

Similar recombination-activating gene (*RAG*) mutations result in similar immunobiological effects but in different clinical phenotypes

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Background: V(D)J recombination takes place during lymphocyte development to generate a large repertoire of T- and B-cell receptors. Mutations in recombination-activating gene 1 (*RAG1*) and *RAG2* result in loss or reduction of V(D)J recombination. It is known that different mutations in *RAG* genes vary in residual recombinase activity and give rise to a broad spectrum of clinical phenotypes.

Objective: We sought to study the immunologic mechanisms causing the clinical spectrum of *RAG* deficiency.

Methods: We included 22 patients with similar *RAG1* mutations (c.519delT or c.368_369delAA) resulting in N-terminal truncated *RAG1* protein with residual recombination activity but presenting with different clinical phenotypes. We studied precursor B-cell development, immunoglobulin and T-cell receptor repertoire formation, receptor editing, and B- and T-cell numbers.

Results: Clinically, patients were divided into 3 main categories: T⁺B⁺ severe combined immunodeficiency, Omenn syndrome, and combined immunodeficiency. All patients showed a block in the precursor B-cell development, low B- and T-cell numbers, normal immunoglobulin gene use, limited B- and T-cell repertoires, and slightly impaired receptor editing.

Conclusion: This study demonstrates that similar *RAG* mutations can result in similar immunobiological effects but different clinical phenotypes, indicating that the level of residual recombinase activity is not the only determinant for clinical outcome. We postulate a model in which the type and moment of antigenic pressure affect the clinical phenotypes of these patients. (*J Allergy Clin Immunol* 2014;133:1124-33.)

Key words: *RAG* deficiency, V(D)J recombination, B- and T-cell receptor repertoire, receptor editing, autoimmunity, next generation sequencing, immune repertoire analysis

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Defects in V(D)J recombination result in a block in B- and T-cell differentiation because formation of immunoglobulin and T-cell receptors (TRs) is perturbed.¹ This results in a combined immunodeficiency (CID) of B and T cells. V(D)J recombination is initiated by the recombination-activating gene (*RAG*) 1 and *RAG2* proteins by creating double-stranded breaks in the immunoglobulin and TR loci. Subsequently, these breaks are processed and repaired by proteins involved in nonhomologous end joining. Thus far, genetic defects have been identified in the *RAG1*, *RAG2*, *Artemis*, ligase IV (*LIG4*), *XLF* (*Cernunnos*), and *DNA-PKcs* genes.²⁻⁸ The immunologic phenotypes and clinical presentations of these mutations are different, depending on the type of genetic defect (ie, null mutations or hypomorphic mutations with residual V(D)J recombination activity). Especially for the *RAG* genes, many different mutations have been described that give rise to residual activity of the mutated *RAG* protein.⁹ Different *RAG* mutations can result in a broad spectrum of clinical phenotypes, including severe combined immunodeficiency (SCID), *RAG* deficiency (*RAGD*) with skin inflammation and $\alpha\beta$ T-cell expansion (classical Omenn syndrome [OS]), *RAGD* with skin inflammation but without T-cell expansion (incomplete OS), *RAGD* with maternofetal transfusion, *RAGD* with $\gamma\delta$ T-cell expansion, late-onset SCID, *RAGD* with granulomas, and *RAGD* with CD4 cytopenia and thymus hypoplasia.^{9,10} This broad spectrum of clinical phenotypes impedes timely recognition of *RAGD* and might thus delay treatment (hematopoietic stem cell transplantation).

Abbreviations used

BM:	Bone marrow
CDR3:	Complementary determining region 3
CID:	Combined immunodeficiency
OS:	Omenn syndrome
PB:	Peripheral blood
RAG:	Recombination-activating gene
RAGD:	Recombination-activating gene deficiency
SCID:	Severe combined immunodeficiency
TR:	T-cell receptor
TRB:	T-cell receptor β
TREC:	T-cell receptor excision circle

In this study we selected 22 patients with RAGD with similar N-terminal truncating *RAG1* mutations to study the effect of a similar mutation on the clinical phenotype. These patients could be divided into 3 main clinical phenotypes (ie, SCID, OS, and CID, which includes the other phenotypes). We studied whether key immunologic parameters (eg, precursor B-cell development, B- and T-cell numbers, and B- and T-cell repertoire) might explain the differences in clinical phenotypes.

METHODS

Cell samples and flow cytometric immunophenotyping

Peripheral blood (PB), bone marrow (BM), and clinical data were obtained according to the guidelines of the Medical Ethics Committee of the Erasmus MC Rotterdam. Flow cytometric analysis was performed, as previously described.^{8,11,12}

RAG analysis and *in vitro* V(D)J recombination assay

The *RAG1* and *RAG2* genes were amplified by means of PCR and sequenced, as previously described.¹³ The level of recombination activity of the *RAG1* expression constructs was determined by using the recombination plasmid pDVG93, as described previously.^{10,13} A TaqMan-based realtime quantitative (RQ)-PCR was used to measure *RAG1* and *RAG2* transcription levels in BM mononuclear cells, as described previously.¹⁴

T-cell receptor β analysis

T-cell receptor β (*TRB*) gene rearrangements were studied, as described previously.¹⁵

Sequence analysis of *V κ* and *J κ* genes

V κ -C κ junctions were amplified in a multiplex PCR by using primers specific for *V κ 1-5* families (*V κ I*: 5'-GTAGGAGACAGAGTCACCATCACT-3', *V κ II*: 5'-TGGAGAGCCGGCTCCA-TCTC-3', *V κ III*: 5'-GGGAAAGAGCCACCCTCTCCTG-3', and *V κ IV*: 5'-GGCGAGAGGGCC-ACCATCAAC-3') and a *C κ* primer (5'-ACTTTGGCTCTCTGGATA-3'). PCR products were cloned in the pGEM-Teasy vector (Promega, Madison, Wis) and prepared for sequencing on the ABI Prism 3130 XL fluorescent sequencer (Applied Biosystems, Foster City, Calif). Obtained sequences were analyzed with the IMG T database (<http://imgt.cines.fr/>) to assign the *V κ* and *J κ* genes.^{16,17} The productive and unique sequences were used to determine the frequency of the *V κ* and *J κ* genes.

Repertoire analysis with next-generation sequencing

The VH-JH junctions were amplified from post-Ficoll PBMCs in a multiplex PCR by using the VH1-6 FR1 and JH consensus BIOMED-2

primers.¹⁵ The primers were adapted for 454 sequencing by adding the forward A or reverse B adaptor, the "TCAG" key, and the multiplex identifier adaptor. PCR products were purified by using gel extraction (Qiagen, Valencia, Calif) and Agencourt AMPure XP beads (Beckman Coulter, Fullerton, Calif). Subsequently, the PCR concentration was measured with the Quant-it Picogreen dsDNA assay (Invitrogen, Carlsbad, Calif). The purified PCR products were sequenced on the 454 GS junior instrument according to the manufacturer's recommendations by using the GS junior Titanium emPCR kit (Lib-A), sequencing kit, and PicoTiterPlate kit (454 Life Sciences; Roche, Branford, Conn). By using the CLC genomic workbench software, the samples were separated based on their multiplex identifier sequence and trimmed, and reads with a quality score of less than 0.05 and less than 250 bp were discarded. The reads were uploaded to IMG T HighV-Quest software.¹⁸ Subsequently, these output files were uploaded to the custom Galaxy platform.¹⁹⁻²¹ Further processing was done in the R programming language²² to generate the tabular and graphic outputs. The complementary determining region 3 (CDR3) amino acid patterns were visualized with WebLogo (<http://weblogo.berkeley.edu/>).^{23,24}

Statistics

Differences in absolute numbers of lymphocyte subsets were analyzed by using the 2-tailed *t* test for independent samples (*P* < .05 was considered significant) in GraphPad Prism software (GraphPad Software, La Jolla, Calif).

RESULTS

Residual RAG1 activity in patients with N-terminal truncating RAG1 mutations

Over the past 10 years, we identified one of the 2 mutations resulting in N-terminal truncating *RAG1* mutations in 22 patients (Tables I and II). These c.519delT (hereafter abbreviated as delT) and c.368_369delAA (hereafter abbreviated as delAA) mutations have been described before in several patients.^{13,25-29} They were found to be hypomorphic^{13,27} because translation can be reinitiated from the alternative start site methionine 202 (M202) or M183, resulting in an N-terminal truncated RAG1 protein¹³ with the same (comparable) residual RAG1 activity (<5% compared with wild type; Fig 1, A).¹³ Sixteen patients were homozygous for the delAA or delT mutation, and 6 patients were compound heterozygous (Table I). Three *RAG1* mutations found on the second allele were also analyzed in the *in vitro* recombination assay, showing no residual RAG1 activity (Fig 1, A). In addition, we determined the presence of polymorphisms in the *RAG1* gene because these might influence the recombination activity of RAG1. The only polymorphism found was p.Arg249His, which was shown not to affect recombination activity.²

N-terminal truncating RAG1 mutations result in a spectrum of clinical phenotypes

Although all patients had similar *RAG1* mutations, resulting in the same N-terminal truncation of the RAG1 protein, the clinical phenotypes varied substantially. The patients could be divided into 3 main clinical phenotypes: "classical" T⁻B⁻ SCID (n = 4), OS (n = 9), and CID (n = 9, Tables I and II). The patients with "classical" SCID were defined as having low B- and T-cell numbers and age at diagnosis before the first year of life. The patients with OS all had generalized and pronounced erythroderma. The patients with CID were given a diagnosis after the first year of life and had greater than 14% $\gamma\delta$ T cells or normal levels of T cells (P17 and P22). Despite the same N-terminal

TABLE I. Clinical data of patients with RAGD

	Onset of infections (mo)	Age at diagnosis (mo)	Infections	Respiratory tract infections	Autoimmunity/erythroderma	Hepatomegaly	Splenomegaly	Lymphadenopathy
SCID								
P1	3							
P2	6	6	BCG	No	ITP	No	No	No
P3	8	8		Pneumonia and upper airway infections				
P4	6	8	BCG					Mild
OS								
P5	0	0.5			Erythroderma	Yes	Yes	Yes
P6 ^a	0	0.5		Recurrent pneumonia	Erythroderma	Yes	No	Yes
P7 ^a	0	0.5	CMV	No	Erythroderma	Yes	Yes	Yes
P8	0	3.5	CMV, <i>Candida</i> species, MRSE	Severe pneumonia	Erythroderma			
P9	0	4			Erythroderma	Yes	No	Yes
P10	1	1			Erythroderma			Yes
P11	1.5	2			Erythroderma			Yes
P12	1	8	BCG	Recurrent pneumonia	Erythroderma	Yes	Yes	Yes
P13	3	6	<i>Candida</i> species, <i>Mycobacterium bovis</i> , coronavirus, rhinovirus	Recurrent upper and lower airway infections	Erythroderma, AIHA, ITP	Yes	No	No
CID								
P14 ^b	9	30	CMV, <i>Candida</i> species	Recurrent bronchopneumonia		Yes	Yes	No
P15 ^b	9	18	CMV	Recurrent bronchopneumonia		Yes	Yes	No
P16 ^c	1	11	CMV	Chronic rhinitis		No	No	No
P17	4	6	CMV, BCG	Pneumonia		Yes	Yes	Yes
P18	18	60	CMV, BCG, rhinovirus	Yes	AIHA, ITP	No	No	No
P19	3	13	<i>Candida</i> species	Chronic rhinitis and bronchitis	AIHA	No	No	No
P20	24	48			AIHA			
P21	13	60	<i>Candida</i> species, aspergillosis	Recurrent pneumonias, bronchitis	AIHA	No	No	No
P22 ^c	0	17		Recurrent pneumonias, bronchitis		No	No	No

Footnote symbols "a," "b," and "c" indicate relatives.

AIHA, Autoimmune hemolytic anemia; CMV, cytomegalovirus; ITP, idiopathic thrombocytopenic purpura; MRSE, methicillin-resistant staphylococcus epidermidis.

truncation of *RAG1* in the 22 patients, the range of clinical phenotypes strongly suggests that factors other than residual *RAG1* activity contribute to the clinical phenotype.

All clinical phenotypes show a block in precursor B-cell development

RAGD results in a block in the precursor B-cell differentiation in BM at the B-cell stages during which V(D)J recombination of the immunoglobulin genes takes place.¹¹ The relative distribution of pro-B, pre-BI, pre-BII, and immature B cells was assessed in BM from 11 of 22 patients to investigate precursor B-cell differentiation. In healthy children pro-B and pre-BI cells constitute 20% to 25% of the precursor B cells (Fig 1, B). All patients with "classical" SCID and OS, except P13, showed a complete block before the pre-BII-cell stage (Fig 1, B), whereas most of the patients with CID had a leaky block with greater than 10% pre-BII and immature B cells (Fig 1, B). *RAG1* and *RAG2* transcription levels were determined in the BM mononuclear cells to exclude that differences in *RAG1* transcription levels caused these difference in precursor B-cell composition. It is known that *RAG1* and *RAG2* transcription levels are correlated, and

that *RAG1* and *RAG2* levels in BM mononuclear cells depend on the number of cells expressing *RAG* (pre-BI and pre-BII cells).³⁰ In all 11 studied patients, the *RAG1* transcription level was correlated to *RAG2* (Fig 1, C), indicating that the differences in severity of the precursor B-cell block were not caused by differences in expression of *RAG1*. B-cell numbers in PB were undetectable or very low in most patients, except P15, who had normal levels (Table II). Correlating the percentage of pre-BII and immature cells in BM with the number of peripheral B cells showed that only patients with greater than 10% pre-BII and immature B cells in BM (P13, P14, P15, P16, and P17) had detectable B cells in PB. Collectively, these data indicate that most patients with CID have a milder block in the precursor B-cell composition and that only patients with a leaky block have detectable B cells in the PB.

Immunoglobulin heavy chain combinatorial repertoire

In those patients with detectable peripheral B cells, we studied the *IGH* V(D)J recombination repertoire. *IGH* gene rearrangements were amplified from mononuclear cells derived

TABLE II. Immunologic data of patients with RAGD

	delT	delAA	Other	CD3 ⁺ T cells, absolute (× 10E9/L)	CD4 ⁺ T cells, absolute (× 10E9/L)	CD8 ⁺ cells, absolute	CD45RA (%)	γδT cells (%)	CD19 ⁺ cells, absolute (× 10E9/L)	NK cells, absolute (× 10E9/L)
SCID										
	P1	Homozygous		0.06 (1.4-8.0)	0.04 (0.9-5.5)	0.01 (0.4-2.3)		21.8	0.03 (0.6-3.1)	0.08 (0.1-1.4)
	P2	Heterozygous	p.P874GfsX82	0.1 (2.4-6.9)	0.06 (1.4-5.1)	0.01 (0.6-2.2)	32.1	24.4	0.01 (0.7-2.5)	0.4 (0.1-1.0)
	P3	Homozygous		0.3 (1.6-6.7)	0.06 (1.0-4.6)	0.3 (0.4-2.1)			0 (0.6-2.7)	0.5 (0.2-1.2)
	P4	Heterozygous	p.R559S	0.3 (1.6-6.7)	0.2 (1.0-4.6)	0.04 (0.4-2.1)	7.7	10	0 (0.6-2.7)	0.1 (0.2-1.2)
OS										
	P5	Homozygous		20.1 (2.3-7.0)	7.56 (1.7-5.3)	12.75 (0.4-1.7)		4	0 (0.6-1.9)	2.59 (0.2-1.4)
	P6 ^a	Homozygous		3.7 (2.3-6.5)	3.1 (1.5-5.0)	0.4 (0.5-1.6)	7.2	5.2	0.03 (0.6-3.0)	0.8 (0.1-1.3)
	P7 ^a	Homozygous		36 (2.3-6.5)	10.7 (1.5-5.0)	24.9 (0.5-1.6)	3.1	1	0.03 (0.6-3.0)	0.4 (0.1-1.3)
	P8	Homozygous		3.93 (2.3-6.5)	1.45 (1.5-5.0)	2.19 (0.5-1.6)	21.4	24.4	0.02 (0.6-3.0)	0.88 (0.1-1.3)
	P9	Homozygous		1.84 (2.3-6.5)	1.48 (1.5-5.0)	0.3 (0.5-1.6)	4.6	3	0.004 (0.6-3.0)	1.64 (0.1-1.3)
	P10	Heterozygous	p.R737H	3.3 (2.3-7.0)	0.32 (1.7-5.3)	2.97 (0.4-1.7)	0.3	0.1	0 (0.6-1.9)	0.34 (0.2-1.4)
	P11	Heterozygous	p.R559S	4.33 (1.6-6.7)	4 (1.0-4.6)	0.27 (0.4-2.1)			0.01 (0.6-2.7)	0.93 (0.2-1.2)
	P12	Homozygous		2.21 (2.3-6.5)	1.34 (1.5-5.0)	0.61 (0.5-1.6)			0.07 (0.6-3.0)	0.56 (0.1-1.3)
	P13	Homozygous		0.6 (2.4-6.9)	0.6 (1.4-5.1)	0.01 (0.6-2.2)	4.8	2.9	0.07 (0.7-2.5)	0.2 (0.1-1.0)
CID										
	P14 ^b	Homozygous		0.3 (0.9-4.5)	0.1 (0.5-2.4)	0.07 (0.3-1.6)	35	49.5	0.4 (0.2-2.1)	0.8 (0.1-1.0)
	P15 ^b	Homozygous		0.5 (1.4-8.0)	0.1 (0.9-5.5)	0.2 (0.4-2.3)	38.9	64.1	0.4 (0.6-3.1)	2.9 (0.1-1.4)
	P16 ^c	Homozygous		0.16 (1.6-6.7)	0.07 (1.0-4.6)	0.02 (0.4-2.1)	26.9	46.3	0.09 (0.6-2.7)	0.32 (0.2-1.2)
	P17	Homozygous		2.7 (1.6-6.7)	0.2 (1.0-4.6)	1.5 (0.4-2.1)	90.7	90.2	0.06 (0.6-2.7)	0.7 (0.2-1.2)
	P18	Heterozygous	p.R759C	0.53 (0.9-4.5)	0.07 (0.5-2.4)	0.12 (0.3-1.6)		57.2	0.12 (0.2-2.1)	1.32 (0.1-1.0)
	P19	Homozygous		0.10 (1.6-6.7)	0.01 (1.0-4.6)	0.10 (0.4-2.1)		97.5	0.04 (0.6-2.7)	0.23 (0.2-1.2)
	P20	Homozygous		0.77 (0.9-4.5)	0.25 (0.5-2.4)	0.24 (0.3-1.6)		41.7	0.02* (0.2-2.1)	0.18 (0.1-1.0)
	P21	Heterozygous	p.A444V	0.12 (0.9-4.5)	0.10 (0.5-2.4)	0.06 (0.3-1.6)		14.4	0.001 (0.2-2.1)	0.25 (0.1-1.0)
	P22 ^c	Homozygous		1.97 (1.6-6.7)	0.42 (1.0-4.6)	1.50 (0.4-2.1)	42		0.29 (0.6-2.7)	0.72 (0.2-1.2)

Numbers in parentheses indicate normal values. Footnote symbols “a,” “b,” and “c” indicate relatives.

NK, Natural killer.

*Under rituximab treatment.

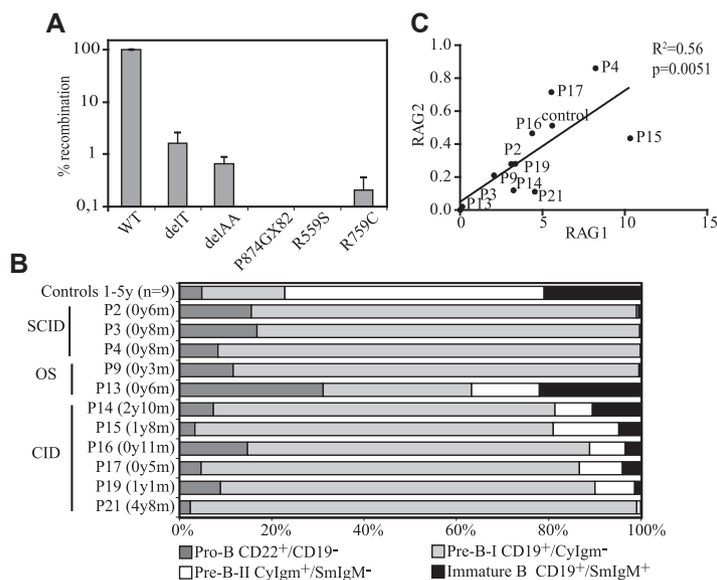


FIG 1. RAG expression and precursor B-cell compartment. **A**, Recombination activity of the c.519delT (delT), c.delA368/A369 (delAA), p.P874GX82, p.R559S, and p.R759C *RAG1* mutations was compared with wild-type (*WT*) *RAG1*. Only the delT and the delAA *RAG1* mutations result in low levels of residual recombination activity. **B**, Composition of the precursor B-cell compartment in control subjects (n = 9), 3 patients with the “classical” SCID phenotype, 2 patients with OS, and 6 patients with CID. **C**, Relative *RAG1* expression levels correlated to *RAG2* expression in all the analyzed RAG patients, as determined by using RQ-PCR.

from PB, BM, or both and subsequently sequenced by using next-generation sequencing in healthy control subjects (PB and BM) and 3 patients with CID (P15, P16, and P18). The frequency of

unique sequences in *IGH* genes was significantly lower in patients with RAGD than in control subjects (Table III), which is a reflection of the low numbers of B cells present in PB.

TABLE III. Number of *IGH* sequences

	All sequences	Unique sequences	Unproductive	Productive
Control BM	35,472	18,241 (51.4)	8,633 (24.3)	26,839 (75.7)
P16 BM	12,195	3,325 (27.3)	1,629 (13.3)	10,566 (86.7)
Control PB	19,294	9,185 (61.2)	4,030 (20.9)	15,003 (77.8)
P15 PB	16,826	7,706 (45.8)	1,047 (6.2)	15,779 (93.8)
P16 PB	14,572	3,763 (25.8)	896 (6.1)	13,676 (93.9)
P18 PB	25,100	3,730 (14.9)	1,488 (5.9)	23,612 (94.1)

Numbers in parentheses indicate percentages. "Unproductive" refers to out-of-frame rearrangements or rearrangements containing a stop codon in the CDR3 region.

Despite the low recombination activity, *IGHV*, *IGHD*, and *IGHJ* gene use was not restricted (Fig 2 and see Fig E1 in this article's Online Repository at www.jacionline.org). Forty-eight of the 57 *IGHV* genes used in control subjects were identified in the patients with RAGD because were the 25 *IGHD* genes and all 6 *IGHJ* genes. *IGHV*, *IGHD*, and *IGHJ* gene uses were similar to those seen in control subjects, although some genes were used with different frequencies (Fig 2 and see Fig E1). Most strikingly, JH6 use was lower whereas JH4 use was higher compared with that seen in control subjects. The patients with RAGD had a significantly lower frequency (5.9% to 6.2% vs 20.9% to 24.3% in control subjects) of unproductive rearrangements (Table III), as reported previously.³¹ Unproductive rearrangements were defined as out-of-frame rearrangements or rearrangements with a stop codon. Therefore even though the patients with RAGD had reduced V(D)J recombination, leading to a limited TR and immunoglobulin repertoire, the *IGH* gene use was similar to that seen in control subjects without preferential use of the proximal or distal genes.

Selection of B cells is slightly impaired

OS is characterized by autoimmune-like clinical features, including severe erythroderma, hepatosplenomegaly, and lymphadenopathy.^{32,33} The immune dysregulation in patients with OS might be caused by the severe abnormalities of thymic architecture and impaired expression of autoimmune regulator and tissue-specific antigens.^{34,35} In addition, hypomorphic Rag mouse models have shown a disturbance in B-cell tolerance.^{36,37} In addition, patients with OS and also 1 patient with "classical" SCID and 4 patients with CID had autoimmunity, and all displayed idiopathic thrombocytopenic purpura, autoimmune hemolytic anemia, or both (Table I). Unfortunately, the thymic architecture and autoimmune regulator and tissue-specific antigen expression could not be studied in our patients, but we were able to evaluate 3 parameters in the *IGH* sequences that are associated with autoimmunity. These are characterized by long CDR3s, and the frequency of *IGHV4-34*, which is known to encode intrinsically self-reactive cold agglutinin antibodies that recognize carbohydrate antigens on erythrocytes.^{38,39} The distribution of the CDR3 length of the unique junctions in BM and PB was similar to that seen in control subjects (Fig 3, A), except patient 18, who seemed to have increased numbers of junctions with a CDR3 of 16 and 22 amino acids. These junctions with a CDR3 length of 22 amino acids displayed high similarity (Fig 3, B). No sequence similarity was found when all 16-amino-acid CDR3s were compared (Fig 3, B), but 18.3% of these junctions used *IGHV6-1*, and all these junctions had a highly similar CDR3 sequence (Fig 3, B), which suggests that they might

recognize a common antigenic determinant. The frequency of long CDR3s (≥ 15 amino acids) was significantly lower in P15 and P16 ($P < .0001$) but not in P18 (Fig 3, C). The frequency of *IGHV4-34* use was significantly higher in P16 ($P < .0001$) and P18 ($P < .0001$; Fig 3, D). From the 3 patients we analyzed, P18 had autoimmunity, which was reflected by the high frequency of *IGHV4-34* use.

In addition to selection against long CDR3s, B-cell tolerance is also generated by receptor editing of self-reactive B cells. These self-reactive B cells are induced to express the RAG proteins and edit their receptor light chains through available upstream $V\kappa$ and downstream $J\kappa$ genes to change the affinity of their receptors. Therefore the $V\kappa$ - $J\kappa$ junctions were amplified from 5 patients with OS and 4 patients with CID. The *IGKV* gene use was not significantly different from that seen in control subjects (Fig 3, E), but less *IGKJ5* genes were used in the patients with RAGD (Fig 3, F). Therefore receptor editing seems partly affected, as deduced from the very low *IGKJ5* use.

Difference between clinical phenotypes in absolute numbers of T cells but not in T-cell repertoire

The hallmark of classical OS is an expansion of autologous T cells with an HLA-DR⁺CD45RO⁺ phenotype and an oligoclonal $\alpha\beta$ T-cell repertoire.⁴⁰ Consistent with this, most of the patients with OS had normal or increased CD3⁺ T-cell numbers; in addition, 2 patients with CID had normal numbers (P17 and P22), whereas all other patients had low absolute numbers of CD3⁺ T cells (Table II). Remarkably, many patients had high percentages ($>14\%$) of $\gamma\delta$ T cells, including 2 patients with "classical" SCID, 1 patient with OS, and 8 patients with CID (Table II). In addition, we determined the T-cell proliferation by determining the δ REC- ψ J α T-cell receptor excision circle (TREC)⁴¹ content per 50 ng of DNA in 3 patients with "classical" SCID, 7 patients with OS, and 5 patients with CID. In 11 patients TRECs were not detectable, and in the other 4 patients (P2, P5, P7, and P18), the number of TRECs/50 ng of DNA was less than 1 compared with 134 ± 75 TRECs/50 ng of DNA in control subjects ($n = 7$; age, 8 months to 11 years; data not shown), meaning that the T cells that were present in these patients showed extensive proliferation. Furthermore, the T-cell repertoire was determined by testing the *TRB* gene rearrangements in 2 patients with "classical" SCID, 3 patients with OS, and 2 patients with CID. In all patients the *TRB* repertoire was clearly restricted (Fig 4). Taken together, the T cells that were present in the patients with RAGD showed extensive proliferation and had a restricted TR repertoire.

DISCUSSION

Many different *RAG1* mutations have been reported to the *RAG* mutation database.⁴² Although most are null mutations, several have been described to result in residual recombinase activity.^{11,13,25,27,43} Previously, it was hypothesized that null mutations in *RAG1* would result in "classical" T⁻B⁻ SCID and that partial reduction of RAG activity would result in OS or an intermediate late-onset SCID or OS phenotype.²⁸ Over the last few years, the spectrum of reported clinical phenotypes of RAGD has broadened and now also includes RAGD with $\gamma\delta$ T-cell expansion, RAGD with skin inflammation but without T-cell expansion (incomplete OS), RAGD with granulomas, RAGD

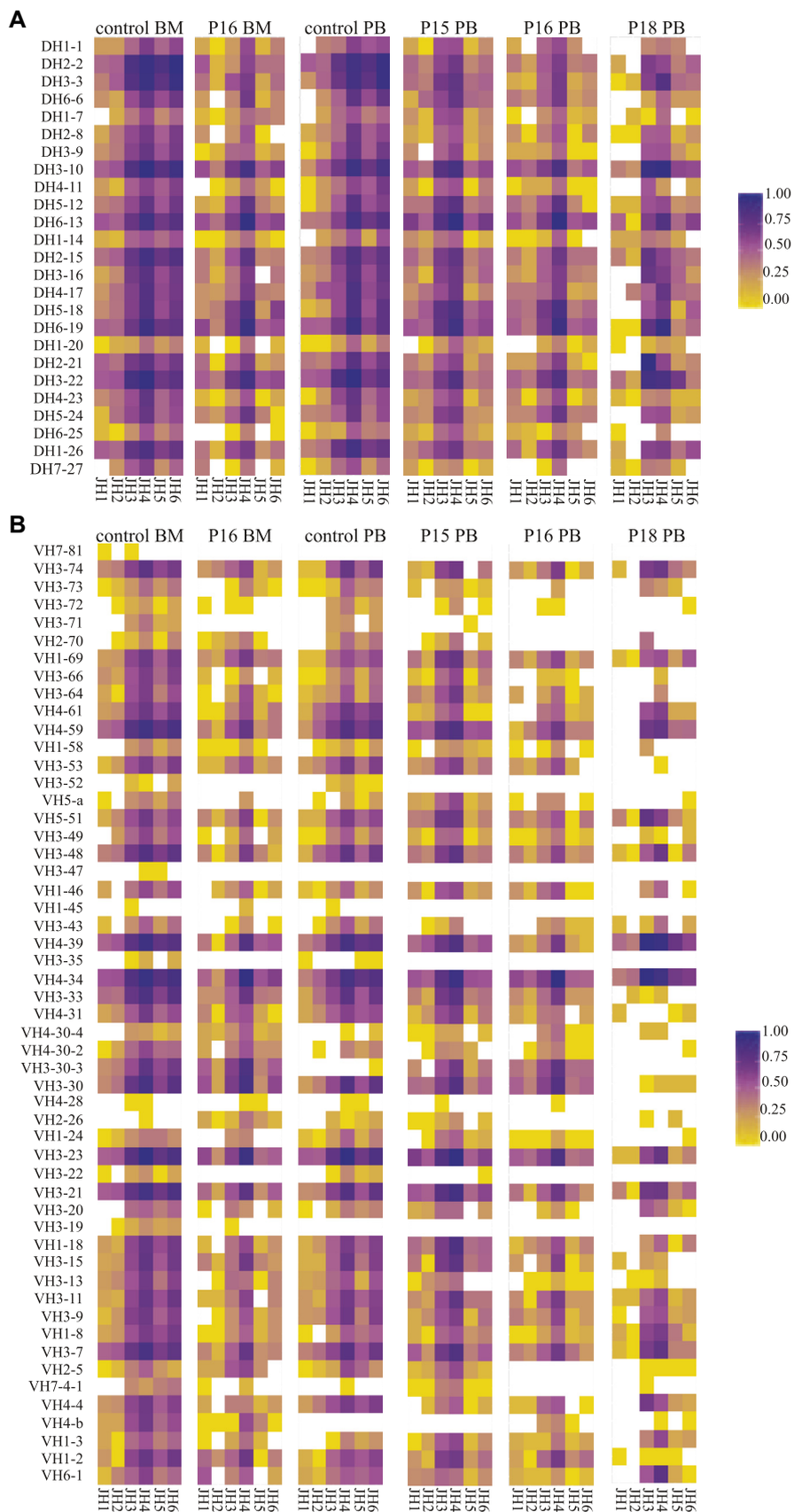


FIG 2. Heat maps of the different combinations of immunoglobulin DH-JH (A) and VH-JH (B), as determined in the unique junctions (defined by the unique combination of VH, DH, JH, and nucleotide sequences of CDR3).

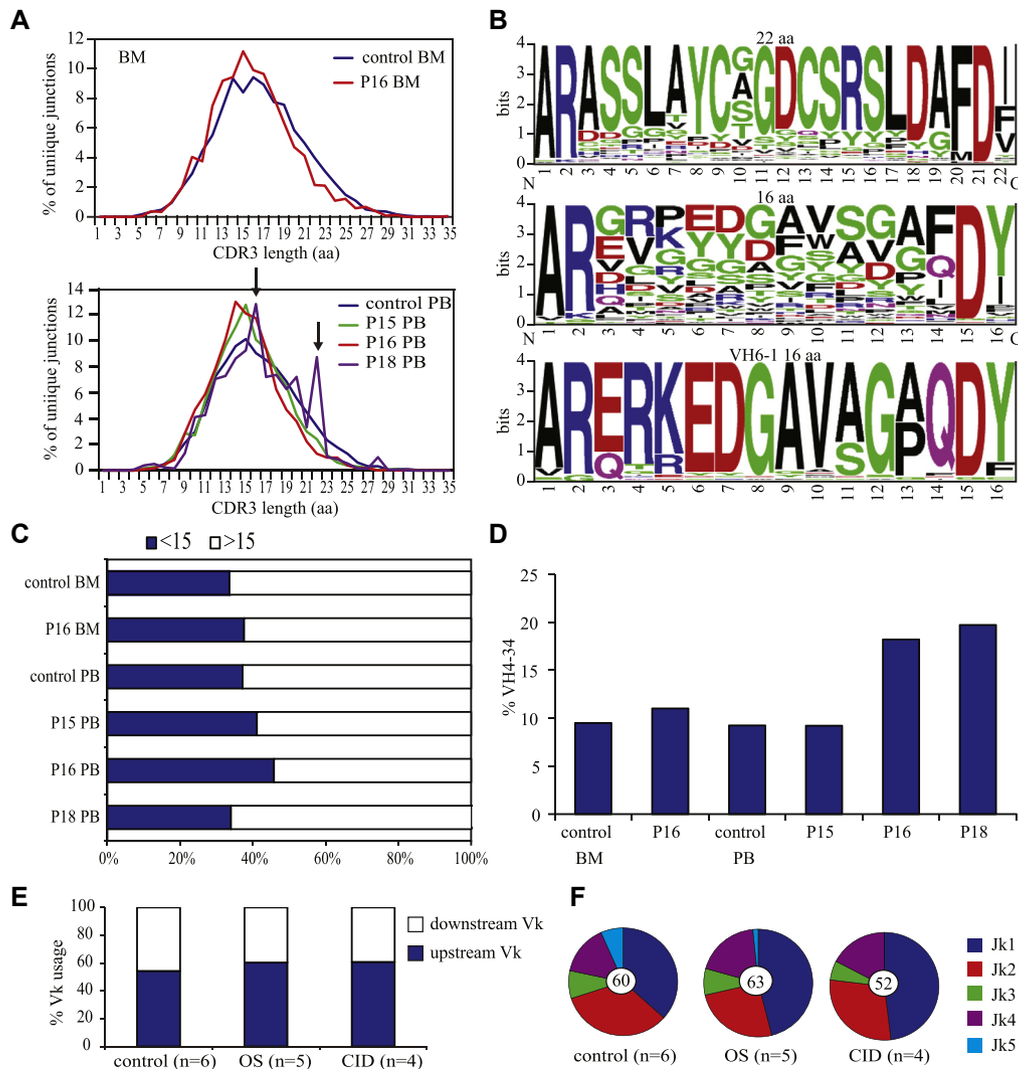


FIG 3. Functional characteristics of *IGH* junctions. **A**, Functional characteristics of the *IGH* junctions were determined in 3 patients with RAGD in PB or BM. Distribution of CDR3 length frequencies in BM and PB was similar in control subjects and patients with RAGD; however, P18 had increased numbers of junctions, with a CDR3 length of 16 and 23 amino acids. **B**, Sequence logo showed no similarity of the 16-amino-acid CDR3s of P23 but high similarity of CDR3s of 16 amino acids using the *IGHV6-1* gene and the 22-amino-acid CDR3s. **C**, The frequency of long CDR3s (≥ 15 amino acids) was decreased in P15 and P16. **D**, *IGHV4-34* use was increased in P16 and P18. **E** and **F**, The percentage of *IGKV* and *IGKJ* genes was determined in 6 control subjects, 5 patients with OS, and 4 patients with CID. *IGKV* use was normal (Fig 3, **E**), but hardly any *IGKJ5* gene was used (Fig 3, **F**).

with maternofetal transfusion, and RAGD with CD4 cytopenia and thymus hypoplasia.^{9,10} A few case reports have shown that the same *RAG* mutation can result in a different clinical phenotype.^{25,29,44,45} This study is the first to report an in-depth immunobiological evaluation of 22 patients with RAGD with similar *RAG1* mutations, resulting in the same N-terminal truncation of the *RAG1* protein. These similar mutations result in 3 different clinical phenotypes, which indicates that a specific mutation does not predict a patient's clinical phenotype.

Because all patients had similar mutations, the residual *RAG1* protein activity was expected to be comparable among all patients. The N-terminally truncated *RAG1* protein is produced through translation starting from an alternative start site (M183 or M202), and hence the amount of protein is dependent on how efficiently these start sites are used. Because the *RAG1*

transcription level correlated with that of *RAG2*, we assume that all patients had similar expression of the mutant *RAG1* protein (Fig 1, **C**). We cannot exclude that epigenetics and modifier genes accounted for small differences in *RAG1* protein expression. Although a previous attempt to identify such modifier genes in human subjects was not successful,⁴⁶ studies in mouse models could shed more light on the contribution of epigenetics and modifier genes.

In our cohort V(D)J recombination was not completely abolished but was strongly reduced because of the low residual activity of the *RAG1* protein. Reduced V(D)J recombination was characterized by normal *IGHV*, *IGHD*, and *IGHJ* gene use, without preferential use of proximal or distal genes. However, as shown previously,³¹ the frequency of unproductive sequences was significantly lower than in healthy control subjects,

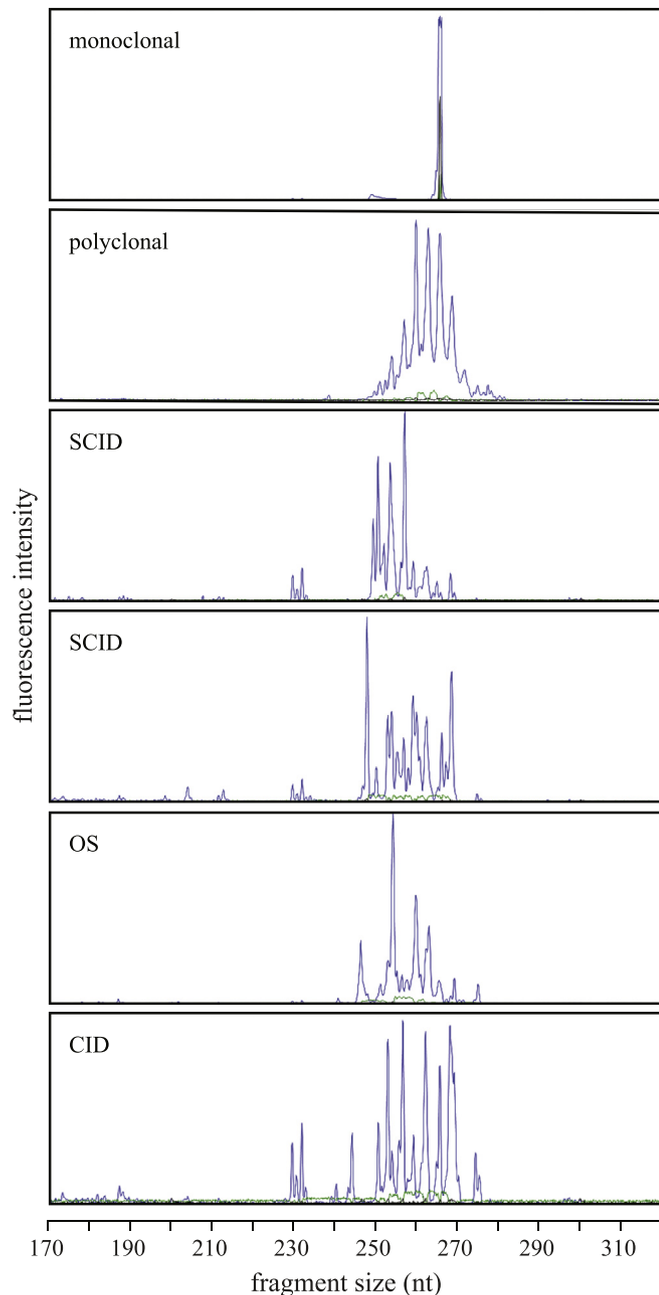


FIG 4. TRB repertoire. TR spectratyping profiles of *TRB* gene rearrangements using the BIOMED-2 TRB tube B. The upper 2 panels show the monoclonal and the polyclonal controls. The patient panels SCID (P2 and P4), OS (P8), and CID (P21) show a restricted TRB repertoire.

indicating that the B cells in the patients with RAGD did not correct unproductive rearrangements by means of recombination of the second *IGH* allele.

As a consequence of the reduced V(D)J recombination, fewer B and T cells with a functional receptor can be produced. The proliferation of the lymphocytes is increased to compensate for low circulating B- and T-cell numbers. This idea is corroborated by the low numbers of TRECs in patients with RAGD. The increased proliferation of T cells might result in normal or increased T-cell counts, especially in the patients with OS; however, the corresponding TR repertoire in all the patients with RAGD remains restricted.

Most patients with RAGD showed clinical signs of immune dysregulation, such as erythroderma, lymphadenopathy, hepatosplenomegaly, idiopathic thrombocytopenic purpura, and autoimmune hemolytic anemia. B cells have been shown to contribute to the immune dysregulation seen in *Rag* mouse models.^{36,37} Sera from these mice contained high-affinity anti-double-stranded DNA and tissue-specific autoantibodies, and B cells displayed impaired receptor editing. In addition, these mice had increased serum B cell-activating factor levels, which might rescue autoreactive B-cell clones. This increase in serum B cell-activating factor levels was also seen in patients with RAG-, Artemis-, and X-linked SCID.³⁷ Similar to observations in mice, most patients with RAGD did not use the *IGKJ5* gene, whereas *IGKV* gene use was normal. This suggests that receptor editing in this group of patients with RAGD was slightly impaired, which can either be a result of reduced recombination activity caused by the *RAG1* mutation or by low B-cell numbers leading to reduced selection against autoreactive B cells. The *IGH* repertoire was investigated for long CDR3s and increased *IGHV4-34* use, which are associated with autoreactive antibodies.^{47,48} From the 3 patients with RAGD we analyzed, only P18 had autoimmunity, which was reflected by an increased *VH4-34* gene use.

The patients divided into the 3 main clinical RAGD groups hardly differed in their immunobiological parameters, and consequently, we could not find any specific pattern that could explain the different clinical phenotypes. On the basis of our results and earlier reported data, we propose an explanatory model for the development of different clinical phenotypes in patients with RAGD with similar mutations (Fig 5). If RAGD results in reduced V(D)J recombination, low B- and T-cell numbers are produced with some (compensatory) clonal expansion. This expansion might increase the B- and T-cell numbers to even normal levels but does not change the limited repertoire. In such limited repertoire the selection against autoreactive cells is impaired. Provided the deficient immune system is not activated, patients with RAGD are asymptomatic. However, when the immune system will be activated by potentially a wide range of different (auto)antigens, the type of antigen and activated effector lymphocyte will have important consequences for the clinical phenotype. In addition, the impaired negative and positive selection of thymic lymphocytes and reduced number of regulatory T cells might result in autoimmunity when patients are exposed to autoantigens. This phenomenon can occur at any early stage, even *in utero*, as shown by the fact that patients with OS can have severe erythroderma already at birth, which is unlikely to be triggered by infections. Additionally, directly after birth, the skin and gastrointestinal tract become colonized by commensal bacteria, which can trigger the chronic diarrhea seen in most patients with RAGD. Key steps in the development of a certain clinical phenotype will be the B- and T-cell repertoire, the type of (auto)antigen exposure, the specificity of the antigen receptors and timing, the cell type involved in the immune activation, and the potential influence of genetic variations in modifier genes. Variability in any of these factors might eventually lead to different clinical phenotypes, despite a similar genetic defect.

In conclusion, this study clearly shows that the type of *RAG1* mutation and the level of residual *RAG1* recombinase activity are not the only determinants predicting the clinical phenotype, as previously assumed. The clinical outcome of an individual

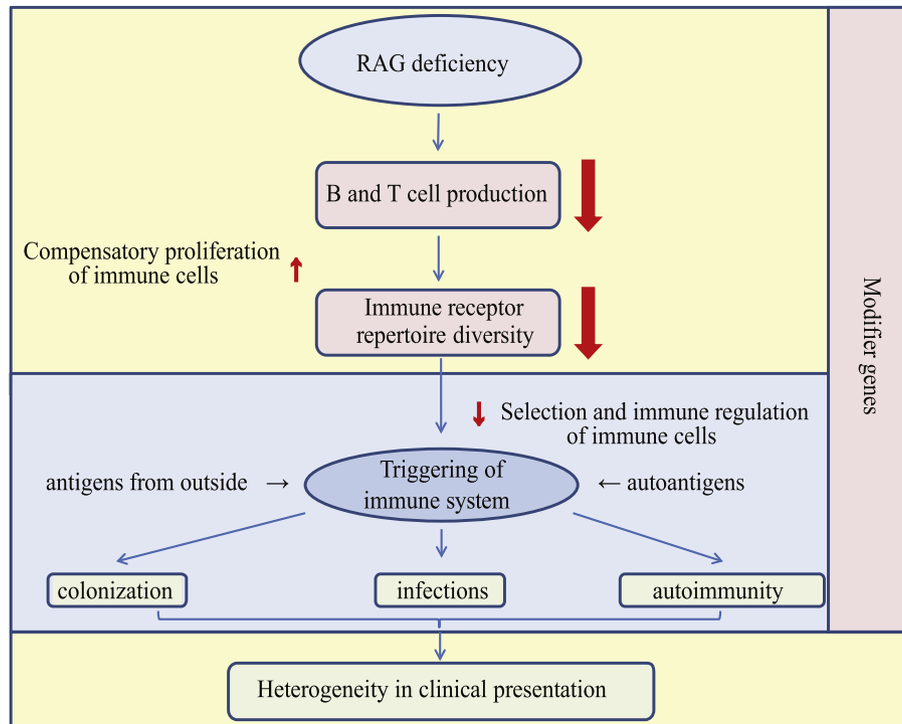


FIG 5. Model for development of clinical phenotype in patients with RAGD. RAGD results in reduced V(D)J recombination, leading to fewer B and T cells with a limited repertoire. In an attempt to compensate for low numbers, B and T cells start to proliferate, but the repertoire remains limited and imbalanced, so that selection and immune regulation are impaired. Most likely the type of antigenic stimulation together with the incomplete and imbalanced repertoire that has been developed will affect the eventual clinical phenotype with immune dysregulation problems.

patient with RAGD depends on a complex interplay between the (limited) immune receptor repertoire, (auto)antigen exposure, the specificity of antigen receptors, and the timing and cell type involved in immune activation. Therefore the clinical outcome of patients with RAGD with similar mutations is extremely difficult to predict.

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Clinical implications: RAGD can result in a broad spectrum of clinical presentations, but the level of residual RAG activity is not always predictive for the clinical outcome.

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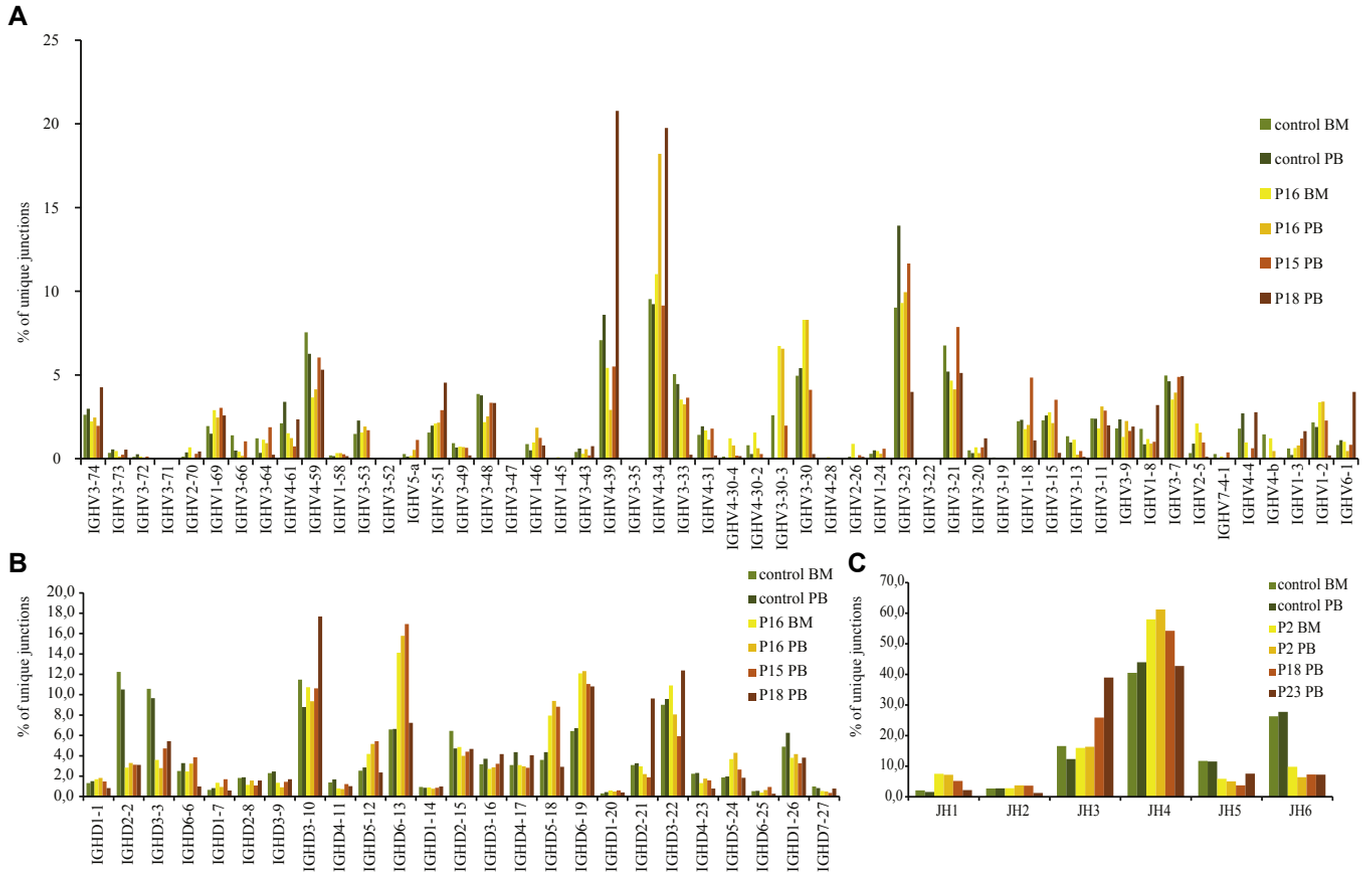


FIG E1. Immunoglobulin heavy chain gene usage. Frequency of *IGHV* (A), *IGHD* (B), and *IGHJ* (C) gene use in control BM and PB and in P16 BM and PB, P15 PB, and P18 PB.