. We observed a relative overrepresentation of mutations affecting the nonamer-binding region (NBR) of hRAG1 (Fig 2, B). Moreover, missense mutations in the NBR and in the heptamer-binding region (HBR) had a significantly lower activity level than mutations in other regions of the RAG1 protein (Fig 2, C), which is consistent with the critical roles played by the NBR and HBR domains in DNA recognition and binding. 1 In contrast, mutations at the C-terminus of RAG1 and some missense mutations in the central region of the molecule showed significant residual VDJ recombination activity (Fig 2, B). These mutations have been identified mostly in patients with CID-G/A (ie, a less severe and often delayed-onset form of the disease), suggesting that mutations in the C-terminus of RAG1 are more tolerable than mutations in the NBR/HBR.

Next, we analyzed correlation between recombination activity of hRAG1 mutant proteins and the patients' clinical and immunologic phenotype (Fig 2, *D*). Recombination activity of those hRAG1 mutants that had been identified in patients belonging to various clinical subgroups were reported for each of the subgroups to avoid a bias in the statistical analysis of the data. VDJ recombination activity levels were greater for *RAG1* mutations identified in patients with CID-G/A and were progressively lower in patients with more severe phenotypes, being extremely low for patients belonging to the T⁻B⁻ SCID subgroup (Fig 2, *D*). Furthermore, significantly lower VDJ recombination activities of hRAG1 mutants were observed in patients with very low to absent circulating B cells (Fig 2, *E*). Altogether, these data indicate that there is a strong, although imperfect, correlation between the VDJ

recombination activity of mutant hRAG1 proteins and the associated clinical and immunologic phenotype.

To determine whether differences in recombination activity of hRAG1 mutants might reflect different levels of protein expression, we performed Western blotting analysis of 41 RAG1 mutants introduced into $Rag 1^{-/-}$ Abl pro-B cells. The same amount of cells (400,000 cells), of which greater than 90% expressed hCD2, were loaded in each lane, and adjusted RAG1 expression was calculated by using the loading control as a reference. Of 27 hRAG1 mutants in the NBR/HBR that were studied, 13 showed very low levels of protein expression (adjusted expression, <0.2; Fig 3, A and B). Variable levels of protein expression were observed for NBR mutants with different amino acid substitutions at the same codon (R394 and R396; Fig 3, A), yet all of them showed very low to undetectable VDJ recombination activity (see Table E1). Among 15 missense mutations located outside of NBR/HBR domains, 4 had very low levels of protein expression, and 7 had expression levels comparable with those of wild-type hRAG1 (Fig 3, C). Overall, levels of protein expression of hRAG1 mutants affecting the NBR/HBR regions were not significantly different than those of mutants affecting other domains (Fig 3, D). Furthermore, we observed only a modest correlation between RAG1 mutant protein expression and recombination activity (Fig 3, *E*).

Comparison of recombination activity and predicted pathogenicity of hRAG1 mutants

Bioinformatics tools might help predict the pathogenicity of DNA variants and missense mutations in particular. To predict the pathogenicity of 63 hRAG1 missense variants, we used the Pathogenic Or Not Pipeline (PON-P), which integrates information provided by Phd-SNP, SNAP, PolyPhen-2, SIFT, and I-Mutant predictors into a single consensus prediction with a reliability score.³⁰ PON-P assigned each hRAG1 variant to a specific class according to probability and reliability scores (P, pathogenic; N, neutral; and UV, unclassified variant). Most variants in class P had absent or negligible recombination activity, whereas most variants in class N had wild-type recombination activity (see Table E1). Overall, correlation between predicted pathogenicity and in vitro recombination activity was not perfect. However, among missense mutations located in the NBR, those affecting residues that are directly involved in DNA binding³¹ were assigned to class P by using PON-P, and they all showed absent or markedly decreased recombination activity (see Fig E2 in this article's Online Repository at www.jacionline.org). In contrast, missense mutations in the more distal α -helix of the NBR, affecting residues that are not directly involved in DNA binding, were mostly assigned to class UV and had low but detectable recombination activity. Therefore PON-P can be informative in validating experimental results and predicting mutations in regions with distinct and imperative activities.

Analysis of rearrangements at the endogenous Ighc locus

Introduction of hRAG1 mutants into $Rag1^{-/-}$ Abl pro-B cells offers the opportunity to analyze VDJ recombination activity in a physiologic setting through the study of rearrangements at the endogenous Ighc locus. For this purpose, we used a semiquantitative deep-sequencing approach based on amplification of mRNA transcripts by using V_H forward primers and C_H reverse primers with a communal barcode (provided by iRepertoire)

and use of the Roche 454 pyrosequencing platform. In particular, we tested the frequency, quality, and diversity of *Ighc* rearrangements induced by transduction of *Rag1*^{-/-} Abl pro-B cells with retroviral vectors encoding either wild-type hRAG1 or mRag1 or 10 distinct hRAG1 mutants (K86Vfs33X, R142X, R314W, C328Y, M435V, G516A, R699Q, R841W, Y912C, and K992E). These mutants were selected because they occur in different domains of the protein, support different levels of VDJ recombination activity (as assessed based on GFP expression), and were identified in patients representative of the phenotypic spectrum of the disease. The total number of reads, as well as total and unique CDR-H3 sequences, were comparable between mRag1, hRAG1, and R699Q, whereas a reduced number of reads or no reads were obtained for all other mutants with lower activity levels than R699Q (see Table E2 in this article's Online Repository at www.jacionline.org).

To assess whether we had collected a sufficient number of reads to cover the diversity of rearrangements induced by each mutant, we analyzed individual rarefaction curves in which the number of species (VDJ rearrangement products) is plotted as a function of the number of sequences obtained, thereby informing on the richness of the reads (see Fig E3, *A*, in this article's Online Repository at www.jacionline.org). Although a very low number of reads was obtained for both the R314W and K992E mutants, the rarefaction curves for these mutants were flatter than for wild-type hRAG and for other mutations of the same sample size, indicating that an increase in sequence reads for R314W and K992E samples would not result in a significant increase in the number of diverse VDJ recombination products (see Fig E3, *B*). Furthermore, analysis of individual V_H to J_H pairing demonstrated progressive restriction of the *Ighc* rearrangement repertoire when moving from hRAG1 mutants with high to those with low VDJ recombination activity (Fig 4, *A*–*F*).

As expected, 18,32 introduction of wild-type hRAGI or mRagI into $RagI^{-/-}$ Abl pro-B cells induced Ighc rearrangements that often use the most proximal V_H gene segment Ighv5-2 (see Fig E4 in this article's Online Repository at www.jacionline. org). In addition to preferential use of this V_H gene segment, skewing in the use of individual V_H , D_H , and J_H genes was observed for all mutants tested and especially for the R314W and K992E mutants (Fig 4, G–I, and see Fig E5 in this article's Online Repository at www.jacionline.org) and for D_H use (Fig 4, H, and see Fig E5, C).

Differences in the composition and length of the CDR-H3 region might affect the ability of immunoglobulins to bind to antigens. Dynamic changes in CDR-H3 composition occur during differentiation from pro-B to mature B cells. In particular, immature B cells enriched for self-reactive specificities tend to have longer CDR-H3 sequences with presence of positively charged amino acid residues. Because immune dysregulation is a common feature in patients with hypomorphic RAG mutations, we sought to determine whether introduction of these mutants into $Rag1^{-/-}$ Abl pro-B cells induces expression of CDR-H3 sequences that differ in length and composition from those induced by wild-type hRAG1. There was a slight but significant increase in the length of CDR-H3 for both the R699Q (35.24 \pm 8.76 nt) and G516A (35.34 \pm 8.54 nt) mutants compared with hRAG1 (34.24 \pm 8.76 nt; P < .0001 and P = .0007, respectively; Fig 4, J), mainly because of a longer DH gene segment (see Fig E6, A and B, in this article's Online Repository at www. jacionline.org). However, the more severe K992E mutant, which is associated with OS,

supported generation of Ighc rearrangements characterized by markedly shorter CDR-H3 sequence (26.57 ± 8.64 nt, P = .0055; Fig 4, J). A reduced number of N nucleotide additions at the VDJ recombination junctions (see Fig E6, A and C), a higher frequency of direct V_{H} -to- J_{H} rearrangements (see Fig E6, D), a significant decrease in the average D_{H} length (see Fig E6, B), and a higher frequency of use of the shorter J_{H} 3 gene segment (Fig 4, I, and see Fig E5, D) all contributed to the shorter CDR-H3 sequences, likely reflecting the impaired function and abnormal VDJ recombination activity of the K992E mutant.

For the R699Q, G516A, and M435V mutants with higher levels of recombination activity, the average index of CDR-H3 hydrophobicity according to Kyte-Doolittle did not deviate much from neutral to slightly hydrophilic, which is similar to what was observed for wild-type hRAG1 (Fig 4, *K*).³³ In contrast, a significant shift toward a charged/hydrophilic profile of CDR-H3 sequences was observed for the more severe R314W and K992E mutants (Fig 4, *K*), mainly because of abundance of arginine, asparagine, and aspartic acid residues (see Fig E7 in this article's Online Repository at www.jacionline.org). The higher frequency of these amino acids resulted mainly from N1 additions in the case of the R314W mutant and from N2 additions for the K992E mutant (see Fig E8 in this article's Online Repository at www.jacionline.org). Moreover, the nature of the RF, which is largely determined by D_H genes, also contributed to the skewed use of these amino acids. In particular, the R314W mutant induced preferential use of the neutrally charged RF1 and reduced use of the hydrophobic RF2, whereas the K992E mutant induced more abundant use of the hydrophilic RF3 (Fig 4, *L*).

Finally, to analyze the diversity of amino acid composition at each residue of the CDR-H3, we used the Shannon entropy index (H), a measure of variability in which the higher the score, the more variable the amino acid use at each position. The CDR-H3 diversity was significantly reduced for the M435V, R314W, and K922E mutants but was only modestly decreased for R699Q and G516A mutants (Fig 4, *M*). Homozygosity for the K992E mutation in human subjects is associated with OS, whereas the R699Q and G516A mutations have been identified only in patients with CID-G/A (Table I). Altogether, these data indicate that restricted diversity in the immune repertoire correlates with severity of impairment of RAG1 function.

DISCUSSION

An important property of the $Rag1^{-/-}$ Abl pro–B-cell system described here is that it allows in-depth analysis of VDJ recombination activity at the endogenous Ighc locus before selection processes. As expected for pro-B cells, we observed that D_H1 and D_H4 (commonly known as DFL16 and DQ52, respectively) genes were preferentially targeted for rearrangement. Furthermore, significant skewing toward D_H4 gene use was observed with the more severe K992E mutant, which reflects impairment of secondary D-replacement rearrangements.

Similarly, the most D_H -proximal V_H^2 and V_H^3 genes were preferentially targeted in V_H^3 -DJ_H rearrangements, as also seen early during B-cell ontogeny. Even for these genes, a skewed use by various hRAG1 mutants was observed. Moreover, increased targeting was

observed for the other D_H -proximal V_H11 and V_H14 genes. Although the association of individual hRAG1 mutations with preferential targeting of specific V_H genes has not been previously reported, significantly reduced D_H - J_H rearrangement in mice homozygous for a hypomorphic Rag1 mutation had been demonstrated. In addition, introduction of an OS-associated mutation into them Rag1 gene induced selective impairment in the ability to target certain coding flanks at V_H , D_H and J_H gene segments. Along with our findings, these data suggest that reduced RAG1 activity might directly alter the quality of endogenous VDJ rearrangements. Overall, hRAG1 mutants with lower recombination activity preferentially targeted the most proximal D_H elements, whereas hRAG1 mutants with relatively higher recombination activity showed the ability to target D_H distal V_H gene segments, thereby allowing generation of a more diverse repertoire. This correlated with a milder clinical and immunologic phenotype $in\ vivo$, with preservation of B-cell development and detectable immunoglobulin levels.

Immune dysregulation is being increasingly recognized as an important clinical phenotype associated with hypomorphic *RAG1* mutations (Table I).^{13–15} The R699Q and G516A mutants induced rearrangements with longer CDR-H3 sequences, a feature that has been previously identified in self-reactive B cells.^{34,35,39} These data suggest that some h*RAG1* mutations might favor generation of a self-reactive B-cell repertoire by skewing the quality of VDJ rearrangements. On the other hand, generation of markedly shorter CDR-H3 sequences caused by increased use of the stop codon–rich RF3 has been observed for the K992E mutant identified in 2 patients with OS.

One caveat of the $Rag 1^{-/-}$ Abl pro–B-cell system described here is that it allows in-depth analysis of the recombination properties of one mutant at a time. In contrast, many patients carry compound heterozygous mutations. Use of polycistronic vectors might allow analysis of the combined effect of 2 distinct mutations.

In conclusion, this study represents the most comprehensive study of the expression and function of hRAG1 mutants identified in patients with a spectrum of clinical and immunologic presentations. Using this platform, we have demonstrated genotype-phenotype correlation and have provided novel molecular insights that might help explain the phenotypic diversity of this disease. The results presented in this study could serve as a reference for recombination activity of known RAG1 variants. Moreover, the fluorescence-activated cell sorting–based assay described here could represent a helpful cellular platform to assess the pathogenicity of newly identified RAG1 variants. This would be especially important in the diagnostic approach to asymptomatic or presymptomatic subjects with RAG1 mutations, including those identified at birth through newborn screening. Note added in proofs: Yu et al⁴⁰ have recently demonstrated restricted TCR β diversity in patients with OS, and preserved diversity, but skewed usage of V, D and J elements in patients with CID-G/A.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations used

A-MuLV Abelson murine leukemia virus

CDR Complementarity-determining region

CID-G/A Combined immune deficiency with granuloma and/or autoimmunity

 $\gamma \delta$ -T SCID with expansion of $\gamma \delta$ T lymphocytes

GFP Green fluorescent protein

HBR Heptamer-binding region

hRAG1 Human RAG1

ICL Idiopathic CD4⁺ T-cell lymphopenia

mRag1 Mouse Rag1

NBR Nonamer-binding region

OS Omenn syndrome

PON-P Pathogenic Or Not Pipeline

RAG Recombination-activating gene

RF Reading frame

RSS Recombination signal sequence

SCID Severe combined immune deficiency

TCR T-cell receptor

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Key messages

 We describe a cellular platform that permits rapid analysis of expression and recombination activity of RAG1 mutant variants.

- RAG1 recombination activity correlates with the severity of the clinical and immunologic phenotype observed *in vivo*.
- Mutant RAG1 proteins differ in the efficiency and quality of V(D)J recombination activity, indicating that skewing of the immune repertoire might occur independently from *in vivo* selection.

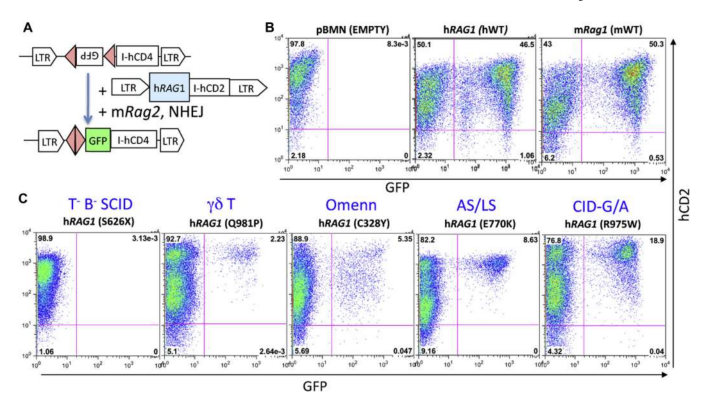


FIG 1.

Schematic representation of the experimental outline and readout. **A**, A mouse Abelson (Abl) virus—transformed pro—B-cell line deficient for m*Rag1* was infected with a retrovirus containing an inverted GFP cassette. Subclones with single-copy stable integrants were transduced with vectors expressing wild-type *RAG1* or various h*RAG1* mutations (see Table E1) and then treated with imatinib to promote cell differentiation and induction of RAG1 activity. B and C, The level of GFP expression indicated the recombinase activity level on imatinib stimulation of *Rag1*—/— Abl pro-B cells transduced with an empty vector, or with vectors encoding either for wild-type hRAG1 (hWT) or wild-type mRag1 (mWT; Fig 1, B) or for one representative hRAG1 mutation for each of the 5 different phenotypic subgroups of the disease (Fig 1, C). LTR, Long terminal repeats.

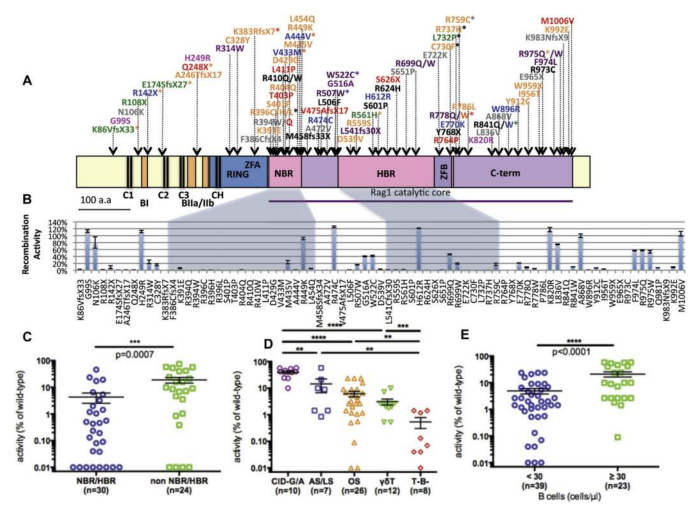
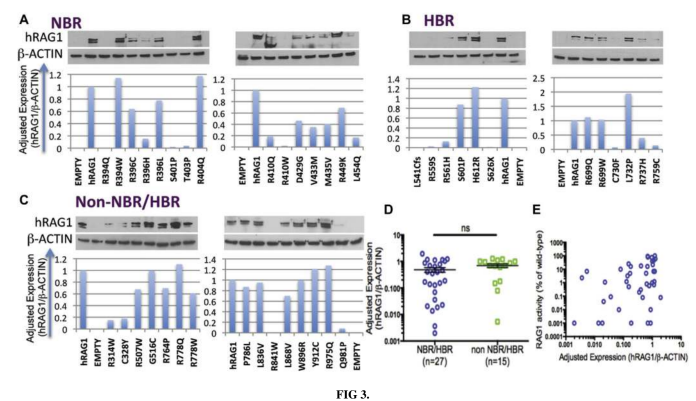


FIG 2.

Activity levels of 79 genetic variants of hRAG1. A, Schematic representation of 79 genetic variants of hRAG1 affecting the various domains: RING, zinc finger RING type domain (amino acids 168–283); ZFA, zinc finger A; NBR (amino acids 387– 461); HBR (amino acids 531–763); ZFB, zinc finger B; and the core domain from amino acids 385 to 1011. The conserved cysteine (C) and histidine (H) residues are marked with black lines, and the basic domains are marked with bars. They are as follows: C1 (103/115); C2 (169/181); C3 (204/214) CH (269/275); CC (905/910); HH (940/945); BI (142–147); BIIa (219– 225); BIIb (234–237); and BIII (244–257). The RAG1 variants are color coded, corresponding to the clinical phenotype of patients in which they were identified ($red = T^-/B^- SCID$, orange = OS, $green = \gamma \delta - T$, blue = atypical/leaky SCID, and purple= CID-G/A). The asterisk marks mutations associated with other phenotypes. Mutations in black correspond to alleles with the lower recombination activity that had been identified in patients who were compound heterozygous for RAG1 mutations. Known polymorphisms are indicated in *pink*, and *gray* is used to identify variants detected in patients for whom incomplete clinical and immunologic information was available. B, Bar diagram representing the activity level of various hRAG1 mutants relative to wild-type hRAG1. Values are expressed as percentages ± SEMs. For each mutant, 3 to 5 independent experiments were performed. Mutations falling in the NBR and HBR are contained in shaded areas. C, Recombination activity of missense mutations falling in the NBR/HBR versus other regions of hRAG1. D, Recombination activity of hRAG1 mutants identified in patients with a distinct clinical and immunologic phenotype. E, Recombination activity of hRAG1 mutants identified in patients with virtual lack of circulating B cells ($\leq 30 \text{ cells/}\mu\text{L}$) and in those with residual B cells ($\leq 30 \text{ cells/}\mu\text{L}$). The Mann-Whitney U

test was performed to demonstrate statistical significance for all the 1-tailed P values in the graphs: **P < .01, ***P < .001, and ****P < .0001.



Protein expression of hRAG1 mutants. **A–C**, Protein expression of hRAG1 mutants affecting NBR (Fig 3, A), HBR (Fig 3, B), and non-NBR/HBR (Fig 3, C) domains. Expression of β -actin was used to normalize the density for each of the h*RAG1* mutants. Results are shown as adjusted density (ImageJ). One representative of 2 immunoblots is shown. **D**, Adjusted density of hRAG1 protein expression of mutants affecting the NBR/HBR or non-NBR/HBR domains of the molecule. ns, Not significant. Statistical analysis was performed with the Mann-Whitney U test. **E**, Correlation between adjusted density of protein expression and recombination activity of RAG1 mutants (Spearman $r_s = 0.351$, P = .023).

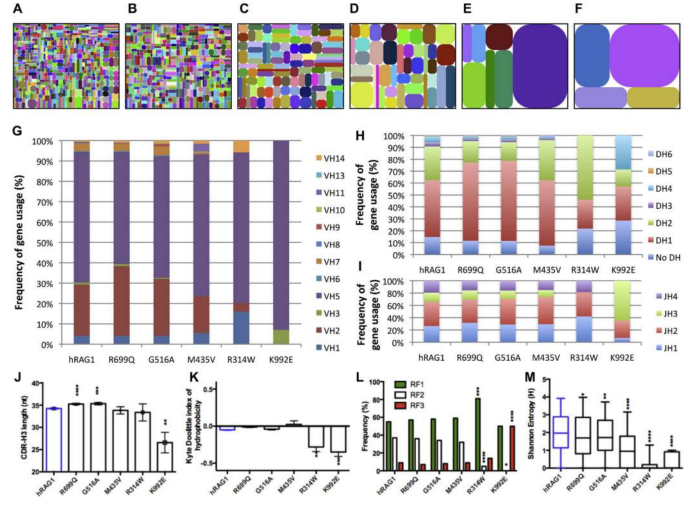


FIG 4.

Diversity and CDR-H3 characteristics and composition of the rearranged *Ighc* repertoire of h*RAG1* mutations. **A–F**, Tree maps (iRepertoire) were generated to depict graphically the diversity and frequency of different V–J pairings induced by various hRAG1 mutants: wild-type hRAG1 (Fig 4, *A*); R699Q (Fig 4, *B*); G516A (Fig 4, *C*); M435V (Fig 4, *D*); R314W (Fig 4, *E*); and K992E (Fig 4, F). Each *dot* represents a unique V_H-J_H recombination, and the size of the *dot* indicates the relative frequency of that specific V-J_H rearrangement. **G–I**, Bar diagrams representing the uses of V_H (Fig 4, *G*), D_H (Fig 4, *H*), and J_H (Fig 4, *I*) genes for wild-type hRAG1 and various hRAG1 mutants. For the V_H genes, only the genes that had positive values are included. **J–M**, Characterization of CDR-H3 sequences: average CDR-H3 length (± SEM; Fig 4, *J*); average index of hydrophobicity (± SEM) according to a normalized Kyte-Doolittle scale (Fig 4, *K*); differential use of 3 RFs (Fig 4, *L*); and average Shannon entropy index (± SEM) of the CDR-H3 loop (Fig 4, *M*). **P* < .05, ***P* < .01, ****P* < .001, and ******P* < .0001 unpaired 2-tailed *t* test.

TABLE 1

List of patients subdivided into 6 clinical groups

PID type	Patient no.	Mutation	Age	T cells/µL	B cells/µL	Eosinophils/µL	IgE (IU/µL)	Autoimmunity	Reference
ICL	1*	a. R474C; b. L506F	8 y	810	480	029	<5	No	16
CID-G/A	2*	a. R975W; b. R778Q	7.5 y	909	132	200	N Q	No	13
	3*	a. R314W; b. R507W/R737H	2.6 y	315	30	21	4.4	No	13
	*4	a. R975Q; b. R396C	4 y	865	51	NA	<30	Yes	24
	.5*	a. W522C; b. L541CfsX30	14 y	1404	552	NA	\$	Yes	14
	9	a. F974L; b. R841Q	0	34	256	N QN	38	Yes	15
	7*	a. G516A; b. S601P	2 y	50	94	2700	ND		
	8	a. R699Q; b. C730F	15 mo	187	51	2100	3.88	Yes	
	97	a. W522C; b. R973C	18 mo	3973	0	100	126	Yes	
	10	a. M435V; b. R699W	5 y	180	06	ND	<5	Yes	
	11	a. R474C; b. K86VfsX33	3 y	327	266	ND	<2	Yes	
	12	a. R841W; b. R841W	18 mo	407	44	0	1	Yes	
AS/LS	13	a. R975Q; b. R396L	4 mo	1350	432	1150	1690		4
	14	a. R841W; b. R410Q	1 mo	202	152	1200	>1000		4
	15	a. A444V; b. A444V	1 mo	822	6	1680	45		4
	16	a. V433M; b. A444V	2 mo	51	8	140	ND		4
	17	a. W896R; b. W896R	4.5 mo	1660	0	ND	<3		25
	$18^{#}$	a. E770K; b E770K	0	36	\$>	ND	ND		26
	19	a. H612R; b. H612R	7.5 y	245	0	33	<2		27
	20	a. R142X; b. unknown	12 mo	778	301	1045	>5000		
γδ-Τ	21	a. R841W; b. R841W	13 mo	4050	200	100	3.6	Yes	11
	22	a. Q981P; b. Q981P	12 mo	3900	2550	100	11	Yes	11
	2383	a. K86VfsX33; b. K86VfsX33	19 mo	1925	420	ND	4>	Yes	11
	24	a. E174SfsX27; b. E174SfsX27	3 mo	420	0	800	ND		11
	25	a. R561H; b. R561H	11 mo	350	24	420	20	Yes	12

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	26	a. R561H; b. R561H	11 mo	135	10	0	ND	Yes	12
	2781	a. K86VfsX33; b. K86VfsX33	13 mo	3400	320	ND	<10	Yes	7
	28	a. R474C; b. L732P	8 mo	294	63	ND QN	ND QN	Yes	26
	29	a. K86VfsX33; b. K86VfsX33	4 y	2175	1535	ND	ND	Yes	
	3083	a. K86VfsX33; b. K86VfsX33	0	135	44	009	ND	Yes	
	31	a. K86VfsX33; b. K86VfsX33	11 mo	1010	310	290	5.6	Yes	
	32	a. R108X; b. R108X	2 mo	2550	193	1400		Yes	
	33	a. E174SfsX27; b. E174SfsX27	2 mo	399	3	0	113	Yes	
SO	34	a. M435V; b. R559S	1 mo	5461	\$	59	ND QN		4
	35	a. Y912C; b. R396C	3 то	21580	<5	2250	>3000		4
	36	a. C328Y; b. C328Y	3 то	371	\$	10	19.9		4
	37	a. K86VfsX33; b. D429G	1.5 mo	96	9>	8	500		4
	38	a. R396C; b. R396C	4 mo	7544	\$	5150	182		4
	39	a. S401P; b. S401P	1.5 mo	2455	\$	1350	166		4
	40	a. A444V; b. A444V	3 mo	310	<>	1700	ND		4
	41	a. R624H; b. K992E	5 mo	1000	0	200	>1000		5
	42	a. R561H; b. R561H	2 mo	245	14	4550	190		12
	43	a. 1956T; b. 1956T	1 mo	12200	0	NA	NA		9
	4481	a. K86VfsX33; b. K86VfsX33	3 mo	820	0	ND	ND		7
	4581	a. K86VfsX33; b. K86VfsX33	18 mo	428	0	N	<10		7
	46	a. R142X; b. M458SfsX34	1 wk	10491	2	N ON	N Q		6
	47	a. L454Q; b. L454Q	NA	1584	26	1970	260		8
	48	a. R559S; b. M435V	3 wk	10755	0	5400	95		
	49	a. M435V/M1006V; b. M435V/M1006V	1 wk	18315	0	2600	ND		
	50	a. K992E; b. K992E	om 9	168	0	203	1		
	51	a. G99S/K391E; b. G99S/K391E	om 9	629	2	150	12		

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53 a. D5 54 a. Q2 55 a. R4 56 a. R5 57 a. R3 57 a. K3 77/B 60 a. M1 77/B 61 a. A4 77/B 63 a. R4	IFSX17 X	1 mo						
54 a. 55 a. 55 a. 56 a. 56 a. 58 a. 58 a. 59 a. 59 a. 60 a. 61 a. 61 a. 63 a. 63 a.	×	OIII I	2025	0	5020	501		
55 a 56 a 57 a a 58 a 58 a 60 a 6	7.)	NA	4792	0	ND	ND		
56 a. 57 a. 58 a. 58 a. 60 a. 60 a. 61 a. 62 a. 63 a. 63 a.		2 mo	36670	0	8640	Increased		
57 a. 58 a. 58 a. 60 a. 60 a. 61 a. 61 a. 63 a. 63 a.	K3393; b. P/86L	om 6	391	4	ND	310	Yes	
58 a. 59 a. 60 a. 61 a. 61 a. 63 a. 63 a.	R396H; b. R396H	2 mo	NA	NA	NA	NA		
59 a. W 60 a. M 61 a. A 62 a. A 63 a. R	K383RfsX7; b. K383RfsX7	om 9	205	31	390	ND		
60 a.N 61 a.A 62 a.A 63 a.R	W959X; b. W959X	3 wk	3424	0	11700	\$	No	
61 a.A 62 a.A 63 a.R	M1006V; b. M1006V	NA	NA	NA	NA	NA		28
62 a. A - 63 a. R	A444V; b. A444V	3 mo	64	0	ND	ND		
- 63 a.R	A444V; b. A444V	0	179	30	ND	ND		
	404Q/R449K; b. R764P	1 mo	28	<20	30	ND	No	
64 a. R3	R394Q; b. R394Q	NA	20	0	ND	ND		
65 a. T40	T403P; b. T403P	3 wk	0	0	350	ND		
66 a. V4	475AfsX17; b. V475AfsX17	2 mo	2	1	26	<2		
67 a.L4	a. L411P; b. L411P	1.5 y	25930	0	006	29464	No	
. 85 a. S6	S626X; b. S626X	2 mo	1836	0	ND	ND		

a, One allele; AS/LS, atypical/leaky SCID; b, the other allele; NA, not available; ND, not determined; PID, primary immunodeficiency.

^{*} Patients with granuloma.

 $^{^{\#}}$ Maternal engraftment.

[§]Siblings.