COURSE CODE: BCH407
COURSE TITLE: Genetic Engineering
NUMBER OF UNITS: 2 Units
COURSE DURATION: Two hours per week

COURSE DETAILS:

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COURSE CONTENT:


COURSE REQUIREMENTS:

READING LIST:
Recombinant DNA and Gene Cloning

Recombinant DNA is DNA that has been created artificially. DNA from two or more sources is incorporated into a single recombinant molecule.

Making Recombinant DNA (rDNA): An Overview

- Treat DNA from both sources with the same restriction endonuclease (BamHI in this case).
- BamHI cuts the same site on both molecules
  
  5' GGATCC 3'
  3' CCTAGG 5'

The ends of the cut have an overhanging piece of single-stranded DNA.
These are called "sticky ends" because they are able to base pair with any DNA molecule containing the complementary sticky end.
In this case, both DNA preparations have complementary sticky ends and thus can pair with each other when mixed.
• a DNA ligase covalently links the two into a molecule of recombinant DNA.

To be useful, the recombinant molecule must be replicated many times to provide material for analysis, sequencing, etc. Producing many identical copies of the same recombinant molecule is called cloning. Cloning can be done in vitro, by a process called the polymerase chain reaction (PCR). Here, however, we shall examine how cloning is done in vivo.

Cloning in vivo can be done in

• unicellular microbes like E. coli
• unicellular eukaryotes like yeast and
• in mammalian cells grown in tissue culture.

In every case, the recombinant DNA must be taken up by the cell in a form in which it can be replicated and expressed. This is achieved by incorporating the DNA in a vector. A number of viruses (both bacterial and of mammalian cells) can serve as vectors. But here let us examine an example of cloning using E. coli as the host and a plasmid as the vector.
Plasmids

Plasmids are molecules of DNA that are found in bacteria separate from the bacterial chromosome.

The desirable properties are:

- are small (a few thousand base pairs)
- usually carry only one or a few genes
- are circular
- have a single origin of replication

Plasmids are replicated by the same machinery that replicates the bacterial chromosome. Some plasmids are copied at about the same rate as the chromosome, so a single cell is apt to have only a few plasmids copied at a high rate and a single cell may have 50 or more of them.

Genes on plasmids with high numbers of copies are usually expressed at high levels. In nature, these genes often encode proteins (e.g., enzymes) that protect the bacterium from one or more antibiotics.

Plasmids enter the bacterial cell with relative ease. This occurs in nature and may account for the rapid spread of antibiotic resistance in
hospitals and elsewhere. Plasmids can be deliberately introduced into bacteria in the laboratory transforming the cell with the incoming genes.

PLASMID CLASSIFICATION.

The most useful classification of naturally occurring plasmids is based on the main characteristic coded by the plasmid genes. The 5 main types of plasmids according to this classification are:

1. Fertility plasmids” F”: Fertility plasmids carry only tra genes (transfer) and have no characteristic beyond the ability to promote conjugal transfer of plasmids.

2. Resistant” R” plasmids: They carry genes conferring on the host bacterium resistance to one or more antibacterial agent such as chloramphenicol, ampicillin and mercury. R plasmids are very important in clinical microbiology as they are spread through natural populations and can have profound consequence in the treatment of bacteria infections. Example RP₄.

3. Col plasmids: They code for colicins. These colicins are proteins that kill other bacteria e.g. colE₁ of E.Coli.

4. Degradative plasmids: They allow the host bacterium to metabolise unusual molecules such as Toluene and Salicylic acid e.g. TOL of Plasmodium putida.

5. Virulence plasmids: These confer pathogenicity on the host bacterium e.g. Ti plasmids of agrobacterium tumefaciens, which induce Crown Gall disease on dicotyledonous plants.

An Example:

pAMP
- 4539 base pairs
- a single replication origin
- a gene (amp\(^r\)) conferring resistance to the antibiotic ampicillin (a relative of penicillin)
- a single occurrence of the sequence

\[
5' \text{GGATCC}\ 3' \\
3' \text{CCTAGG}\ 5'
\]

that, as we saw above, is cut by the restriction enzyme \textbf{BamHI}

- a single occurrence of the sequence

\[
5' \text{AAGCTT}\ 3' \\
3' \text{TTCGAA}\ 5'
\]

that is cut by the restriction enzyme \textbf{HindIII}

Treatment of pAMP with a \textbf{mixture} of BamHI and HindIII produces:

- a fragment of \textbf{3755} base pairs carrying both the \textbf{amp}^r gene and the replication origin
- a fragment of \textbf{784} base pairs
- both fragments have sticky ends

\textbf{pKAN}

- 4207 base pairs
- a single replication origin
- a gene (kan\(^r\)) conferring resistance to the antibiotic \textbf{kanamycin}.
- a single site cut by \textbf{BamHI}
- a single site cut by \textbf{HindIII}

Treatment of pKAN with a \textbf{mixture} of BamHI and HindIII produces:
- a fragment of 2332 base pairs
- a fragment of 1875 base pairs with the kan$^r$ gene (but no origin of replication)
- both fragments have sticky ends

These fragments can be visualized by subjecting the digestion mixtures to electrophoresis in an agarose gel. Because of its negatively-charged phosphate groups, DNA migrates toward the positive electrode (anode) when a direct current is applied. The smaller the fragment, the farther it migrates in the gel.

**Ligation Possibilities**

If you remove the two restriction enzymes and provide the conditions for DNA ligase to do its work, the pieces of these plasmids can rejoin (thanks to the complementarity of their sticky ends).

Mixing the pKAN and pAMP fragments provides several (at least 10) possibilities of rejoined molecules. Some of these will not produce functional plasmids (molecules with two or with no replication origin cannot function).
One interesting possibility is the joining of

- the 3755-bp pAMP fragment (with \textit{amp}^r and a replication origin) with the
- 1875-bp pKAN fragment (with \textit{kan}^r)

Sealed with \textbf{DNA ligase}, these molecules are functioning plasmids that are capable of conferring resistance to \textit{both} ampicillin and kanamycin. They are molecules of \textbf{recombinant DNA}.

Because the replication origin, which enables the molecule to function as a plasmid, was contributed by pAMP, pAMP is called the \textbf{vector}.

**Transforming E. coli**

Treatment of E. coli with the mixture of religated molecules will produce some colonies that are able to grow in the presence of both ampicillin and kanamycin.

- A suspension of E. coli is treated with the mixture of religated DNA molecules.
- The suspension is spread on the surface of agar containing both ampicillin and kanamycin.
- The next day, a few cells — resistant to both antibiotics — will have grown into visible colonies containing billions of transformed cells.
- Each colony represents a \textbf{clone} of transformed cells.

However, E. coli can be simultaneously transformed by more than one plasmid, so we must demonstrate that the transformed cells have acquired the recombinant plasmid.

Electrophoresis of the DNA from doubly-resistant colonies (clones) tells the story.
Plasmid DNA from cells that acquired their resistance from a **recombinant plasmid** only show only the 3755-bp and 1875-bp bands (Clone 1, lane 3).

- **Clone 2** (Lane 4) was simultaneous transformed by religated pAMP and pKAN. (We cannot tell if it took up the recombinant molecule as well.)
- **Clone 3** (Lane 5) was transformed by the recombinant molecule as well as by an intact pKAN.

### Cloning other Genes

The recombinant vector described above could itself be a useful tool for cloning other genes. Let us assume that within its **kanamycin resistance gene** (kan') there is a single occurrence of the sequence

\[
5' \text{GAATTC} \quad 3'
\]

\[
3' \text{CTTAAG} \quad 5'
\]

This is cut by the restriction enzyme **EcoRI**, producing sticky ends.

If we treat any other sample of DNA, e.g., from human cells, with EcoRI, fragments with the same sticky ends will be formed. Mixed with EcoRI-treated plasmid and DNA ligase, a small number of the human molecules will become incorporated into the plasmid which can then be used to transform E. coli.

But how to detect those clones of E. coli that have been transformed by a plasmid carrying a piece of human DNA?
The key is that the EcoRI site is within the kan\textsuperscript{r} gene, so when a piece of human DNA is inserted there, the gene's function is destroyed.

All E. coli cells transformed by the vector, whether it carries human DNA or not, can grow in the presence of ampicillin. But E. coli cells transformed by a plasmid carrying human DNA will be unable to grow in the presence of kanamycin.

So,

- Spread a suspension of treated E. coli on agar containing ampicillin only
- Grow overnight
- With a sterile toothpick transfer a small amount of each colony to an identified spot on agar containing kanamycin
- (do the same with another ampicillin plate)
- Incubate overnight

All those clones that continue to grow on ampicillin but fail to grow on kanamycin (here, clones 2, 5, and 8) have been transformed with a piece of human DNA.

**Some recombinant DNA products being used in human therapy**

Using procedures like this, many human genes have been cloned in E. coli or in yeast. This has made it possible — for the first time — to
produce unlimited amounts of human proteins in vitro. Cultured cells (E. coli, yeast, mammalian cells) transformed with a human gene are being used to manufacture more than 100 products for human therapy. Some examples:

- **insulin** for diabetics
- **factor VIII** for males suffering from hemophilia A
- **factor IX** for hemophilia B
- **human growth hormone** (HGH)
- **erythropoietin** (EPO) for treating anemia
- several types of **interferons**
- several **interleukins**
- **granulocyte-macrophage colony-stimulating factor** (GM-CSF) for stimulating the bone marrow after a bone marrow transplant
- **granulocyte colony-stimulating factor** (G-CSF) for stimulating neutrophil production (e.g., after chemotherapy) and for mobilizing hematopoietic stem cells from the bone marrow into the blood.
- **tissue plasminogen activator** (TPA) for dissolving blood clots
- **adenosine deaminase** (ADA) for treating some forms of **severe combined immunodeficiency** (SCID)
- **parathyroid hormone**
- many **monoclonal antibodies**
- **hepatitis B surface antigen** (HBsAg) to vaccinate against the hepatitis B virus
- **C1 inhibitor** (C1INH) used to treat hereditary angioedema.

**The Polymerase Chain Reaction (PCR): Cloning DNA in the Test Tube**

The polymerase chain reaction is a technique for quickly "cloning" a particular piece of DNA in the test tube (rather than in living cells like E. coli). Thanks to this procedure, one can make virtually unlimited copies of a single DNA molecule even though it is initially present in a mixture containing many different DNA molecules.
The procedure

- In order to perform PCR, you must know at least a portion of the sequence of the DNA molecule that you wish to replicate.
- You must then synthesize **primers**: short oligonucleotides (containing about two dozen nucleotides) that are precisely complementary to the sequence at the 3' end of each strand of the DNA you wish to amplify.
- The DNA sample is heated to separate its strands and mixed with the primers.
- If the primers find their complementary sequences in the DNA, they bind to them.
- Synthesis begins (as always 5' -> 3') using the original strand as the template.
- The reaction mixture must contain
  - all four deoxynucleotide triphosphates (dATP, dCTP, dGTP, dTTP)
  - a DNA polymerase. It helps to use a DNA polymerase that is not denatured by the high temperature needed to separate the DNA strands.
- Polymerization continues until each newly-synthesized strand has proceeded far enough to contain the site recognized by the other primer.
- Now you have two DNA molecules identical to the original molecule.
- You take these two molecules, heat them to separate their strands, and repeat the process.
- Each cycle doubles the number of DNA molecules.
Using automated equipment, each cycle of replication can be completed in less than 5 minutes. After 30 cycles, what began as a single molecule of DNA has been amplified into more than a billion copies \((2^{30} = 1.02 \times 10^9)\).

With PCR, it is routinely possible to amplify enough DNA from a single hair follicle for DNA typing. Some workers have successfully amplified DNA from a single sperm cell. The PCR technique has even made it possible to analyze DNA from microscope slides of tissue preserved years before. However, the great sensitivity of PCR makes contamination by extraneous DNA a constant problem.
Genetic Recombination in Bacteria

Bacteria have no sexual reproduction in the sense that eukaryotes do. They have:

- no alternation of diploid and haploid generations
- no gametes
- no meiosis

But the essence of sex is genetic recombination, and bacteria do have three mechanisms to accomplish that:

- transformation
- conjugation
- transduction

Transformation

Many bacteria can acquire new genes by taking up DNA molecules (e.g., a plasmid) from their surroundings. The ability to deliberately transform the bacterium *E. coli* has made possible the cloning of many genes — including human genes — and the development of the biotechnology industry.

The first demonstration of bacterial transformation was done with *Streptococcus pneumoniae* and led to the discovery that DNA is the substance of the genes. The path leading to this epoch-making discovery began in 1928 with the work of an English bacteriologist, Fred Griffith.
The cells of *S. pneumoniae* (also known as the pneumococcus) are usually surrounded by a gummy capsule made of a polysaccharide. When grown on the surface of a solid culture medium, the capsule causes the colonies to have a glistening, smooth appearance. These cells are called "S" cells.

**Streptococcus pneumoniae** (pneumococci) growing as colonies on the surface of a culture medium. Left: The presence of a capsule around the bacterial cells gives the colonies a glistening, smooth (S) appearance. Right: Pneumococci lacking capsules have produced these rough (R) colonies. (Courtesy of Robert Austrian, *J. Exp. Med.* 98:21, 1953.)

However, after prolonged cultivation on artificial medium, some cells lose the ability to form the capsule, and the surface of their colonies is wrinkled and rough ("R"). With the loss of their capsule, the bacteria also lose their virulence. Injection of a single *S* pneumococcus into a mouse will kill the mouse in 24 hours or so. But an injection of over 100 million (100 x 10⁶) *R* cells is entirely harmless.

Encapsulated (left) and nonencapsulated (right) pneumococci. The encapsulated forms produce smooth
The reason? The capsule prevents the pneumococci from being engulfed and destroyed by scavenging cells — neutrophils and macrophages — in the body. The R forms are completely at the mercy of phagocytes.

Pneumococci also occur in over 90 different types: I, II, III and so on. The types differ in the chemistry of their polysaccharide capsule.

Unlike the occasional shift of S -> R, the type of the organism is constant. Mice injected with a few S cells of, say, Type II pneumococci, will soon have their bodies teeming with descendant cells of the same type.

However, Griffith found that when living R cells (which should have been harmless) and dead S cells (which also should have been harmless) were injected together, the mouse became ill and living S cells could be recovered from its body. Furthermore, the type of the cells recovered from the mouse's body was determined by the type of the dead S cells. In the experiment shown, injection of

- living R-I cells and
dead S-II cells produced a dying mouse with its body filled with living S-II pneumococci.

The S-II cells remained true to their new type. Something in the dead S-II cells had made a permanent change in the phenotype of the R-I cells. The process was named **transformation**.

**Oswald Avery** and his colleagues at The Rockefeller Institute in New York City eventually showed that the "something" was **DNA**.

In pursuing Griffith's discovery, they found that they could bring about the same kind of transformation **in vitro** using an **extract** of the bacterial cells.

Treating this extract with

- enzymes to destroy all polysaccharides (including the polysaccharide of the capsule)
- a lipase to destroy any lipids
- proteases to destroy all proteins
- RNase to destroy RNA

did not destroy the ability of their extracts to transform the bacteria.

But treating the extracts with **DNase** to destroy the DNA in them did abolish their transforming activity. So DNA was the only material in the dead cells capable of transforming cells from one type to another. **DNA was the substance of genes.**

View an electron micrograph showing DNA entering a pneumococcus.

Although the chemical composition of the capsule is determined by genes, the relationship is indirect. DNA is **transcribed** into RNA and RNA is **translated** into proteins. The phenotype of the pneumococci — the chemical composition of the polysaccharide capsule — is
determined by the particular enzymes (proteins) used in polysaccharide synthesis.

**Conjugation**

Some bacteria, *E. coli* is an example, can transfer a portion of their chromosome to a recipient with which they are in direct contact. As the donor replicates its chromosome, the copy is injected into the recipient. At any time that the donor and recipient become separated, the transfer of genes stops. Those genes that successfully made the trip replace their equivalents in the recipient's chromosome.

**Features:**

- Can only occur between cells of opposite mating types.
  - The donor (or "male") carries a fertility factor (F⁺).
  - The recipient ("female") does not (F⁻).
- **F**
  - is a set of genes originally acquired from a plasmid and now integrated into the bacterial chromosome;
  - establishes the origin of replication for the chromosome.
  - A portion of F is the "locomotive" that pulls the chromosome into the recipient cell.
  - The rest of it is the "caboose".
- In *E. coli*, about one gene gets across each second that the cells remain together. (So, it takes about 100 min for the entire genome (4377 genes) to make it. But,
  - the process is easily interrupted so
    - it is more likely that host genes close behind the leading F genes ("locomotive") will make it than those farther back
    - The "caboose" seldom makes it so failing to receive a complete F factor, the recipient cell continues to be "female"
- The DNA that makes it across finds the homologous region on the female chromosome and replaces it (by a double crossover).
- By deliberately separating the cells (in a kitchen blender) at different times, the order and relative spacing of the genes can be determined. In this way, a genetic map — equivalent to the
genetic maps of eukaryotes — can be made. But here the map intervals are seconds, not centimorgans (cM).

**Demonstration**

The figure shows the mechanism of conjugation in *E. coli* cells where

- The "male" lacks functional genes needed to synthesize the vitamin biotin and the amino acid methionine (Bio\(^-\), Met\(^-\)) so these must be added to its culture medium.
- The "female" has those genes (Bio\(^+\), Met\(^+\)) but has nonfunctional (mutant) genes that prevent it from being able to synthesize the amino acids threonine and leucine (Thr\(^-\), Leu\(^-\)) so these must be added to its culture medium.
- When cultured together, some female cells receive the functional Thr and Leu genes from the male donor.
- A double crossover enables them to replace the nonfunctional alleles.
- Now the cells now can grow on a "minimal" medium containing only glucose and salts.

**Transduction**

**Bacteriophages** are viruses that infect bacteria. In the process of assembling new virus particles, some host DNA may be incorporated in them.

The virion head can hold only so much DNA so these viruses

- while still able to infect new host cells
- may be unable to lyse them.

Instead the hitchhiker bacterial gene (or genes) may be inserted into the DNA of the new host, replacing those already there and giving the host an altered phenotype. This phenomenon is called **transduction**.
Significance of genetic recombination in bacteria.

Transformation, conjugation, and transduction were discovered in the laboratory. How important are these mechanisms of genetic recombination in nature? We don't really know, but

Some thoughts:

- The completion of the sequence of the entire genome of a variety of different bacteria (and archaea) suggest that genes have in the past moved from one species to another. This phenomenon is called lateral gene transfer (LGT).
- The remarkable spread of resistance to multiple antibiotics may have been aided by the transfer of resistance genes within populations and even between species.
- Many bacteria have enzymes that enable them to destroy foreign DNA that gets into their cells. It seem unlikely that these would be needed if that did not occur in nature. In any case, these restriction enzymes have provided the tools upon which the advances of molecular biology and the biotechnology industry depend.

Reductionism

The understanding of complex systems almost always has to await unraveling the details of some simpler system. You may feel that trying to find out how one type of pneumococcus could be converted into another was an exceedingly specialized and esoteric pursuit. But Avery and his coworkers realized the broader significance of what they were observing and, in due course, the rest of the scientific world did as well. By electing to work with a well-defined system: the conversion of R forms of one type into S forms of a different type, these researchers made a discovery that has revolutionized biology and medicine.

Attempting to understand the workings of complex systems by first understanding the workings of their parts is called reductionism. Some scientists (and many nonscientists) question the value of
reductionism. They favor a holistic approach emphasizing the workings of the complete system.

But the record speaks for itself. From skyscrapers to moon walks, to computer chips to the advances of modern medicine, progress comes from first understanding the properties of the parts that make up the whole.

The late George Wald, who won the 1967 Nobel Prize in Physiology or Medicine for his discoveries of the molecular basis of detecting light [Link], once worried that his work was overly specialized — studying not vision, not the eye, not the whole retina, not even their rods and cones, but just the chemical reactions of their rhodopsins. But he came to realize "it is as though this were a very narrow window through which at a distance one can see only a crack of light. As one comes closer, the view grows wider and wider, until finally through this same window one is looking at the universe. I think this is the way it always goes in science, because science is all one. It hardly matters where one enters, provided one can come closer...."

**Gene Therapy I**

Many human diseases are caused by defective genes.

A few common examples:

<table>
<thead>
<tr>
<th>Disease</th>
<th>Genetic defect</th>
</tr>
</thead>
<tbody>
<tr>
<td>hemophilia A</td>
<td>absence of clotting factor VIII</td>
</tr>
<tr>
<td>cystic fibrosis</td>
<td>defective chloride channel protein</td>
</tr>
<tr>
<td>muscular dystrophy</td>
<td>defective muscle protein (dystrophin)</td>
</tr>
<tr>
<td>sickle-cell disease</td>
<td>defective beta globin</td>
</tr>
<tr>
<td>hemophilia B</td>
<td>absence of clotting factor IX</td>
</tr>
<tr>
<td>severe combined immunodeficiency (SCID)</td>
<td>any one of several genes fail to make a protein essential for T and B cell function</td>
</tr>
</tbody>
</table>
All of these diseases are caused by a defect at a **single gene locus**. (The inheritance is recessive so both the maternal and paternal copies of the gene must be defective.) Is there any hope of introducing functioning genes into these patients to correct their disorder? Probably.

Other diseases also have a genetic basis, but it appears that several genes must act in concert to produce the disease phenotype. The prospects of gene therapy in these cases seems far more remote.

**Case study: severe combined immunodeficiency (SCID)**

SCID is a disease in which the patient has neither

- cell-mediated immune responses nor
- is able to make antibodies.

It is a disease of young children because, until recently, the absence of an immune system left them prey to infections that ultimately killed them.

About 25% of the cases of SCID are the result of the child being homozygous for a defective gene encoding the enzyme **adenosine deaminase (ADA)**. The normal catabolism of purines is deficient, and this is particularly toxic for **T cells** and B cells.

**Treatment Options:**

- Raise the child in a strictly **germfree environment**: all food, water, and air to be sterilized. David, the "bubble boy" from Houston, survived this way until he was 12 years old.
- Give the child a transplant of **bone marrow** from a normal, histocompatible, donor. Ideally, this would give the child a continuous source of ADA$^+$ T and B cells. However,
  - even though the child cannot reject the transplant (the child has no immune system), T cells in the transplant (unless the donor was an identical twin) can attack the cells of the child producing **graft-versus-host disease**.
the donor cells may be infected with a virus which could overwhelm the recipient before his or her immune system was restored. (David received a bone marrow transplant from his sister, but she, like many people, had been infected earlier with the *Epstein-Barr virus* (the cause of "mono"). The virus was still present in the cells she donated, and killed her brother.

- Give **injections of ADA** (the enzyme is currently extracted from cows). When conjugated with polyethylene glycol (PEG) to delay its breakdown in the blood, ADA-PEG injections have kept SCID patients reasonably healthy. But just like the insulin injections of a diabetic, they must be repeated at frequent intervals. So,
- what about giving the patient functioning ADA genes; that is, **gene therapy**?

**Gene Therapy: requirements**

- The gene must be identified and **cloned**. This has been done for the ADA gene.
- It must be inserted in cells that can take up long-term residence in the patient. So far, this means removing the **patient's own cells**, treating them in tissue culture, and then returning them to the patient.
- It must be inserted in the DNA so that it will be expressed adequately; that is, **transcribed** and **translated** with sufficient efficiency that worthwhile amounts of the enzyme are produced.

All these requirements seem to have been met for SCID therapy using a **retrovirus** as the gene **vector**. Retroviruses have several advantages for introducing genes into human cells.

- Their **envelope protein** enables the virus to infect human cells.
- RNA copies of the human ADA gene can be incorporated into the retroviral genome using a packaging cell.

**Packaging cells** are treated so they express:
• an RNA copy of the human ADA gene along with
  o a **packaging signal** (P) needed for the assembly of fresh virus particles
  o **inverted repeats** (“R”) at each end; to aid insertion of the DNA copies into the DNA of the target cell.
• an RNA copy of the retroviral **gag**, **pol**, and **env** genes but with no **packaging signal** (so these genes cannot be incorporated in fresh viral particles).

Treated with these two genomes, the packaging cell produces a crop of **retroviruses** with:

• the envelope protein needed to infect the **human target cells**
• an RNA copy of the human ADA gene, complete with R sequences at each end
• **reverse transcriptase**, needed to make a DNA copy of the ADA gene that can be inserted into the DNA of the target cell
• none of the genes (gag, pol, env) that would enable the virus to replicate in its new host.

Once the virus has infected the target cells, this RNA is reverse transcribed into DNA and inserted into the chromosomal DNA of the host.

**What to use for target cells?**

**T cells**
The first attempts at gene therapy for SCID children (in 1990), used their own T cells (produced following ADA-PEG therapy) as the target cells.

The T cells were:

- placed in tissue culture
- stimulated to proliferate (by treating them with the *lymphokine*, Interleukin 2 (IL-2))
- infected with the retroviral vector
- returned, in a series of treatments, to the child

The children developed improved immune function but:

- the injections had to be repeated because T cells live for only 6–12 months in the blood
- the children also continued to receive ADA-PEG so the actual benefit of the gene therapy was unclear

**Stem cells**

Blood ("hematopoietic") stem cells:

- produce (by mitosis) all the types of blood cells, including T and B lymphocytes
- produce (by mitosis) **more stem cells**, thus ensuring an inexhaustible supply

In June of 2002, a team of Italian and Israeli doctors reported on two young SCID patients that were treated with their own blood stem cells that had been transformed in vitro with a retroviral vector carrying the ADA gene. After a year, both children had fully-functioning immune systems (T, B, and NK cells) and were able to live normal lives without any need for treatment with ADA-PEG or immune globulin (IG). The doctors attribute their success to first destroying some of the bone marrow cells of their patients to "make room" for the transformed cells.
Nine years later (August 2011) these two patients are still thriving and have been joined by 28 other successfully-treated children most of whom no longer need to take ADA-PEG.

**Gene Therapy for X-linked SCID**

Gene therapy has also succeeded for 20 baby boys who suffered from another form of severe combined immunodeficiency called X-linked SCID because it is caused by a mutated X-linked gene encoding a subunit — called γc (gamma-c) — of the receptor for several interleukins, including interleukin-7 (IL-7).

IL-7 is essential for converting blood stem cells into the progenitors of T cells. [View]. Boys with X-linked SCID can make normal B cells, but because B cells need T-helper cells to function, these boys could make neither cell-mediated nor antibody-mediated immune responses and had to live in a sterile bubble before their treatment.

Their doctors

- isolated blood stem cells from the bone marrow of each infant;
- treated the cells with a retroviral vector containing the normal gene for the γc interleukin receptor subunit;
- returned the treated cells to each donor.

The results: Now after as long as 11 years, 19 of these boys

- are able to live normal lives at home instead of inside a sterile "bubble";
- have normal (with some exceptions*) numbers of T cells of both the CD4 and CD8 subsets;
- have responded to several childhood immunizations, including diphtheria, tetanus and polio by producing both T cells and antibodies specific for these agents.
- Antibody production is sufficiently good that most of the boys have no need for periodic infusions of immune globulin (IG).

* Five of the little boys developed leukemia (one has died):
• in one case caused by a proliferating clone of γδ T cells in which the vector has inserted itself in a gene (on chromosome 11) implicated in some cases of acute lymphoblastic leukemia (ALL).
• in a second case, the leukemia was of αβ T cells.

Gene Therapy for β-thalassemia

β-thalassemia is an inherited disease. The most severe cases result from mutations in both copies of the gene encoding the beta chain of hemoglobin. Many causative mutations have been identified, and most lead to a failure to make any beta chains. The resulting hemoglobin functions poorly and the victim requires frequent blood transfusions.

In the 16 September 2010 issue of Nature, Cavazzana-Calvo (and many colleagues) report a single case of successful gene therapy for this disorder. Their patient was an 18-year old male.

Their procedure:

• Harvest blood stem cells from the patient.
• Expose them to a retroviral vector that contained
  o a human gene for beta-hemoglobin complete with its promoter, enhancer, and other control elements;
  o alterations to the vector to make it safe.
• After sufficient chemotherapy to "make room" for them, the patient was injected with these cells.
The result: Almost three years later, the patient is well and no longer requires periodic blood transfusions. One-third of his hemoglobin is now manufactured by the red-cell precursors descended from the gene-altered stem cells.

**Adenovirus Vectors**

Adenoviruses are human pathogens responsible for some cases of the human "cold". Modified versions of two strains are currently being used as vectors in gene therapy trials.

Advantages:

- Unlike retroviral vectors, they do not integrate into the host genome and thus should not be able to disrupt host genes (It was such disruption that caused some X-linked SCID patients treated with a retroviral vector to develop leukemia).
- They can infect nondividing cells with high efficiency.

Disadvantages:

- They elicit a powerful immune response, both by T cells and by B cells (antibodies) so repeated doses soon lose their effectiveness.
- Many people already have antibodies against the virus from earlier "colds", and these can inactivate the vector at the outset. A recent trial of an HIV vaccine using an adenovirus as the vector was halted when it was found not only not to be effective but, in people with preexisting high levels of anti-adenovirus antibodies, may have even increased their susceptibility to HIV.