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Loss-of-function nuclear factor κB subunit 1 (NFKB1) variants are the most common monogenic cause of common variable immunodeficiency in Europeans

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GRAPHICAL ABSTRACT

Whole Genome Sequencing:
Heterozygous pathogenic NFKB1 variants in 4% of European CVID cohort (n = 390)

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Background: The genetic cause of primary immunodeficiency disease (PID) carries prognostic information.

Objective: We conducted a whole-genome sequencing study assessing a large proportion of the NIHR BioResource–Rare Diseases cohort.

Methods: In the predominantly European study population of principally sporadic unrelated PID cases (n = 846), a novel Bayesian method identified nuclear factor κB subunit 1 (NFKB1) as one of the genes most strongly associated with PID, and the association was explained by 16 novel heterozygous truncating, missense, and gene deletion variants. This accounted for 4% of common variable immunodeficiency (CVID) cases (n = 390) in the cohort. Amino acid substitutions predicted to be pathogenic were assessed by means of analysis of structural protein data. Immunophenotyping, immunoblotting, and ex vivo stimulation of lymphocytes determined the functional effects of these variants. Detailed clinical and pedigree information was collected for genotype-phenotype cosegregation analyses.

Results: Both sporadic and familial cases demonstrated evidence of the noninfective complications of CVID, including massive lymphadenopathy (24%), unexplained splenomegaly (48%), and autoimmune disease (48%), features prior studies correlated with worse clinical prognosis. Although partial penetrance of clinical symptoms was noted in certain pedigrees, all carriers have a deficiency in B-lymphocyte differentiation.

Detailed assessment of B-lymphocyte numbers, phenotype, and function identifies the presence of an increased CD21 B-cell population. Combined with identification of the disease-causing variant, this distinguishes between healthy subjects, asymptomatic carriers, and clinically affected cases.

Conclusion: We show that heterozygous loss-of-function variants in NFKB1 are the most common known monogenic cause of CVID, which results in a temporally progressive defect in the formation of immunoglobulin-producing B cells. (J Allergy Clin Immunol 2018;nnn:nnn-nnn.)

Key words: B cells, common variable immunodeficiency, nuclear factor κB

Common variable immunodeficiency (CVID; MIM 607594), which occurs in approximately 1:25,000 persons, is a clinically and genetically heterogeneous disorder characterized by susceptibility to sinopulmonary infections, hypogammaglobulinemia, and poor vaccine responses. CVID is the most common primary immune deficiency, requiring lifelong clinical follow-up, and the clinical course is highly variable, with substantial excess mortality. Affected subjects present frequently with recurrent respiratory tract infections, as well as immune dysregulatory features. The antibody deficiency is often not as marked as the agammaglobulinemia seen in patients with genetically defined conditions leading to B-lymphocyte aplasia. Conversely, although patients with B-lymphocyte aplasia have a favorable prognosis on adequate replacement immunoglobulin treatment, the response of patients with CVID is highly variable.

Past studies focused on familial cases with CVID and used techniques ranging from traditional linkage analysis to more recent exome sequencing to characterize the genetic cause. This has revealed that monogenic gene dysfunction accounts for 10% of cases. Several of the variants in these genes have been characterized as partially penetrant; it remains unclear whether genetic or environmental factors determine disease onset. Multiple recent studies identified variants in nuclear factor κB subunit 1 (NFKB1) as a monogenic cause of CVID and reported on the clinical features of these cases.

As part of this NIHR BioResource–Rare Diseases study, we sequenced the genomes of 846 unrelated patients with predominantly sporadic primary immunodeficiency disease (PID) who were recruited from across the United Kingdom and by European collaborators. Application of the recently developed statistical method BeviMed to the 846 PID cases and more than 5000 control genomes identified NFKB1 as the gene with the highest probability of association with the disease and with the largest number of cases explained by variants in that gene. Further investigations revealed a series of 16 heterozygous loss-of-function (LOF) variants in NFKB1 as the most common genetic cause of CVID.

Mutations in genes that affect nuclear factor κB (NF-κB)–dependent signaling are associated with a number of immunodeficiencies. NF-κB is a ubiquitous transcription factor member of the Rel proto-oncogene family. NF-κB regulates the expression of several genes involved in inflammatory and immune responses. The classical activated form of NF-κB consists of a heterodimer of the p50/p65 protein subunits. The NF-κB family of transcription factors comprises 5 related proteins, c-Rel, p65 (RelA), RelB, p50 (NF-κB1) and p52 (NF-κB2), which interact to form homodimers and heterodimers with distinct gene regulatory functions.

Each Rel NF-κB protein has a conserved 300-amino-acid N-terminal Rel homology domain (RHD) that encompasses sequences needed by NF-κB proteins to bind DNA motifs (κB elements), form dimers, interact with regulatory inhibitor IκB proteins, and enter the nucleus. The 10 different NF-κB dimers identified have distinct transcriptional properties. In most cells NF-κB is retained in the cytosol in a latent state through interaction with the IκB proteins (e.g., IκBα, IκBβ, and IκBε), a family of proteins with ankyrin repeats that mediate IκB interaction with the RHD of NF-κB, masking the nuclear localization sequence and DNA-binding domains. Signal-dependent activation of an IκB kinase complex comprising catalytic (α and B) and regulatory (NF-κB essential modulator) subunits induces the phosphorylation and degradation of IκB, which permits NF-κB factors to enter the nucleus and regulate gene expression.

We show that variants in NFKB1 culminate in a progressive humoral immunodeficiency indistinguishable from CVID, with a highly variable penetrance. We demonstrate the utility of an in silico protein prediction model for validating the predicted
disease-causing substitutions, and we report on the clinical spectrum, immunologic phenotype, and functional consequences of heterozygous NFKB1 variants.

METHODS

Cohort

The NIHR BioResource–Rare Diseases (NIHRBR-RD) study was established in the United Kingdom to further the clinical management of patients with rare diseases by providing a national resource of whole-genome sequence data. All participants provided written informed consent, and the study was approved by the East of England–Cambridge South national institutional review board (13/EE/0325). At the time of our analysis, the NIHRBR-RD study included whole-genome sequence data from 8066 subjects, of whom 1299 were part of the PID cohort. These were predominantly singleton cases, but additional affected and/or unaffected family members of some of the patients were also sequenced; in total, there were 846 unrelated index cases.

Patients with PIDs were recruited by specialists in clinical immunology (either trained in pediatrics or internal medicine) from 26 hospitals in the United Kingdom and a smaller number came from The Netherlands, France, and Italy. The recruitment criteria included the following: clinical diagnosis of CVID according to the European Society for Immunodeficiencies ESID criteria (ESID Registry–Working Definitions for Clinical Diagnosis of PID, 2014, latest version: April 25, 2017), extreme autoimmunity, or recurrent (and/or unusual) infections suggestive of severely defective innate or cell-mediated immunity. Exclusion of known causes of PID was encouraged, and some of the patients were screened for 1 or more PID genes before enrollment in the PID cohort. The ethnic makeup of the study cohort represented that of the general United Kingdom population: 82% were European, 6% were Asian, 2% were African, and 10% were of mixed ethnicity based on the patients’ whole-genome data.

Given that PID is a heterogeneous disease, with overlap in phenotypes and genetic causes across different diagnostic categories, we decided to perform an unbiased genetic analysis of all 846 unrelated index cases. Whole-genome sequence data were additionally available for 63 affected and 345 unaffected relatives. Within a broad range of phenotypes, CVID is the most common disease category, comprising 46% of the NIHRBR-RD PID cohort (n = 390 index cases; range, 0-93 years of age).

Sequencing and variant filtering

Whole-genome sequencing of paired-end reads was performed by Illumina on their HiSeq X Ten system (Illumina, San Diego, Calif). Reads of 100, 125, or 150 bp in length were aligned to the GRCh37 genome build by using the Isaac aligner, variants across the samples were jointly called with the AGG tool, and large deletions were identified by using Canvas and Manta algorithms (all software by Illumina), as described previously. Average read depth was 35, with 95% of the genome covered by at least 20 reads.

Single nucleotide variants and small insertions/deletions were filtered based on the following criteria: passing standard Illumina quality filters in greater than 80% of the genomes sequenced by the NIHRBR-RD study, having a variant effect predictor (VEP) effect of either moderate or high, having a frequency of less than 0.01 in the NIHRBR-RD genomes were included in the analysis.

Gene and variant pathogenicity estimation

To evaluate genes for their association with PID, we applied the BeviMed inference procedure to the NIHRBR-RD whole-genome data set. BeviMed (https://CRAN.R-project.org/package=BeviMed) evaluates the evidence for association between case/control status of unrelated subjects and allele counts at rare variant sites in a given locus. The method infers the posterior probabilities of association under dominant and recessive inheritance and, conditional on such an association, the posterior probability of pathogenicity of each considered variant in the locus. BeviMed was applied to rare variants and large rare deletions in each gene, treating the 846 unrelated PID index cases as cases and the 5097 unrelated subjects from the rest of the NIHRBR-RD cohort as control subjects. All genes were assigned the same prior probability of association with the disease of .01, regardless of their previously published

FIG 1. Overall BeviMed results showing that NFKB1 has the highest posterior probability of association with disease in the NIHRBR-RD PID cohort. Genes with variants previously reported to cause PIDs are highlighted in red. Genes with posterior probabilities of greater than .05 are shown.
FIG 2. Plot of rare missense, truncating, and gene deletion NFKB1 variants identified in the NIHRBR-RD genomes of unrelated subjects and their location relative to NFKB1 domains. Tracks from left to right show the following: number of unrelated case (red) and control (black) subjects in whom each variant was observed; the 4 major NFKB1 domains; each exon in transcript ENST00000226574 (gray bars); variant annotation relative to transcript ENST00000226574 and genomic location of large deletions, with VEP high-effect variants and large deletions highlighted in blue; Combined Annotation Dependent Depletion (CADD) scores of all variants with pathogenic probabilities greater than 0.2. Variants are color-coded by CADD score, with high-effect variants highlighted in blue. The figure also includes a table with variant annotations and their positions relative to the NFKB1 gene.
associations with an immune deficiency phenotype. Variants with a VEP effect labeled as high were assigned higher prior probabilities of pathogenicity than variants with a moderate effect, as described previously.\textsuperscript{12}

**Immunophenotyping and B- and T-cell functional assays**

PBMCs were isolated by using standard density gradient centrifugation techniques with Lymphoprep (Nycomed, Oslo, Norway). Absolute numbers of lymphocytes, T cells, B cells, and natural killer cells were determined with Multitest 6-color reagents (BD Biosciences, San Jose, Calif), according to the manufacturer’s instructions. For PBMC immunophenotyping, we refer to the Methods section in this article’s Online Repository at www.jacionline.org.

PBMCs were resuspended in PBS at a concentration of 5 to 10×10^6 cells/mL and labeled with 0.5 μmol/L carboxyfluorescein succinimidyl ester (CFSE; Molecular Probes, Eugene, Ore), as described previously\textsuperscript{33} and in the Methods section in this article’s Online Repository, to analyze the ex vivo activation of T and B cells. Proliferation of B and T cells was assessed by measuring CFSE dilution in combination with the same mAbs used for immunophenotyping. Analysis of cells was performed with a FACSCanto II flow cytometer (BD Biosciences) and FlowJo software (TreeStar, Ashland, Ore). Patient samples were analyzed simultaneously with PBMCs from healthy control subjects.

**ELISA**

Secretion of immunoglobulins by mature B cells was assessed by testing supernatants for secreted IgM, IgG, and IgA with an in-house ELISA using polyclonal rabbit anti-human IgM, IgG, and IgA reagents and a serum protein calibrator, all from Dako (Glostrup, Denmark), as described previously.\textsuperscript{33}

**SDS-PAGE and Western blot analysis**

Blood was separated into neutrophils and PBMCs. Neutrophils (5×10^8) were used for protein lysates, separated by means of SDS-PAGE, and transferred onto a nitrocellulose membrane. Individual proteins were detected with antibodies against NF-κB p50 (mouse mAb E-10; Santa Cruz Biotechnology, Dallas, Tex), against IκBα (rabbit polyclonal antiserum C-21; Santa Cruz Biotechnology), and against human glyceraldehyde-3-phosphate dehydrogenase (mouse mAb; Merck Millipore, Darmstadt, Germany).

Secondary antibodies were either goat anti-mouse-IgG IRDye 800CW, goat anti-rabbit IgG IRDye 680CW or goat anti-mouse IgG IRDye 680LT (LI-COR Biosciences, Lincoln, Neb). Relative fluorescence quantification of bound secondary antibodies was performed on an Odyssey Infrared Imaging system (LI-COR Biosciences) and normalized to glyceraldehyde-3-phosphate dehydrogenase.

**NF-κB1 protein structure**

A previously resolved crystal structure of the p50 homodimer (A43-K353) bound to DNA was used to gain structural information on the NF-κB1 RHD.\textsuperscript{34} Ankyrin repeats of NF-κB1 (Q498-D802) were modeled by using comparative homology modeling (Modeller 9v16) with the ankyrin repeats crystal structure of NF-κB2 as a template.\textsuperscript{35,36} There is no structural information on the region between the sixth and seventh ankyrin repeats (F751-V771),\textsuperscript{37} and therefore these were omitted in the model.

**Statistical analysis of lymphocyte data**

Differences between groups with 1 variable were calculated with a nonpaired Student t test or 1-way ANOVA with the Bonferroni post hoc test, differences between groups with 2 or more variables were calculated with 2-way ANOVA with the Bonferroni post hoc test by using GraphPad Prism 6 software (GraphPad Software, La Jolla, Calif). A P value of less than .05 was considered significant.

**RESULTS**

**Pathogenic variants in NFKB1 are the most common monogenic cause of CVID**

In an unbiased approach to analysis, we obtained BeviMed posterior probabilities of association with PID for every individual gene in all 848 unrelated patients with PID in the NIHRBR-RD study. Genes with posterior probabilities of greater than .05 are shown in Fig 1, showing that NFKB1 has the strongest prediction of association with disease status (1.000). All 13 high-effect variants (large deletion, nonsense, frameshift, and splice site variants) in NFKB1 were observed in cases only, resulting in the very high posterior probabilities of pathogenicity (mean, 0.99) for this class of variants (Fig 2). On the other hand, moderate-effect variants (missense substitutions) were observed both in cases and control subjects. The majority had near-zero probability of pathogenicity, but 3 substitutions were observed in the patients with PID only and had posterior probabilities of greater than 0.15 (Fig 2), suggesting their potential involvement in the disease. Genomic variants with a high Combined Annotation Dependent Depletion score were found within both the PID and control cohorts, suggesting that this commonly used metric of variant deleteriousness cannot reliably distinguish disease-causing from benign variants in NFKB1. All 16 predicted likely pathogenic variants were private to the PID cohort, and further investigation revealed that all 16 subjects were within the diagnostic criteria of CVID (Table I).

Assessment of all 390 CVID cases in our cohort for pathogenic variants showed that the next most commonly implicated genes are NFKB2 and Bruton tyrosine kinase (BTK), with 3 explained cases each (see Fig E2 in this article’s Online Repository at www.jacionline.org). Importantly, based on the gnomAD data set of 135,000 predominantly healthy subjects, none of the NFKB1 variants reported here are observed in a single gnomAD subject, even though 90% of our CVID cohort and all of the NFKB1-positive cases had European ancestry. Therefore our results suggest that LOF variants in NFKB1 are the most commonly identified monogenic cause of CVID in the European population, with 16 of 390 patients with CVID explaining up to 4.1% of our cohort. None of the variants

nonsense, frameshift, splice and missense variants; Exome Aggregation Consortium (ExAC) allele frequencies; and conditional probability of variant pathogenicity inferred by using BeviMed. Only variants labeled as being of moderate or high effect relative to the canonical transcript ENST00000226574 are shown. The initial inference that formed part of the genome-wide analysis included variant chr4:1034232325G>A, which was observed in 1 control sample. This variant is intronic (low effect) relative to ENST0000022674 but is a splice variant (high effect) relative to the minor transcript ENST00000050648. Because variants were filtered based on the highest-effect variant annotation against any Ensembl transcript, this variant was originally included in the inference. For this plot, the inference was rerun, including only missense, truncating, and gene deletion variants relative to the canonical transcript.
identified here had been reported in the previously described NFKB1 cases.5–11

The NFKB1 gene encodes the p105 protein, which is processed to produce the active DNA-binding p50 subunit.12,13 The 16 potentially pathogenic variants we identified were all located in the N-terminal p50 part of the protein (Fig 2). The effects of the 3 rare substitutions on NF-kB1 structure were less clear than those of the truncating and gene deletion variants, and therefore we assessed their position in the crystal structure of the p50 protein. Their location in the inner core of the RHD (Fig 3, A) suggested a potential effect on protein stability, whereas other rare substitutions in the NHRBR-RD cohort were found in locations less likely to affect this (Figs 2 and 3, A, and see Fig E3 in this article’s Online Repository at www.jacionline.org).

NFKB1 LOF as the disease mechanism

Twelve patients with truncating variants (Arg284*, His513Glnfs*28, c.160-1G>A, and Asp451*), 1 patient with gene deletion (del 103370996-103528207), and 3 patients with putative protein destabilizing missense variants (Ile281Met, Val98Asp, and Ile87Ser) were investigated for evidence of reduced protein level. Assessment of the NF-kB1 protein level in PBMCs or monocytes demonstrated a reduction in all subjects (38% ± 4.3% mean ± SEM) compared with healthy control subjects. There was no difference between clinically affected and clinically unaffected subjects (36% ± 4.4% vs 42% ± 10.1%, n = 11 vs n = 5, P = .50). Our observations indicate that the pathogenic NFKB1 variants result in LOF of the NF-kB1 p50 subunit because reduction in protein levels was seen in all carriers regardless of their clinical phenotype and was absent in family members who were noncarriers.

**Variable disease manifestations in NFKB1 LOF subjects**

Seven subjects had evidence of familial disease (Table I), prompting us to investigate genotype-phenotype cosegregation and disease penetrance in cases for which pedigree information and additional family members were available (Fig 4 and see Tables E1 to E3 in this article’s Online Repository at www.jacionline.org). The age at which hypogammaglobulinemia becomes clinically overt is highly variable (see Fig E5 in this article’s Online Repository at www.jacionline.org), as shown by pedigree C in which grandchildren carrying the c.160-1G>A splice-site variant had IgG subclass deficiency (C:III-3 and C:III-4), in one case combined with an IgA deficiency (C:III-3). Although not yet overtly immunodeficient, the clinical courses of their fathers (C:II-3 and C:II-5) and grandmother (C:I-2) predict this potential outcome and warrant long-term clinical follow-up of these children.

We also observed variants in subjects who were clinically asymptomatic. Pedigree A highlights variable disease penetrance: the healthy mother (A:II-1) carries the same Arg284* variant as 2 of her clinically affected children (A:III-2 and A:III-3). Identification of this nonsense variant prompted clinical assessment of the extended kindred and demonstrated that her sister (A:II-4) had recurrent sinopulmonary disease and nasal polyps with serum hypogammaglobulinemia consistent with a CVID diagnosis. Overall, based on the clinical symptoms observed at the time of this study across 6 pedigrees, the penetrance of NFKB1 variants with respect to the clinical manifestation of CVID is incomplete (about 60% in our cohort, 11 affected subjects among 18 variant carriers), with varied expressivity not only of age at disease onset but also of specific disease manifestations, even within the same pedigree.

The clinical disease observed among the NFKB1 variant carriers is characteristic of progressive antibody deficiency associated with recurrent sinopulmonary infections (100% of clinically affected subjects) by encapsulated microbes, such as Streptococcus pneumoniae and Haemophilus influenzae (see

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**Table I. Summary of the CVID patients’ clinical presentation and NFKB1 variants**

| Case ID | Sporadic/familial | Infections | Autoimmunity | Malignancy | Chromosome 4 position (GRCh37) | Nucleotide change | Type of variant | cDNA (NM_003998.3); Amino acid |
|---------|-------------------|------------|--------------|------------|--------------------------------|------------------|----------------|----------------------------------|
| A       | Familial          | ●          |              |            | 103504037                       | C>T              | Nonsense       | c.850C>T; Arg284*                 |
| B       | Familial          | ●          | ●           | ○          | 103518717                       | delCATGC         | Frameshift     | c.1539_1543del; His513Glnfs*28    |
| C       | Familial          | ●          | ●           | ○          | 103459014                       | G>A              | Splice acceptor | c.160-1G>A                     |
| D       | Familial          | ●          | ●           | ○          | 103518801                       | delGA            | Nonsense       | c.1621_1622del; Asp541*           |
| E       | Sporadic          | ●          | ●           | ○          | 103504030                       | C>G              | Missense       | c.293T>G; Val98Asp                |
| F       | Sporadic          | ●          | ●           | ○          | 103488178                       | T>A              | Missense       | c.843C>G; Ile281Met               |
| G       | Sporadic          | ●          | ●           | ○          | 103488115                       | T>G              | Missense       | c.260T>G; Ile87Ser                |
| H       | Familial          | ●          | ●           | ○          | 103501798                       | T>C              | Missense       | c.835+2T>C                     |
| I       | Sporadic          | ●          | ●           | ○          | 103370996-103528207             | delG             | Missense       | c.1423del; Ala475Profs*10         |
| J       | Familial          | ●          | ●           | ○          | 103517841                       | delG             | Missense       | c.137del; Ala475Profs*10         |
| K       | Sporadic          | ●          | ●           | ○          | 103436974-10365265              | delG             | Missense       | c.1036del; Arg369Profs*10         |
| L       | Familial          | ●          | ●           | ●          | 103459041                       | delG             | Frameshift     | c.187del; Glu63Lysfs*64           |
| M       | Sporadic          | ●          | ●           | ●          | 103501790                       | insA             | Frameshift     | c.830dup; Lys278Glufs*3           |
| N       | Sporadic          | ●          | ●           | ●          | 103504086                       | insT             | Frameshift     | c.904dup; Ser302Profs*7          |
| O       | Sporadic          | ●          | ●           | ●          | 103588130                       | C>T              | Nonsense       | c.295C>T; Gln99*                  |
| P       | Sporadic          | ●          | ●           | ●          | 103505914                       | delG             | Frameshift     | c.1005del; Arg369Profs*10         |

●, Presence of symptoms in index patient; ○, presence of symptoms in family member of index patient.
The clinical spectrum of \( \text{NFkB1} \) LOF variants includes massive lymphadenopathy (24%), unexplained splenomegaly (48%), and autoimmune disease (48%), either organ-specific and/or hematologic in nature (mainly autoimmune hemolytic anemia and idiopathic thrombocytopenic purpura, see Table E1). The percentage of autoimmune complications is based on the presence of autoimmune cytopenias (autoimmune hemolytic anemia, idiopathic thrombocytopenic purpura [\(<50-75 \times 10^9/mL\), autoimmune neutropenia, and Evans syndrome), alopecia areata/totalis, vitiligo, and Hashimoto thyroiditis among the clinically affected cases. Granulomatous-lymphocytic interstitial lung disease and splenomegaly were considered lymphoproliferation. Enteropathy, liver disease, colitis, and a mild decrease in platelet count (\(>100 \times 10^9/mL\)) were neither included in those calculations nor scored separately. Histologic assessment of liver disease found in 3 patients showed no evidence of autoimmune or granulomatous liver disease, although fibrosis and cirrhosis were observed in these male patients. Finally, the number of oncological manifestations, predominantly hematologic, was noticeable. There were 2 cases with solid tumors (parathyroid adenoma and breast cancer) and 4 cases with hematologic malignancies (B-cell non-Hodgkin lymphoma, diffuse large B-cell lymphoma, follicular lymphoma, and peripheral T-cell lymphoma), which add up to 6 (28.6%) of 21 cases.

### B-cell phenotype in subjects with \( \text{NFkB1} \) LOF mutations and immune cell activation

Index cases and family members carrying \( \text{NFkB1} \) variants were approached for repeat venipuncture for further functional assessment. In clinically affected subjects, B-cell numbers and phenotypes were indistinguishable from those described for patients with CVID (Fig 5 and see Fig E6 in this article’s Online Repository at www.jacionline.org). However, in clinically unaffected subjects the absolute B-cell count was often normal or increased (Fig 5, A). In all subjects with \( \text{NFkB1} \) LOF variants, the numbers of switched memory B cells were reduced (Fig 5, B-D), whereas a broad range of nonswitched memory B cells was observed. This demonstrates that although the clinical phenotype of \( \text{NFkB1} \) LOF variants is partially penetrant, all carriers have a deficiency in class-switched memory B-cell generation. The presence of increased numbers of the CD21\textsuperscript{low} population described in patients with CVID discriminates between clinically affected and unaffected subjects with \( \text{NFkB1} \) LOF variants (Fig 5, E). B cells from subjects with \( \text{NFkB1} \) LOF variants demonstrated impaired proliferative responses to anti-IgM/anti-CD40/IL-21 and CpG/IL-2 (Fig 6, A); this corresponded with the inability to generate plasmablasts (CD38\textsuperscript{+}/CD27\textsuperscript{+}), which was most pronounced in the more extreme phenotypes (ie, clinically affected cases; Fig 6, B, and see Fig E7, B, in this article’s Online Repository at www.jacionline.org). Similarly, \( \text{ex vivo} \) IgG production was reduced in subjects with LOF variants, whereas IgM levels in the supernatants were normal (Fig 6, C and D, and see Fig E7, C), which is compatible with hypogammaglobulinemia.

### T-cell phenotype in subjects \( \text{NFkB1} \) LOF variants

The T-cell phenotype was largely normal in the subset distribution (see Figs E8 and E9 in this article’s Online Repository at www.jacionline.org). Similar to the knockout mouse model,\(^8\) we found an aberrant number of invariant natural killer T cells in clinically affected subjects (see Fig E8). T-cell proliferation was intact on anti-CD3/anti-CD28 or IL-15 activation (see Fig E10 in this article’s Online Repository at www.jacionline.org). Because invariant natural killer T cells have been implicated in diverse immune reactions,\(^9\) this deficiency might contribute to...
DISCUSSION

In our study we show that LOF variants in NFKB1 are present in 4% of our cohort of patients with CVID, being the most commonly identified genetic cause of CVID. Furthermore, we highlight specific features of these patients that distinguish them within the diagnostic category of CVID, which otherwise applies to an indiscrete phenotype acquired over time that is termed common and variable. The majority of the genetic variants we report here truncate or delete 1 copy of the gene; together with pedigree cosegregation analyses, protein expression, and B-cell functional data, we conclude that NFKB1 LOF variants cause autosomal dominant haploinsufficiency. This has now been recognized as the genetic mode of inheritance for at least 17 known PIDs, including those associated with previously reported variants in NFKB1. In monogenic causes of PID, incomplete penetrance has been more frequently described in haploinsufficient relative to dominant negative PID disease, having been reported in more than half of the monogenic autosomal dominant haploinsufficient immunologic conditions described. This might be because dominant negative gain-of-function mutations cause disease by expression of an abnormal protein at any level, whereas, as seen in this study, haploinsufficiency is predicted to lead to 50% residual function of the gene product. By definition, incomplete penetrance of a genetic illness will be associated with substantial variation in the clinical spectrum of disease, and the spectrum seen in this study is consistent with prior reports; in 3 pedigrees with 20 subjects harboring heterozygous mutant NFKB1 alleles, the age of onset varied from 2 to 64 years, with a high variety of disease severities, including 2 mutation carriers who were completely healthy at the ages of 2 and 43 years.

It is important to temper skepticism of partial penetrance of immune genetic lesions with our knowledge that individual immune genes might have evolved in response to selection...
pressure for host protection against specific pathogens. Consequently, within the relatively pathogen-free environment of developed countries, the relevant pathogen for triggering disease might be scarce, and reports documenting partial penetrance of the clinical phenotype will increase. This makes the traditional approaches to genetics for determining causality difficult. The BeviMed algorithm used in this study prioritized both the gene NFKB1 and individual variants within NFKB1 for contribution to causality; the power of methods like this will increase with greater data availability. Identification of a number of rare NFKB1 variants with high Combined Annotation Dependent Depletion scores in both the PID and control data sets highlights the potential for false attribution of disease causality when the genetics of an individual case are considered outside the context of relevant control data.

Currently healthy family members carrying the same NFKB1 LOF variant demonstrated similar reductions in p50 expression and low numbers of switched memory B cells as their relatives with CVID. The longitudinal research investigation of these subjects could help identify the additional modifiers, including epigenetic or environmental factors, that influence the clinical penetrance of these genetic lesions. The similarity of results seen in patients with large heterozygous gene deletions and in those with more discrete substitutions is consistent with haploinsufficiency as the shared disease mechanism.

In patients with mild antibody deficiency, it is often difficult to decide when to initiate replacement immunoglobulin therapy; this might be the case for subjects and their family members identified with LOF NFKB1 variants. Two measures seem to correlate well with clinical disease. First, the class-switch defect and lower IgG and IgA production \textit{ex vivo} was examined. Immunoglobulin class-switching is known to be regulated by NF-κB. Mutations in NF-κB essential modulator cause a class-switch–defective hyper-IgM syndrome in human subjects,\textsuperscript{20} as well as in p50 knockout mice.\textsuperscript{13,44,45} Haploinsufficiency of NF-κB might result in defective...
class-switch recombination because of poor expression of activation-induced cytidine deaminase, a gene regulated by NF-κB, which, when absent, is also associated with immunodeficiency. 46

Second, the ability to measure the CD21low B-cell population is widespread in diagnostic immunology laboratories, and our study identifies this marker to correlate with NF-κB disease activity. Although the function of these cells remain to be fully elucidated, 47 this laboratory test might be useful for longitudinal assessment of clinically unaffected subjects identified with LOF NFKB1 variants.

Apart from having recurrent and severe infections (including viral disease) for which these patients had been given a diagnosis of PID in the first place, autoimmunity and unexplained splenomegaly are very common manifestations in our patient cohort similar to the other heterozygous NFKB1 cases described. 6-11 Although autoimmunity has been subject to variable percentages per cohort study, 3,48,49 it seems that these complications occur more frequently in NFKB1 haploinsufficient patients compared with unselected CVID cohorts. In contrast to IKAROS defects but similar to cytotoxic T lymphocyte–associated protein 4 (CTLA4) haploinsufficiency, we observed that NFKB1 haploinsufficiency can also result in chronic and severe viral disease, as noted for cytomegalovirus and JC virus infections in 3 of our patients. In the study of Maffucci et al, 11 one of the NFKB1-affected patients also experienced Pneumocystis jirovecii infection and progressive multifocal leukoencephalopathy (PML), which is suggestive for JC virus infection. Whether the B-cell defect in NFKB1 haploinsufficiency is responsible for these nonbacterial infections is unclear. 50,51 PML is most often discovered in the context of an immune reconstitution inflammatory syndrome, as seen in patients with HIV receiving antiretroviral therapy and in patients with multiple sclerosis after natalizumab discontinuation. 52 Although the exact contribution of B-cell depletion in PML pathogenesis is unknown, the increased PML risk in rituximab-treated patients 53 suggests a protective role for B cells.

Three subjects in this cohort had liver failure, and an additional 3 had transaminitis. Although autoimmunity is suspected, a nonhematopoietic origin of liver disease cannot be excluded in the absence of autoantibodies and nodular regenerative disease. Mouse models have suggested a nonimmune role for NF-κB signaling in patients with liver failure. 13,54-56

![Fig 6](image_url) Ex vivo class-switch recombination defect of subjects carrying NFKB1 LOF variants is linked to the more extreme phenotype. Six-day culture of CFSE-labeled lymphocytes normalized for B-cell numbers: unstimulated, CpG/IL-2 stimulated (T cell–independent activation), or anti-IgM/anti-CD40/IL-21 stimulated (T cell–dependent activation). A, Percentage of divided B cells, as measured based on CFSE dilution. B, Percentage of CD27+ plasmablasts. The gating strategy is shown in Fig E7, A, C, and D, IgM and IgG production in supernatants of 6-day cultures. CA, Clinically affected subjects with an LOF variant in NFKB1; CU, clinically unaffected subject; HD, healthy donor. Only subjects with sufficient B cells could be analyzed. P values were determined by using 2-way ANOVA with the Bonferroni post hoc test. ns, Not significant. **P<.01 and ***P<.001.
In the cohort of patients with NFKB1 variants, we identified a number of malignancies. Malignancies in patients with PID have been cited as the second-leading cause of death after infection, and murine models have demonstrated that haploinsufficiency of NF-κB1 is a risk factor for hematologic malignancy. In a large CVID registry study of 2212 patients, 9% had malignancies, with one third being lymphomas, some presenting before their CVID diagnosis. Despite the fact that our cohort is relatively small, we found oncologic manifestations in 29% of our cases (two thirds being lymphoma), suggesting that malignancies in patients with NFKB1 haploinsufficiency can occur more often than in unselected patients with CVID. In a study in 176 patients with CVID, among the 626 relatives of patients with CVID, no increase in cancer risk was observed, suggesting that when this does occur, as in this study (3/7), it might be due to a shared genetic lesion. Therefore in a pedigree with an LOF variant in NFKB1, any relatives with cancer should be suspected of sharing the same pathogenic variant. In conclusion, previous publications have suggested that CVID is largely a polygenic disease. Our results provide further evidence that LOF variants in NFKB1 are the most common monogenic cause of disease to date, even in seemingly sporadic cases. In these patients there is a clear association with complications, such as malignancy, autoimmunity, and severe nonimmune liver disease; this is important because the excess mortality seen in patients with CVID occurs in this group. The screening for defined pathogenic NFKB1 variants accompanied by B-cell phenotype assessment has prognostic value and is effective in stratifying these patients.

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Key messages

- Pathogenic variants in NFKB1 are currently the most common known monogenic cause of CVID.
- There is a clear association with complications, such as autoimmunity and malignancy, features associated with worse prognosis.
- These patients can be stratified by NFKB1 protein level and B-cell phenotype.

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