A general review of advances in the treatment of Primary Immunodeficiencies (PID) has been performed.

Treatment with immunoglobulins is indicated in cases of humoral immunodeficiencies and in selected cases of combined immunodeficiencies.

The use of intramuscular immunoglobulins in the treatment of PID was abandoned after obtaining the intravenous immunoglobulins, since these are much more effective and have fewer adverse effects. Now subcutaneous immunoglobulins are also available. Immunoglobulins help to keep the patients free of symptoms and infections as these substances are able to neutralise infectious agents, modulate and promote the immune response and favour phagocytosis. Adverse effects have been reported in 5-15 % of patients receiving IV Ig, and patients with deficiencies of subclasses of IgG with IgA deficiency and/or anti-IgA antibodies are at risk of severe reactions.

No severe adverse effects of subcutaneous immunoglobulins have been reported and the medication can be self-administered. The efficacy and safety of IV Ig and SC Ig are similar and SC Ig administered at home is associated with better quality of life.

Stem Cell Transplantation (SCT) in Primary Immunodeficiencies is aimed at restoring the number and/or function of lymphocytes or phagocytes. Matched, related or unrelated donors, or related haploidentical donors are selected. HLA class II mismatched unrelated donors are avoided owing to the risk of severe graft versus host disease (GVHD).

Stem cells are obtained from bone marrow, cord blood or peripheral blood. Prophylactic immunosuppression (as well as donor T lymphocyte depletion in haploidentical and unrelated donors) is performed to avoid or minimize GVHD. Less toxic "reduced intensity" protocols now exist for pre-transplantation conditioning, indicated to avoid graft rejection if there is residual T-lymphocyte immunity in the host. In the majority of Severe Combined Immunodeficiencies (SCID), SCT results in T lymphocytes graft and the antibody immunodeficiency persists in many cases. The results are better the earlier it is performed, with the absence of previous infections, and with the degree of matching. The patient must be maintained in a laminar flow room with broad anti-infectious prophylaxis and with the intravenous administration of gammaglobulin for a variable period. Many other complications can be expected.

Gene therapy. Patients with PID are ideal candidates, as they are monogenic, the haematopoietic cells are easily obtained and virus replication is easy within them. Vectors (viruses) “infect” the stem cells of the patient’s bone marrow, producing the transfection of the wild (healthy) gene in these cells. Encouraging results have been achieved in X-linked SCID as there are a number of patients who are considered “cured”, although neoplastic processes have
occurred due to the activation of proto-oncogenes close to the point of insertion of the external gene, using retroviruses as vectors; there are now trials with adenovirus, physical methods (direct injection...) and chemical methods (viral modification, artificial viruses...). Gene therapy has also been performed in patients with Chronic Granulomatous Disease and trials will improve in the future with changes in protocols used in oncology and infectious diseases.

**Key words:** Primary immunodeficiency. Severe combined immunodeficiency. Gammaglobulin therapy. Stem cell transplantation. Gene therapy.

**INTRODUCTION**

Knowledge on etiology, genetics, clinical aspects and management have deeply improved the outcome of patients affected of Primary Immunodeficiencies (PID) in last years. This article reviews the main modalities of management currently used in the field, and it is the result of an activity of the Immunology Working Group, Spanish Society of Pediatric Clinical Immunology and Allergy (SEICAP).

**IMMUNOglobulin**

In 1809, Behring and Kitasato showed the utility of immune sera for providing protection against infections; decades later, their utility in the prevention of infections such as measles, tetanus, diphtheria and hepatitis A was demonstrated. In 1952, the first treatment with human serum immunoglobulin (Ig) was performed in a case of Bruton-type Congenital Agammaglobulinemia; since then, and up to 1981, the use of intramuscular Ig became standard treatment, being replaced subsequently by intravenous Ig.

Igs are obtained by alcoholic fractionation of a pool of human sera derived from Cohn fraction II; this procedure eliminates other proteins and live viruses (hepatitis B virus, HIV, HCV), giving rise to a sterile product for intramuscular or subcutaneous injection. The preparations obtained the WHO guidelines contain thiomersal as the preservative, glycerol as the stabiliser and have a pH of 6.8, yielding a product with 95% IgG, at a concentration of 16.5% (165 mg/ml), containing all the IgG subclasses and multiple IgG allotypes (Gm and Km), with traces of IgM, IgA and other serum proteins. This preparation contains a broad spectrum of antibodies to viruses and bacteria.

**Intramuscular immunoglobulin**

In vitro, it has been demonstrated that IMIg produces aggregates of IgG with high molecular weights that activate the complement system and are responsible for the systemic reactions that are sometimes observed. The incidence is higher if the person to whom it is administered has previously received IgG or if it is accidentally administered intravenously – these preparations are therefore contraindicated by this route.

Today, in our setting, IMIg is practically never used for the treatment of the primary immunodeficiencies and its use is limited to the prevention of several infections (HAV, HBV, measles, tetanus, rabies and Central European tick-borne encephalitis).

**Intravenous immunoglobulins**

Treatment with IVIg is the most widely used for the treatment of primary and secondary immunodeficiencies (table I). It has significant advantages, including the easy administration of large doses, rapid onset of action, absence of proteolysis of the product and its administration is painless. All the available preparations approved by the FDA and EMEA have a half-life of 18 to 25 days, contain all the IgG subclasses, have minimal anti-complement activity, have a broad spectrum of antibodies and are free of hepatitis B and C viruses and HIV. HCV infection did occur in 88 of 137 patients who received an IVIg product in October 1994, 58 % of whom had a PID. This obliged the manufacturers to employ other methods to inactivate HCV and other viruses.

The recommended dose to avoid infections or hospitalisations and to improve lung function is 400 to 600 mg/kg, aiming to maintain an IgG level of...

<table>
<thead>
<tr>
<th>Immunodeficiencies that can be treated with SCIg or IVIg</th>
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<tr>
<td><strong>Antibody deficiencies</strong></td>
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<tr>
<td>X-linked agammaglobulinemia</td>
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<tr>
<td>Common variable immunodeficiency</td>
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<tr>
<td>HyperIgM syndrome</td>
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<tr>
<td>Functional antibody deficiencies</td>
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<td>IgG subclass deficiencies with or without IgA deficiency</td>
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<tr>
<td><strong>Combined deficiencies</strong></td>
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<tr>
<td>Severe combined immunodeficiency</td>
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<tr>
<td>Wiskott-Aldrich syndrome</td>
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<tr>
<td>Ataxia telangiectasia</td>
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<td>X-linked lymphoproliferative syndrome</td>
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over 500 mg/ml or 350 mg/ml over the baseline level. In patients with chronic lung disease, chronic diarrhoea or growth failure, the doses must be higher until the clinical situation is controlled.

Administration requires a venous access and an infusion over several hours (between 2 and 4). The infusion must be started at a rate of 0.5 mg/kg/minute or 0.001 ml/kg/minute, with the rate of administration being doubled every half-hour if no adverse reaction occurs, up to a maximum of 2-3 mg/kg/minute or 0.004-0.006 ml/kg/minute.

The adverse effects of the IV infusion of gammaglobulin can be due to an excessively rapid administration to patients who receive it for the first time, patients with infection at the time of the infusion, or if more than four to six weeks have passed since the previous dose was administered.

The adverse effects reported include: headache, nausea, vomiting, rigors, joint pain and/or abdominal pain. These occur in 5-15 % of patients receiving IVIG and can be minimised by pre-treatment with oral paracetamol or antihistamines. Allergic reactions such as urticaria and anaphylaxis rarely occur. Severe reactions require treatment with adrenaline, corticosteroids and antihistamines.

When a severe reaction has occurred, pre-treatment must be given with paracetamol, antihistamines and an IV infusion of hydrocortisone 6 mg/kg (with a maximum dose of 100 mg) administered one hour before the infusion of the IVIG, with the dose being repeated at four hours if the infusion of Ig has not finished.

Although rare, some late reactions have been reported with high doses of IVIG used as immune modulators. These include aseptic meningitis, cerebral thrombosis, disseminated intravascular coagulation, renal and respiratory failure and haemolytic anaemia.

IVIG is contraindicated in patients with a history of anaphylactic reactions to IVIG or other blood products. It must be administered with great caution in cases of patients with deficiencies of subclasses of IgG with IgA deficiency and/or anti-IgA antibodies, as these patients are at risk of severe reactions.

Subcutaneous immunoglobulins

SCIg, used for years in the north of Europe as prophylaxis in humoral immunodeficiencies, is an alternative to IVIG for the treatment of primary immunodeficiencies. As the technique for inserting a small perfusion needle subcutaneously is simple and there are no reports of adverse effects, the medication can be self-administered into the abdominal wall or thighs. The injections are well tolerated. The local reactions are minimal and include erythema and/or pain; systemic reactions are rare.

The monthly dose used is the same but is divided over four weeks. Initially, the concentration of the SCIg is 16 %, to be infused at a rate of 0.05 to 0.2 ml/kg/hour or 1-3 ml/hour, with the aid of a small, battery-powered perfusion pump.

The standard doses used are of 100 to 160 mg/kg/week, corresponding to a dose of 45-60 ml of a 16 % solution for a 70-kg patient. Injection sites are the four abdominal quadrants and the lateral region of the thighs or forearms; the infusion of more than 15 to 25 ml of solution is not recommended at any site and many patients therefore give themselves infusions of 20 ml per site at intervals of one hour and find this perfectly satisfactory. The change of treatment from an IVIG to SCIg must be performed with an interval of one week between the final dose of IVIG and the first dose of SCIg. Patients diagnosed for the first time and who have not previously been treated with Ig of any type can start the replacement treatment directly with SCIg with daily doses of 100 mg/kg for five consecutive days, followed by weekly maintenance doses.

In children and infants, the subcutaneous infusion of 2-3 ml per site can be performed directly, every 5 minutes, without a perfusion pump. Depending on the size of the child, one or two doses per week can be sufficient.

Comparative studies of the efficacy and safety of IVIG and SCIg for replacement therapy found no significant differences with respect to the number of infections or adverse reactions, and recent studies have shown that SCIg administered at home is associated with better quality of life than IVIG administered in the hospital; it also gives the patients and their families greater independence and greater control over aspects of treatment and of their daily life. Learning how to administer the SCIg is easy and represents an alternative to the IVIG.

This route has been used by a number of authors in patients who had previously suffered anaphylactic reactions with IM or IVIG, when venous access is difficult or when an episode of aseptic meningitis has occurred after the use of IVIG.

The Scandinavian experience has demonstrated that an SC dose of 100 mg/kg/week of Ig has fewer adverse effects (30 from 3,232 doses, 0.9 %) than IM administration (442/1,893, 23 %) or IVIG (178/387, 46 %) (10 and 11). The serum levels of IgG achieved were similar to those reached with
IVlg. A number of products have been marketed specifically for the subcutaneous route and to avoid the Ig preparations with thiomersal in order to prevent possible mercury intoxication; some authors recommend the use of 10 % IVlg, stating that this is well tolerated by the subcutaneous route.

**STEM CELL TRANSPLANTATION**

That is a procedure aimed at restoring the number or function of haematopoietic cells (Primary Immunodeficiencies and other congenital haemopathies, or after bone marrow aplasia resulting from the treatment of acquired neoplastic or autoimmune processes), of the cells of the macrophage-monocyte system (haemophagocytic syndromes) or to provide a source for the replacement of enzymes (congenital metabolic diseases)\(^{12-16}\).

It is used in those PIDs of T lymphocytes and phagocytes which would otherwise almost certainly be fatal, with high morbidity or with a negative impact on quality of life. Severe combined immunodeficiency is a true diagnostic and therapeutic emergency in paediatrics\(^{17}\).

The SCT is indicated in T lymphocyte PID (combined or isolated) and phagocyte defects (table II)\(^{15,16}\).

### Donor selection

Syngenic identical related (homozygotic twin) or allogenic transplant. Matched unrelated allogenic transplant: Partially matched transplant: haploidentical related donor\(^{19-22}\).

For SCT in PID, unmatched unrelated donors are not used due to the high risk of severe GVHD caused by this situation. The choice is based on availability, condition of the patient and urgency, as well as on the underlying disease.

### Sources of stem cells

1. Bone marrow.
2. Cord blood: this is a rich source and produces hundreds of times more CD34 + CD38- stem cells in culture than bone marrow or peripheral blood, and these cells have different immunological properties, causing less GVHD. Since 1988, more than 4000 SCT have been performed using cord blood and there are more than 150,000 cords stored in the different bone marrow banks throughout the world\(^{23}\).

<table>
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<td><strong>Indications of SCT in PID</strong></td>
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| Severe Combined Immunodeficiency (SCID): X-linked due to a defect in the cytokine receptor common gamma chain, AR due to a mutation in RAG1/2, JAK3, ADA, IL7Ralpha, ...
| Other Combined Immunodeficiencies (CID): Omenn’s Syndrome, ADA and PNP deficiency, HLA deficiency type II, Hyper-IgM due to defects in the CD40 Ligand or in the CD40
| Complex Immunodeficiencies: Wiskott-Aldrich Syndrome, Di George Syndrome
| Phagocyte disorders: Kostmann agranulocytosis, severe congenital neutropenia
| Chronic granulomatous disease, Leukocyte Adhesion Deficiency type II
| X-linked lymphoproliferative syndrome, familial haemophagocytic lymphohistiocytosis
| Chediak-Higashi Syndrome, Griscelli Syndrome
| IFN-gamma Receptor Deficiency type I
| 3. Peripheral blood (mobilisation of CD34 + stem cells to the periphery using G-CSF, with leukapheresis on days 5-7)\(^{24}\).
| Thymus transplantation (foetal or postnatal) is not exactly a stem cell transplantation. It is indicated in the Di George Syndrome\(^{25}\).

### Pretransplant preparation

Laminar flow room. Low germ diet. Central venous line. Ensure nutrition (if necessary, use Total Parenteral Nutrition or Continuous Enteral Nutrition).


### Graft-versus-host-disease (GVHD)\(^{26,27}\)

This is due to the recognition of the host tissues by the donor T lymphocytes ("reverse rejection"). If an HLA Class II mismatch exists, the disorder is usually very severe or even fatal. The minor histocompatibility antigens give rise to a greater or lesser degree of GVHD despite the matching (even if "complete") in the majority of patients transplanted for PID. It is more common and/or intense with un-
related donors due to the possible higher degree of mismatch. It may be:

Acute: in the first three months post-transplant. It affects the skin (rash), bowel (diarrhoea) and liver (dysfunction). It has four grades of severity.

Chronic: from 100 days post-transplant. This occurs in many patients who suffer moderate or severe acute GVHD. Predominance of “autoimmune” manifestations (cytopenias).

Prevention: Selection of the best-matched donor possible. Immunosuppression prior to SCT (calcineurin inhibitors: cyclosporine or mycophenolate).

Depletion of mature donor T lymphocytes: reduces the GVHD but increases rejection and infections. Produces “mixed chimerism” (cells of different origin, donor T lymphocytes and host B and myeloid cells).

Methods of depletion: destruction (monoclonal antibodies) or extraction of the T lymphocytes (separation by rosettes), extraction of CD34+ stem cells (by magnetic microbeads; positive selection).

Treatment of GVHD: Increase the immunosuppression (corticosteroids and calcineurin inhibitors,...), which also further increases the risk of infection.

Pretransplantation conditioning

Indicated when there is residual T-lymphocyte immunity (combined immunodeficiencies, phagocyte disease) to prevent graft rejection. The protocols used in SCT for PID are still not well standardised, and must be adapted in each centre to the characteristics of the individual patient and to the advances that are taking place in this field; effective “reduced intensity” regimens now exist with lower toxicity than those used previously. For this reason, it is important that SCT is performed in centres with extensive experience in these types of patients, with adherence to the current recommendations of the experts in the field. The reader is referred to the literature references.

Graft

A Chimera is a cell line that is foreign to the host and that comes from the donor. Its existence and grade can be determined by various methods (the most simple being karyotyping if the donor and host are of different sexes, and up to PCR microsatellite amplification). The degree of chimerism that is curative varies depending on the genetic disease for which SCT is performed. In the majority of the PIDs, as there is no T-cell function, ablative conditioning is not performed (or is of low intensity) and the B lymphocytes and myeloid series principally of host origin persist in co-existence with the chimera of donor T lymphocytes; for this reason, the persistence of the host B lymphocytes in many cases of SCID means that antibody immunity does not recover. In other cases, the thymic microenvironment in children with SCID is able to induce the differentiation and functional maturation of the donor stem cells into T lymphocytes that can co-operate with apparently normal B lymphocytes from the host in order to form antibodies.

Graft kinetics

The T lymphocyte chimera and its onset of function can develop between the second and fourth week after the transplant in matched cases, and after the second month with a maximum between the third and fourth month in haploidentical transplants, with functional normalisation in the fourth to seventh month. In many cases there is a non-chimeric persistence of the host B lymphocytes, though the normalisation of their function (antibody formation) often takes more than two years, if it happens at all.

Prognostic factors

Age: early transplantation (three to six months of life in SCID, under five years in Wiskott-Aldrich syndrome) is substantially better. Intrauterine SCT has been performed but the results do not appear to be better than those performed in the newborn, and the procedure is more complicated.

Degree of matching: DR (maximum importance), minor antigens.

Related-unrelated

Active infections (CMV, RSV...) prior to the SCT

Possibly, in the future, the performance of related haploidentical SCT, performed early because of availability, with the better prognosis after improvements in the techniques of T lymphocyte depletion, may represent a highly effective therapeutic option.

Post-transplantation problems


   – Immediate post-transplantation phase (up to 30 days). The neutropenia and breakdown of the mucocutaneous barrier (catheters, mucositis) favour bacterial infections (gram positive and negative).
All the cells of our body contain the complete human genome (and, hence, the defective gene in patients with genetic defects). However, not all cells are essential for performing certain functions. Thus, in the case of haematopoietic cell diseases, such as the primary immunodeficiencies (PID)\(^1\), the genetic defects in these cells must be corrected. This has made the PIDs the first diseases in which gene therapy has been used\(^2\)\(^-\)\(^4\) and the first in which the efficacy of this treatment has been demonstrated\(^2\)\(^5\).

**Methods**

In order to introduce a healthy gene into the cells that we wish to “correct”, vehicles or vectors, usually in the form of viruses, particularly retroviruses\(^5\)\(^-\)\(^7\), are used after their manipulation to prevent their replication within the cells, though maintaining their “infective” capacity. These vectors enable the healthy gene to be incorporated into the cell genome and thus achieve the stable expression of the gene.

The transfection is performed “ex-vivo” by the incubation of the cells in culture with the viral vector containing the desired gene and their subsequent reintroduction into the patient (around 300 million corrected cells) by a simple transfusion. In some cases, the direct administration by systemic injection or injection into the affected organ is being tested\(^8\).

The insertion of the healthy gene into the cell genome “close” to other genes such as those responsible for tumours (proto-oncogenes) gave rise to an uncontrolled proliferation of these cells and the appearance of leukaemias\(^5\)\(^-\)\(^7\) in some of the cases of severe combined immunodeficiency (SCID) treated in the Necker Hospital in Paris.

These problems have not been observed in the protocols performed by other groups but, to avoid them, a number of research studies are under way on how to control and regulate the insertion of the gene, what are the most suitable types of vector, what quantities must be used, etc., which will help to perfect the techniques and obtain better results, avoiding the adverse effects\(^9\).

Other undesirable effects can include the inactivation of essential tumour suppressor genes. However, these effects would be beneficial in the case of gene therapy for tumour processes\(^1\)\(^6\), blocking activated oncogenes, replacing inactivated tumour suppressor genes or adding cell apoptosis-facilitating genes, etc.

Other methods used:

1. The use of other viruses such as adenoviruses (EBV), or adeno-associated viruses (AAV), smaller...
viruses not associated with disease, and herpes viruses. However, it is known that the majority of people have antibodies to adenoviruses, reducing the efficacy of the technique. Furthermore, a case of death has been reported in the USA, possibly due to excessive doses of the adenovirus giving rise to an uncontrolled immune reaction.

2. Physical methods: direct microinjection, electroporation, etc.

3. Chemical methods: using chemically-modified viral vectors and even by the synthesis of artificial viruses, or using synthetic substances such as liposomes, polymers, etc.

RESULTS AND DISCUSSION

There are many candidate diseases for gene therapy and a number are in clinical trials. The majority are haematological diseases, such as thalassaemia and haemophilia, metabolic diseases, etc. However, there is also great hope for the treatment of certain tumours and for the treatment of AIDS by the suppression of viral genes, for example.

Why are severe PID ideal candidates for gene therapy? PIDs are diseases caused by molecular defects, each due to a single defective gene (monogenic diseases) responsible for the functional abnormality of the haematopoietic cells. These cells are the precursors of all the cells of the immune response and, as mature cells, they emigrate to the whole lymphoid system and are able to perform the functions specific to each one of them. In addition, haematopoietic cells are easy to obtain and culture, and maintain their capacity of replication, ensuring a high level of efficacy.

The first treatments performed in humans in the 1990s, in SCID due to ADA deficiency, were not very positive: the synthesis of this enzyme decreased very rapidly and the immunological defect was not corrected. More recently, after many experimental studies, a protocol has been initiated for the treatment of severe combined immunodeficiency (SCID), specifically in the form with an X-linked inheritance (a defect in the cytokine receptor common gamma chain) for patients who do not have a matched donor for stem cell transplant (SCT). A number of treatments have now been performed in various European centres (Paris, Milan and London) with slightly varying protocols. Treatment has also been performed in cases of ADA deficiency.

The problems encountered by the Paris group, the appearance of three cases of ALL in a total of 11 children treated, were caused by the capacity of the vector to activate proto-oncogenes or inactivate tumour suppressor genes after their insertion into the genome. This led to the interruption of the treatments, though they have now been restarted after intense research into the points of insertion of these vectors, the amount of vector used, etc.

In 2006, the results of the treatments with gene therapy in two adults suffering X-linked chronic granulomatous disease (X-CGD) were published by the Institute of Biomedical Research in Frankfurt. The initial results were very good but the neutrophils did not maintain their bactericidal activity indefinitely. The results, despite the consequences described above, are very encouraging. Firstly, it has been shown that gene therapy works, that it is able to provide a permanent reconstruction of the function of lymphoid cells (presence of the full cytokine receptor), that it has “cured” the defect, and that some patients are still alive after four or five years with no problems. Despite the difficulties that have occurred, the reconstitution of normality has been demonstrated in these cases (with normalised lymphocyte function) and many studies are being performed on the different regimens and different indications in very diverse diseases, which will come to fruition in the near future.

In conclusion, therapy of PID has changed a lot in the last 15-20 years and a near normal life is obtained in most patients, if diagnosis is made correctly and therapy measures are precociously established; in addition, we can expect big advances in next years which will probably improve even more the prognosis of PID patients.

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