

ESID Online Registry: Diseases and Genes

as of 11.03.2024 - Please send ideas and questions to registry@esid.org

Main Category (IUIS Table)	Sub Category	IEI Diagnosis (PID Diagnosis)	Gene
Combined immunodeficiencies	Atypical Severe Combined Immunodeficiency (Atypical SCID)	Atypical Severe Combined Immunodeficiency (Atypical SCID)	ADA Artemis CARD11 CD3d CD3e CD3z Coronin-1A IL2RG IL21R IL7Ralpha JAK3 PNP RAG1 RAG2
	CD4-deficiency	Selective CD4 cell deficiency	CD4
	CD8-deficiency	CD8 deficiency	CD8A
	Combined immunodeficiency (CID)	Combined immunodeficiency	ADA ARPC1B Artemis CARD11 Caspase 8 CD3G CD27 CD70 Cernunnos C-REL CTLA-4 DNA-PKcs DOCK2 DOCK8 IKBKB IL21R IL2Ralpha IRF4

Combined immunodeficiencies	Combined immunodeficiency (CID)	Combined immunodeficiency	ITK LCK LIG4 LRBA MAGT1 MALT1 MSN (moesin) MST1 (STK4) NFKB1 Orai1 (TMEM142A) OX40 PGM3 PIK3CD (PI3K-delta) PIK3CG PIK3R1 PRKDC PNP RECQL4 RhoH RLTPR (CARMIL2) RAG1 RAG2 RELB RIPK1 STAT1 STAT5 STAT5a STAT5b STIM1 TACI TBX1 TRAC TTC7A UNC119D ZAP70
		Activated PI3K-delta syndrome (APDS)	PIK3CD (PI3K-delta) PIK3R1
		DIAPH1 deficiency	DIAPH1
		EXTL3 deficiency	EXTL3
		ICOS deficiency	ICOS
		NCKAPL1 deficiency	NCKAPL1
		PI4KA deficiency	PI4KA
		Polymerase d 2 deficiency	POLD2
		SASH3 deficiency	SASH3
	HLA class I deficiency	HLA class I deficiency	TAP1 TAP2 TAPBP
	HLA class II deficiency	HLA class II deficiency	MHC2TA RFX5 RFXANK RFXAP
	Kabuki syndrome 1	Kabuki syndrome 1	KMT2D (MLL2)
	Kabuki syndrome 2	Kabuki syndrome 2	KDM6A
	NUDE/SCID	Winged-helix nude deficiency (FOXN1)	FOXN1
	Omenn syndrome	Omenn Syndrome	ADA Artemis CD45 Coronin-1A Del 22q11.2 IL2RG IL21R IL2Ralpha

Combined immunodeficiencies	Omenn syndrome	Omenn Syndrome	<i>IL7Ralpha</i> <i>JAK3</i> <i>RAG1</i> <i>RAG2</i> <i>RMRP</i> <i>TBX1</i>
	Otofaciocervical syndrome type 2	PAX1 deficiency	<i>PAX1</i>
	Severe combined immunodeficiency (SCID)	Severe combined immunodeficiency (SCID)	<i>ADA</i> <i>Artemis</i> <i>CD3d</i> <i>CD3e</i> <i>CD3z</i> <i>CD45</i> <i>Coronin-1A</i> <i>Del 22q11.2</i> <i>DNA-PKcs</i> <i>IL2RG</i> <i>IL21R</i> <i>IL7Ralpha</i> <i>JAK3</i> <i>LIG4</i> <i>NHEJ1</i> <i>PGM3</i> <i>RAG1</i> <i>RAG2</i> <i>TBX1</i> <i>ZAP70</i>
		Cernunnos/XLF deficiency	<i>NHEJ1</i>
		ITPKB deficiency	<i>ITPKB</i>
		Reticular Dysgenesis - AK2 (SCID)	<i>AK2</i>
		Reticular Dysgenesis - UNK (SCID)	none
	T-B- SCID	DNA ligase IV deficiency	<i>LIG4</i>
	Combined immunodeficiency with associated or syndromic features	AIOLOS deficiency	<i>IKZF3</i>
		Chromosome 11q deletion syndrome (Jacobsen syndrome)	11q23del
Primary antibody deficiencies		BCL11B deficiency	<i>BCL11B</i>
		EPG5 deficiency	<i>EPG5</i>
		FNIP1- associated IEI	<i>FNIP1</i>
		Immunodeficiency, developmental delay and hypohomocysteinemia, IMDDHH	<i>NFE2L2</i>
		MIRAGE syndrome	<i>SAMD9 (GOF)</i>
		HOIP deficiency	<i>RNF31</i>

Predominantly antibody disorders (III)	Agammaglobulinemias	Agammaglobulinemia	BLNK/SLP65 BTK CD79A CD79B IGHM IGLL1 PIK3R1 TCF3
	Class switch recombination defects (CSR) / HIGM syndromes	CSR defects and Hyper IgM (HIGM) syndromes	AID CD40 (TNFRSF5) CD40L (CD154) PMS2 UNG
Hypogammaglobulinemias		ADA2 Deficiency (DADA2)	CECR1
		ARHGEF1-deficiency	ARHGEF1
		Common variable immunodeficiency disorders (CVID)	ATP6AP1 BACH2 BAFFR CD19 CD20 CD21 CD81 CTLA-4 ICOS IKZF1 IRF2BP2 LRBA MOGS NFKB1 NFKB2 PTEN TACI TRNT1 TTC37 TWEAK
		Deficiency of specific IgG (Specific antibody deficiency - SPAD)	none NFKBIA
		IgA with IgG subclass deficiency	TACI
		Immunoglobulin chain deficiencies	Heavy chain Kappa light chain Lambda light chain
		Isolated IgG subclass deficiency	BAFFR CD19 CD21 CD81 ICOS TACI
		Other immunoglobulin gene deletions	none
		Selective IgA deficiency	TACI
		Selective IgM deficiency	none
		Thymoma with immunodeficiency	none
		Transcobalamin II deficiency	Transcobalamine II
		Transient hypogammaglobulinemia of infancy	none
		NFKB1 deficiency	NFKB1
		TACI deficiency	TNFRSF13B
		CARD11 GOF	CARD11
		Unclassified antibody deficiency	none

Diseases of immune dysregulation (IV)	Autoimmune lymphoproliferative syndrome (ALPS)	Autoimmune lymphoproliferative syndrome (ALPS)	Caspase 10 (ALPS IIA) FAS (TNFRSF6) germline CD95 (germline – ALPS IA) FAS (TNFRSF6) somatic CD95 (somatic – ALPS Ia) FAS (TNFRSF6) germline FAS (TNFRSF6) FASL biallelic FASL (CD178) (ALPS IB) FADD (combined germline and somatic)
		ALPS-like disease	LRBA FADD
		Ras associated lymphoproliferative disease (RALD)	K-Ras N-Ras
	Autoimmunity with or without lymphoproliferation	JAK1 GOF Tripeptidyl-Peptidase II deficiency	JAK1 GOF TPP2
	Autoimmune polyendocrinopathy candidiasis ectodermal dystrophy (APECED)	Autoimmune polyendocrinopathy candidiasis ectodermal dystrophy (APECED)	AIRE ITCH
	Early-onset inflammatory bowel disease	Early-onset inflammatory bowel disease	ILR10 ILR10 receptor alpha chain ILR10 receptor beta chain
		IBD-like	LRBA
	Hemophagocytic Lymphohistiocytosis (HLH)	CD27 deficiency CD70-deficiency Chediak Higashi syndrome Familial hemophagocytic lymphohistiocytosis syndromes (FHLH)	CD27 CD70 LYST PRF1 STX11 STXBP2 (Munc 18-2) UNC13D
		Griselli syndrome type 2 Hermansky-Pudlak syndrome ITK deficiency (HLH phenotype) X-linked lymphoproliferative syndrome (XLP)	RAB27A AP3B1 ITK BIRC4/XIAP (XLP2) SH2D1A (XLP1)
	IPEx	FOXP3 deficiency (IPEx) Interleukin 2 receptor alpha deficiency (CD25) (IPEx phenotype) IPEx-like disease	FOXP3 IL2Ralpha LRBA
Type 1 interferonopathies		Type 1 interferonopathies	ACP5 ADAR1 RNASEH2A RNASEH2B RNASEH2C SAMHD1 TREX1 STING (TMEM173)
	Early-onset multi-organ autoimmune disease	Early-onset multi-organ autoimmune disease	CTLA-4 SOCS1 STAT3 GOF
		Infancy onset STING-associated vasculopathy	STING (TMEM173)
	Regulatory T-cell defects	LRBA-deficiency	LRBA
		DEF6 deficiency	DEF6
Susceptibility to EBV and lymphoproliferative conditions		BACH2 deficiency	BACH2
		CTPS1 deficiency	CTPS1
		PRKCD deficiency	PRKCD
		CD137 deficiency (41BB)	TNFRSF9
Unclassified disorders of immune dysregulation		Unclassified disorders of immune dysregulation	none

Other well defined PIDs	Asplenia syndrome	Asplenia syndrome (Ivemark syndrome)	none
		Isolated congenital asplenia	RPSA
	Cartilage hair hypoplasia	Cartilage hair hypoplasia	RMRP
	CHARGE syndrome	CHARGE syndrome	CHARGE-CHD7
			SEMA3E
	Congenital sideroblastic anemia with immunodeficiency, fevers, and developmental delay (SIFD)	Congenital sideroblastic anemia with immunodeficiency, fevers, and developmental delay (SIFD)	TRNT1
	Defects of Vitamin B12 and Folate metabolism	Defects of Vitamin B12 and Folate metabolism	MTHFD1
			SLC46A1
			TCN2
	DiGeorge syndrome	DiGeorge syndrome	Del 10p
			Del 22q11.2
			TBX1
	DNA-breakage disorder	AT-like disorder	MRE11
		Ataxia telangiectasia (ATM)	ATM
		Bloom syndrome	BLM Helicase
			none
		DNA-ligase 1 ATP-dependent deficiency (LIG1)	LIG1
		Immunodeficiency centromeric instability facial anomalies syndrome (ICF)	CDCA7
			DNMT3B
			HELLS
			ZBTB24
		MCM4 deficiency	MCM4
		Nijmegen breakage syndrome (NBS1)	NBS1
		Other DNA-breakage disorder	none
		Post-Meiotic Segregation 2 (PMS2) deficiency	PMS2
		RNF168 deficiency	RNF168
		Seckel syndrome	none
	Dyskeratosis congenita	Dyskeratosis congenita	DKC1
			GAR1
			NAF1
			NHP2
			NOP10
			PARN
			RTEL1
			TERC
			TERT
			TINF2
		Hoyeraal-Hreidarsson syndrome	APOLLO (snm1b def)
			DKC1
			NHP2
			NOP10
			PARN
			RTEL1
			TERC
			TERT
			TINF2
	Fc receptor deficiencies	Fc receptor deficiencies	FCGR1A
			FCGR2A
			FCGR2B
			FCGR3A
			FCGR3B
			FCGRT
	FILS syndrome	Facial dysmorphism, immunodeficiency, livedo, and short stature (FILS syndrome)	POLE1
	FOXN1 Haploinsufficiency	FOXN1 Haploinsufficiency	FOXN1

Other well defined PIDs	GINS1 deficiency	GINS1 deficiency	GINS1
Hyper IgE syndromes	Hyper IgE syndrome (HIES)		DOCK8 IL6ST STAT3 DN Tyk2 ZNF341
IKAROS deficiency	IKAROS deficiency		IKAROS
Immunodeficiencies with multiple intestinal atresias	Immunodeficiencies with multiple intestinal atresias		TTC7A
MonoMAC (WILD)	Monocytopenia and mycobacterial infection (MonoMAC (WILD))		GATA2
MYSM1 deficiency	MYSM1 deficiency		MYSM1
Netherton syndrome	Netherton syndrome		SPINK5
Roifman syndrome	Roifman syndrome		RNU4ATAC
Schimke disease	Schimke disease		SMARCAL1
Trichohepatoenteric syndrome (Giraud syndrome)	Trichohepatoenteric syndrome		SKIV2L TTC37
Unclassified syndromic immunodeficiencies	Unclassified syndromic immunodeficiencies		none
SIFD	Congenital sideroblastic anemia with immunodeficiency, fevers, and developmental delay		TNRT1
VODI	Hepatic venoocclusive disease with immunodeficiency (VODI)		SP110
Wiskott-Aldrich syndrome (WAS)	WIP deficiency Wiskott-Aldrich syndrome (WAS) X-linked thrombocytopenia with mutations in WAS		WIPF1 WAS WAS

Phagocytic disorders (V)	Actin beta deficiency	Actin beta deficiency (ACTB)	Actin beta
	Barth syndrome	Barth syndrome	TAZ
	Chronic granulomatous disease (CGD)	Chronic granulomatous disease (CGD)	GP91-phox (CYBB) P22-phox (CYBA) CYBC1 EROS P40-phox (NCF4) P47-phox (NCF1) P67-phox (NCF2)
	Clericuzio-type poikiloderma with neutropenia syndrome	Clericuzio-type poikiloderma with neutropenia syndrome	C16orf57 USB1
	COHEN syndrome	COHEN syndrome	COH1 VPS13B
	Congenital neutropenia	Congenital neutropenia	CLPB CSF3R ELA2 G6PC3 GATA2 GFI1 HAX1 JAGN1 P14 SRP54 WAS
		VPS45 deficiency (SCN5)	VPS45
	Cyclic neutropenia	Cyclic neutropenia	ELA2
	Defects of Motility	MKL1 deficiency	MKL1
	Defects with susceptibility to mycobacterial infection (MSMD)	Defects with susceptibility to mycobacterial infection (MSMD)	GP91-phox (CYBB) IFNGR1 IFNGR2 IL12B IL12R beta-1 IL18 IL23-alpha IRAK4 IRF8 ISG15 SPPL2A STAT1 STAT5
	Glycogen storage disease type 1b (GS1b)	Glycogen storage disease type 1b (GS1b)	G6PT1
	Leukocyte adhesion deficiency (LAD)	Leukocyte adhesion deficiency (LAD)	LAD1 / ITGB2 LAD2 / FUCT1 LAD3
	Localized juvenile periodontitis	Localized juvenile periodontitis	Formyl peptide receptor
	Myeloperoxidase deficiency (MPO)	Myeloperoxidase deficiency (MPO)	MPO
	Neutrophil glucose-6-phosphate dehydrogenase	Glucose-6-phosphate dehydrogenase deficiency (G6PD)	G6PD
	Papillon-Lefevre syndrome	Papillon-Lefevre syndrome	CTSC
	PID with partial albinism	Partial albinism and immunodeficiency syndrome	none
	Pulmonary alveolar proteinosis	Pulmonary alveolar proteinosis	CSF2RA
	RAC2-GTPase defect	RAS-related C3 Bolutinin toxin substrate 2 deficiency (RAC2)	RAC2
	Shwachman-Diamond-syndrome	Shwachman-Diamond-syndrome	DNAJC21 EFL1 SBDS SRP54
	Specific granule defect	CCAAT/enhancer binding protein epsilon deficiency (CEBPE)	CEBPE
	Unclassified phagocytic disorders	Unclassified phagocytic disorders	none

Defects in innate immunity (VI)	CARD 9 deficiency	CARD9 deficiency	CARD9
	Chronic mucocutaneous candidiasis (CMC)	Chronic mucocutaneous candidiasis (CMC)	ACT1 AIRE IL-17 F IL-17 receptor alpha STAT1 STAT1 (GOF)
	DBR1 deficiency	DBR1 deficiency	DBR1
	Defects of TLR/NFkappa-B signalling	Defects of TLR/NFkappa-B signalling	IKK-gamma (NEMO, IKBKG) IRAK4 MyD88 NFKBIA TIRAP
		IRAK1 deficiency	MECP2
	Epidermolytic hyperkeratosis	Epidermolytic hyperkeratosis	TMC6 (EVER1) TMC8 (EVER2) CIB1 (EVER3)
	Herpetic encephalitis	Herpetic encephalitis (HSE)	DBR1 IRF3 TBK1 TLR3 TRAF3 TRIF UNC93B1
		TRIF deficiency	TICAM1 none
	HOIL1 deficiency	HOIL1 deficiency	HOIL1 , RBCK1
	IRF9 deficiency	IRF9 deficiency	IRF9
	NBAS deficiency	Acute liver failure due to NBAS deficiency	NBAS
	Predisposition to severe viral infection	Predisposition to severe viral infection	MCM4 STAT2
		IFNAR1 deficiency	IFNAR1
		IRF7 deficiency	IRF7
		ZNFX1 deficiency	ZNFX1
	Trypanosomiasis	Trypanosomiasis	APOL-I
	Unclassified defects in innate immunity	Unclassified defects in innate immunity	none
	Warts hypogammaglobulinemia infections and myelokathexis (WHIM)	Warts hypogammaglobulinemia infections and myelokathexis (WHIM)	CXCR4

Autoinflammatory disorders (VII)	Blau syndrome	Caspase recruitment domain-containing protein 15 deficiency (CARD15)	CARD15
	CINCA syndrome	CINCA syndrome	CIAS1 (NLRP3)
	Covid-19 hyperinflammation	Covid-19 related hyperinflammation syndrome	none
	Familial cold autoinflammatory syndrome	Familial cold autoinflammatory syndrome	CIAS1 (NLRP3)
	Familial mediterranean fever (FMF)	Familial mediterranean fever defect (FMF)	MEFV
	Familial periodic fever	Hyper IgD syndrome (MVK)	MVK
		TNF-receptor associated periodic fever syndrome (TRAPS)	TNFRSF1A
	Muckle-Wells syndrome	Muckle-Wells syndrome	CIAS1 (NLRP3)
	Other autoinflammatory diseases with known genetic defect	Other autoinflammatory diseases with known genetic defect	CAD14 CECR1 HOIL1 IL1RN IL36RN LPIN2 PLCG2 PSMB8 SH3BP2 SLC29A3 A20 deficiency ADA2 deficiency (DADA2) CDC42 deficiency
			TNFAIP3 CECR1 CDC42
	Pyogenic sterile arthritis pyoderma gangrenosum and acne	Proline/serine/threonine phosphatase-interacting protein 1 deficiency (PSTPIP1)	PSTPIP1
	Non-Inflammasome Related Conditions	COPA defect	COPA
		Tim-3 deficiency	HAVCR2
	Type 1 Interferonopathies	Aicardi-Goutieres syndrome 7 (AGS7)	IFIH1 (GOF)
	Unclassified autoinflammatory diseases	Unclassified autoinflammatory diseases	none

Complement deficiencies (VIII)	Complement deficiency	Acquired angioedema	none
		C3b inactivator deficiency	C3b inactivator
		CD59 antigen P18-20 deficiency (CD59)	CD59
		Complement component 1q deficiency	C1Q-alpha C1Q-beta C1Q-gamma Q, subunit unknown
		Complement component 1r deficiency	C1r
		Complement component 1s deficiency	C1s
		Complement component 2 deficiency	C2
		Complement component 3 deficiency	C3
		Complement component 4 deficiency	C4
		Complement component 5 deficiency	C5
		Complement component 6 deficiency	C6
		Complement component 7 deficiency	C7
		Complement component 8 deficiency	C8
		Complement component 9 deficiency	C9
		Complement factor B deficiency	Factor B
		Complement factor H deficiency	Factor H
		Decay-accelerating factor for complement deficiency (DAF CD55)	CD55
		Factor D deficiency	Factor D
		Factor I deficiency	Factor I
		Ficolin3 deficiency	FCN3
		Heditary Angioedema (C1inh)	C1 Inhibitor
		Mannan-binding lectin serine protease (MASP) deficiency	COLEC11 MASP1
		Mannan-binding lectin serine protease (MASP) deficiency	MASP2
		Membrane Attack Complex Inhibitor (CD59) deficiency	CD59
		Membrane Cofactor Protein (CD46) deficiency	CD46
		Properdin P factor complement deficiency (PFC)	Properdin
		Thrombomodulin deficiency	THBD
		Factor H Related Protein Deficiency	CFHR1 CFHR2 CFHR3 CFHR4 CFHR5
	Mannose-binding lectin (MBL)	Mannose-binding lectin deficiency (MBL)	MBL
	Unclassified complement deficiencies	Unclassified complement deficiencies	none
Bone marrow failure (IX)	Bone marrow failure	BMFS2 (Hebo deficiency)	ERCC6L2
Unclassified Immunodeficiencies	Unclassified immunodeficiencies	Unclassified immunodeficiencies	none