

# Primary Immunodeficiency Diseases: an update on the Classification from the International Union of Immunological Societies Expert Committee for Primary Immunodeficiency

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Mary Ellen Conley, M.D. Department of Pediatrics Univ of Tennessee College of Medicine Department of Immunology MS 351 St. Jude Children's Research Hospital 262 Danny Thomas Place Memphis, TN 38105 Tel: (901)-595-2576 FAX: (901)-595-3977 Maryellen.conley@stjude.org Luigi D. Notarangelo, M.D. Division of Immunology and The Manton Center for Orphan Disease Research Children's Hospital Boston Karp Research Building, Room 20217 1 Blackfan Circle Boston, MA 02115 Tel: (617)-919-2277 FAX: (617)-730-0709 Iuigi.notarangelo@childrens.harvard.edu The International Union of Immunological Societies (IUIS) Expert Committee on Primary Immunodeficiency met in New York City, May 31-June 1, 2011 to update the classification of human primary immunodeficiencies (PIDs). Novel developments in gene discovery and increased knowledge in the mechanisms that govern immune system development and function have resulted in the identification of several novel PIDs in the last 2 years.

The classification of PIDs provides a framework to help in the diagnostic approach to patients. As in recent classifications, eight major groups of PIDs have been included in the Tables; however the order of the Tables has been changed with Table II now describing the "Well-defined syndromes with immunodeficiency" (previously Table III) to reflect the immunological similarity between the disorders included in this Table and those in Table I, "Combined immunodeficiencies".

Any classification of human disorders is somewhat arbitrary, and the classification of PIDs is no exception. Some disorders might well belong to more than one group. CD40 ligand deficiency, for example, is reported both in Table I and in Table III ("Predominantly antibody deficiencies"), to reflect the facts that failed B cell isotype switching was historically the most prominent feature of this condition (originally named Hyper-IgM syndrome) and that some patients survive into adulthood without significant opportunistic infections and do well with only immunoglobulin replacement therapy. Explanatory notes provided after each Table offer additional information (particularly where a condition appears in more than one Table) and indicate which new disorders have been added to that Table.

Although this updated classification reports on the most typical immunological findings and associated clinical and genetic features for the various PIDs, there is extensive clinical, immunological and molecular heterogeneity that can not be easily recapitulated in a brief summary. To facilitate a more rigorous analysis of each disease, a column has been added on the right to refer to its catalog number in the Online Mendelian Inheritance in Man (OMIM) publicly accessible database (www.omim.org) of human genetic disorders. It is suggested that the reader consult this regularly updated and fully referenced resource.

The prevalence of the various PIDs varies in different countries. For this reason, in this new classification, we have elected to avoid giving a comment on the relative frequency of PID disorders. However, an asterisk has been placed in the first column, after the disease name, to identify disorders for which fewer than 10 unrelated cases have been reported in the literature. Some of these forms of PID can be considered extremely rare. Others have only recently been identified and it may be that more patients will be detected over time.

Finally, it is increasingly recognized that different mutations in the same gene may result in different phenotypes and may be associated with different patterns of inheritance. This concept of clinical, immunological and genetic heterogeneity is assuming foremost importance. Notes in the text or in the footnotes identify such heterogeneity, when known.

The scope of the IUIS Expert Committee on Primary Immunodeficiency is to increase awareness, facilitate recognition and promote optimal treatment for patients with Primary Immunodeficiency disorders worldwide. For this reason, in addition to

periodically revising the Classification of Primary Immunodeficiencies, the Expert Committee is also actively involved in the development of diagnostic criteria and in providing, upon request, advice with regard to therapeutic guidelines.

Disease	Circulating T cells	Circulating B cells	Serum Ig	Associated Features	Inheritance	Genetic defect/ Presumed pathogenesis	OMIM number
1. TB <sup>+</sup> Severe Combined Immunodeficiency (SCID)							
(a) γc deficiency	Markedly decreased	Normal or increased	Decreased	Markedly decreased NK cells; leaky cases may present with low to normal T and/or NK cells or Omenn syndrome	XL	Defect in γ chain of receptors for IL-2, -4, -7, - 9, -15, -21	<u>300400</u>
(b) JAK3 deficiency	Markedly decreased	Normal or increased	Decreased	Markedly decreased NK cells; leaky cases may present with variable T and/or NK cells	AR	Defect in Janus activating kinase 3	<u>600173</u>
(c) IL7R $\alpha$ deficiency	Markedly decreased	Normal or increased	Decreased	Normal NK cells	AR	Defect in IL-7 receptor $\alpha$ chain	<u>146661</u>
(d) CD45 deficiency*	Markedly decreased	Normal	Decreased	Normal γ/δ T cells	AR	Defect in CD45	<u>151460</u>
(e) CD3δ*/CD3ε*/CD3ζ* deficiency	Markedly decreased	Normal	Decreased	Normal NK cells No γδT cells	AR	Defect in CD3δ, CD3ε or CD3ζ chains of T cell antigen receptor complex	<u>186790,</u> <u>186830,</u> <u>186740</u>
(f) Coronin-1A deficiency*	Markedly decreased	Normal	Decreased	Detectable thymus	AR	Defective thymic egress of T cells and defective T cell locomotion	<u>605000</u>
1. T'B'SCID							
(a) RAG 1/2 deficiency	Markedly decreased	Markedly decreased	Decreased	May present with Omenn syndrome, expanded γδT cells, autoimmunity and/or granulomas	AR	Defective VDJ recombination; defect of recombinase activating gene (RAG) 1 or 2	<u>601457</u>
(b) DCLRE1C (Artemis) deficiency	Markedly decreased	Markedly decreased	Decreased	Defective VDJ recombination, radiation sensitivity; may present with Omenn syndrome	AR	Defective VDJ recombination; defect in Artemis DNA recombinase-repair protein	<u>602450</u>
(c) DNA PKcs deficiency*	Markedly decreased	Markedly decreased	Decreased	[widely studied <i>scid</i> mouse defect]	AR	Defective VDJ recombination; defect in DNAPKcs Recombinase repair protein	<u>600899</u>

### Table I – Combined immunodeficiencies

(d) Adenosine deaminase (ADA) deficiency	Absent from birth (null mutations) or progressive decrease	Absent from birth of progressive decrease	Progressive decrease	Decreased NK cells, often with costochondral junction flaring, neurological features, hearing impairment, lung and liver manifestations; partial ADA deficiency may lead to delayed or milder presentation	AR	Absent ADA activity, elevated lymphotoxic metabolites (dATP, S- adenosylhomocysteine)	102700
(e) Reticular dysgenesis, AK2 deficiency	Markedly decreased	Decreased or normal	Decreased	Deficiency of T, B and NK cells with granulocytopenia, deafness	AR	Defective maturation of lymphoid and myeloid cells (stem cell defect) Defect in mitochondrial adenylate kinase 2.	<u>103020</u>
3. Omenn syndrome	Present; restricted heterogeneity	Normal or decreased	Decreased, except increased IgE	Erythroderma, eosinophilia, adenopathies, hepatosplenomegaly	AR	Hypomorphic mutations in RAG1/2, Artemis, IL- 7R $\alpha$ , RMRP, ADA, DNA Ligase IV, $\gamma$ c, or associated with DiGeorge syndrome; some cases have no defined gene mutation	<u>603554</u>
4. DNA ligase IV deficiency	Decreased	Decreased	Decreased	Microcephaly, facial dysmorphisms, radiation sensitivity; may present with Omenn syndrome or with a delayed clinical onset.	AR	DNA ligase IV defect, impaired nonhomologous end joining (NHEJ)	<u>601837</u>
5. Cernunnos/NHEJ1 deficiency*	Decreased	Decreased	Decreased	Microcephaly, in utero growth retardation, radiation sensitivity	AR	Cernunnos (NHEJ1) defect, impaired non- homologous end joining	<u>611291</u>
6. CD40 ligand deficiency	Normal; may progressively decrease	IgM <sup>+</sup> and IgD <sup>+</sup> B cells present, other isotypes absent	lgM increased or normal, other isotypes decreased	Neutropenia, thrombocytopenia; hemolytic anemia, biliary tract and liver disease, opportunistic infections	XL	Defects in CD40 ligand (CD40L) cause defective isotype switching and impaired dendritic cell signaling	<u>300386</u>
7. CD40 deficiency*	Normal	IgM <sup>*</sup> and IgD <sup>*</sup> B cells present, other isotypes absent	IgM increased or normal, other isotypes decreased	Neutropenia, gastrointestinal and liver/biliary tract disease, opportunistic infections	AR	Defects in CD40 cause defective isotype switching and impaired dendritic cell signaling	<u>109535</u>
8. Purine nucleoside phosphorylase (PNP) deficiency	Progressive decrease	Normal	Normal or decreased	Autoimmune haemolytic anemia, neurological impairment	AR	Absent PNP, T cell and neurologic defects from elevated toxic metabolites, especially dGTP	<u>164050</u>

9. CD3γ deficiency*	Normal, but reduced TCR expression	Normal	Normal		AR	Defect in CD3 γ	<u>186740</u>
10. CD8 deficiency*	Absent CD8, normal CD4 cells	Normal	Normal		AR	Defects of CD8 $\alpha$ chain	<u>186910</u>
11. ZAP-70 deficiency	Decreased CD8, normal CD4 cells	Normal	Normal		AR	Defects in ZAP-70 signaling kinase	<u>176947</u>
12. Ca <sup>++</sup> channel deficiency							
(a) ORAI-I deficiency*	Normal number, but defective TCR mediated activation	Normal	Normal	Autoimmunity, anhydrotic ectodermic dysplasia, non- progressive myopathy	AR	Defect in ORAI-1, a Ca <sup>++</sup> release-activated channel (CRAC) modulatory component	<u>610277</u>
(b) STIM-1 deficiency*	Normal number, but defective TCR mediated activation	Normal	Normal	Autoimmunity, anhydrotic ectodermic dysplasia, non- progressive myopathy	AR	Defect in STIM-1, a stromal interaction molecule Ca <sup>++</sup> sensor	<u>605921</u>
13. MHC class I deficiency	Decreased CD8, normal CD4	Normal	Normal	Vasculitis	AR	Mutations in <i>TAP1</i> , <i>TAP2</i> or <i>TAPBP</i> (tapasin) genes giving MHC class I deficiency	<u>604571</u>
14. MHC class II deficiency	Normal number, decreased CD4 cells	Normal	Normal or decreased	Failure to thrive, diarrhea, respiratory tract infections	AR	Mutation in transcription factors for MHC class II proteins ( <i>CIITA</i> , <i>RFX5</i> , <i>RFXAP</i> , <i>RFXANK</i> genes)	<u>209920</u>
15. Winged helix deficiency (Nude)*	Markedly decreased	Normal	Decreased	Alopecia, abnormal thymic epithelium, impaired T cell maturation [widely studied nude mouse defect]	AR	Defects in forkhead box N1 transcription factor encoded by <i>FOXN1</i> , the gene mutated in nude mice	<u>600838</u>
16. Complete DiGeorge syndrome	Profoundly decreased	Low to normal	Decreased	Lymphoproliferation (lymphadenopathy, hepatosplenomegaly), autoimmunity (may resemble IPEX syndrome), impaired T-cell proliferation	AD	Deletion of chromosome 22q11.2 or in a minority of cases other chromosomal regions, including 10p; heterozygous defects in transcription factor TBX1	<u>188400</u>
17. STAT5b deficiency*	Modestly decreased	Normal	Normal	Growth-hormone insensitive dwarfism, dysmorphic features, eczema, lymphocytic interstitial pneumonitis, autoimmunity	AR	Defects of STAT5b, impaired development and function of vot cells, Treg and NK cells, impaired T-cell proliferation	<u>604260</u>
18. ITK deficiency*	Modestly decreased	Normal	Normal or decreased		AR	Defects in ITK, EBV associated lymphoproliferation	<u>613011</u>
19. MAGT1 deficiency*	Decreased CD4 cells	Normal	Normal	EBV infection, lymphoma; viral infections, respiratory and GI infections	XL	Mutations in MAGT1, Impaired Mg <sup>++</sup> flux leading to impaired TCR signaling	<u>300715</u>

20. DOCK8 deficiency	Decreased	Decreased	Low IgM, increased IgE	Low NK cells, hypereosinophilia,	AR	Defect in DOCK8	<u>243700</u>
				recurrent infections; severe atopy, extensive cutaneous viral and bacterial (staph.) infections, susceptibility to			
				cancer			

XL, X-linked inheritance; AR, autosomal recessive inheritance; AD, autosomal dominant inheritance; SCID, Severe Combined Immune Deficiencies; EBV, Epstein Barr Virus; Ca<sup>++</sup>, calcium; MHC, Major Histocompatibility Complex

\*Ten or fewer unrelated cases reported in the literature

Notes: Two disorders have been added to Table I: DOCK8 deficiency and MAGT1 deficiency.

Infants with SCID who have maternal T cells engraftment may have T cells 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. Both of these disorders can be associated with higher numbers of T cells and reduced rather than absent activation responses when compared with typical SCID caused by null mutations. A spectrum of clinical findings including typical SCID, OS, leaky SCID, and granulomas with T lymphopenia can be found with *RAG* gene defects. RAC2 deficiency is a disorder of leukocyte motility and is reported in Table V; however, one patient with RAC2 deficiency was found to have absent T cell receptor excision circles (TRECs) by newborn screening, but T cell numbers and mitogen responses were not impaired. For additional syndromic conditions with T cell lymphopenia, such as DNA repair defects, cartilage hair hypoplasia and NEMO syndrome, see Table II and VI; however, it should be noted that individuals with the most severe manifestations of these disorders could have clinical signs and symptoms of SCID. Severe folate deficiency (such as with malabsorption due to defects in folate carrier or transporter genes *SLC10A1* or *PCFT*) and some metabolic disorders, such as methylmalonic aciduria, may present with reversible profound lymphopenia in addition to their characteristic presenting features.

Table II: Well-defined syndromes with immunodeficiency

Disease	Circulating T cells	Circulating B cells	Serum Ig	Associated features	Inheritance	Genetic defect/ Presumed pathogenesis	OMIM number
1. Wiskott-Aldrich syndrome (WAS)	Progressive decrease, Abnormal lymphocyte responses to anti-CD3	Normal	Decreased IgM: antibody to polysaccharides particularly decreased; often increased IgA and IgE	Thrombocytopenia with small platelets; eczema; lymphoma; autoimmune disease; IgA nephropathy; bacterial and viral infections. XL thrombocytopenia is a mild form of WAS, and XL neutropenia is caused by missense mutations in the GTPase binding domain of WASP	XL	Mutations in WAS; cytoskeletal and immunologic synapse defect affecting haematopoietic stem cell derivatives	<u>301000</u>
2. DNA repair defects (other than those in Table 1)							
(a) Ataxia-telangiectasia	Progressive decrease	Normal	Often decreased IgA, IgE and IgG subclasses; increased IgM monomers; antibodies variably decreased	Ataxia; telangiectasia; pulmonary infections; lymphoreticular and other malignancies; increased alpha fetoprotein and X-ray sensitivity; chromosomal instability	AR	Mutations in <i>ATM</i> ; disorder of cell cycle check-point and DNA double- strand break repair	<u>208900</u>
(b) Ataxia-telangiectasia-like disease (ATLD)*	Progressive decrease	Normal	Antibodies variably decreased	Moderate ataxia; pulmonary infections; severely increased radiosensitivity	AR	Hypomorphic mutations in <i>MRE11</i> ; disorder of cell cycle checkpoint and DNA double- strand break repair	<u>604391</u>
(c) Nijmegen breakage syndrome	Progressive decrease	Variably reduced	Often decreased IgA, IgE and IgG subclasses; increased IgM; antibodies variably decreased	Microcephaly; bird-like face; lymphomas; solid tumors; ionizing radiation sensitivity; chromosomal instability	AR	Hypomorphic mutations in NBS1 (Nibrin); disorder of cell cycle checkpoint and DNA double- strand break repair	<u>251260</u>
(d) Bloom Syndrome	Normal	Normal	Reduced	Short stature; bird like face; sun- sensitive erythema; marrow failure; leukemia; lymphoma; chromosomal instability	AR	Mutations in <i>BLM</i> ; RecQ like helicase	<u>210900</u>
(e) Immunodeficiency with centromeric instability and facial anomalies (ICF)	Decreased or normal; Responses to PHA may be decreased	Decreased or normal	Hypogammaglobulinemia; variable antibody deficiency	Facial dysmorphic features; macroglossia; bacterial/opportunistic infections; malabsorption; cytopenias; malignancies; multiradial configurations of chromosomes 1, 9, 16; no DNA breaks	AR	Mutations in DNA methyltransferase <i>DNMT3B</i> (ICF1) resulting in defective DNA methylation; or in <i>ZBTB24</i> (ICF2)	<u>242860</u>

<ul> <li>(f) PMS2 Deficiency (Class Switch recombination deficiency due to impaired mismatch repair)</li> <li>(g) Riddle Syndrome*</li> </ul>	Normal	Switched and non-switched B cells are reduced Normal	Low IgG and IgA, elevated IgM, abnormal antibody responses Low IgG	Recurrent infections; café-au-lait spots ; lymphoma, colorectal carcinoma, brain tumor Mild motor control and learning difficulties, mild facial dysmorphism,	AR	Mutations in <i>PMS2</i> , resulting in defective CSR-induced DNA double strand breaks in Ig switch regions Mutations in RNF168, resulting in defective	<u>600259</u> <u>611943</u>	
				and short stature		DNA double-strand break repair		
3. Thymic defects								
DiGeorge anomaly (Chromosome 22q11.2 deletion syndrome)	Decreased or normal	Normal	Normal or decreased	Hypoparathyroidism, conotruncal malformation; abnormal facies; large deletion (3Mb) in 22q11.2 (or rarely a deletion in 10p)	De novo defect or AD	Contiguous gene defect in 90% affecting thymic development; mutation in <i>TBX1</i>		<b>Formatted:</b> Position: Horizontal: 1.13", Relative to: Page, Vertical: 0.3", Relative
4. Immune-osseous dysplasias							to	o: Paragraph, Horizontal: 0.13", Wrap
(a) Cartilage hair hypoplasia	Decreased or 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	AR	Mutations in <i>RMRP</i> (RNase MRP RNA) Involved in processing of mitochondrial RNA and cell cycle control	230230	vround
(b) Schimke syndrome	Decreased	Normal	Normal	Short stature, spondiloepiphyseal dysplasia, intrauterine growth retardation, nephropathy; bacterial, viral, fungal infections; may present as SCID; bone marrow failure	AR	Mutations in SMARCAL1 Involved in chromatin remodeling	<u>242900</u>	
5. Comel-Netherton Syndrome	Normal	Switched and non-switched B cells are reduced	Elevated IgE and IgA Antibody variably decreased	Congenital ichthyosis, bamboo hair, atopic diathesis, increased bacterial infections, failure to thrive	AR	Mutations in <i>SPINK5</i> resulting in lack of the serine protease inhibitor LEKTI, expressed in epithelial cells	<u>256500</u>	
6. Hyper-IgE syndromes (HIES)								
(a) AD-HIES (Job Syndrome)	Normal Th-17 cells decreased	Normal (Switched and non-switched memory B cells are reduced; BAFF level increased)	Elevated IgE; specific antibody production decreased	Distinctive facial features (broad nasal bridge), eczema, osteoporosis and fractures, scoliosis, failure/delay of shedding primary teeth, hyperextensible joints, bacterial infections (skin and pulmonary abscesses, pneumatoceles)due to <i>Staphylococcus aureus</i> , candidiasis	AD Often <i>de</i> <i>novo</i> defect	Dominant-negative heterozygous mutations in <i>STAT3</i>	<u>147060</u>	

(b) AR-HIES				No skeletal and connective tissue	AR		<u>243700</u>	
(i) Tyk2 deficiency*	Normal, but Multiple cytokine signaling defect	Normal	(+/-) Elevated IgE	abnormalities; no pneumatoceles Susceptibility to intracellular bacteria (Mycobacteria, Salmonella), fungi and viruses		Mutation in TYK2	611521 611432	Formatted: Position: Horizontal: 1.13", Relative to: Page, Vertical: 0.3", Relative to: Paragraph, Horizontal: 0.13", Wrap Around
(ii) DOCK8 deficiency	Reduced	Reduced	(+/-) Elevated IgE, low IgM	Recurrent respiratory infections; extensive cutaneous viral and staphylococcal infections, increased risk of cancer, severe atopy with anaphylaxis		Mutation in DOCK8		
(iii) Unknown origin	Normal	Normal	Elevated IgE	CNS hemorrhage, fungal and viral infections		Unknown		
7. Hepatic veno-occlusive disease with immunodeficiency (VODI)	Normal (Decreased memory T cells)	Normal (Decreased memory B cells)	Decreased IgG, IgA, IgM Absent germinal centers Absent tissue plasma cells	Hepatic veno-occlusive disease; Pneumocystis jiroveci pneumonia; Susceptibility to CMV, candida; thrombocytopenia; hepatosplenomegaly	AR	Mutations in SP110	235550	
8. Dyskeratosis congenita (DKC)								
(a) XL-DKC (Hoyeraal- Hreidarsson Syndrome)	Progressive decrease	Progressive decrease	Variable	Intrauterine growth retardation, microcephaly, nail dystrophy, recurrent infections, digestive tract involvement, pancytopenia, reduced number and function of NK cells	XL	Mutations in Dyskerin (DKC1)	<u>305000</u>	
(b) AR-DKC	Abnormal	Variable	Variable	Pancytopenia, sparse scalp hair and eyelashes, prominent periorbital telangiectasia, and hypoplastic/dysplastic nails	AR	Mutation in NOLA2 (NHP2) Mutation in NOLA3 (NOP10)	224230	
(c) AD-DKC	Variable	Variable	Variable	Reticular hyperpigmentation of the skin, dystrophic nails, osteoporosis, premalignant leukokeratosis of the mouth mucosa, palmar hyperkeratosis, anemia, pancytopenia	AD	Mutation in <i>TERC</i> Mutation in <i>TERT</i> Mutation in <i>TINF2</i>	<u>127550</u>	
9. IKAROS deficiency*	Normal, but Impaired Iymphocyte proliferation	Absent	Presumably decreased	Anemia, neutropenia, thrombocytopenia	AD de novo	Mutation in <i>IKAROS</i>	<u>603023</u>	

SCID = Severe Combined Immune Deficiencies; XL= X-linked inheritance; AR= autosomal recessive inheritance; AD = autosomal dominant inheritance; MSMD, Mendelian susceptibility of mycobacterial disease

\*Ten or fewer unrelated cases reported in the literature

Notes: Three disorders listed in Table II, complete DiGeorge anomaly, cartilage hair hypoplasia and AR-HIES caused by DOCK 8 deficiency, are also included in Table I as they are characterized by striking T and B cell abnormalities. While not all DOCK-8 deficient patients have elevated serum IgE, most have recurrent viral infections and malignancies as a result of combined immunodeficiency. AR-HIES due to Tyk2 deficiency is also described in Table VI, because of its association with atypical mycobacterial disease resulting in MSMD. Because Riddle syndrome is caused by mutations in a gene involved in DNA double-strand break repair and is associated with hypogammaglobulinemia, we have added this rare syndrome to Table II. Chronic mucocutaneous candidiasis (CMC) has been moved to Table VI. Autosomal dominant and autosomal recessive forms of Dyskeratosis congenita, caused by mutations of recently identified genes, have been included in this table. Finally, we added IKAROS deficiency, observed in a single case, a prematurely born infant, who died at the age of 87 days. He had absent B and NK cells and non-functional T cells, suggesting combined immunodeficiency.

Disease	Serum Ig	Associated features	Inheritance	Genetic defect/ Presumed pathogenesis	OMIM number
1. Severe reduction in all serum immunoglobulin isotypes with profoundly decreased or absent B cells					
(a) BTK deficiency	All isotypes decreased in majority of patients; some patients have detectable immunoglobulins	Severe bacterial infections; normal numbers of pro-B cells	XL	Mutations in <i>BTK</i> , a cytoplasmic tyrosine kinase activated by crosslinking of the BCR	<u>300300</u>
(b) $\mu$ heavy chain deficiency	All isotypes decreased	Severe bacterial infections; normal numbers of pro-B cells	AR	Mutations in $\boldsymbol{\mu}$ heavy chain	<u>147020</u>
(c) $\lambda 5$ deficiency*	All isotypes decreased	Severe bacterial infections; normal numbers of pro-B cells	AR	Mutations in $\lambda 5$ ; part of the surrogate light chain in the pre-BCR	<u>146770</u>
(d) $Ig\alpha$ deficiency*	All isotypes decreased	Severe bacterial infections; normal numbers of pro-B cells	AR	Mutations in Ig $\alpha$ ( <i>CD79a</i> ); part of the pre-BCR and BCR	<u>112205</u>
(e) Igβ deficiency*	All isotypes decreased	Severe bacterial infections; normal numbers of pro-B cells	AR	Mutations in Ig $\beta$ ( <i>CD79b</i> ); part of the pre-BCR and BCR	<u>147245</u>
(f) BLNK deficiency*	All isotypes decreased	Severe bacterial infections; normal numbers of pro-B cells	AR	Mutations in <i>BLNK</i> ; a scaffold protein that binds to BTK	<u>604615</u>
(g) Thymoma with immunodeficiency	One or more isotypes may be decreased	Bacterial and opportunistic infections; autoimmunity; decreased number of pro-B cells	None	Unknown	
(h) Myelodysplasia with hypogammaglobulinemia	One or more isotypes may be decreased	Infections; decreased number of pro-B cells	Variable	May have monosomy 7, trisomy 8 or dyskeratosis congenita	
2. Severe reduction in at least 2 serum immunoglobulin isotypes with normal or low number of B cells					
(a) Common variable immunodeficiency disorders	Low IgG and IgA and/or IgM	Clinical phenotypes vary: most have recurrent infections, some have polyclonal lymphoproliferation, autoimmune cytopenias and/or granulomatous disease	Variable	Unknown	
(b) ICOS deficiency*	Low IgG and IgA and/or IgM		AR	Mutations in ICOS	<u>604558</u>
(c) CD19 deficiency*	Low IgG and IgA and/or IgM	May have glomerulonephritis	AR	Mutations in <i>CD19</i> ; transmembrane protein that amplifies signal through BCR	<u>107265</u>
(d) CD81 deficiency*	Low IgG, low or normal IgA and IgM	May have glomerulonephritis	AR	Mutations in <i>CD81</i> ; transmembrane protein that amplifies signal through BCR	<u>186845</u>
(e) CD20 deficiency*	Low IgG, normal or elevated IgM and IgA		AR	Mutations in CD20	<u>112210</u>

## Table III. Predominantly antibody deficiencies

(f) TACI deficiency	Low IgG and IgA and/or IgM	Variable clinical expression	AD or AR or complex	Mutations in TNFRSF13B (TACI)	<u>604907</u>
(g) BAFF receptor deficiency*	Low IgG and IgM	Variable clinical expression	AR	Mutations in TNFRSF13C (BAFF-R)	606269
3. Severe reduction in serum IgG and IgA with normal/elevated IgM and normal numbers of B cells					
(a) CD40L deficiency	IgG and IgA decreased; IgM may be normal or increased; B cell numbers may be normal or increased	Opportunistic infections, neutropenia, autoimmune disease	XL	Mutations in <i>CD40LG</i> (also called <i>TNFSF5</i> or <i>CD154</i> )	<u>300386</u>
(b) CD40 deficiency*	Low IgG and IgA; normal or raised IgM	Opportunistic infections, neutropenia, autoimmune disease	AR	Mutations in <i>CD40</i> (also called <i>TNFRSF5</i> )	<u>109535</u>
(c) AID deficiency	IgG and IgA decreased; IgM increased	Enlarged lymph nodes and germinal centers	AR	Mutations in AICDA gene	<u>605257</u>
(d) UNG deficiency	IgG and IgA decreased; IgM increased	Enlarged lymph nodes and germinal centers	AR	Mutations in UNG	<u>191525</u>
4. Isotype or light chain deficiencies with normal numbers of B cells					
(a) Ig heavy chain mutations and deletions	One or more IgG and/or IgA subclasses as well as IgE may be absent	May be asymptomatic	AR	Mutation or chromosomal deletion at 14q32	
(b) $\kappa$ chain deficiency*	All immunoglobulins have lambda light chain	Asymptomatic	AR	Mutations in Kappa constant gene	<u>147200</u>
(c) Isolated IgG subclass deficiency	Reduction in one or more IgG subclass	Usually asymptomatic; a minority may have poor antibody response to specific antigens and recurrent viral/bacterial infections	Variable	Unknown	
(d) IgA with IgG subclass deficiency	Reduced IgA with decrease in one or more IgG subclass	Recurrent bacterial infections in majority	Variable	Unknown	
(e) Selective IgA deficiency	IgA decreased/absent	Usually asymptomatic; may have recurrent infections with poor antibody responses to carbohydrate antigens; may have allergies or autoimmune disease. A very few cases progress to CVID, others coexist with CVID in the family	Variable	Unknown	
5. Specific antibody deficiency with normal Ig concentrations and normal numbers of B cells	Normal	Reduced ability to make antibodies to specific antigens	Variable	Unknown	
6. Transient hypogammaglobulinemia of infancy with normal numbers of B cells	IgG and IgA decreased	Normal ability to make antibodies to vaccine antigens, usually not associated with significant infections	Variable	Unknown	

XL, X-linked inheritance; AR, autosomal recessive inheritance; AD, autosomal dominant inheritance; BTK, Bruton tyrosine kinase; BLNK, B cell linker protein; AID, activation-induced cytidine deaminase; UNG, uracil-DNA glycosylase; ICOS, inducible costimulator; Ig(κ), immunoglobulin or κ light-chain type;

\*Ten or fewer unrelated cases reported in the literature

Notes: Two new autosomal recessive disorders that might previously have been called CVID have been added to Table III. CD81 is normally coexpressed with CD19 on the surface of B cells. Like CD19 mutations, mutations in CD81 result in normal numbers of peripheral blood B cells, low serum IgG and an increased incidence of glomerulonephritis. A single patient with a homozygous mutation in CD20 has been reported.

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. Alterations  $\Box$  in *TNFRSF13B (TACI)* and *TNFRSF13C (BAFF-R)* sequences may represent disease modifying mutations rather than disease causing mutations. CD40L and CD40 deficiency are included in Table 1 as well as this table. A small minority of patients with XLP (Table IV), WHIM syndrome (Table VI), ICF (Table II), VOD1 (Table II), 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. Patients with GATA2 mutations (Table V) may have markedly reduced numbers of B cells, as well as decreased monocytes and NK cells and a predisposition to myelodysplasia but they do not have an antibody deficiency.

Disease	Circulating T Cells	Circulating B cells	Serum Ig	Associated Features	Inheritance	Genetic defect/Presumed pathogenesis	OMIM number
1. Immunodeficiency with hypopigmentation							
(a) Chediak-Higashi syndrome	Normal	Normal	Normal	Partial albinism, recurrent infections, late-onset primary encephalopathy, increased lymphoma risk. Neutropenia, Giant lysosomes, low NK and CTL activities, elevation of acute-phase markers,	AR	Mutations in <i>LYST</i> , impaired lysosomal trafficking	<u>214500</u>
(b) Griscelli syndrome, type2	Normal	Normal	Normal	Partial albinism, elevation of acute phase markers, encephalopathy in some patients. Low NK and CTL activities,	AR	Mutations in <i>RAB27A</i> encoding a GTPase that promotes docking of secretory vesicles to the cell membrane	<u>607624</u>
(c) Hermansky-Pudlak syndrome, type 2*	Normal	Normal	Normal	Partial albinism, increased bleeding. Neutropenia, low NK and CTL activity,	AR	Mutations in the <i>AP3B1</i> gene, encoding for the $\beta$ subunit of the AP-3 complex	<u>608233</u>
2. Familial hemophagocytic lymphohistiocytosis (FHL) syndromes							
(a) Perforin deficiency, FHL2	Normal	Normal	Normal	Severe inflammation, persistent fever, cytopenias, splenomegaly. Hemophagocytosis, decreased to absent NK and CTL activities	AR	Mutations in <i>PRF1</i> ; perforin, a major cytolytic protein	<u>603553</u>
(b) UNC13D (Munc13-4) deficiency, FHL3	Normal	Normal	Normal	Severe inflammation, persistent fever, splenomegaly, Hemophagocytosis, decreased NK and CTL activities	AR	Mutations in UNC13D* required to prime vesicles for fusion (*as named in OMIM). Note that also in OMIM the "official" name is UNC13D deficiency with the alternative title of MUNC13D deficiency	<u>608898</u>
(c) Syntaxin 11 deficiency, FHL4	Normal	Normal	Normal	Severe inflammation, persistent fever, splenomegaly. Hemophagocytosis, decreased to absent NK activity	AR	Mutations in <i>STX11</i> , required for fusion of secretory vesicles with the cell membrane and release of contents	<u>603552</u>
(d) STXBP2 (Munc 18-2) deficiency, FHL5	Normal	Normal	Normal or low	Severe inflammation, fever, splenomegaly, hemophagocytosis possible bowel disease. Decreased NK and CTL activities with partial restoration after IL-2	AR	Mutations in <i>STXBP2</i> , required for fusion of secretory vesicles with the cell membrane and release of contents	<u>613101</u>

## TABLE IV. Diseases of immune dysregulation

		1		stimulation			
3. Lymphoproliferative syndromes							
(a) SH2D1A deficiency, XLP1	Normal	Normal or reduced	Normal or low	Clinical and immunologic abnormalities triggered by EBV infection, including hepatitis, hemophagocytic syndrome, aplastic anaemia and lymphoma. Dysgammaglobulinemia or hypogammaglobulinemia, low to absent NKT cells	XL	Mutations in SH2D1A encoding an adaptor protein regulating intracellular signals	<u>308240</u>
(b) XIAP deficiency, XLP2	Normal	Normal or reduced	Normal or low	Clinical and immunologic abnormalities triggered by EBV infection, including splenomegaly, hepatitis, hemophagocytic syndrome colitis	XL	Mutations in <i>XIAP</i> encoding an inhibitor of apoptosis	<u>300635</u>
4. Syndromes with autoimmunity							
(a) Autoimmune lymphoproliferative syndrome (ALPS)							
(i) ALPS-FAS	Increased CD4 <sup>-</sup> CD8 <sup>-</sup> double negative (DN) T cells	Normal	Normal or increased	Splenomegaly, adenopathies, autoimmune cytopenias, increased lymphoma risk. Defective lymphocyte apoptosis.	AD (AR cases are rare and severe, ALPS)	Mutations in <i>TNFRSF6</i> , cell surface apoptosis receptor; in addition to germline mutations, somatic mutations cause a similar phenotype (ALPS SFAS)	<u>601859</u>
(ii) ALPS-FASG	Increased DN T cells	Normal	Normal	Splenomegaly, adenopathies, autoimmune cytopenias, SLE defective lymphocyte apoptosis,	AD AR	Mutations in <i>TNFSF6</i> , ligand for CD95 apoptosis receptor	<u>134638</u>
(iii) ALPS-CASP10*	Increased DN T cells	Normal	Normal	Adenopathies, splenomegaly, autoimmunity. Defective lymphocyte apoptosis	AD	Mutations in CASP10, intracellular apoptosis pathway	<u>603909</u>
(iv) Caspase 8 defect*	Slightly increased DN T cells	Normal	Normal or decreased	Adenopathies, splenomegaly, recurrent bacterial and viral infections. Defective lymphocyte apoptosis and activation, hypogammaglobulinemia	AD	Mutations in CASP8, intracellular apoptosis and activation pathways	<u>607271</u>
(v) Activating N-Ras defect, Activating Kras defect*	Increased or normal DN T cells	Elevation of CD5 B cells	Normal	Adenopathies, splenomegaly, leukemia, lymphoma. Defective lymphocyte apoptosis following IL-2 withdrawal	Sporadic	Somatic mutations in NRAS encoding a GTP binding protein with diverse signaling functions; activating mutations impair mitochondrial apoptosis	<u>164790</u>
(vi) FADD deficiency*	Increased DN T cells	Normal	Normal	Functional hyposplenism, recurrent bacterial and viral infections, recurrent episodes of encephalopathy and liver	AR	Mutations in FADD encoding an adaptor molecule interacting with FAS, and promoting apoptosis,	<u>613759</u>

				dysfunction. Defective lymphocyte apoptosis,		inflammation and innate immunity	
(b) APECED (APS-1), autoimmune polyendocrinopathy with candidiasis and ectodermal dystrophy	Normal	Normal	Normal	Autoimmunity, particularly of parathyroid, adrenal and other endocrine organs, chronic candidiasis, dental enamel hypoplasia and other abnormalities	AR	Mutations in <i>AIRE</i> , encoding a transcription regulator needed to establish thymic self-tolerance	<u>240300</u>
(c) IPEX, immune dysregulation, polyendocrinopathy, enteropathy (X-linked)	Lack of (and/or impaired function of) CD4 <sup>+</sup> CD25 <sup>+</sup> FOXP3 <sup>+</sup> regulatory T cells	Normal	Elevated IgA, IgE	Autoimmune enteropathy, early onset diabetes, thyroiditis hemolytic anemia, thrombocytopenia, eczema	XL	Mutations in <i>FOXP3</i> , encoding a T cell transcription factor	<u>304790</u>
(d) CD25 deficiency	Normal to modestly decreased	Normal	Normal	Lymphoproliferation, autoimmunity. Impaired T cell proliferation	AR	Mutations in IL-2R $\alpha$ chain	<u>606367</u>
(e) ITCH deficiency*	Not assessed (Th2 skewing in <i>Itch</i> -deficient mice)	Not assessed (B cells are dysfunctional in <i>Itch</i> - deficient mice)	Not assessed (elevated in <i>Itch</i> - deficient mice)	Multi-organ autoimmunity, chronic lung disease, failure to thrive, developmental delay, macrocephaly	AR	Mutations in <i>ITCH</i> , an E3 ubiquitin ligase	<u>613385</u>

XL, X-linked inheritance; AR, autosomal recessive inheritance; AD, autosomal dominant inheritance; DN, double-negative; SL, systemic lupus erythematosus

\*Ten or fewer unrelated cases reported in the literature

Notes: STXBP2/Munc18-2 deficiency has been added as the cause of "FHL5", a new form of FHL. Of note, "FHL1" has not yet received a genetic/molecular identification. FADD deficiency is classified among the causes of ALPS. It should be stressed however that FADD deficiency is a more complex syndrome that encompasses hyposplenism, hence bacterial infections, as well as a brain and liver primary dysfunction. EBV-driven lymphoproliferation is also observed in ITK deficiency and in MAGT1 deficiency (Table I).

Disease	Affected cells	Affected function	Associated features	Inheritance	Genetic defect/ Presumed pathogenesis	OMIM number
1.Defects of Neutrophil Differentiation						
(a) Severe congenital neutropenia 1 (ELANE deficiency)	N	Myeloid differentiation	Subgroup with myelodysplasia	AD	ELANE: misfolded protein response	<u>202700</u>
(b) SCN2 * (GFI 1 deficiency)	N	Myeloid differentiation	B/T lymphopenia	AD	GFI1: loss of repression of ELANE	<u>613107</u>
(c) SCN3 (Kostmann Disease)	N	Myeloid differentiation	Cognitive and neurological defects in some patients	AR	HAX1: control of apoptosis	<u>610738</u>
(d) SCN4 (G6PC3 deficiency)	N + F	Myeloid differentiation, chemotaxis, $O_2^{-}$ production	Structural heart defects, urogenital abnormalities, and venous angiectasias of trunks and limbs	AR	<i>G6PC3</i> : abolished enzymatic activity of glucose-6- phosphatase, aberrant glycosylation, and enhanced apoptosis of N and F	<u>612541</u>
(e) Glycogen storage disease type 1b	N + M	Myeloid differentiation, chemotaxis, $O_2^{-}$ production	Fasting hypoglycemia, lactic acidosis, hyperlipidemia, hepatomegaly	AR	<i>G6PT1</i> : Glucose-6- phosphate transporter 1	<u>232220</u>
(f) Cyclic neutropenia	N	?	Oscillations of other leukocytes and platelets	AD	ELANE: misfolded protein response	<u>162800</u>
(g) X-linked neutropenia/ * myelodysplasia	N + M	Mitosis	Monocytopenia	XL	WAS: Regulator of actin cytoskeleton (loss of autoinhibition)	<u>300299</u>
(h) P14 deficiency *	N+L Mel	Endosome biogenesis	Neutropenia Hypogammaglobulinemi a ↓CD8 cytotoxicity Partial albinism Growth failure	AR	ROBLD3: Endosomal adaptor protein 14	<u>610389</u>
(i) Barth Syndrome	N	Myeloid differentiation	Cardiomyopathy, growth retardation	XL	Tafazzin ( <i>TAZ</i> ) gene: Abnormal lipid structure of mitochondrial membrane	<u>302060</u>
(j) Cohen syndrome	N	Myeloid differentiation	Retinopathy, developmental delay, facial dysmorphisms	AR	COH1 gene: Pg unknown	<u>216550</u>

## TABLE V. Congenital defects of phagocyte number, function, or both

(k) Poikiloderma with neutropenia	N	Myeloid differentiation, $O_2^-$ production	Poikiloderma, MDS	AR	C16orf57 gene: Pg unknown	<u>604173</u>
2. Defects of Motility						
(a) Leukocyte adhesion deficiency type 1 (LAD1)	N + M + L + NK	Adherence, Chemotaxis, Endocytosis, T/NK cytotoxicity	Delayed cord separation, skin ulcers Periodontitis Leukocytosis	AR	INTGB2: Adhesion protein (CD18)	<u>116920</u>
(b) Leukocyte adhesion deficiency type 2 (LAD2)*	N + M	Rolling, chemotaxis	Mild LAD type 1 features plus hh-blood group plus mental and growth retardation	AR	FUCT1: GDP-Fucose transporter	<u>266265</u>
(c) Leukocyte adhesion deficiency type 3 (LAD3)	N + M + L + NK	Adherence, chemotaxis	LAD type 1 plus bleeding tendency	AR	KINDLIN3: Rap1-activation of $\beta$ 1-3 integrins	<u>612840</u>
(d) Rac 2 deficiency*	N	Adherence, chemotaxis O <sub>2</sub> <sup>-</sup> production	Poor wound healing, leukocytosis	AD	<i>RAC2</i> : Regulation of actin cytoskeleton	<u>602049</u>
(e) β-actin deficiency*	N + M	Motility	Mental retardation, short stature	AD	ACTB: Cytoplasmic Actin	<u>102630</u>
(f) Localized juvenile Periodontitis	N	Formylpeptide induced chemotaxis	Periodontitis only	AR	FPR1: Chemokine receptor	<u>136537</u>
(g) Papillon-Lefèvre Syndrome	N + M	Chemotaxis	Periodontitis, palmoplantar hyperkeratosis in some patients	AR	CTSC: Cathepsin C activation of serine proteases	<u>245000</u>
(h) Specific granule deficiency*	Ν	Chemotaxis	Neutrophils with bilobed nuclei	AR	<i>C/EBPE</i> : myeloid transcription factor	<u>245480</u>
(i) Shwachman-Diamond Syndrome	N	Chemotaxis	Pancytopenia, exocrine pancreatic insufficiency, chondrodysplasia	AR	SBDS: Defective ribosome synthesis	<u>260400</u>

3. Defects of Respiratory Burst						
(a) X-linked chronic granulomatous disease (CGD)	N + M	Killing (faulty O <sub>2</sub> <sup>-</sup> production)	McLeod phenotype in patients with deletions extending into the contiguous Kell locus	XL	CYBB: Electron transport protein (gp91phox)	<u>306400</u>
(b-e) Autosomal CGD's	N + M	Killing (faulty O <sub>2</sub> <sup>-</sup> production)		AR	CYBA: Electron transport protein (p22phox) NCF1: Adapter protein (p47phox) NCF2: Activating protein (p67phox) NCF4: Activating protein (p40 phox)	233690 233700 233710 601488
4. MSMD						
<ul> <li>(a) IL-12 and IL-23 receptor β1 chain deficiency</li> </ul>	L + NK	IFN-γ secretion	Susceptibility to <i>Mycobacteria</i> and <i>Salmonella</i>	AR	<i>IL12RB1</i> : IL-12 and IL-23 receptor β1 chain	<u>601604</u>
(b) IL-12p40 deficiency	М	IFN-γ secretion	Susceptibility to <i>Mycobacteria</i> and <i>Salmonella</i>	AR	IL12B : subunit of IL12/IL23	<u>161561</u>
(c) IFN-γ receptor 1 deficiency	M + L	IFN-γ binding and signaling	Susceptibility to <i>Mycobacteria</i> and <i>Salmonella</i>	AR, AD	<i>IFNGR1</i> : IFN-γR ligand binding chain	<u>107470</u>
(d) IFN-γ receptor 2 deficiency	M + L	IFN-γ signaling	Susceptibility to <i>Mycobacteria</i> and <i>Salmonella</i>	AR	<i>IFNGR2</i> : IFN-γR accessory chain	<u>147569</u>
(e) STAT1 deficiency (AD form)*	M + L	IFN-γsignaling	Susceptibility to Mycobacteria, Salmonella	AD	STAT1	<u>600555</u>
(f) Macrophage gp91 phox deficiency*	Mø only	Killing (faulty O <sub>2</sub> <sup>-</sup> production)	Isolated susceptibility to mycobacteria	XL	CYBB: Electron transport protein (gp 91 phox)	<u>306400</u>
(g) IRF8-deficiency (AD form)*	CD1c+ MDC	Differentiation of CD1c+ MDC subgroup	Susceptibility to <i>Mycobacteria</i>	AD	<i>IRF8:</i> IL12 production by CD1c <sup>+</sup> MDC	<u>601565</u>

5. Other Defects						
(a) IRF 8-deficiency (AR form)*	Monocytes peripheral DC	Cytopenias	Susceptibility to <i>Mycobacteria</i> , Candida, Myeloproliferation	AR	IRF8: IL12 production	
(b) GATA2 deficiency (Mono MAC Syndrome)	Monocytes peripheral DC +NK+B	Multilineage cytopenias	Susceptibility to <i>Mycocbacteria</i> , Papilloma Viruses, Histoplasmosis, Alveolar proteinosis, MDS/AML/CMML	AD	GATA-2: loss of stem cells	<u>137295</u>
(c) Pulmonary alveolar proteinosis*	Alveolar macrophage s	GM-CSF signaling	Alveolar proteinosis	Biallelic mutations in pseudoautosomal gene	CSF2RA	<u>306250</u>

XL, X-linked inheritance; AR, autosomal recessive inheritance; AD, autosomal dominant inheritance; ACTB, Actin beta; B, B lymphocytes; CEBPE, CCAAT/Enhancer-binding protein epsilon; CMML, chronic myelomonocytic leukaemia; CTSC, cathepsin C; CYBA, cytochrome b alpha subunit; CYBB, cytochrome b beta subunit; DC, Dendritic cells; ELANE elastase neutrophil-expressed; GATA2, GATA binding protein 2; IFN, interferon; IFNGR1, interferon-gamma receptor subunit 1;IFNGR2, interferon-gamma receptor subunit; L12B, interleukin-12 beta subunit;IL12RB1, interleukin-12 receptor beta 1; IFR8, interferon regulatory factor 8; F, fibroblasts; FPR1, formylpeptide receptor 1; FUCT1, fucose transporter 1; GFI1, growth factor independent 1; HAX1, HLCS1-associated protein X1; ITGB2, integrin beta-2; L, lymphocytes; M, monocytes-macrophages; MDC, myeloid dendritic cells; MDS, myelodysplasia; Mel, melanocytes; Mol, macrophages; MSMD, Mendelian susceptibility to mycobacterial disease; N, neutrophils; NCF1, neutrophil cytosolic factor 1; NCF2, neutrophil cytosolic factor 2; NCF4, neutrophil cytosolic factor 4; NK, natural killer cells; ROBLD3: roadblock domain containing 3; SBDS, Shwachman-Bodian-Diamond syndrome; STAT, signal transducer and activator of transcription.

#### \*Ten or fewer unrelated cases reported in the literature

Notes: Table V includes seven newly described genetic defects of phagocyte number and/or function including Barth-syndrome, Cohen syndrome and Poikiloderma with neutropenia. In these three clinically well-known diseases the genetic defects have been elucidated, although their molecular pathogenesis remains ill-defined. A new cause of autosomal recessive chronic granulomatous disease, namely a deficiency of the cytosolic activating protein p40 phox, has now been found in two CGD patients and is included under defects of respiratory burst. Under the heading of Mendelian susceptibility of mycobacterial disease (MSMD) two new entities were added: a) a *subgroup of X-linked gp91 phox deficiency* with isolated susceptibility to mycobacteria and a defect of the respiratory burst in macrophages only; b) an autosomal dominant form of IRF8 deficiency, resulting from a lack of CD1c+ myeloid dendritic cells that would normally secrete IL12. The clinical phenotype of MSMD may vary, depending on the nature of the genetic defect. Finally GATA2 deficiency was recently identified as the cause of the Mono MAC syndrome, with multilineage cytopenias (of monocytes, peripheral dendritic cells, NK- and B-lymphocytes) resulting in opportunistic infections (including mycobacteria), alveolar proteinosis and malignancy.

Table VI: Defects in innate immunity	Table	VI: Defects	in innate	immunity
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Disease	Affected Cell	Functional Defect	Associated Features	Inheritance	Genetic defect/Presumed pathogenesis	OMIM number
1. Anhidrotic ectodermal dysplasia with immunodeficiency (EDA-ID)						
(a) EDA-ID, X-linked (NEMO deficiency)	Lymphocytes + Monocytes	NFκB signaling pathway	anhidrotic ectodermal dysplasia + specific antibody deficiency (lack of Ab response to polysaccharides) + various infections (mycobacteria and pyogens)	XL	Mutations of NEMO ( <i>IKBKG</i> ), a modulator of NF-κB activation	<u>300291,</u> <u>300584,</u> <u>300301</u>
(b) EDA-ID, autosomal-dominant*	Lymphocytes + Monocytes	NFκB signaling pathway	Anhidrotic ectodermal dysplasia + T cell defect + various infections	AD	Gain-of-function mutation of <i>IKBA</i> , resulting in impaired activation of NF-□B	<u>612132</u>
2. IRAK4 deficiency	Lymphocytes + Monocytes	TIR-IRAK signaling pathway	Bacterial infections (pyogens)	AR	Mutation of <i>IRAK4</i> , a component of TLR- and IL-1R-signaling pathway	<u>607676</u>
3. MyD88 deficiency	lymphocytes + Monocytes	TIR-MyD88 signaling pathway	Bacterial infections (pyogens)	AR	Mutation of <i>MYD88</i> , a component of the TLR and IL-1R signaling pathway	<u>612260</u>
4. WHIM (Warts, Hypogammaglobulinemia, infections, Myelokathexis) syndrome	Granulocytes + Lymphocytes	Increased response of the CXCR4 chemokine receptor to its ligand CXCL12 (SDF-1)	Hypogammaglobulinemia, reduced B cell number, severe reduction of neutrophil count, warts/HPV infection	AD	Gain-of-function mutations of <i>CXCR4</i> , the receptor for CXCL12	<u>193670</u>
5. Epidermodysplasia verruciformis	Keratinocytes and leukocytes		Human Papilloma virus (group B1) infections and cancer of the skin	AR	Mutations of EVER1, EVER2	<u>226400</u>
6. Herpes simplex encephalitis (HSE)*						
(a) TLR3 deficiency*	Central nervous system (CNS) resident cells and fibroblasts	TLR3-dependent IFN- $\alpha$ , - $\beta$ , and – $\lambda$ induction	Herpes simplex virus 1 encephalitis	AD	(b) Mutations of <i>TLR3</i>	<u>613002</u>
(b) UNC93B1 deficiency	CNS resident cells and fibroblasts	UNC-93B-dependent IFN- $\alpha$ , - $\beta$ , and - $\lambda$ induction	Herpes simplex virus 1 encephalitis	AR	(a) Mutations of UNC93B1	<u>610551</u>
(c) TRAF3 deficiency	CNS resident cells and fibroblasts	TRAF3-dependent IFN- $\alpha$ , - $\beta$ , and - $\lambda$ induction	Herpes simplex virus 1 encephalitis	AD	(c) Mutation of TRAF3	
7. Predisposition to fungal diseases*	Mononuclear phagocytes	CARD9 signaling pathway	Invasive candidiasis and peripheral dermatophytosis	AR	Mutations of CARD9	212050
8. Chronic mucocutaneous candidiasis (CMC)						
(a) IL-17RA deficiency*	Epithelial cells,	IL-17RA signaling pathway	CMC	AR	(a) Mutation in IL17RA	<u>605461</u>

	fibroblasts, mononuclear phagocytes					
(b) IL-17F deficiency*	T cells	IL-17F-containing dimers	CMC	AD	(b) Mutation in IL17F	<u>606496</u>
(c) STAT1 gain-of-function	T cells	Gain-of-function STAT1 mutations that impair the development of IL-17- producing T cells	CMC	AD	(c) Mutations in STAT1	Not in OMIM yet
9. Trypanosomiasis*		APOL-I	Trypanosomiasis	AD	Mutation in APOL-I	<u>603743</u>

XL, X-linked inheritance; AR, autosomal recessive inheritance; AD, autosomal dominant inheritance; NF-kB, nuclear factor Kappa B; TIR, Toll and Interleukin 1 Receptor; IFN, interferon; HP, human papilloma virus; TLR, Toll-like receptor; IL: interleukin.

\*Ten or fewer unrelated cases reported in the literature

Notes: Four new disorders have been added to Table VI. AD TRAF3 deficiency is a new genetic etiology of HSE that has been diagnosed in a single patient. A new entry in the Table is CMC, for which three genetic etiologies have been discovered. AR IL-17RA deficiency and AD IL-17F deficiency have been found in one kindred each. Gain-of-function mutations in STAT1 have been found in over 50 patients with AD CMC. The mechanism of CMC in these patients involves impaired development of IL-17-producing T cells, due to the hyperactivity of STAT1-dependent signals.

XR-EDA-ID is highly heterogeneous clinically, both in terms of developmental features (some patients display osteopetrosis and lymphedema, in addition to EDA, while others do not display any developmental features) and infectious diseases (some display multiple infections, viral, fungal and bacterial, while others display a single type of infection). The various OMIM entries correspond to these distinct clinical diseases.

Table VII: Autoinflammatory disorder	Table VII:	Autoinflammator	v disorders
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Disease	Affected cells	Functional defects	Associated Features	Inheritance	Genetic defect/ Presumed	OMIM number
1. Defects effecting the inflamm		Functional defects	Associated Features	Inneritance	pathogenesis	
(a) Familial Mediterranean Fever	Mature granulocytes, cytokine-activated monocytes.	Decreased production of pyrin permits ASC-induced IL-1 processing and inflammation following subclinical serosal injury; macrophage apoptosis decreased.	Recurrent fever, serositis and inflammation responsive to colchicine. Predisoposes to vasculitis and inflammatory bowel disease.	AR	Mutations of <i>MEFV</i>	<u>249100</u>
(b) Hyper IgD syndrome		Mevalonate kinase deficiency affecting cholesterol synthesis; pathogenesis of disease unclear	Periodic fever and leukocytosis with high IgD levels	AR	Mutations of <i>MVK</i>	<u>260920</u>
(c) Muckle-Wells syndrome	PMNs Monocytes	Defect in cryopyrin, involved in leukocyte apoptosis and NFkB signaling and IL-1 processing	Urticaria, SNHL, amyloidosis.	AD	Mutations of CIAS1 (also called PYPAF1 or NALP3)	<u>191900</u>
(d) Familial cold autoinflammatory syndrome	PMNs, monocytes	same as above	Non-pruritic urticaria, arthritis, chills, fever and leukocytosis after cold exposure.	AD	Mutations of CIAS1 Mutations of NLRP12	<u>120100</u>
<ol> <li>Neonatal onset multisystem inflammatory disease (NOMID) or chronic infantile neurologic cutaneous and articular syndrome (CINCA)</li> </ol>	PMNs, chondrocytes	same as above	Neonatal onset rash, chronic meningitis, and arthropathy with fever and inflammation.	AD	Mutations of CIAS1	<u>607115</u>
2. Non inflammasome-related co	onditions					
<ul> <li>(a) TNF receptor-associated periodic syndrome (TRAPS)</li> </ul>	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	AD	Mutations of TNFRSF1A	<u>142680</u>
(b) Early onset inflammatory bowel disease	Monocyte/macrophag e, activated T cells	Mutation in IL-10 or IL-10 receptor leads to increase of TNF $\gamma$ and other proinflammatory cytokines	Early onset enterocolitis enteric fistulas, perianal abscesses, chronic folliculitis.	AR	Mutations in <i>IL-10,</i> <i>IL10RA</i> or <i>IL10RB</i>	<u>146933</u>
(c) Pyogenic sterile arthritis, pyoderma gangrenosum, acne (PAPA) syndrome	Hematopoietic tissues, upregulated in activated T-cells	Disordered actin reorganization leading to compromised physiologic signaling during inflammatory response	Destructive arthritis, inflammatory skin rash, myositis	AD	Mutations of <i>PSTPIP1</i> (also called C2BP1)	<u>604416</u>

(d) Blau syndrome	Monocytes	Mutations in nucleotide binding site of CARD15, possibly disrupting interactions with lipopolysaccharides and NF- κB signaling	Uveitis, granulomatous synovitis, camptodactyly, rash and cranial neuropathies, 30% develop Crohn's disease	AD	Mutations of <i>NOD2</i> (also called CARD15)	<u>186580</u>
<ol> <li>Chronic recurrent multifocal osteomyelitis and congenital dyserythropoietic anemia (Majeed syndrome)*</li> </ol>	Neutrophils, bone marrow cells	undefined	Chronic recurrent multifocal osteomyelitis, transfusion- dependent anemia, cutaneous inflammatory disorders	AR	Mutations of <i>LPIN2</i>	<u>609628</u>
11. DIRA (Deficiency of the Interleukin 1 Receptor Antagonist)*	PMNs, Monocytes	Mutations in the IL1 receptor antagonist allows unopposed action of Interleukin 1	Neonatal onset of sterile multifocal osteomyelitis, periostitis and pustulosis.	AR	Mutations of <i>IL1RN</i>	<u>612852</u>

AR, autosomal recessive inheritance; AD, autosomal dominant inheritance; PMN, polymorphonuclear cells; ASC, apoptosis-associated speck-like protein with a caspase recruitment domain; CARD, caspase recruitment domain; CD2BP1, CD2 binding protein-1; PSTPIP1, Proline/serine/threonine phosphatase-interacting protein 1; SNHL, sensorineural hearing loss; CIAS1- cold-induced autoinflammatory syndrome 1

\*Ten or fewer unrelated cases reported in the literature

Notes: Autoinflammatory diseases are clinical disorders marked by abnormally increased inflammation, mediated predominantly by the cells and molecules of the innate immune system, with a significant host predisposition. While the genetic defect of one of the most common autoinflammatory conditions, PFAPA, is not known, recent studies suggest that it is associated with activation of IL-1 pathway and response to IL-1 beta antagonists.

Muckle-Wells syndrome, familial cold autoinflammatory syndrome and neonatal onset multisystem inflammatory disease (NOMID) which is also called chronic infantile neurologic cutaneous and articular syndrome (CINCA) are caused by similar mutations in CIAS1 mutations. The disease phenotype in any individual appears to depend on modifying effects of other genes and environmental factors.

## Table VIII Complement deficiencies

Disease	Functional Defect	Associated Features	Inheritance	Genetic defect/Presumed pathogenesis	OMIM number
C1q deficiency	Absent CH50 hemolytic activity, Defective MAC Faulty dissolution of immune complexes Faulty clearance of apoptotic cells	SLE-like syndrome, rheumatoid disease, infections	AR	Mutations in <i>C1QA</i> , <i>C1QB</i> , <i>C1QC</i> and loss of early complement activation	<u>120550; 601269;</u> <u>120575</u>
C1r deficiency	Absent CH50 hemolytic activity, Defective MAC Faulty dissolution of immune complexes	SLE–like syndrome, rheumatoid disease, multiple autoimmune diseases, infections	AR	Mutations in C1r and loss of early complement activation	<u>216950</u>
C1s deficiency	Absent CH50 hemolytic activity	SLE-like syndrome; multiple autoimmune diseases	AR	Mutations in C1s and loss of early complement activation	<u>120580</u>
C4 deficiency	Absent CH50 hemolytic activity, Defective MAC Faulty dissolution of immune complexes Defective humoral immune response to carbohydrate antigens in some patients	SLE-like syndrome, rheumatoid disease, infections <i>C4A</i> ; homozygous; SLE, type I diabetes <i>C4B</i> : homozygous: bacterial meningitis	AR	Mutations in C4A and C4B and loss of early complement activation	<u>120810; 120820</u>
C2 deficiency	Absent CH50 hemolytic activity, Defective MAC Faulty dissolution of immune complexes	SLE-like syndrome, vasculitis, atherosclerosis, polymyositis, pyogenic infections; glomerulonephritis	AR	Mutations in <i>C2</i> and loss of early complement activation	<u>217000</u>
C3 deficiency	Absent CH50 and AP50 hemolytic activity, Defective MAC Defective Bactericidal activity Defective humoral immune response	Life threatening pyogenic infections; SLE-like disease; glomerulonephritis; Atypical Hemolytic-uremic syndrome; Selected SNPs with age related macular degeneration	AR	Mutations in <i>C3</i> and loss of complement activation by classical and alternative pathways	120700
C5 deficiency	Absent CH50 and AP50 hemolytic activity, Defective MAC Defective Bactericidal activity	Neisserial infections, SLE	AR	Mutations in $C5\alpha$ or $C5\beta$ and loss of complement activation	<u>120900</u>
C6 deficiency	Absent CH50 and AP50 hemolytic activity, Defective MAC Defective Bactericidal activity	Neisserial infections, SLE	AR	Mutations in <i>C6</i> and loss of complement activation	<u>217050</u>
C7 deficiency	Absent CH50 and AP50 hemolytic activity, Defective MAC Defective Bactericidal activity	Neisserial infections, SLE, vasculitis	AR	Mutations in <i>C7</i> and loss of terminal complement activation	<u>217070</u>
C8a deficiency	Absent CH50 and AP50 hemolytic activity, Defective MAC Defective Bactericidal activity	Neisserial infections, SLE	AR	Mutations in $C8\alpha$ and loss of terminal complement activation	<u>120950</u>
C8b deficiency	Absent CH50 and AP50 hemolytic activity, Defective MAC Defective Bactericidal activity	Neisserial infections, SLE	AR	Mutations in $C8\beta$ and loss of terminal complement activation	<u>120960</u>
C9 deficiency	Reduced CH50 and AP50 hemolytic activity, Defective MAC Defective Bactericidal activity	Neisserial infections, weaker association than in C5, C6, C7 or C8 deficiency	AR	Mutations in <i>C9</i> and loss of terminal complement activation	<u>120940</u>

C1 inhibitor	Spontaneous activation of the complement	Hereditary angioedema	AD	Mutations in C1 inhibitor	606860
deficiency	pathway with consumption of C4/C2 Spontaneous activation of the contact system with generation of bradykinin from high molecular weight kininogen			and loss of regulation of proteolytic activities of complement C1	
Factor B deficiency	Failure of activation of the alternative complement pathway with consumption of C3	Severe Neisserial infections, atypical hemolytic-uremic syndrome; some SNRs with reduced age related macular degeneration	AR	Mutations in Factor B ( <i>CFB</i> ), impair alternative pathway complement activation resulting in susceptibility to Neisserial infections	<u>138470</u>
Factor D deficiency	Absent AP50 hemolytic activity	Severe Neisserial infection	AR	Mutations in Factor D ( <i>CFD</i> ), impairing alternative complement activation	<u>134350</u>
Properdin deficiency	Absent AP50 hemolytic activity	Severe Neisserial infection	XL	Mutations in properdin ( <i>PFC</i> ), impairing alternative complement activation	<u>312060</u>
Factor I deficiency	Spontaneous activation of the alternative complement pathway with consumption of C3	Recurrent pyogenic infections, glomerulonephritis, SLE; hemolytic- uremic syndrome; Selected SNPS: Severe pre-eclampsia	AR	Mutations in Factor I ( <i>CFI</i> ), leading to accelerated catabolism of C3	<u>610984</u>
Factor H deficiency	Spontaneous activation of the alternative complement pathway with consumption of C3	Hemolytic-uremic syndrome, membranoproliferative glomerulonephritis; Neisserial infections; Selected SNPS: Severe pre-eclampsia	AR	Mutations in Factor H ( <i>CFH</i> ), leading to continuous activation of the alternative complement pathway and C3 deposition in tissues	<u>609814</u>
MASP1 deficiency	Potential loss of embryonic cell migration signals	A developmental syndrome of facial dysmorphism, cleft lip and/or palate, craniosynostosis, learning disability and genital, limb and vesicorenal anomalies.	AR	Mutations in <i>MASP1</i> leading to impaired complement pathway through the mannan- binding lectin serine proteases.	<u>600521</u>
3MC syndrome COLEC11 deficiency	Potential loss of embryonic cell migration signals	A developmental syndrome of facial dysmorphism, cleft lip and/or palate, craniosynostosis, learning disability and genital, limb and vesicorenal anomalies.	AR	Gene product CL-K1, a C- type lectin that may serve as a chemoattractant	<u>612502</u>
MASP2 deficiency*	Absent hemolytic activity by the lectin pathway	Pyogenic infections; Inflammatory lung disease	AR	Mutations in MASP2 leading to impaired complement pathway through the mannan- binding lectin serine proteases	<u>605102</u>
Complement Receptor 3 (CR3) deficiency	See LAD1 in Table V		AR	Mutations in INTGB2	<u>116920</u>
Membrane Cofactor Protein (CD46)	Inhibitor of complement alternate pathway, decreased C3b binding	Glomerulonephritis, atypical hemolytic uremic syndrome;	AD	Mutations in <i>MCP</i> leading to loss of the cofactor	<u>120920</u>

deficiency		Selected SNPS: Severe pre-eclampsia		activity needed for the Factor I-dependent cleavage of C3B and C4B	
Membrane Attack Complex Inhibitor (CD59) deficiency	Erythrocytes highly susceptible to complement- mediated lysis	Hemolytic anemia, thrombosis	AR	Mutations in <i>CD5</i> 9 leading to loss of this membrane inhibitor of the membrane attack complexes	<u>107271</u>
Paroxysmal nocturnal hemoglobinuria	Complement-mediated hemolysis	Recurrent hemolysis; hemoglobinuria, abdominal pain, smooth muscle dystonias, fatigue, and thrombosis	Acquired X- linked mutation	Disease results from the expansion of hematopoietic stem cells bearing mutations in PIGA and subsequent loss of biosynthesis of glycosylphosphatidylinositol (GPI) a moiety that attaches proteins to the cell surface.	<u>300818</u>
Immunodeficiency associated with Ficolin 3 deficiency*	Absence of complement activation by the Ficolin 3 pathway.	Recurrent severe pyogenic infections mainly in the lungs; necrotizing enterocolitis in infancy; selective antibody defect to pneumococcal polysaccharides	AR	Mutations in <i>FCN3</i> , leading to impaired complement deposition	<u>604973</u>

XL, X-linked inheritance; AR, autosomal recessive inheritance; AD, autosomal dominant inheritance; MAC, Membrane attack complex; SLE, systemic lupus erythematosus; MBP, Mannose binding Protein; MASP-2, MBP associated serine protease 2.

\*Ten or fewer unrelated cases reported in the literature

Notes: New entities added to Table VIII demonstrate the important role of complement regulators in a group of well-described inflammatory disorders. In particular, we have added mutations in membrane bound as well as surface attached soluble complement regulatory proteins recognized in hemolytic uremic syndrome, age-related macular degeneration and preeclampsia. The connecting theme of these otherwise unrelated clinical events is excessive activation or insufficient regulation of C3; these events lead to recruitment of leukocytes and permit secretion of inflammatory and anti-angiogenic mediators that disrupt the vascular bed of the target organ. Alterations  $\Box$  in the genes for Factor B (*CFB*), Factor I (*CFI*), Factor H (*CFH*) and *CD46* act as susceptibility genes rather than disease causing mutations. Population studies reveal no detectable increase in infections in MBP (also known at mannose binding lectin – MBL) deficient adults. The 3MC syndrome, a developmental syndrome, has been variously called Carnevale, Mingarelli, Malpuech and Michels syndrome.