CLINICAL TRIALS AND OBSERVATIONS

Pediatric Evans syndrome is associated with a high frequency of potentially damaging variants in immune genes

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KEY POINTS

- At least 65% of cases of pES may be genetically determined.
- Genetic findings have prognostic significance and may guide the physician's choice of a targeted treatment.

Evans syndrome (ES) is a rare severe autoimmune disorder characterized by the combination of autoimmune hemolytic anemia and immune thrombocytopenia. In most cases, the underlying cause is unknown. We sought to identify genetic defects in pediatric ES (pES), based on a hypothesis of strong genetic determinism. In a national, prospective cohort of 203 patients with early-onset ES (median [range] age at last follow-up: 16.3 years ([1.2-41.0 years]) initiated in 2004, 80 nonselected consecutive individuals underwent genetic testing. The clinical data were analyzed as a function of the genetic findings. Fifty-two patients (65%) received a genetic diagnosis (the M+ group): 49 carried germline mutations and 3 carried somatic variants. Thirty-two (40%) had pathogenic mutations in 1 of 9 genes known to be involved in primary immunodeficiencies (*TNFRSF6*, *CTLA4*, *STAT3*, *PIK3CD*, *CBL*, *ADAR1*, *LRBA*, *RAG1*, and *KRAS*), whereas 20 patients (25%) carried probable pathogenic variants in 16 genes that had not previously been reported in the context of

autoimmune disease. Lastly, no genetic abnormalities were found in the remaining 28 patients (35%, the M- group). The M+ group displayed more severe disease than the M- group, with a greater frequency of additional immunopathologic manifestations and a greater median number of lines of treatment. Six patients (all from the M+ group) died during the study. In conclusion, pES was potentially genetically determined in at least 65% of cases. Systematic, wideranging genetic screening should be offered in pES; the genetic findings have prognostic significance and may guide the choice of a targeted treatment. (*Blood.* 2019;134(1):9-21)

Introduction

Autoimmune diseases are numerous and common in adults.¹ The underlying mechanism is complex, and involves both environmental and polygenic factors.¹ A few monogenic autoimmune diseases have nevertheless been identified, mostly in children, as isolated or combined manifestations in the context of primary immunodeficiencies (PIDs).^{2,3} These diseases include autoimmune lymphoproliferative syndrome (ALPS, caused by mutations in the Fas apoptotic pathway),⁴ autoimmune polyendocrinopathy candidiasis ectodermal dystrophy (due to AIRE mutations),⁵ immunodysregulation polyendocrinopathy enteropathy X-linked (caused by FOXP3 mutations),⁶ and many others; indeed, variants in >150 different genes have been found to drive autoimmunity.7 The analysis of these genes has provided a wealth of information on the mechanisms that control reactivity to self in humans, including central negative selection, antigen receptor gene editing, regulatory T-cell count numbers and functions, peripheral elimination of autoreactive T and B cells, the clonal redemption of B cells from self-reactivity, and cis-acting regulatory elements of antigen-specific immune responses.8-11

It has been suggested that the occurrence of autoimmune disease in children (where is it relatively rare, compared with adults) may be caused by a high-risk predisposition gene. This hypothesis is based on the fact that (i) autoimmune conditions are present in relatively high proportion of PIDs (\sim 25%) and (ii) autoimmune conditions occur between 10 and 80 times more frequently in children with a PID than in an age-matched nonclinical population.³ For immune thrombocytopenic purpura (ITP) and autoimmune hemolytic anemia (AIHA), the relative risk factor is particularly high (up to 120 in childhood)³; however, no large series of nonselected patients with hematologic autoimmune disease have been comprehensively explored using next-generation sequencing (NGS). This prompted us to screen the French OBS'CEREVANCE cohort of patients with pediatric Evans syndrome (pES, a rare severe disease characterized by the simultaneous or sequential development of ITP and AIHA^{12,13}) for the presence of potentially damaging mutations.

Patients and methods

Patient selection and data collection

As of 1 April 2018, a total of 203 children (ie, patients under the age of 18 years) with pES had been consecutively included in the OBS'CEREVANCE French national observational cohort. The inclusion and exclusion criteria are summarized in supplemental Table 1 (available on the Blood Web site). For each patient, relevant aspects of the family medical history, and clinical, laboratory and treatment-related data from birth to last followup were prospectively collected, coded, and integrated into a database. Lymphoproliferation was defined by clinically significant, persistent lymphadenopathy (>1 cm), and/or clinically significant splenomegaly detected below the costal margin in the absence of active hemolysis. Long-term follow-up data on clinical events were also collected. During the whole followup period, lines of treatments, treatment outcomes, and the presence of absence of additional immunopathologic manifestations and severe or recurrent infections were prospectively registered (supplemental Table 1). Written, informed consent was obtained from the patients' parents and then (when old enough) from the patients themselves. The cohort database was registered with the French national data protection authority (CNIL) on 9 November 2009. Since 2015, 80 patients from the cohort (the study group; Figure 1) have undergone genetic testing. These 80 patients were consecutively included in the present genetic study on request by their attending physicians, once the latter had become aware of the study. Patients with a documented PID before genetic testing were excluded (n = 3). We have previously reported on a small subgroup of these patients (n = 18).¹⁴ The median (range) length of follow-up following the first observation of cytopenia and following the diagnosis of ES were 9.1 years (0.2-26.6 years) and 6.7 years (0.2-25 years), respectively. To avoid selection bias, we compared the study group of 80 patients with the 123 nontested members of the cohort (Table 1). With the exception of slightly but significantly higher frequencies of consanguinity, ITP as the first symptom, and neutropenia, there were no significant intergroup differences, notably with regard to associated immunopathologic manifestations, the median length of follow-up, the need for second-line therapy, and death.

Genetic analysis

Samples of DNA were prepared from the patients' whole peripheral blood, using standard extraction methods.¹⁵ For patients with typical clinical profiles and laboratory data suggestive of a specific PID, Sanger sequencing was performed on the most likely candidate genes. All patients with negative Sanger sequencing results (n = 69) underwent targeted NGS (tNGS) of 203 genes or (in selected cases with consanguinity or multiplex families) whole-exome sequencing. The methods used for genetic analysis and the custom panel used for tNGS are described in supplemental Appendix and supplemental Table 2, respectively. The 80 patients were then classified into 3 groups, depending on the results of the genetic analysis: (i) patients with a "pathogenic" mutation, that is, the identified mutation had been described previously in the context of autoimmune disease; (ii) patients with a "probably pathogenic" mutation, that is, the identified mutation was likely to be pathogenic but had not been described previously in the context of autoimmune disease; and (iii) patients lacking an identified genetic defect. The pathogenicity of "probably pathogenic" mutations was defined according to the following criteria: the frequency in the general population (minor allele frequency <0.01 in the GnomAD database; http://gnomad.broadinstitute.org), a literature review of the coded protein's function, and algorithms predicting the pathogenicity of missense mutations, a Sorting Intolerant from Tolerant (SIFT) score (http://sift.jcvi.org/www/SIFT_seq_submit2. html) below 0.05 and a PolyPhen2 score (http://genetics.bwh. harvard.edu/pph2/) above 0.85, or a Combined Annotation-Dependent Depletion (CADD) score above 20 (http://cadd.gs. washington.edu/home).

Statistical analysis

Continuous variables were described as the median (range), and categorical variables were described as the number (percentage). Quantitative variables were compared using the χ^2 test or (for a small sample size) the Fisher exact test. Qualitative variables were compared using the Mann-Whitney nonparametric test (for 2 variables) or an analysis of variance (for 3 variables). The percentage of patients free of additional immunopathologic manifestations was analyzed from the date of birth to the date of



Figure 1. Summary of clinical presentations in the study group.

the first nonhematological immunopathologic manifestation or censored at the date of last follow-up. Survival estimates were calculated using the Kaplan-Meier method. Differences in survival estimates were assessed using the log-rank test. All statistical analyses were performed using R software (version 3.3.2; R Foundation for Statistical Computing, Vienna, Austria).

Results

Genetic analysis

Fifty-two of the 80 patients in the study group (65%, forming the M+ group) obtained a molecular diagnosis. Details of the mutations, the diagnostic method, and the predicted impact at the protein level are given in Tables 2 and 3. Thirty-two patients (40%) had pathogenic mutations in 9 genes known to be involved in PIDs: 23 had autosomal heterozygous germline mutations in TNFRSF6 (n = 6), CTLA4 (n = 8), STAT3 (n = 6), PIK3CD (n = 1), CBL (n = 1) or ADAR1 (n = 1), and 6 patients had autosomal homozygous disease with a homozygous LRBA mutation (n = 3), a large homozygous LRBA deletion (n = 1), or a compound heterozygous RAG1 mutation (n = 2). Three of the M+ patients had somatic heterozygous mutations, including a TNFRSF6 mutation detected in double-negative T cells (n = 1), and a KRAS mutation (n = 2). All of the variants in the "pathogenic" group have been previously described¹⁶⁻²⁶ or otherwise functionally validated. TNFRSF6 variants have been validated by the observation of an apoptosis defect in T cells, with CTLA4 variants validated by low CTLA4 expression, LRBA variants validated by the absence of protein expression in a western blot analysis, and a *STAT3* variant validated by elevated *SOCS3* RNA transcript levels (detected using a quantitative polymerase chain reaction assay).²⁷ Patients with *RAG1* mutations developed naive T-cell lymphopenia, which was suggestive of a combined immunodeficiency.

Twenty patients (25%) had "probably pathogenic" variants in 16 genes that had not previously been described in the context of autoimmune diseases and/or were suggestive of new molecular mechanisms (Tables 2 and 3). All but one of the variants met our predicted pathogenicity criteria (a CADD score >20, or PolyPhen2 and SIFT scores >0.85 and <0.05, respectively). The NFATC1 variant P49 had a lower CADD score but had a demonstrated impact on exon splicing with complementary DNA (cDNA; data not shown). Furthermore, another patient (P48) had a pathogenic variant in NFATC1. No other variants in immune-related genes were detected in these 20 patients. These variants mainly involve immune cell receptors (with a probable loss of function [LOF] in TNFR2, TRAF3, IFNAR1, and TGFBR2, n = 1 for each), intracellular signaling (JAK1, with a probable gain of function [GOF] in 2 cases, and JAK2 with a probable GOF, PLCG2 with a probable GOF, CARD11 with a probable GOF, PARP4 with a probable LOF, ARGHEF4 with a probable GOF, and PTPN11 with a probable GOF, n = 1 for each), the regulation of apoptosis (RIPK2 with a probable LOF in 2 cases, and APAF1, with a probable LOF in 1 case), and lastly, transcriptional factors in immune cells (IKZF1 with a probable GOF and NFATC1 with a probable GOF in 2 cases each, and *IKZF2* with a probable LOF in 1 case).

Table 1. Demographical and clinical description of 203 pES patients, and the study group of 80 patients

	pES patients, N = 203	Study group, N = 80	Others, N = 123	P, study group vs others
Median age at first episode of cytopenia (min- max), y	5.9 (0.2-17.4)	5.9 (0.3-15.6)	5.9 (0.2-17.4)	
Median age at ES diagnosis (min-max), y	9.1 (0.2-19.4)	8.7 (0.3-17.9)	9.5 (0.2-19.4)	
Sex ratio (female/male)	0.8 (91/112)	0.68 (32/48)	0.90 (59/64)	
Consanguinity, % (n)	9 (16/177)	13 (11)	5 (5)	.047
Immune manifestations in first-degree relatives, % (n)	31 (57/181)	30 (24)	32 (33)	
Immune manifestations in extended relatives, % (n)	38 (69/181)	42 (34)	37 (37)	
Occurrence of ITP and AIHA Simultaneous, % (n) Sequential, % (n) AIHA first, % (n) ITP first, % (n)	42 (85) 58 (117) 23 (47) 35 (70)	40 (32) 60 (48) 14 (11) 46 (37)	43 (53) 57 (69) 30 (36) 27 (33)	.01 .004
Immune neutropenia, % (n)	32 (64)	44 (35)	24 (29)	.002
Associated immunopathologic manifestations, % (n)	75 (153)	82 (66)	71 (87)	
Need for second-line therapy, % (n)	72 (146)	75 (60)	70 (86)	
Rituximab, % (n)	44 (90)	51 (41)	40 (49)	
Splenectomy, % (n)	19 (39)	17 (14)	20 (25)	
Median follow-up from first cytopenia (min-max), y	7.9 (0.1-29.0)	9.1 (0.2-26.6)	7.1 (0.1-29.0)	
Median follow-up from ES diagnosis (min-max), y	5.8 (0.1-29.1)	6.7 (0.2-25.0)	5.5 (0.1-29.1)	
Deaths, % (n)	10 (21)	7 (6)	12 (15)	

max, maximum; min, minimum.

In light of recently published results, some of the variants merit further comment. IKZF1 LOF mutations have been described in patients with common variable immunodeficiency (haploinsufficiency mutations)²⁸ and combined immunodeficiency (dominant-negative mutations).²⁹ In contrast, we suggest that the 2 patients studied here carry a previously undescribed GOF mutation leading to an autoimmune phenotype; this GOF hypothesis is supported by preliminary data, that is, greater binding of Ikaros to specific DNA targets in an electrophoretic mobility shift assay.³⁰ Various types of CARD11 mutation can lead to distinct phenotypes; for example, biallelic null mutations lead to severe T- and B-cell immune deficiencies,³¹ whereas germline GOF mutations give rise to "B-cell expansion with NFкВ and T-cell anergy" (BENTA) disease.³² Lastly, hypomorphic/ dominant-negative mutations predispose to atopic phenotypes and variable immunodeficiency.^{33,34} The present CARD11 mutation has never been reported; functional data are not yet available, and the phenotype differs from those reported. A recent study indicated that hypomorphic CARD11 mutations are associated with a variety of immunologic phenotypes and (in some cases) atopic disease,³⁵ including autoimmunity in 20% of patients. PLCG2 deletions are responsible for the PLCG2-associated antibody deficiency and immune dysregulation

(PLAID) syndrome (cold urticaria, immunodeficiency, and autoimmunity³⁶), and missense GOF mutations have been associated with a dominantly inherited autoinflammatory disease with immunodeficiency.37,38 The PLCG2 mutation (I875M) reported here has never been described. The fact that it is located near to the mutation reported by Neves et al³⁸ (L848P) prompts us to suspect a GOF mutation. JAK1 GOF mutations have been recently described in patients with severe atopic dermatitis and hypereosinophilic syndrome without autoimmune manifestations.³⁹ Given the importance of the JAK-STAT pathway in PIDs with autoimmunity (such as STAT1 and STAT3 GOF mutations), a GOF mutation leading to hyperactivation of the JAK-STAT pathway and a specific clinical phenotype is likely. Germline JAK2 mutations have previously been linked to myeloproliferative neoplasms but not to autoimmune manifestations. By analogy with the JAK1 variants, we suggest that a JAK2 GOF mutation is present. Homozygous IFNAR1 mutations have been recently described in patients with increased susceptibility to viral infections⁴⁰; our patient's clinical phenotype (ie, autoimmunity) is quite different. Preliminary data have evidenced the impaired induction of interferon-stimulated genes (n = 6) in the patient's peripheral blood mononuclear cells (PBMCs; relative to PBMCs from healthy controls) following

Table 2. Results of genetic testing in the study group(80 patients with pES)

Gene (no. of patients)	Mutation type and consequences
Pathogenic mutations, n = 32, 40% TNFRSF6 (6) CTLA4 (8) STAT3 (6) PIK3CD (1) CBL (1) ADAR1 (1) LRBA (4) RAG 1 (2) TNFRSF6 somatic (1) KRAS somatic (2)	Heterozygous/LOF Heterozygous/LOF Heterozygous/GOF Heterozygous/LOF Heterozygous/LOF Homozygous/LOF Compound heterozygous/LOF Heterozygous/LOF Heterozygous/LOF
Probably pathogenic mutations, n = 20, 25% Immune cell receptors <i>IFNAR1</i> (1) <i>TNFR2</i> (1) <i>TGFBR2</i> (1) Intracellular signaling <i>JAK1</i> (2) <i>JAK2</i> (1) <i>PLCG2</i> (1) <i>TRAF3</i> (1) <i>CARD11</i> (1) <i>ARHGEF4</i> (1) <i>PTPN11</i> (1) <i>PARP4</i> (1)	Homozygous/likely LOF Heterozygous/likely GOF Heterozygous/likely GOF Heterozygous/likely GOF Heterozygous/likely GOF Heterozygous/likely GOF Heterozygous/likely GOF Heterozygous/likely GOF Heterozygous/likely GOF Compound heterozygous/likely LOF
Apoptosis regulation <i>RIPK2</i> (2) <i>APAF1</i> (1) Transcription factors <i>IKZF1</i> (2) <i>NFATC1</i> (2) <i>IKZF2</i> (1)	Heterozygous/likely LOF Heterozygous/likely GOF Heterozygous/likely GOF Heterozygous/likely GOF Heterozygous/likely LOF
No genetic abnormalities, n = 28, 35%	

GOF, gain of function; LOF, loss of function.

stimulation with interferon- α , thus suggesting a LOF mutation (data not shown). Germline mutations in *PTPN11* are usually associated with Noonan syndrome, which is characterized by a dysmorphic syndrome, congenital heart disease, and coagulation defects.⁴¹ However, dysregulation of the MAPK pathway due to a somatic mutation in *NRAS* has been linked to an ALPS-like syndrome (RAS-associated ALPS-like disease [RALD^{25,42}]). We suggest that a GOF mutation in *PTPN11* in the present case might lead to hyperactivation of the MAPK pathway and thus the onset of autoimmunity. *TGFBR2* mutations have been linked to Marfan syndrome–related disorders.⁴³ The patient described here does not present a Marfan phenotype. A LOF mutation that impairs regulatory T-cell activity could conceivably lead to autoimmunity. All of the other germline mutations detected in the "probably pathogenic" group have never been described in human diseases; all are now undergoing extensive functional validation, in order to precisely characterize the mechanism leading to autoimmunity.

Lastly, neither tNGS (n = 20) nor whole-exome sequencing (n = 8) detected any genetic abnormalities in 28 of the 80 patients (35%, forming the M- group).

Clinical phenotypes and outcomes

The clinical presentations of the 80 children in the study group are described in detail in Table 1. The median (range) duration of follow-up was 9.1 years (0.2-26.6 years). pES was associated with additional immunopathologic manifestations in 66 of the patients (82%). These manifestations were present before the first episode of cytopenia in 18 cases (27%, with a median [range] time interval of 3.6 years [0.4-14 years]), and arose at the same time as the first episode of cytopenia in 22 cases (33%), or arose afterward in 26 cases (39%, with a median [range] time interval of 3 years [0.1-11.5]). Forty of the 80 patients (50%) had lymphoproliferation (8 had isolated splenomegaly, 8 had isolated lymph node enlargement, and 24 had both), 37 (46%) had hypogammaglobulinemia (after anti-CD20 antibody treatment in 25 of these), and 38 (47%) had various autoimmune/ autoinflammatory manifestations (mainly liver, digestive tract, and lung manifestations). The median (range) number of additional immunopathologic manifestations per patient was 2 (0-8). Severe or recurrent infections occurred in 47% of the patients, although we were not able to determine whether these were disease-related or treatment-related.

We next compared the M+ and M- groups with regard to the phenotype (Figure 1; Table 4). The M+ group was more likely to present with additional immunopathologic manifestations (P = .007) (Figure 2) and hypogammaglobulinemia (P = .02). There was a nonsignificant trend toward a greater frequency of lymphoproliferation (P = .06) in the M+ group. No significant differences were observed between the patients with a known pathogenic mutation and the patients with a probable pathogenic variant, with the exception of the occurrence of gut immunopathologic manifestations (P = .008) and a trend toward the more frequent occurrence of lung manifestations (P = .07) and ALPS biomarkers (P = .07) in the "pathogenic" group (Table 4). Overall, these results suggest that pES tends to be more severe when it occurs in the context of genetic abnormalities, as illustrated by a higher median number of lines of treatment. The time to the first immunopathologic manifestation was shorter in the M+ group than in the M- group (Figure 3), while the shape of the curves suggests that the genetic impact on the occurrence of additional immunopathologic manifestations is greater in the first 5 years of life than later in life. Six of the 80 patients in the study group died during the study period (supplemental Figure 1; supplemental Table 3). It is noteworthy that all 6 belonged to the M+ group. Death occurred at a median (range) age of 18.9 years (3.9-25 years), and the median (range) time interval between the initial diagnosis and death was 10.7 years (0.9-24.6 years). Three deaths were disease-related (cerebral hemorrhage, juvenile myelomonocytic leukemia, and fulminant hepatitis) and 3 were disease- or treatment-related (sepsis, Epstein-Barr virus lymphoproliferation, and pneumococcal meningitis). Three asplenic patients died, although asplenic sepsis was the cause of death in only 1 patient.

Table 3. Detailed genetic results with in silico analysis in 52 pES

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MAF	c	þ	0	0	c	D	0	C		0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	4.066×10^{-6}
CADD	VIV		NA	41	,	23.6	NA	22.4		27	36	32	21	36	14.4	ΝA	23.9	21.8	18.4	35	25.3	24.2	25.6	31	25	34	25.9	NA	NA	NA	26.4
SIFT	VIV		NA	NA	500	0.01	AN	AA		0	0	0	AN	0	0.17	NA	0	0.02	0.13	0	0.42	0.34	0.11	0.01	0.06	0.05	NA	NA	NA	AN	0
PolyPhen2	< Z		NA	NA		-	NA	AA		-	0.999	0.999	AN	0.999	0.653	AN	0.998	0.997	0.995	-	0.986	0.999	-	0.999	0.536	-	AN	AN	AN	AN	0.974
OMIM no.	267861	10040	134637	134637		13463/	134637	134637		123890	123890	123890	123890	123890	123890	123890	123890	102582	102582	102582	102582	102582	102582	602839	165360	146920	606453	606453	606453	606453	179615
Method	Canaor	Jaliger	Sanger	Sanger	,	Sanger	Sanger	Sanger	0	TNGS	Sanger	TNGS	TNGS	TNGS	Sanger	Sanger	TNGS	TNGS	WB	TNGS	TNGS	TNGS									
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Protein level mutation	Ž		D269fsX277	R250X		A23/P	D269fsX279	AN		P137L	R51X	R70W	NA	R51X	T106P	p.M38Kfs*22	C58G	T663I	T716M	G421R	G419R	Y640F	P715L	E1021K	A474T	G1007R	E789fsX792	del exon 18 to 30	L564RfsX25	A2541YfsX2542	S37Y
cDNA mutation	32E 120~G		808_815del	748C>T		/0AG>C	806_807del	506-2A>G		410C>T	151C>T	208C>T	110-2A>G	151C>T	316A>C	c.109 + 1092_568- 512del	172T>G	1988 C>T	2147C>T	1261G>A	1255G>C	1919A>T	2144C>T	3061G>A	1420G>A	3019G>A	2450+1C>T	del exon 18 to 30	1691delT	7620_7621 insT	110C>A
Genetic	TNIEDCEA	dermline	TNFRSF6 aermline	TNFRSF6	germline	I NFKSF6 germline	TNFRSF6	TNFRSF6	germline	CTLA4	CTLA4	STAT3	STAT3	STAT3	STAT3	STAT3	STAT3	PIK3CD	CBL	ADAR1	LRBA	LRBA	LRBA	LRBA	RAG1						
Patients	Pathogenic		P2	P3	Č	P4	P5	P6		P7	P8	P9	P10	P11	P12	P13	P14	P15	P16	P17	P18	P19	P20	P21	P22	P23	P24	P25	P26	P27	P28

CADU, Compiled Amoration-Dependent, Depredont, CUNA, complementary DNA, IMAY, IMPA, IMPA, Imper allele requency, INA, not applicable, UMINI, UMINE N sequencing, WB, westem blot; WES, whole-exome sequencing. See Table 2 for expansion of other abbreviations. *Patient PBMCs were shown to have impaired induction of interferon-stimulated genes (n = 6) after stimulation by IFNα comparatively to control cells.

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GnomeAD	- 0	0 0	D	0	0		0	2	4	2	0	ю	0	0	Ŋ	0	. 	0	0	0	0	2	0	. 	0	0	
MAF	8.144×10^{-6} 0	0 0	D	0	0		0	8.408×10^{-6}	1.638×10^{-5}	7.226×10^{-6}	0	1.228×10^{-5}	0	0	1.805×10^{-5}	0	4.063×10^{-6}	0	0	0	0	8.122×10^{-6}	0	8.527×10^{-6}	0	0	3-01 / 02V V
CADD	17.8 29.2	17.49	33	33	33		23.4	23.2	26.6	23.8	23.6	24	22.1	28.3	24	27.5	26.1	ΔN	17.1	28.2	23.3	34	21	6.6	33	22	
SIFT	0.02 NA	AN C	C	0	0		0	0.12	0	0.1	0.04	0.06	0.06	0	0.04	0.07	0.03	AN	0.04	0	0.14	0	0.5	AN	0	0.12	Ĺ
PolyPhen2	0.036 0.953	0.978		~	~		0.998	0.286	0.989	0.661	0.011	0.851	0.316	-	0.375	0.995	0.001	NA	0.971	-	0.751	0.999	0.629	AN	0.998	0.154	
OMIM no.	179615		13463/	190070	190070		191191	107450	190182	147795	147795	147796	600220	601896	607210	605216	176876	607519		603455	603455	602233	600489	600489	603023	603023	
Method	TNGS	ţ	sanger	TNGS	Sanger		TNGS	WES	TNGS	TNGS	TNGS	TNGS	TNGS	TNGS	TNGS	TNGS	TNGS	TNGS		TNGS	TNGS	TNGS	TNGS	TNGS	TNGS	TNGS	
Protein function impact	LOF	LOF	LOF	GOF	GOF		Likely GOF	Likely LOF	Likely LOF	Likely GOF	Likely GOF	Likely GOF	Likely GOF	Likely GOF	Likely GOF	Likely GOF	Likely GOF	Likely LOF	Likely LOF	Likely LOF	Likely LOF	Likely GOF	Likely GOF	Likely GOF	Likely GOF	Likely GOF	
Zygosity	Heterozygous Compound	Heterozygous	Heterozygous	Heterozygous	Heterozygous		Heterozygous	Homozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Compound	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	Heterozygous	
Protein level mutation	M886T R112L	H753Q	200528	G13C	G13C		C161Y	A24V	Y284C	p.V661M	pl6T	p.S3271	1875M	A531S	D1137V	443L	R220H	NA	H1142Q	Y198C	T233S	R226H	p.E205K	NA	R183H	Q500R	
cDNA mutation	2657T>C 335G>T	2259T>A	1490×A	37 G>T	37 G>T		c.482G>A	71C>T	851A>G	c.1981G>A	c.17T>C	c.980G>T	2625C>G	1591G>T	3410A>T	1328C>T	659G>A	3284_3285delAG - 2ex 26	3426C>G	593A>G	698C>G	677G>A	c.613G>A	c.1226+8G>A	548G>A	1499A>G	
Genetic	RAG1		INFRSF0 somatic	KRAS somatic	KRAS somatic		TNFR2	IFNAR1*	TGFBR2	JAK1	JAK1	JAK2	PLCG2	TRAF3	CARD11	ARHGEF4	PTPN11	PARP4		RIPK2	RIPK2	APAF1	NFATC1	NFATC1	IKZF1	IKZF1	
Patients	P29		P30	P31	P32	Probably pathogenic	P33	P34	P35	P36	P37	P38	P39	P40	P41	P42	P43	P44		P45	P46	P47	P48	P49	P50	P51	050

CADD, Combined Annotation-Dependent Depletion; CDNA, complementary DNA; MAF, minor allele frequency; NA, not applicable; OMIM, Online Mendelian Inheritance in Man; SIFT, Sorting Intolerant from Tolerant; TNGS, targeted next-generation sequencing; WB, western blot; WES, whole-exome sequencing. See Table 2 for expansion of other abbreviations. *Patient PBMCs were shown to have impaired induction of interferon-stimulated genes (n = 6) after stimulation by IFN« comparatively to control cells.

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Table 3. (continued)

	M+, N = 52	M-, N = 28	P, M+ vs M-	Pathogenic, N = 32	Probably pathogenic, N = 20	P, pathogenic vs probably pathogenic
Consanguinity, % (n)	16 (9)	7 (2)		19 (6)	15 (3)	
Immune manifestations in first-degree relatives, $\%$ (n)	37 (19)	18 (5)	.08	44 (14)	25 (5)	
Immune manifestations in extended relatives, $\%$ (n)	50 (26)	30 (8)	90.	57 (18)	38 (8)	
Sex ratio (F/M)	0.7	0.6		0.8	0.7	
Median age at first episode of cytopenia (min-max), y	5.2 (0.5-15.2)	7.1 (0.3-15.6)		6.8 (0.5-15.2)	4.1 (0.8-14.7)	
Associated immunopathological manifestations, % (n)	92 (48)	64 (18)	.001	97 (31)	85 (17)	
Median age at first immunopathologic manifestation (min-max), y	5.3 (0.3-14.9)	8.6 (1.6-17.2)		5.0 (0.3-14.9)	5.5 (0.8-17.7)	
Median no. of immunopathologic manifestations (min-max)	2.5 (0-8)	1 (0-6)	.007	3 (0-8)	2 (0-4)	
Median age at immunological disease onset (ES or IM) (min-max), y	3.3 (0.3-14.7)	6.0 (0.3-15.6)		3.9 (0.3-14.1)	2.6 (0.8-14.7)	
Lymphoproliferation, % (n)	58 (30)	35 (10)	.06	63 (20)	50 (10)	
Hypogammaglobulinemia, % (n)	56 (29)	29 (8)	.02	60 (19)	50 (10)	
Autoimmuna/autoinflammatoru organ dicasca % (n)	56 (20)	37 (0)	50	43 (20)	15 (0)	
Truction of directive tract manifestations %	JO (27) 13	02 (V) 2	190	10 (20)		800
Live manifectations %	<u>0</u> C	2 C	- 00.	<u>4</u> 0		. 000
Lung manifestations, & Neurologic manifestations. %	<u>></u> ∞	2 2		~ \C	- m	0.
Skin manifestations, %	6	4		ъ Ю	-	
Rheumatologic manifestations, %	т	۲-		2	1	
Endocrinologic manifestations, %	3	0		2	1	
Atopy, % (n)	25 (13)	7 (2)	.07	28 (9)	20 (4)	
SLE biomarkers,* % (n)	17 (9)	36 (10)		16 (5)	20 (4)	
ALPS biomarkerst % (n)	19 (10)	7 (3)		28 (9)	2 (1)	20.
Severe or recurrent infections, % (n)	50 (26)	43 (12)		57 (18)	40 (8)	
Median follow-up from first cytopenia (min-max), y	9.6 (0.9-25.0)	7.0 (0.2-26.5)		9.2 (0.9-25.0)	10.2 (4.2-20.8)	
Need for second-line treatments, % (n)	81 (42)	63 (18)		88 (28)	(14) 22	
Median no. of second-line treatments (min-max)	2 (0-9)	1 (0-5)	.02	2 (0-9)	5 (0-6)	
Deaths, % (n)	12 (6)	(0) 0	.086	12 (4)	10 (2)	

Table 4. Clinical presentation of 80 pES patients according to genetic subgroups

Bold values represent P < .05, considered statistically significant.

ALPS, autoimmune lymphoproliferative syndrome; F, female; IM, immunopathological manifestation; M, male; M+, patients with mutations; M-, patients without detected mutations; SLE, systemic lupus erythematosus. *SLE biomarkers: antituclear antibodies titer >1/160 on 2 separate samples, isolated significant autoantibodies.

TALPS biomarkers: persistent hypergammaglobulinemia (at least 2 SD over the mean for age), high counts of circulating TCR ab CD4-CD8- double-negative T lymphocytes

Discussion

The results of the present prospective study showed that pathogenic or probably pathogenic genetic variants (most of which were germline mutations) can be detected in the majority of children with ES. The generation of these novel data was made possible by accessing a large national prospective cohort of patients with pES and NGS resources.44,45 It was already known that children with monogenic inherited disorders of the immune system (ie, PIDs) are 80- to 120-fold more likely to develop chronic immune cytopenia than an age-matched population is.^{3,46,47} In adult patients, Michel et al⁴⁸ have shown that ES was a secondary condition in 56% of cases (mainly as a results of malignancies and autoimmune diseases); a PID was identified in only 9% of cases. Given the estimated prevalence of PIDs, it has been predicted that at least one-third of children with chronic immune cytopenia have a causal genetic disorder.³ We therefore decided to perform a genetic analysis in a large group of patients from the French OBS'CEREVANCE pES cohort. We did not apply any a priori selection criteria other than consent by the family and/or patient. The study group was similar (but not identical) to the overall pES cohort. Our results show that in 65% of the patients, pES was strongly associated with the detection of significant genetic variants. These results will be addressed for each of 3 observed groups, that is, patients with known genetic variants, patients with new potentially pathogenic genetic variants, and patients with no detected pathogenic or probably pathogenic genetic variants.

Unsurprisingly, a number of patients carried mutations known to be strong risk factors for autoimmune cytopenia. These included germline and somatic TNFRSF6 mutations causing ALPS,^{4,49} mutations in the CTLA4 and LRBA genes,^{16,20} GOF mutations in STAT3⁵⁰ or PiK3CD,¹⁷ hypomorphic LOF mutations in RAG1,⁵¹ and somatic mutations in CBL and KRAS associated with RALD.^{52,53} Despite the presence of associated clinical manifestations in some of the patients, it is noteworthy that none of them (with the exception of cases of ALPS) had previously received a firm diagnosis. Although autoimmune cytopenia is not uncommon in these conditions, it is not always present.^{16-18,20,27,50-55} It is also known that some TNFRSF6 and CTLA4 mutations do not have full penetrance.^{18,54} Hence, additional predisposing factors (possibly environmental and/ or genetic factors) may favor the occurrence of ES in these children.^{1,2} In this respect, the high overall heritability of the genomic variations in many pediatric autoimmune conditions (although not in autoimmune cytopenia) identified in genome-wide association studies might represent a "favorable" background on which a critical gene variant (CTLA4, LRBA) causes autoimmunity.² The high proportion of patients with a family history of autoimmune disease (42% when considering first-degree relatives, and 57% when considering extended families, values that are greater than expected for diseases with autosomal-dominant inheritance) fits with this view. Based on these results, performance of the same genetic analysis in the patients' relatives is an obvious next step.

The putative causal role of the probably pathogenic variants identified in the second ("probably pathogenic") group of patients must be scrutinized with caution. Strict criteria were used (in terms of both allele frequency and predicted pathogenicity) to select the detected variants. The gene product's



Figure 2. A box plot representation of the number of immunopathologic manifestations per patient, as a function of the genetic subgroup in 80 pES patients (the M+, M-, "pathogenic" and "probably pathogenic" groups). The bottom of the box marks the 25th percentile, the median line marks the 50th percentile, and the top of the box marks the 75th percentile. The bottom of the vertical line indicates the minimum and the maximum. The vertical lines at the top and bottom indicate outlying data points.

function (if known from the literature) was also considered. It is noteworthy that variants in 5 genes were found in unrelated families, further suggesting causality. Other variants in some of these genes are known to be pathogenic in contexts other than autoimmunity.^{28,29,39,56} The observation that the phenotypic characteristics of this patient group were very similar to those of patients carrying known pathogenic variants also argues in favor of the variants' pathogenicity. Relative to the M- group, patients in the M+ group were more likely to have more additional immunopathologic events. Twenty-five patients developed hypogammaglobulinemia after rituximab treatment, as previously reported.⁵⁷ Still, given its efficacy, rituximab should probably be considered as the first-line treatment in pES. However, a careful evaluation is warranted before rituximab initiation and during follow-up. All of the associated immunopathologic events occurred earlier in the M+ group, required more frequent immunosuppressive therapy, and tended to be associated with a poorer prognosis (ie, poorer survival). However, it must be noted there were some phenotypic differences between patients with known mutations and patients with probably pathogenic variants; the latter group were slightly (but not significantly) less likely to have a family history of immunopathology, a personal history of digestive tract disease and/or ALPS biomarkers. The latter findings may be explained, respectively, by (i) the presence of a number of ALPS patients in the first group, and (ii) the presence of cases at risk of inflammatory bowel disease in the same group (ie, LRBA and CTLA4 deficiencies, and STAT3 GOF mutations).18,20,27 We therefore hypothesize that the "probably pathogenic" variants detected here (see below) are risk factors that are mostly (but not exclusively) related to the control or occurrence of antibody-mediated autoimmunity.

The detection of "probably pathogenic" variants may broaden the field of genetic predisposition to autoimmunity. Several



mechanisms can be considered: (i) impaired suppression of conventional T cell proliferation by regulatory T cells,⁵⁸⁻⁶⁰ (ii) GOF or LOF variants that increase immune effector function (in various ways) downstream of cytokines and antigen-specific B- or T-cell receptors or co-receptors, or (iii) pathologic changes in immune responses in the context of infection or the control of skin and gut microbiotas.^{28,32,39,61-69} A fourth potential mechanism relates to lymphocyte apoptosis, which is likely to be defective in patients with *RIPK2* and *APAF1* mutations.^{70,71} All of these variants must now undergo extensive functional validation, in order to characterize the mechanism predisposing to auto-immunity. It will probably also be of value to extend screening for the variants described here to other children with ES and to pediatric cohorts with other autoimmune diseases.

The present results do not imply that the known genetic variants (within the M+ group) caused an autoimmune disease with monogenic Mendelian inheritance. The familial segregation of these mutations will now also have to be studied. It is important to bear in mind that no other significant immune gene variants were detected in these patients. Nevertheless, this observation does not rule out a role of other gene variants in general and regulatory variants (expressed quantitative trait loci) in particular.⁷²

Lastly, genetic factors may be less critical in the third group of patients with pES, that is, those in whom no significant genetic variants were found with our genetic testing strategy. Overall, this group of patients is distinct from the other 2 groups because its members were less likely to present with additional immunopathologic features (including allergy) and had somewhat less severe ES. Nevertheless, a family history of autoimmunity was not uncommon, suggesting that polygenic genetic variations are involved, given the high heritability of autoimmunity in children.²

Our genetic approach had some limitations, and may have failed to detect variants in some cases (ie, false-negatives). Most of the patients were screened using tNGS; we cannot rule out the possibility that mutations in genes not included in the panel, intronic variants or somatic mutations of known genes not detected by the present testing strategy could be involved in the M- group. Further analyses (such as whole-genome sequencing or the in-depth sequencing of candidate genes in lymphocyte subsets⁷³) should therefore be performed before a genetic cause is ruled out.

In conclusion, our results suggest that wide-ranging genetic screening should be offered to children with ES because the findings have prognostic significance and may thus influence treatment choices. Indeed, we identified 29 patients (36%) with mutations that might help the physician to choose a targeted treatment of severe autoimmune cytopenia, as shown (for instance) by the use of (i) a mechanistic target or rapamycin inhibitors in patients with ALPS⁷⁴ or an activated phosphoinositide 3-kinase delta (PIK38) syndrome,75 (ii) CTLA-4 fusion protein therapy in CTLA-4- and LRBA-deficient patients,18,20 (iii) the potential use of JAK inhibitors to targeting GOF variants in JAK1 or JAK2,³⁹ and possibly (iv) calcineurin inhibitor therapy in patients with NFATC1 variants.⁷⁶ The present study raises new questions about the genetic background of pES and, more widely, autoimmune diseases in children. It is now justifiable to extend genetic testing to all children with chronic multilineage cytopenia, their family members, and other cohorts of pediatric patients with autoimmune diseases. Our study also provides a basis for gaining further insights into the mechanistic aspects of controlling reactivity to self, along with the many checkpoints already known.77,78 We expect this type of study to provide additional insight into the pathophysiology of autoimmunity, together with important patient-level information with prognostic and therapeutic value.

Figure 3. Patients with pES free of additional immunopathologic manifestations, as a function of the genetic subgroup.

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Authorship

Contribution: F.R.-L., A.F., and B.N. designed the study; J.H., N.A., H.F., and F.R.-L. acquired the data; J.H., N.A., H.F., B.N., A.F., and F.R.-L. drafted the manuscript; and all of the authors analyzed and interpreted the data and revised the manuscript for critical content.

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A complete list of the members of the French Reference Center For Pediatric Autoimmune Cytopenia (CEREVANCE) appears in the supplemental Appendix.

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Footnotes

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