Clinical flow cytometry in the diagnosis of primary immunodeficiencies

Clinical Immunodiagnostic and Research Laboratory
A Jeffrey Modell Diagnostic Center for Primary Immunodeficiencies
Jeffrey Modell Foundation Mission

Vicki and Fred Modell established the Jeffrey Modell Foundation, a 501(c)(3) nonprofit organization, in 1987, in memory of their son Jeffrey, who died at the age of 15 from complications of primary immunodeficiency – a genetic condition that is chronic, serious and often fatal. JMF is a global patient organization devoted to early and precise diagnosis, meaningful treatments and ultimately, cures through clinical and basic research, physician education, patient support, advocacy and public awareness. The Foundation has developed a global network of more than 100 Jeffrey Modell Diagnostic and Research Centers worldwide – consisting of 459 expert physicians at 195 academic institutions in 58 countries and spanning 6 continents – and continues to expand globally.

The Foundation’s website, info4pi.org, offers resources to patients, families and health care providers.
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Introduction

The purpose of this booklet is to inform physicians of the commonly used flow cytometric tests used for the evaluation of primary immune deficiency disorders (PIDDs). PIDDs can present at any age and are characterized by recurrent infections, severe infections requiring hospitalization or intravenous antibiotics, and infections caused by opportunistic or unusual organisms. The ability to characterize and define these disorders has improved greatly as our understanding of human immunology has progressed.

The evaluation of primary immune deficiency disorders has benefited greatly from the use of flow cytometry. Flow cytometry utilizes antibodies or reagents that emit fluorescence to enumerate the subsets of peripheral blood leukocytes and characterize the functional capacity of these cells. By staining peripheral blood leukocytes with antibodies that are specific for defined antigens, detailed assessments of the different components of the immune system are possible. This educational pamphlet will give a brief overview of PIDDs, and the specific flow cytometric tests that can be used to diagnose these disorders.

Introduction to flow cytometry

Flow cytometry is a technique where fluorescently labeled cells flow through a cytometer a single cell at a time. The fluorescent compounds are excited with a laser, and detectors measure the light emitted from these compounds. Different fluorescent compounds emit light at different wavelengths, which allows for the discrimination of several different proteins on a cell. When multiple fluorescent compounds are used to analyze a cell population, the results are typically depicted in two-dimensional diagrams (two-parameter dot plot or one-parameter histogram).

For example, assume a heterogeneous mixture of cells is stained with fluorescently labeled antibodies specific for two proteins, then analyzed on a cytometer, with results shown below. A cell that expresses only one protein detected with an antibody (labeled with green fluorescence) will shift along the X-axis but not the Y-axis. A cell that expresses only one protein detected with an antibody (labeled with red fluorescence) will shift along the Y-axis but not the X-axis. A cell expressing both antigens will shift along both X- and Y-axes. These cells are referred to as “double positive” cells. Cells that do not express either antigen will not shift along either X- or Y-axis, which can be seen as a population in the lower left corner. These cells are referred to as “double negative” cells.
Autoimmune Lymphoproliferative Syndrome (AILYMP)

Utility
• Diagnostic screen for autoimmune lymphoproliferative syndrome

Clinical indication/general description
Autoimmune lymphoproliferative syndrome (ALPS, also known as Canale-Smith syndrome) is caused by a defect in apoptosis (programmed cell death) of lymphocytes via the FAS pathway, leading to the abnormal accumulation of lymphocytes. Patients with ALPS present clinically with lymphadenopathy, hepatosplenomegaly and autoimmunity (autoimmune cytopenias and other autoimmune disorders) and have an increased, long-term risk to develop lymphomas.

Detection methodology
Normally, less than 1% of T cells that express the T cell receptor alpha and beta chain (TCRαβ+) do not express either the CD4 or the CD8 co-receptor. These T cells are termed double negative T cells (DNT). In ALPS, the number of TCRαβ+DNT cells is increased. Additionally, the TCRαβ+DNT in ALPS express an isoform of CD45 that is usually expressed only on B cells, the B220 antigen. An increased number of B220+, TCRαβ+ DNT cells is found in all characterized forms of ALPS.

Markers
CD3/CD4/CD8/B220/CD19/CD21/CD27/HLA-DR/IgD/CD16/CD56/CD14/CD45/IgM/ TCRαβ/ TCRγδ/ CD45RA/CD45RO
**Bruton’s Tyrosine Kinase (BTK)**

**Utility**
- Diagnostic screen for X-linked agammaglobulinemia (XLA)
- Detection of carrier status in female relative of XLA
- Evaluation of hypogammaglobulinemia in male patients

**Clinical indication/general description**
X-linked agammaglobulinemia (XLA), also known as Bruton’s agammaglobulinemia, is characterized by a marked reduction or absence of peripheral blood B cells and profound hypogammaglobulinemia of all isotypes (IgG, IgA, IgM, and IgE). Patients with XLA present in early childhood with recurrent infections, especially with encapsulated bacteria, as well as chronic enteroviral infections and enteric bacterial infections. XLA is caused by mutations in the Bruton’s Tyrosine Kinase (BTK) gene, which is essential for the development of B cells. Some mutations in BTK result in a milder clinical and laboratory phenotype and are therefore described as leaky.

**Detection methodology**
X-linked agammaglobulinemia (XLA) presents with severe reduction or absence of B cells (CD19+). Therefore, BTK protein expression is determined in CD14+ monocytes since these cells also express BTK. In XLA, monocytes express either no or very low amounts of BTK protein. Women who carry the mutated allele express normal numbers of B cells that express normal levels of BTK due to non-random X inactivation. However, only 50% of monocytes express the BTK protein, and this observation can be used to determine carrier status of relatives of affected children.

**Markers**
CD3/CD4/CD8/CD19/BTK/CD16/CD56/CD14/CD45
Common variable immunodeficiency (CVID)

Utility
• Diagnostic screen for common variable immunodeficiency (CVID)
• Assess B cell response to immune therapeutics
• Assess B cell subset reconstitution post-stem cell or bone marrow transplant

Clinical indication/general description
Common variable immunodeficiency (CVID) is characterized by a low serum IgG and either a
low IgA, a low IgM or both a low IgA and IgM, along with an impaired ability to make specific
antibodies in response to immunization. CVID may occur at any age, and such patients have
recurrent respiratory tract infections with encapsulated bacteria and Mycoplasma sp. B cell
numbers may be normal or decreased, and T cell numbers may also be reduced. Patients with
CVID are at increased risk to develop granulomatous or lymphocytic interstitial lung disease
(GLILD), autoimmunity (autoimmune thrombocytopenic purpura, autoimmune hemolytic
anemia) and lymphomas.

Detection methodology
The CVID assay analyzes B cell maturation, which occurs in a sequential immunophenotypic
pattern (illustrated below). B cells that are naïve (antigen inexperienced) do not express CD27.
Upon engagement with antigen, B cells express CD27 and are termed memory B cells. Memory
B cells are further classified as unswitched memory B cells (express CD27 and IgD) and switched
memory B cells (express CD27 but do not express IgD). Patients with CVID that have a markedly
reduced number of switched memory B cells are at
increased risk to have a more severe clinical phenotype
and to develop GLILD. Low
numbers of switched memory
B cells can also be seen in
other disorders affecting B
cell maturation, such as hyper-
IgM syndrome. This assay
also defines B cell subsets as
detailed by the EUROclass
criteria, including CD38, IgM,
etc.

Markers
CD3/CD4/CD8/CD45RA/
CD45RO/CD21/CD27/HLA-
DR/IgD/CD16/CD56/CD14/
CD45/CD38/IgM/CD24
Cytokine IBD (CYTIBD)

Utility
- Diagnostic screen for inflammatory bowel disease (IBD)
- Evaluates IL-10 receptor function
- Evaluates STAT3 function (Hyper IgE)
- Determines the ability of anti-inflammatory cytokine IL-10 to suppress endotoxin-induced cytokine storm

Clinical indication/general description
The inability of IL-10 to mitigate the synthesis and secretion of the pro-inflammatory cytokine IL-6 to lipopolysaccharide (LPS)/endotoxin stimulation is an indirect indicator of defective IL-10 receptor-mediated signaling in a patient’s lymphocytes. Patients with IL-10 receptor defects do not suppress IL-6 production. Since IL-10R uses STAT3 to signal, this also tests function of STAT3 and can be used to detect hyper IgE syndrome.

Detection methodology
Peripheral blood mononuclear cells (PBMCs) are stimulated with varying concentrations of IL-10 in the presence of LPS. The cell culture supernatants are collected and stored. The cell culture supernatants are then analyzed using a flow cytometric bead array to measure IL-6 production. Patients with IL-10 receptor defects will have markedly increased levels of IL-6.
Cytotoxicity/Apoptosis (CYTAPO)

Utility
- Diagnostic screen for hemophagocytic lymphohistiocytosis
- Functional evaluation of natural killer (NK) function

Clinical indication/general description
Hemophagocytic lymphohistiocytosis (HLH) is a rare, life-threatening disorder characterized by excessive lymphocytic activation and cytokine secretion, macrophage activation, subsequent hemophagocytosis of blood cells, and organ dysfunction. This disorder is usually triggered by viral infections, and typically presents at a young age. All known genetic mutations associated with HLH occur in genes encoding proteins required to kill virally infected cells. Perforin, a pore-forming protein stored in cytotoxic granules and secreted by NK cells and cytotoxic CD8+ lymphocytes, is required to kill virally infected and malignant cells. Mutations in perforin or other granule-associated proteins cause HLH. Individuals with HLH exhibit defective natural killer (NK) cell function. In addition, defective NK cell function may be seen in patients with recurrent, severe viral infections, particularly infections with herpes viruses. In addition, CD107 (LAMP-1) is upregulated upon NK degranulation.

Detection methodology
A functional flow cytometric-based assay is used to evaluate the ability of a patient’s natural killer (NK) cells to induce apoptosis of a target cell population. Target cells (K562) are fluorescently labeled to differentiate them from a patient’s PBMCs. PBMCs containing NK cells (effectors) are cultured with targets at different effector to target cell ratios, and target cell apoptosis is measured by incorporation of the fluorescent dye 7-AAD, which binds to the DNA in apoptotic cells. The Cytotoxicity/Apoptosis assay will detect defects in NK cell function in patients with clinical symptoms of HLH and can also be used to test the function of NK cells in patients with severe or chronic viral infections.

Markers
CD8/CD56/CD16/CD107a
**Functional Asplenia/Howell-Jolly Body Detection (FAHJB)**

**Utility**
- Aid in the diagnosis of splenic dysfunction by analyzing erythrocytes to detect the presence of micronuclei (Howell-Jolly bodies)
- Patients with hyposplenism are unable to filter out micronucleated red blood cells (RBCs); thus, these cells are found in the peripheral circulation in increased numbers.

**Clinical indication/general description**
The loss of splenic function can result in life-threatening complications, such as infections. There are a number of conditions that could result in splenic dysfunction, such as heterotaxy, and it can be hard to detect splenic issues in these cases.

**Detection methodology**
Determination of functional asplenia is performed using the reagent propidium iodide (PI) to detect small fragments of DNA known as Howell-Jolly bodies (HJB) within mature erythrocytes. Normally, as red blood cells transition from the reticulocyte stage to a mature red blood cell, the remaining nuclear material is extruded from the cell. Patients with a spleen that is not functioning properly will exhibit an increase in circulating mature red blood cells staining positive for PI, indicating the presence of DNA within the cells. In this assay, we stain the cells with CD45 to exclude white blood cells, CD71 to stain immature red blood cells and CD61 to exclude platelets.

**Markers**
CD45/CD61/CD71
Hyper IgM (HIGM)

Utility
- Diagnostic screen for X-linked (CD40L) and autosomal recessive (CD40) hyper IgM syndrome
- Detection of carrier status in female relative of X-linked hyper IgM syndrome

Clinical indication/general description
Hyper IgM (HIGM) syndrome is a group of primary immunodeficiencies characterized by the absence of immunoglobulin class switching. Patients with HIGM syndrome exhibit low serum IgG, IgA, and IgE levels with normal or elevated serum IgM levels. HIGM syndrome can be caused by mutations in CD40LG (Type 1, X-linked HIGM syndrome), CD40 (Type 3, autosomal recessive HIGM syndrome), activation-induced cytidine deaminase (AID, Type 2, autosomal recessive), or uracil-DNA glycosylase (UNG, Type 5, autosomal recessive). T lymphocytes upregulate CD40L upon activation that interacts with CD40 on B cells, resulting in immunoglobulin class switching, a process that requires AID and UNG. CD40L also interacts with CD40 on monocytes, resulting in activation of cell-mediated immune responses. HIGM patients with mutations in CD40 or CD40LG are prone to a variety of bacterial and viral infections as well as opportunistic infections with Pneumocystis jirovecii or Cryptosporidium. HIGM patients, due to mutations in AID or UNG, suffer from recurrent infections but do not show susceptibility to opportunistic infections.

Detection methodology
X-linked hyper IgM (XL-HIGM, Type 1) presents with the inability of activated T helper cells (CD3+CD4+) to upregulate CD40L (CD154) surface expression. Peripheral blood cells are activated pharmacologically and the expression of CD40L is determined by flow cytometry. Female carriers show two populations: one with CD40L expression and one without expression. In cases of patient mutations in CD40L, the protein can be expressed but doesn’t bind CD40. To address this, we measure CD40L expression by antibodies against CD40L as well as CD40-Ig fusion protein. This assay also detects CD40 expression on B cells and monocytes (autosomal recessive, Type 3).
Note: This assay does not detect defects in UNG or AID.

Markers
CD3/CD4/CD8/CD154/
CD40-Fc/CD19/CD40/
CD16/CD56/CD14/CD15

![Graph showing CD40 and CD154 expression](image-url)
**LRBA**

**Utility**
- To assess for expression of LRBA by flow cytometry

**Clinical indication/general description**
LRBA is required for efficient recycling of CTLA4, and defects in LRBA can look like CTLA4 deficiency. These patients suffer from a variety of autoimmune complications, such as inflammatory bowel disease. They can also have antibody deficiencies. This assay detects intracellular LRBA and is useful when LRBA variants are detected by genetic testing.

**Detection methodology**
Intracellular protein expression of LRBA in peripheral blood lymphocytes is determined by flow cytometry. Deficient expression of LRBA in neutrophils is used as a negative control.

**Markers**
LRBA
Mendelian Susceptibility to Mycobacterial Diseases (MSMD)

Utility
- Evaluates susceptibility to intracellular bacterial pathogens such as Mycobacterium, Leishmania, Listeria species, non-pathogenic (atypical) mycobacteria (M. tuberculosis, M. africanum, M. microti, M. bovis) and certain viruses (vaccinia virus)
- Detects IFNγR1, IFNγR2, IL12Rγ1, IL12p40, STAT1 and STAT4 defects

Clinical indication/general description
Patients with IFNγR1, IFNγR2, IL12Rγ1, IL12p40, NEMO, STAT1 and STAT4 defects have been identified and present with defective monocyte/macrophage/NK cell activation and impaired mycobacterial killing due to reduced IFNγ and TNFγ production. Autosomal recessive and autosomal dominant mutations leading to complete or partial deficiencies in the cytokine receptors or signaling molecules have been reported. This assay assesses the integrity of the IFNγ signaling pathway and cell surface expression of IFNγR (CD119) on monocytes and the integrity of the IL12 signaling pathway and IL12R (CD212) on natural killer (NK) cells.

Detection methodology
This assay utilizes recombinant IFNγ to activate Signal Transducer and Activator of Transcription 1 (STAT1) in CD14+ monocytes. Recombinant IL12 is used to activate Signal Transducer and Activator of Transcription 4 (STAT4) in CD56+/CD3- NK cells. IFNγ and LPS are used to activate IL12 in CD14+ monocytes. Additionally, expression of IFNγR (CD119) and IL12R (CD212) on monocytes and NK cells, respectively, is determined to rule out abnormal receptor expression.

Markers
CD3/CD4/CD8/CD45RA/CD45RO/CD19/CD16/CD56/CD212/CD14/CD45/HLA-DR/CD119/IL-12/pSTAT1/pSTAT4
Neutrophil Phenotype and Function (NPF)*

Utility

- Functional evaluation of neutrophil oxidative burst potential
- Diagnostic screen for chronic granulomatous disease (CGD)
- Detection of carrier status in female relative of CGD patient
- Leukocyte adhesion deficiencies

Clinical indication/general description

Chronic granulomatous disease (CGD) is a group of disorders characterized by a defective oxidative Burst, resulting in an inability to generate toxic oxygen radicals (superoxide) that are required to kill bacteria. Patients affected by this disorder present with recurrent bacterial infections or abscesses, particularly of the skin, subcutaneous areas or regional lymph nodes. In CGD, microbial killing is defective due to mutations in one of four known components of the NADPH oxidase system; one is X-linked (gp91-phox) and three are autosomal recessive (p22-phox, p47-phox, and p67-phox). Leukocyte adhesion deficiency Type I (LAD-I), also known as LFA-1 deficiency, is caused by a decreased expression of CD18. Leukocyte adhesion deficiency Type II (LAD-II) is caused by the decreased expression of CD15.

Detection methodology

This functional flow cytometric assay is used to assess the ability of neutrophils to produce an oxidative burst. Neutrophils are loaded with dihydrorhodamine (DHR) dye and then activated with phorbol-12-myristate-13 acetate (PMA). Normal activated neutrophils produce superoxides that reduce DHR, resulting in the emission of fluorescence, which is quantitated by flow cytometry. Neutrophils from patients with CGD cannot generate superoxide and, therefore, do not reduce DHR. This assay also examines the expression of CD18 and CD15 on neutrophils.

Markers

CD15/CD18/CD45

*Formerly NEUOXB
Perforin-Granzyme (PERGRA)

Utility
• Determination of intracellular perforin and granzyme B with cytolytic lymphocytes

Clinical indication/general description
Hemophagocytic lymphohistiocytosis (HLH) is a rare, life-threatening disorder characterized by excessive lymphocytic activation and cytokine secretion, macrophage activation, subsequent hemophagocytosis of blood cells, and organ dysfunction. This disorder is usually triggered by viral infections and typically presents at a young age. All known genetic mutations associated with HLH occur in genes encoding proteins in NK cells and cytotoxic lymphocytes that are required to kill virally infected cells. Perforin, a pore-forming protein stored in cytotoxic granules and secreted by NK cells and cytotoxic lymphocytes, is required to kill virally infected and malignant cells. Mutations in perforin as well as other granule-associated proteins have been shown to cause HLH. Individuals with HLH exhibit defective natural killer (NK) cell function.

Detection methodology
The assay is designed to enumerate the percentage of natural killer cells (CD56+CD16-) and T cytotoxic cells (CD56-CD8+) expressing perforin and granzyme B. This assay detects patients with HLH that lack perforin and is used to evaluate the expression of cytotoxic molecules (perforin and granzyme B) in NK cells.

Markers
CD3/CD4/CD8/
CD19/CD56/CD16/
CD14/CD45/
Granzyme B/
Perforin
**PHOX**

**Utility**
- To assess expression of the NADPH oxidase complex
- Confirmation of oxidative burst defects and genetic defects (important for the p47 subunit, which is difficult to sequence)

**Clinical indication/general description**
Assessment of expression of the NADPH oxidase subunits is used to confirm loss of expression when functional testing (DHR) is abnormal. This can also be used to assess any genetic defects in this pathway. It's important to note that the p47-PHOX subunit has a pseudogene, which can make sequencing difficult.

**Detection methodology**
Whole blood is stained intracellularly with antibodies against the subunits and staining in neutrophils assessed.

**Markers**
CD14/CD45/p40/p47/p67/p22/p90
Primary Immunodeficiency 1 (PID1)

Utility
- General evaluation for T, B and natural killer (NK) cell populations

Clinical indication/general description
PIDDs and secondary immunodeficiencies present with recurrent upper and lower respiratory tract infections (encapsulated and atypical bacteria), deep-seated infections, recurrent or deep-seated abscesses, intractable diarrhea and failure to thrive. In addition, PIDDs can also present with autoimmune manifestations and malignancies. Evaluation of patients with these manifestations includes enumeration of the different types of lymphocytes (T cells, B cells, NK cells) because an absolute lymphocyte count from a complete blood count differential can miss important deficiencies in specific subsets of lymphocytes.

The PID1 assay enumerates the numbers of helper (CD4) and cytotoxic (CD8) T cells, B cells and NK cells. Numerous immunodeficiencies associated with decreased numbers of T cells, B cells or NK cells can be detected with this assay, including DiGeorge syndrome (low T cell numbers will be detected), AIDS (low CD4 cell counts will be detected), X-linked agammaglobulinemia (low B cells will be detected) and NK cell deficiencies (low NK cell numbers will be detected).

Detection methodology
The assay is designed to enumerate the percent and absolute cell counts of T helper cells (CD3+CD4+), T cytotoxic cells (CD3+CD8+), B cells (CD19+), and natural killer cells (CD3-CD16+/CD56+).

Markers
- CD3/CD4/CD8/CD19/
- CD16/CD56/CD14/CD45
**Primary Immunodeficiency 2 (PID2)**

**Utility**
- Comprehensive initial screening for cellular and humoral inherited immune deficiency
- Confirmatory test of severe combined immunodeficiency (SCID)
- Includes panels from PID1

**Clinical indication/general description**
Evaluation of patients with suspected primary immunodeficiency includes enumeration of the different types of lymphocytes (T cells, B cells, NK cells) because an absolute lymphocyte count from a complete blood count differential can miss important deficiencies in specific subsets of lymphocytes.

The PID2 is a more extensive enumeration of the different subsets of lymphocytes. The PID2 assay also detects the increased activation of CD4 T cells or CD8 T cells that may be caused by an ongoing viral infection or immune dysregulation. Activated human T cells express the HLA class II protein (CD4+DR+ or CD8+DR+) and an isoform of the CD45 antigen known as CD45RO. Numerous immunodeficiencies associated with decreased numbers of T cells, B cells or NK cells can be detected with this assay, including DiGeorge syndrome (low T cell numbers will be detected), AIDS (low CD4 cell counts will be detected), X-linked agammaglobulinemia (low B cells will be detected) and NK cell deficiencies (low NK cell numbers will be detected). This assay also measures central and effector memory cells as well as T follicular helper cells.

Severe combined immunodeficiency (SCID) is characterized by profound impairment of both cellular and humoral immunity due to the absence or markedly diminished number of T cells and variably decreased numbers of B cells or NK cells. Most infants are detected at birth with newborn screening, but confirmatory testing is necessary and must include T, B and NK cells as well as naive and memory T cells for instances of maternal engraftment or Omen's syndrome.

**Detection methodology**
The assay is designed to enumerate the percent and absolute cell counts of T helper cells (CD3+CD4+), T cytotoxic cells (CD3+CD8+), B cells (CD19+) and natural killer cells (CD3-CD16+/CD56+). The assay enumerates the numbers of T cells that are activated (DR+CD4 or CD8 cells and CD45RO+CD4 cells). CD45RO+CD4 cells are also referred to as memory T cells. Effector and central memory T cells are reported out by expression of CD45RO+, CD62L, CD197. Tfh cells expressing CD4+ CXCR5+ are also reported.

**Markers**
Normal

CD4

HLA-DR

CD45RO

44.3%

53%

CD45RA

13.5%

CD4

CD62L

CXCR5

CCR7
**STAT Gain of Function (SGOF)**

**Utility**
- Tests the phosphorylation of Signal Transducer and Activator of Transcription 1 (STAT1) to determine if there is evidence of enhanced phosphorylation or delayed dephosphorylation
- Useful to determine if variants in STAT1 are gain of function

**Clinical indication/general description**
STAT1 is involved in mounting both innate and adaptive immune responses to viruses and bacteria. It can be activated by several ligands, including INFγ, INFγ, EGF, PDGF and IL-6. Loss-of-function mutations are associated with susceptibility to mycobacterial, salmonella and herpesvirus infections. Gain-of-function mutations in STAT1 can lead to chronic mucocutaneous candidiasis, other invasive fungal infections and autoimmune disorders such as psoriasis, SLE and alopecia.

**Detection methodology**
This assay utilizes recombinant IL-6 to activate STAT1 in CD4+ T cells. The cells are stimulated for 15 minutes, washed, rested and analyzed for pSTAT1 expression at 15, 30, 60 and 120 minutes post-stimulation to determine the levels of pSTAT1 and total STAT1 expression.

**Markers**
CD3/CD4/CD8/CD45RA/CD45RO/CD19/CD16/CD56/CD14/CD45/HLA-DR/pSTAT1/STAT1
T Cell Mitogen (TMITO)/ T Interleukin Proliferation (TINTL)

Utility
- Functional evaluation of T cell proliferation to mitogens

Clinical indication/general description
Once a defect in T cells is detected, or defective T cell function is suspected, further evaluation involves examination of T cell proliferation in response to mitogens. Activation of T cells with antibodies to the T cell receptor or with plant lectins results in the proliferation of these cells over the next 7-10 days. Diminished or absent proliferative response to T cell stimuli is consistent with a primary (SCID) or secondary immunodeficiency disease that affects T lymphocytes (cellular immunity). This assay tests the proliferative function of T cells to T cell receptor antibodies and a variety of mitogens.

Detection methodology
A functional flow cytometric-based assay provides a semi-quantitative assessment of lymphocyte proliferation in response to concanavalin A (ConA)/IL-2, phytohemagglutinin (PHA)/IL-2, phorbol esters and soluble CD3. Lymphocytes are labeled with the fluorescent dye CFSE and activated with mitogens. As lymphocytes divide, the fluorescent label is diluted in half, which can be seen on flow diagrams as peaks of decreasing fluorescence. Lymphocyte proliferative response is demonstrated by an increase in FSC/SSC and a progressive two-fold reduction in a proliferation tracking dye.

Markers
CD4/CD8

TINTL: Using the same methodology as above, this assay uses several cytokines to induce growth (IL2, IL7, IL15). This is useful when evaluating for CD25 deficiency. Since all lymphocytes require endogenous IL2 for growth in vitro, in cases of CD25 deficiency, the IL2 response will be abnormal while the IL7 and IL15 responses will be normal.
T Helper IL17 (THIL17)

Utility
• Diagnostic screen for hyper-immunoglobulin E syndrome (HIES)

Clinical indication/general description
Hyper-immunoglobulin E syndrome (HIES), also known as Job syndrome, is characterized by pulmonary infections, staphylococcal abscesses, eczema, and abnormalities of bone and connective tissue. IgE levels are typically very high. HIES can look very similar to severe eczema; thus, a clinical test to be able to differentiate these syndromes is clinically useful. The defects in hyper-IgE syndrome are caused by mutations in the transcription factor STAT3. STAT3 is required to induce T cells to produce IL-17, a cytokine that is important for the elicitation of an effective immune response to several bacteria and fungi. The THIL17 assay is a functional assay that measures the ability of T cells to make IL-17, which is defective in patients with HIES.

Detection methodology
Peripheral blood mononuclear cells (PBMCs) are activated in vitro with PMA and ionomycin to induce the expression of IL-17 in normal T helper cells (CD3+CD4+), which is measured by flow cytometry using an antibody that specifically recognizes IL-17. Simultaneously, IFN-γ is measured within the T cytotoxic cell (CD3+CD8+) as a control to ensure adequate activation of T cells. Extremely low percentage of IL-17+CD4+ T cells is associated with HIES, whereas a mildly decreased percentage of IL-17+CD4+ cells is normal or slightly reduced in eczema.

Markers
CD3/CD4/CD8/CD19/CD16/CD56/CD14/CD45/INF-γ/IL-17
Toll-Like Receptor (TLREC)

Utility
- Functional evaluation of toll-like receptors (TLRs)
- Screen for MYD88, IRAK-4, NFKB1A and XIAP

Clinical indication/general description
Toll-like receptors (TLRs) recognize a variety of molecules conserved in microorganisms that are not present in humans, such as lipopolysaccharides in bacteria and double-stranded RNA in viruses. Defects in MYD88 and IRAK-4, molecules required for TLR4 signaling, have been detected in patients with recurrent, invasive pneumococcal or staphylococcal infections. Infants and young children are particularly susceptible to infections when they have defects in TLRs since the adaptive immune system has not developed at this age to offer protection.

Defects in the X-linked inhibitor of apoptosis (XIAP) protein are associated with the development of X-linked lymphoproliferative syndrome-2 (XLP-2), hemophagocytic lymphohistiocytosis (HLH) and very early onset inflammatory bowel disease (VEO-IBD). XIAP is critical for downstream signaling of nucleotide-binding oligomerization domain-containing protein 2 (NOD2). The interaction of NOD2 with its ligand MDP can be utilized to determine if XIAP deficiency is indicated.

Detection methodology
A functional flow cytometric-based assay tests TLR4 function by assessing the ability of monocytes to produce tumor necrosis factor-alpha (TNFα). Peripheral blood is incubated with lipopolysaccharide or L18-Muramyl DiPeptide (NOD2 ligand) and TNFα production is measured. This assay detects defects in MYD88 and IRAK-4, and IκBα. Patients with a NOD2 signaling defect will demonstrate a compromised ability to induce expression of TNFα to L18-Muramyl DiPeptide. This assay also detects defects in XIAP by measuring TNF-α levels.

Markers
CD14/TNF-α
T Regulatory-FOXP3 (TREG)

Utility
- Determine CD4+CD25+FoxP3+ regulatory T cells (TREG) in the peripheral blood
- Diagnostic screen for the presence of immunodysregulation polyendocrinopathy enteropathy X-linked (IPEX) syndrome by assessing T regulatory cells
- Measures CD25 and CTLA-4 expression

Clinical indication/general description
IPEX is an X-linked recessive disorder causing widespread autoimmune manifestations. Mutations in the forkhead transcription factor FoxP3 are responsible for this disease. FoxP3 is crucial to the development, survival and effector function of regulatory T (Treg) cells, a cell population essential to immune regulation. Treg cells account for approximately 5 to 10% of peripheral blood CD4+ T cells. Treg cells also express high levels of CD25, the high affinity binding alpha subunit of the IL-2 receptor. IPEX-like diseases can also be caused by deficiencies in CD25 or STAT5b, key signal transduction subunits of the IL-2 receptor. Treg cells also express CTLA-4, a known inhibitor of immune response.

Detection methodology
Use of antibodies to CD4, CD25, CD3, CTLA-4 and FoxP3 allows for the determination of the percentage of Treg cells in peripheral blood of patients suspected of IPEX or IPEX-like diseases.

Markers
**X-Linked Lymphoproliferative Syndrome (XLP)**

**Utility**
- Evaluate the presence of SLAM-associated protein (SAP) and X-linked inhibitor of apoptosis (XIAP) in peripheral blood
- Diagnostic screen for X-linked lymphoproliferative syndrome (XLP), types 1 and 2
- Detection of carrier status in female relatives of XLP patients

**Clinical indication/general description**
XLP, also called Duncan disease, is a rare X-linked disorder affecting predominantly male patients. Mutations in SH2D1A gene encoding the signaling lymphocyte activation molecule (SLAM)-associated protein (SAP) cause XLP1. XLP type 2 (XLP2) is caused by mutations in BIRC4 gene encoding the XIAP protein.

**Detection methodology**
Intracellular protein expression of SAP and XIAP in peripheral blood lymphocytes is determined by flow cytometry. Deficient expression of SAP and XIAP in lymphocytes is associated with XLP type 1 or XLP type 2, respectively.

**Markers**
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Primary Immunodeficiency Diseases and Treatment

While identification of primary immunodeficiency diseases can be difficult, timely diagnosis and treatment prevent complications and save lives. We can help.

Clinical Immunodiagnostic and Research Laboratory (CIRL)

Our lab
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- Technical staff members are trained experts in the use of flow cytometric methodologies.
- Meets all the requirements of the Clinical Laboratory Improvement Amendments

For information
Phone: (414) 955-4165 | Fax: (414) 955-6488 | Email: cirl@mcw.edu
Medical director: Dr. James Verbsky, MD, PhD, jverbsky@mcw.edu
mcw.edu/CIRL (includes link to lab request forms)
Note: specimen requirements vary depending on test type, patient’s WBC, absolute lymphocyte count, the number of immature or abnormal cells present and the patient’s history.

Specimen delivery address
Clinical Immunodiagnostic and Research Laboratory
The Medical College of Wisconsin
MACC Fund Research Center
8701 Watertown Plank Rd., Room 5072
Milwaukee, WI 53226

Our clinic
- Evaluates children and adults and offers a full range of treatments
- Provides multidisciplinary care from experts in Allergy and Clinical Immunology, Rheumatology, Hematology, Oncology, Blood and Marrow Transplant and Infectious Disease
- Conducts research to improve diagnostic testing and care

For more information, visit childrenswi.org/pip.

Jeffery Modell Diagnostic and Research Center for Primary Immunodeficiency

James Verbsky, MD, PhD
Professor of Pediatrics (Rheumatology)
Director of Clinical Immunodiagnostic Research Laboratory
Director of Clinical and Translational Research

Our program is a nationally recognized referral center for Jeffery Modell Foundation Diagnostic and Research Centers of Excellence in the country.

Our physicians worked with the Wisconsin State Laboratory to make Wisconsin the first state to pilot a newborn screening test for severe combined immunodeficiency. The test identifies SCID early enough to help treat children impacted by this often-fatal disease.