Towards the first recombinant drug

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Insulin

Ells – European Learning Laboratory for the Life Sciences
Towards the first recombinant drug

Insulin
Over the last century, enormous progress has been made in preventing, alleviating and curing diseases. These developments can be linked to progress in the fields of molecular biology and genetic engineering. Insulin is one of the best examples of how the combined knowledge and application of these scientific disciplines has lead to successful disease therapy.

The discovery and isolation of insulin in 1921, by Banting and Best, paved the way for the development of a therapy which has saved the life of many diabetes patients. Insulin was the first protein to be sequenced in 1951, marking an important milestone in molecular biology. In 1976, insulin became the first recombinant drug – obtained using newly developed genetic engineering techniques – to be used for therapy in humans. Five researchers have earned Nobel Prizes for research related to insulin.

The aim of this activity is to follow the pharmaceutical history of insulin using IT tools. After describing the characteristics of the hormone, the different types of diabetes will be explained. Historically, two different approaches have been used to treat diabetes that require insulin therapy:

- Early on insulin was extracted from pigs or bovines. We will use bioinformatics tools to compare the protein sequence of insulin from different animals, in order to establish which one is most closely related to the human isoform and, therefore, best suited for treating diabetes.
- More recently the production of recombinant human insulin has been adopted. After explaining the major genetic engineering achievements behind recombinant human insulin, a lab activity will guide the students through the most significant steps in the preparation of this drug.
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Insulin is a protein hormone. Its name is derived from the Latin “insula” (island) as it is secreted by groups of endocrine cells called the islets of Langerhans in the pancreas. In general, insulin enables glucose metabolism by activating glycolysis, it promotes the accumulation of glucose in the liver in the form of glycogen and favours the storage of fat.

1.1 Short history of the discovery of insulin

While studying the structure of the pancreas under the microscope, in 1869 the young medical student Paul Langerhans identified small areas of clear cells throughout the organ (islets of Langerhans). It was thought that they produced a secretion involved in digestion. Since then researchers have been trying to understand the role of these cells.

In 1889, Oscar Minkowski and Joseph von Mering found that if they removed the pancreas from dogs, the animals would develop diabetes. This meant that this gland and the islets of Langerhans in particular (as was established later in 1901 by Eugene Opie) were involved in regulating sugar levels in the blood.

In the following years, several attempts were made to isolate the secretion from the islets of Langerhans. But it was only in 1920 that Frederick Banting came up with a successful method after reading an article by Minkowski. He saw that the internal secretion from the islets of Langerhans could be destroyed by the exocrine secretion of the pancreas and, therefore, it was necessary to remove the latter. During the following summer while working at the University of Toronto under the supervision of John James Rickard Macleod and with the help of medical student Charles Best, he came up with the idea of blocking the pancreatic ducts in dogs by tying them up, causing death and re-absorption of the digestive cells by the immune system, but leaving the islets of Langerhans untouched. They extracted the secretion from these cells and named it “isletin” (later insulin). When administered to dogs deprived of a pancreas, the extract lowered the glucose level in their blood. In the months that followed, Banting and Best tried to improve and speed up the extraction process (it took 6 weeks to extract insulin from an animal!). They used fetal bovine pancreas, in which the digestive glands were undeveloped, and later collaborated with the biochemist James Bertram Collip to purify the extract. In 1922, clinical trials began: doses of purified insulin were successfully injected into diabetic children. In November of the same year, the pharmaceutical company Eli Lilly started to produce purified insulin.

Fig. 1.1 Left: Charles Best and Frederick Banting (source: www.lillydiabete.it). Right, top: J. J. R. Macleod (source: www.uhealth-care.com); bottom: James B. Collip (source: www.medicalhistory.uwo.ca)
In 1923, Banting and Macleod were awarded the Nobel Prize for Medicine, which they chose to share with Best and Collip. 

In 1958, the Nobel Prize for Chemistry was awarded to the British molecular biologist Frederick Sanger for determining the protein sequence of insulin. In 1969, Dorothy Crowfoot Hodgkin, already a Nobel Laureate in Chemistry, determined the spatial structure of insulin by X-ray diffraction. In 1977, Rosalyn Sussman Yalow received the Nobel Prize in Medicine for developing a Radio Immuno Assay (RIA) for insulin diagnostics.

1.2 Insulin Production

In mammals insulin is synthesized in the pancreas, which is an exocrine gland that produces digestive juices. Only 2% of its total mass has an endocrine function. This portion is represented by the islets of Langerhans, which are composed of at least four types of cells:

- β-cells (65-80%) which secrete insulin
- α-cells (15-20%) which secrete glucagon (a hormone involved in the release of glucose in the bloodstream from the stores of hepatic glycogen)
- γ-cells (3-10%) which secrete somatostatin (a hormone that inhibits the release of insulin and glucagon, as well as the synthesis of the pituitary growth hormone)
- PP cells (about 1%) which secrete the pancreatic peptide (a hormone that regulates the exocrine secretion of the pancreas).
1.3 Structure of insulin

Insulin is formed from two polypeptide chains (A- and B-chains), held together by two disulphide bridges; a third disulphide bridge is situated within the A-chain. The A-chain is composed of 21 amino acids, the B-chain of 30 amino acids.

![Fig. 1.4 Structure of insulin (source: www.chemistryexplained.com)](image)

Insulin is initially synthesized as preproinsulin, a 110 amino acid polypeptide that contains additional sequences:

- A “pre” amino-terminal sequence (signal peptide of 24 amino acids), which enables the secretion of the protein
- A central “pro” sequence (the C peptide of 35 amino acids) which determines the correct folding of the protein.

After preproinsulin is translated in the endoplasmic reticulum, an enzyme cuts off the 24 amino-terminal amino acids, leaving proinsulin, which in turn folds and allows the formation of the disulphide bonds between cysteine residues. At this stage, the protein passes into the Golgi apparatus, where the C peptide is removed, forming mature insulin, which is then stored in the Golgi vescicles.

![Fig. 1.5 Insulin post-translational modifications. Cell components and molecules are not to scale (adapted from www.en.wikipedia.org/wiki/Insulin)](image)
The **secondary structure** (Fig. 1.6) is determined by two α-helices in the A-chain, which surround a central β-sheet region. The B-chain has a major α-helix section and folds sharply around the A-chain.

The **tertiary structure** (Fig. 1.7) is stabilized by disulphide bridges. The external part of the protein is polar, while internally it is mostly hydrophobic.

As far as the **quaternary structure** is concerned (Fig. 1.8), insulin tends to form dimers in solution, owing to the formation of hydrogen bonds between the C-terminal ends of the B-chain.

In the presence of zinc ions, insulin forms hexamers (groups of 6 molecules, Fig. 1.9), resulting in a torus-like (or “doughnut”) shape. Insulin is stored in β-cells and secreted in the bloodstream as a hexamer. However, the active form is a monomer.

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**Fig. 1.6** Structure of insulin: The A-chain is in purple, the B-chain is in blue (adapted from www.ncbi.nlm.nih.gov/sites/entrez)

**Fig. 1.7** Structure of insulin: the disulphide bridges are indicated in orange (adapted from www.ncbi.nlm.nih.gov/sites/entrez)

**Fig. 1.8** Insulin dimers: the cylindrical elements indicate the alpha helices (adapted from www.ncbi.nlm.nih.gov/sites/entrez)

**Fig. 1.9** Insulin hexamer: the cylindrical elements indicate the alpha helices (adapted from http://www.ncbi.nlm.nih.gov/sites/entrez)
1.4 Mechanism of action of insulin

Insulin acts, whether directly or indirectly, on most body tissues, except the encephalon. Its effects are summarized below:

- It increases glucose uptake in cells, stimulating facilitated diffusion: this causes an increase in glycolysis, and the synthesis of fat and glycogen.
- It increases the synthesis of glycogen, not only by increasing facilitated diffusion of glucose in the cells, but also by stimulating the activity of the enzyme which catalyzes the synthesis.
- It increases the synthesis of fatty acids and the formation of triglycerides, as it stimulates fat cells to scavenge lipids present in the bloodstream and inhibits the activity of the enzyme that catalyzes the demolition of triglycerides.
- It promotes the active transport of amino acids in cells, especially muscle cells, thereby increasing the concentration of metabolites necessary for protein synthesis and enhancing synthesis; in addition, it reduces proteolysis.

- It stimulates the enzymes involved in glucose metabolism and inhibits those involved in glucose synthesis (decrease of gluconeogenesis).
- It stimulates potassium uptake in cells.
- It favours the relaxation of the smooth muscle cells in the artery walls, thereby increasing the blood flow, especially in the smaller capillaries.

Note: in the figure above the glucose transporter is called GLUT4 (Glucose Transporter-4): this is a protein found in the plasma membrane of fat and striated muscle cells (i.e. cardiac and skeletal muscle). In the absence of insulin, it is stored...
inside the cell. The binding of insulin to its receptor induces the distribution of GLUT4 in the membrane, where it enables the facilitated diffusion of glucose.

Muscle and fat (which represent on average two-thirds of all the cells of the human body) are the two tissues in which glucose uptake is influenced by insulin. Although insulin plays a primary role in glucose uptake, it does not modify the absorption of glucose in the cells of the encephalon, nor its active transport in the kidney tubule and gastrointestinal epithelium.

### 1.5 Regulation of insulin production

The secretion of insulin is mainly regulated by the concentration of glucose in the blood that permeates the pancreas, and the β-cells in particular, via a negative feedback mechanism (Fig. 1.11). An increase in glucose enhances the secretion of the hormone, which causes an increase of glucose uptake in the cells and a decrease of glucose concentration in the plasma. The decrease of glucose inhibits further secretion of insulin.

The specific mechanism by which insulin is secreted is still unclear. However, a model can be deduced based on what is currently known: glucose enters β-cells of the pancreas mediated by GLUT2, which is present only in the membrane of β-cells, and cells
of the liver, the hypothalamus, the small intestine and the renal tubule. The sugar is used for glycolysis and cell respiration, coupled to the production of ATP. The increase of ATP determines the closure of potassium channels and results in membrane depolarization. As a consequence, calcium ions enter the cell and are further released in the cytoplasm from the endoplasmic reticulum. This increase of calcium is thought to be responsible for the release of insulin accumulated in the secretory vesicles of the Golgi apparatus. In addition, the increase of glucose in the β-cells seems to activate other calcium-independent metabolic pathways that also play a role in insulin secretion. Intracellular glucose also stimulates the transcription of the insulin gene and the translation of its mRNA.

However, glucose is not the only factor that controls insulin release (although it is the most important).

The ingestion of food in general, not only carbohydrates, determines increased secretion. In addition, the nervous system (via visual and taste stimuli triggered by food) and hormones (such as gastrointestinal hormones) also have an influence on insulin secretion.

1.6 Insulin in other animals

The protein sequence of insulin differs from one species to another (sometimes additional amino acids are present). However, there are highly conserved regions, which include the position of the disulphide bridges, both ends of the A-chain, and the C-terminal end of the B-chain. Therefore, the three-dimensional structure is very similar. Mammalian insulin is highly conserved: e.g. the bovine hormone differs from the human one by only three amino acids, and the porcine one by a single amino acid! Even within vertebrates, in general, there are certain similarities: in some species of fish, insulin is sufficiently similar to be efficient in humans. In contrast, the C-peptide of proinsulin differs greatly from one species to another.
The term *diabetes mellitus* means “sweet flow” in ancient Greek. The name originates from the time when doctors used to taste a patient’s urine to diagnose diabetes. The sweet flavour of the urine would be indicative of diabetes mellitus as opposed to *diabetes insipidus* (which means “tasteless” in Latin), another disorder characterized by frequent urination. There are several forms of diabetes mellitus and the feature that they all have in common is an increase of glucose levels in the bloodstream.

### 2.1 Classification

The most frequent forms of diabetes are type 1 and type 2 diabetes. Type 1 diabetes normally has an early onset during childhood or adolescence. However, adults can develop this form of diabetes, in some cases even at the age of forty or fifty. Type 2 diabetes typically develops in adulthood. However, due to the increase in obesity, many young people, even adolescents develop this condition. Other forms of diabetes include gestational diabetes (which occurs during pregnancy), diabetes due to pancreas resection, and some rare forms of diabetes due to genetic mutations.

### 2.2 Epidemiology

About 90% of diabetes mellitus patients suffer from type 2 diabetes. It has been estimated that 150 million people in the world are affected by the disease, and the number is expected to double over the next 20 years. The increase is expected to occur in developing countries, such as India and China. In the United States, where the incidence of the disease is already high, it is expected that one person out of three will develop type 2 diabetes.

### 2.3 Pathophysiology

#### 2.3.1 Insulin resistance

Most patients with type 2 diabetes, as well as prediabetic individuals, are characterized by insulin resistance. In these patients, although insulin is present at normal or even high levels, it is less efficient compared to insulin in normal individuals. Peripheral insulin resistance lowers the efficiency of insulin in mediating glucose uptake in muscle cells. In the liver, insulin is unable to inhibit the production of new glucose (gluconeogenesis) and the lysis of glyco-gen. As a result, the level of glucose in blood rises, leading to hyperglycemia.

#### 2.3.2 Type 1 diabetes

Type 1 diabetes is an autoimmune disease. In genetically predisposed individuals, probably following a viral infection, an inflammation of pancreatic β-cells takes place. Since the β-cells are the only cells that secrete insulin, insulin deficiency occurs as a result. Therefore, patients who suffer from type 1 diabetes require insulin therapy.
2.3.3 Type 2 diabetes

Type 2 diabetes is typically caused by a combination of genetic and environmental factors. The genetic component is stronger than in type 1 diabetes: the twin of a patient affected by type 2 diabetes will almost certainly develop the disease. Other determining factors are diet and physical exercise. When food is scarce, for instance, the incidence of type 2 diabetes is very low.

A good example of the complementarity between genetic background and lifestyle is provided by the Pima Indians. Those who live in Mexico have a diabetes incidence rate of about 8%, while those who emigrated to the United States, where life is more sedentary with access to high energy food (i.e. fat), have a diabetes incidence of up to 50%.

The major risk factor in type 2 diabetes is obesity. Epidemiological studies have shown that compared to lean individuals, obese men and women (with a body mass index >35) have, respectively, a 60- and 90-times greater chance of developing type 2 diabetes. In genetic terms, type 2 diabetes is a multifactorial disease, as its onset cannot be attributed to a single gene.

In contrast to patients with type 2 diabetes, prediabetic patients (who suffer from insulin resistance) do not develop hyperglycemia when fasting. However, when subjected to a glucose tolerance test (oGTT), involving the ingestion of 75 g of glucose, these patients are characterized by very high levels of blood glucose.

For a brief period of time, the pancreatic $\beta$-cells produce a high amount of insulin to counteract the insulin resistance. This is why many prediabetic patients display high insulin levels in their plasma. However, in most cases, the percentage of death of $\beta$-cells overrides that of regeneration of new cells, resulting in a decrease of insulin-producing $\beta$-cells. When insulin productivity in the pancreas is unable to meet the increased requirement of the hormone due to insulin resistance, the prediabetic patient develops type 2 diabetes.

Three main factors contribute to hyperglycemia:
1. Insulin resistance in the muscle, which causes a decrease of glucose uptake from the bloodstream;
2. Defective insulin secretion by the pancreas;
3. Increased production of glucose in the liver due to hepatic insulin resistance.

2.4 Diagnosis

2.4.1 Symptoms

Early symptoms of diabetes include fatigue, general discomfort and a greater tendency to develop infections, such as infection of the bladder. When hyperglycemia is more pronounced, patients begin to expel glucose in the urine, and produce more urine in general. This situation determines the onset of the typical symptoms of diabetes, which are also the typical initial symptoms of type 1 diabetes: frequent urination.
Diabetes Mellitus

(polyuria), with a consequent increased thirst (polydipsia) and, as a result, dehydration and weight loss.

2.4.2 Screening

During the initial phases of type 2 diabetes, the symptoms may be very subtle, and the disease can often remain silent for several years. Unfortunately, the complications of diabetes (reported below) often develop during this phase. It is therefore crucial to regularly screen patients at risk, such as obese individuals, those who have a family history of diabetes and women with previous experience of gestational diabetes. The screening can be done either by determining blood glucose levels when fasting, or by oGTT (described above). The most important criteria for diagnosing diabetes are listed in Table 1.

<table>
<thead>
<tr>
<th>Fastig glucose levels</th>
<th>Normal</th>
<th>Altered glucose tolerance or altered fasting glucose levels</th>
<th>Diabetes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&lt;110 mg/dl</td>
<td>110-125 mg/dl</td>
<td>≥126 mg/dl</td>
</tr>
<tr>
<td></td>
<td>&lt;6.1 mmol/l</td>
<td>6.1-6.9 mmol/l</td>
<td>≥7.0 mmol/l</td>
</tr>
<tr>
<td>2h after oGTT</td>
<td>&lt;140 mg/dl</td>
<td>140-199 mg/dl</td>
<td>140-199 mg/dl</td>
</tr>
<tr>
<td></td>
<td>&lt;7.8 mmol/l</td>
<td>7.8-11.1 mmol/l</td>
<td>7.8-11.1 mmol/l</td>
</tr>
</tbody>
</table>

Table 1: Diagnostic criteria for type 2 diabetes

The most important factor for monitoring the progress of diabetes is glycosylated haemoglobin (HbA1c). The higher the blood glucose, the higher the levels of HbA1C. Since haemoglobin is carried by the red blood cells, which have a half-life of about 120 days, the HbA1c value reflects a control of glucose levels in the 3 months previous to the analysis. A HbA1c value below 6.1% is considered normal. Diabetic patients typically have an HbA1c value of about 7% but it can be as low as 6.5%.

2.5 Treatment

Fundamental for the cure of diabetes is patient education about the causes and mechanism of the disease, the complications, diet and pharmacological treatment.

Diabetic patients must be informed about diet, the importance of physical exercise and the necessity, in most cases, to lose weight. Those patients affected by type 2 diabetes who are able to radically change their life style and lose weight have a good chance of keeping the disease symptoms under control. Unfortunately, only a very small percentage of patients succeed.

If the level of HbA1c exceeds 7%, diet and physical exercise are no longer sufficient and pharmacological treatment becomes necessary. There are many commercially available antidiabetic drugs that counteract hyperglycemia. Metformin reduces hepatic production of glucose, sulphonylurea derivatives increase the secretion of pancreatic insulin and glitazones reduce the peripheral resistance to insulin.
If the level of HbA1c exceeds 7% despite pharmacological treatment with oral anti-diabetic drugs, insulin-based therapy is required. The therapy usually begins by administering long-term insulin during the night. Eventually, many type 2 diabetes patients, similarly to type 1 diabetes patients, require a complete insulin therapy. This consists either in administering a combination of long- and short-term insulin twice daily, or a more intense therapy based on long-term insulin injections, during the night or in the morning, and short-term insulin injections at meals. Insulin must always be given in association with food intake, in order to avoid hyper- or hypoglycemic shocks.

Owing to the high risk of cardiovascular complications (see below), it is crucial not only to keep glucose-dependent symptoms under control, but also cardiovascular risk factors, such as high blood pressure and high cholesterol levels.

2.6 Complications

Because of the high frequency of complications, diabetes lowers a patient’s life expectancy. Disorders of the small blood vessels (microvascular disorders), type 2 diabetes is the major cause of blindness in adults (diabetic retinopathy), renal failure (diabetic nephropathy) and amputation (diabetic foot) in the industrialized world. The most common microvascular complication is diabetic neuropathy, a disorder that generally involves distal sensory neurons, altering perception of vibrations, temperature and pain in the hands and feet. At advanced stages, diabetic neuropathy can cause intense pain.

Type 2 diabetes is often associated with larger blood vessels such as arteries (macrovascular disorder) and a 2-5 fold increase of the risk of developing cardiovascular diseases: myocardial infarction (heart attack) and stroke.
Classical Therapy for Diabetes

Diabetes is a chronic disease and there are currently no effective therapies that can cure it completely. Therapy is mostly necessary for preventing serious complications that arise if diabetes is not kept under control which may lead to death of the patient. The therapy for juvenile insulin-dependent diabetes mellitus is “substitutive”, meaning that insulin is administered from the “outside”. Since insulin was discovered in 1922, huge progress has been made in developing insulin therapy, starting with the classical substitutive therapy using insulin extracted from animal pancreas and progressing, from the 1970s onward, to therapies using recombinant insulin.

3.1 Therapy with animal Insulin

For a long time after its discovery, insulin was extracted from animal tissues and purified as a drug to treat diabetes. Later on, researchers, driven by the need to find alternative sources of insulin in order to supply the increasing demand of diabetic patients, resolved to produce human insulin via two simple strategies: the chemical synthesis of insulin from single amino acids (a feasible process, albeit expensive), and conversion of pig insulin into human insulin by the substitution of a single amino acid. In the 1970s, following the second strategy, the pharmaceutical company Novo Nordisk A/S started to produce “semi-synthetic” insulin by substituting an alanine residue on the B-chain with a threonine.
Activity I – Bioinformatics

Insulin and molecular evolution

The aim of this bioinformatics activity is to guide the student through the world of protein databases, by searching for the protein sequence of human insulin and sequences from other species. The sequences are compared to highlight sequence homologies and possible phylogenetic connections between species. Based on these comparisons, it is possible to determine which animal is best suited for the extraction of the hormone to treat human diabetes. A part of this activity has been developed and re-adapted using bioinformatics material obtained from the web site http://www.gtac.edu.au.

Al.1 Main reference sites

www.expasy.org

ExPASy, or “Expert Protein Analysis System”, is the proteomics server of the (Swiss Institute of Bioinformatics SIB), dedicated to molecular biology, and particularly to the analysis of protein sequences and structures. Created in 1993, ExPASy was one of the first servers for biology on the web. Since then it has been constantly updated and improved. It provides access to several databases in Geneva, such as Swiss-Prot, PROSITE, SWISS-3DPAGE and ENZYME, as well as other databases, such as EMBL/GenBank/DBJ, OMIM, Medline, FlyBase, ProDom, SGD, SubtiList. In addition, it grants access to several analytical tools for the identification of proteins, the analysis of their sequence and the prediction of their tertiary structure. ExPASy also offers documents related to the field, with links to the main information sources on the web.

In particular, Swiss-Prot (www.expasy.org/sprot) is a well-annotated collection of protein sequences that aims to provide high-standard information (such as the description of the function of a protein, its structure, its post-translational modifications, variants, etc.), keeping redundant information to a minimum and guaranteeing a high level of integration with other databases.
http://workbench.sdsc.edu

**Biology WorkBench** is a web interface that allows anyone to readily use bioinformatics tools for research, revision or teaching. It has been available since June 1996 and provides a portal to databases on the web, data collection and various other applications. It has the advantage of having a very simple interface, reducing many passages of data processing with a ‘point and click’. The current version contains a vast collection of databases and calculation tools, among the most useful for molecular biology are the tools which illustrate the relationship between protein sequences and nucleic acids.


National Center for Biotechnology Information (NCBI), created in 1988, creates public databases, allows bioinformatics searches, develops software to analyse genomic data and disseminates biomedical information. The aim is to improve the comprehension of molecular processes related to human health and disease. The site contains databanks for the human genome and other organisms, nucleotide and protein sequences, molecular structures and scientific publications (such as “PubMed”, the major public bibliographic databank with open access in the biomedical field). **NCBI-Entrez** includes a database of experimentally determined 3D biomolecular structures: Molecular Modeling Database (MMDB). These structures are obtained mainly by X-ray crystallography and nuclear magnetic resonance spectroscopy (NMR), and they provide information on the biological function, the evolutionary history and macromolecular interactions. This database is naturally smaller than a protein or nucleotide database (the 3D structure of only a fraction of all known proteins has been determined), but many proteins are homologous to those already present.

**AI.2 Searching for the protein sequence**

- Go to the web site: [www.expasy.org/sprot](http://www.expasy.org/sprot)
  
  Go to the **Access to the UniProt Knowledgebase**: UniProt (Universal Protein Resource) is the most complete worldwide catalogue of protein information. It is the central archive of protein sequences, which brings together the information contained in Swiss-Prot, TrEMBL (i.e. Translated EMBL, the database derived by the translation of the genetic information contained in the EMBL database) and PIR (Protein Information Resource, a public and integrated bioinformatics resource, for genomic and proteomic research, based at the Georgetown University Medical Center – GUMC).

- Click on **SRS** (Sequence Retrieval System).
• Then click **Start** (Start a new SRS session).

• At this point, a query must be entered to search for insulin protein sequences from different animals. Select SWISS-PROT and TREMBL and then click **Continue**.

• In the first box at the top left select **GeneName** from the dropdown menu and then type “ins”, for insulin.

• On the second line select **Organism** and type “chordata” next to it.

• For “Entry list in chunks of” make sure that 200 is selected. Then click **Do Query**.
• A list of proteins belonging to different animals will appear: it is necessary to select only the ones of interest (note that other proteins that contain the text “ins” in their GeneName will appear!) The proteins we are interested in have the initials INS followed by the code for the animal species (go to Appendix I for the species acronyms). For example you can select: INS1_MOUSE, INS_CHICK, INS_HORSE, INS_HUMAN, INS_HYDCO, INS_MYXGL, INS_ONCKE, INS_PANTR, INS_PIG, INS_RABIT, INS_SHEEP, INS_BOVIN. Under “Perform operation on” choose selected, then select FastaSeqs next to the word “with” and click Save.

• A confirmation will appear. Click Save once again, after making sure that in the “Use view” box FastaSeqs is selected.

• The list of protein sequences for every selected protein will appear. Copy the text into a .txt file and save (with the name “ins_sequences” for instance). For amino acid symbols see Appendix II.
AI.3 Comparing the protein sequences

- Go to the site: http://workbench.sdsc.edu. Registration is necessary (it is for free), so click on **Register**.

- After filling in the required fields, click **Register** once more.

- Back to the initial page, select **Click to Enter the Biology Workbench**, insert your **User ID** and **password** and enter.

- In the new page, scroll down and click on **Session Tools** (note: it is also possible to choose the background colour: grey, pink or blue).
• In the list that appears in the white box select **Start New Session** and then click **Run**.

![Image of software interface](image1.png)

• Give a name to the new session (e.g. “insulin”) and click **Start New Session** once again.

![Image of software interface](image2.png)

• The previous page will appear again with the name of the new session and the date. Now click on **Protein Tools**. From the list that appears in the white box select **Add New Protein Sequence** and then click **Run**.

![Image of software interface](image3.png)

• With the entry **Browse** search for the previously saved file (insulin_sequences.txt), select it so that it appears in the box, and then click **Upload File**.

![Image of software interface](image4.png)
• The sequences should now be imported, one by one, in Workbench (please verify by scrolling down). Then click **Save**.

• At this point, the initial page of “Protein Tools” is updated with the name of the session and the imported protein sequences. First of all, we will try comparing two sequences, e.g. the human and chick. They can be selected by ticking the box next to their name. In the central white box, scroll down to the entry **CLUSTALW – Multiple Sequence Alignment** and select it. Then click **Run**.

• The resulting page will show the selected sequences: confirm the choice by clicking **Submit**. Otherwise, go back to the previous page and repeat the selection.
Questions

1) Look at the human insulin: the A-chain is formed by 21 amino acids, the B-chain by 30. In the given sequence, are there really 51 amino acids in all?

2) Initially, insulin is synthesized in a form called “preproinsulin”, a 110 amino acid long polypeptide, which contains additional sequences that will eventually be eliminated: an amino-terminal “pre” sequence (signal peptide, 24 amino acids), which enables the secretion of the protein, and a “pro” sequence (C peptide) that determines the correct folding of the polypeptide chain. On the polypeptide chain the sequences are placed in the following order: signal peptide – B-chain – C peptide – A-chain. Identify the different segments in the human and chick protein sequence.

3) What is the percentage of homology in the different segments? Complete the table:
4) Which sequences are more conserved? Which are less conserved? Give the reasons why you think some sequences are more conserved than others.

5) Why do you think dashes (−) have been inserted in the INS_CHICK sequence alignment? What could be the evolutionary interpretation?

6) You can compare the human insulin sequence with that of another animal. For example, INS_HUMAN vs INS_PANTER, or INS_HUMAN vs INS_PIG, etc. Calculate the percentage of conservation of the different segments.

7) In several regions of the proteins from different animals, the sequences are not conserved. To what segments do they correspond?

8) In which segments of the polypeptide chain do the asterisks appear?

9) Why do you think there is more sequence variability in these segments?

Scroll down to observe the section “Clustal W Dendrogram”. Based on the differences of the protein sequences, the program builds a phylogenetic tree (dendrogram). In a phylogenetic tree there are “branches” that develop from “nodes” (an evolutionary divergence from a common ancestor) that terminate in “leaves”, which correspond to the sequences present in each taxon.
The length of a branch indicates the estimated time since the divergence took place. There are two types of phylogenetic trees: “rooted trees” and “unrooted trees”. In the first case, an event that is evolutionarily distant from the others is taken as the starting point (the root); in the second case, this reference is absent, and the tree can be useful for examining the phylogenetic distance between defined groups of organisms.

- Observe the phylogenetic tree and answer the following questions.

Questions

10) Examining the unrooted tree: which two species are most related to each other? Explain why.

11) The lengths of the “branches” give an idea of the evolutionary distance: after printing out the dendrogram, use a ruler to determine the evolutionary distance in mm between man and the other animals:
- man and horse: ...
- man and chimera: ...
- man and rabbit: ...
- man and pig: ...
- man and hagfish: ...
- man and cow: ...
- man and sheep: ...
- man and chick: ...
- man and salmon: ...
- man and chimpanzee: ...
- man and mouse: 

12) Is the sheep insulin phylogenetically closer to the chick or to the human insulin?

13) What evolutionary relationship is there between pig, sheep and cow?

14) Based on what you now know, comment on the significance of these results.

- Now scroll down to the bottom of the page and click on Import Alignments.

- Tick the box on the left of CLUSTALW – Protein and then select, in the central white box, DRAWGRAM – Draw Rooted Phylogenetic Tree from Alignment. Click Run.
On the following page confirm with **Submit**. A rooted phylogenetic tree will appear. Answer the following questions based on this tree.

**Questions**

15) *Which species is phylogenetically closer to the cow? Which is closer to salmon?*

16) *Observe the pairs “chimpanzee-human”, “salmon hagfish”, “sheep-cow”. Each pair is differentiated by a single “node”. What differences do you notice in the divergence of these animals?*

17) *In biology the term “similarity” indicates a quantitative feature that correlates two or more sequences (such as the percentage of identity, percentage of conservative mutations...). Identify the two species that share the highest percentage of identity for the amino acid sequence of insulin.*

18) *Note: it is important to understand that phylogenetic trees generated using bioinformatics tools are based on data related to the single nucleotide or protein sequences. Besides sequence homology, other methods must be used to determine the evolutionary relationship between species. Can you name any?*

At this point, let’s move back to the initial purpose of this activity: **Which animal would you choose to extract insulin for therapeutic use that is most similar to human insulin?**

**Questions**

19) *Based on insulin’s sequence homology, which animal species would you consider best suited for extracting this hormone?*
20) Do you think the use of this animal could raise ethical and commercial issues (consider whether this species is widespread, easy to breed, or an endangered species, etc.)?

21) Based on these considerations, and keeping homology in mind, which animal could then be the ideal “donor”?

AI.4 Visualizing the tridimensional structure

- There are databases for the 3D structures of proteins. Go to the site: http://www.ncbi.nlm.nih.gov/ and select Structure on the blue toolbar at the top.

- On this page, at the tab “Search Entrez Structure/MMDB” type in the protein whose structure you want to visualize (in this case type “insulin structure” in quotation marks) and click GO.

- The search will provide different models, each identified by a code (e.g. 2AIY). Using the program Cn3D (downloadable for free from the site) it is possible to visualize the 3D structure of proteins. By clicking on the protein code and Related Structures additional information can be obtained. For example:

- The related scientific article (Reference): it is useful to look up the type of insulin (whether recombinant or not), the technique and the conditions used to analyse the 3D structure of the protein (sometimes the article mentions that the hormone is in solution with some other substance, e.g. phenol, that then appears in the structure image).
- Information on the polypeptide chains that are viewed (Biopolymer chains page). Pointing to the different chains, the number of amino acids will appear. In addition, the chain is put in the context of the protein family it belongs to (red sequence). For example, insulin belongs to the “IGF_like family” (protein family similar to the “Insulin-like Growth Factor”), in the subgroup “insulin-like” of vertebrates. The site shows that this family includes insulin and the insulinlike growth factors I and II, which are peptides with multiple functions in control processes such as metabolism, growth, differentiation and reproduction. At the cellular level, they regulate the cell cycle, apoptosis, cell migration and differentiation. With the exception of Insulin-like Growth Factor, the active forms of these peptide hormones are made of two chains (A and B) bound by two disulphide bridges. In particular, in the A-chain the position of the four cysteine residues is highly conserved: Cys1 forms a disulphide bridge inside the chain with Cys3, while Cys2 and Cys4 bind to cysteine residues of the B-chain. In all cases, the two chains are formed from one propeptide.

- Indication of the solvent that may be present, at the entry Ligand (e.g. phenol).

By clicking directly on the image, the 3D structure of insulin is displayed (2AIY – Cn3D 4.1) and, in a separate window, the protein sequences of the imaged chains (2AIY – Sequence/Alignment Viewer) are reported. Specifically, the chosen image 2AIY shows the 3D structure of a hexamer, i.e. of the molecular complex obtained from the association of six insulin monomers, each one formed by the A- and B-chains. As the complex has trigonal symmetry, it can also be interpreted as the association of three insulin dimers.

By clicking and moving the mouse over any part of the figure while keeping the left mouse button pressed, the image can be rotated in any direction.
• Two options are possible:

- By going to Style _ Rendering Shortcuts in the menu, the type of model can be chosen (ball and stick, tubular, etc.). On selecting worms a wormlike structure, which is easy to interpret, is obtained.

- By going to Style _ Coloring Shortcuts the type of colouring can be selected: Secondary Structure highlights with different colours the α-helix and β-sheet structures; Molecule associates a different colour to each polypeptide chain; Hydrophobicity highlights the hydrophilic and hydrophobic regions, Charge the charges, etc.

- Using Style _ Favorites _ Add/Replace, the image can be saved with a name in the favourites.

- Using Style _ Edit Global Style, the global settings can be viewed and modified: in Settings the model type, the colours of chains and background can be modified. You can also decide whether to view external molecules (e.g. zinc ions, phenol, etc.), the objects that represent the secondary structure, and the disulphide bridges. To get familiar with the various style options, select and deselect them and observe how the image changes. In Labels the names of the amino acids can be viewed (under “Spacing” you can decide whether to label each amino acid, or one, two, three, etc.). In Details the sizes of the model type can be changed (e.g. the diameter of the worm-like structure).

- In Show/Hide _ Pick Structures you can decide whether to view all of the chains, or just a few. This can be useful for identifying the A- and B-chains in each monomer.

- With View _ Animation _ Spin the model will rotate automatically.

• We propose the following activity.

Questions

22) In Figure 2AIY select Style _ Coloring Shortcuts and choose the colour mode molecule. With Show/Hide _ Pick Structures ... view only a single monomer of insulin (A- and B- chain), and using Style _ Edit Global Style eliminate external molecules and the objects indicating the α-helix. Orient the monomer so as to be able to view it comfortably. In the window 2AIY - Sequence/Alignment Viewer select the last 3 aa of the B-chain (P-K-T, i.e. proline-lysine-threonine): they will be highlighted in yellow on the image. Without closing the image, go back to http://www.ncbi.nlm.nih.gov/Structure/ to search for the structure of a recombinant insulin, the Lyspro insulin for instance. To do this you can type “Lys pro – human insulin” (in quotation marks!). Several mutant insulin molecules will appear. The one we are interested in has the code 1LPH: Lys(B28)pro(B29)-Human Insulin, which means human recombinant insulin obtained by placing a ly-
sine residue in position 28 of the B-chain and a proline in position 29 of the B-chain. Open the 3D image, view one monomer only as for the 2AIY insulin, and in the box 1LPH - Sequence/Alignment Viewer highlight the last 3 aa of the B-chain (K-P-T). Compare the two proteins: to get a better view of the angles, select the tubular model in Style _ Rendering Shortcuts; going on Show/Hide _ Show Selected Residues you can delete the chains except for the selected amino acids.

23) What difference do you observe?

24) It is possible to repeat this exercise comparing the human insulin with another recombinant obtained from the database search.

25) Open http://workbench.sdsc.edu/ once again to go back to the “insulin” session (Session Tools, tick the session, choose Resume Session in the central white box, and then click Run). In Protein Tool select human and pig insulin and then CLUSTALW – Multiple Sequence Alignment (Run and Submit) to compare the protein sequences.

Identify the A- and B-chains and annotate the difference. In http://www.ncbi.nlm.nih.gov/sites/entrez search “porcine insulin” (in quotation marks!). View the 3D structure (it should be the 7INS) and, as before, keep only the A- and B-chains, highlight the residue that is different from the human insulin and compare.
The drawback of using animal insulin for therapy, besides not being able to satisfy the demand, was that it triggered an immune response in some patients. The production of semi-synthetic insulin only partly solved the problem, as it still depended on the availability of porcine insulin. The possibility to obtain unlimited quantities of human insulin would, therefore, make a strong impact on the pharmaceutical market.

4.1 Towards recombinant Insulin

Some fundamental scientific discoveries of the past century led to the production of human recombinant insulin and revolutionized the treatment of diabetes.

4.1.1 Restriction enzymes

In 1962 Werner Arber, a Swiss biochemist, proved the existence of what he called “molecular scissors”, i.e. proteins capable of cutting DNA. Arber showed that the E. coli bacterium is equipped with an enzymatic immune system capable of recognizing and destroying exogenous DNA and modifying its own native DNA to avoid its degradation. Arber and his colleagues (Daniel Nathans and Hamilton Smith) called this group of proteins “restriction enzymes” and in 1978 they received the Nobel Prize in Medicine for their discovery. They also showed that the activity of these enzymes is controlled: each enzyme targets a specific DNA sequence (also called consensus site or restriction site), which it recognizes and cuts.

4.1.2 Ligases

Shortly after Arber’s discovery, Arthur Kornberg discovered that it was possible to join together fragments of DNA by using an enzyme that he called “DNA ligase”.

Kornberg was trying to construct an artificial viral DNA from fragments of viral DNA, but until then had been unable to synthesize biologically active molecules. However, after adding the ligase, he realized that the enzyme was capable of joining together fragments of DNA and re-establishing phosphodiester bonds between the nucleotides.

The artificial DNA was biological active, as it was able to replicate autonomously. Therefore, Kornberg was considered by the scientific community to be the scientist who was “able to generate life in a test tube”.
4.1.3 Cloning

By the end of the 1960s, the techniques to cut and ligate DNA had been refined. However, scientists were still searching for a mechanism to copy DNA, in order to obtain sufficient amounts to work with.

The discovery finally came in 1971, when scientists developed bacterial transformation. This technique consists in enhancing the introduction of small circular molecules of DNA (called plasmids, Fig. 4.1) in bacteria (such as *E. coli*, in Fig. 4.2), a process which occurs in Nature, albeit seldom. Transformation is achieved by modifying some chemical-physical properties of the bacterial membrane using chemical substances (CaCl₂) associated with rapid heat shock, or an electric shock at high voltage (electroporation). The bacteria become momentarily permeabilized to the exogenous DNA.

Plasmids are extrachromosomal structures present in Nature in some bacteria. They contain a gene that confers antibiotic resistance to the bacterium, as well as sequences of origin and end of replication. Plasmid DNA can be modified by inserting fragments of foreign DNA into especially created insertion sites. Once introduced into bacteria, the plasmid (and thus the foreign DNA) is rapidly amplified thanks to the high frequency of replication of bacteria (Fig. 4.3).

4.1.4 Recombinant DNA technology

Scientists now understood how to cut, ligate and copy DNA. While refining these techniques, they started to think of possible applications.
The term “recombinant DNA” (rDNA) defines a DNA sequence which has been artificially obtained by combining genetic material from different organisms, as is the case for a plasmid containing a gene of interest.

Recombinant DNA technology is based on the discovery (by Herbert W. Boyer and Stanley Cohen at the beginning of the 1970s) that genes – in their study human genes – can be inserted into plasmids and then into bacteria, where they are continuously activated producing functional proteins. Recombinant DNA heralded a new era in biology.

The concept that DNA continues to function when it is transferred from one organism to another opened up infinite possibilities. In 1976, biotechnology became a reality with the convergence of the techniques described above in a single experiment designed to produce a human protein from a recombinant DNA (Fig. 4.4). The idea was to use restriction enzymes to insert the human insulin gene into a plasmid that would then be transformed in bacteria. By expressing the human gene in bacteria, it would be possible to obtain unlimited amounts of hormone.

Insulin became the first commercial product of the biotech industry: instead of receiving insulin extracted from pig or calf pancreas, diabetes patients could now receive recombinant insulin identical to the insulin normally produced in man.

Following the discovery of recombinant DNA, scientists questioned the safety of this new technology. Could the ability to join genes together have the same impact as the atomic bomb or the telescope?

These issues were discussed in 1975 at a conference in Asilomar, California, where scientists, lawyers and the media came together for an open debate.
The meeting ended in the dissemination of guidelines that allowed researchers to use only certain types of bacteria considered to be “safe” and restricted the use of mammalian DNA, measures equivalent to today’s restrictions on the use of the Ebola virus. Five years later these restrictions were amended, allowing considerable progress to be made in the field of mammalian research.

4.2 Synthesis of human recombinant insulin

In the 1970s two biotech companies — Genentech and Biogen — accepted the challenge to synthesize insulin using recombinant DNA technology. Although the idea appeared simple, there were substantial problems. Nobody knew the nucleotide sequence of human insulin and, as previously mentioned, after the Asilomar conference severe restrictions had been imposed on the production of human recombinant DNA.

Two strategies were used to obtain the nucleotide sequence of insulin and transfer it into bacteria. In the end, only one strategy and one of the two companies was successful.

4.2.1 Genentech strategy

In order to synthesize human insulin using recombinant DNA technology, the sequence of insulin DNA was necessary. The amino acid sequence of insulin was known. Genentech scientists deduced the nucleotide sequence of human insulin from the amino acid sequence. The deduced sequence was then generated, inserted into a plasmid and introduced into bacteria by transformation to produce insulin.

In this way, scientists assembled the sequence of human insulin without even using human DNA, allowing them to circumvent the restrictions on human recombinant DNA from the Asilomar conference.

4.2.2 Biogen strategy

To obtain human insulin, Walter Gilbert and his group at Biogen decided to isolate the DNA sequence that encodes human insulin. The sequence of rat insulin was already known. The sequences of human and rat insulin are very similar. It was therefore possible to use the rat sequence as a probe to isolate the human sequence. To do this, the researchers used a “library” of human DNA sequences enclosed in bacteriophages. The rat sequence was labelled with radioactivity and used as a probe to recognize the phage that contained the sequence of human insulin. At this point, the DNA sequence was isolated and introduced into bacteria.

Although promising, this second strategy failed. The sequence that had been amplified by the scientists was not that of human insulin, but rat insulin, owing to a contamination during the course of the experiments.
4.3 Production of recombinant insulin

To produce insulin (or any other eukaryotic protein) in bacteria, several factors need to be considered: the genes of eukaryotes (such as man) contain sequences called introns that do not encode any protein. Bacteria do not have introns in their genes, and therefore do not possess the biochemical apparatus necessary to remove the introns.

Another point to consider is that some eukaryotic proteins are processed after translation. This is the case for insulin, which is initially translated as preproinsulin and through subsequent passages of maturation reaches the final sequence and conformation. In bacteria such maturation processes do not take place, therefore alternative solutions had to be found.

4.3.1 Insulin from bacteria

The production of recombinant proteins can be carried out in different systems: bacteria, yeast, insect cells and mammalian cells. Bacterial cells have the advantage of being simple to handle; they have a short replication time and produce high yields at low production costs.

To produce insulin from bacteria, scientists used two different methods:

a) Method of the two chains

This was the first method used by Genentech in collaboration with Eli Lilly and Company. As previously mentioned, one DNA strand was synthesized based on the amino acid sequence of the A- and B-chains. An enzyme called DNA polymerase was then used to synthesize the complementary strand of DNA. In this way, the two DNA fragments that needed to be inserted into the plasmids were generated.

Each fragment is then inserted into a plasmid inside the gene that encodes β-galactosidase (LacZ), so that the bacteria produce high quantities of fusion protein containing the insulin sequence fused to the end of the β-galactosidase enzyme. The plasmid also contains the gene for the resistance to the antibiotic tetracycline. Plasmids are transformed in the bacteria, and tetracycline is added to inhibit the growth of nontransformed bacteria.

The transformed bacteria grow and the β-galactosidase-insulin fusion protein is extracted and purified. The insulin chains are separated from the β-galactosidase by treating them with cyanogen bromide, a chemical that cuts peptide bonds following methionine residues. Since a methionine residue is inserted between the β-galactosidase and the insulin chains in the fusion proteins, the treatment with cyanogen bromide produces intact insulin chains, detached from the fusion proteins. Finally, the two protein chains are mixed together, and under ideal conditions the disulphide bridges form. Thus, functional human insulin was synthesized in bacteria.
b) The proinsulin method

The strategy of the two chains was later modified by producing a single β-galactosidase-insulin fusion protein, which can be cut in a single step, releasing mature insulin. This method consists in the synthesis of the proinsulin cDNA from its mRNA, modifying its sequence by adding a codon at the 5' end coding for methionine. The modified cDNA is then inserted in a plasmidic gene (such as LacZ, used in the previous method) and amplified in bacteria. The proinsulin, which is produced in the bacteria, is then separated from the β-galactosidase enzyme by degrading the methionine residue. The chain is then induced to refold, which allows the formation of the disulphide bridges, and the C peptide is cut off by an enzymatic reaction that produces the mature protein.
4.3.2 Insulin from yeast

In order to simplify the purification process, scientists decided to devise a method in which insulin would be directly secreted in the culture media after its production inside the cell. For this they used yeast rather than bacteria.

Yeast is a eukaryote, i.e. an organism that has a nucleus and is therefore capable of completing the maturation process of human insulin. The proinsulin sequence is inserted into a plasmid and the recombinant plasmid is transformed in yeast. The yeast can, at this point, produce proinsulin, which is processed in the same way as in humans (Fig. 4.7). The proinsulin folds into a closed structure and the disulphide bridges between the cysteine residues can form normally. The pro-sequence of 33 amino acids is removed normally releasing the mature protein. Novo Nordisk S/A was the first company to obtain proinsulin secreted from the yeast *Saccharomyces cerevisiae*. Today there are different categories of insulin: rapid-acting, short-acting or regular, intermediate-acting, long-acting and premixed. Although the principle of action is the same, the rate at which they are absorbed are different.

![Insulin production from yeast](source: www.accessexcellence.org)
From a clinical point of view, monomers and dimers diffuse more rapidly in the blood compared to the hexameric form. Insulin preparations containing a majority of hexamers are absorbed more slowly. Therefore, researchers aimed at obtaining molecules of recombinant insulin with reduced tendency to form dimers and hexamers. This is the case for “Lispro” or “intelligent insulin”, which is obtained by inverting lysine and proline residues at the C-terminal end of the B-chain (Fig. 4.8): this modification allows a more rapid absorption of the hormone without altering the site for the receptor.

Fig. 4.8 Lispro Insulin (source: www.minerva.unito.it)

In the past few years, while optimizing the administration procedures of the hormone, new schemes for rational treatment have been developed. The possibility of combining insulins with different absorption rate allowed the achievement of good glycemic control. A crucial point is to use rapid-acting insulin at meal times, and long-acting insulin to cover the nocturnal insulin requirements and the hyperglycemic periods during the day.
Activity II – Molecular Biology
Simulation of recombinant insulin synthesis

All.1 Introduction

Today, the insulin gene can be isolated from the genome using the technique called Polymerase Chain Reaction (PCR).

Up to the mid 1980s, the only way to obtain DNA copies was to insert different DNA fragments into bacteria and select the desired fragment amongst countless colonies grown on a plate. In 1985, Kary Mullis invented a new, extremely precise, method to select and amplify a DNA segment. It is commonly called PCR, and Mullis was awarded the Nobel Prize for Chemistry in 1993 for his groundbreaking invention.

PCR enables the amplification of nucleic acid fragments, provided that the initial and terminal nucleotide sequences of these fragments are known. PCR amplification allows rapid, in vitro production of large amounts of genetic material. Once isolated, the insulin gene can be modified at its ends by adding on sequences recognized by a restriction enzyme. The same enzyme also recognizes a specific sequence on the plasmid. The restriction enzyme cuts both the plasmid and the insulin gene ends in a specific and asymmetric manner, producing sticky ends that facilitate the insertion of the gene into the plasmid (Fig. All.1).

Fig. All.1 Construction of the recombinant plasmid
The recombinant plasmid is then integrated in the bacterial cell (through the process called transformation Fig. AII.2). The plasmid contains all the necessary elements for the transcription of the insulin gene, which will then be translated by the bacterial translational machinery.

**Fig. AII.2 Integration of the recombinant plasmid in the bacterial cells**

**AII.2 Protocol**

For this activity, each group will receive a test tube (**TUBE A**) containing the different recombinant plasmids obtained by the insertion of the insulin gene into a plasmid, after they have been digested by the same restriction enzyme.

Different interactions between gene and plasmid are possible:

1st case – the plasmid closes upon itself without interacting with the gene

2nd case – the gene inserts into the plasmid in the wrong direction

3rd case – the gene inserts into the plasmid in the right direction

**Fig. AII.3 Different possibilities of interaction between gene and plasmid**

Our goal is to select the case in which the gene has inserted correctly into the plasmid, producing functional protein. In order to select the correct recombinant plasmid, it is necessary to introduce the ligation product (containing all the combinations of interaction between gene and plasmid) in bacteria by transformation. During this process some of the bacterial cells will allow the recombinant plasmid to enter.
First of all, each group will have to do a transformation to introduce the different recombinant plasmids (TUBE A) into the bacterial cells (TUBE B). The next step consists of increasing the temperature suddenly to enhance the entrance of the plasmids into the bacteria.

1. Take 3 μl from TUBE A and mix with 50 μl bacteria.
2. Shake the tube by tapping the side with finger, and incubate on ice for 30 min.

The next step consists of increasing the temperature suddenly to enhance the entrance of the plasmids into the bacteria.

3. After incubation on ice, transfer the tube to 42°C for 2 min.
4. Then immediately transfer the tube to ice once more for 2 min.
5. Add 500 μl LB buffer to the tube and incubate at 37°C for 30 min. This helps the transformed bacteria recover from the heat shock and gives them time to express the gene for antibiotic resistance.
6. Centrifuge the tube at 3000 rpm for 1 min. Remove 500 μl supernatant. Mix the remaining content of the tube thoroughly.
7. Transfer 50 μl of the tube content onto an agar plate (containing the antibiotic) and spread onto the surface of the plate with a cell spreader.
8. Incubate the plate (upside down) at 37°C, overnight.

The plasmid contains a gene that confers the bacteria with resistance to a specific antibiotic. As a result, the bacteria that have incorporated the plasmid will be able to grow in the presence of the antibiotic. Growing on the plate, the bacteria form colonies of identical cells. Each colony is derived from a different cell. Therefore, each colony contains a different recombinant plasmid. Our goal is to find a bacterial colony that contains the plasmid with the gene inserted correctly: these cells will be able to produce recombinant insulin.

On the following day, you will be able to see how many colonies have grown on the plate after transformation.

At this point, we can analyse the colonies and select those that incorporated the correct plasmid during transformation. To do this, the colonies must be grown separately in order to let the bacteria amplify in suspension.

9. Each member of the group must pick a colony from the plate and transfer it to a tube containing 3 ml LB buffer containing the antibiotic.
10. Incubate the bacteria at 37°C overnight to let them grow.
The next day, the plasmid DNA must be isolated and purified from the bacterial suspension.

11. Pour 2 ml bacterial suspension in a 2 ml tube.
12. Centrifuge at 14000 rpm for 15 s to harvest the cells.
13. Eliminate the supernatant.
14. Re-suspend (using the vortex) the bacterial pellet in 300 μl P1 buffer.
15. Add 300 μl P2 buffer and invert tube gently 3-5 times.
16. Leave for 5 min on the bench.

The P2 buffer disrupts the bacterial membrane and RNA, isolating the DNA from the bacteria. The incubation time is necessary to ensure that the plasmid is released from the bacteria, while the chromosomal DNA is retained.

17. Add 300 μl P3 buffer and gently shake the tube.
18. Incubate on ice for 10 min.

The P3 buffer precipitates all the cell debris, while the recombinant plasmid remains in the supernatant.

19. Centrifuge at 14000 rpm for 10 min.
20. Transfer 900 μl supernatant (which contains the recombinant plasmid into a new 1.5-ml tube.
21. Add 630 μl isopropanol to precipitate the DNA.
22. Centrifuge at 14000 rpm for 20 min.
23. Remove the supernatant and add 200 μl 70% ethanol to wash the DNA.
24. Centrifuge at 14000 rpm for 2 min.
25. Remove the supernatant and let the DNA pellet dry for a few minutes.
26. Re-suspend the DNA in 50 μl water.

The DNA can now be analysed to determine whether the bacterial culture that is being examined has incorporated the right recombinant plasmid during transformation.

For this purpose, the DNA must be digested with another restriction enzyme (called BamHI) that will produce fragments of different length, depending on the gene’s orientation in the plasmid.
27. Prepare the reaction mix for the enzymatic reaction. For each DNA sample you need:
   - 1 μl enzyme (BamHI)
   - 2 μl BamHI buffer
   - 13.9 μl water
   - 0.1 μl BSA 100X

28. Add to each tube 17 μl reaction mix and 3 μl DNA

29. Incubate at 37°C.

The enzymatic reaction takes about 1 h.
Each group will then load the digestion product onto a 2% agarose gel.

30. Weigh 1 g agarose and dissolve it in a flask with 50 ml TAE 1x buffer. Boil the solution in the microwave and mix gently until the solution is clear.

31. Prepare an electrophoresis tray with scotch tape around the edges and insert the comb.

32. Pour the agarose solution in the electrophoresis tray. Leave to set (polymerize).

A control (C) will have to be loaded on the same gel. The control is a closed undigested plasmid, and it is necessary to make sure that the enzymatic reaction has worked and that the plasmid has been digested. If our sample in the gel looks like the control, this means that the recombinant plasmid is still closed and the enzyme has not worked.
To load the samples on the gel, a loading buffer (LB) must first be added to every tube. The loading buffer makes the sample heavier, so that it sinks to the bottom of the well. In addition, to determine the size of the DNA fragments generated by the enzymatic reaction, a molecular weight marker (S), consisting of a mixture of DNA fragments of known lengths, must also be loaded on the gel.

33. Remove the tape around the gel. Put the gel in the chamber and fill the chamber with TAE 1X buffer (the gel must be covered by the liquid).

34. Take 10 μl digestion product (TUBE C) and put in a new tube (TUBE D).

35. Add 4 μl Loading Buffer (LB) to the digestion product (TUBE D) and to the control (C).

36. Load 14 μl control (C) in the first well.

37. Load 14 μl TUBE D in the second well.

38. Load 5 μl marker (S) in the third well.

39. Run the gel at 100 V for about 40 min.

40. Observe the size of the DNA fragments using UV light.

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Fig. AII.5 Picture of the samples on the agarose gel. Control C (closed plasmid); Sample 1 (empty plasmid); Sample 2 (plasmid with the gene in the wrong direction); Sample 3 (plasmid with the gene in the right direction); Size reference S.
Comparing the size of the fragments in the sample with the marker DNA (size expressed in base pairs - bp), we will be able to determine whether our bacterial colony has incorporated the right recombinant plasmid or not. The bacterial colonies containing the plasmid with the insulin gene inserted in the right direction will be amplified further and, after an appropriate stimulation, they will produce recombinant insulin.

All.3 Reagents and instruments

All.3.1 Description of the products used for this activity

- **TUBE A**: contains the different products of the interaction between the plasmid (pCR® -Blunt II-TOPO®, 3519 base pairs, commercial product distributed by Invitrogen) and the insert (a sequence of 1146 base pairs amplified in the lab)
- **TUBE B**: *Escherichia coli* strain DH5α. These bacteria have been treated with calcium chloride in order to make them chemically competent.
- **LB buffer (for 1 liter of solution)**: 10 g triptone, 5 g yeast extract, 10 g NaCl, pH 7.0
- **Antibiotic**: kanamycin, used at a final concentration of 25 μg/ml
- **Buffer P1**: Tris-Cl 50 mM, pH 8.0; EDTA 10 mM; RNase A 100 μg/ml
- **Buffer P2**: NaOH 200 mM, SDS 1%
- **Buffer P3**: potassium acetate 3 M, pH 5.5
- **BamHI**: restriction enzyme that recognizes the sequence (GGATCC)
- **BamHI buffer**: NaCl 150 mM, Tris-HCl 10 mM, MgCl2 10 mM, DTT 1 mM, BSA 100 μg/ml
- **DNA intercalating agent**: SYBR Safe™ DNA Gel Stain in TAE 1X, commercial product distributed by Invitrogen
- **TAE 1X buffer**: Tris-acetate 0.04 M, EDTA 0.001 M
- **Loading Buffer (LB) 6X**: Tris-HCl 10 mM, pH 7.5; EDTA 50 mM, Ficoll 400 10%, Bromophenol Blue 0.25%
- **Control (C)**: closed plasmid derived from the fusion of plasmid and insert (4665 bp)
- **Marker (S)**: 1 Kb DNA Ladder, commercial product distributed by Invitrogen

All.3.2 Materials and solutions necessary for the activity

- Gloves
- Pipettes (P2, P200, P1000)
- Tips
- Eppendorf tubes (1.5 ml, 2 ml)
- Ice buckets
- Timers
- Glass flasks
- Loops for bacteria
- Tubes for bacteria
- Sterile water
- Isopropanol
- 70% ethanol
- Ice
- Agarose
- Agar plates with antibiotic

AII.3.3 Instruments needed for the activity
- Electrophoresis chambers and power supply
- Microwave oven
- Vortex
- Heat block
- Incubator at 37°C
- Ultracentrifuge
- UV transilluminator

AII.4 General laboratory safety measures

AII.4.1 General rules

Do not work alone in the lab.

Smoking in the lab is forbidden.

It is forbidden to eat or drink inside the lab, or to store food in the fridges where laboratory material is stored.

Lab coats must be worn inside the laboratory. The lab coat must be removed before entering rest areas, the library or before meals.

To wash laboratory material never use acids, but appropriate detergents.

Before using dangerous substances or materials it is mandatory to be aware of the precautions that must be taken. Every substance, in its original packaging, is marked by a symbol that indicates the hazard involved in the handling (e.g., toxic, irritant, corrosive, explosive, etc.).

Every lab must be equipped with a first aid kit, with instructions for adequate first aid procedures in case of accident.

The use of splintered or cracked glassware must be avoided. When dangerous procedures are being executed, disposable gloves must be worn and discarded after use.
The last person leaving the lab in the evening is in charge of checking that everything is in order (solvents, equipment, chemicals, cultures, etc.).

### All.4.2 Safety measures when using electrical equipment

Never use multiple socket adaptors to connect more than one instrument.

Report faulty electrical equipment, and worn or damaged cables and plugs to the technical service.

If electrical equipment is found faulty, it must be immediately disconnected from the power supply and technical assistance must be sought.

Appliances with electrical motors should not be placed next to inflammable material or explosives. High voltage appliances (e.g. electrophoresis apparatus) can be lethal. The appliance should be marked with appropriate danger signs.

### All.4.3 Safety measures for handling toxic substances

Toxic substances in the lab should be marked with appropriate signs. Toxic substances can be harmful either through ingestion, skin contact or inhalation. To avoid ingestion of these substances, eating and drinking in the lab is forbidden.

Always wear gloves of appropriate quality, rinse before removing.

To avoid contamination of other areas, gloves should be taken off before leaving the lab.

Volatile substances should always be handled under the hood.

If necessary, a face mask can be used.

### All.4.4 Safety measures for working with biological material

Infectious agents include parasites, protozoa, helminthes, fungi, bacteria and viruses. Contact with animals, or tissues, secretions, blood and urine derived from animals, can involve risk of infection.

Extra care must be taken when handling biological material of human origin (blood, tissues, cell lines). The safety measures in these cases must correspond to risk group 2 policy.

Each lab must be provided with a good disinfectant for first aid purpose. It is forbidden to mouth-pipette, eat, drink, or smoke in the lab.

It is mandatory to wash hands before leaving the lab.
All.4.5 Safety measures for using ultraviolet radiations

Ultraviolet (UV) light sources are commonly used in the lab (e.g. lights for looking at electrophoretic DNA gels or chromatograms, or germicide lamps). The light source must have a screen, or UV-specific protective goggles must be worn.

Short-wave ultraviolet radiation causes the formation of ozone via a photochemical reaction with oxygen in the air. Ozone concentration in the air over 0.1 ppm can cause irritation to the eyes, nose and throat. To avoid ozone-related risks, the room containing the UV light source should be well ventilated.

All.4.6 Safety measures for using centrifuges

Personnel handling centrifuges should be perfectly aware of how they work. Before use, they should make sure that:

- All centrifuges have a safety device, which prevents their functioning if the lid is open;
- The maximum speed, also in relation to the density of the material to spin, is not exceeded;
- All samples to be centrifuged are balanced;
- The centrifuge and rotor manuals are available in the proximity of the machine.

In addition, the person handling the centrifuge must:

- Leave the centrifuge and rotor perfectly clean after use, so that they can be used again without the risk of contamination in case toxic or infective material was used;
- Immediately stop the centrifuge if it vibrates or emits abnormal sounds.

All.4.7 Use of the microwave oven

Do not turn the oven on if it is empty.

Do not use the oven with inflammmable material.

Do not use the oven with sealed containers (they may explode): unscrew bottle caps, remove lids.

Do not use the oven with metal or metal-plated objects (e.g. bottles wrapped in foil) and tin foil.

Do not fill the containers to the brim: the liquid may spill when boiling.

Apply power and time of heating in proportion to the water volume of the object that is being heated. In particular, for aqueous solutions, the liquid may overheat beyond boiling point without the formation of bubbles. This can cause sudden spill of the boiling liquid. To avoid this, mix the solution before placing it in the oven, and let it stand a few minutes in the oven before taking it out.

Use padded gloves to take hot containers out of the oven.
Activity II – Molecular Biology

After use, clean the oven with paper sprinkled with glass detergent.
In case of fire inside the oven, keep the oven door closed, turn it off, disconnect the plug and let the fire extinguish itself by suffocation.

All.4.8 Handling of the apparatus for electrophoresis
Make sure the power supply is off before connecting the cables.
Make sure the chamber lid is correctly positioned before connecting the cables.
After the run, before removing the chamber lid, turn off the power supply and disconnect the cables.
Appendix I: Species of animals utilized for compared analysis of insulin

**BOVIN – *Bos Taurus* (cow)**
- Kingdom: Animalia
- Phylum: Chordata
- Subphylum: Vertebrata
- Class: Mammalia
- Order: Artiodactyla
- Family: Bovidae
- Genus: *Bos*
- Species: *B. taurus*

**CHICK – *Gallus gallus***
- Kingdom: Animalia
- Phylum: Chordata
- Subphylum: Vertebrata
- Class: Aves
- Order: Galliformes
- Family: Phasianidae
- Genus: *Gallus*
- Species: *G. gallus*

**HORSE – *Equus caballus***
- Kingdom: Animalia
- Phylum: Chordata
- Subphylum: Vertebrata
- Class: Mammalia
- Order: Perissodactyla
- Suborder: Hyppomorpha
- Family: Equidae
- Genus: *Equus*
- Species: *E. caballus*
HUMAN – *Homo sapiens*
Kingdom: Animalia
Phylum: Chordata
Subphylum: Vertebrata
Class: Mammalia
Order: Primates
Superfamily: Hominoidea
Family: Hominidae
Genus: *Homo*
Species: *H. sapiens*

HYDCO – *Hydrolagus colliei* (chimaera)
Kingdom: Animalia
Phylum: Chordata
Subphylum: Vertebrata
Class: Chondrichthyes
Order: Chimaeriformes
Family: Chimaeridae
Genus: *Hydrolagus*
Species: *H. Colliei*

MOUSE – *Mus musculus*
Kingdom: Animalia
Phylum: Chordata
Subphylum: Vertebrata
Class: Mammalia
Order: Rodentia
Family: Muridae
Genus: *Mus*
Species: *M. musculus*

MYXGL – *Myxine glutinosa* (hagfish)
Kingdom: Animalia
Phylum: Chordata
Subphylum: Craniata
Order: Myxiniformes
Family: Myxinidae
Genus: *Mixine*
Species: *M. glutinosa*
ONCKE – *Oncorhynchus keta* (chum salmon)
Kingdom: Animalia
Phylum: Chordata
Subphylum: Vertebrata
Class: Osteichthyes
Subclass: Actinopterygii
Order: Salmoniformes
Family: Salmonidae
Genus: *Oncorhynchus*
Species: *O. keta*

PANTR – *Pan troglodytes* (chimpanzee)
Kingdom: Animalia
Phylum: Chordata
Subphylum: Vertebrata
Class: Mammalia
Order: Primates
Superfamily: Hominoidea
Family: Hominidae
Genus: *Pan*
Species: *M. troglodytes*

PIG – *Sus scrofa*
Kingdom: Animalia
Phylum: Chordata
Subphylum: Vertebrata
Class: Mammalia
Order: Artiodactyla
Suborder: Suiformes
Family: Suidae
Genus: *Sus*
Species: *S. scrofa*

RABIT – *Oryctolagus cuniculus* (rabbit)
Kingdom: Animalia
Phylum: Chordata
Subphylum: Vertebrata
Class: Mammalia
Order: Lagomorpha
Family: Leporidae
Genus: *Oryctolagus*
Species: *O. cuniculus*
**SHEEP – *Ovis aries***

Kingdom: Animalia  
Phylum: Chordata  
Subphylum: Vertebrata  
Class: Mammalia  
Order: Artiodactyla  
Family: Bovidae  
Subfamily: Caprinae  
Genus: *Ovis*  
Species: *O. aries*

Image source: [http://gattivity.blogspot.it/](http://gattivity.blogspot.it/)
# Appendix II: Amino acids and their symbols

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>1 letter</th>
<th>3 letters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alanine</td>
<td>A</td>
<td>Ala</td>
</tr>
<tr>
<td>Arginine</td>
<td>R</td>
<td>Arg</td>
</tr>
<tr>
<td>Asparagine</td>
<td>N</td>
<td>Asn</td>
</tr>
<tr>
<td>Aspartic acid</td>
<td>D</td>
<td>Asp</td>
</tr>
<tr>
<td>Asn+Asp</td>
<td>B</td>
<td>Asx</td>
</tr>
<tr>
<td>Cysteine</td>
<td>C</td>
<td>Cys</td>
</tr>
<tr>
<td>Glutamine</td>
<td>Q</td>
<td>Gin</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>E</td>
<td>Glu</td>
</tr>
<tr>
<td>Gln+Glu</td>
<td>Z</td>
<td>Glx</td>
</tr>
<tr>
<td>Glycine</td>
<td>G</td>
<td>Gly</td>
</tr>
<tr>
<td>Histidine</td>
<td>H</td>
<td>His</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>I</td>
<td>Ile</td>
</tr>
<tr>
<td>Leucine</td>
<td>L</td>
<td>Leu</td>
</tr>
<tr>
<td>Lysine</td>
<td>K</td>
<td>Lys</td>
</tr>
<tr>
<td>Methionine</td>
<td>M</td>
<td>Met</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>F</td>
<td>Phe</td>
</tr>
<tr>
<td>Proline</td>
<td>P</td>
<td>Pro</td>
</tr>
<tr>
<td>Serine</td>
<td>S</td>
<td>Ser</td>
</tr>
<tr>
<td>Threonine</td>
<td>T</td>
<td>Thr</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>W</td>
<td>Trp</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>Y</td>
<td>Tyr</td>
</tr>
<tr>
<td>Valine</td>
<td>V</td>
<td>Val</td>
</tr>
</tbody>
</table>
Appendix III: Answers to the questions of the bioinformatics activity

1) Look at the human insulin: the A-chain is formed by 21 amino acids, the B-chain by 30. In the given sequence, are there really 51 amino acids in all?

No, there are more: 110.

2) Initially, insulin is synthesized in a form called “preproinsulin”, a 110 amino acid long polypeptide, which contains additional sequences that will eventually be eliminated: an amino-terminal “pre” sequence (signal peptide, 24 aa), which enables the secretion of the protein, and a “pro” sequence (C peptide) that determines the correct folding of the polypeptide chain. On the polypeptide chain the sequences are placed in the following order:

<table>
<thead>
<tr>
<th>Signal peptide</th>
<th>B Chain</th>
<th>C Peptide</th>
<th>A Chain</th>
</tr>
</thead>
</table>

Identify the different segments in the human and chick protein sequence.

3) What is the percentage of homology in the different segments? Complete the table:

<table>
<thead>
<tr>
<th></th>
<th>Signal peptide</th>
<th>B Chain</th>
<th>C Peptide</th>
<th>A Chain</th>
</tr>
</thead>
<tbody>
<tr>
<td>n°</td>
<td>%</td>
<td>n°</td>
<td>%</td>
<td>n°</td>
</tr>
<tr>
<td>INS_HUMAN VS INS_CHICK</td>
<td>15</td>
<td>62.5</td>
<td>26</td>
<td>86.7</td>
</tr>
</tbody>
</table>

4) Which sequences are more conserved? Which are less conserved? Give the reasons why you think some sequences are more conserved than others?

The sequences corresponding to the A- and B-chains are more conserved. The signal peptide and the C peptide in particular are less conserved. The divergence between the human and chick insulin mostly concern the signal peptide and the C peptide. The mutations that may have appeared in these segments were probably not eliminated by natural selection because they do not directly influence the active form of insulin.
5) Why do you think dashes (-) have been inserted in the INS_CHICK sequence alignment? What can be the evolutionary interpretation?

To obtain the best statistical value for the sequence alignment. One could suppose, for example, that evolution caused the insertion of some amino acids in the human insulin.

6) You can compare the human insulin sequence with that of another animal. For example, INS_HUMAN vs INS_PANTR, or INS_HUMAN vs INS_PIG, etc. Calculate the percentage of conservation of the different segments.

<table>
<thead>
<tr>
<th>Signal peptide</th>
<th>B Chain</th>
<th>C Peptide</th>
<th>A Chain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n°</td>
<td>%</td>
<td>n°</td>
</tr>
<tr>
<td>INS_HUMAN VS INS_MOUSE</td>
<td>16</td>
<td>66,7</td>
<td>27</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Signal peptide</th>
<th>B Chain</th>
<th>C Peptide</th>
<th>A Chain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n°</td>
<td>%</td>
<td>n°</td>
</tr>
<tr>
<td>INS_HUMAN VS INS_HORSE</td>
<td>0</td>
<td>0</td>
<td>29</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Signal peptide</th>
<th>B Chain</th>
<th>C Peptide</th>
<th>A Chain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n°</td>
<td>%</td>
<td>n°</td>
</tr>
<tr>
<td>INS_HUMAN VS INS_HYDCO</td>
<td>0</td>
<td>0</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td>Signal peptide</td>
<td>B Chain</td>
<td>C Peptide</td>
</tr>
<tr>
<td>------------------</td>
<td>----------------</td>
<td>---------</td>
<td>-----------</td>
</tr>
<tr>
<td></td>
<td>n° *</td>
<td>%</td>
<td>n° *</td>
</tr>
<tr>
<td>INS_HUMAN VS INS_MYXGL</td>
<td>8</td>
<td>30,8</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>11</td>
<td>45,8</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td>22</td>
<td>91,7</td>
<td>30</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>83,3</td>
<td>29</td>
</tr>
</tbody>
</table>
7) In several regions of the proteins from different animals, the sequences are not conserved. To what segments do they correspond?

*Mostly to the signal peptide and the C peptide.*

8) In which segments of the polypeptide chain do the asterisks appear?

*In the segments corresponding to the A- and B-chains.*
9) Why do you think there is more sequence variability in these segments?

*Evolution has probably allowed a greater variability between species in the signal peptide and in the C peptide. The mutations that may have appeared in these segments were probably not eliminated by natural selection because they do not directly influence the active form of insulin.*

10) Examining the ‘unrooted tree’: which two species are most related to each other? Explain why.

*Man and chimpanzee. They are separated by a single node, and their branches are very short. They have therefore differentiated from a common ancestor relatively recently.*

11) The lengths of the “branches” give an idea of the evolutionary distance: after printing out the dendrogram, use a ruler to determine the evolutionary distance in mm between man and the other animals:

*Note: depending on the size of the printout, different values can be measured. We will therefore simply list the pairs in order of increasing distance:*

- human and chimpanzee
- human and pig
- human and horse
- human and cow
- human and sheep
- human and rabbit
- human and mouse
- human and chimaera
- human and chick
- human and salmon
- human and hagfish.

12) Is the sheep insulin phylogenetically closer to the chick or to the human insulin?

*Considering the number of nodes that separate the different species and the length of the branches, we can deduce that sheep insulin is more similar to the human one.*

13) What evolutionary relationship exists between pig, sheep and cow?

*The evolutionary line of the insulin variants in the three species differentiated at one point into a new line that produced the pig, while the other branch split only later differentiating to cow and sheep. These two species have a relatively recent common ancestor, and therefore seem to be phylogenetically closer to each other. On the other hand, the common ancestor for sheep-cow-pig is more distant in time.*

14) Based on what you now know, comment on the significance of these results.

*As expected, the fish and the hagfish are placed at the opposite end of the dendrogram compared to the other animals, which are all terrestrial. The chick, which is a bird, has differentiated from all the other animals, which are mammals.*
15) Which species is phylogenetically closer to the cow? Which is closer to salmon?

*The sheep is closer to the cow, the hagfish is closer to the salmon.*

16) Observe the pairs “chimpanzee-human”, “salmon hagfish”, “sheep-cow”. Each pair is differentiated by a single “node”. What differences do you notice in the divergence of these animals?

*The divergence for the pair “salmon-hagfish” took place further back in time, while it is relatively recent for the pair “chimpanzee-human”.*

17) In biology the term “similarity” indicates a quantitative feature that correlates two or more sequences (such as the percentage of identity, percentage of conservative mutations...). Identify the two species that share the highest percentage of identity for the amino acid sequence of insulin.

*Man and chimpanzee.*

18) Note: it is important to understand that these phylogenetic trees generated using bioinformatics tools are based on data related to the single nucleotide or protein sequences. Besides sequence homology, other methods must be used to determine the evolutionary relationship between species. Can you name any?

*Using bioinformatics tools, more reliable phylogenetic trees can be obtained by integrating the sequence homology of more genes and/or proteins. Traditionally, animal phylogenetics is based on differences in body structure, anatomical and physiological differences, embryonal development, modes of reproduction, etc.*

At this point, let's move back to the initial purpose of this activity: **which animal would you choose to extract insulin for therapeutic use that is most similar to human insulin?**

19) Based on insulin’s sequence homology, which animal species would you consider best suited for extracting this hormone?

*The chimpanzee.*

20) Do you think the use of this animal could raise ethical and commercial issues (consider whether this species is widespread, easy to raise, not an endangered species, etc.)?

*The chimpanzee is certainly the animal with insulin most similar to human insulin. However, the chimpanzee is not a widespread animal, and to breed it in captivity would raise objections on the grounds of animal rights. There are less than 200,000 animals in the tropical rain forests, and wet savannas of Africa, spanning from the West coast of the continent to the Eastern regions including Uganda, Rwanda, Burundi and Tanzania. Like other apes (gorilla, orangutan and bonobo), chimpanzees are an endangered species, via loss of their natural habitat, being hunted for their meat and through illegal trafficking of infants separated from the mothers which are killed. Chimpanzees are also bred in captivity for research purposes. Chimpanzees in Africa have been classified as animals “at risk of extinction”, while those in captivity in the United States are considered “endangered”. A recent report (“Reuters” - 24 May, 2007 - Will Dunham) communicated that the U.S. National Institute of Health (NIH), which supports a variety of biochemical*
studies on animals, will stop breeding chimpanzees belonging to the government for research purposes. Based on these considerations, it is not feasible to breed chimpanzees for the extraction of insulin for therapeutic purposes. Pig insulin also has a high degree of homology to the human hormone. As opposed to chimpanzees, pigs are widespread and easy to breed (there are 120 million pigs in the European Union alone!) and every day many are killed for food. The “sacrifice” of pigs for scientific or medical purposes certainly raises fewer ethical problems than the use of primates. As a matter of fact, the pig is also used for xenotransplants. The horse also produces an insulin protein that is very similar to human insulin. However, breeding horses is more expensive and less “productive” than breeding pigs.

21) Based on these considerations, and keeping homology in mind, which animal could then be the ideal “donor”?

The pig.
Watson JD, Gilman M, Witkowski J, Zoller M, *DNA ricombinante*, Zanichelli Editore


http://www.dnaj.org/b/index.html
http://en.wikipedia.org/wiki/Insulin
http://www.vivo.colostate.edu
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http://www.scienceinschool.org
http://www.gtac.edu.au
http://www.cusmibio.unimi.it

http://www.med.unibs.it
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