Hematopoietic stem cell gene therapy: selecting only the best

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Hematopoietic stem cell (HSC) gene therapy can potentially cure a variety of human hematopoietic diseases, such as sickle cell disease. Selection and expansion of gene-corrected HSCs has now been accomplished for the first time using HSC from large animals — dogs and humans — with a novel drug-resistance gene, MGMT, which is not expressed in normal HSCs (see the related articles beginning on pages 1561 and 1581). Highly efficient lentiviral transfer and expression of MGMT into relatively few HSCs led to repopulation of most of the hematopoietic compartment with gene-corrected cells following suitable drug treatment. This selection system may be useful in human clinical trials to permit gene therapy in autologous and allogeneic bone marrow transplantation settings.


Curing human diseases using gene transfer into human hematopoietic stem cells (HSCs) has long been a goal of scientists and physicians. HSCs remain among the most promising target cells for gene therapy because of their unique differentiation and expansion capabilities, which ensure that the transfer and expression of genes added to a small number of cells result in gene correction of much greater numbers of cells by self-renewal and differentiation. Targeting genes to cells more differentiated than HSCs does not result in long-term expression of the transferred genes, since these cells further differentiate and die rather than self-renew. In order to transduce HSCs it is necessary to use vectors that integrate into chromosomal DNA and are faithfully transferred to daughter cells. Currently, oncoretroviruses and lentiviruses are used as suitable vectors; adenoviruses and other transfer vectors are either carried in the cell as episomes or integrate poorly. HSCs are relatively rare populations and are often quiescent, which makes it difficult to transfer genes into these cells with high efficiency. Lentiviral vectors have an advantage over oncoretroviruses in that they target nondividing cells including quiescent HSC.

Lentiviral gene transfer

High-titer lentiviruses have been successful in curing mouse models of sickle cell disease and β-thalassemia (1, 2). Gene transfer into the HSCs of larger animals, including humans, has been much more problematic. Although human gene therapy trials have demonstrated successful transduction of human HSCs, the numbers of gene-corrected human HSCs observed following transplantation have been relatively low. To expand this population of gene-corrected cells, selectable marker genes have been used. The most successful clinical gene therapy trials to date used the human multidrug resistance 1 gene (MDR1), the P-glycoprotein product of which is required for the removal of certain drugs from cells (3, 4). P-glycoprotein expression permits the selection and expansion of the gene-corrected HSCs and their progeny by drug treatment. In these trials, selection with anthracyclines and taxanes, drugs toxic to untransduced cells without significant endogenous MDR1 activity, led to a modest selection and expansion of the MDR1-transduced human HSCs (3, 4).

A potent HSC selection system

More recently, a powerful selection system has been exploited to select for transduced HSCs using a mutant form of the enzyme O6-methylguanine-DNA methyltransferase (MGMT), encoded by the MGMT gene, the expression of which is markedly reduced in normal HSCs. MGMT confers resistance to the cytotoxic effects of the anticancer drugs the nitrosoureas (NUs). In murine studies, transduction of very few HSCs with the mutant MGMT vector results in profound selection and expansion of gene-corrected cells following exposure to NU and O6-benzylguanine (O6BG) (5–8). In this issue of the JCI, two independent studies demonstrate for the first time the power of lentiviral vectors expressing the mutant MGMT gene in HSC selection and expansion in dogs and in human HSC (9, 10). In the study by Neff et al. (9), donor dog HSCs transduced with mutant MGMT-expressing vectors were strongly selected in vivo after transfer into normal recipients in an allogeneic bone marrow setting after treatment with 1,3-bis-(2-chloroethyl)-1-nitrosourea (BCNU) and O6BG. In the second study, by Zielske et al. (10), human CD34+ cells (among which the majority of human HSCs are included) were transduced with the same mutant MGMT vector, and selection and expansion of gene-corrected human HSCs was demonstrated using NOD/SCID mice to measure engraftment and expansion of human HSCs and their progeny (Figure 1).
The MGMT cell selection system is already being tested for protection of marrow from the toxic effects of NU treatment in a human phase 1 clinical trial in patients being treated with NUs (S. Gerson, unpublished observations). One future possibility is the use of this system to permit successful human allogeneic transplantation by MGMT gene transfer into umbilical cord blood that may have insufficient HSCs to fully engraft adults. Subsequent in vivo selection and expansion of the transduced donor HSCs would be achieved using NU and O6BG administration, analogous to the experiment performed in dogs and described in this issue by Neff et al. (9). Another potential application in humans would be the addition of the MGMT gene to retroviral or lentiviral vectors containing a nonselectable gene such as the human β-globin gene in an autotransplantation setting, followed by selection for the gene-corrected cells after transplantation.

**Gene therapy today**
Despite the recent development of leukemia in two of the nine subjects enrolled in a human clinical trial of retroviral gene therapy for X-linked SCID (11), the benefits afforded to almost all patients in this trial, in addition to the lack of significant toxicity reported in other similar trials, lend weight to the general opinion that human retroviral gene therapy trials using HSCs should continue in the hope of increasing therapeutic efficacy and safety (12, 13). Both reports in this issue of the *JCI* also show that relatively low numbers of integrations of the MGMT gene result in sufficient expression of the enzyme to successfully select and expand the gene-corrected HSCs (9, 10). This is important because of the presumed potential for increased insertional mutagenesis with increased numbers of integrations. In the X-linked SCID gene therapy trial, both patients who developed leukemia demonstrated insertion of an unregulated γC receptor gene into the locus of a known proliferative gene, LMO2. In both cases, there was a 2- to 3-year interval between the gene-correction event and the development of leukemia. These circumstances suggest that the leukemic clones were produced by the rare event of gene insertion at the LMO2 locus and were selected for proliferation over time. Methods that may be employed in future studies to reduce the type of insertional mutagenesis associated with these two individuals include the use of (a) insulators, which would prevent the upstream or downstream activation of the inserted gene(s); and (b) gene- and tissue-specific promoters, such as the human β-globin gene promoter in human β-globin gene therapy trials, as alternatives to more generally expressed promoters such as the retroviral long-terminal repeats used in the X-linked SCID trial. These modifications and the use of HSC selection as described in the two papers in this issue of the *JCI* (9, 10) may permit safe and effective HSC gene therapy of hematological disorders in the near future.
Therapeutic CD154 antibody for lupus: promise for the future?

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Systemic lupus erythematosus (SLE) is a prototypical systemic autoimmune disease characterized by the production of pathogenic autoantibodies. A new study (see the related article beginning on page 1506) demonstrates that passive antibody specific for the TNF family member, CD154, ameliorates disease by reducing levels of self-reactive antibody in the serum. This study demonstrates a substantial potential for anti-CD154 antibody in the treatment of humoral autoimmunity.


The autoimmune disease systemic lupus erythematosus (SLE) is characterized by the production of high-affinity IgG antibodies against double-stranded DNA (dsDNA). Typically, these pathogenic autoantibodies are encoded by mutated Ig genes, and the frequency and pattern of mutations strongly suggest that the source of autoreactive antibodies is B cells that have participated in the germinal center (GC) reaction. Indeed, GCs appear spontaneously in many strains of lupus-prone mice (1), and extraordinary numbers of GC B cells and their progeny are present in the blood of SLE patients (2).

GCs are organized collections of antigen-activated T and B lymphocytes in secondary lymphoid tissues (3) or sites of chronic inflammation (4, 5). Although transient GC-like reactions can be elicited by thymus-independent pathways (6–8), most GC responses are thymus-dependent. GC B cells require ongoing survival and proliferation signals that depend on CD154–CD40 signaling (9). CD154, a member of the TNF family, is inductively expressed on the surface of CD4 T lymphocytes, whereas its ligand, CD40, is constitutively present on B lymphocytes (reviewed in ref. 10).

T cell–dependent GCs begin with the separate activation of T and B cells by antigen (11). In secondary lymphoid tissues, binding of antigen modifies B cell chemotaxis, resulting in migration toward the T cell zone (12), where cognate T cell–B cell interaction expands both lymphocyte populations. Soon afterward, activated T and B cells emigrate from T cell areas into the reticula of follicular dendritic cells (FDCs) that define the B cell follicle (Figure 1). The immigrant B lymphocytes proliferate in the FDC reticulum to generate nascent GCs and, in the process, acquire distinctive phenotypes, including expression of the CD69 activation antigen and several differentiation markers, including CD38 and CD27. Remarkably, human CD38+ GC B cells can also express variable levels of CD154, especially under conditions of chronic stimulation (13, 14), and appear to have the capacity for continued self-activation.

GCs become polarized into histologic dark and light zones (DZ, LZ). The DZ is proximal to the T cell area and contains rapidly dividing B cells called centroblasts that express little or no surface immunoglobulin. The more distal LZ contains the bulk of the activated FDC network, antigen-specific CD4 T cells, and nondividing B cells known as centrocytes. Centrocytes express surface immunoglobulin and are thought to be the progeny of DZ centroblasts. In turn, selected centrocytes likely reenter the DZ and regain the centroblast form.

The enzyme AID (activation-induced cytidine deaminase) drives both somatic hypermutation (SHM) and immunoglobulin class switch recombination (CSR) in GC B cells (15). SHM introduces point mutations and occasional small deletions into the