The development of immune defense mechanisms begins early during fetal life but is not completely achieved at birth, leading to a peculiar susceptibility of the neonate to infectious diseases. The earliest hematopoietic stem cells are detectable in the yolk sac around 6 weeks of gestation, but at that time they are only able to differentiate into myelomonocytic cells. Thereafter, the hematopoiesis takes place in the liver from the 7th week of gestation until the 6th month, thereafter in the bone-marrow. A unique precursor is at the origin of all hematopoietic cell lines.

NORMAL SPECIFIC IMMUNITY DEVELOPMENT

A schematic representation of the T and B lymphocyte differentiation is shown in Figure 1.

Cellular Immunity

Lymphoid cells with the surface phenotype of pro-T (CD7+) are found in the fetal liver by 7 weeks of gestation, and the fetal thymus is colonized by 8.5 weeks of gestation. Pro-T cells (from fetal liver in early gestation, thereafter from bone-marrow) are attracted into the thymus by chemotactic agents secreted by thymic epithelial cells. Shortly after, thymocytes express the marker CD2, as well as both markers CD4 and CD8 (pre-T cells). Then, the pre-T cells acquire the specific T-cell antigen receptor (TCR) associated with the complex CD3 and express either CD4 or CD8 molecules (mature T cells). The TCR which results from stochastic gene rearrangements specifically recognizes antigenic peptides which have been processed by antigen presenting cells only if presented by self HLA molecules; the diversity of the TCR repertoire is progressively acquired by positive (leading to amplification of T-cell clones able to recognize foreign antigens) and negative (allowing the destruction of autoreactive T-cell clones) selection mechanisms occurring in the thymus by close contact of pre-T lymphocytes with dendritic and epithelial cells. Mature CD4+ T cells recognize by their TCR foreign antigenic peptides presented by HLA class II molecules, expressed on B cells and monocytes, and, by the way, are essentially helper T cells; in contrast, mature CD8+ T cells recognize foreign antigenic peptides presented by HLA class I molecules and, thus, are essentially cytotoxic suppressor T cells. CD4+ T cells represent 65% and CD8+ 35% of mature T cells. Mature T cells are able to go out of the thymus, to circulate in the blood and to reach the lymphoid organs (spleen and lymph nodes) in which the immune response occurs.

By 13 weeks of gestation, each of the major T-cell subsets are present in cord blood and normally functional for mitogenic and allogeneic-stimulation. However, fetal and neonatal T cells are virgin T cells because of the lack of prior exposure to antigen; this status explains the selective impairment of lymphokine production which likely contributes to the immune deficiency observed during the neonatal period: contrasting with a normal production of IL2, the production of other cytokines are either moderately (TNFa, GM-CSF) or markedly reduced (IL-3, IL-4, IL-5, IFNγ). Moreover, neonatal T lymphocytes are less efficient for B cell help to immunoglobulin (Ig) synthesis, likely because of an impaired expression of the T-cell activation marker CD40-ligand. During fetal life, infections can also occur when TCR repertoire is

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still limited; this situation can explain the severity of *in utero* early infections by *Toxoplasma gondii* or CMV.\(^2\)

**Humoral Immunity**

Precursors of B cells are first detected in fetal liver at 8 weeks of gestation and thereafter in bone-marrow. The maturation of pre-B cells into mature B cells occurs in these hematopoietic organs. Immature B cells expressing a surface immunoglobulin (the B Cell Receptor or BCR) of the IgM isotype are detected at 12 weeks of gestation.\(^{12}\) As for T cells, the diversity of the BCR repertoire is progressively acquired with positive and negative (deletion of auto-reactive B cells) selection mechanisms occurring in the bone-marrow. The main difference between TCR and BCR is the fact that BCR can recognize native antigens (and not processed antigenic peptides) without any interaction with HLA molecules. The mature B cell bearing on its membrane IgM+IgD leaves the bone-marrow and reaches the spleen and lymph nodes in which the immune response occurs. In germinal centers of lymphoid organs, other mechanisms take place, requiring a close relation with T cells: the immunoglobulin switching which allows a B cell to produce an immunoglobulin IgG, IgA or IgE with the same antigen specificity as the IgM previously produced and the positive selection of high affinity BCR bearing B cells.

Although specific antibody synthesis, essentially of the IgM isotype, is detectable as early as 20–24 weeks of gestation in an infected fetus, little IgM and IgA are present in the healthy neonate because of the lack of antigen exposure in normal conditions, and IgG is only from maternal origin (the transplacental transfer progressively increases from the 6th month of gestation). Immunoglobulin levels increase progressively to reach adult levels at 4 years of life. The neonate is able to produce specific antibodies to protein antigens but at a lesser degree than adults. This impaired response is likely related to the virgin status of B cells and to a defect in T-B cell cooperation. The response of infants to polysaccharidic antigens, a response which is supposed to be less T-cell dependent than...
the response to protein antigens, is severely impaired during the two first years of life.\textsuperscript{13}

**Nonspecific Immunity Development**

**Natural Killer Activity**

Cells with a Natural Killer (NK) phenotype are detected as early as 6 weeks of gestation in the fetal liver. NK activity increases progressively during fetal life; it is about 50\% of adult NK activity at birth and reaches normal values at age of 4 years.\textsuperscript{14}

**Antigen Presenting Cells**

Macrophages are detectable very early (4 weeks of gestation) in the yolk sac and found in lymphoid organs at 13 weeks. However, at birth, their number in tissues appears reduced, especially in the lungs. This can be related to an impaired chemotaxis. The production of monokines has been reported as normal (IL-1) or slightly diminished (IL-6, TNF\textalpha\ and IL-8). The processing of native antigens and presentation to T cells are also efficient.\textsuperscript{15}

**Polymorphonuclear Cells**

The most obvious defect is the diminished storage pool and the impaired delivery of polymorphonuclear cells at sites of infections, a defect which can be reported to an impaired chemotaxis. Neutrophils precursors are first detected in the yolk sac at 6 weeks of gestation, then in the liver and bone-marrow. Functional mature polymorphonuclear cells are present at 12–14 weeks of gestation but represent a very low percentage of circulating white blood cells. Their number sharply increases after birth; however, preterm or severely infected neonates often present with a severe neutropenia.\textsuperscript{16}

**Complement System**

Synthesis of the different components of complement begins at the 8th week of gestation and progressively increases until birth. At that time, levels of the different factors are around 50\% of adult values.\textsuperscript{17}

**Immune Deficiencies**

**Acquired Immunodeficiencies**

Most of the abnormalities impairing the fetal immune system development are acquired in utero. It is especially the case for infections occurring during pregnancy, such as rubella, cytomegalovirus (CMV) or human immunodeficiency virus (HIV). In these situations, T cells are defective both in their number and in their function.

Preterm neonates also present with a severe, but transitory, immune deficiency characterized by a hypogammaglobulinemia, a neutropenia and a defect of the different complement factors.

An alloimmunisation of the mother directed against paternal antigens present on fetal polymorphonuclear cells may result in a severe, but transitory, neutropenia.

Finally, a defect of humoral immunity in the mother will result in a transitory hypogammaglobulinemia in the neonate.

**Inherited Immune Deficiencies**

**Specific Immunity Deficiencies**

Numerous blockades can occur on the maturation pathway of T and/or B lymphocytes resulting in different inherited immunodeficiencies (Figure 2):

\( a \) = A defect of the lymphoid precursor results in a complete lack of T and B lymphocytes leading to a severe combined immunodeficiency (SCID), the transmission of which is autosomal recessive. An enzymatic defect (deficiency in the adenosine deaminase, an enzyme necessary for purine metabolism, the gene of which is located on chromosome 20) is found in around 50\% of cases.\textsuperscript{18}

\( b \) = In other cases, a defect of enzymes involved in TCR or Ig gene recombination is suspected.\textsuperscript{19}

\( c \) = A selective defect of the T-cell precursors is defined by a complete absence of T lymphocytes with normal B cells. However, the B cells cannot produce antibodies without T-cell help. Thus, this immune deficiency is a SCID. One form is due to a mutation of the gene coding for a chain (the g chain) of several receptors for lymphokines involved in T-cell differentiation. This gene is located on chromosome X.\textsuperscript{20} Another form, autosomal recessive, is secondary to a defect of an enzyme (jak3) which is known to transmit intracellular signals of the g chain.\textsuperscript{21}

\( d \) = Other abnormalities can occur on the T-cell maturation pathway, including the defect of thymus. This immune deficiency, called Di George syndrome, is characterised by a complete absence of T cells (resulting in SCID) when the thymus is absent or by a partial T-cell defect in incomplete forms. The immune deficiency is associated with...
cardiopathy and hypoparathyroidism. The mode of transmission is autosomal dominant, and a microdeletion of chromosome 22 is constantly found.\textsuperscript{22} 

e, f = Other abnormalities can also impair T-cell immunity development or function, as observed in defective expression of membrane antigens, resulting in combined immunodeficiencies: indeed, the defect in the T-cell pathway induces a secondary B cell deficiency. For example, in the X-linked hyper-IgM syndrome, a defect of expression of the molecule CD40- ligand on activated T cells (\textsuperscript{=f}) results in a complete lack of immunoglobulin switching.\textsuperscript{23} 
g = The B cell maturation pathway can also be affected, leading to a humoral immune defect. The best known is the X-linked agammaglobulinemia secondary to an early blockade on B lymphocyte differentiation. This abnormality has been reported to a defect in an enzyme (\textit{hbk}) necessary for B cell development.\textsuperscript{24} 

Two genes responsible when mutated for relatively frequent and severe immune deficiencies have been recently cloned: on X chromosome, the gene responsible for the Wiskott-Aldrich syndrome which associates an immune deficiency, eczema and thrombopenia (\textsuperscript{=h})\textsuperscript{25} and on chromosome 11 the gene responsible for ataxia telangiectasia characterised by a T-cell defect and progressive neurological problems.\textsuperscript{26} 

\textbf{Nonspecific Immunity Deficiencies} 

Inherited defects can also impair nonspecific immunity: the chronic granulomatous disease is due to a genetic defect of cytochrome b involved in oxidative metabolism of polymorphonuclear cells. This disease is either X-linked or autosomal recessive, depending of the affected chain of cytochrome b.\textsuperscript{27} 

Other defects can affect the complement factors. 

The recent cloning of different genes involved in maturation or function of specific or nonspecific immune cells has greatly improved our understanding of development of the immune system in normal human fetuses and neonates. In pathological conditions, this genetic approach allows a greater accuracy in postnatal diagnosis and is peculiarly useful for prenatal diagnosis. It also opens the way for gene therapy in the near future.
REFERENCES
