Monoclonal antibodies in Pediatrics: use in prevention and treatment

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SUMMARY

An update is provided on monoclonal antibodies (MAbs): concept, production, indications for the diagnosis and treatment of neoplastic diseases and autoimmune disorders, prevention of transplant rejection, and treatment of allergic diseases, autoimmune disease and other noninflammatory disorders such as coronary disease. Mention is also made of MAb use in the prevention of respiratory syncytial virus (RSV) infection.

A more extensive account is provided of the use of MAb in B cell lymphomas (anti-CD20) and T cell leukemias (anti-IL-2 R). Likewise, mention is made of the use of MAbs in autoimmune disorders, such as anti-TNF-alfa in application to chronic arthritis, Crohn’s disease and psoriasis, anti-IFN-5 in the treatment of chronic arthritis, uveitis, systemic lupus erythematosus, and autoimmune hemolytic anemia. Anti-KT 3 MAb is used to treat acute rejection and graft versus host disease, while anti-IL-2 R alpha and anti-IL-2 R gamma are used for the prevention of acute transplant rejection. Anti-IgE MAb (omalizumab) is used to treat asthma and allergic rhinitis refractory to other treatments. Anti-L5 (mepolizumab), anti-IL-4, anti-TNF and anti-inflammatory cytokine mediator MAbs all have indications in asthma and severe allergic rhinitis, and in intense atopic dermatitis refractory to other treatments. As to the MAbs used for the prevention of RSV infection, mention is made of anti-epitope A of the F protein of the virus.


INTRODUCTION

The great advances in science have provided insight to aspects in immunity that were practically not even suspected only a few years ago. The evolution of knowledge is so fast that the developments prove difficult to assimilate, and are soon forgotten - replaced by new findings.

Immune modulation in general is used to change immune responses that prove deleterious for humans. Such negative effects are usually the result of an excessive production of anomalous antibodies and sensitized cells that cause disease of an autoimmune or allergic nature. However, in some cases it is also of interest to modulate or change a normal response such as the rejection of a transplanted organ or tissue that the host immune system recognizes as foreign, in order to ensure tolerance of the transplant. It is also possible to modulate responses to improve immune reaction against infections, or as prophylaxis in children with immune defects such as either immaturity or primary immune deficiencies.
MONOCLONAL ANTIBODIES (MAbs)

Monoclonal antibodies (MAbs) are proteins produced by hybrid cells (hybridomas) resulting from the fusion of a B lymphocyte (BL) capable of producing specific antibodies against a certain antigen, though in a limited manner, and a myeloma cell capable of unlimited production of nonspecific antibodies. The fusion between a BL and myeloma cell results in a mixture of cells from which a selection is made of only those cells that produce specific antibodies in an almost unlimited manner. These cells are called hybridomas, and produce MAbs that are all identical and specifically targeted to a concrete antigen. The rest of the cells in the mixture are destroyed.

The laboratory production of these MAbs is simple. A rodent (rat, hamster, rabbit) is immunized with an antigen, after which B lymphocytes are extracted from the spleen or a lymph node. These cells produce specific antibodies against the antigen to which the rodent was immunized. These BLs are then fused with myeloma cells, followed by the selection of those fused cells or hybridomas that form MAbs on an unlimited basis, targeted against the antigen to which the rodent was immunized. The capacity to produce MAbs is enormous, and is only dependent upon the identification and isolation of the antigenic proteins, and immunization of the animal (rodent) with the latter (fig. 1).

The MAbs thus obtained are contaminated by DNA sequences of the rodent used to generate the sensitized BLs. As a result, on administering them to humans, they are recognized as foreign and trigger the production of human anti-mouse antibodies, with the induction of hypersensitivity phenomena and annuling the planned function of the MAbs. In order to avoid this problem, genetic engineering technology has been used to modify the MAb chains. The resulting part responsible for binding to the antigen (variable region) is minimal and of rodent origin, while the constant supporting part or region is of human origin. Consequently, tolerance of these modified MAbs is much greater. These modified molecules are known as humanized MAbs, since they contain 90% of human material. These are the antibodies used in clinical practice, while non-humanized MAbs are used mainly in laboratories.

The capacity of a hybridoma to produce MAbs persists indefinitely, and they can be stored frozen and reconstituted at any time.

MAbs are used in a great variety of situations requiring sensitivity, high affinity and precision in relation to medical diagnoses and treatments, in veterinary practice, the food industry, in agriculture, etc. In medicine, MAbs are used to identify cells by detecting cell surface antigens, identify human hematopoietic cell lines, and diagnose and treat viral or bacterial infections. They are also used to diagnose and treat tumors, prepare vaccines, modulate immune hyper-responsiveness, and ensure transplant tolerance. Further applications remain to be discovered.

Some aspects of MAbs to be revised in this article refer to immune diagnosis, the diagnosis and treatment of neoplasms, the treatment of autoimmune diseases and allergic disorders, the prevention of respiratory syncytial virus (RSV) infection, and asthma in the nursing infant (table I).

MABS FOR DIAGNOSTIC PURPOSES

Identification of phenotypic markers

MAbs have been produced against all the cell surface markers and proteins, thus allowing the identification and classification of a broad range of cells, as well as the investigation of their functions. In the year 2000, a total of 247 cell antigens had been produced and officially acknowledged. The first MAbs...
against cell surface determinants were referred to with the letters “CD”, followed by a number identifying basic cells of the immune system. Thus, anti-CD16 corresponds to a MAb that identifies natural killer cells (NK); anti-CD3 identifies T lymphocytes; anti-CD19-20 identifies BL; anti-CD4 identifies T helper cells; and anti-CD8 MAb identifies suppressor T cells. Binding of the specific MAb to the cell receptor is evidenced by immunofluorescent techniques, among other methods.

**Diagnosis of neoplastic diseases**

MAbs have been produced against surface antigens of tumor cells and molecules secreted by such cells, corresponding to a broad range of neoplasms. This has improved our knowledge of the biology of tumor cells, and has contributed to the development of new classification and diagnostic methods, the location of tumor cells, and the identification of specific tumor antigens that are exclusive of neoplastic cells. These tumor-associated antigens are phenotypical markers of cell function that have allowed the classification of leukemias and lymphomas.

**MAbs USED IN THERAPY**

The therapeutic application of MAbs is a developing field, though a number of promising results already have been obtained, such as the antibodies used in application to cancer treatment, allergic diseases and graft rejection in transplant patients.

**Neoplasms**

MAbs combined with antineoplastic drugs are specifically targeted to tumor cell receptors, thus avoiding damage to the healthy cells. MAbs also generate biological responses in the immune system; as a result, and in addition to direct cytotoxic action, they can induce antitumor responses via indirect mechanisms.

Examples of MAbs applied to neoplastic disease include the following: anti-CD20 (BL) (rituximab) is used to treat non-Hodgkin lymphomas and chronic lymphocytic leukemia. Anti-CD38 MAb is bound to a toxin in the treatment of multiple myeloma. Anti-IL-2-alfa MAb, bound to diphtheria toxin at low doses, is used to treat T cell leukemias and T cell lymphomas. The binding of a MAb to a toxin that inhibits tumor protein synthesis yields a so-called immunotoxin.

<table>
<thead>
<tr>
<th>Specific MAb</th>
<th>Type of MAb</th>
<th>Neoplasm</th>
</tr>
</thead>
<tbody>
<tr>
<td>RIL2 alfa</td>
<td>H murine</td>
<td>T lymphoma</td>
</tr>
<tr>
<td>CD33</td>
<td>H murine</td>
<td>AM leukemia</td>
</tr>
<tr>
<td>Her-2/Neu</td>
<td>H murine</td>
<td>Breast cancer</td>
</tr>
<tr>
<td>CD20 (LB)</td>
<td>H murine</td>
<td>B lymphoma</td>
</tr>
<tr>
<td>CD10</td>
<td>H murine</td>
<td>B lymphoma</td>
</tr>
<tr>
<td>CEA Kar Aigi</td>
<td>H murine</td>
<td>Lung and dig ca</td>
</tr>
<tr>
<td>CD38</td>
<td>HM + gen toxins</td>
<td>Multiple myeloma</td>
</tr>
<tr>
<td>CA125</td>
<td>M urine</td>
<td>Ovarian cancer</td>
</tr>
<tr>
<td>L nodes. CD3</td>
<td>H murine</td>
<td>Melanoma</td>
</tr>
</tbody>
</table>

Anti-idiotypic MAbs have been developed and used to treat B cell lymphomas that express surface immunoglobulins with concrete idiotypes. Antitumor MAbs have also been applied to eliminate neoplastic cells from the bone marrow of cancer patients before autotransplantation. In this context, the bone marrow of the patient is extracted and treated with immunotoxins. Once cleared of neoplastic cells, the marrow is returned to the patient to reconstruct the hematopoietic system destroyed as a consequence of chemo- or radiotherapy. Table II shows some examples of MAbs used to treat cancer.

**Autoimmune diseases**

Autoimmune diseases are produced as a consequence of anomalous T lymphocyte (TL) activation. In order to block such activation, MAbs can be used in different ways. Thus, anti-CD4 MAbs inhibit T helper cell function (T4), though it is also possible to block binding between CD40 and CD40 ligand (CD-40L). In this way, anti-CD40-L MAbs prevent binding to CD40 and therefore also co-stimulation, which is greatly increased in autoimmune diseases and in graft rejection. Moreover, tumor necrosis factor (TNF) is released in the pathogenesis of autoimmune diseases, and anti-TNF MAbs are able to exert blocking action in this context.

At present, the use of MAbs in application to autoimmune diseases has become quite widespread in both adults and in children, and constitutes a useful alternative in patients refractory to other forms of treatment. These MAbs are administered via the intravenous route on a hospital basis, and tolerance is generally excellent. Table III reports the MAbs used
to treat autoimmune diseases, together with the precise indications.

This sense, experience has been gained in the treatment of acute anterior uveitis, idiopathic posterior uveitis, uveitis associated to Behçet’s disease and chronic childhood arthritis refractory to other treatment modalities, using anti-TNF-alfa MAbs. The response is rapid and satisfactory in such cases.

In refractory Crohn’s disease, anti-TNF-alfa MAbs have obtained good results, associated to methotrexate. Tolerance is generally good - headaches and an increase in infections having been reported as side effects.

**Table III**

<table>
<thead>
<tr>
<th>MAb</th>
<th>Name</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-TNF-alfa</td>
<td>Infliximab</td>
<td>Crohn’s disease</td>
</tr>
<tr>
<td></td>
<td>Etanercept</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td></td>
<td>Adalimumab</td>
<td>Psoriasis</td>
</tr>
<tr>
<td>Anti-C5</td>
<td>HS1-1-MAb</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td></td>
<td>Eculizumab</td>
<td>Lupus</td>
</tr>
<tr>
<td>Anti-CD20</td>
<td>Rituximab</td>
<td>Acute hemolytic anemia</td>
</tr>
<tr>
<td></td>
<td>(Mabthera)</td>
<td>Acute hemolytic anemia associated to SLE</td>
</tr>
<tr>
<td>Anti-CD52</td>
<td>Alentuzumab</td>
<td>Refractory acute hemolytic anemia</td>
</tr>
</tbody>
</table>

**Table IV**

<table>
<thead>
<tr>
<th>Specific MAb</th>
<th>Name</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>OKT 3</td>
<td>Muronobab (Orthoclone)</td>
<td>Treatment acute rejection GVHD</td>
</tr>
<tr>
<td>R-IL-2-alfa</td>
<td>Basilimab (Simulect)</td>
<td>Prevention acute rejection</td>
</tr>
<tr>
<td>R-IL-2-gamma</td>
<td>Daclizumab (Zenapax)</td>
<td>Prevention acute rejection</td>
</tr>
</tbody>
</table>

**Table V**

<table>
<thead>
<tr>
<th>Anti-IgE MAb. Omalizumab</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>• Humanized MAb (5% murine, 95% human)</td>
<td></td>
</tr>
<tr>
<td>• Binds to C3 epsilon domain of free IgE. Does not bind to IgE bound to cells</td>
<td></td>
</tr>
<tr>
<td>• Patients &gt; 12 years of age with asthma and/or rhinitis</td>
<td></td>
</tr>
<tr>
<td>• Dose: 0.016-0.50 mg/kg/IU/ml of IgE every 2-4 weeks</td>
<td></td>
</tr>
<tr>
<td>• Intravenous and subcutaneous routes</td>
<td></td>
</tr>
<tr>
<td>• Side effects: headache, mild urticaria</td>
<td></td>
</tr>
</tbody>
</table>

Allergic diseases produced as a result of type I hypersensitivity reactions and resistant to habitual treatment can be treated with humanized MAbs selectively targeted to the mediators of the allergic reaction: IgE, IL-5, IL-4, TNF-alfa, and others.

At present, humanized MAbs (5% murine, 95% human) targeted to IgE (omalizumab) are used in clinical practice with special indications and in patients over 12 years of age. The anti-IgE MAb mu MAb-E25, known as omalizumab (Xolair®), selectively binds to the C3 epsilon domain of free IgE, blocking binding to the high-affinity receptor. As a result, IgE does not bind to mast cells and basophils, and these cells therefore do not release their mediators. The MAb only binds to circulating IgE, not to other immunoglobulins, forming immune complexes that neither precipitate nor cause disease. The drug can be administered intravenously or via the subcutaneous route at variable doses according to the levels of IgE in serum and the weight of the patient. The dosage ranges according to different authors between 0.016-0.50 mg/kg every 2-4 weeks. These MAbs have been used in asthma and allergic rhinitis, producing a rapid decrease in serum IgE levels, in correlation with improvement of the clinical manifestations and patient quality of life. Tolerance is good - headaches and mild urticaria having been reported as side effects (table V).

**Treatment of graft rejection in transplant patients**

In the treatment of acute rejection in transplant patients, use is made of MAbs targeted to TL surface antigens, with the purpose of inhibiting their function and thus avoiding graft rejection. The most widely used antibody is OKT3 MAb, which specifically binds to TL surface antigen CD3. Complement activation and destruction of the T cell results upon binding.

Another MAb used in clinical practice is targeted to CD25, which is the alfa-subunit of the IL-2 receptor. In this context IL-2 binding to the T cells is blocked, and the latter are therefore not activated.

Both anti-CD3 and anti-CD25 are humanized murine MAbs that are widely used in clinical practice (table IV).
In relation to humanized anti-IL-5 MAb (mepolizumab), it should be pointed out that IL-5 is essential for the recruitment, differentiation and maturation of eosinophils. These cells in turn play a key role in allergic reactions, including respiratory allergy and food and skin allergies. Anti-IL-5 MAb has been used in clinical trials in asthma, rhinitis, and atopic dermatitis. The result in terms of the reduction of peripheral blood eosinophil counts proved excellent, though this reduction was not correlated to clinical improvement of the patients\(^\text{13,14}\).

Other MAbs used to treat allergic diseases are indicated in Table VI.

### Table VI

**MAbs in allergic diseases**

- Anti-IgE. Omalizumab (Xolair)
- Anti-IL-5. Mepolizumab
- Anti-TNF
- Other MA bs against other mediators (Chemokines and MoA)

**Indications:** Moderate-severe asthma  
Severe allergic rhinitis  
Severe atopic dermatitis

**Mediated by IgE and refractory to other treatments**

### Table VII

**Prevention of RSV infection with palivizumab. Indications**

- Patients under 2 years of age with chronic lung disease  
- Patients under 2 years of age with congenital heart disease and hemodynamic alterations  
- Under 12 months of age, premature infants with < 28 weeks of gestation  
- Under 6 months of age, premature infants with 29-32 weeks of gestation

With strict clinical criteria and conducted on a hospital basis, treatment with class IgG humanized MAbs (palivizumab) specifically inhibits the antigenic A epitope of RSV protein F (fusion protein). This prevents the virus from fusing with and infecting the cells for posterior intracellular replication\(^\text{15}\).

The drug is administered via the intramuscular route, though in premature infants with very low body weights the intravenous route can be used, at a dose of 15 mg/kg/month, in the months prior to the season in which the infection appears. A total of 4-5 doses are administered. This treatment does not interfere with the vaccination calendar. Tolerance and efficacy are excellent, avoiding the infection in most cases and reducing the recurrent wheezing episodes in the first year of life, as well as the number of hospital admissions (tables VII and VIII).

### Table VIII

**Prevention with palivizumab. Form of administration**

- Hospital pharmacy  
- Storage and transport 2-8 °C  
- Once reconstituted, leave to rest 20 min.; administer in 3 hours, via i.m. route  
- Dose of 15 mg/kg/month from before start of epidemic (October) until end (February). Total: 5 doses

**Non-inflammatory diseases**

Anti-CD40 ligand (CD40-L) MAbs have been used in coronary disease, with partial results. In advanced coronary disease, associated to myocardial infarction, or unstable angina, anti-platelet aggregation factor MAbs (anti-GPIIb/IIIa MAbs) have been tested with good results in terms of the prevention of myocardial infarction\(^\text{16}\).

### REFERENCES


