ORIGINAL ARTICLE



Human Inborn Errors of Immunity: 2022 Update on the Classification from the International Union of Immunological Societies Expert Committee

Stuart G. Tangye^{1,2} • Waleed Al-Herz³ · Aziz Bousfiha⁴ · Charlotte Cunningham-Rundles⁵ · Jose Luis Franco⁶ · Steven M. Holland⁷ · Christoph Klein⁸ · Tomohiro Morio⁹ · Eric Oksenhendler¹⁰ · Capucine Picard^{11,12} · Anne Puel^{13,14} · Jennifer Puck¹⁵ · Mikko R. J. Seppänen¹⁶ · Raz Somech¹⁷ · Helen C. Su⁷ · Kathleen E. Sullivan¹⁸ · Troy R. Torgerson¹⁹ · Isabelle Meyts²⁰

Received: 20 March 2022 / Accepted: 2 May 2022 / Published online: 24 June 2022 © The Author(s) 2022

Abstract

We report the updated classification of inborn errors of immunity, compiled by the International Union of Immunological Societies Expert Committee. This report documents the key clinical and laboratory features of 55 novel monogenic gene defects, and 1 phenocopy due to autoantibodies, that have either been discovered since the previous update (published January 2020) or were characterized earlier but have since been confirmed or expanded in subsequent studies. While variants in additional genes associated with immune diseases have been reported in the literature, this update includes only those that the committee assessed that reached the necessary threshold to represent novel inborn errors of immunity. There are now a total of 485 inborn errors of immunity. These advances in discovering the genetic causes of human immune diseases continue to significantly further our understanding of molecular, cellular, and immunological mechanisms of disease pathogenesis, thereby simultaneously enhancing immunological knowledge and improving patient diagnosis and management. This report is designed to serve as a resource for immunologists and geneticists pursuing the molecular diagnosis of individuals with heritable immunological disorders and for the scientific dissection of cellular and molecular mechanisms underlying monogenic and related human immune diseases.

 $\textbf{Keywords} \ \ Inborn\ errors\ of\ immunity\cdot immune\ dysregulation\cdot primary\ immunodeficiencies\cdot autoinflammatory\ disorders\cdot IUIS\ Committee\ update$

Introduction

Inborn errors of immunity (IEI) are caused by damaging germline variants in single genes. IEI present clinically as increased susceptibility to infections, autoimmunity, autoinflammatory diseases, allergy, bone marrow failure, and/or malignancy. While individually rare, the aggregated number of individuals with an IEI represents a significant health burden [1]. Genetic variants cause disease by altering the encoded gene product, such as by abolishing or reducing protein expression and function (null/hypomorphic) or modifying the protein to acquire gain-of-function (GOF) [2–5]. Mechanisms of disease in IEI depend on the nature of the

variant as well as the mode of inheritance. Thus, monoallelic variants can cause disease by haploinsufficiency, negative dominance, or GOF. In contrast, biallelic genetic lesions (homozygous, compound heterozygous) cause autosomal recessive (AR) traits by loss of expression, loss of function (LOF), GOF, or even neomorphic function of the encoded protein, while X-linked recessive traits arise from LOF or GOF variants on the X chromosome, either in hemizygosity in males, or homozygous state in females.

The fact that some monogenic variants are pathogenic clearly highlights the non-redundant and fundamental roles of individual genes and proteins, and associated pathways and cell types, in the development and function of leukocytes and non-hematopoietic cells that contribute to immune homeostasis and host defense [6, 7]. Thus, IEI represent an elegant model linking defined monogenic defects with clinical phenotypes of immune dysregulation. IEI have also revealed mechanisms of disease pathogenesis in, and

Stuart G. Tangye s.tangye@garvan.org.au

Extended author information available on the last page of the article



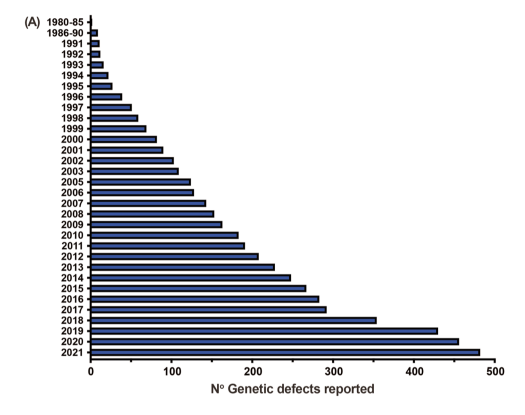
enabled the implementation of gene- or pathway-specific therapies for the treatment of, rare and common conditions and established fundamental aspects of human immunology [8–10]. Thus, the study of IEI has enabled profound advances in molecular medicine and human biology.

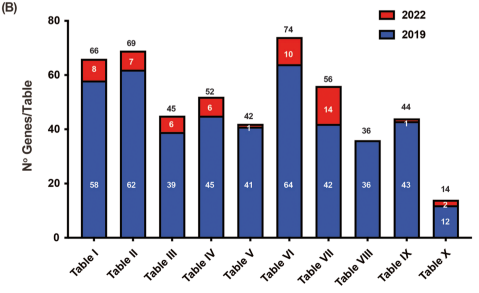
Since 1970, an international expert committee comprising pediatric and adult clinical immunologists, clinician/scientists and researchers in basic immunology — initially under

the auspices of the World Health Organization and currently the International Union of Immunological Societies (IUIS) — has provided the clinical and research communities with an update of genetic causes of immune deficiency and dysregulation https://iuis.org/committees/iei/ (Fig. 1A).

IEI are currently categorized into 10 Tables, with subtables segregating groups of disorders into overlapping phenotypes. These tables describe the following: combined

Fig. 1 Accumulative discovery of novel inborn errors of immunity: 1980-2022. (A) The number of genetic defects underlying monogenic immune disorders as reported in the indicated year. (B) The number of pathogenic variants listed in each Table of the IUIS IEI committee 2022 report. The numbers in each column correspond to the number of genes reported in the 2019 IUIS update (blue bars) [4, 5], the number of new genes for each Table contained in this report (red bars), and the total number of genes for each Table (black number). Note: The 14 conditions listed for Table 10 are either phenocopies of germline IEI due to somatic variants or neutralizing autoAbs. Somatic variants in UBA1 are also listed in Table 10, although there is currently no IEI resulting from germline UBA1 variants [97]







immunodeficiencies (Table 1, 3 subtables); combined immunodeficiencies with syndromic features (Table 2; 9 subtables); predominantly antibody deficiencies (Table 3; 3 subtables); diseases of immune dysregulation (Table 4; 7 subtables); congenital defects of phagocytes (Table 5; 4 subtables); defects in intrinsic and innate immunity (Table 6; 9 subtables); autoinflammatory diseases (Table 7; 3 subtables); complement deficiencies (Table 8); bone marrow failure (Table 9), and phenocopies of inborn errors of immunity (Table 10) (Fig. 1B) [5].

The committee strives to publish an updated report approximately every 2 years to consolidate advances and catalog current IEIs (Fig. 1A) [5]. While COVID-19 has delayed producing this report in the desired timeframe, it has also uncovered several new IEI — some of these are highlighted below. Many genetic variants related to IEI have been reported recently. Rather than including every candidate gene reported in the peer-reviewed scientific literature, the committee applies stringent criteria to classify gene defects as novel causes of IEI [11]. These criteria include:

- The patient's candidate genotype is monogenic and does not occur in individuals without the clinical phenotype (acknowledging that some conditions have incomplete penetrance).
- Experimental studies establish that the genetic variant impairs, destroys, or alters expression or function of the gene product.
- 3. The causal relationship between the candidate genotype and the clinical phenotype must be confirmed via a relevant cellular phenotype, including where possible rescue of a functional defect [11].

These criteria can be met by publication of multiple cases from unrelated kindreds, including detailed immunologic data, or publication of very few — even single — cases for whom compelling mechanistic data are provided, often revealed from complementary studies in animal or cell culture models. We also considered whether sufficient justification was provided to exclude alternative candidate gene variants identified in single cases, the depth of the clinical descriptions of affected individuals, and the level of immune and mechanistic characterization. This 2022 update and the accompanying "Phenotypical IUIS Classification" publications are intended as resources for clinicians and researchers, as well as guiding the design of panels used for targeted gene sequencing to facilitate genetic diagnoses of IEI. Here, we summarize data on the genetic cause of 55 novel IEI, and 1 phenocopy due to autoantibodies, that have been assessed since the previous update [5] (Supplementary Table 1). Remarkably, 15 of the 55 novel IEI have come from the identification and extensive work-up of single patients. Two themes that are expanded in this new set of genes are narrow infection susceptibility and immune dysregulation, which collectively account for over half of the phenotypes associated with these new genetic etiologies of IEI. This paper increases the number of known genetic defects identified as causing IEI to 485 (Fig. 1A, B; see all Tables and Supplementary Table 1).

Novel Inborn Errors of Immunity

Novel gene defects have been found for most categories of IEI, including novel causes of:

- Combined immunodeficiencies (*LCP2* (SLP76) [12], *PAX1* [13, 14], *ITPKB* [15]; *SASH3* [16, 17], *MAN2B2* [18], *COPG1* [19], *IKZF2* [20–23], *CHUK* [24], *IKZF3* [25, 26], *CRACR2A* [27], *CD28* [28]) (Table 1; Supplementary Table 1);
- Combined immunodeficiencies with syndromic features (MCM10 [29, 30], IL6ST [31–33], DIAPH1 [34]) (Table 2; Supplementary Table 1);
- B cell deficiencies, agammaglobulinemia, or hypogammaglobulinemia (FNIP1 [35, 36], SP1I [37], PIK3CG [38, 39], POU2AF1 [40], CTNNBL1 [41], TNSRSF13 [42]) (Table 3; Supplementary Table 1);
- Immune dysregulation (*RHOG* [43], *SOCS1* [44–46], *PDCD1* [47], *ELF4* [48, 49], *TET2* [50], *CEBPE* [51], *IKZF1* GOF [52]) (Table 4; Supplementary Table 1)
- neutropenia *CXCR2* [53, 54] (Table 5, Supplementary Table 1)
- innate immune defects resulting in susceptibility to mycobacterial/bacterial (*TBX21* [55, 56], *IFNG* [57], *TLR8* [58, 59]), viral (*NOS2* [60], *SNORA31* [61], *ATG4A*, *MAP1LC3B2* [62], *ZNFX1* [63–65], *TLR7* [66–68]), and/or fungal infections (*MAPK8* [69]) (Table 6; Supplementary Table 1);
- Autoimmune/autoinflammatory disorders (*TMEM173* [70], *LSM11*, *RNU7-1* [71], *CDC42* [72–78], *STAT2* [79, 80], *ATAD3A* [81], AR *TBK1* [82], *C2orf69* [83, 84], *RIPK1* [85, 86], *NCKAP1L* [87–89], *SYK* [90], *HCK1* [91], *IKBKG* [92–94]); *PSMB9* [95, 96]; and somatic variants in *UBA1* [97]) (Table 7, 10, Supplementary Table 1);



- Bone marrow failure (MECOM1) [98, 99] (Table 9; Supplementary Table 1); and
- Phenocopies of IEI (somatic variants in *TLR8* [58], autoAbs against type 1 IFNs [100–104]) (Table 10; Supplementary Table 1).

Novel IEI Phenocopy Known IEI, Confirming Critical Pathways for Immune Function

Some of these novel genetic findings link common clinical phenotypes that converge on a shared pathway. Examples in this update include:

- SLP76, encoded by LCP2, is part of the TCR signalo-some, interacting with or being downstream of ZAP70, LCK, LAT and ITK [105]. Thus, the phenotype of AR SLP76 deficiency overlaps substantially with that of individuals with mutations in these genes [12].
- MCM10 is a component of the DNA replication machinery of mammalian cells and forms part of multimeric/multiprotein "replisome" complexes [106]. Thus, biallelic mutations in MCM10 result in a clinical phenocopy of AR MCM4 or GINS1 variants [29, 30], which also encode key proteins involved in DNA replication [106].
- The non-redundant role of IFNγ-mediated immunity in protection against mycobacterial infection was established by identifying individuals with mutations in not only *IFNG* itself [57], but also *TBX21* [55], the transcription factor that regulates IFNγ, who develop Mendelian susceptibility to mycobacterial disease. T-bet deficiency also resulted in upper airway inflammation and Th2 dysregulation [56], further highlighting immune regulation mediated by opposing functions of transcription factors in T cells with distinct fates (Th1 vs Th2).
- Individuals with complete gp130-deficiency due to bi-allelic mutations of *IL6ST* [33], or dominant negative heterozygous variants of *IL6ST* [31], present with eczema, hyper-IgE, and eosinophilia, similar to individuals with AD hyper-IgE syndrome due to dominant negative mutations in *STAT3* or AR mutation in *ZNF341* [107]. These findings from the different genotypes indicate a key role for IL-6 signaling, via STAT3/ZNF341, in regulating hyper-IgE and atopy.

- Store-operated calcium entry via Ca²⁺-release activated Ca²⁺ channels (CRAC) enable transfer of Ca²⁺ across cell membranes following activation of surface receptors, thereby eliciting Ca²⁺ flux and initiation of key intracellular signals [108]. Bi-allelic LOF variants in *STIM1* or *ORA1* disrupt Ca²⁺ flux, thereby impairing lymphocyte activation following engagement of antigen receptors, resulting in combined immunodeficiencies [108]. The first report of an individual with compound heterozygous inactivating variants in *CRACR2A* provides further insight into the importance of Ca²⁺-dependent signaling in immune cells [27].
- The IKAROS family of proteins IKAROS, AIOLOS, and HELIOS interacts with one another as homodimers, heterodimers, or heterotrimers to regulate immune cell development and function [109]. While variants in *IKZF1* encoding IKAROS have been previously reported [5, 109], individuals have now been identified with pathogenic variants in *IKZF2* (HELIOS) [20–23] and *IKZF3* (AIOLOS) [25, 26], as well as GOF variants in *IKZF1* [52]. While these genotypes present with some distinct clinical phenotypes, there is also substantial overlap, such as B cell deficiency, hypoor agammaglobulinemia, recurrent infections, and predisposition to B cell malignancy.

One Gene, Several Phenotypes

The discovery of novel IEI continues to demonstrate that distinct types of variants (GOF, LOF, mono-allelic, biallelic, exon splicing) in the same gene can cause disparate clinical conditions. This update includes AR and AD forms of IKZF2 (HELIOS) [20–23] and IL6ST [31–33] deficiency, as well as AD *RIPK1* LOF [85, 86], AR GOF *TMEM173/* STING [70], AR LOF TBK1 [82], and mono-allelic IKZF1 GOF [52] variants which complement previous reports of AR RIPK1 deficiency, AD GOF TMEM173/STING, AD TBK1 deficiency, and mono-allelic IKZF1 inactivating variants, respectively [5]. AR GOF variants in *CEBPE* also represent a novel IEI [51]. Notably, these variants resulted in neomorphic function of the C/EBPE transcription factor, causing dysregulated expression of >400 genes, ~15–20% of which are not normally targeted by C/EBPe [51]. This may represent the prototype for neomorphic variants causing IEI.



Intriguingly, specific variants in STAT2 or IKBKG — which are already well-known to cause IEIs — have recently been reported that cause very distinct phenotypes from those previously associated with pathogenic variants in these genes. STAT2 plays a ying/yang role in type 1 IFN signalling. Thus, it is responsible for not only inducing, but also restraining, responses elicited via IFNαR1/2 complexes [110]. This regulatory role of STAT2 is mediated by binding to and recruiting USP18 to IFNαR2, which then prevents further recruitment of JAKs to type 1 IFN receptors, thereby attenuating IFN α signalling [110]. Bi-allelic variants in STAT2 that specifically affect amino acid R148 (STAT2R148Q/W) have now been reported [79, 80]. These STAT2^{R148Q/W} variants are LOF for binding to USP18 [79, 80, 110]. Consequently, STAT2^{R148Q/W} prevents USP18-mediated restraint of type 1 IFN signalling. It is important to appreciate that while STAT2R148Q/W is not intrinsically GOF, the net outcome of loss of STAT2-mediated regulation of type 1 IFN signalling is reminiscent of other Mendelian IFN-opathies. Indeed, STAT2R148Q/W is a phenocopy of USP18 deficiency [110], which is clearly distinct from severe susceptibility to some live attenuated viral vaccines and viral infections typical of individuals with null/nonsense mutations in STAT2 [110]. Lastly, unique variants in *IKBKG* that result in deletion of exon 5 were found to cause an autoinflammatory disease which is also very different from ectodermal dysplasia and immunodeficiency that is typically associated with hypomorphic IKBKG variants that impair NEMO expression and/ or function [92–94].

Somatic/mosaic disease-causing mutations in *TLR8* [58] and *UBA1* [97] have also been identified, even though the pathogenic alleles were detected in only 5–30% of most blood cells (*TLR8*) [58] or 50–85% of myeloid cells but not in lymphocytes of fibroblasts (*UBA1*) [97]. These findings are an important reminder to consider the nature of genetic variants identified from unbiased next-generation sequencing, recognizing multiple mechanisms of pathogenicity for the same gene. This is highlighted by at least 40 genes having multiple entries in the current update to reflect these distinct modes of disease pathogenesis (Supplementary Table). This also emphasizes the crucial need to undertake indepth in vitro functional validation of any variant considered to be potentially pathogenic. Alternatively, it

signifies the difficulty in excluding a candidate pathogenic variant without functional testing. It also underscores the need to consider variants detected at low allelic frequencies that may represent somatic/mosaic, rather than germline, variants. These findings also predict that somatic variants in key immune genes will be frequently discovered as causes of novel IEI in the nottoo distant future [111].

IEI Define Specific Roles for Known Genes and Reveal Immune-Specific Functions of Novel Genes

One of most profound outcomes of discovering the genetic cause of an IEI is the ability to ascribe unequivocally non-redundant, as well as redundant, functions to a specific gene in human immunity. Classic examples of this are the fundamental requirement for *IL2RG* in humans for the development of T and NK cells, but not B cells, and the essential role of STAT3 for CD4⁺ T cell differentiation into Th17 cells and subsequent host defense against fungal infections, but not for the generation of most other CD4+ T cell effector populations [112]. Findings included in this update confirm data from mice on the importance of FNIP1 and SPI1 (encoding PU.1) during human B cell development [35-37] and the fundamental regulatory role of PD-1 (encoded by *PDCD1*) in human immune function [47]. However, and perhaps counter to all expectations and immunology dogma relating to T cell co-stimulation, CD28 is required for host defense against HPV but is largely redundant in the face of other infectious pathogens [28]. Who would have thought!

The latest IEI have also revealed critical roles for genes not previously strongly associated with immune regulation and/or host defense. For instance, we have now learned that:

- The SH3-domain containing protein SASH3 contributes to B and T cell developments [16, 17].
- *ZNFX1*, a member of an RNA helicase superfamily, plays a dual role in human immunity, including in innate immune responses against viruses, bacteria, mycobacteria, and fungi, as well as in restraining type 1 IFN-mediated inflammation [63–65].



- The small nucleolar RNA *SNORA31* plays a critical role in CNS-intrinsic immunity against HSV-2 infection, likely via production of type 1 IFN, yet the exact mechanism remains unknown [61].
- The hitherto uncharacterized protein-coding gene *C2orf69* has a multitude of roles across numerous biological systems, including regulating autoinflammation [83, 84].

The discovery of these novel IEIs provides opportunities to further extend our understanding of human immunity and immune regulation.

SARS-CoV2 and Inborn Errors of Immunity

The emergence of novel pathogens poses potential health risks to the general population due to the lack of substantial pre-existing immune memory. More critically though, individuals with specific germline genetic variants — causing known and unknown IEIs - may be at greater risk of experiencing more severe disease following infection than the general population. The COVID-19 pandemic has indeed revealed genes and pathways essential for anti-SARS-CoV2 immunity. Genomic studies discovered that ~2–3% of cases of severe life-threatening SARS-CoV2 infection resulted from germline LOF/LOE variants in the type 1 IFN signaling pathway: TLR3, UNC93B1, TICAM1, TBK1, IRF3, IRF7, IFNAR1, and IFNAR2 [113]. These findings are reminiscent of earlier studies that identified variants in these genes in individuals susceptible to life-threatening infections with other viruses, including influenza virus, HSV-1, and live viral vaccines [114]. Hemizygous deleterious variants have also been identified in TLR7 in ~1% of males who developed severe/fatal COVID-19 [66-68]. Thus, X-linked TLR7 deficiency represents a novel IEI predisposing to severe COVID-19.

The importance of type 1 IFN in anti-SARS-CoV2 immunity was also realized by the finding that ~10–20% of patients with severe COVID-19 have high levels of neutralizing serum autoantibodies (autoAbs) against type 1 IFNs; these were not detected in asymptomatic infected individuals [100–104]. Collectively, these studies defined a non-redundant role for type 1 IFNs in host defense against SARS-CoV2 infection and established that autoAbs against type 1 IFN phenocopy an IEI.

Conclusions

The goals of the IUIS Expert Committee on IEI are to increase awareness, facilitate recognition, promote optimal treatment, and support research in the field of clinical immunology. Since the last IEI update, we have continued to witness the ongoing rapid identification, and molecular, biochemical, and cellular characterization, of genetic variants that cause human diseases by disrupting host defense or immune regulation. The 55 novel gene defects reported here bring to total number of IEI to 485 (Fig. 1A, B), thus underscoring the power of next-generation sequencing technologies and sophisticated functional validation of candidate pathogenic variants to (1) identify novel gene defects underlying human disease, (2) elucidate mechanisms of disease pathogenesis, (3) define non-redundant functions of key genes in human immune cell development, host defense and immune regulation, (4) expand the immunological and clinical phenotypes of IEI, and (5) implement gene-specific therapies. These fundamental discoveries continue to highlight the critical contributions of IEI to our broader understanding of basic, translational, and clinical immunology, as well as molecular medicine. And we will no doubt observe novel insights into basic and clinical immunology with the next wave of novel IEIs.



Table 1 Immunodeficiencies affecting cellular and humoral immunity

	1. T-B+ Severe Combined Immune Deficiency (SCID)											
Disease	Genetic defect	Inheritance	ОМІМ	T cells	B cells	lg	Associated features					
γc deficiency (common gamma chain SCID, CD132 deficiency)	IL2RG	XL	308380	Very low	Normal to high	Low	Low NK					
JAK3 deficiency	JAK3	AR	600173	Very low	Normal to high	Low	Low NK					
IL7Rα deficiency	IL7R	AR	146661	Very low	Normal to high	Low	Normal NK					
CD45 deficiency	PTPRC	AR	151460	Very low	Normal	Low	Normal γ/δ T cells					
CD3δ deficiency	CD3D	AR	186790	Very low	Normal	Low	Normal NK, no γ/δ T cells					
CD3 ₈ deficiency	CD3E	AR	186830	Very low	Normal	Low	Normal NK, no γ/δ T cells					
CD3ζ deficiency	CD3Z	AR	186780	Very low	Normal	Low	Normal NK, no γ/δ T cells					
Coronin-1A deficiency	CORO1A	AR	605000	Very low	Normal	Low	Detectable thymus					
LAT deficiency	LAT	AR	602354	Normal to low	'Normal to low	High	Typical SCID or combined immunodeficiency, the latter with adenopathy, splenomegaly, recurrent infections, autoimmunity					
SLP76 deficiency (1 patient)	LCP2	AR	619374	Reduced	Normal	High IgM, low IgA	Early-onset skin abscesses, rash, recurrent infections, autoimmunity					

	2. T-B- SCID												
Disease	Genetic defect	Inheritance	ОМІМ	T cells	B cells	lg	Associated features						
RAG deficiency	RAG1	AR	<u>179615</u>	Verslau	Variani	Desired	Normal NK cell number, but increased						
	RAG2	AR	<u>179616</u>	Very low	Very low	Decreased	risk of graft rejection, possibly due to activated NK cells						
DCLRE1C (Artemis) deficiency	DCLRE1C	AR	605988	Very low	Very low	Decreased	Normal NK cell number, but increased risk of graft rejection, possibly due to activated NK cells, radiation sensitivity						
DNA PKcs deficiency	PRKDC	AR	<u>615966</u>	Very low	Very low	Variable	Normal NK, radiation sensitivity, microcephaly						
Cernunnos/XLF deficiency	NHEJ1	AR	<u>611290</u>	Very low	Very low	Decreased	Normal NK, radiation sensitivity, microcephaly						
DNA ligase IV deficiency	LIG4	AR	601837	Very low	Very low	Decreased	Normal NK, radiation sensitivity, microcephaly						
Adenosine deaminase (ADA) deficiency	ADA	AR	608958	Very low	Low, decreasing	Low, decreasing	Low NK, bone defects, may have pulmonary alveolar proteinosis, cognitive defects						
AK2 defect	AK2	AR	103020	Very low	Very Low	Decreased	Reticular dysgenesis with neutropenia; deafness						
Activated RAC2 defect	RAC2	AD GOF	602049	Very low	Very Low	Low, poor specific antibody responses	Recurrent bacterial and viral infections, lymphoproliferation; neutropenia						

	3. Combin	ed Immunodefic	ciency (CID),	Generally Less			
Disease	Genetic defect	Inheritance	ОМІМ	T cells	B cells	lg	Associated features
CD40 ligand (CD154) deficiency	CD40LG	XL	308230	Normal to low	slgM*lgD* naïve B cells present; IgG*, IgA*, IgE* memory B cells	IgM normal or high, other Ig isotypes low	Severe and opportunistic infections, idiopathic neutropenia; hepatitis and cholangitis, Cryptosporidium infections, cholangiocarcinoma; autoimmune blood cytopenias; peripheral neuroectodermal tumors
CD40 deficiency	CD40	AR	606843	Normal	absent		Neutropenia, opportunistic infections, gastrointestinal and biliary tract and liver disease, Cryptosporidium infections
ICOS deficiency	ICOS	AR	<u>604558</u>	Normal	Normal	Low	Recurrent infections, autoimmunity, gastroenteritis, granulomas
ICOSL deficiency	ICOSLG	AR	<u>605717</u>	Low	Low	Low	Recurrent bacterial and viral infections, neutropenia
$\text{CD3}\gamma\text{deficiency}$	CD3G	AR	<u>186740</u>	Normal number, but low TCR expression	Normal	Normal	Immune deficiency and autoimmunity of variable severity
CD8 deficiency	CD8A	AR	186910	Absent CD8, Normal CD4	Normal	Normal	Recurrent infections, may be asymptomatic
ZAP-70 deficiency (ZAP70 LOF)	ZAP70	AR	<u>269840</u>	Low CD8 number, normal CD4 number but with poor function	Normal	Normal	May have immune dysregulation, autoimmunity
ZAP-70 combined hypomorphic and activating mutations	ZAP70	AR (LOF/GOF)	617006	Decreased CD8, normal or decreased CD4 cells	Normal or decreased	Normal IgA, low IgM, low/normal IgG; protective Ab responses to vaccines	Severe autoimmunity (bullous pemphigoid, inflammatory colitis
	TAP1	AR	<u>170260</u>				.,
	TAP2	AR	<u>170261</u>	Low CD8, normal			Vasculitis, pyoderma gangrenosum
MHC class I deficiency	TAPBP	AR	601962	CD4, absent MHC I	Normal	Normal	gangronosam
	В2М	AR	109700	on lymphocytes			Sinopulmonary infections, cutaneous granulomas. Absent β2m associated proteins MHC- CD1a, CD1b, and CD1c
	CIITA	AR	600005	Low CD4+ T cells,			Failure to thrive, respiratory and
MHC class II deficiency group A, B, C, D	RFXANK	AR	603200	reduced MHC II expression on	Normal	Normal to low	gastrointestinal infections, liver/biliary tract disease
	RFX5	AR	601863	lymphocytes	Noma	140iiiai to iow	ivon billiary tract disease
, -, -, -	RFXAP	AR	<u>601861</u>	7			
IKAROS deficiency	IKZF1	AD DN	603023	no memory T cells	no memory B cells	Low Ig,	recurrent sinopulmonary infections, pneumocystis early CID onset



Table 1 (continued)

DOCK8 deficiency	DOCK8	AR	243700	T cell lymphopenia, reduced naïve CD8 T cells, increased exhausted CD8+ T _{EM} cells, reduced MAIT, NKT cells, increased γδ T cells; poor proliferation; few Treg with poor function	increased total B cells, reduced memory B cells Poor peripheral B cell tolerance.	Low IgM, normal/high IgG and IgA, very high IgE, poor antibody responses	Low NK cells with poor function. Eosinophilia, recurrent infections, cutaneous viral, fungal and staphylococcal infections, severe atopy/allergic disease, cancer diathesis
DOCK2 deficiency	DOCK2	AR	603122	Low	Normal	IgG normal or low, poor antibody responses	Early invasive herpes viral, bacterial infections, Normal NK cell number, but defective function. Poor interferon responses in hematopoietic and non-hematopoietic cells
Dehamenes S deficiency	POLD1	AR	<u>174761</u>	Low CD4 T cells	Low B cells but normal	Low igG	Recurrent respiratory tract infections, skin infections, warts
Polymerase δ deficiency	POLD2	AR	<u>600815</u>		maturation	_	and molluscum, short stature, intellectual disability
RHOH deficiency	RHOH	AR	602037	Normal, few naïve T cells, restricted repertoire, poor proliferation to CD3	Normal	Normal	HPV infection, lung granulomas, molluscum contagiosum, lymphoma
STK4 deficiency	STK4	AR	<u>614868</u>	CD4 lymphopenia, reduced naïve T cells, increased TEM and TEMRA cells, poor proliferation	Reduced memory B cells	Reduced IgM, increased IgG, IgA, IgE; impaired Ab responses	Intermittent neutropenia, bacterial, viral (HPV, EBV, molluscum), candidal infections, lymphoproliferation, autoimmune cytopenias, lymphoma, congenital heart disease
TCRα deficiency	TRAC	AR	<u>615387</u>	Absent TCRαβ except for a minor CD3-dim TCRαβ population; most T cells γδ; poor proliferation	Normal	Normal	Recurrent viral, bacterial, fungal infections, immune dysregulation and autoimmunity, diarrhea
LCK deficiency	LCK	AR	<u>615758</u>	Low CD4*, low Treg, restricted T cell repertoire, poor TCR signaling	Normal	Normal IgG and IgA, high IgM	Recurrent infections, immune dysregulation, autoimmunity
ITK deficiency	ITK	AR	<u>186973</u>	Progressive CD4 T cell lymphopenia; reduced T cell activation	Normal	Normal to low serum Ig	EBV associated B cell lymphoproliferation, lymphoma, immune dysregulation
MALT1 deficiency	MALT1	AR	<u>615468</u>	Normal number, poor proliferation	Normal	Normal levels, poor specific antibody response	Bacterial, fungal and viral infections
CARD11 deficiency	CARD11	AR LOF	<u>615206</u>	Normal number, predominantly naïve T-cells, poor proliferation	Normal, transitional B cell predominance	Absent/low	Pneumocystis jirovecii pneumonia, bacterial and viral infections
BCL10 deficiency	BCL10	AR	<u>616098</u>	Normal number, few memory T and Treg cells, poor antigen and anti-CD3 proliferation	Normal number, decreased memory and switched B cells	Low	Recurrent bacterial and viral infections, candidiasis, gastroenteritis
IL-21 deficiency	IL21	AR	<u>615767</u>	Normal number, normal/low function	Low, decreased memory and switched B cells	Hypogammaglob ulinemia, poor specific antibody	Severe early onset colitis, recurrent sinopulmonary infections
IL-21R deficiency	IL21R	AR	<u>615207</u>	Normal number, low cytokine production, poor antigen proliferation	Normal, decreased memory and switched B cells	responses; increased lgE	Recurrent infections, Pneumocystis jiroveci, Cryptosporidium infections, liver disease
OX40 deficiency	TNFRSF4	AR	615593	Normal numbers, low antigen specific memory CD4+	Normal numbers, low memory B cells	Normal	Impaired immunity to HHV8, Kaposi's sarcoma
IKBKB deficiency	IKBKB	AR	<u>615592</u>	Normal number, absent Treg and γ/δ T cells, impaired TCR activation	Normal number, poor function	Low	Recurrent bacterial, viral, fungal infections, opportunistic infections
NIK deficiency	MAP3K14	AR	<u>604655</u>	Normal number, poor proliferation to antigen	Low, low switched memory B cells	Low Ig's	Low NK number and function, recurrent bacterial, viral and Cryptosporidium infections
ReIB deficiency	RELB	AR	<u>604758</u>	Normal number, poor diversity, reduced proliferation to mitogens; no response to Ag	Marked increase in B cell number	Normal Ig levels but Impaired specific antibody responses	Recurrent infections
RelA haploinsufficiency	RELA	AD	<u>618287</u>	Normal/increased	Normal	Normal	Chronic mucocutaneous ulceration, Impaired NFkB activation; reduced production of inflammatory cytokines
Moesin deficiency	MSN	XL	300988	Normal number, defective migration, proliferation	Low number	Low Ig's over time	Recurrent infections with bacteria, varicella, neutropenia
TFRC deficiency	TFRC	AR	<u>616740</u>	Normal number, poor proliferation	Normal number, low memory B cells	Low	Recurrent infections, neutropenia, thrombocytopenia



Table 1 (continued)

c-Rel deficiency	REL	AR	<u>164910</u>	Normal, decreased memory CD4, poor proliferation	Low, mostly naïve; few switched memory B cells, impaired proliferation	Low, poor specific antibody responses	Recurrent infections with bacteria, mycobacteria, salmonella and opportunistic organisms. Defective innate immunity
FCHO1 deficiency	FCHO1	AR	613437	Low, poor proliferation	Normal number	Normal	Recurrent infections (viral, mycobacteria, bacterial, fungal), lymphoproliferation, failure to thrive, increased activation-induced T-cell death, defective clathrin-mediated endocytosis
PAX1 deficiency (8 patients)	PAX1	AR	615560	severe T cell lymphopenia, low TRECs	Normal number	Normal	Omenn-like syndrome (erythroderma, lymphocytosis, eosinophilia, severe/recurrent infections), no thymus, T cell deficiency not corrected by HSCT. Otofaciocervical syndrome type 2, ear abnormalities
ITPKB deficiency (1 patient)	ITPKB	AR	<u>NA</u>	Very few T cells	Normal	Normal IgM, A; low IgG	FTT, recurrent bacterial/fungal infections, pan-leukopenia, anemia, thrombocytopenia
SASH3 deficiency (5 patients)	SASH3	XL	<u>NA</u>	T/NK cell lymphopenia	B cell lymphopenia	Low, poor specific antibody responses	Recurrent sinopulmonary, cutaneous and mucosal infections, refractory autoimmune cyto-/neutropenia
MAN2B2 deficiency (1 patient)	MAN2B2	AR	<u>NA</u>	Low T cells	Low B cells	Normal/low	recurrent infections, vasculitis, arthritis, FTT, microcephaly, microdevelopmental delay; congenital disorder of glycosylation
COPG1 deficiency (5 patients)	COPG1	AR	<u>NA</u>	T cell lymphopenia	Normal	Normal but poor lg response to vaccines	recurrent pneumonia, viral respiratory infections, chronic EBV, CMV viremia, FTT, bronchiectasis
HELIOS deficiency	IKZF2	AD AR	<u>NA</u>	Increased activated T cells	Normal number; reduced memory	Reduced	recurrent upper respiratory infections/pneumonia, thrush, mucosal ulcers, chronic lymphadenopathy, SLE, ITP, AIHA (Evan's syndrome), EBV-associated HLH, lymphoma
IKKα deficiency (1 patient)	сник	AR	<u>NA</u>	Normal	Reduced	Low	recurrent bacterial, viral, fungal infections, absent secondary lymphoid tissues; skeletal abnormalities, FTT

SCID/CID spectrum: Infants with SCID who have maternal T cell engraftment may have T cells in normal numbers that do not function normally; these cells may cause autoimmune cytopenias or graft versus host disease. Hypomorphic mutations in several of the genes that cause SCID may result in Omenn syndrome (OS), or "leaky" SCID, or still less profound combined immunodeficiency (CID) phenotypes. Both OS and leaky SCID can be associated with >300 autologous T cells/uL of peripheral blood and reduced, rather than absent, proliferative responses when compared with typical SCID caused by null mutations. A spectrum of clinical findings including typical SCID, OS, leaky SCID, CID, granulomas with T lymphopenia, autoimmunity and CD4 T lymphopenia can be found in an allelic series of *RAG1/2* and other SCID-associated genes. There can be clinical overlap between some genes listed here and those listed in Table 7

Total number of mutant genes: 66. New inborn errors of immunity: 8 (SLP76 [12], PAX1 [13, 14], ITPKB [15]; SASH3 [16, 17], MAN2B2 [18], COPG1 [19], IKZF2 [20–23], CHUK [24])

SCID severe combined immunodeficiency, CID combined immunodeficiency, EBV Epstein-Barr virus, MHC major histocompatibility complex, HPV human papillomavirus, Treg T regulatory cell, XL X-linked inheritance, AR autosomal recessive inheritance, AD autosomal dominant inheritance, LOF loss-of-function, GOF gain-of-function, FTT failure to thrive



 Table 2 Combined immunodeficiencies with associated or syndromic features

	1. Immunodeficiency with Congenital Thrombocytopenia											
Disease	Genetic defect	Inheritance	OMIM	T cells	B cells	lg	Associated features					
Wiskott-Aldrich syndrome (WAS LOF)	WAS	XL	300392	Progressive decrease in numbers, abnormal lymphocyte responses to anti-CD3	Normal numbers	Low IgM and antibody responses to polysaccharides, often high IgA and IgE	Thrombocytopenia with small platelets, eczema, recurrent bacterial/viral Infections, bloody diarrhea, lymphoma, autoimmune disease, IgAn enphropathy. Patients with XL-thrombocytopenia have later onset of complications and more favourable life expectancy but eventually develop similar complications as observed in WAS					
WIP deficiency	WIPF1	AR	602357	Reduced, defective lymphocyte responses to anti-CD3	Normal or low	Normal, except for high IgE	Thrombocytopenia with or without small platelets, recurrent bacterial and viral infections, eczema, bloody diarrhea; WAS protein absent					
Arp2/3-mediated filament branching defect	ARPC1B	AR	604223	Normal	Normal numbers	Normal except for high IgA and IgE	Mild thrombocytopenia with normal sized platelets, recurrent invasive infections; colitis, vasculitis, autoantibodies (ANA, ANCA), eosinophilia; defective Arp2/3 filament branching					

		2. DNA	Repair	Defects Other	Than Tho	se Listed in Table 1	
Disease	Genetic	Inheritance	ОМІМ	T cells	B cells	lg	Associated features
Ataxia-telangiectasia	ATM	AR	607585	Progressive decrease, poor proliferation to mitogens; may have low TRECs and T cells by newborn screening (NBS)	Normal	Often low IgA, IgE and IgG subclasses, increased IgM monomers; antibodies variably decreased	Ataxia, telangiectasia especially of sclerae; pulmonary infections; lymphoreticular and other malignancies; increased alpha fetoprotein; increased radiosensitivity, chromosomal instability and chromosomal translocations
Nijmegen breakage syndrome	NBS1	AR	602667	Progressive decrease; may have low TRECs and T cells by NBS	Variably reduced	Often low IgA, IgE, and IgG subclasses, increased IgM; antibodies variably decreased	Microcephaly, dysmorphic facies; lymphomas and solid tumors; increased radiosensitivity;, chromosomal instability
Bloom syndrome	BLM	AR	604610	Normal	Normal	Low	Short stature, dysmorphic facies sun-sensitive erythema; marrow failure; leukemia, lymphoma; chromosomal instability
Immunodeficiency with	DNMT3B	AR	602900	Decreased or normal, responses to PHA may be decreased	Decreased	Hypogammaglobulinemia or	Facial dysmorphic features, developmental delay, macroglossia; bacterial/opportunistic infections; malabsorption; cytopenias; malignancies; multiradial configurations of chromosomes 1, 9, 16
centromeric instability and facial anomalies (ICF types 1, 2, 3, 4)	ZBTB24 CDCA7	AR AR	614064 609937	Decreased or normal Decreased or normal; responses	or normal	agammaglobulinemia, variable antibody deficiency	Facial dysmorphic features, macroglossia; bacterial/opportunistic infections; malabsorption; cytopenias; maliqnancies; multiradial
	HELLS	AR	603946	to PHA may be decreased Decreased or normal			configurations of chromosomes 1, 9, 16
PMS2 Deficiency	PMS2	AR	600259	Normal	Low B cells, switched and non-switched	Low IgG and IgA, high IgM, abnormal antibody responses	Recurrent infections; café-au-lait spots; lymphoma, colorectal carcinoma, brain tumors
RNF168 deficiency (Radiosensitivity, Immune Deficiency, Dysmorphic features, Learning difficulties [RIDDLE] syndrome)	RNF168	AR	612688	Normal	Normal	Low IgG or IgA	Short stature, mild defect of motor control to ataxia; normal intelligence to learning difficulties; mild facial dysmorphism to microcephaly; increased radiosensitivity
MCM4 deficiency	MCM4	AR	602638	Normal	Normal	Normal	NK cells: low number and function; viral infections (EBV, HSV, VZV); short stature; B cell lymphoma; adrenal failure
X-linked reticulate pigmentary disorder (POLA1 deficiency)	POLA1	XL	301220	Not assessed	Not assessed	Not assessed	Hyperpigmentation, characteristic facies, lung and GI involvement
POLE1 (Polymerase ε subunit 1) deficiency (FILS syndrome)	POLE1	AR	174762	Normal; decreased T cell proliferation	Low memory B cells	Low IgG2 and IgM, lack of antibody to PPS	Recurrent respiratory infections, meningitis; facial dysmorphism, livedo, short stature
POLE2 (Polymerase ε subunit 2) deficiency	POLE2	AR	602670	Lymphopenia, lack of TRECS at NBS, absent proliferation in response to antigens	Very low	Hypogammaglobulinemia	Recurrent infections, disseminated BCG infections; autoimmunity (type 1 diabetes), hypothyroidism, facial dysmorphism
Ligase I deficiency	LIG1	AR	126391	Lymphopenia, increased γδ T cells, decreased mitogen response	Normal	Hypogammaglobulinemia, Reduced antibody responses	Recurrent bacterial and viral infections; growth retardation; sun sensitivity, radiation sensitivity; macrocytic red blood cells
NSMCE3 deficiency	NSMCE3	AR	608243	Decreased number, poor responses to mitogens and antigens	Normal	Normal IgG, IgA, normal to elevated IgM; decreased antibody responses to PPS	Severe lung disease (possibly viral); thymic hypoplasia; chromosomal breakage, radiation sensitivity
ERCC6L2 (Hebo deficiency)	ERCC6L2	AR	615667	Lymphopenia	Low	Normal	Facial dysmorphism, microcephaly; bone marrow failure
GINS1 deficiency	GINS1	AR	610608	Low or normal	Low or normal	High IgA, low IgM and IgG	Neutropenia; IUGR; NK cells very low
MCM10 deficiency (1 patient)	MCM10	AR	619313	Low or normal	Low	Normal IgM, IgA, decreased IgG	severe (fatal) CMV infection, HLH-like, phenocopies GINS1 and MCM4 deficiencies; ↓ NK cells and NK function



 Table 2 (continued)

		3. Thy	mic De	fects with Addition	nal Cong	genital Anomalie	s	
Disease	Genetic defect	Inheritance	OMIM	T cells	B cells	lg	Associated features	
DiGeorge/velocardio- facial syndrome Chromosome 22q11.2 deletion syndrome (22q11.2DS)	Large deletion (3Mb) typically in chromosome 22 (TBX1)	AD	602054	Decreased or normal, 5% have low TRECs at NBS and <1500 CD3T cells/µL in neonatal period			Hypoparathyroidism; conotruncal cardiac	
DiGeorge/velocardio- facial syndrome	Unknown	Sporadic		Decreased or normal	Normal	Normal or decreased	malformation, velopalatal insufficiency; abnormal facies; intellectual disability	
TBX1 deficiency	TBX1	AD	602054	Decreased or normal, may have low TRECs at NBS				
	CHD7	AD	608892	Decreased or normal,	l		Coloboma of eye; heart anomaly; choanal atresia;	
CHARGE syndrome	SEMA3E	AD	608166	may have low TRECs at NBS; response to PHA	Normal	Normal or decreased	intellectual disability; genital and ear anomalies, CNS malformation: some are SCID-like	
	Unknown			may be decreased				
Winged helix nude FOXN1 deficiency	FOXN1	AR	<u>601705</u>	Very low	Normal	Decreased	Severe infections; abnormal thymic epithelium, immunodeficiency; congenital alopecia, nail dystrophy; neural tube defect	
FOXN1 haploinsufficiency	FOXN1	AD	600838	Severe T cell lymphopenia at birth, normalised by adulthood	Normal/ low	Not assessed	Recurrent, viral and bacterial respiratory tract infections; skin involvement (eczema, dermatitis), nail dystrophy	
Chromosome 10p13- p14 deletion syndrome (10p13-p14DS)	Del10p13-p14	AD	601362	Normal, rarely lymphopenia and decreased lymphoproliferation to mitogens and antigens; hypoplastic thymus may be present	Normal	Normal	Hypoparathyroidism; renal disease; deafness; growth retardation; facial dysmorphism; cardiac defects may be present; recurrent infections +/-	
Chromosome 11q deletion syndrome (Jacobsen syndrome)	11q23del	AD	<u>147791</u>	Lymphopenia; low NK cells	Decreas ed B cells and switched memory B cells	Hypogammaglobuline mia, decreased antibody responses	Recurrent respiratory infections; multiple warts; facial dysmorphism, growth retardation	

	4. Immuno-osseous Dysplasias											
Disease	Genetic defect	Inheritance	ОМІМ	T cells	B cells	lg	Associated features					
Cartilage hair hypoplasia (CHH)	RMRP	AR	<u>157660</u>	Varies from severely decreased (SCID) to normal; impaired lymphocyte proliferation	Normal	Normal or reduced, antibodies variably decreased	Short-limbed dwarfism with metaphyseal dysostosis, sparse hair, bone marrow failure; autoimmunity; susceptibility to lymphoma and other cancers; impaired spermatogenesis; neuronal dysplasia of the intestine					
Schimke Immuno-osseous dysplasia	SMARCAL1	AR	606622	Decreased	Normal	Normal	Short stature, spondyloepiphyseal dysplasia, intrauterine growth retardation; nephropathy; bacterial, viral, fungal infections; may present as SCID; bone marrow failure					
MYSM1 deficiency	MYSM1	AR	612176	T cell lymphopenia, reduced naïve T cells, low NK cells	B-cell deficiency	Hypogammaglobulinemia	Short stature; recurrent infections; congenital bone marrow failure, myelodysplasia; immunodeficiency affecting B-cells and granulocytes; skeletal anomalies; cataracts; developmental delay					
MOPD1 Deficiency (Roifman syndrome)	RNU4ATAC	AR	601428	Decreased NK cell function	Decreased total and memory B cells	Hypogammaglobulinemia, variably decreased specific antibodies	Recurrent bacterial infections; lymphadenopathy; spondyloepiphyseal dysplasia, extreme intrauterine growth retardation; retinal dystrophy; facial dysmorphism; may present with microcephaly; short stature					
Immunoskeletal dysplasia with neurodevelopmental abnormalities (EXTL3 deficiency)	EXTL3	AR	<u>617425</u>	Decreased	Normal	Decreased to normal	Short stature; cervical spinal stenosis, neurodevelopmental impairment; eosinophilia; may have early infant mortality					



Table 2 (continued)

				5. Hyper IgE Sync	Iromes (HIES)		
Disease	Genetic defect	Inheritance	OMIM	T cells	B cells	Ig	Associated features
AD-HIES STAT3 deficiency (Job syndrome)	STAT3	AD LOF (dominant negative)	<u>147060</u>	Normal overall; Th17, T follicular helper, MAIT, NKT cells decreased, Tregs may be increased; impaired responses to STAT3-activating cytokines	Normal, reduced memory B cells, BAFF expression increased, impaired responses to STAT3-activating cytokines	Very high IgE, specific antibody production decreased	Distinctive facial features (broad nasal bridge); bacterial infections (boils, pulmonary abscesses, pneumatoceles) due to Saureus, pulmonary Aspergillus, Pneumocystis Jirovecii; eczema, mucocutaneous candidiasis; hyperektensible joints, osteoporosis and bone fractures, scollosis, retained primary teeth; coronary and cerebral aneurysms
IL6 receptor deficiency	IL6R	AR	<u>147880</u>	Normal/increased; normal responses to mitogens	Normal total and memory B; reduced switched memory B	Normal/low serum IgM, G, A. Very high IgE; specific antibody production low	Recurrent pyogenic infections, cold abscesses; high circulating IL-6 levels
IL6 signal transducer (IL6ST) deficiency (partial)	IL6ST	AR	618523	Decreased Th17 cells	Reduced switched and non-switched memory B cells	High IgE, specific antibody production variably affected	Bacterial infections, boils, eczema, pulmonary abscesses, pneumatoceles; bone fractures; scoliosis; retention of primary teeth; craniosynostosis
IL6ST deficiency (partial) (12 patients)	ILEST	AD	619752	Normal, increased naïve, increased Th2	Normal total but reduced memory	Normal IgM, G, A; hyper- IgE	Dermatitis/eczema, eosinophilia, recurrent skin infections, pneumonia, bronchiectasis, pneumonia, bronchiectasis, pneumatoceles with severe secondary pulmonary aspergillosis, connective tissue defects (scoliosis, face, joints, fractures, palate, tooth retention). Phenocopies aspects of IL6R and IL11R deficiencies (due to unresponsiveness to these cytokines), as well as STAT3 DN/AR ZNF341
IL6ST deficiency (complete) (6 patients)	IL6ST	AR	<u>619751</u>	ND death in utero or in ne individuals)	onatal period occurred fo	or most affected	Fatal Stuve-Wiedemann-like syndrome; skeletal dysplasia, osteoporosis, hyperextensibility, lung dysfunction, renal abnormalities, thrombocytopenia, dermatitis, eczema. Defective acute phase response. Completely unresponsive to IL-6 family cytokines
ZNF341 deficiency AR-HIES	ZNF341	AR	<u>618282</u>	Decreased Th17 and NK cells	Normal, reduced memory B cells, impaired responses to STAT3-activating cytokines	High IgE and IgG, specific antibody production decreased	Phenocopy of AD-HIES; mild facial dysmorphism; early onset eczema, MCC, bacterial skin infections, abscesses, recurrent bacterial respiratory infections (S. aureus), lung abscesses and pneumatoceles; hyperextensible joints; bone fractures and retention of primary teeth
ERBIN deficiency	ERBB2IP	AD	606944	Increased circulating Treg	Normal	Moderately increased IgE	Recurrent respiratory infections, susceptibility to S. aureus, eczema; hyperextensible joints, scoliosis; arterial dilatation in some patients
Loeys-Dietz syndrome (TGFBR deficiency)	TGFBR1	AD	609192	Normal	Normal	Elevated IgE	Recurrent respiratory infectons; eczema, food allergies; hyper- extensible joints, scoliosis, retention
	TGFBR2		<u>610168</u>		*	5	of primary teeth; aortic aneurisms.
Comel-Netherton syndrome	SPINK5	AR	605010	Normal	Low switched and non- switched B cells	High IgE and IgA, Antibody variably decreased	Congenital ichthyosis, bamboo hair, atopic diathesis; increased bacterial infections; failure to thrive
PGM3 deficiency	Р СМЗ	AR	<u>172100</u>	CD8 and CD4 T cells may be decreased	Low B and memory B cells	Normal or elevated IgG and IgA, most with high IgE, eosinophilia	Severe atopy, autoimmunity, bacterial and viral infections; skeletal anomalies/dysplasia: short stature, brachydactyly, dysmorphic facial features; intellectual disability and cognitive impairment, delayed CNS myelination in some affected individuals
CARD11 deficiency (heterozygous DN)	CARD11	AD LOF	<u>617638</u>	Normal overall, but defective T cell activation and proliferation; skewing toward Th2	Normal to low	High IgE, poor specific antibody production; impaired activation of both NF-kB and mTORC1 pathways	Variable atopy, eczema, food allergies, eosinophilia; cutaneous viral infections, recurrent respiratory infections; lymphoma; CID

	6. Defects of Vitamin B12 and Folate Metabolism											
Disease	Genetic defect	Inheritance	OMIM	T cells	B cells	lg	Associated features					
Transcobalamin 2 deficiency	TCN2	AR	613441	Normal	Variable	Decreased	Megaloblastic anaemia, pancytopenia; if untreated (B12) for prolonged periods results in intellectual disability					
SLC46A1/PCFT deficiency causing hereditary folate malabsorption	SLC46A1	AR	229050	Variable numbers and activation profile	Variable	Decreased	Megaloblastic anaemia, failure to thrive; if untreated for prolonged periods results in intellectual disability					
Methylene-tetrahydrofolate dehydrogenase 1 (MTHFD1) deficiency	MTHFD1	AR	172460	Low thymic output, normal in vitro proliferation	Low	Decreased/poor antibody responses to conjugated polysaccharide antigens	Recurrent bacterial infection, Pneumocystis jirovecii; megaloblastic anaemia; failure to thrive; neutropenia; seizures, intellectual disability; folate-responsive					

	7. Anhidrotic Ectodermodysplasia with Immunodeficency (EDA-ID)											
Disease	Genetic defect	Inheritan ce	ОМІМ	T cells	B cells	lg	Associated features					
EDA-ID due to NEMO /IKBKG deficiency (ectodermal dysplasia, immune deficiency)	IKBKG	XL	300248	Normal or decreased, TCR activation impaired	Normal; Low memory and isotype switched B cells	Decreased, some with elevated IgA, IgM, poor specific antibody responses, absent antibodies to polysaccharide antigens	Anhidrotic ectodermal dysplasia (in some); various infections (bacteria, mycobacteria, viruses, fungi); colitis; conical teeth, variable defects of skin, hair and teeth; monocyte dysfunction					
EDA-ID due to IKBA GOF mutation	NFKBIA	AD GOF	<u>164008</u>	Normal total T cells, TCR activation impaired	Normal B cell numbers, impaired BCR activation, low memory and isotype switched B cells	Decreased IgG and IgA, elevated IgM, poor specific antibody responses, absent antibody to polysaccharide antigens	Anhidrotic ectodermal dysplasia; various infections (bacteria, mycobacteria, viruses, fungi); colitis; variable defects of skin, hair and teeth; T cell and monocyte dysfunction					



Table 2 (continued)

EDA-ID due to IKBKB GOF mutation	IKBKB	AD GOF	618204	Decreased T cells, impaired TCR activation	Normal number, poor function	Reduced	Recurrent bacterial, viral, fungal infections; variable ectodermal defects

	8. Calcium Channel Defects										
Disease	Genetic defect	Inheritance	OMIM	T cells	B cells	lg	Associated features				
ORAI-1 deficiency	ORAI1	AR	610277	Normal, defective TCR mediated activation	Normal	Normal	Autoimmunity; EDA; non-progressive myopathy				
STIM1 deficiency	STIM1	AR	605921								
CRACR2A deficiency (1 patient)	CRACR2A	AR	NA	Mild reduction in T cell numbers	Normal	Low	Later onset, chronic diarrhea, recurrent lower respiratory tract infections, including pneumonia				

				9. Other Defec	ets		
Disease	Genetic defect	Inheritance	OMIM	T cells	B cells	lg	Associated features
Purine nucleoside phosphorylase (PNP) deficiency	PNP	AR	<u>164050</u>	Progressive decrease	Normal	Normal or low	Autoimmune haemolytic anaemia; neurological impairment
Immunodeficiency with multiple intestinal atresias	TTC7A	AR	609332	Variable, but sometimes absent or low TRECs at NBS; may have SCID phenotype at birth	Normal or low	Markedly decreased IgG, IgM, IgA	Bacterial (sepsis), fungal, viral infections; multiple intestinal atresias, often with intrauterine polyhydramnios and early demise
Tricho-Hepato-Enteric Syndrome (THES)	TTC37 SKIV2L	AR	<u>222470</u> <u>614602</u>	Impaired IFN γ production	Variably low numbers of switched memory B cells	Hypogammaglobulin emia, may have low antibody responses	Respiratory infections; IUGR; facial dysmorphic features, wooly hair; early onset intractable diarrhea, liver cirrhosis; platelet abnormalities
Hepatic veno-occlusive disease with immunodeficiency (VODI)	SP110	AR	604457	Normal (decreased memory T cells)	Normal (decreased memory B cells)	Decreased IgG, IgA, IgM, absent germinal center and tissue plasma cells	Hepatic veno-occlusive disease; susceptibility to <i>Pneumocystis jirovecii</i> pneumonia, CMV, candida; thrombocytopenia; hepatosplenomegaly; cerebrospinal leukodystrophy
BCL11B deficiency	BCL11B	AD	617237	Low, poor proliferation	Normal	Normal	Congenital abnormalities, neonatal teeth, dysmorphic facies; absent corpus callosum, neurocognitive deficits
EPG5 deficiency (Vici syndrome)	EPG5	AR	615068	Profound depletion of CD4+ cells	Defective	Decreased (particularly IgG2)	Agenesis of the corpus callosum; cataracts; cardiomyopathy; skin hypopigmentation; intellectual disability; microcephaly; recurrent infections, chronic mucocutaneous candidiasis
HOIL1 deficiency	RBCK1	AR	610924	Normal numbers	Normal, decreased memory B cells	Poor antibody responses to polysaccharides	Bacterial infections; autoinflammation; amylopectinosis
HOIP deficiency	RNF31	AR	612487	Normal numbers	Normal, decreased memory B cells	decreased	Bacterial infections; autoinflammation; amylopectinosis; lymphangiectasia
Hennekam-lymphangiectasia-	CCBE1	AR	612753	Low/variable	Low/variable	decreased	Lymphangiectasia and lymphedema with facial abnormalities and other dysmorphic features
lymphedema syndrome	FAT4	AR	<u>612411</u>	Low/variable	Low/variable	decreased	Lymphangiectasia and lymphedema with facial abnormalities and other dysmorphic features
Activating de novo mutations in nuclear factor, erythroid 2- like (NFE2L2)	NFE2L2	AD	<u>617744</u>	Not reported	Decreased switched memory B cells	Hypogammaglobulin emia, decreased antibody responses	Recurrent respiratory and skin infections; growth retardation, developmental delay; white matter cerebral lesions; increased level of homocysteine; increased expression of stress response genes
STAT5b deficiency	STAT5B	AR	245590	Modestly decreased, reduced Treg number and function	Normal	hypergammaglobulin emia, increased IgE	Growth-hormone insensitive dwarfism; dysmorphic features; eczema; lymphocytic interstitial pneumonitis; prominent autoimmunity
STAT5b deficiency	STAT5B	AD (dominant negative)	604260	Normal	Normal	Increased IgE	Growth-failure; eczema (no immune defects compared to AR STAT5 deficiency)
Kabuki syndrome	KMT2D	AD	602113			Low IgA and	Typical facial abnormalities, cleft or high arched palate, skeletal abnormalities, short
(type 1 and 2)	KDM6A	XL (females may be affected)	300128	Normal	Normal	occasionally low IgG	stature; intellectual disability; congenital heart defects; recurrent infections (otitis media, pneumonia) in 50% of patients; autoimmunity may be present
KMT2A deficiency (Wiedemann-Steiner syndrome)	KMT2A	AD	<u>605130</u>	Normal	Decreased switched and non- switched memory B cells	Hypogammaglobulin emia, decreased antibody responses	Respiratory infections; short stature; hypertelorism; hairy elbows; developmental delay, intellectual disability
DIAPH1 deficiency (7 patients)	DIAPH1	AR	616632	Reduced naïve T cells	Decreased memory B cells	Low IgM, normal IgG	Seizures, cortical blindness, microcephaly syndrome (SCBMS); recurrent bacterial, viral, fungal infections; B-lymphoma (3/7)
AIOLOS deficiency (7 patients)	IKZF3	AD	619437	Normal	Reduced; impaired development	Very low	EBV susceptibility, recurrent sinopulmonary & respiratory infections, Pneumocystis jirovecii, warts (HPV), M avium, B cell malignancy
CD28 deficiency (3 patients)	CD28	AR	NA	Normal	Normal	Normal	Susceptibility to HPV infection only

Total number of mutant genes in Table 2: 69. New inborn errors of immunity: 7 (MCM10 [29, 30], AR and AD IL6ST [31–33], CRACR2A [27], DIAPH1 [34], IKZF3 [25, 26], CD28 [28]). Unknown cause of DiGeorge syndrome, unknown cause of CHARGE syndrome, unknown gene(s) within 10p13-14 deletion responsible for phenotype

EDA ectodermal dysplasia anhidrotic, HSV herpes simplex virus, VZV varicella zoster virus, BCG Bacillus Calmette-Guerin, NBS newborn screen, TREC T cell receptor excision circle (biomarker for low T cells used in NBS), IUGR intrauterine growth retardation



Table 3 Predominantly antibody deficiencies

Disease	Genetic defect	Inheritance	OMIM	lg	Associated features		
BTK deficiency, X-linked agammaglobulinemia (XLA)	втк	XL	300300	All isotypes decreased in majority of patients, some patients have detectable immunoglobulins	Severe bacterial infections, normal numbers of pro-B cells		
μ heavy chain deficiency	IGHM	AR	147020				
λ5 deficiency	IGLL1	AR	146770		Severe bacterial infections.		
lgα deficiency	CD79A	AR	112205		normal numbers of pro-B cells		
lgβ deficiency	CD79B	AR	147245				
BLNK deficiency	BLNK	AR	604515				
p110∂ deficiency	PIK3CD	AR	602839		Severe bacterial infections; autoimmune complications (IBD)		
p85 deficiency	PIK3R1	AR	615214		Severe bacterial infections, cytopenias, decreased or absent pro-B cells		
	TCF3	AD	616941		Recurrent bacterial infections		
E47 transcription factor deficiency	TCF3	AR	<u>147141</u>	All isotypes decreased	Severe, recurrent bacterial infections, failure to thrive		
SLC39A7 (ZIP7) deficiency	SLC39A7	AR	601416		Early onset infections, blistering dermatosis, failure to thrive, thrombocytopenia		
Hoffman syndrome/TOP2B deficiency	TOP2B	AD	126431		Recurrent infections, facial dysmorphism limb anomalies		
FNIP1 deficiency (6 patients)	FNIP1	AR	619705		Early onset recurrent infections, bronchiectasis, fibrosis, interstitial pneumoniae; neutropenia (severe or intermittent); Crohn disease (one patient); congenital heart defects, muscular hypotonia; developmental delav		
PU1 deficiency	SPI1	AD	619707	1	Sinopulmonary infections with encapsulated bacteria, viral infections		

2. Severe Reduction in at Leas	t 2 Serum In		ulin Isotyp enotype	oes with Normal or Lo	w Number of B Cells, CVID
Disease	Genetic defect	Inheritance	OMIM	lg	Associated features
Common variable immune deficiency with no gene defect specified (CVID)	Unknown	Variable		Low IgG and IgA and/or IgM	Clinical phenotypes vary: most have recurrent infections, some have polyclonal lymphoproliferation, autoimmune cytopenias and/or granulomatous disease
Activated p1108 syndrome (APDS)	PIK3CD GOF	AD	615513 (APDS1)	Normal/increased IgM,	Severe bacterial infections; reduced memory B cells and increased transitional B cells, EBV ± CMV viremia, lymphadenopathy/splenomegaly, autoimmunity, lymphoproliferation,
	PIK3R1	AD	616005 (APDS2)	reduced IgG and IgA	Jymphoma Severe bacterial infections, reduced memory B cells and increased transitional B cells, lymphadenopathy/splenomegaly, lymphoproliferation, lymphoma; developmental delay
PTEN Deficiency (LOF)	PTEN	AD	<u>158350</u>	Normal/Decreased	Recurrent infections, Lymphoproliferation, Autoimmunity; developmental delay
CD19 deficiency	CD19	AR	107265	Low IgG and IgA and/or IgM	Recurrent infections, may have
CD81 deficiency	CD81	AR	<u>186845</u>	Low IgG, low or normal IgA and IgM	glomerulonephritis (CD81 mutation abolishes expression of CD19, thereby phenocopying CD19 mutations)
CD20 deficiency	CD20	AR	112210	Low IgG, normal or elevated IgM and IgA	Recurrent infections
CD21 deficiency	CD21	AR	<u>120650</u>	Low IgG, impaired anti- pneumococcal response	Recurrent infections
TACI deficiency [#]	TNFRSF13B	AR or AD	604907	Low IgG and IgA and/or IgM	Variable clinical expression and penetrance for monoallelic variants
BAFF receptor deficiency	TNFRSF13C	AR	606269	Low IgG and IgM,	Variable clinical expression
TWEAK deficiency	TNFSF12	AD	602695	Low IgM and A, lack of anti- pneumococcal antibody	Pneumonia, bacterial infections, warts, thrombocytopenia. neutropenia
TRNT1 deficiency	TRNT1	AR	612907	B cell deficiency and hypogammaglobulinemia	congenital sideroblastic anemia, deafness developmental delay
NFKB1 deficiency	NFKB1	AD	<u>164011</u>	Normal or low IgG, IgA, IgM, low or normal B cells, low memory B cells	Recurrent sinopulmonary infections, COPD, EBV proliferation, autoimmune cytopenias, alopecia and autoimmune thyroiditis
NFKB2 deficiency	NFKB2	AD	<u>615577</u>	Low serum IgG, A and M; low B cell numbers	Recurrent sinopulmonary infections, alopecia and endocrinopathies
IKAROS deficiency	IKZF1	AD (haploinsuffici ency)	603023	Low IgG, IgA, IgM, Iow or normal B cells; B cells and Ig levels reduce with age	Decreased pro-B cells, recurrent sinopulmonary infections; increased risk o ALL, autoimmunity, CVID phenotype
IRF2BP2 deficiency	IRF2BP2	AD	615332	Hypogammaglobulinemia, absent IgA	Recurrent infections, possible autoimmunity and inflammatory disease
ATP6AP1 deficiency	ATP6AP1	XL	300972	Variable immunoglobulin findings	Hepatopathy, leukopenia, low copper
ARHGEF1 deficiency	ARHGEF1	AR	618459	Hypogammaglobulinemia; lack of antibody	Recurrent infections, bronchiectasis
SH3KBP1 (CIN85) deficiency	SH3KBP1	XL	300310	IgM, IgG deficiency; loss of antibody	Severe bacterial infections
SEC61A1 deficiency	SEC61A1	AD	609213	Hypogammaglobulinemia	Severe recurrent respiratory tract infections
RAC2 deficiency	RAC2	AR	602049	Low IgG, IgA, IgM, Iow or normal B cells; reduced Ab responses following vaccination	Recurrent sinopulmonary infections, selective IgA deficiency; poststreptococca glomerulonephritis; urticaria
Mannosyl-oligosaccharide glucosidase deficiency	MOGS	AR	601336	Low IgG, IgA, IgM, increased B cells; poor Ab responses following vaccination	Bacterial and viral infections; severe neurologic disease; also known as congenital disorder of glycosylation type Ilb (CDG-Ilb)
PIK3CG deficiency (2 patients)	PIK3CG	AR	619802	Reduced memory B cells, hypogammaglobulinemia	Recurrent infections, Cytopenia /lymphopenia, eosinophilia, splenomegaly, lymphadenopathy, HLH- like
BOB1 deficiency (1 patient)	POU2AF1	AR	<u>NA</u>	Reduced memory B cells, agammaglobulinemia	Recurrent respiratory infections, possible chronic viral infection of CNS with progressive tetraparesia



Table 3 (continued)

3. Severe Reduction in Serum	IgG and IgA	with Norm	al/Elevate	d IgM and Normal I	Numbers of B cells, Hyper IgM
Disease	Genetic defect	Inheritance	ОМІМ	lg	Associated features
AID deficiency		AR	6055258	IgG and IgA decreased, IgM increased; normal memory B cells but lacking somatic hypermutation	Bacterial infections, enlarged lymph nodes and germinal centers; autoimmunity
AD deliciency	AICDA	AD	605257	IgG absent or decreased, IgA undetected, IgM increased; normal memory B cells with intact somatic hypermutation	Bacterial infections, enlarged lymph nodes and germinal centers. Mutations uniquely localise to the nuclear export signal.
UNG deficiency	UNG	AR	<u>191525</u>	IgG and IgA decreased, IgM increased	Enlarged lymph nodes and germinal centers
INO80 deficiency	INO80	AR	<u>610169</u>	IgG and IgA decreased, IgM increased	Severe bacterial infections
MSH6 deficiency	MSH6	AR	600678	Variable IgG, defects, increased IgM in some, normal B cells, low switched memory B cells, Ig class switch recombination and somatic hypermutation defects	Family or personal history of cancer
CTNNBL1 deficiency (1 patient)	CTNNBL1	AR	<u>NA</u>	Reduced memory B cells, Ig class switch recombination and somatic hypermutation defects, progressive hypogammaglobulinemia	CVID, autoimmune cytopenias, recurrent infections, hyperplastic germinal centers
APRIL deficiency (1 patient)	TNFSF13	AR	<u>NA</u>	Normal total B cell counts, Reduced memory B cells, hypogammaglobulinemia	CVID, chronic but mild infections, alopecia areata

4. Isotype, Light Cha	in, or Function	onal Defici	encies wit	h Generally Normal Nur	nbers of B Cells
Disease	Genetic defect	Inheritance	ОМІМ	lg .	Associated features
lg heavy chain mutations and deletions	Mutation or chromosomal deletion at 14q32	AR		One or more IgG and/or IgA subclasses as well as IgE may be absent	May be asymptomatic
Kappa chain deficiency	IGKC	AR	147200	All immunoglobulins have lambda light chain	Asymptomatic
Isolated IgG subclass deficiency	Unknown	?		Reduction in one or more IgG subclass	Usually asymptomatic, a minority may have poor antibody response to specific antigens and recurrent viral/bacterial infections
IgG subclass deficiency with IgA deficiency	Unknown	?		Reduced IgA with decrease in one or more IgG subclass	Recurrent bacterial infections May be asymptomatic
Selective IgA deficiency	Unknown	?		Absent IgA with other isotypes normal, normal subclasses and specific antibodies	May be asymptomatic Bacterial infections, autoimmunity mildly increased
Specific antibody deficiency with normal Ig levels and normal B cells	Unknown	?		Normal	Reduced ability to produce antibodies to specific antigens
Transient hypogammaglobulinemia of infancy	Unknown	?		IgG and IgA decreased	Normal ability to produce antibodies to vaccine antigens, usually not associated with significant infections
CARD11 GOF	CARD11	AD GOF	<u>616452</u>	polyclonal B cell lymphocytosis due to constitutive NF-κB activation	Splenomegaly, lymphadenopathy, poor vaccine response
Selective IgM deficiency	Unknown	?		Absent serum IgM	Pneumococcal / bacterial

Common variable immunodeficiency disorders (CVID) include several clinical and laboratory phenotypes that may be caused by distinct genetic and/or environmental factors. Some patients with CVID and no known genetic defect have markedly reduced numbers of B cells as well as hypogammaglobulinemia. Identification of causal variants can assist in defining treatment. In addition to monogenic causes on this table, a small minority of patients with XLP (Table 4), WHIM syndrome (Table 6), ICF (Table 2), VODI (Table 2), thymoma with immunodeficiency (Good syndrome) or myelodysplasia are first seen by an immunologist because of recurrent infections, hypogammaglobulinemia and normal or reduced numbers of B cells

Total number of mutant genes in Table 3: 45. New inborn errors of immunity: 6 (FNIP1 [35, 36], SP1I [37], PIK3CG [38, 39], POU2AF1 [40], CTNNBL1 [41], TNSRSF13 [42])

EBV Epstein-Barr virus, COPD chronic obstructive pulmonary disease

*Heterozygous variants in TNFRSF13B have been detected in healthy individuals, thus such variants are likely to be disease-modifying rather than disease-causing



 Table 4 Diseases of immune dysregulation

	1. Familial H	emophago	cytic Lymp	hohistiocytos	is (FHL syı	ndromes)	
Disease	Genetic defect	Inheritance	ОМІМ	Circulating T Cells	Circulating B cells	Functional defect	Associated Features
Perforin deficiency (FHL2)	PRF1	AR	170280	Increased activated T cells	Normal	Decreased to absent NK and CTL activities cytotoxicity	Fever, HSM, hemophagocytic lymphohistiocytosis (HLH), cytopenias
UNC13D / Munc13-4 deficiency (FHL3)	UNC13D	AR	608897			Decreased to absent NK	Fever, HSM, HLH,
Syntaxin 11 deficiency (FHL4)	STX11	AR	<u>605014</u>	Increased activated T cells	Normal	and CTL activities (cytotoxicity and/or	cytopenias,
STXBP2 / Munc18-2 deficiency (FHL5)	STXBP2	AR or AD	601717	donvatou i cono		degranulation)	
FAAP24 deficiency	FAAP24	AR	610884	Increased activated T cells	Normal	Failure to kill autologous EBV transformed B cells. Normal NK cell function	EBV-driven lymphoproliferative disease
SLC7A7 deficiency	SLC7A7	AR	222700	Normal	Normal	Hyper-inflammatory response of macrophages Normal NK cell function	Lysinuric protein intolerance, bleeding tendency, alveolar proteinosis
RHOG deficiency (1 patient)	RHOG	AR	<u>NA</u>	Normal	Slightly reduced	Impaired CTL and NK cell cytotoxicity	HLH (hemophagocytosis, hepatosplenomegaly, fever, cytopenias, low hemoglobin, hyper- triglyceridemia, elevated ferritin, sCD25)

	2. F	HL Syndro	mes with	Hypopigment	tation		
Disease	Genetic defect	Inheritance	ОМІМ	Circulating T Cells	Circulating B cells	Functional defect	Associated Features
Chediak-Higashi syndrome	LYST	AR	606897	Increased activated T cells	Normal	Decreased NK and CTL activities (cytotoxicity and/or degranulation)	Partial albinism, recurrent infections, fever, HSM, HLH, giant lysosomes, neutropenia, cytopenias, bleeding tendency, progressive neurological dysfunction
Griscelli syndrome, type 2	RAB27A	AR	603868	Normal	Normal	Decreased NK and CTL activities (cytotoxicity and/or degranulation)	Partial albinism, fever, HSM, HLH, cytopenias
Hermansky-Pudlak syndrome, type 2	AP3B1	AR	603401	Normal	Normal	Decreased NK and CTL activities (cytotoxicity and/or degranulation)	Partial albinism, recurrent infections, pulmonary fibrosis, increased bleeding, neutropenia, HLH
Hermansky-Pudlak syndrome, type 10	AP3D1	AR	<u>617050</u>	Normal	Normal	Decreased NK and CTL activities (cytotoxicity and/or degranulation)	Oculocutaneous albinism, severe neutropenia, recurrent infections, seizures, hearing loss and neurodevelopmental delay
CEBPE neofunction (3 patients)	CEBPE	AR GOF	<u>245480</u>	Mild reduction	Not done	Autoinflammasome activation/ ↑ IFN gene expression, altered chromatin occupancy of mutant CEBPE, and transcriptional changes	Recurrent abdominal pain, aseptic fever, systemic inflammation; abscesses, ulceration, infections; mild bleeding diathesis

		3. Reg	ulatory T	Cell Defects	S		
Disease	Genetic defect	Inheritance	ОМІМ	Circulating T Cells	Circulating B cells	Functional defect	Associated Features
IPEX, immune dysregulation, polyendocrinopathy, enteropathy X-linked	FOXP3	XL	300292	Normal	Normal	Lack of (and/or impaired function of) CD4* CD25* FOXP3* regulatory T cells (Tregs)	Autoimmune enteropathy, early onset diabetes, thyroiditis hemolytic anemia, thrombocytopenia, eczema, elevated IgE and IgA
CD25 deficiency	IL2RA	AR	<u>147730</u>	Normal to decreased	Normal	No CD4+C25+ cells with impaired function of Tregs cells	Lymphoproliferation, autoimmunity, impaired T cell proliferation in vitro
CD122 deficiency	IL2RB	AR	<u>618495</u>	Increased memory CD8 T cells, decreased Tregs	Increased memory B cells	Diminished IL2Rβ expression, dysregulated signaling in response to IL-2/IL-15; increased immature NK cells	Lymphoproliferation, lymphadenopathy, hepatosplenomegaly, autoimmune hemolytic anemia, dermatitis, enteropathy, hypergammaglobulinemia, recurrent viral (EBV, CMV) infections
CTLA4 haploinsufficiency (ALPS-V)	CTLA4	AD	123890	Decreased	Decreased	Impaired function of Tregs.	Autoimmune cytopenias, enteropathy, interstitial lung disease, extra-lymphoid lymphocytic infiltration, recurrent infections
LRBA deficiency	LRBA	AR	606453	Normal or decreased CD4 numbers T cell dysregulation	Low or normal numbers of B cells	Reduced IgG and IgA in most	Recurrent infections, inflammatory bowel disease, autoimmunity
DEF6 deficiency	DEF6	AR	610094	Mild CD4 and CD8 lymphopenia	Low or normal numbers of B cells	Impaired Treg function	Enteropathy, hepatosplenomegaly, cardiomyopathy, recurrent infections



 Table 4 (continued)

STAT3 GOF mutation	STAT3	AD GOF	102582	Decreased	Decreased	Enhanced STAT3 signaling, leading to increased Th17 cell differentiation, lymphoproliferation and autoimmunity. Decreased Tregs and impaired function	Lymphoproliferation, solid organ autoimmunity, recurrent infections
BACH2 deficiency	BACH2	AD	605394	Progressive T cell lymphopenia	Impaired memory B cell development	Haploinsufficiency for a critical lineage specification transcription factor	Lymphocytic colitis, sinopulmonary infections
FERMT1 deficiency	FERMT1	AR	<u>173650</u>	Normal	Normal	Intracellular accumulation of IgG, IgM, IgA, and C3 in colloid bodies under the basement membrane	Dermatosis characterized by congenital blistering, skin atrophy, photosensitivity, skin fragility, and scaling
IKAROS GOF (8 patients)	IKZF1	AD GOF	<u>NA</u>	Normal	Normal/mil d decrease	Increased binding of mutant IKAROS to DNA/target genes	Multiple autoimmune features (diabetes, colitis, thyroiditis), allergy, lymphoproliferation, plasma cell expansion (IgG4*), Evans Syndrome, recurrent infections

	4	4. Autoimm	nunity wit	h or without Lyr	nphoprolifera	ation	
Disease	Genetic defect	Inheritance	ОМІМ	Circulating T Cells	Circulating B cells	Functional defect	Associated Features
APECED (APS-1), autoimmune polyendocrinopathy with candidiasis and ectodermal dystrophy	AIRE	AR or AD	240300	Normal	Normal	AIRE serves as check- point in the thymus for negative selection of autoreactive T cells and for generation of Tregs	Autoimmunity: hypoparathyroidism, hypothyroidism, adrenal insufficiency, diabetes, gonadal dysfunction and other endocrine abnormalities; dental enamel hypoplasia, alopecia areata enteropathy, pernicious anemia; chronic mucocutaneous candidiasis
ITCH deficiency	ITCH	AR	606409			Itch deficiency may cause immune dysregulation by	Early-onset chronic lung disease (interstitial pneumonitis), autoimmunity (thyroiditis, type I
				Not assessed	Not assessed	affecting both anergy induction in auto- reactive effector T cells and generation of Tregs	diabetes, chronic diarrhea/enteropathy, and hepatitis), failure to thrive, developmental delay, dysmorphic facial features
Tripeptidyl-Peptidase II Deficiency	TPP2	AR	<u>190470</u>	Decreased	Decreased	TPP2 deficiency results in premature immunosenescence and immune dysregulation	Variable lymphoproliferation, severe autoimmune cytopenias, hypergammaglobulinemia, recurrent infections
JAK1 GOF	JAK1	AD GOF	<u>147795</u>	Not assessed	Not assessed	Hyperactive JAK1	HSM, eosinophilia, eosinophilic enteritis, thyroid disease, poor growth, viral infections
Prolidase deficiency	PEPD	AR	613230	Normal	Normal	Peptidase D	Autoantibodies common, chronic skin ulcers, eczema, infections
SOCS1 haploinsufficiency (15 patients)	SOCS1	AD	619375	Decreased	Reduced switched memory B cells	↑ pSTAT1, ↑ type I/II IFN signature	Early onset severe multisystemic autoimmunity, neutropenia, lymphopenia, ITP, AIHA, S.L.E, GN, hepatosplenomegaly, psoriasis, arthritis, thyroiditis, hepatitis; recurrent bacterial infections. Incomplete penetrance
PD-1 deficiency (1 patient)	PDCD1	AR	<u>NA</u>	Mostly intact	Normal	Lack of PD-1 on patient PBMCs, reduced IFNy production in response to mycobacterial stimuli	Tuberculosis, autoimmunity (T1D, hypothyroidism, JIA), fatal pulmonary autoimmunity, hepatosplenomegaly

	5	. Immune	Dysregula	tion with (Colitis		
Disease	Genetic defect	Inheritance	ОМІМ	Circulating T Cells	Circulating B cells	Functional defect	Associated Features
IL-10 deficiency	IL10	AR	<u>124092</u>	Normal	Normal	No functional IL-10 secretion	Inflammatory bowel disease (IBD) Folliculitis, recurrent respiratory diseases, arthritis,
	IL10RA	AR	<u>146933</u>	Normal	Normal	Leukocytes unresponsive to IL-10	IBD, Folliculitis, recurrent
IL-10R deficiency	IL10RB	AR	123889	Normal	Normal	Leukocytes unresponsive to IL-10, and IL-22, IL-26, IL-28A, IL-28B and IL-29	respiratory diseases, arthritis, lymphoma
NFAT5 haploinsufficiency	NFAT5	AD	604708	Normal	Normal	Decreased memory B cells and plasmablasts	IBD, recurrent sinopulmonary infections
TGFB1 deficiency	TGFB1	AR	618213	Normal	Normal	Decreased T cell proliferation in response to anti-CD3	IBD, immunodeficiency, recurrent viral infections, microcephaly, and encephalopathy
RIPK1	RIPK1	AR	<u>618108</u>	Reduced	Normal/ Reduced	Reduced activation of MAPK, NFkB pathways to	Recurrent infections, early- onset IBD, progressive polyarthritis
ELF4 deficiency (3 patients)	ELF4	XL	301074	Normal	Normal	hyper inflammatory macrophages	Early onset IBD/mucosal autoinflammation, fevers, ulcers, Responded to IL-1, TNF or IL-12p40 blockade



Table 4 (continued)

6. Autoim	mune Lymph	oprolifera	tive Syndr	ome (ALP	S, Canale-	Smith syndrome)	
Disease	Genetic defect	Inheritance	ОМІМ	Circulating T Cells	Circulating B cells	Functional defect	Associated Features
		AD		Increased TCR α/β+	Normal, low memory	Apoptosis defect FAS mediated	Splenomegaly, adenopathies, autoimmune cytopenias,
ALPS-FAS	TNFRSF6	AR <u>11</u>	<u>134637</u>	CD4 CD8 double negative (DN) T cells	B cells		increased lymphoma risk, IgG and A normal or increased, elevated serum FasL, IL-10, vitamin B12
ALPS-FASLG	TNFSF6	AR	<u>134638</u>	Increased DN T cells	Normal	Apoptosis defect FASL mediated	Splenomegaly, adenopathies, autoimmune cytopenias, SLE, soluble FasL is not elevated
ALPS-Caspase10	CASP10	AD	601762	Increased DN T cells	Normal	Defective lymphocyte apoptosis	Adenopathies, splenomegaly, autoimmunity
ALPS-Caspase 8	CASP8	AR	601763	Slightly increased DN T cells	Normal	Defective lymphocyte apoptosis and activation	Adenopathies, splenomegaly, bacterial and viral infections, hypogammaglobulinemia
FADD deficiency	FADD	AR	602457	Increased DN T cells	Normal	Defective lymphocyte apoptosis	Functional hyposplenism, bacterial and viral infections, recurrent episodes of encephalopathy and liver dysfunction

			l			•	dystunction
	7. Susceptib			phoprolife Circulating	rative Con		
Disease	Genetic defect	Inheritance	OMIM	T Cells	B cells	Functional defect	Associated Features
SAP deficiency (XLP1)	SH2D1A	XL	300490	Normal or Increased activated T cells	Reduced Memory B cells	Reduced NK cell and CTL cytotoxic activity	Clinical and immunologic features triggered by EBV infection: HLH, Lymphoproliferation, Aplastic anaemia, Lymphoma. Hypogammaglobulinemia, Absent iNKT cells
XIAP deficiency (XLP2)	XIAP	XL	300079	Normal or Increased activated T cells; Iow/normal iNK T cells	Normal or reduced Memory B cells	Increased T cells susceptibility to apoptosis to CD95 and enhanced activation-induced cell death (AICD)	EBV infection, Splenomegaly, lymphoproliferation HLH, Colitis, IBD, hepatitis Low iNKT cells
CD27 deficiency	CD27	AR	615122	Normal	No memory B cells	hypogammaglobulinemia; poor Ab responses to some vaccines/infections	Features triggered by EBV infection, HLH, aplastic anemia, low iNKT cells, B-lymphoma
CD70 deficiency	CD70	AR	602840	Normal number, low Treg, poor activation and function	Decreased memory B cells	hypogammaglobulinemia; poor Ab responses to some vaccines/infections	EBV susceptibility, Hodgkin lymphoma; autoimmunity in some patients
CTPS1 deficiency	CTPS1	AR	<u>615897</u>	Normal to low, but reduced activation, proliferation	Decreased memory B cells	Normal/high IgG poor proliferation to antigen	Recurrent/chronic bacterial and viral infections (EBV, VZV), EBV lymphoproliferation, B-cell non-Hodgkin lymphoma
CD137 deficiency (41BB)	<u>TNFRSF9</u>	AR	<u>602250</u>	Normal	Normal	Low IgG, low IgA, poor responses to T cell-dependent and T cell independent antigens, decreased T cell proliferation,IFNy secretion, cytotoxicity	EBV lymphoproliferation, B- cell lymphoma, chronic active EBV infection
RASGRP1 deficiency	RASGRP1	AR	<u>603962</u>	Poor activation, proliferation, motility. Reduced naïve T cells	Poor activation, proliferation, motility	Normal IgM, IgG, increased IgA	Recurrent pneumonia, herpesvirus infections, EBV associated lymphoma Decreased NK cell function
RLTPR deficiency	CARMIL2	AR	610859	Normal number, high CD4, increased naïve CD4* and CD8*, low Treg and MAIT, poor CD28- induced function	Normal B cell numbers, reduced memory B cells	Normal to low, poor T dependent antibody response	Recurrent bacterial, fungal and mycobacterial infections, viral warts, molluscum and EBV lymphoproliferative and other malignancy, atopy
X-linked magnesium EBV and neoplasia (XMEN)	MAGT1	XL	300853	Low CD4 Low recent thymic emigrant cells, inverted CD4/CD8 ratio, reduced MAIT cells, poor proliferation to CD3	Normal but decreased memory B cells	Progressive hypogarmaglobulinemia Reduced NK cell and CTL cytotoxic activity due to impaired expression of NKG2D	EBV infection, lymphoma, viral infections, respiratory and GI infections Glycosylation defects
PRKCD deficiency	PRKCD	AR	615559	Normal	Low memory B cells, high CD5 B cells	Apoptotic defect in B cells	Recurrent infections, EBV chronic infection, lymphoproliferation, SLE-like autoimmunity (nephrotic and antiphospholipid syndromes), low lgG
TET2 deficiency (3 patients)	TET2	AR	619126	Increased CD4·CD8· T cells	Low memory B cells	DNA hypermethylation, defective FAS-mediated apoptosis	ALPS-like, recurrent viral infections, EBV viremia, lymphadenopathy, hepatospienomegaly, autoimmunity, B-lymphoma, FTT, developmental delay

Total number of mutant genes in Table IV: 52. New inborn errors of immunity: 7 (RHOG [43], CEBPE [51], AD GOF IKZF1 [52], SOCS1 [44-46], PDCD1 [47], ELF4 [48], TET2 [50])

FHL familial hemophagocytic lymphohistiocytosis, HLH hemophagocytic lymphohistiocytosis, HSM hepatosplenomegaly, DN double-negative, SLE systemic lupus erythematous, IBD inflammatory bowel disease



 Table 5
 Congenital defects of phagocyte number or function

		1. Con	genital Ne	eutropenias		
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features
Elastase deficiency (Severe congenital neutropenia [SCN] 1)	ELANE	AD	130130	N	Myeloid differentiation	Susceptibility to MDS/leukemia Severe congenital neutropenia or cyclic neutropenia
GFI 1 deficiency (SCN2)	GFI1	AD	<u>600871</u>	N	Myeloid differentiation	B/T lymphopenia
HAX1 deficiency (Kostmann Disease) (SCN3)	HAX1	AR	605998	N	Myeloid differentiation	Cognitive and neurological defects in patients with defects in both HAX1 isoforms, susceptibility to MDS/leukemia
G6PC3 deficiency (SCN4)	G6PC3	AR	<u>611045</u>	N	Myeloid differentiation, chemotaxis, O ₂ - production	Structural heart defects, urogenital abnormalities, inner ear deafness, and venous angiectasias of trunks and limbs
VPS45 deficiency (SCN5)	VPS45	AR	<u>610035</u>	N	Myeloid differentiation, migration	Extramedullary hematopoiesis, bone marrow fibrosis, nephromegaly
Glycogen storage disease type 1b	G6PT1	AR	602671	N + M	Myeloid differentiation, chemotaxis, O ₂ - production	Fasting hypoglycemia, lactic acidosis, hyperlipidemia, hepatomegaly
X-linked neutropenia/myelodysplasia	WAS	XL GOF	300299	N	Differentiation, mitosis. Results from GOF mutations in GTPase binding domain of WASp	Neutropenia, myeloid maturation arrest, monocytopenia, variable lymphoid anomalies
P14/LAMTOR2 deficiency	LAMTOR2	AR	610389	N + M	Endosomal biogenesis	Neutropenia Hypogammaglobulinemia ↓CD8 cytotoxicity, partial albinism, growth failure
Barth Syndrome (3-Methylglutaconic aciduria type II)	TAZ	XL	300394	N+L Mel	Mitochondrial function	Cardiomyopathy, myopathy, growth retardation, neutropenia
Cohen syndrome	VPS13B	AR	607817	N	Myeloid differentiation	Dysmorphism, mental retardation, obesity, deafness, neutropenia
Clericuzio syndrome (Poikiloderma with neutropenia)	USB1	AR	<u>613276</u>	N	Myeloid differentiation	Retinopathy, developmental delay, facial dysmorphisms, poikiloderma
JAGN1 deficiency	JAGN1	AR	616012	N	Myeloid differentiation	Myeloid maturation arrest, osteopenia
3-Methylglutaconic aciduria	CLPB	AR	616254	N	Myeloid differentiation Mitochondrial protein	Neurocognitive developmental aberrations, microcephaly, hypoglycemia, hypotonia, ataxia, seizures, cataracts, IUGR
G-CSF receptor deficiency	CSF3R	AR	<u>138971</u>	N	Stress granulopoiesis disturbed	
SMARCD2 deficiency	SMARCD2	AR	601736	N	Chromatin remodeling, Myeloid differentiation and neutrophil functional defect	Neutropenia, developmental aberrations, bones, hematopoietic stem cells, myelodysplasia
Specific granule deficiency	CEBPE	AR	<u>189965</u>	N	Terminal maturation and global dysfunction	Neutropenia, Neutrophils with bilobed nuclei
Shwachman-Diamond Syndrome	SBDS	AR	607444	N	Neutrophil maturation,	Pancytopenia, exocrine pancreatic insufficiency, chondrodysplasia
	DNAJC21	AR	<u>617052</u>	N + HSC	chemotaxis, ribosomal	Pancytopenia, exocrine
	EFL1	AR	617941	N + HSC	biogenesis	pancreatic insufficiency
HYOU1 deficiency	HYOU1	AR	<u>601746</u>	N	Unfolded protein response	Hypoglycemia, inflammatory complications
SRP54 deficiency	SRP54	AD	604857	N	Protein translocation to ER, myeloid differentiation and neutrophil functional defect	Neutropenia, exocrine pancreatic insufficiency
CXCR2 deficiency (6 patients)	CXCR2	AR	<u>619407</u>	N	Reduced expression of CXCR2 on patient cells, impaired responses to CXCL8	Profound neutropenia, myelokathexis, recurrent gingivitis, oral ulcers, hypergammaglobulinemia

	2. Defects of Motility										
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features					
Leukocyte adhesion deficiency type 1 (LAD1)	ITGB2	AR	600065	N + M + L + NK	Adherence, Chemotaxis, Endocytosis, T/NK cytotoxicity	Delayed cord separation, skin ulcers, periodontitis, leukocytosis					
Leukocyte adhesion deficiency type 2 (LAD2)	SLC35C1	AR	605881	N + M	Rolling, chemotaxis	Mild LAD type 1 features with hh-blood group, growth retardation, developmental delay					
Leukocyte adhesion deficiency type 3 (LAD3)	FERMT3	AR	607901	N + M + L + NK	Adherence, chemotaxis	LAD type 1 plus bleeding tendency					
Rac2 deficiency	RAC2	AD LOF	608203	N	Adherence, chemotaxis O ₂ - production	Poor wound healing, leukocytosis					
β actin deficiency	ACTB	AD	102630	N + M	Motility	Mental retardation, short stature					
Localized juvenile periodontitis	FPR1	AR	136537	N	Formylpeptide induced chemotaxis	Periodontitis only					
Papillon-Lefèvre Syndrome	CTSC	AR	602365	N + M	Chemotaxis	Periodontitis, palmoplantar hyperkeratosis in some patients					
WDR1 deficiency	WDR1	AR	604734	N	Spreading, survival, chemotaxis	Mild neutropenia, poor wound healing, severe stomatitis, neutrophil nuclei herniate					
Cystic fibrosis	CFTR	AR	602421	M only	Chemotaxis	Respiratory infections, pancreatic insufficiency, elevated sweat chloride					
Neutropenia with combined immune deficiency due to MKL1 deficiency	MKL1	AR	606078	N + M +L + NK	Impaired expression of cytoskeletal genes	Mild thrombocytopenia					



Table 5 (continued)

	3.Defects of Respiratory Burst											
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features						
X-linked chronic granulomatous disease (CGD), gp91phox	CYBB	XL	306400			Infections, autoinflammatory phenotype, IBD McLeod phenotype in patients with deletions extending into the contiguous Kell locus						
	CYBA		608508	N + M	Killing (faulty O ₂ - production)							
	CYBC1		618334			Infections, autoinflammatory						
Autosomal recessive CGD	NCF1	AR	608512			phenotype						
,	NCF2		<u>608515</u>									
	NCF4		613960									
G6PD deficiency class I	G6PD	XL	305900	N	Reduced O2- production	Infections						

4. Other Non-Lymphoid Defects										
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features				
GATA2 deficiency	GATA2	AD	<u>137295</u>	Monocytes + peripheral DC	Multi lineage cytopenias	Susceptibility to mycobacteria, HPV, histoplasmosis, alveolar proteinosis, MDS/AML/CMML, lymphedema				
Pulmonary alveolar proteinosis	CSF2RA	XL (Biallelic mutations in pseudo- autosomal gene)	300770	Alveolar macrophages	GM-CSF signaling	Alveolar proteinosis				
	CSFR2B	AR	<u>614370</u>]						

Total number of mutant genes in Table 5: 42. New inborn errors of immunity: 1 (CXCR2 [53, 54]). Removed: Cyclic neutropenia was merged with elastase deficiency

MDS myelodysplastic syndrome, IUGR intrauterine growth retardation, LAD leukocyte adhesion deficiency, AML acute myelogenous leukemia, CMML chronic myelomonocytic leukemia, N neutrophil, M monocyte, MEL melanocyte, L lymphocyte, NK natural killer



 Table 6
 Defects in intrinsic and innate immunity

	1. Men	delian Sus	ceptibility	to mycobac	cterial disease (MSMD)
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features
IL-12 and IL-23 receptor β1 chain deficiency	IL12RB1	AR	<u>601604</u>	L + NK		
IL-12p40 (IL-12 and IL-23) deficiency	IL12B	AR	<u>161561</u>	M	IFN-y secretion	
IL-12Rβ2 deficiency	IL12RB2	AR	<u>601642</u>	L + NK	IFIN-γ Secretion	Susceptibility to mycobacteria and Salmonella
IL-23R deficiency	IL23R	AR	<u>607562</u>	L +NK		
IFN-γ receptor 1 deficiency	IFNGR1	AR	<u>209950</u>	M + L		
	IFNGRI	AD	615978	M + L	IFN-γ binding and signaling	
IFN-γ receptor 2 deficiency	IFNGR2	AR	<u>147569</u>	M + L	IFN-γ signaling	
STAT1 deficiency	STAT1	AD LOF	<u>614892</u>	M + L		
Macrophage gp91 phox deficiency	CYBB	XL	300645	Macrophage only	Killing (faulty O ₂ - production)	Isolated susceptibility to mycobacteria
		AD	614893	M + L	Impaired development of cDCs and Th1* cells	Susceptibility to mycobacteria
IRF8 deficiency	IRF8	AR	226990	M	Lack of circulating monocytes and DCs, reduced NK cell numbers and function reported in some patients	Susceptibility to mycobacteria and multiple other infectious agents including EBV
SPPL2a deficiency	SPPL2A	AR	608238	M + L	Impaired development of cDCs and Th1* cells	Susceptibility to mycobacteria and Salmonella
Tyk2 deficiency	TYK2	AR	<u>611521</u>	M + L	Impaired cellular responses to IL-10, IL-12, IL-23, and type I IFNs	Susceptibility to intracellular bacteria (mycobacteria, Salmonella), and viruses
P1104A TYK2 homozygosity	TYK2	AR	176941	L	Impaired cellular responses to IL-23	MSMD or tuberculosis
ISG15 deficiency	ISG15	AR	<u>147571</u>		IFNγ production defect	Susceptibility to mycobacteria (BCG), brain calcification
RORγt deficiency	RORC	AR	602943	L + NK	Lack of functional RORyT protein, IFNy production defect, complete absence of IL-17A/F-producing T cells	Susceptibility to mycobacteria and candida
JAK1 deficiency	JAK1	AR LOF	<u>147795</u>	N + L	Reduced JAK1 activation to cytokines, Reduced IFN _γ production	Susceptibility to mycobacteria and viruses, urothelial carcinoma
T-bet deficiency (1 patient)	TBX21	AR	<u>619630</u>	L	↓ IFN-γ and TNF-α production by γδ T cells, MAIT cells, iNKT cells, NK cells, and CD4* T cells	Susceptibility to mycobacteria
IFNγ deficiency (2 patients)	IFNG	AR	<u>618963</u>	L	No IFN-γ production by patient T and NK cells	Susceptibility to mycobacteria

	2. Epidermodysplasia verruciformis (HPV)									
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features				
EVER1 deficiency	TMC6		605828		EVER1, EVER2 and CIB1 form a complex in	Human papillomavirus (HPV) (group B1)				
EVER2 deficiency	TMC8	AR	<u>605829</u>	Keratinocytes	keratinocytes	infections and cancer of the skin (typical EV)				
CIB1 deficiency	CIB1		<u>618267</u>	,		, ,				
WHIM (Warts, Hypogammaglobulinemia, infections, Myelokathexis) syndrome	CXCR4	AD GOF	<u>162643</u>	Leukocytes	Increased response of the CXCR4 chemokine receptor to its ligand CXCL12 (SDF-1)	Warts (HPV) infection, neutropenia, low B cell number, hypogammaglobulinemia				

			3. Pre	disposition to Severe	Viral Infection		
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features	
STAT1 deficiency	STAT1	AR LOF	600555	Leukocytes and other cells	STAT1-dependent IFN- α/β γ and λ responses	Severe viral infections, mycobacterial infection	
STAT2 deficiency	STAT2	AR	600556	Leukocytes and other cells	STAT2-dependent IFN-α/β and λ response	Severe viral infections (disseminated vaccine- strain measles)	
IRF9 deficiency	IRF9	AR	<u>147574</u> *	Leukocytes and other cells	IRF9- and ISGF3-dependent IFN- α/β and λ responses		
IRF7 deficiency	IRF7	AR	605047	Leukocytes, plasmacytoid dendritic cells, non- hematopoietic cells	IFN- α , β and γ production and IFN- λ production	Severe influenza disease	
IFNAR1 deficiency	IFNAR1	AR	<u>107450</u> *	Leukocytes and other cells	IFNAR1-dependent responses to IFN-α/β	Severe disease caused by Yellow Fever vaccine and Measles vaccine	
IFNAR2 deficiency	IFNAR2	AR	602376	Broadly expressed	IFNAR2-dependent responses to IFN-α/β	Severe viral infections (disseminated vaccine- strain measles, HHV6)	
CD16 deficiency	FCGR3A	AR	146740	NK cells	Altered NK cells function	Severe herpes viral infections, particularly VZV, Epstein Barr virus (EBV), and (HPV)	
MDA5 deficiency	IFIH1	AR LOF	606951	Broadly expressed	Viral recognition and IFN induction	Rhinovirus and other RNA viruses	
NOS2 deficiency (1 patient)	NOS2	AR	<u>NA</u>	Myeloid cells	Mutant NOS2 failed to induce nitrous oxide	Severe (fatal) susceptibility to CMV- induced disease; pneumocystis pneumonia secondary to CMV; intact responses to infection with other herpes viruses (EBV, VZV, HSV)	
ZNFX1 deficiency (28 patients)	ZNFX1	AR	<u>619644</u>	Broadly expressed	↑ ISG in response to poly I/C	Severe infections by RNA/DNA viruses, mycobacteria; early-onset severe inflammation affecting liver, brain, kidneys, lungs; virally triggered inflammatory episodes, hepatosplenomegaly, lymphadenopathy	
RNA polymerase III	POLR3A	AD	614258	1	Impaired viral recognition and IFN	Severe VZV infection	
deficiency	POLR3C POLR3F	AD AD	617454 617455	Leukocytes and other cells	induction in response to VZV or poly I:C	Severe vzv intection	



Table 6 (continued)

			4. Herpes	s Simplex Encephalitis (HSE)		
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features	
		AD			TLR3-dependent IFN- α , β and γ response	Herpes simplex virus 1 encephalitis (incomplete clinical penetrance for all	
TLR3 deficiency	TLR3	AR	613002			etiologies listed here); severe pulmonary influenza; VZV	
UNC93B1 deficiency	UNC93B1	AR	608204		UNC-93B-dependent IFN- α , β and γ response		
TRAF3 deficiency	TRAF3	AD	601896		TRAF3-dependent IFN- α , β and γ response		
TRIF deficiency	TICAM1	AD AR	<u>607601</u>	Central nervous system (CNS) resident cells and fibroblasts	TRIF-dependent IFN- α , β and γ response	Herpes simplex virus 1 encephalitis	
TBK1 deficiency	TBK1	AD	604834		TBK1-dependent IFN- α , β and γ response		
IRF3 deficiency	IRF3	AD	<u>616532</u>		Low IFN –α/β production in response to HSV1 and decreased IRF3 phosphorylation		
DBR1 deficiency	DBR1	AR	607024		Impaired production of anti-viral IFNs	HSE of the brainstem. Other viral infections of the brainstem.	
SNORA31 deficiency (5 patients)	SNORA31	AD	<u>619396</u>		Impaired production of anti-viral IFNs	Forebrain HSV1 encephalitis	
ATG4A deficiency (1 patient	ATG4	45		Central nervous system (CNS)	Impaired HSV2-induced autophagy → increased viral replication and apoptosis of patient fibroblasts	Mollaret's meningitis (recurrent lymphocytic meningitis) due to HSV2	
MAP1LC3B2 deficiency (1 patient	MAP1LC3B2	AD	<u>NA</u>	resident cells and fibroblasts			

5. Predisposition to INVASIVE Fungal Diseases										
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features				
CARD9 deficiency	CARD9	AR	607212	Mononuclear phagocytes	CARD9 signaling pathway	Invasive candidiasis infection, deep dermatophytoses, other invasive fungal infections				

		6. Pre	edisposit	tion to Mucocutaneo	us Candidiasis	
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Affected function	Associated features
IL-17RA deficiency	IL17RA	AR	605461	Epithelial cells, fibroblasts, mononuclear phagocytes	IL-17RA signaling pathway	CMC, folliculitis
IL-17RC deficiency	IL17RC	AR	610925		IL-17RC signaling pathway	CMC
IL-17F deficiency	IL17F	AD	606496	T cells	IL-17F-containing dimers	CMC, folliculitis
STAT1 GOF	STAT1	AD GOF	600555	T cells, B cells, monocytes	Gain-of-function STAT1 mutations that impair the development of IL-17-producing T cells	CMC, various fungal, bacterial and viral (HSV) infections, auto-immunity (thyroiditis, diabetes, cytopenias), enteropathy
ACT1 deficiency	TRAF3IP2	AR	607043	T cells, fibroblasts	Fibroblasts fail to respond to IL-17A and IL-17F, and their T cells to IL-17E	CMC, blepharitis, folliculitis and macroglossia
JNK1 haplo- insufficiency (3 patients)	MAPK8	AD	NA	T cells, fibroblasts	↓ Th17 cells ex vivo, in vitro, ↓ responses of fibroblasts to IL-17A, IL-17F, ↓ c-Jun/ATF-2- dependant TGF β signaling	CMC, connective tissue disorder (similar to Ehlers-Danlos syndrome)

	7.	TLR Signali	ng Pathwa	ay Deficiency with	Bacterial Susceptibility	
Disease	Genetic defect	Inheritance	OMIM	Affected cells	Affected function	Associated features
IRAK4 deficiency	IRAK4	AR	606883	Lymphocytes + Granulocytes+ Monocytes	TIR-IRAK4 signaling pathway	
MyD88 deficiency	MYD88	AR	602170	Lymphocytes + Granulocytes+ Monocytes	TIR-MyD88 signaling pathway	Bacterial infections (pyogenes)
IRAK1 deficiency	IRAK1	XL	300283	Lymphocytes + Granulocytes+ Monocytes	TIR-IRAK1 signaling pathway	Bacterial infections, X-linked MECP2 deficiency-related syndrome due to a large de novo Xq28 chromosomal deletion encompassing both MECP2 and IRAK1
TIRAP deficiency	TIRAP	AR	614382	Lymphocytes + Granulocytes+ Monocytes	TIRAP- signaling pathway, TLR1/2, TLR2/6, and TLR4 agonists were impaired in the fibroblasts and leukocytes	Staphylococcal disease during childhood
TLR7 deficiency	TLR7	XL	<u>301051</u>	Lymphocytes, Myeloid cells	impaired responses to TLR7 ligands; reduced production of type 1 IFN	Severe COVID19 infection
TLR8 GOF	TLR8	XL	<u>NA</u>	Myeloid cells	Elevated proinflammatory serum cytokines; increased pro- inflammatory responses of patient myeloid cells to TLR8 agonists; reduced ability of mutant TLR8 to attenuate TLR7 signalling	Early onset, severe cytopenias, hepatosplenomegaly, lymphadenopathy; progressive autoinflammatory disease

8.	Other Inborn E	Errors of In	nmunity Re	lated to Non	-Hematopoietic Tissue	s
Disease	Genetic defect	Inheritance	Gene OMIM	Affected cells	Affected function	Associated features
Isolated congenital asplenia (ICA)	RPSA	AD	<u>271400</u>	No spleen	RPSA encodes ribosomal protein SA, a component of the small subunit of the ribosome	Bacteremia (encapsulated bacteria)
	НМОХ	AR	141250	Macrophages	HO-1 regulates iron recycling and heme-dependent damage occurs	Hemolysis, nephritis, inflammation
Trypanosomiasis	APOL1	AD	603743	Somatic	Pore forming serum protein	Trypanosomiasis



Table 6 (continued)

Acute liver failure due to NBAS deficiency	NBAS	AR	608025	Somatic and hematopoietic	ER stress	Fever induces liver failure
Acute necrotizing encephalopathy	RANBP2	AR	<u>601181</u>	Ubiquitous expression	Nuclear pore	Fever induces acute encephalopathy
	CLCN7	AR	602727		Secretory lysosomes	Osteopetrosis with hypocalcemia, neurologic features
	SNX10	AR	614780			Osteopetrosis with visual impairment
Osteopetrosis	OSTM1	AR	607649	Osteoclasts		Osteopetrosis with hypocalcemia, neurologic features
	PLEKHM1	AR	<u>611466</u>			Osteopetrosis
	TCIRG1	AR	604592			Osteopetrosis with hypocalcemia
	TNFRSF11A	AR	603499		Osteoclastogenesis	Osteopetrosis
	TNFSF11	AR	602642	Stromal	Osteoclastogenesis	Osteopetrosis with severe growth retardation
	NCSTN	AD	605254		Notch signaling/ Gamma- secretase in hair follicle	Verneuil's disease/ Hidradenitis suppurativa with acne
Hidradenitis suppurativa	PSEN	AD	<u>613737</u>	Epidermis	regulates keratinization	Verneuil's disease/ Hidradenitis suppurative with cutaneous hyperpigmentation
	PSENEN	AD	613736			Verneuil's disease/ Hidradenitis suppurativa

	9. Other Inborn Errors of Immunity Related to Leukocytes										
Disease Genetic defect Inheritance Gene OMIM Affected cells Affected function Associated features											
IRF4 haploinsufficiency	IRF4	AD	601900	L+M	IRF4 is a pleiotropic transcription factor	Whipple's disease					
IL-18BP deficiency	IL18BP	AR	604113	Leukocytes and other cells	IL-18BP neutralizes secreted IL- 18	Fulminant viral hepatitis					

Total number of mutant genes in Table 6: 74. New inborn errors of immunity: 10 (TBX21 [55], IFNG [57], NOS2 [60], ZNFX1 [63–65], SNORA31 [61], ATG4A, MAP1LC3B2 [62], MAPK8 [69], TLR7 [66–68], TLR8 [58, 59])

NF-κB nuclear factor kappa B, TIR Toll and interleukin 1 receptor, IFN interferon, TLR Toll-like receptor, MDC myeloid dendritic cell, CNS central nervous system, CMC chronic mucocutaneous candidiasis, HPV human papillomavirus, VZV varicella zoster virus, EBV Epstein-Barr virus



Table 7 Autoinflammatory disorders

			1	. Type 1 Int	erferonopa	thies	
Disease	Genetic defect	Inheritance	ОМІМ	T Cells	B cells	Functional defect	Associated Features
AD STING-associated vasculopathy, infantile- onset (SAVI)	TMEM173 (STING)	AD	<u>612374</u>	Not assessed	Not assessed	STING activates both the NF- kappa-B and IRF3 transcription pathways to induce expression of IFN	Skin vasculopathy, inflammatory lung disease, systemic autoinflammation and ICC, FCL
AR STING-associated vasculopathy, infantile- onset (SAVI)	TMEM173 (STING)	AR GOF	<u>615934</u>	Not assessed	Not assessed	STING activates both the NF- kappa-B and IRF3 transcription pathways to induce expression of IFN	FTT, early onset rash, fever, dyspnea, interstitial lung disease/pneumonitis, polyarthritis, autoAbs, increased inflammatory markers, IFN gene signature. Phenocopy of SAVI due to AD GOF TMEM173
ADA2 deficiency	ADA2	AR	<u>607575</u>	Not assessed	Not assessed	ADAs deactivate extracellular adenosine and terminate signaling through adenosine receptors	Polyarteritis nodosa, childhood-onset, early-onset recurrent ischemic stroke and fever; some patients develop hypogammaglobulinemia
TREX1 deficiency, Aicardi-Goutières syndrome 1 (AGS1)	TREX1	AR AD	<u>606609</u>	Not assessed	Not assessed	Intracellular accumulation of abnormal ss DNA species leading to increased type I IFN production	Classical AGS, SLE, FCL
RNASEH2B deficiency, AGS2	RNASEH2B	AR	610326	Not assessed	Not assessed	Intracellular accumulation of	Classical AGS, SP
RNASEH2C deficiency, AGS3	RNASEH2C	AR	610330	Not assessed	Not assessed	abnormal RNA-DNA hybrid species leading to increased type	Classical AGS
RNASEH2A deficiency, AGS4	RNASEH2A	AR	606034	Not assessed	Not assessed	I IFN production	Classical AGS
SAMHD1 deficiency, AGS5	SAMHD1	AR	606754	Not assessed	Not assessed	Controls dNTPs in the cytosol, failure of which leads to increased type I IFN production	Classical AGS, FCL
ADAR1 deficiency, AGS6	ADAR1	AR	146920	Not assessed	Not assessed	Catalyzes the deamination of adenosine to inosine in dsRNA substrates, failure of which leads to increased type I IFN production	Classical AGS, BSN, SP
Aicardi-Goutières syndrome 7 (AGS7)	IFIH1	AD GOF	<u>615846</u>	Not assessed	Not assessed	IFIH1 gene encodes a cytoplasmic viral RNA receptor that activates type I interferon signaling through the MAVS adaptor molecule	Classical AGS, SLE, SP, SMS
DNAse II deficiency	DNASE2	AR	126350	Not assessed	Not assessed	DNAse II degrades and eliminates DNA. Loss of DNase II activity induces type I interferon signaling	AGS
LSM11 deficiency (2 patients)	LSM11	AR	619486	Not assessed	Not assessed	Increased IFN signaling in fibroblasts	AGS, type 1 IFN-opathy
RNU7-1 deficiency (16 patients)	RNU7-1	AR	619487	Not assessed	Not assessed	Increased IFN signaling in fibroblasts	AGS, type 1 IFN-opathy
Pediatric systemic lupus erythematosus due to DNASE1L3 deficiency	DNASE1L3	AR	<u>614420</u>			DNASE1L3 is an endonuclease that degrades extracellular DNA. DNASE1L3 deficiency decreases clearance of apoptotic cells	Very early onset SLE, reduced complement levels, autoantibodies (dsDNA, ANCA), lupus nephritis, hypocomplementemic urticarial vasculitis syndrome
Spondyloenchondro- dysplasia with immune dysregulation (SPENCD)	ACP5	AR	<u>171640</u>	Not assessed	Not assessed	Upregulation of IFN through mechanism possibly relating to pDCS	Short stature, SP, ICC, SLE, thrombocytopenia and autoimmune hemolytic anemia, possibly recurrent bacterial and viral infections
X-linked reticulate pigmentary disorder	POLA1	XL	301220	Not assessed	Not assessed	POLA1 is required for synthesis of cytosolic RNA:DNA and its deficiency leads to increase production of type I interferon	Hyperpigmentation, characteristic facies, lung and GI involvement
USP18 deficiency	USP18	AR	607057	Not assessed	Not assessed	Defective negative regulation of ISG15 leading to increased IFN	TORCH like syndrome
OAS1 deficiency	OAS1	AD GOF	<u>164350</u>		Low	Increased interferon from recognition of RNA	Pulmonary alveolar proteinosis, skin rash
CDC42 deficiency (15 patients)	CDC42	AD	<u>616737</u>	Normal/ decreased	Normal/ decreased	↑ serum levels of IL1, IL18, IFN- γ, ferritin, sCD25, CRP etc. Mutation affects actin function, ↓ NK cell cytotoxicity	Neonatal onset: pancytopenia, fever, rash, hepatosplenomegaly, multisystemic inflammation, myelofibrosis/proliferation, HLH, enterocolitis; Recurrent GIT/URT infections; neurodevelopmental delay, FTT
STAT2 R148 LOF/regulation (3 patients)	STAT2	AR	<u>616636</u>	Increased	Normal	Patient cells hyper-sensitive to IFN-a, GOF for induction of the late (not early) response to type 1 IFNs due to impaired interaction of mutant STAT2 with USP18, a negative regulator of type 1 IFN responses	Severe fatal early onset autoinflammation, ↑ serum IFN-α, IL6, TNFα, phenocopy of USP18 deficiency
ATAD3A deficiency (8 patients)	ATAD3A	AD/AR	617183	Not assessed	Not assessed	Elevated ISG expression, increased serum type 1 IFNs	Predominantly neurological defects (development delay, spasticity)

2. Defects Affecting the Inflammasome										
Disease	Associated Features									
Familial Mediterranean fever	MEFV	AR LOF	<u>249100</u>	Mature granulocytes, cytokine-activated monocytes.	Increased inflammasome-mediated induction of IL1β.	Recurrent fever, serositis and inflammation responsive to colchicine. Predisposes to vasculitis and inflammatory bowel disease.				



Table 7 (continued)

		AD	<u>134610</u>	Mature granulocytes, cytokine-activated monocytes.	Usually M694del variant.	
Mevalonate kinase deficiency (Hyper IgD syndrome)	MVK	AR	260920	Somatic and hematopoietic	affecting cholesterol synthesis, pathogenesis of disease unclear	Periodic fever and leukocytosis with high IgD levels
Muckle-Wells syndrome		AD GOF	<u>191900</u>	PMNs Monocytes		Urticaria, SNHL, amyloidosis.
Familial cold autoinflammatory syndrome 1	NLRP3	AD GOF	120100	PMNs, monocytes	Defect in cryopyrin, involved in	Non-pruritic urticaria, arthritis, chills, fever and leukocytosis after cold exposure.
Neonatal onset multisystem inflammatory disease (NOMID) or chronic infantile neurologic cutaneous and articular syndrome (CINCA)		AD GOF	607115	PMNs, chondrocytes	leukocyte apoptosis and NFkB signaling and IL-1 processing	Neonatal onset rash, chronic meningitis, and arthropathy with fever and inflammation.
Familial cold autoinflammatory syndrome 2	NLRP12	AD GOF	611762	PMNs, monocytes		Non-pruritic urticaria, arthritis, chills, fever and leukocytosis after cold exposure.
NLRC4-MAS (macrophage activating syndrome) Familial cold autoinflammatory syndrome 4	NLRC4	AD GOF	616050 616115	PMNs monocytes macrophages	Gain of function mutation in NLRC4 results in elevated secretion of IL-1β and IL-18 as well as macrophage	Severe enterocolitis and macrophage activation syndrome
PLAID (PLC ₂ 2 associated antibody deficiency and immune dysregulation) Familial cold autoinflammatory syndrome 3 or APLAID (c2120A>C)	PLCG2	AD GOF	614878 614468	B cells, NK, Mast cells	activation Mutations activate IL-1 pathways	Cold urticaria hypogammaglobulinemia, impaired humoral immunity, autoinflammation
NLRP1 deficiency	NLRP1	AR	617388	leukocytes	Systemic elevation of IL-18 and caspase 1, suggesting involvement of NLRP1 inflammasome	Dyskeratosis, autoimmunity and arthritis
NLRP1 GOF	NLRP1	AD GOF	<u>615225</u>	Keratinocytes	Increased IL1β	Palmoplantar carcinoma, corneal scarring; recurrent respiratory papillomatosis
RIPK1 deficiency (12 patients)	RIPK1	AD	618852		↑ inflammatory markers and pro- inflammatory cytokines/gene signature	Autoinflammatory disorder: regular/prolonged fevers, lymphadenopathy, spleno/hepatomegaly, ulcers, arthralgia, Gl features,

			3. Non-Inf	lammasome Rela	ted Conditions	
Disease	Genetic defect	Inheritance	ОМІМ	Affected cells	Functional defects	Associated Features
TNF receptor-associated periodic syndrome (TRAPS)	TNFRSF1A	AD	142680	PMNs, monocytes	Mutations of 55-kD TNF receptor leading to intracellular receptor retention or diminished soluble cytokine receptor available to bind TNF	Recurrent fever, serositis, rash, and ocular or joint inflammation
Pyogenic sterile arthritis, pyoderma gangrenosum, acne (PAPA) syndrome, hyperzincemia and hypercalprotectinemia	PSTPIP1	AD	604416	Hematopoietic tissues, upregulated in activated T-cells	Disordered actin reorganization leading to compromised physiologic signaling during inflammatory response	Destructive arthritis, inflammatory skin rash, myositis
Blau syndrome	NOD2	AD	<u>186580</u>	Monocytes	Mutations in nucleotide binding site of CARD15, possibly disrupting interactions with lipopolysaccharides and NF-kB signaling	Uveitis, granulomatous synovitis, camptodactyly, rash and cranial neuropathies, 30% develop Crohn colitis
ADAM17 deficiency	ADAM17	AR	614328	Leukocytes and epithelial cells	Defective TNFα production	Early onset diarrhea and skin lesions
Chronic recurrent multifocal osteomyelitis and congenital dyserythropoietic anemia (Majeed syndrome)	LPIN2	AR	609628	Neutrophils, bone marrow cells	Undefined	Chronic recurrent multifocal osteomyelitis, transfusion-dependent anemia, cutaneous inflammatory disorders
DIRA (Deficiency of the Interleukin 1 Receptor Antagonist)	IL1RN	AR	612852	PMNs, Monocytes	Mutations in the IL1 receptor antagonist allow unopposed action of Interleukin 1	Neonatal onset of sterile multifocal osteomyelitis, periostitis and pustulosis.
DITRA (Deficiency of IL- 36 receptor antagonist)	IL36RN	AR	<u>614204</u>	Keratinocytes, leukocytes	Mutations in IL-36RN leads to increase IL-8 production	Pustular psoriasis
SLC29A3 mutation	SLC29A3	AR	602782	Leukocytes, bone cells		Hyperpigmentation hypertrichosis, histiocytosis-lymphadenopathy plus syndrome
CAMPS (CARD14 mediated psoriasis)	CARD14	AD	602723	Mainly in keratinocytes	Mutations in CARD14 activate the NF-kB pathway and production of IL-8	Psoriasis
Cherubism	SH3BP2	AD	<u>118400</u>	Stroma cells, bone cells	Hyperactive macrophage and increase NF - kB	Bone degeneration in jaws



Table 7 (continued)

CANDLE (chronic atypical neutrophilic	PSMB8*	AR and AD	<u>256040</u>	Keratinocytes, B cell adipose cells	Mutations cause increased IFN signaling	Contractures, panniculitis, ICC, fevers	
dermatitis with lipodystrophy)	PSMG2	AR	609702	Lymphocytes	through an undefined mechanism	Panniculitis, lipodystrophy, autoimmune hemolytic anemia	
COPA defect	COPA	AD	6011924	PMN and tissue specific cells	Defective intracellular transport via the coat protein complex I (COPI)	Autoimmune inflammatory arthritis and interstitial lung disease with Th17 dysregulation and autoantibody production	
Otulipenia/ORAS	OTULIN	AR	615712	Leukocytes	Increase LUBAC induction of NF-KB activation leading to high proinflammatory cytokines levels.	Fever, diarrhea , dermatitis	
A20 deficiency	TNFAIP3	AD	<u>616744</u>	Lymphocytes	Defective inhibition of NF-KB signaling pathway	Arthralgia, mucosal ulcers, ocular inflammation	
AP1S3 deficiency	AP1S3	AR	<u>615781</u>	Keratinocytes	Disrupted TLR3 translocation	Pustular psoriasis	
ALPI deficiency	ALPI	AR	<u>171740</u>	Intestinal epithelial cells	Deficient inhibition of LPS in intestine	Inflammatory bowel disease	
TRIM22	TRIM22	AR	606559	Macrophages, intestinal epithelial cells	Granulomatous colitis	Inflammatory bowel disease	
T-cell lymphoma subcutaneous panniculitis-like (TIM3 deficiency)	HAVCR2	AR	<u>618398</u>	Leukocytes	Increased inflammasome activity due to defective checkpoint signaling	Panniculitis, HLH, polyclonal cutaneous T cell infiltrates or T-cell lymphoma	
C2orf69 deficiency (28 patients)	C2orf69	AR	619423			Early onset severe autoinflammation disorder, often fatal. Global developmental delay, with recurrent seizures, Muscle weakness. Liver dysfunction,	
NCKAP1L deficiency (9 patients	NCKAP1L	AR	618982	Lymphocytes	Hyperinflammation and cytokine overproduction (↑ Th1), ↓ T cell proliferation, cytoskeletal defects	Recurrent URTI, skin rashes/abscesses/ atopy, ulcers, lymphoproliferation/ lymphadenopathy, hyperinflammation, anti dsDNA Abs, fever, FTT	
SYK GOF (6 patients)	syk	AD GOF	<u>619381</u>	Lymphocytes	Increased SYK phosphorylation, enhance downstream signaling	Recurrent infections, multi-organ inflammation/inflammatory disease (gut, skin, CNS, lung, liver), B cell lymphoma (2 pts)	
HCK GOF (1 patient)	нск	AD GOF	<u>NA</u>		Increased kinase activity of HCK mutant in vitro; ↑ production of inflammatory cytokines (IL-1β, IL-6, IL-8, TNF-α), ROS	cutaneous vasculitis, inflammatory leukocyte infiltration of the lungs (pulmonary fibrosis) and skin, anemia, hepatosplenomegaly	
PSMB9 GOF (3 patients)	PSMB9	AD GOF	<u>617591</u>	Mild pancytopenia; Leukocytes	Elevated levels of inflammatory cytokines (IL-6, IL-18, IP-10, IFN α), liver enzymes in blood and CSF (IFN α), hyperactivation of IFN- α , pSTAT1, reduced proteasome activities	Severe autoinflammatory phenotype (neonatal-onset fever, skin rash, myositis, severe pulmonary hypertension, basal ganglia calcification), periodic inflammatory exacerbation; immunodeficiency. Partial phenocopy of PRAAS	
IKBKG (NEMO exon 5 deletion (5 patients)	IKBKG	XL	<u>NA</u>	Leukocytes	Mutant NEMO lacked exon 5 (NEMO- Aex5), failed to bind TBK1; NEMO-Aex5 stabilized IKKi, increasing type 1 IFN production	fever, skin rash, systemic autoinflammation, infections, CNS involvement, panniculitis, uveitis, hepatosplenomegaly, ectodermal dysplasia	
TBK1 deficiency (4 patients)	TBK1	AR	<u>NA</u>	Leukocytes	Autoinflammation driven by TNF-induced RIPK1-dependent cell death	Chronic systemic autoinflammation (polyarthritis, vasculitis, rash); delayed neurocognitive development	

Total number of disorders in Table 7: 56. New inborn errors of immunity: 14 (AR GOF *TMEM173* [70], *LSM11*, *RNU7-1* [71], *CDC42* [72–78], *STAT2* [79, 80], *ATAD3A* [81], *C2orf69* [83, 84], *RIPK1* [85, 86], *NCKAP1L* [87–89], *SYK* [90], *HCK1* [91], *PSMB9* [95, 96], *IKBKG* NEMO-Δex5, AR *TBK1* [82])

IFN interferon, HSM hepatosplenomegaly, CSF cerebrospinal fluid, SLE systemic lupus erythematosus, TORCH toxoplasmosis, other, rubella, cytomegalovirus, and herpes infections, SNHL sensorineural hearing loss, AGS Aicardi-Goutières syndrome, BSN bilateral striatal necrosis, FCL familial chilblain lupus, ICC intracranial calcification, IFN interferon type I, pDCs plasmacytoid dendritic cells, SP spastic paraparesis, SMS Singleton-Merten syndrome, ss single-stranded DNA

*Variants in *PSMB4*, *PSMB9*, *PSMA3*, and *POMP* have been proposed to cause a similar CANDLE phenotype in compound heterozygous monogenic (*PSMB4*), digenic (*PSMB3/PSMB8*, *PSMB9/PSMB4*, *PSMB4/PSMB8*) and AD monogenic (*POMP*) models [115]



 Table 8 Complement deficiencies

Crase Control defects Control A R Control Control Record Control Record Control Record Control Record			Compleme	ent Deficie	encies	
C10 deficiency due to defects C10C AR 1909/5 C10C AR 1909/5 C10C AR 1909/5 AR 1909/5 AR 1909/5 AR 1909/5 AR 1909/5 AR 1909/5 C1r deficiency C1r deficiency C1r deficiency C1r deficiency C1r AR 1919/5 C1r deficiency C1r deficiency C1r AR 1919/5 C1r deficiency C1r deficiency	Disease					Associated features
Circ deficiency Circ deficiency Circ deficiency Circ deficiency Circ Periodontal Ehlers-Danlos Circ And And Circ deficiency Circ Periodontal Ehlers-Danlos Circ And Cir			AR	120550		
C1rd efficiency	C1a deficiency due to defects	C1QB	AR	120570		organisms
C1r deficiency C1R AR 013785 Absent C150 hemolytic actively operations of the complete C4 deficiency C1s deficiency C1s Periodontal Enter-Danlos C1s AR 013785 AR 0137	, , , , , , , , , , , , , , , , , , , ,	C1QC	AR	120575		
Ctr Periodontal Ehlers-Danios C1R AD GOF 01328 Normal CH80 Normal CH80 SELE infections, with triagilly Selection of the displayment						SLE, infections with encapsulated
C1s deficiency C1s Periodontal Ehlers-Danios C1s AD GOF S13255 C1s deficiency C1s Periodontal Ehlers-Danios C1s Periodontal Eh	C1r deficiency	C1R	AR	<u>613785</u>	defective activation of the	
Cris deficiency Cris deficiency Cris Periodontal Ehlors-Danios Cris AD GOF					classical pathway	I have a maintain and a fine fine silitar
C1s Periodontal Ehlers-Danios C7s AD GOF C1s Periodontal Ehlers-Danios C7s AD GOF C3s AD GOF C44+C4B AR L20810 C2 deficiency C2 deficiency C3 deficiency C3 deficiency C3 deficiency C3 deficiency C3 deficiency C44+C4B AR L20810 C3 deficiency C3 deficiency C3 deficiency C44+C4B AR L20810 C3 deficiency C3 deficiency C3 deficiency C3 deficiency C44+C4B AR L20810 C3 deficiency C3 deficiency C3 deficiency C3 deficiency C3 deficiency C3 deficiency C4 deficiency C5 deficiency C5 deficiency C5 deficiency C6 AR L20800 Demonstrate deficiency bundle activity, defective activation of the complement of the comple	C1r Periodontal Ehlers-Danlos	C1R	AD GOF	<u>613785</u>		
Cts Periodontal Ehlers-Danios Cts AD GOF G13286 Normal CH80 Inmobile activity of Complete C4 deficiency C44-C48 AR 120810 Absent CH60 hemolytic activity of C44-C48 Arc C48 Arc C4	C1s deficiency	C1S	AR	613785		
Complete C4 deficiency C44+C4B AR 120810 ABABRIC H50 hemolyte activity, defective activation of the classical pathway, complete deficiency requires beliable of the conference of the C4A and C4B and appears to have deficiency requires beliable of the characteristic beliable of the c4A and C4B and appears to he classical pathway, complete deficiency requires beliable of the c4A and C4B and appears to he classical pathway, complete deficiency requires beliable of the c4A and C4B and appears to he classical pathway, complete deficiency requires beliable of the c4A and C4B and appears to hemolyte activity, defective activation of the classical pathway and the case of the	o to demoisine,	0.0	/ " "	0.000		organisms, Emere Barnes priemetype
Complete C4 deficiency C3 AR 217000 C3 deficiency (CF) C3 AR C4 17000 Assert CH50 hemolytic activity, defective organisms, partial deficiency is counted and c18 control of the desical pathway. C5 deficiency (LOF) C3 AR C4 17000 C3 deficiency (LOF) C3 AR C4 17000 AR C5 deficiency (LOF) C5 deficiency (LOF) C6 deficiency C7 AR C7 AR C7 AR C7 AR C7 AR C7 AR C8 4 17000 C8 deficiency C6 deficiency C7 AR C8 4 17000 C8 deficiency C8 AR C9 4 17000 C8 deficiency C8 AR C9 4 17000 Assert CH50 and AH50 Assert C	C1s Periodontal Ehlers-Danlos	C1S	AD GOF	<u>613785</u>	Normal CH50	Hyperpigmentation, skin fragility
Complete C4 deficiency C2 AR 217000 ABant C160 hemolytic activity C3 deficiency (LOF) C3 AR 120700 Absent C160 and AP60 C3 AD G0F C3 AR 120700 C4 deficiency C5 deficiency C6 deficiency C7 deficiency C8 AR C9 17000 C6 deficiency C8 AR C9 17000 C6 deficiency C8 AR C9 17000 C6 deficiency C8 AR C9 17000 C7 deficiency C8 AR C9 17000 C8 deficiency C9 AR C9 17000 Absent C160 and AP60 C9 deficiency C8 AR C9 17000 Absent C160 and AP60 C9 deficiency C8 AR C9 17000 Absent C160 and AP60 C9 deficiency C8 AR C9 17000 Absent C160 and AP60 C9 deficiency C8 AR C9 17000 Absent C160 and AP60 C9 deficiency C8 AR C9 17000 Absent C160 and AP60 C9 deficiency C8 AR C9 17000 Absent C160 and AP60 C9 deficiency C8 AR C9 17000 Absent C160 and AP60 C9 deficiency C8 AR C9 17000 Absent C160 and AP60 C9 deficiency C8 AR C9 17000 Absent C160 and AP60 C9 deficiency C9 AR C9 AR C9 deficiency ABSP2 deficiency ABSP2 deficiency Ficolin 3 deficiency Ficolin 3 deficiency C1 inhibitor deficiency C6 Deficiency C7 BAR C9 12024 Absent C160 and AP60 C9 deficiency C9 AR C9 deficiency C9 AR C9 deficiency C1 inhibitor deficiency C6 AR C7 deficiency C7 BAR C9 deficiency C8 AR C9 deficiency C9 AR C9 deficiency C1 inhibitor deficiency C6 BAR C9 deficiency C7 BAR C9 deficiency C7 BAR C9 deficiency C1 inhibitor deficiency C7 BAR C9 deficiency C7 BAR C9 deficiency C7 BAR C9 deficiency C1 inhibitor deficiency C7 BAR C9 deficiency C7 BAR C9 deficiency C7 BAR C9 deficiency C1 inhibitor deficiency C7 BAR C9 deficiency C7 BAR C9 deficiency C9 AR C1 inhibitor deficiency C7 BAR C9 deficiency C9 AR C9						
Complete L4 deficiency Complete L4 deficie						
C2 deficiency	Complete C4 deficiency	C4A+C4B	AR	<u>120810</u>		
C2 deficiency C3 AR 12000 C3 AR C3 AR C3 AR C3 AR C3 AR C4 AR C5 AR C5 AR C5 AR C5 AR C5 AR C6 AR C7 AR C7 AR C7 AR C8 AR C9 AR C					mutations/deletions/conversions	
C2 deficiency (LOF) C3 AR 120700 C3 AR 120700 C3 AR 120700 C3 AP 20700 C3 AP 20700 C3 AP 20700 C5 deficiency (LOF) C5 deficiency C5 deficiency C6 AR 120800 C5 deficiency C7 AR 217650 C6 deficiency C7 AR 217650 C7 deficiency C8A AR 120800 C8 deficiency C8B AR 120800 C9 deficiency C9B AR 120800 C1 inhibitor deficiency C9B AR 13350 Factor B deficiency CPB AR 13450 Factor B defi						
C3 deficiency (LOF) C3 AR 120700 ABROFICE CHSG and AH50 Infections, glomerulonephrilis, atypical hemolytic activity, defactive production, directive humans in fractions, glomerulonephrilis, atypical hemolytic activity. defactive humans in fractions, glomerulonephrilis, atypical hemolytic activity. defactive humans in fractions, glomerulonephrilis, atypical hemolytic-uremic syndrome with GOF mutations. C5 deficiency C5 AR 120900 Absent CH5G and AH50 Abrest CH5G and AH	C2 deficiency	C2	A D	217000		
ABSENC LH50 and AH50 Intercelos y glomerulone-phritis, atyrized herrollytic activity, defective opportunation, defective humoral minute resignates Intercelos y complement particular	C2 deliciency	02	AIX	217000		organisms, ameroscierosis
C3 OF C3 AD GOF 120700 C3 AD GOF 120700 C3 AD GOF 120700 C5 deficiency C5 AR 120900 C6 deficiency C6 AR 120900 C7 deficiency C7 AR 120900 C7 deficiency C8A AR 120900 C8 deficiency C8B AR 120900 C8 deficiency C9 AR 120900 C8 de					Absent CH50 and AH50	Infections , glomerulonephritis, atypical
C3 GOF C3 AD GOF 120700 Immunity responses of the complement and the complement pathway with consumption of the contact system with encessed sportatory. When the lection of the contact system with encested sportatory infections, absoesses and efficiency C1 inhibitor deficiency C6 AR C12090 AR C1	C3 deficiency (LOF)	C3	AR	120700		
C3 GoF C5 deficiency C6 AR C7 deficiency C7 AR C8 deficiency C8 AR C8 deficiency C9 AR C9 AR C9 deficiency C1 inhibitor deficiency C2 inhibitor deficiency C2 inhibitor deficiency C2 inhibitor deficiency C3 AR C1 inhibitor deficiency C4 AR C5 inhibitor deficiency C6 AR C6 inhibitor deficiency C6 AR C7 inhibitor deficiency C7 inhi	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,					mutations.
C5 deficiency C6 AR 120900 C7 deficiency C7 deficiency C8 AR 120900 C8 gefficiency C9 AR 120900 C9 deficiency C9 AR 120900 C9 AR 1			45.005	100700		Atypical hemolytic-uremic syndrome
C5 deficiency C6 AR 210900 Defective bactericidal activity De	C3 GOF	C3	AD GOF	<u>120700</u>	complement	3,
C6 deficiency 6 6 AR 217050 C7 deficiency 7 C8A AR 120950 C8 deficiency 9 C8A AR 120950 C8 p deficiency 9 C8A AR 120950 C8 p deficiency 9 C8B AR 120950 C8 p deficiency 9 C9 AR 120950 C9 deficiency 9 C9 AR 120950 MASP2 AR 605102 C9 deficiency 9 C8B AR 120950 C9 deficiency 9 AR 120950 C9 AR 1	OF deficiency	05	4.5	400000		
C7 deficiency C7 AR 217070	C5 deficiency	C5	AR	120900		
C7 deficiency C8	C6 deficiency	C6	AR	217050	Defective Busicinsidal delivity	
C8α deficiency C8A AR 120950 120950 120950 ARSENT CHOL Bit A PROJECTION (and APSO) bedrictions of bemolytic activity, deficiently addressed infections. Dissertinated treasseral infections of bemolytic activity. Dissertinated treasseral infections of bemolytic activity. Mild susceptibility to disseminated neisseral infections. Dissertinated releasement and protein deficiency Mild susceptibility to disseminated neisseral infections. Dissertinated releasement and protein deficiency Mild susceptibility to disseminated neisseral infections. Mild susceptibility to disseminated neisseral infections. Dissertinated releasement and protein deficiency. ARSENCE of Complement activity, deficient bactericidal activity. Deficient activity, deficient neisseral infections. Mild susceptibility to disseminated neisseral infections. Proportion of the lectin activity, deficient neisseral infections. All services of complement activity, deficient neisseral infections. Proportion of the lectin activity, deficient neisseral infections. All services of complement activity, deficient neisseral infections. Proportion of the lectin activity. Absence of complement activity of the lectin activity of the propertion of the complement pathway with complement pathway with complement pathway with generation of the complement pathway with generation of the alternative activation of the alternative pathway. Applical hemolytic-uremic syndrome Applical hemolytic-uremic syndrome, propertion deficiency Applical hemolytic-uremic syndrome. Applical hemolytic-uremic syndrome. <t< td=""><td>-</td><td></td><td></td><td></td><td>-</td><td></td></t<>	-				-	
C8 y deficiency C8 deficiency C9 deficiency C9 deficiency C9 AR 120940 MASP2 deficiency MASP2 AR 605102 Ficolin 3 deficiency FCN3 AR C1 inhibitor deficiency C1 inhibitor deficiency C1 inhibitor deficiency C2 deficiency C3 AR C4 deficiency C5 AR C5 deficiency C6 deficiency C7 AR C6 deficiency C8 deficiency C9 AR C8 deficiency FCN3 AR C8 deficiency FCN4 AR C8 deficienc						Disseminated neisserial infections
C8 B deficiency C9 AR 120940 MASP2 deficiency MASP2 deficiency MASP2 deficiency MASP2 deficiency MASP2 deficiency Ficolin 3 deficiency Ficolin 3 deficiency Ficolin 3 deficiency C1 inhibitor deficiency C5 AR C60850 C1 inhibitor deficiency C6 AR C7 Inhibitor deficiency C7 Inhibitor deficiency C8 AR C8 B05102 C8 AR C8 B05102 C9 AR C8 B05102 C1 inhibitor deficiency C1 inhibitor deficiency C2 Inhibitor deficiency C3 AR C60850 C6 B06850 C6 C6 B06850 C6 C	-					
C9 deficiency MASP2 deficiency MASP2 AR 605102 Deficient activity, deficient bactericidal activity of bemotylic activity of the lectin activation of the lectin activation by the Ficolin 3 pallway. Spontaneous activation of the configuration of CAIC2, spontaneous activation of the configuration of CAIC2, spontaneous activation of the configuration of CAIC2, spontaneous activation of the configuration of the administration of the adm					- Basis Holdar assivity	
C9 deficiency C9	C8 B deficiency	C8B	AR	120960	Reduced CHEO and AREO	Mild augentibility to discominated
Ficolin 3 deficiency FCN3 AR S04973 AR AR S04973 AR S04973 AR S04973 AR AR AR S04973 AR AR AR S04973 AR AR AR S04973 AR AR AR AR AR AR AR AR AR A	C9 deficiency	C9	AR	120940	hemolytic activity, deficient	
Ficolin 3 deficiency SERPING1 AD S06860 Spontaneous activation by the Ficolin 3 pathway. Spontaneous activation of the complement pathway with consumption of C4/C2, spontaneous activation of the contact system with generation of bradykinin from high contact system with generation of the alternative activation of the alternative activation of the alternative activation of the alternative activation of the alternative complement pathway with consumption of C3	MASP2 deficiency	MASP2	AR	605102	Deficient activation of the lectin activation pathway	disease, autoimmunity
Pathway with consumption of CAIC constraints of the conficiency SERPING1 AD Sontaneous activation of the complement pathway with consumption of CAIC constraints of the confice system with generation of bradykinin from high molecular weight kininges of CFB AD GOF S12924 Sign-of-function mutation with increased spontaneous AH50 Applical hemolytic-uremic syndrome infections with encapsulated organisms at least to pathway with consumption of Large constraints and the confice of the confice system with generation of the alternative pathway Altypical hemolytic-uremic syndrome infections with encapsulated organisms at least to pathway with consumption of C3 AB confidence of CFB AR Sontaneous activation of the alternative pathway with consumption of C3 AB confidence of CFB AR Sontaneous activation of the alternative complement pathway with consumption of C3 Sontaneous activation of the alternative complement pathway with consumption of C3 Sontaneous activation of the alternative complement pathway with consumption of C3 Sontaneous activation of the alternative complement pathway with consumption of C3 Sontaneous activation of the alternative complement pathway with consumption of C3 Sontaneous activation of the alternative complement pathway with consumption of C3 Sontaneous activation of the alternative complement pathway with consumption of C3 Sontaneous activation of the alternative complement pathway with consumption of C3 Sontaneous activation of the alternative complement pathway with consumption of C3 Sontaneous activation of the pathway with consumption of C3 Sontaneous activation of the pathway with consumption of C3 Sontaneous activation of the pathway with consumption of C3 Sontaneous activation of the pathway with consumption of C3 Sontaneous activation of the pathway with consumption of C3 Sontaneous activation of the pathway with consumption of C3 Sontaneous activation of the pathway with consumption of C3 Sontaneous activation of the pathway w	u o . c .	50410		004070		Respiratory infections, abscesses
C1 inhibitor deficiency SERPING1 AD 606860 Spontaneous activation of the complement pathway with consumption of C4/C2, spontaneous activation of the contact system with generation of bradykinin from high molecular weight kininogen deiden activation of the alternative curvation of the alternative activation of the alternative complement pathway with consumption of C3 and Spontaneous Attavation of the alternative complement pathway with consumption of C3 and Spontaneous activation of the alternative complement pathway with consumption of C3 and Spontaneous Attavation of the alternative complement pathway with consumption of C3 and Spontaneous Attavation of the alternative complement pathway with consumption of C3 and Spontaneous Attavation of the alternative complement pathway with consumption of C3 and Spontaneous Attavation of the alternative pathway with consumption of C3 and Spontaneous Attavation o	Ficolin 3 deficiency	FCN3	AR	604973		
C1 inhibitor deficiency SERPING1 AD 606860 SERPING1 AD 606860 Consumption of CA/ÉZ, spontaneous activation of the contact system with generation of bradykinin from high molecular weight kininogen white molecular weight kininogen white molecular weight kininogen molecular weight kininom of the alternative pathway with consumption of C3 Spontaneous activati						Hereditary angioedema
C1 inhibitor deficiency SERPING1 AD 606860 Spontaneous activation of the contact system with generation of bradykinin from high molecular weight kininogen Gain-of-function mutation with increased spontaneous AH50 Factor B deficiency Factor B deficiency Factor D deficiency Factor I deficiency Factor I deficiency Factor I deficiency Factor H deficiency Fact						
CFB AD GOF Gain-of-function mutation with increased spontaneous AH50 Deficient activation of the alternative pathway	C1 inhibitor deficiency	SEDDING1	_ AD	606860		
Factor B GOF CFB AR 615561 Factor B deficiency CFB AR 615561 Factor D deficiency Factor I deficiency Factor I deficiency Factor H deficiency CFH AR CFHR1 CFHR2 CFHR3 CFHR4 CFHR5 CFHR5 AR or AD	C i illilibitor deliciency	SERFINGI	AD	<u>606860</u>		
Factor B deficiency Factor D deficiency Factor I deficiency Factor I deficiency Factor I deficiency Factor I deficiency Factor H -related protein deficiency Thrombomodulin deficiency Thrombomodulin deficiency Membrane Cofactor Protein (CD46) deficiency CFB AR AR AD GOF AR 615561 AR 615661 AR 61566					of bradykinin from high	
Factor B deficiency Factor D deficiency Factor D deficiency Factor I deficiency Factor I deficiency Factor I deficiency Factor H						
Factor B deficiency CFB AR 615561 Deficient activation of the alternative pathway Infections with encapsulated organisms alternative pathway Neisserial infections	Factor B GOF	CFB	AD GOF	612924		Atypical hemolytic-uremic syndrome
Factor D deficiency Factor I deficiency Factor I deficiency CFD AR 134350 Absent AH50 hemolytic activity Neisserial infections Neisserial infections Neisserial infections Neisserial infections Spontaneous activation of the alternative complement pathway with consumption of C3 Factor H deficiency CFH AR or AD CFHR1 CFHR2 Factor H -related protein deficiencies CFHR3 CFHR4 CFHR5 Thrombomodulin deficiency AR or AD Membrane Cofactor Protein (CD46) deficiency AR AR AR AR AR AR AR AR AR A	Factor B deficiency	CEB	A.D.	615564		Infections with encapsulated organisms
Properdin deficiency CFP XL 300383 Absent AH50 hemolytic activity Spontaneous activation of the alternative complement pathway with consumption of C3 Factor H deficiency CFH AR or AD CFHR1 CFHR2 CFHR3 CFHR4 CFHR4 CFHR4 CFHR5 Factor H -related protein deficiencies Factor H -related protein deficiency Thrombomodulin deficiency Thrombomodulin deficiency Membrane Cofactor Protein (CD46) deficiency Membrane Attack Complex Inhibitor (CD59) AR A A A A A A A A A A A A A A A A A A	ractor b deficiency					·
Factor I deficiency CFH AR or AD CFH AR or AD AR or AD CFHR1 CFHR2 CFHR4 CFHR4 CFHR4 CFHR5 CFHR5 CFHR5 CFHR5 CD46 AD AR or AD AR or AD Spontaneous activation of the alternative complement pathway with consumption of C3 Spontaneous activation of the alternative complement pathway with consumption of C3 Spontaneous activation of the alternative complement pathway with consumption of C3 Normal CH50, AH50, autoantibodies to Factor H., linked deletions of one or more CFHR genes leads to susceptibility autoantibody-mediated aHUS Thrombomodulin deficiency THBD AD 120920 Membrane Attack Complex Inhibitor (CD59) Membrane Attack Complex Inhibitor (CD59) AR AR or AD 217030 Spontaneous activation of the alternative complement alternative complements alternate pathway with consumption of C3 Spontaneous activation of the alternative complement alternative complement alternative pathway with consumption of C3 Spontaneous activation of the alternative complement alternative complement alternative pathway with consumption of C3 Spontaneous activation of the alternative complement alternative complement, pathway with consumption of C3 Normal CH50, AH50 Older onset atypical hemolytic-uremic syndrome, infections Spontaneous activation of the alternative complement alternative complement, pathway with consumption of C3 Normal CH50, AH50, AH50 Older onset atypical hemolytic-uremic syndrome, of CFHR genes leads to susceptibility autoantibody-mediated aHUS Normal CH50, AH50 Atypical hemolytic-uremic syndrome, infections, preeclampsia Atypical hemolytic-uremic syndrome, infections, preeclampsia	Factor D deficiency				, ,	
Factor I deficiency CFH AR or AD CFH AR or AD AR or AD CFHR1 CFHR2 CFHR4 CFHR4 CFHR5 CFHR5 CFHR5 AR or AD AR or	Properdin deficiency	CFP	XL	<u>300383</u>	• •	Neisserial infections
Factor H deficiency CFH AR or AD CFHR1 CFHR2 CFHR3 CFHR4 CFHR5 CFHR6 CFHR6 CFHR6 CFHR6 CFHR7 C						
Factor H deficiency CFH AR or AD AR or AD CFHR1 CFHR2 CFHR3 CFHR4 CFHR4 CFHR5 Thrombomodulin deficiency Thrombomodulin deficiency Membrane Cofactor Protein (CD46) deficiency CFH AR or AD CFHR1 CFHR3 CFHR4 CFHR5 CFHR4 CFHR5 AR or AD AD 134370 Spontaneous activation of the alternative complement pathway with consumption of C3 autoantibodies to Factor H., linked deletions of one or more consistent of the susceptibility autoantibody-mediated aHUS mediated aHUS Normal CH50, AH50, autoantibody-mediated aHUS susceptibility autoantibody-mediated aHUS Inhibitor of complement alternate pathway, decreased C3b binding Membrane Attack Complex Inhibitor (CD59) AB CFHR1 CFHR2 CFHR2 CFHR3 CFHR4 CFHR5 AR or AD AR or AD AR or AD AR or AD 134371, Normal CH50, AH50, autoantibodies to Factor H., linked deletions of one or more consistent of the syndrome, disseminated neisserial infections infections, atypical Hemolytic-uremic syndrome, dispension of the alternate pathway, decreased C3b binding Membrane Attack Complex Inhibitor (CD59) AB AR or AD BROWN AR or AD BROWN AR or AD BROWN AR or AD AR or AD BROWN AR OR	Factor I deficiency	CFI	AR	<u>217030</u>		Infections, disseminated neisserial
Factor H deficiency CFH AR or AD 134370 CFHR1 CFHR2 CFHR3 CFHR4 CFHR4 CFHR4 CFHR4 CFHR5 CFHR5 CFHR5 CFHR5 CFHR5 CFHR6 CFHR7 CFHR6 CFHR6 CFHR7 CFHR6 CFHR6 CFHR7 CFHR6 CFHR6 CFHR6 CFHR7 CFHR7 CFHR6 CFHR7 CFHR6 CFHR7 CFHR7 CFHR6 CFHR7 CFHR						
Factor H deficiency CFHR1 CFHR2 CFHR3 CFHR4 CFHR4 CFHR4 CFHR5 CFHR5 CFHR5 CFHR5 CFHR6 CFHR7 CFHR6 CFHR7 CFH					Spontaneous activation of the	
Factor H -related protein deficiencies CFHR1	Factor H deficiency	CFH	AR or AD	134370		
Factor H -related protein deficiencies Factor H -related protein deficiencies AR or AD AR or AD Thrombomodulin deficiency ThBD AD 134371, 600889, 605336, 605336, 605336, 608536, 608533, 608533, 608533, 608533, 608533, 608533, 608593 Thrombomodulin deficiency ThBD AD 188040 AD 120920 Membrane Cofactor Protein (CD46) deficiency Membrane Attack Complex Inhibitor (CD59) Membrane Attack Complex Inhibitor (CD59) AR AR or AD 134371, 600889, autoantibodies to Factor H., 60089, aut	_					
Factor H -related protein deficiencies CFHR3 CFHR4 CFHR5 CFHR7 Susceptibility autoantibody-mediated aHUS Normal CH50, AH50 Atypical hemolytic-uremic syndrome Atypical hemolytic-uremic syndrome, infections, preeclampsia C3b binding Membrane Attack Complex Inhibitor (CD59) Membrane Attack Complex Inhibitor (CD59) AR CFHR3 CFHR3 CFHR3 CFHR3 CFHR3 CFHR4 CFHR5 CFHR3 CFHR3 CFHR4 CFHR5 CFHR3 CFHR5 CFHR3 CFHR3 CFHR4 CFHR4 CFHR5 CFHR3 CFHR4 CFHR4 CFHR5 CFHR3 CFHR4 CFHR4 CFHR5 CFHR4 CFHR5 CFHR4 CFHR5 CFHR4 CFHR5 CFHR5 CFHR5 CFHR4 CFHR5 CFHR5 CFHR5 CFHR4 CFHR5 CFHR4 CFHR5 CFHR5 CFHR5 CFHR5 CFHR4 CFHR5 CFHR5 CFHR5 CFHR5 CFHR6 CFHR4 CFHR5 CFHR6 CFHR7		CFHR1		<u>134371</u> ,	Normal CH50, AH50,	
CFHR4]			
CFHR5 608593 susceptibility autoantibody-mediated aHUS	Factor H –related protein deficiencies		AR or AD			
Thrombomodulin deficiency THBD AD 188040 Normal CH50, AH50 Atypical hemolytic-uremic syndrome Inhibitor of complement alternate pathway, decreased C3b binding Membrane Attack Complex Inhibitor (CD59) AB AB Mediated aHUS Normal CH50, AH50 Atypical hemolytic-uremic syndrome, infections, preeclampsia C3b binding Hemolytic anemia, polyneuropathy			1			miocaons
Membrane Cofactor Protein (CD46) deficiency AD 120920 Membrane Attack Complex Inhibitor (CD59) Membrane Attack Complex Inhibitor (CD59) AB 120920 AB 120920 C3b binding Erythrocytes highly susceptible Hemolytic anemia, polyneuropathy		CFHR5		<u>608593</u>	mediated aHUS	
deficiency CD46 AD 120920 alternate pathway, decreased C3b binding Membrane Attack Complex Inhibitor (CD59) AB 107374 Erythrocytes highly susceptible Hemolytic anemia, polyneuropathy	Thrombomodulin deficiency	THBD	AD	<u>188040</u>	· · · · · · · · · · · · · · · · · · ·	1
deficiency AD 12/92/20 alternate pathway, decreased C3b binding C3b binding Membrane Attack Complex Inhibitor (CD59) AB 40/2734 Erythrocytes highly susceptible Hemolytic anemia, polyneuropathy	Membrane Cofactor Protein (CD46)	05.15		400		
Membrane Attack Complex Inhibitor (CD59) CD50 AB 407274 Erythrocytes highly susceptible Hemolytic anemia, polyneuropathy		CD46	AD	120920		intections, preeclampsia
	Membrane Attack Complex Inhibitor (CD59)	05		10777		Hemolytic anemia, polyneuropathy
	deficiency	CD59	AR		to complement-mediated lysis	.,, 20,,
CD55 deficiency (CHAPEL disease) CD55 AR 125240 Hyperactivation of complement on endothelium Protein losing enteropathy, thrombosis	CD55 deficiency (CHAPEL disease)	CD55	AR	125240		Protein losing enteropathy, thrombosis

Total number of mutant genes in Table 8: 36. New disorders: Nil *MAC* membrane attack complex, *SLE* systemic lupus erythematosus



Table 9 Bone marrow failure

				В	one Marrow	Failure			
Disease	Genetic defect	Inheritance	Gene OMIM	T cells	B cells	Other affected cells	Associated features	Major Category	Subcategory
Fanconi Anemia	FANCA	AR	227650						
Type A									
Fanconi Anemia	FANCB	XLR	300514	_					
Type B Fanconi Anemia	FANCC	AR	227645	-					
Type C									
Fanconi Anemia Type D1	BRCA2	AR	605724						
Fanconi Anemia Type D2	FANCD2	AR	227646						
Fanconi Anemia Type E	FANCE	AR	<u>600901</u>						
Fanconi Anemia Type F	FANCF	AR	603467						
Fanconi Anemia Type G	XRCC9	AR	614082				normal to low NK, CNS,	Bone marrow	
Fanconi Anemia Type I	FANCI	AR	609053	normal to low	normal to low	HSC	normal to low NK, CNS, skeletal, skin, cardiac, GI, urogenital anomalies, increased chromosomal breakage	failure with immune	Fanconi Anem
Fanconi Anemia Type J	BRIP1	AR	609054	1				deficiency	
Fanconi Anemia	FANCL	AR	614083	-					
Type L Fanconi Anemia	FANCM	AR	<u>618096</u>	-					
Type M Fanconi Anemia Type N	PALB2	AR	610832	-					
Fanconi Anemia	RAD51C	AR	613390	-					
Type O Fanconi Anemia	SLX4	AR	<u>613951</u>						
Type P Fanconi Anemia	ERCC4	AR	<u>615272</u>	-					
Type Q Fanconi Anemia	RAD51	AR	617244	_					
Type R									
Fanconi Anemia Type S	BRCA1	AR	<u>617883</u>						
Fanconi Anemia Type T	UBE2T	AR	<u>616435</u>						
Fanconi Anemia Type U	XRCC2	AR	617247						
Fanconi Anemia Type V	MAD2L2	AR	617243						
Fanconi Anemia Type W	RFWD3	AR	<u>617784</u>						
MIRAGE myelodysplasia, infection, restriction of growth, adrenal hypoplasia, genital phenotypes, enteropathy)	SAMD9	AD GOF	617053	Not reported	Not reported	HSC, myeloid cells	Intrauterine growth retardation, gonadal abnormalities, adrenal failure, MDS with chromosome 7 aberrations, predisposition to infections, enteropathy, absent spleen		
Ataxia Pancytopenia Syndrome	SAMD9L	AD GOF	611170	Normal	low	HSC, myeloid cells	MDS, neurological features		
DKCX1	DKC1	XL	305000				Bone marrow failure, pulmonary and hepatic fibrosis,	1	
DKCA1	TERC	AD	<u>127550</u>	1			nail dystrophy, leukoplakia, reticulate skin pigmentation;		
DKCA2	TERT	AD	<u>187270</u>				microcephaly, neurodevelopmental delay		
DKCA3	TINF2	AD	604319						
DKCA4	RTEL1	AD	616373						
DKCA5	TINF2	AD	<u>268130</u>						
DKCA6	ACD	AD	616553	Normal	Normal to !	1100			Duolearet
DKCB1	NOLA3	AR	224230	Normal to low	Normal to low	HSC			Dyskeratosis Congenita
DKCB2	NOLA2	AR	<u>613987</u>	1					
DKCB3	WRAP53	AR	613988	1					



 Table 9 (continued)

DKCB4	TERT	AR	<u>613989</u>			
DKCB5	RTEL1	AR	615190		low	nail dystrophy, leukoplakia, bone marrow failure, severe B- cell immunodeficiency, intrauterine growth retardation, growth retardation, microcephaly, cerebellar hypoplasia, and esophageal dysfunction
DKCB6	PARN	AR	616353		Normal to low	developmental delay, microcephaly, and cerebellar hypoplasia
DKCB7	ACD	AR	616553		Normal to low	Bone marrow failure, pulmonary and hepatic fibrosis, nail dystrophy, leukoplakia, reticulate skin pigmentation; microcephaly, neurodevelopmental delay
BMFS1 (SRP72- deficiency)	SRP72	AD	602122	NA	NA	Bone marrow failure and congenital nerve deafness
BMFS2	ERCC6L2	AR	<u>615667</u>	NA	NA	Bone marrow failure, learning difficulties, microcephaly
BMFS5	TP53	AD	<u>618165</u>	NA	low B	Erythroid hypoplasia, B-cell deficiency
Coats plus syndrome	STN1	AR	613129	Normal	Normal	Intrauterine growth retardation, premature aging, pancytopenia, hypocellular
	CTC1	AR	617053	Not reported	Not reported	bone marrow, gastrointestinal hemorrhage due to vascular ectasia, intracranial calcification, abnormal telomeres
MECOM deficiency	MECOM	AD	<u>616738</u>	Not reported	B cell deficiency	Bone marrow failure, thrombocytopenia/pancytopeni a, radioulnar synostosis, clinodactyly, cardiac and renal malformations

Total number of mutant genes in Table 9: 44. New Inborn errors of immunity: 1 (MECOM1) [98, 99])

HSC hematopoietic stem cell, NK natural killer, CNS central nervous system, GI gastrointestinal, MDS myelodysplastic syndrome, DKCX X-inked dyskeratosis congenital, DKCA autosomal dominant dyskeratosis congenita, DKCB autosomal recessive dyskeratosis congenita, BMFS bone marrow failure syndrome



 Table 10
 Phenocopies of inborn errors of immunity

1. Phenocopies of Inborn Errors of Immunity											
Disease	Genetic defect/presumed pathogenesis	Circulating T cells	Circulating B cells	Serum Ig	Associated features/similar PID						
Associated with somatic mutations											
Autoimmune lymphoproliferative syndrome (ALPS–SFAS)	Somatic mutation in TNFRSF6	Increased CD4·CD8- double negative (DN) $\alpha\beta$ T cells	Normal, but increased number of CD5+ B cells	Normal or increased	Splenomegaly, lymphadenopathy, autoimmune cytopenias, Defective lymphocyte apoptosis/ALPS-FAS (=ALPS type lm)						
RAS-associated autoimmune leukoproliferative disease (RALD)	Somatic mutation in KRAS (GOF)	Normal	B cell lymphocytosis	Normal or increased	Splenomegaly, lymphadenopathy, autoimmune cytopenias, granulocytosis, monocytosis/ALPS-like						
RAS-associated autoimmune leukoproliferative disease (RALD)	Somatic mutation in NRAS (GOF)	Increased CD4-CD8-double negative (DN) T alpha/beta cells	Lymphocytosis	Normal or increased	Splenomegaly, lymphadenopathy, autoantibodies/ALPS-like						
Cryopyrinopathy, (Muckle-Wells /CINCA/NOMID-like syndrome)	Somatic mutation in NLRP3	Normal	Normal	Normal	Urticaria-like rash, arthropathy, neurological signs						
Hypereosinophilic syndrome due to somatic mutations in STAT5b	Somatic mutation in STAT5B (GOF)	Normal	Normal	Normal	Eosinophilia, atopic dermatitis, urticarial rash, diarrhea						
VEXAS (vacuoles, E1 enzyme, X-linked, autoinflammatory, somatic) syndrome	Somatic mutation in <i>UBA1</i> (XL)		Reduced		Late onset treatment- refractory inflammatory syndrome (fevers, cytopenias, dysplastic bone marrow, interstitial nephritis, chondritis, vasculitis).						
TLR8 GOF (5 patients)	Somatic mutation in <i>TLR8</i>	↑(mild) CD4+, CD8+ T cells, effector/memory subsets; ↓NK cells	Normal B cells/subsets, ↓ pDCs	Normal/I ow igG, ↑lgM/lgA	Severe cytopenias,, hepatosplenomegaly, lymphadenopathy; recurrent infections; hypocellular bone marrow, elevated proinflammatory serum cytokines						
Associated with autoantibodies					-						
Chronic mucocutaneous candidiasis	AutoAb to IL-17 and/or IL-22	Normal	Normal	Normal	Endocrinopathy, chronic mucocutaneous candidiasis/CMC						
Adult-onset immunodeficiency with susceptibility to mycobacteria	AutoAb to IFNγ	Decreased naive T cells	Normal	Normal	Mycobacterial, fungal, Salmonella VZV infections/MSMD, or CID						
Recurrent skin infection	AutoAb to IL-6	Normal	Normal	Normal	Staphylococcal infections/STAT3 deficiency						
Pulmonary alveolar proteinosis	AutoAb to GM-CSF	Normal	Normal	Normal	Pulmonary alveolar proteinosis, cryptococcal meningitis, disseminated nocardiosis/CSF2RA deficiency						
Acquired angioedema	AutoAb to CI inhibitor	Normal	Normal	Normal	Angioedema/C1 INH deficiency (hereditary angioedema)						
Atypical Hemolytic Uremic Syndrome	AutoAb to Complement Factor H	Normal	Normal	Normal	aHUS = Spontaneous activation of the alternative complement pathway						
Thymoma with hypogammaglobulinemia (Good syndrome)	AutoAb to various cytokines	Increased CD8+ T cells	No B cells	Decrease d	Invasive bacterial, viral or opportunistic infections, autoimmunity, PRCA, lichen planus, cytopenia, colitis, chronic diarrhea						
Severe COVID-19	AutoAb to type 1 IFNs (IFNα, IFNω)				Severe, life-threatening infection with SARS-CoV-2						

Total number of conditions for Table 10: 15 (7 due to somatic mutations; 8 due to autoAbs). New phenocopies: 3 (somatic variants in *UBA1* [97], *TLR8* [58]; autoAbs against type 1 IFNs [100–104])

aHUS atypical hemolytic uremic syndrome, XL X-linked inheritance, AR autosomal recessive inheritance, AD autosomal dominant inheritance, LOF loss-of-function, GOF gain-of-function, PRCA pure red cell aplasia



Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s10875-022-01289-3.

Author Contribution SGT wrote the drafts of the manuscript, prepared the tables, and revised the original manuscripts for resubmission. All co-authors contributed to and edited drafts of the original and revised manuscripts and tables and approved the final submitted version.

Funding The Inborn Errors of Immunity Expert Committee received funding from the International Union of Immunological Societies; CSL Behring, Baxalta, and Shire/Takeda provided educational grants to enable us to compile this interim update to novel causes of immune diseases. This work was also supported in part by the Intramural Research Program of the NIAID, NIH. SGT is supported by an Investigator Grant (Level 3) awarded by the National Health and Medical Research Council of Australia. IM is a senior clinical investigator of FWO Vlaanderen (EBD-D8974-FKM)

Data Availability Not applicable

Declarations

Ethics Approval This work is a summary of recently reported genetic variants that represent novel inborn errors of immunity. No human research studies were performed to produce this summary. Thus, no approvals by appropriate institutional review boards or human research ethics committees were required to undertake the preparation of this report.

Consent to Participate Not applicable.

Consent for Publication The authors consent to publish the content of this summary. However, as noted above, as this is a summary of recently-reported genetic variants that represent novel inborn errors of immunity, we did not require consent to publish from participants.

Conflict of Interest The authors declare no competing interests.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

- Zhang Q, Frange P, Blanche S, Casanova JL. Pathogenesis of infections in HIV-infected individuals: insights from primary immunodeficiencies. Curr Opin Immunol. 2017;48:122–33. https://doi.org/10.1016/j.coi.2017.09.002.
- Picard C, Bobby Gaspar H, Al-Herz W, Bousfiha A, Casanova JL, Chatila T, et al. International Union of Immunological Societies: 2017 Primary Immunodeficiency Diseases Committee report on inborn errors of immunity. J Clin Immunol. 2018;38(1):96–128. https://doi.org/10.1007/s10875-017-0464-9.

- Bousfiha A, Jeddane L, Picard C, Ailal F, Bobby Gaspar H, Al-Herz W, et al. The 2017 IUIS phenotypic classification for primary immunodeficiencies. J Clin Immunol. 2018;38(1):129–43. https://doi.org/10.1007/s10875-017-0465-8.
- Bousfiha A, Jeddane L, Picard C, Al-Herz W, Ailal F, Chatila T, et al. Human inborn errors of immunity: 2019 update of the IUIS Phenotypical Classification. J Clin Immunol. 2020;40(1):66–81. https://doi.org/10.1007/s10875-020-00758-x.
- Tangye SG, Al-Herz W, Bousfiha A, Chatila T, Cunningham-Rundles C, Etzioni A, et al. Human inborn errors of immunity: 2019 update on the classification from the International Union of Immunological Societies Expert Committee. J Clin Immunol. 2020;40(1):24–64. https://doi.org/10.1007/s10875-019-00737-x.
- Casanova JL, Abel L. Human genetics of infectious diseases: unique insights into immunological redundancy. Semin Immunol. 2018;36:1–12. https://doi.org/10.1016/j.smim.2017.12.008.
- Fischer A, Rausell A. What do primary immunodeficiencies tell us about the essentiality/redundancy of immune responses? Semin Immunol. 2018;36:13–6. https://doi.org/10.1016/j.smim. 2017.12.001.
- Picard C, Fischer A. Contribution of high-throughput DNA sequencing to the study of primary immunodeficiencies. Eur J Immunol. 2014;44(10):2854–61. https://doi.org/10.1002/ejj.201444669.
- Leiding JW, Forbes LR. Mechanism-based precision therapy for the treatment of primary immunodeficiency and primary immunodysregulatory diseases. J Allergy Clin Immunol Pract. 2019;7(3):761–73. https://doi.org/10.1016/j.jaip.2018.12.017.
- Ma CS, Tangye SG. Flow cytometric-based analysis of defects in lymphocyte differentiation and function due to inborn errors of immunity. Front Immunol. 2019;10:2108. https://doi.org/10. 3389/fimmu.2019.02108.
- Casanova JL, Conley ME, Seligman SJ, Abel L, Notarangelo LD. Guidelines for genetic studies in single patients: lessons from primary immunodeficiencies. J Exp Med. 2014;211(11):2137–49. https://doi.org/10.1084/jem.20140520.
- Lev A, Lee YN, Sun G, Hallumi E, Simon AJ, Zrihen KS, et al. Inherited SLP76 deficiency in humans causes severe combined immunodeficiency, neutrophil and platelet defects. J Exp Med. 2021;218(3). https://doi.org/10.1084/jem.20201062.
- 13. Yamazaki Y, Urrutia R, Franco LM, Giliani S, Zhang K, Alazami AM, et al. PAX1 is essential for development and function of the human thymus. Sci Immunol. 2020;5(44). https://doi.org/10.1126/sciimmunol.aax1036.
- Paganini I, Sestini R, Capone GL, Putignano AL, Contini E, Giotti I, et al. A novel PAX1 null homozygous mutation in autosomal recessive otofaciocervical syndrome associated with severe combined immunodeficiency. Clin Genet. 2017;92(6):664–8. https://doi.org/10.1111/cge.13085.
- Almutairi A, Wallace JG, Jaber F, Alosaimi MF, Jones J, Sallam MTH, et al. Severe combined immunodeficiency caused by inositol-trisphosphate 3-kinase B (ITPKB) deficiency. J Allergy Clin Immunol. 2020. https://doi.org/10.1016/j.jaci.2020.01.014.
- Delmonte OM, Bergerson JRE, Kawai T, Kuehn HS, McDermott DH, Cortese I, et al. SASH3 variants cause a novel form of X-linked combined immunodeficiency with immune dysregulation. Blood. 2021;138(12):1019–33. https://doi.org/10.1182/blood.2020008629.
- Labrador-Horrillo M, Franco-Jarava C, Garcia-Prat M, Parra-Martinez A, Antolin M, Salgado-Perandres S, et al. Case report: X-Linked SASH3 deficiency presenting as a common variable immunodeficiency. Front Immunol. 2022;13:881206. https://doi.org/10.3389/fimmu.2022.881206.
- Verheijen J, Wong SY, Rowe JH, Raymond K, Stoddard J, Delmonte OM, et al. Defining a new immune deficiency syndrome: MAN2B2-CDG. J Allergy Clin Immunol. 2020;145(3):1008–11. https://doi.org/10.1016/j.jaci.2019.11.016.



- Bainter W, Platt CD, Park SY, Stafstrom K, Wallace JG, Peters ZT, et al. Combined immunodeficiency due to a mutation in the gamma1 subunit of the coat protein I complex. J Clin Invest. 2021;131(3). https://doi.org/10.1172/JCI140494.
- Hetemaki I, Kaustio M, Kinnunen M, Heikkila N, Keskitalo S, Nowlan K, et al. Loss-of-function mutation in IKZF2 leads to immunodeficiency with dysregulated germinal center reactions and reduction of MAIT cells. Sci Immunol. 2021;6(65):eabe3454. https://doi.org/10.1126/sciimmunol.abe3454.
- Shahin T, Kuehn HS, Shoeb MR, Gawriyski L, Giuliani S, Repiscak P, et al. Germline biallelic mutation affecting the transcription factor Helios causes pleiotropic defects of immunity. Sci Immunol. 2021;6(65):eabe3981. https://doi.org/10.1126/sciimmunol.abe3981.
- 22. Hadjadj J, Aladjidi N, Fernandes H, Leverger G, Magerus-Chatinet A, Mazerolles F, et al. Pediatric Evans syndrome is associated with a high frequency of potentially damaging variants in immune genes. Blood. 2019;134(1):9–21. https://doi.org/10.1182/blood-2018-11-887141.
- Shahin T, Mayr D, Shoeb MR, Kuehn HS, Hoeger B, Giuliani S, et al. Identification of germline monoallelic mutations in IKZF2 in patients with immune dysregulation. Blood Adv. 2021. https:// doi.org/10.1182/bloodadvances.2021006367.
- Bainter W, Lougaris V, Wallace JG, Badran Y, Hoyos-Bachiloglu R, Peters Z, et al. Combined immunodeficiency with autoimmunity caused by a homozygous missense mutation in inhibitor of nuclear factor B kinase alpha (IKKalpha). Sci Immunol. 2021;6(63):eabf6723. https://doi.org/10.1126/sciimmunol.abf6723.
- Yamashita M, Kuehn HS, Okuyama K, Okada S, Inoue Y, Mitsuiki N, et al. A variant in human AIOLOS impairs adaptive immunity by interfering with IKAROS. Nat Immunol. 2021;22(7):893–903. https://doi.org/10.1038/s41590-021-00951-z.
- Kuehn HS, Chang J, Yamashita M, Niemela JE, Zou C, Okuyama K, et al. T and B cell abnormalities, pneumocystis pneumonia, and chronic lymphocytic leukemia associated with an AIOLOS defect in patients. J Exp Med. 2021;218(12). https://doi.org/10.1084/jem.20211118.
- 27. Wu B, Rice L, Shrimpton J, Lawless D, Walker K, Carter C, et al. Biallelic mutations in calcium release activated channel regulator 2A (CRACR2A) cause a primary immunodeficiency disorder. Elife. 2021;10. https://doi.org/10.7554/eLife.72559.
- Beziat V, Rapaport F, Hu J, Titeux M, Bonnet des Claustres M, Bourgey M et al. Humans with inherited T cell CD28 deficiency are susceptible to skin papillomaviruses but are otherwise healthy. Cell. 2021;184(14):3812-28 e30. doi:https://doi.org/10. 1016/j.cell.2021.06.004.
- Mace EM, Paust S, Conte MI, Baxley RM, Schmit MM, Patil SL, et al. Human NK cell deficiency as a result of biallelic mutations in MCM10. J Clin Invest. 2020. https://doi.org/10.1172/JCI134966.
- Baxley RM, Leung W, Schmit MM, Matson JP, Yin L, Oram MK, et al. Bi-allelic MCM10 variants associated with immune dysfunction and cardiomyopathy cause telomere shortening. Nat Commun. 2021;12(1):1626. https://doi.org/10.1038/s41467-021-21878-x.
- Beziat V, Tavernier SJ, Chen YH, Ma CS, Materna M, Laurence A, et al. Dominant-negative mutations in human IL6ST underlie hyper-IgE syndrome. J Exp Med. 2020;217(6). https://doi.org/ 10.1084/jem.20191804.
- Monies D, Abouelhoda M, Assoum M, Moghrabi N, Rafiullah R, Almontashiri N, et al. Lessons learned from large-scale, first-tier clinical exome sequencing in a highly consanguineous population. Am J Hum Genet. 2019;104(6):1182–201. https://doi.org/ 10.1016/j.ajhg.2019.04.011.
- 33. Chen YH, Grigelioniene G, Newton PT, Gullander J, Elfving M, Hammarsjo A, et al. Absence of GP130 cytokine receptor

- signaling causes extended Stuve-Wiedemann syndrome. J Exp Med. 2020;217(3). https://doi.org/10.1084/jem.20191306.
- Kaustio M, Nayebzadeh N, Hinttala R, Tapiainen T, Astrom P, Mamia K, et al. Loss of DIAPH1 causes SCBMS, combined immunodeficiency, and mitochondrial dysfunction. J Allergy Clin Immunol. 2021;148(2):599–611. https://doi.org/10.1016/j. jaci.2020.12.656.
- Niehues T, Ozgur TT, Bickes M, Waldmann R, Schoning J, Brasen J, et al. Mutations of the gene FNIP1 associated with a syndromic autosomal recessive immunodeficiency with cardiomyopathy and pre-excitation syndrome. Eur J Immunol. 2020;50(7):1078–80. https://doi.org/10.1002/eji.201948504.
- Saettini F, Poli C, Vengoechea J, Bonanomi S, Orellana JC, Fazio G, et al. Absent B cells, agammaglobulinemia, and hypertrophic cardiomyopathy in folliculin interacting protein 1 deficiency. Blood. 2020. https://doi.org/10.1182/blood.2020006441.
- Le Coz C, Nguyen DN, Su C, Nolan BE, Albrecht AV, Xhani S, et al. Constrained chromatin accessibility in PU.1-mutated agammaglobulinemia patients. J Exp Med. 2021;218(7). https://doi.org/10.1084/jem.20201750.
- Takeda AJ, Maher TJ, Zhang Y, Lanahan SM, Bucklin ML, Compton SR, et al. Human PI3Kgamma deficiency and its microbiota-dependent mouse model reveal immunodeficiency and tissue immunopathology. Nat Commun. 2019;10(1):4364. https://doi.org/10.1038/s41467-019-12311-5.
- Thian M, Hoeger B, Kamnev A, Poyer F, Kostel Bal S, Caldera M, et al. Germline biallelic PIK3CG mutations in a multifaceted immunodeficiency with immune dysregulation. Haematologica. 2020. https://doi.org/10.3324/haematol.2019.231399.
- Kury P, Staniek J, Wegehaupt O, Janowska I, Eckenweiler M, Korinthenberg R, et al. Agammaglobulinemia with normal B-cell numbers in a patient lacking Bob1. J Allergy Clin Immunol. 2021;147(5):1977–80. https://doi.org/10.1016/j.jaci.2021.01. 027.
- 41. Kuhny M, Forbes LR, Cakan E, Vega-Loza A, Kostiuk V, Dinesh RK, et al. Disease-associated CTNNBL1 mutation impairs somatic hypermutation by decreasing nuclear AID. J Clin Invest. 2020. https://doi.org/10.1172/JCI131297.
- Yeh TW, Okano T, Naruto T, Yamashita M, Okamura M, Tanita K, et al. APRIL-dependent life-long plasmacyte maintenance and immunoglobulin production in humans. J Allergy Clin Immunol. 2020. https://doi.org/10.1016/j.jaci.2020.03.025.
- Kalinichenko A, Perinetti Casoni G, Dupre L, Trotta L, Huemer J, Galgano D, et al. RhoG deficiency abrogates cytotoxicity of human lymphocytes and causes hemophagocytic lymphohistiocytosis. Blood. 2021;137(15):2033–45. https://doi.org/10.1182/ blood.2020008738.
- Lee PY, Platt CD, Weeks S, Grace RF, Maher G, Gauthier K, et al. Immune dysregulation and multisystem inflammatory syndrome in children (MIS-C) in individuals with haploinsufficiency of SOCS1. J Allergy Clin Immunol. 2020. https://doi. org/10.1016/j.jaci.2020.07.033.
- Thaventhiran JED, Lango Allen H, Burren OS, Rae W, Greene D, Staples E, et al. Whole-genome sequencing of a sporadic primary immunodeficiency cohort. Nature. 2020;583(7814):90–5. https:// doi.org/10.1038/s41586-020-2265-1.
- Hadjadj J, Castro CN, Tusseau M, Stolzenberg MC, Mazerolles F, Aladjidi N, et al. Early-onset autoimmunity associated with SOCS1 haploinsufficiency. Nat Commun. 2020;11(1):5341. https://doi.org/10.1038/s41467-020-18925-4.
- Ogishi M, Yang R, Aytekin C, Langlais D, Bourgey M, Khan T, et al. Inherited PD-1 deficiency underlies tuberculosis and autoimmunity in a child. Nat Med. 2021;27(9):1646–54. https://doi.org/10.1038/s41591-021-01388-5.
- Tyler PM, Bucklin ML, Zhao M, Maher TJ, Rice AJ, Ji W, et al. Human autoinflammatory disease reveals



- ELF4 as a transcriptional regulator of inflammation. Nat Immunol. 2021;22(9):1118–26. https://doi.org/10.1038/s41590-021-00984-4.
- Sun G, Qiu L, Yu L, An Y, Ding Y, Zhou L, et al. Loss of function mutation in ELF4 causes autoinflammatory and immunode-ficiency disease in human. J Clin Immunol. 2022. https://doi.org/10.1007/s10875-022-01243-3.
- Stremenova Spegarova J, Lawless D, Mohamad SMB, Engelhardt KR, Doody G, Shrimpton J, et al. Germline TET2 loss of function causes childhood immunodeficiency and lymphoma. Blood. 2020;136(9):1055–66. https://doi.org/10.1182/blood. 2020005844.
- Goos H, Fogarty CL, Sahu B, Plagnol V, Rajamaki K, Nurmi K, et al. Gain-of-function CEBPE mutation causes noncanonical autoinflammatory inflammasomopathy. J Allergy Clin Immunol. 2019;144(5):1364–76. https://doi.org/10.1016/j.jaci.2019. 06.003.
- Hoshino A, Boutboul D, Zhang Y, Kuehn HS, Hadjadj J, Ozdemir N, et al. Gain-of-function IKZF1 variants in humans cause immune dysregulation associated with abnormal T/B cell late differentiation. Sci Immunol. 2022;7(69):eabi7160. https:// doi.org/10.1126/sciimmunol.abi7160.
- Marin-Esteban V, Youn J, Beaupain B, Jaracz-Ros A, Barlogis V, Fenneteau O, et al. Biallelic CXCR2 loss-of-function mutations define a distinct congenital neutropenia entity. Haematologica. 2021. https://doi.org/10.3324/haematol.2021.279254.
- 54. Auer PL, Teumer A, Schick U, O'Shaughnessy A, Lo KS, Chami N, et al. Rare and low-frequency coding variants in CXCR2 and other genes are associated with hematological traits. Nat Genet. 2014;46(6):629–34. https://doi.org/10.1038/ng.2962.
- Yang R, Mele F, Worley L, Langlais D, Rosain J, Benhsaien I, et al. Human T-bet governs innate and innate-like adaptive IFNgamma immunity against mycobacteria. Cell. 2020;183(7):1826– 47 e31. https://doi.org/10.1016/j.cell.2020.10.046.
- Yang R, Weisshaar M, Mele F, Benhsaien I, Dorgham K, Han J, et al. High Th2 cytokine levels and upper airway inflammation in human inherited T-bet deficiency. J Exp Med. 2021;218(8). https://doi.org/10.1084/jem.20202726.
- Kerner G, Rosain J, Guerin A, AlKhabaz A, Oleaga-Quintas C, Rapaport F, et al. Inherited human IFNgamma deficiency underlies mycobacterial disease. J Clin Invest. 2020. https://doi.org/10. 1172/JC1135460.
- Aluri J, Bach A, Kaviany S, Chiquetto Paracatu L, Kitcharoensakkul M, Walkiewicz MA, et al. Immunodeficiency and bone marrow failure with mosaic and germline TLR8 gain of function. Blood. 2021;137(18):2450–62. https://doi.org/10.1182/blood. 2020009620.
- Fejtkova M, Sukova M, Hlozkova K, Skvarova Kramarzova K, Rackova M, Jakubec D, et al. TLR8/TLR7 dysregulation due to a novel TLR8 mutation causes severe autoimmune hemolytic anemia and autoinflammation in identical twins. Am J Hematol. 2022;97(3):338–51. https://doi.org/10.1002/ajh.26452.
- Drutman SB, Mansouri D, Mahdaviani SA, Neehus AL, Hum D, Bryk R, et al. Fatal cytomegalovirus infection in an adult with inherited NOS2 deficiency. N Engl J Med. 2020;382(5):437–45. https://doi.org/10.1056/NEJMoa1910640.
- Lafaille FG, Harschnitz O, Lee YS, Zhang P, Hasek ML, Kerner G, et al. Human SNORA31 variations impair cortical neuronintrinsic immunity to HSV-1 and underlie herpes simplex encephalitis. Nat Med. 2019;25(12):1873–84. https://doi.org/ 10.1038/s41591-019-0672-3.
- 62. Hait AS, Olagnier D, Sancho-Shimizu V, Skipper KA, Helleberg M, Larsen SM, et al. Defects in LC3B2 and ATG4A underlie HSV2 meningitis and reveal a critical role for autophagy in antiviral defense in humans. Sci Immunol. 2020;5(54). https://doi.org/10.1126/sciimmunol.abc2691.

- Vavassori S, Chou J, Faletti LE, Haunerdinger V, Opitz L, Joset P, et al. Multisystem inflammation and susceptibility to viral infections in human ZNFX1 deficiency. J Allergy Clin Immunol. 2021;148(2):381–93. https://doi.org/10.1016/j.jaci.2021.03.045.
- 64. Le Voyer T, Neehus AL, Yang R, Ogishi M, Rosain J, Alroqi F, et al. Inherited deficiency of stress granule ZNFX1 in patients with monocytosis and mycobacterial disease. Proc Natl Acad Sci U S A. 2021;118(15). https://doi.org/10.1073/pnas.2102804118.
- Alawbathani S, Westenberger A, Ordonez-Herrera N, Al-Hilali M, Al Hebby H, Alabbas F, et al. Biallelic ZNFX1 variants are associated with a spectrum of immuno-hematological abnormalities. Clin Genet. 2022;101(2):247–54. https://doi.org/10.1111/cge.14081.
- 66. Asano T, Boisson B, Onodi F, Matuozzo D, Moncada-Velez M, Maglorius Renkilaraj MRL, et al. X-linked recessive TLR7 deficiency in ~1% of men under 60 years old with life-threatening COVID-19. Sci Immunol. 2021;6(62). https://doi.org/10.1126/sciimmunol.abl4348.
- van der Made CI, Simons A, Schuurs-Hoeijmakers J, van den Heuvel G, Mantere T, Kersten S, et al. Presence of genetic variants among young men with severe COVID-19. JAMA. 2020;324(7):663–73. https://doi.org/10.1001/jama.2020.13719.
- Abolhassani H, Vosughimotlagh A, Asano T, Landegren N, Boisson B, Delavari S, et al. X-linked TLR7 deficiency underlies critical COVID-19 pneumonia in a male patient with ataxiatelangiectasia. J Clin Immunol. 2021. https://doi.org/10.1007/ s10875-021-01151-y.
- Li J, Ritelli M, Ma CS, Rao G, Habib T, Corvilain E, et al. Chronic mucocutaneous candidiasis and connective tissue disorder in humans with impaired JNK1-dependent responses to IL-17A/F and TGF-beta. Sci Immunol. 2019;4(41). https://doi.org/10.1126/ sciimmunol.aax7965.
- Lin B, Berard R, Al Rasheed A, Aladba B, Kranzusch PJ, Henderlight M, et al. A novel STING1 variant causes a recessive form of STING-associated vasculopathy with onset in infancy (SAVI). J Allergy Clin Immunol. 2020;146(5):1204–8 e6. https://doi.org/10.1016/j.jaci.2020.06.032.
- Uggenti C, Lepelley A, Depp M, Badrock AP, Rodero MP, El-Daher MT, et al. cGAS-mediated induction of type I interferon due to inborn errors of histone pre-mRNA processing. Nat Genet. 2020;52(12):1364–72. https://doi.org/10.1038/ s41588-020-00737-3.
- Verboon JM, Mahmut D, Kim AR, Nakamura M, Abdulhay NJ, Nandakumar SK, et al. Infantile myelofibrosis and myeloproliferation with CDC42 dysfunction. J Clin Immunol. 2020. https:// doi.org/10.1007/s10875-020-00778-7.
- Lam MT, Coppola S, Krumbach OHF, Prencipe G, Insalaco A, Cifaldi C, et al. A novel disorder involving dyshematopoiesis, inflammation, and HLH due to aberrant CDC42 function. J Exp Med. 2019;216(12):2778–99. https://doi.org/10.1084/jem.20190147.
- 74. Gernez Y, de Jesus AA, Alsaleem H, Macaubas C, Roy A, Lovell D, et al. Severe autoinflammation in 4 patients with C-terminal variants in cell division control protein 42 homolog (CDC42) successfully treated with IL-1beta inhibition. J Allergy Clin Immunol. 2019;144(4):1122–5 e6. https://doi.org/10.1016/j.jaci. 2019.06.017.
- Bucciol G, Pillay B, Casas-Martin J, Delafontaine S, Proesmans M, Lorent N, et al. Systemic inflammation and myelofibrosis in a patient with Takenouchi-Kosaki syndrome due to CDC42 Tyr64Cys mutation. J Clin Immunol. 2020. https://doi.org/10. 1007/s10875-020-00742-5.
- Bekhouche B, Tourville A, Ravichandran Y, Tacine R, Abrami L, Dussiot M, et al. A toxic palmitoylation of Cdc42 enhances NF-kappaB signaling and drives a severe autoinflammatory syndrome. J Allergy Clin Immunol. 2020. https://doi.org/10.1016/j.jaci.2020.03.020.



- He T, Huang Y, Ling J, Yang J. A new patient with NOCARH syndrome due to CDC42 defect. J Clin Immunol. 2020;40(4):571–5. https://doi.org/10.1007/s10875-020-00786-7.
- Szczawinska-Poplonyk A, Ploski R, Bernatowska E, Pac M. A novel CDC42 mutation in an 11-year old child manifesting as syndromic immunodeficiency, autoinflammation, hemophagocytic lymphohistiocytosis, and malignancy: a case report. Front Immunol. 2020;11:318. https://doi.org/10.3389/fimmu.2020.00318.
- Duncan CJA, Thompson BJ, Chen R, Rice GI, Gothe F, Young DF et al. Severe type I interferonopathy and unrestrained interferon signaling due to a homozygous germline mutation in STAT2. Sci Immunol 2019;4(42). https://doi.org/10.1126/sciim munol.aav7501.
- Gruber C, Martin-Fernandez M, Ailal F, Qiu X, Taft J, Altman J, et al. Homozygous STAT2 gain-of-function mutation by loss of USP18 activity in a patient with type I interferonopathy. J Exp Med. 2020;217(5). https://doi.org/10.1084/jem.20192319.
- Lepelley A, Della Mina E, Van Nieuwenhove E, Waumans L, Fraitag S, Rice GI, et al. Enhanced cGAS-STING-dependent interferon signaling associated with mutations in ATAD3A. J Exp Med. 2021;218(10). https://doi.org/10.1084/jem.20201560.
- Taft J, Markson M, Legarda D, Patel R, Chan M, Malle L, et al. Human TBK1 deficiency leads to autoinflammation driven by TNF-induced cell death. Cell. 2021;184(17):4447–63 e20. https://doi.org/10.1016/j.cell.2021.07.026.
- 83. Wong HH, Seet SH, Maier M, Gurel A, Traspas RM, Lee C, et al. Loss of C2orf69 defines a fatal autoinflammatory syndrome in humans and zebrafish that evokes a glycogen-storage-associated mitochondriopathy. Am J Hum Genet. 2021;108(7):1301–17. https://doi.org/10.1016/j.ajhg.2021.05.003.
- Lausberg E, Giesselmann S, Dewulf JP, Wiame E, Holz A, Salvarinova R, et al. C2orf69 mutations disrupt mitochondrial function and cause a multisystem human disorder with recurring autoinflammation. J Clin Invest. 2021;131(12). https://doi.org/10.1172/JCI143078.
- Tao P, Sun J, Wu Z, Wang S, Wang J, Li W, et al. A dominant autoinflammatory disease caused by non-cleavable variants of RIPK1. Nature. 2020;577(7788):109–14. https://doi.org/10.1038/ s41586-019-1830-y.
- Lalaoui N, Boyden SE, Oda H, Wood GM, Stone DL, Chau D, et al. Mutations that prevent caspase cleavage of RIPK1 cause autoinflammatory disease. Nature. 2020;577(7788):103–8. https://doi.org/10.1038/s41586-019-1828-5.
- 87. Cook SA, Comrie WA, Poli MC, Similuk M, Oler AJ, Faruqi AJ, et al. HEM1 deficiency disrupts mTORC2 and F-actin control in inherited immunodysregulatory disease. Science. 2020;369(6500):202–7. https://doi.org/10.1126/science.aay5663.
- 88. Salzer E, Zoghi S, Kiss MG, Kage F, Rashkova C, Stahnke S, et al. The cytoskeletal regulator HEM1 governs B cell development and prevents autoimmunity. Sci Immunol. 2020;5(49). https://doi.org/10.1126/sciimmunol.abc3979.
- Castro CN, Rosenzwajg M, Carapito R, Shahrooei M, Konantz M, Khan A, et al. NCKAP1L defects lead to a novel syndrome combining immunodeficiency, lymphoproliferation, and hyperinflammation. J Exp Med. 2020;217(12). https://doi.org/10.1084/jem.20192275.
- Wang L, Aschenbrenner D, Zeng Z, Cao X, Mayr D, Mehta M, et al. Gain-of-function variants in SYK cause immune dysregulation and systemic inflammation in humans and mice. Nat Genet. 2021;53(4):500–10. https://doi.org/10.1038/s41588-021-00803-4.
- Kanderova V, Svobodova T, Borna S, Fejtkova M, Martinu V, Paderova J, et al. Early-onset pulmonary and cutaneous vasculitis driven by constitutively active SRC-family kinase HCK. J Allergy Clin Immunol. 2021. https://doi.org/10.1016/j.jaci.2021. 07.046.

- 92. de Jesus AA, Hou Y, Brooks S, Malle L, Biancotto A, Huang Y, et al. Distinct interferon signatures and cytokine patterns define additional systemic autoinflammatory diseases. J Clin Invest. 2020;130(4):1669–82. https://doi.org/10.1172/JCI129301.
- Hegazy S, Marques MC, Canna SW, Goldbach-Mansky R, de Jesus AA, Reyes-Mugica M, et al. NEMO-NDAS: a panniculitis in the young representing an autoinflammatory disorder in disguise. Am J Dermatopathol. 2022. https://doi.org/10.1097/DAD. 00000000000002144.
- 94. Lee Y, Wessel AW, Xu J, Reinke JG, Lee E, Kim SM, et al. Genetically programmed alternative splicing of NEMO mediates an autoinflammatory disease phenotype. J Clin Invest. 2022;132(6). https://doi.org/10.1172/JCI128808.
- 95. Kataoka S, Kawashima N, Okuno Y, Muramatsu H, Miwata S, Narita K, et al. Successful treatment of a novel type I interferonopathy due to a de novo PSMB9 gene mutation with a Janus kinase inhibitor. J Allergy Clin Immunol. 2021;148(2):639–44. https://doi.org/10.1016/j.jaci.2021.03.010.
- Kanazawa N, Hemmi H, Kinjo N, Ohnishi H, Hamazaki J, Mishima H, et al. Heterozygous missense variant of the proteasome subunit beta-type 9 causes neonatal-onset autoinflammation and immunodeficiency. Nat Commun. 2021;12(1):6819. https:// doi.org/10.1038/s41467-021-27085-y.
- Beck DB, Ferrada MA, Sikora KA, Ombrello AK, Collins JC, Pei W, et al. Somatic mutations in UBA1 and severe adult-onset autoinflammatory disease. N Engl J Med. 2020. https://doi.org/ 10.1056/NEJMoa2026834.
- 98. Niihori T, Ouchi-Uchiyama M, Sasahara Y, Kaneko T, Hashii Y, Irie M, et al. Mutations in MECOM, encoding oncoprotein EVII, cause radioulnar synostosis with amegakaryocytic thrombocytopenia. Am J Hum Genet. 2015;97(6):848–54. https://doi.org/10.1016/j.ajhg.2015.10.010.
- Germeshausen M, Ancliff P, Estrada J, Metzler M, Ponstingl E, Rutschle H, et al. MECOM-associated syndrome: a heterogeneous inherited bone marrow failure syndrome with amegakaryocytic thrombocytopenia. Blood Adv. 2018;2(6):586–96. https:// doi.org/10.1182/bloodadvances.2018016501.
- Bastard P, Rosen LB, Zhang Q, Michailidis E, Hoffmann HH, Zhang Y, et al. Autoantibodies against type I IFNs in patients with lifethreatening COVID-19. Science. 2020;370(6515). https://doi.org/ 10.1126/science.abd4585.
- 101. Bastard P, Gervais A, Le Voyer T, Rosain J, Philippot Q, Manry J, et al. Autoantibodies neutralizing type I IFNs are present in ~4% of uninfected individuals over 70 years old and account for ~20% of COVID-19 deaths. Sci Immunol. 2021;6(62). https://doi.org/10.1126/sciimmunol.abl4340.
- 102. Abers MS, Rosen LB, Delmonte OM, Shaw E, Bastard P, Imberti L, et al. Neutralizing type-I interferon autoantibodies are associated with delayed viral clearance and intensive care unit admission in patients with COVID-19. Immunol Cell Biol. 2021;99(9):917–21. https://doi.org/10.1111/imcb.12495.
- 103. Troya J, Bastard P, Planas-Serra L, Ryan P, Ruiz M, de Carranza M, et al. Neutralizing autoantibodies to type I IFNs in >10% of patients with severe COVID-19 pneumonia hospitalized in Madrid, Spain. J Clin Immunol. 2021;41(5):914–22. https://doi.org/10.1007/s10875-021-01036-0.
- 104. Solanich X, Rigo-Bonnin R, Gumucio VD, Bastard P, Rosain J, Philippot Q, et al. Pre-existing autoantibodies neutralizing high concentrations of type I interferons in almost 10% of COVID-19 patients admitted to intensive care in Barcelona. J Clin Immunol. 2021;41(8):1733–44. https://doi.org/10.1007/s10875-021-01136-x.
- Koretzky GA, Abtahian F, Silverman MA. SLP76 and SLP65: complex regulation of signalling in lymphocytes and beyond. Nat Rev Immunol. 2006;6(1):67–78. https://doi.org/10.1038/nri1750.



- Bellelli R, Boulton SJ. Spotlight on the replisome: aetiology of dna replication-associated genetic diseases. Trends Genet. 2021;37(4):317–36. https://doi.org/10.1016/j.tig.2020.09.008.
- Chen YH, Spencer S, Laurence A, Thaventhiran JE, Uhlig HH. Inborn errors of IL-6 family cytokine responses. Curr Opin Immunol. 2021;72:135–45. https://doi.org/10.1016/j.coi.2021.04.007.
- Lacruz RS, Feske S. Diseases caused by mutations in ORAI1 and STIM1. Ann N Y Acad Sci. 2015;1356:45–79. https://doi.org/10. 1111/nyas.12938.
- Yamashita M, Morio T. Inborn errors of IKAROS and AIOLOS. Curr Opin Immunol. 2021;72:239

 –48. https://doi.org/10.1016/j.coi.2021.06.010.
- Duncan CJA, Hambleton S. Human disease phenotypes associated with loss and gain of function mutations in STAT2: viral susceptibility and type I interferonopathy. J Clin Immunol. 2021;41(7):1446–56. https://doi.org/10.1007/s10875-021-01118-z.
- Van Horebeek L, Dubois B, Goris A. Somatic variants: new kids on the block in human immunogenetics. Trends Genet. 2019;35(12):935–47. https://doi.org/10.1016/j.tig.2019.09.005.

- 112. Casanova JL, Holland SM, Notarangelo LD. Inborn errors of human JAKs and STATs. Immunity. 2012;36(4):515–28. https://doi.org/10.1016/j.immuni.2012.03.016.
- Zhang Q, Bastard P, Liu Z, Le Pen J, Moncada-Velez M, Chen J, et al. Inborn errors of type I IFN immunity in patients with life-threatening COVID-19. Science. 2020;370(6515). https://doi.org/10.1126/science.abd4570.
- Moens L, Meyts I. Recent human genetic errors of innate immunity leading to increased susceptibility to infection. Curr Opin Immunol. 2020;62:79–90. https://doi.org/10.1016/j.coi.2019.12.002.
- Brehm A, Liu Y, Sheikh A, Marrero B, Omoyinmi E, Zhou Q, et al. Additive loss-of-function proteasome subunit mutations in CANDLE/PRAAS patients promote type I IFN production. J Clin Invest. 2015;125(11):4196–211. https://doi.org/10.1172/JCI81260.

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Authors and Affiliations

Stuart G. Tangye^{1,2} · Waleed Al-Herz³ · Aziz Bousfiha⁴ · Charlotte Cunningham-Rundles⁵ · Jose Luis Franco⁶ · Steven M. Holland⁷ · Christoph Klein⁸ · Tomohiro Morio⁹ · Eric Oksenhendler¹⁰ · Capucine Picard^{11,12} · Anne Puel^{13,14} · Jennifer Puck¹⁵ · Mikko R. J. Seppänen¹⁶ · Raz Somech¹⁷ · Helen C. Su⁷ · Kathleen E. Sullivan¹⁸ · Troy R. Torgerson¹⁹ · Isabelle Meyts²⁰

- Garvan Institute of Medical Research, Darlinghurst, Sydney, NSW 2010, Australia
- St Vincent's Clinical School, Faculty of Medicine & Health, UNSW Sydney, Darlinghurst, NSW, Australia
- Department of Pediatrics, Faculty of Medicine, Kuwait University, Kuwait City, Kuwait
- ⁴ Laboratoire d'Immunologie Clinique, d'Inflammation et d'Allergy LICIA Clinical Immunology Unit, Casablanca Children's Hospital, Ibn Rochd Medical School, King Hassan II University, Casablanca, Morocco
- Departments of Medicine and Pediatrics, Mount Sinai School of Medicine, New York, NY, USA
- ⁶ Grupo de Inmunodeficiencias Primarias, Facultad de Medicina, Universidad de Antioquia UdeA, Medellin, Colombia
- ⁷ Laboratory of Clinical Immunology & Microbiology, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, MD, USA
- Dr von Hauner Children's Hospital, Ludwig-Maximilians-University Munich, Munich, Germany
- Department of Pediatrics and Developmental Biology, Tokyo Medical and Dental University, Tokyo, Japan
- Department of Clinical Immunology, Hôpital Saint-Louis, APHP, Université Paris Diderot, Sorbonne Paris Cité, Paris, France
- Study Center for Primary Immunodeficiencies, Necker Hospital for Sick Children, APHP, Paris, France

- Laboratory of Lymphocyte Activation and Susceptibility to EBV, INSERM UMR1163, Imagine Institute, Necker Hospital for Sick Children, Université Paris Cité, Paris, France
- Laboratory of Human Genetics of Infectious Diseases, INSERM U1163, Necker Hospital, 75015 Paris, France
- ¹⁴ Université Paris Cité, Imagine Institute, 75015 Paris, France
- Department of Pediatrics, University of California San Francisco and UCSF Benioff Children's Hospital, San Francisco, CA, USA
- Adult Immunodeficiency Unit, Infectious Diseases, Inflammation Center and Rare Diseases Center, Children's Hospital, University of Helsinki and Helsinki University Hospital, Helsinki, Finland
- Pediatric Department and Immunology Unit, Sheba Medical Center, Tel Aviv, Israel
- Division of Allergy Immunology, Department of Pediatrics, Children's Hospital of Philadelphia, University of Pennsylvania Perelman School of Medicine, Philadelphia, PA, USA
- Allen Institute for Immunology, Seattle, WA, USA
- Department of Immunology and Microbiology, Laboratory for Inborn Errors of Immunity, Department of Pediatrics, University Hospitals Leuven and KU Leuven, 3000 Leuven, Belgium

