

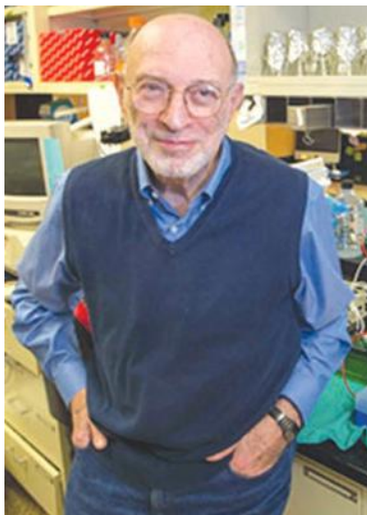
UNIT II

Genome Engineering

Chapter 5: Genome Technology and Engineering

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Stanley Norman Cohen
17 February 1935—present

Stanley Cohen was born in Perth Amboy, New Jersey. After graduating from Rutgers College in 1956, Cohen went to the University of Pennsylvania School of Medicine. He finished his medical degree in 1960. He accepted a position at Stanford University's Medical School in 1968 and began experimenting with plasmids. Cohen is a Professor of Genetics at Stanford University. He is a member of the National Academy of Sciences, and in 1980, won the Albert Lasker Basic Medical Research Award. He was awarded the National Medal of Science in 1988. Stanley Cohen is well-known for his innovation with Herbert Boyer of recombinant DNA technology, which they published in 1973. Stanley Cohen and Herbert Boyer were the first two scientists to transplant genes from one living organism to another, a fundamental discovery for genetic engineering.



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Gradual developments in the area of biotechnology and molecular biology, especially in genome mapping and its implication, has led to the opening of new frontiers for research as well as its applications in many fields. Advancement in genome research has opened up scope for understanding of transcriptome and proteome of any organism on one hand, and application of genomics and related engineering in prediction and treatment of diseases on the other. Applications in crop improvement, gene therapy and remediation of environmental hazards are yet another frontier in which genome technology has tremendous potential and scope. In this chapter, concepts related to genome, its sequencing, analysis and its impact on understanding the biology of organisms have been detailed. In Chapter 7 of Class XI, students have already studied as to how genetic information is encoded in DNA and the processes, like transcription and translation.

5.1 MAPPING OF GENOME: GENETIC AND PHYSICAL

Genome of an organism is the complete genetic information present in the cell of an organism. For prokaryotes, it includes

- 5.1 *Mapping of the Genome: Genetic and Physical*
- 5.2 *High-throughput DNA sequencing*
- 5.3 *Other genome-related technology*
- 5.4 *Genome Engineering*
- 5.5 *Structural, functional and comparative genomics*
- 5.6 *Protein engineering*

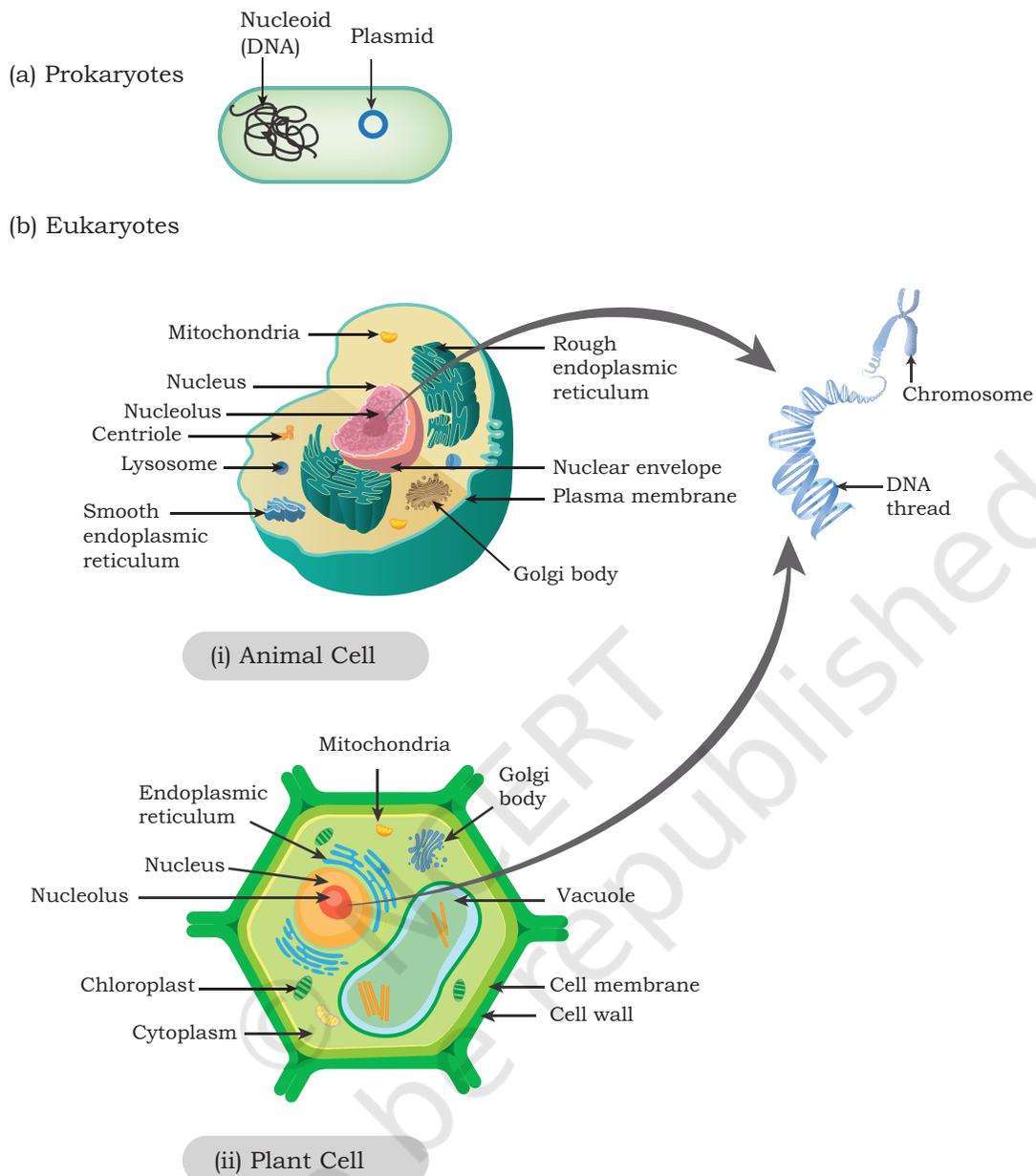


Fig. 5.1: Genome represents DNA content present in Prokaryotes or Eukaryotes. (a) For prokaryotes, it includes the DNA present in the nucleoid and plasmid (b) For eukaryotes, it includes the DNA present in the chromosomes of the nucleus as well as in organelles, like mitochondria and chloroplast

the DNA present in the nucleoid region and plasmid (as it is present in bacteria); and for eukaryotes, it includes the DNA present in the chromosome of the nucleus as well as in organelles, like mitochondria and the plastids, e.g. Protists and all multicellular organisms (Fig. 5.1).

One of the important tasks for many biologists is to compare genomes or DNA of two organisms for various purposes including the understanding of conserved genes or DNA sequences and relationships, etc. In simple words, from multiple samples of a biological organism, a scientist wants to know whether DNA samples from two or more sources are similar or different. If DNA from two sources are different, then how are these different from each other. Further, if an interesting phenotype is noticed, the scientist might also be interested to know the genetic basis (genes and mutations in it) of a particular phenotype. Therefore, a set of techniques is required to answer such questions. Ideally, if we know the complete DNA sequence of each sample, we can compare them and answer all such questions.

In genome research, it is extremely important and crucial to create a map of the genome of an organism from the point of view of identification and relative position of genes or loci. Maps of genome can be created by two different approaches, i.e., the genetic approach and the physical approach, giving rise to genetic and physical mapping techniques for genome mapping in biological organisms (Fig. 5.2).

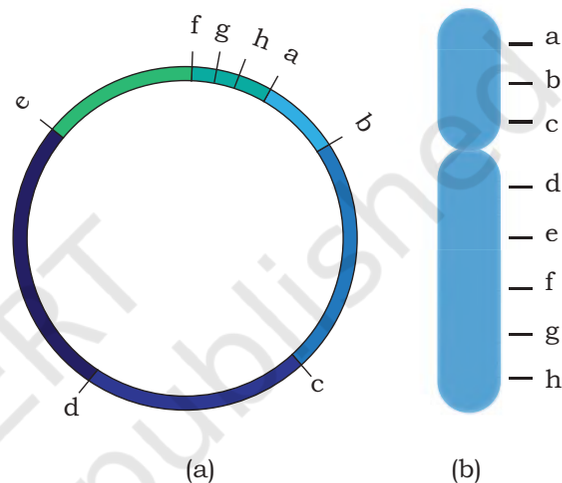


Fig. 5.2: Genome map (a) Map of a prokaryotic/organelle genome. (b) Map of a eukaryotic genome, where a, b, c, d, e, f, g, h are genetic loci or physical sites like SSLP, STS arranged based on genetic distances

5.1.1 Genetic Mapping

Genetic mapping experiments give us an estimate of distances between two or more genetic loci responsible for a set of well-known phenotypes. A genetic map, as shown in Fig. 5.2, is created by carrying out crossover analysis. You have already studied about the phenomenon of crossing over in Class XI (refer section 6.2, Chapter 6). Rate of crossing over (also referred to as recombination frequency) between two or more genes or loci has been successfully used to measure relative genetic distance between them. One map unit is equal to one per cent of observed

cross-over (recombinants). The unit centimorgan (cM) is used to denote genetic distance between genes based on offspring phenotype frequency. Chances of segregation of two loci by recombination are directly proportional to the distances between them. Therefore, decreased recombination frequency is an indication that distance between said loci is less. When genetic loci are arranged based on the genetic distances between them, it creates a genetic map of the organism. The main drawback of this approach is the availability of limited number of genetic loci responsible for a set of well-known phenotypes. This makes genetic maps sparsely populated or coarsely grained and therefore, it limits on the utility of such maps for fine mapping of new phenotypes to the genome.

5.1.2 Physical Mapping

An alternative to genetic mapping is physical mapping of the genome in which specific locations on the genome are identified using DNA based map features. The presence of restriction enzyme sites (restriction mapping), Simple Sequence Length Polymorphism (SSLP), and Sequence Tagged Sites (STS) have been used as map features (Fig. 5.2). These features are helpful in creating a high resolution physical map of genome. Let us briefly understand these genome map features.

Restriction Fragment Length Polymorphism (RFLP):

RFLP is a mapping technique that uses a DNA endonuclease enzyme called restriction enzymes (refer Section 3.3, Chapter 3) that cuts DNA at a specific base sequence resulting in the fragmentation of DNA. The fragmentation pattern of the digested DNA can be visualised when the fragments are separated based on their sizes by agarose gel electrophoresis. Typically, a single cut in a circular DNA yields one fragment of DNA, while the dual cut leads to two fragments. For linear DNA, a single cut results in two fragments of DNA while the dual cut results in three fragments. The DNA is digested with some of the enzymes such as a EcoRI, BamHI, etc., either singly or sequentially

to create a map of restriction sites based on the size of fragments they produce. This exercise results in a high-resolution physical map of restriction sites of the genome. Such a map can easily be compared with others noticing the gain or loss of certain restriction sites resulting in observed RFLP from DNA samples by the changes in length of DNA fragments (Fig. 5.3). For example, in a family with a history of certain disease, comparison of these maps among the diseased and disease-free members may help to identify specific RFLP locus associated with the disease. Association mapping can also help to identify the genetic divergence between populations. However, it is also important here that a mammalian genome may have several sites for different restriction enzymes and it is quite possible that a restriction enzyme like EcoRI may cut a mammalian genome in lakhs of fragments and these cannot be distinguished based on the electrophoresis. This problem is addressed by the technique of Southern hybridisation in which specific DNA probe between different RFLP sites are used. (Fig. 5.4).

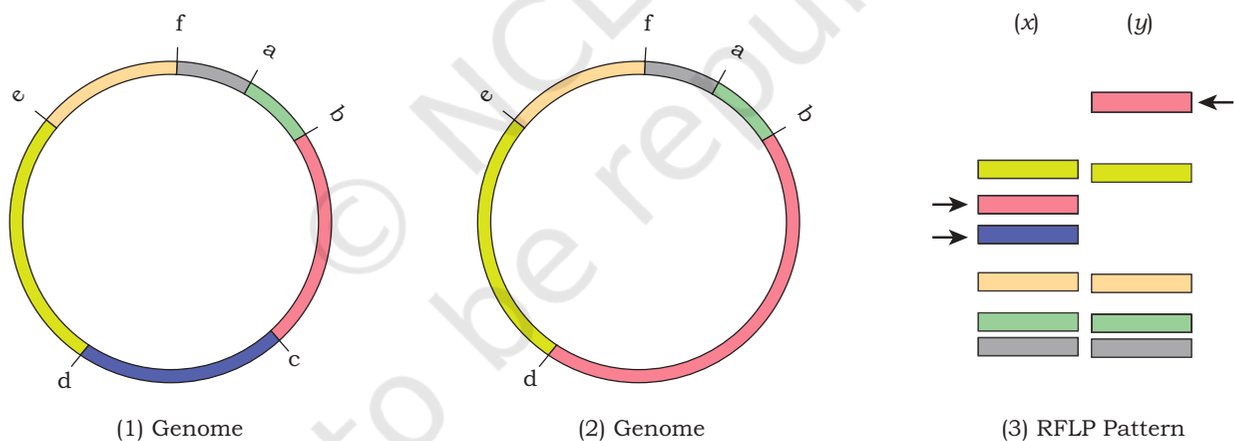


Fig. 5.3: Comparing physical genome maps. The two maps differ at site c. The restriction site c is absent or lost in genome (2) with respect to genome (1) or the restriction site c is present or gained in genome (1) with respect to genome (2). Physical map of a prokaryotic/organelle genome (1) and genome (2), where a, b, c, d, e, f are physical sites (such as restriction enzyme sites) arranged based on physical distances, (3) RFLP pattern of genome (1) and genome (2) obtained by restriction digestion of both the genomes and separating on agarose gel electrophoresis. Presence of restriction site c in genome (1) results in two smaller unique bands in (x) whereas absence of restriction site c in genome (2) gives one larger unique band in (y).

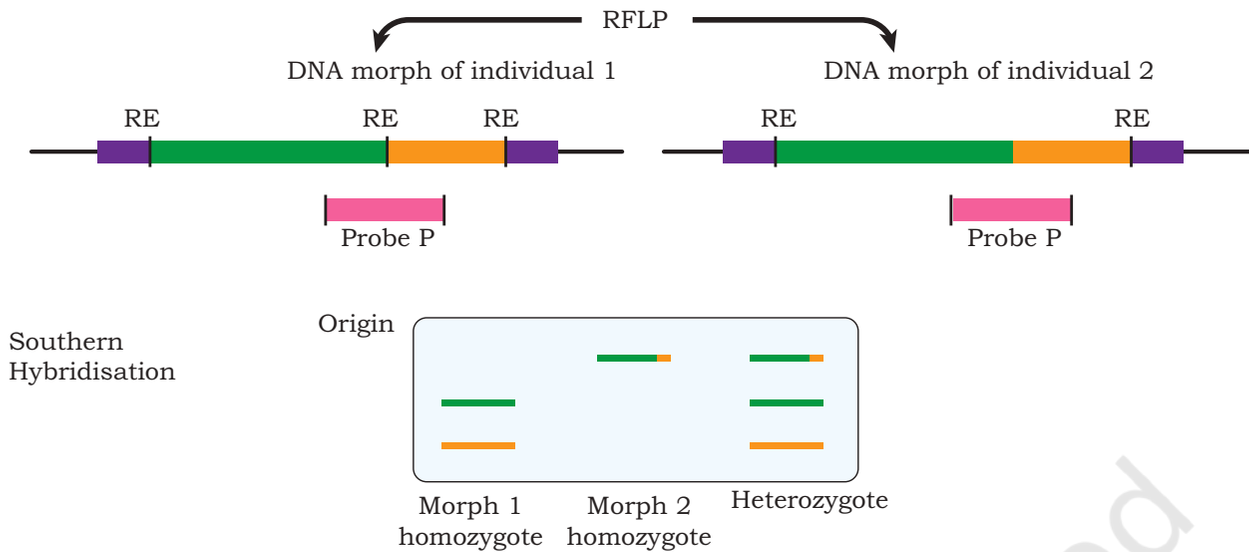


Fig. 5.4: Physical mapping using restriction fragment length polymorphism—A probe P detects two DNA morphs when the DNA is cut by a certain restriction enzyme (RE).

Simple Sequence Length Polymorphism (SSLP) map:

A large genome contains arrays of different repeat sequences, which may be of varying length. Microsatellites and minisatellites are such repeat sequences that can also be used for physical mapping of genome based on its length variant (refer Section 4.1 in Chapter 4 of this book).

Physical Mapping using Sequence Tagged Sites (STS):

Sequence tagged sites are a useful addition to physical mapping. STS are unique DNA sequence (200–500 bp) that occur once in a genome and whose unique location in the genome is known. The presence or absence of STS site in a given fragment of DNA can be easily demonstrated using PCR. Hence, the presence of these sites serve as a useful location landmark in the creation of physical map of a genome. They are often combined with the restriction maps to improve the resolution and effectiveness of association mapping.

5.2 HIGH-THROUGHPUT DNA SEQUENCING

While we can compare the physical and genetic map to differentiate and classify organisms, it is highly desirable to get the complete DNA sequence of the organism. In earlier

days, sequencing complete genomes was prohibitively expensive and therefore, less expensive techniques were used to narrow down the region of interest to small locus of DNA, that is, statistically associated with the phenotype of interest. Only this small region of interest is used for sequencing to avoid the prohibitive cost of full genome sequencing. However, this approach requires a map of genome that could be referred to or compared with others. Advances in sequencing technology have now made it possible to obtain complete DNA sequence of any organism, if we can get its DNA. Initially, the cost of DNA sequencing per base was prohibitive to carry out full DNA sequencing of eukaryotic organisms including human beings. However, this has changed drastically at present. In this section, we will see as to how the DNA sequence technology has evolved over last three decades.

5.2.1 First generation DNA sequencing technology

Initial or first phase of genome sequencing was an elaborate and a multistep procedure. It involved separation of chromosome on agarose gel using Pulse Field Gel Electrophoresis (PFGE). Separated chromosome was then subjected to restriction digestion and the digested fragment were ligated to high-capacity cloning vectors, such as BAC, YAC, PAC, etc. (Refer Section 2.3, Chapter 2). The DNA present in these clones are often overlapping and the clones from such exercises can be organised in a set of minimum clones that carry overlapping set of DNA from the organism.

DNA segments from these clones are physically mapped and sub-cloned into sequencing plasmid for sequencing the cloned DNA using the first generation DNA sequencing techniques.

The first generation DNA sequencing technique used a method called sequencing by chain termination (refer Section 12.9.1 of Class XI *Biotechnology* textbook). In this method, a ladder of single stranded DNA is created, each terminating at a specific base (identified by a specific fluorescence tagged to the terminating base) location. These fragments are separated based on their size using capillary gel electrophoresis technique (in



which the electrophoresis of the charged molecule, i.e., DNA fragments, is done in a capillary filled with gel). A fluorescence detector is positioned near the bottom end of the capillary to read the fluorescence of the single stranded DNA passing in front of the detector. Output of this technology is a chromatogram of fluorescence, where each peak represents a base position and colour of the peak indicates which DNA base is present there. Each capillary in a single run, sequences about 800-1000 base-long DNA. A single DNA sequencer can have as many as 96 capillaries (with 96 different sequencing reactions loaded) operating in parallel in one machine. This type of setup can provide up to 96000 bases per run, per machine.

One of the drawbacks of this technology is that it is time consuming and labour intensive which leads to a prohibitively high cost of genome sequencing. To overcome this challenge, the next generation of DNA sequencing technology was developed.

5.2.2 Next (second) generation DNA sequencing technology

Next generation DNA sequencing technology uses a massively parallel (millions of parallel sequencing reactions in one machine) DNA sequencing approach. This technology does not require time consuming cloning and subcloning steps and therefore, reduces cost required for the genome sequencing project. Compared to the first generation sequencing methods, next generation methods provide advantages, such as simple workflow high data accuracy (due to high depth of coverage) and a broad range of application.

Although there are many approaches to this strategy, we will look at working of one of the most popular Next generation sequencing technologies platform, i.e., **Illumina Sequencing Technology** (Fig. 5.5). It uses a flow cell (glass slide) where DNA fragments are amplified through a special method called **Bridge PCR Amplification**. A flow cell is a solid support, where single stranded small oligonucleotides (8–10 bases) are attached. Initially, DNA is fragmented into small pieces (~1–2 kb) and small oligonucleotide adaptors are attached to these DNA fragments. These adaptors are

complementary to the oligonucleotides on the flow cell and hence, help in the attachment of DNA fragments to flow cell. These adaptors are used as primers and new DNA strands are synthesised. During this process, fragments

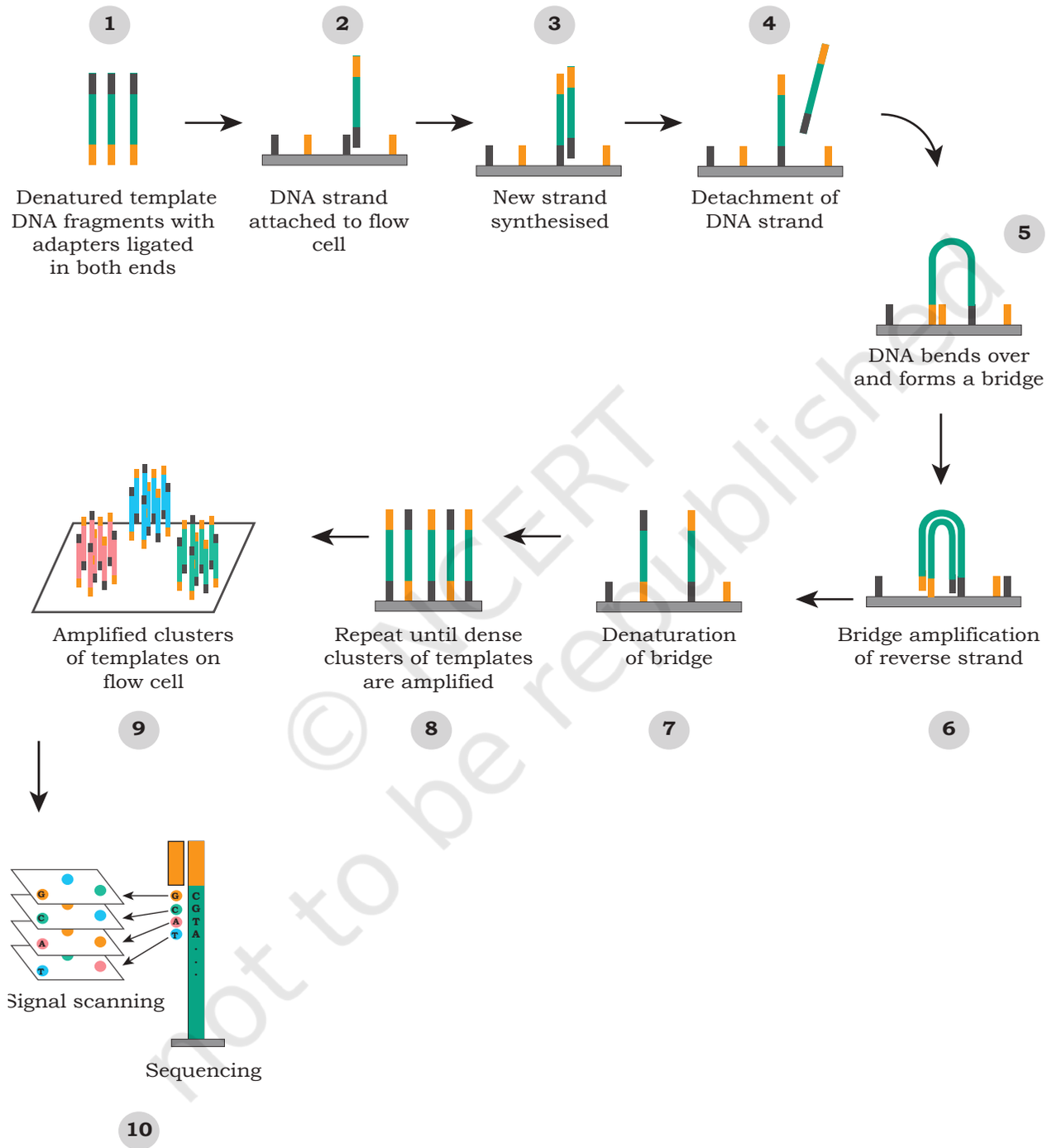


Fig. 5.5: A popular Next generation sequencing technology

make a bridge like structure because of complementarity of adaptors with the oligonucleotides attached to the flow cell. Therefore, it is called **bridge amplification**. It results in a cluster of similar fragments at the same location. After the PCR amplification, sequencing primer is used, which binds to one of the adaptors and each fragment is sequenced at the same time. For sequencing, fluorescently tagged dNTPs are used for the synthesis of complementary strand and each of the bases, i.e., dATPs, dTTPs, dGTPs or dCTPs, is attached with different fluorescent tags. Only one base is added at one time and incorporated bases are read based upon the type of its fluorescence. This is followed by the addition of next base and fluorescence is imaged. In this way, millions of fragments are sequenced parallelly at the same time.

The sequencing reaction is polymerase based and it is carried out in the flow cell by passing labelled base (these bases have unique fluorescence tag). These fluorescent bases would get incorporated depending on the DNA template initially attached to the flow cells. The fluorescence of the newly incorporated base reveal the sequence of the DNA fragment attached on the flow cells (Fig. 5.5). After recording the image of the flow cell, the fluorescent tag of the incorporated base is removed and then next base is passed through the flow cell (Fig. 5.5). The whole sequencing is recorded as an array of fluorescent images, where the DNA sequencing is recorded as fluorescence changes at a given coordinate or spot. Such an array of images reveals the sequence of millions of such spots in parallel.

Although this technology is capable of sequencing millions of DNA in parallel, it generates a very short (75 to 300 base long) length of sequences compared to the first generation technology. Also, additional sequencing strategies are available to overcome some of the limitations posed by short sequencing output.

5.2.3 Some recent advances in DNA sequencing technology

One of the recent advances in DNA sequencing technology is the use of nanopores for DNA sequencing. This is also

considered a third generation sequencing technology. Here, sequencing is carried out by measuring the pattern change in electric current across a nanopore when a specific base in single stranded DNA passes through a nanopore (Fig. 5.6). The current implementation of this technology makes use of two proteins—a DNA helicase and a porins like molecule that is assembled to form nanopore on a synthetic membrane. The DNA helicase captures and unwinds a double stranded DNA and pushes one of its strand through the nanopore. An ionic current is maintained across the nanopore, and it is monitored continuously. When the single stranded DNA bases pass through the pore, the porins like molecule interacts differentially with DNA bases and it disrupts the ionic current. The disruption caused by each base is

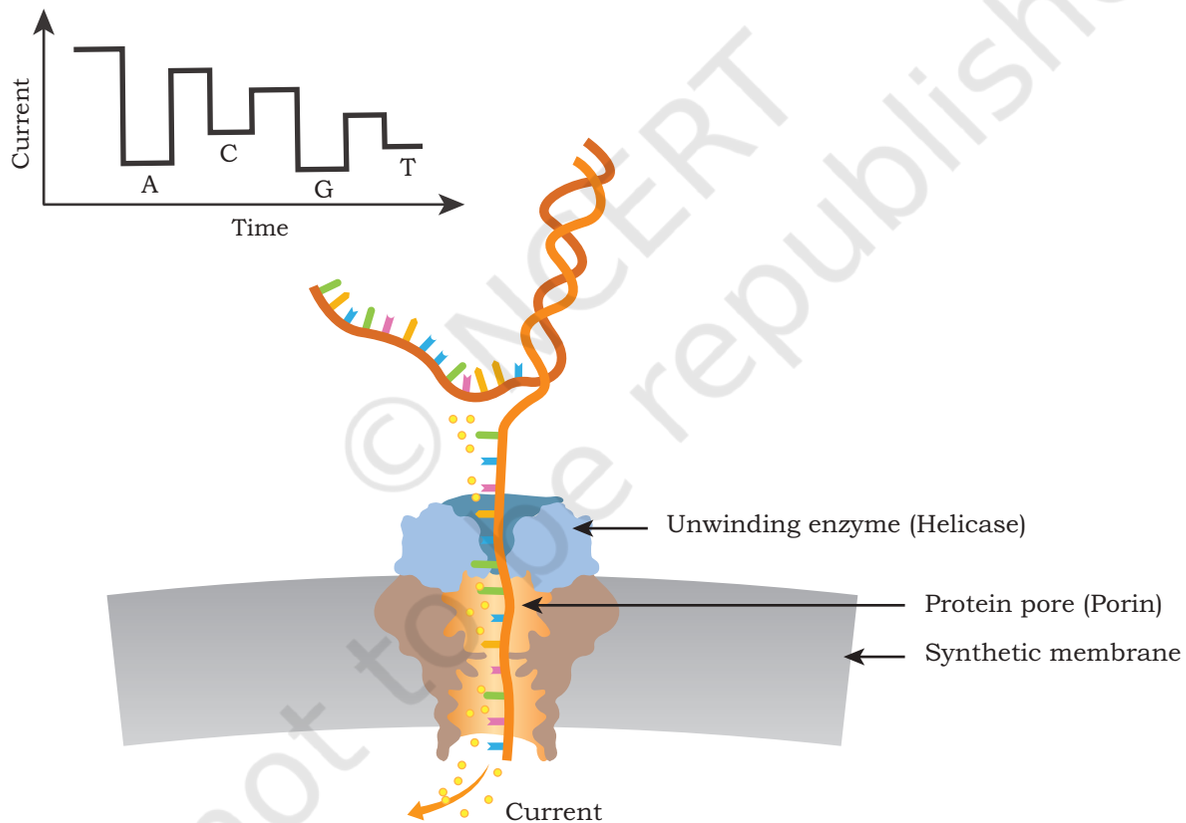


Fig. 5.6: Nanopore based sequencing technology. Yellow dots are ions flowing through a pore creating current across nanopore formed by protein pore. Passage of each base through the nanopore results in specific changes in the ionic current. These changes in the ionic current are measured and the corresponding bases are identified. The sequence read here is ACGT. The unwinding enzyme efficiently captures the DNA, docks on the protein pore and drives one strand of DNA through the nanopore.

very specific and by measuring these signatures of electric current changes, corresponding bases can be identified. The nanopore based sequencing technology offers a rapid and simple sample processing and has the potential to display sequencing results in real-time, making it highly useful for genotyping even in the field setup. Another major advantage of the nanopore sequencing is the production of very long sequencing reads (as long as 1 Mb or even more) at a relatively low cost.

5.3 OTHER GENOME-RELATED TECHNOLOGIES

Genome sequencing comes in multiple flavours or variations. This includes whole genome sequencing (WGS), targeted sequencing, metagenomics etc., as described in the following sections.

5.3.1 Whole Genome Sequencing (WGS)

WGS is a comprehensive method of determining the DNA sequence of the entire genome of an organism. Using WGS, the first genome of *Haemophilus influenzae* was sequenced. The availability of the whole genome sequences is helpful in understanding the genetic regulation of inheritance, identification of the genetic disorders that are inherited and so on. Whole genome sequencing of human genome started in 1990 and complete sequence was published in 2003. With advances in sequencing technologies (as discussed in the previous section), now it is possible to sequence the entire genome in a few days. Whole genome sequence of a large number of microbes, animals and plants has already been obtained. Availability of the WGS can help to create personalised plans to treat human diseases or reading the variations in the cancer cells can help the physician to select the best chemotherapy.

When a whole genome sequencing uses a reference genome sequence (already sequenced and phylogenetically related genome sequence) to assemble its individual sequencing reads into genome sequence it is called reference based genome sequencing. When no such reference is used, it is called *de novo* genome sequencing.

5.3.2 Targeted sequencing

Targeted sequencing project is done to know as to whether targeted regions of a genome have variation or not. Such projects make use of already sequenced genome as reference genome sequence in order to assemble the newly sequenced genome. Targeted sequencing focuses on the sequencing and analysis of a set of selected genes or genomic regions of the genome. It can be used to compare the smaller set of the genomic regions among different genomes to identify variations like mutations, insertions or deletions, etc. For this purpose, target genomic regions is selected, isolated and then sequenced. Usually, PCR is used for this purpose.

Clinical exome sequencing is also very similar to targeted sequencing. Clinical exome sequencing is done for genes that are known to be disease associated (as reported and curated from databases, like OMIM, HGMD, etc.). Genetic diagnostics tests based on clinical exome sequencing are cost effective than whole genome sequencing.

With the development of high-throughput DNA sequencing, DNA sequencing technology itself has been adapted to do much more than just reveal the genome sequence of an organism. One can use this technology to study the biology of an organism at genome level. Some of the advance applications of sequencing technologies are ChipSeq, RNASeq, etc.

ChipSeq sequencing projects are used to map genome-wide DNA binding site of a transcription regulator. Since transcription regulators play an important role in controlling the gene expression of one or more genes, ChipSeq projects are used to know genome-wide DNA binding sites (as well as target genes) of a transcription regulator.

RNASeq projects are carried out to study the global gene expression profile (transcriptome) of an organism, a tissue or a sample. Total RNA is extracted from the target sample and cDNA is synthesised, which is sequenced using next generation sequencing technologies.

The sequencing read thus generated mapped to various genes on the genome and count of such reads mapping to those genes are in proportion to their expression reads.



5.3.3 Metagenomics: Sequencing of DNA or cDNA present in a microbial community

Metagenomics projects are used to study the genomics of community of organisms. Here, we do not separate the organisms or culture them individually. Metagenomics, in fact, is the study of the total genetic material directly obtained from the microorganisms living in a particular environment, like microbes living in the animal gut or in any particular ecological environment. It has several applications in medical microbiology, agriculture as well as environmental microbiology. In particular, it helps to study the diversity of microbial communities and how the diversity of these microbes changes with changes in the environment. Metagenomics may also help in identifying novel genes or enzymes with the significant industrial applications. For example, the study of microbial communities living in extreme environments, like high temperature sulphur springs can help to identify the novel enzymes that are resistant and functional at such high temperatures.

Thus, metagenomics has applications in understanding the genomes of gut microbes, throat microbes, microbes present on a toilet seat, and many more. It provides an insight about the genome of diverse forms of microbes and their correlations. For example, study of genomes of diverse forms of viruses may provide an understanding about virus-host interaction, epidemiology and also evolution of viruses. Presence of multiple genomes in metagenomics sample creates special challenges in data analysis, hence they require a different set of computing algorithms.

5.4 GENOME ENGINEERING

This technology is used to modify the genome. It is often carried out to introduce or remove one or more genes. The goal of this engineering is to introduce a new functionality or to modify (or remove) the existing functionality of a biological organism. In this section, we would get a glimpse of the approaches that had been used to engineer a genome. One of the approaches uses transposon or jumping genes to inactivate or delete a gene(s) in a genome. Another approach is to edit the genome without having any additional unwarranted changes in the genome.

5.4.1 Knock-out and knock-in of a gene by transposon insertion

Transposons are DNA sequences also known as ‘Jumping Genes’ that move from one location to another on the genome found in almost all organisms (both prokaryotes and eukaryotes). DNA transposon generally moves by a ‘cut and paste’ mechanism in which the transposon is excised from one location and integrated elsewhere in the genome. Many a times, such movement of DNA segment results in the insertional inactivation of a gene located at the site of insertion. This behaviour of transposon is exploited to *knock-out* an existing gene. In such a strategy, the transposon is engineered, so that it recognises a specific sequence of a target gene, where it gets inserted. Once the transposon is inserted at the target site, the coding frame of target gene is disrupted and therefore, it can no longer produce the original transcript required to code the gene product (Fig. 5.7).

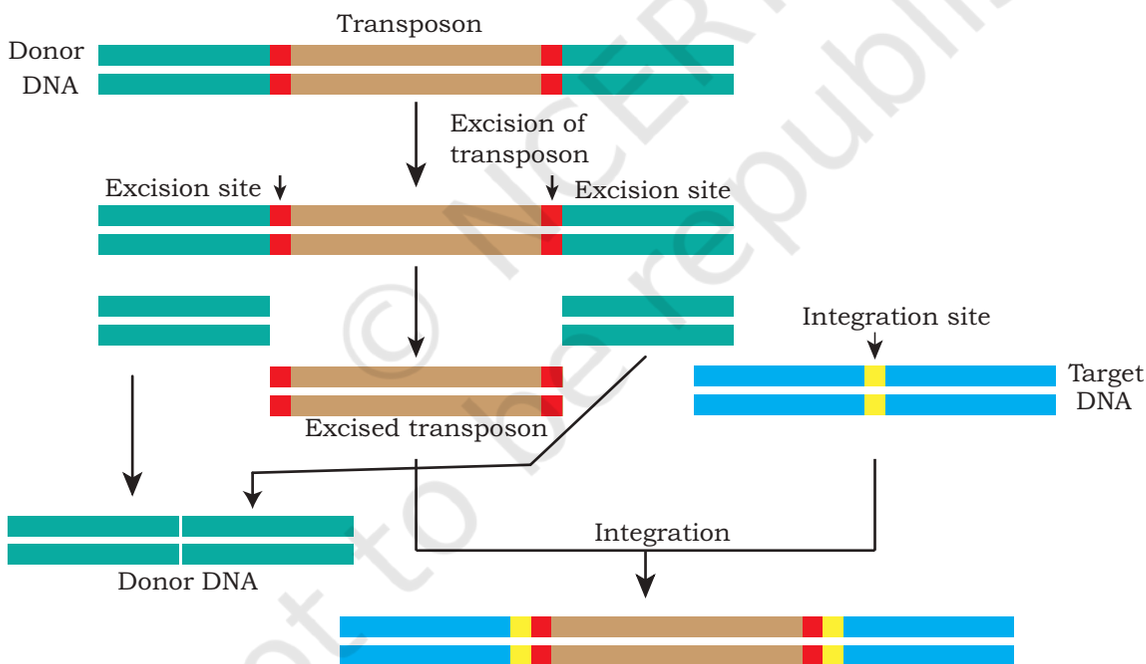


Fig. 5.7: Genome engineering through transposition

In certain cases, when a transposon gets inserted at a particular site, it alters the genetic locus of interest, resulting in the addition of DNA sequence, that was not present on the said genetic locus. This property of

transposon is used to *knock-in* of a functional gene. This *knock-in* and *knock-out* technology helps in the generation of animal/plant models to understand the molecular basis and development of new drugs for the disease.

5.4.2 Genome editing using Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR-Cas9)

Genome editing is a set of techniques or technologies that can be used to change the DNA of an organism by adding, removing or altering the specific DNA sequence of the genome. Like restriction enzymes, this technique is a biotechnological adaptation or repurposing of a naturally occurring genome editing system present in bacteria. In its natural context, this system is used to memorise the DNA signature of past viral attacks so that any subsequent viral infection with matching DNA signature present on the viral DNA can be countered or destroyed effectively.

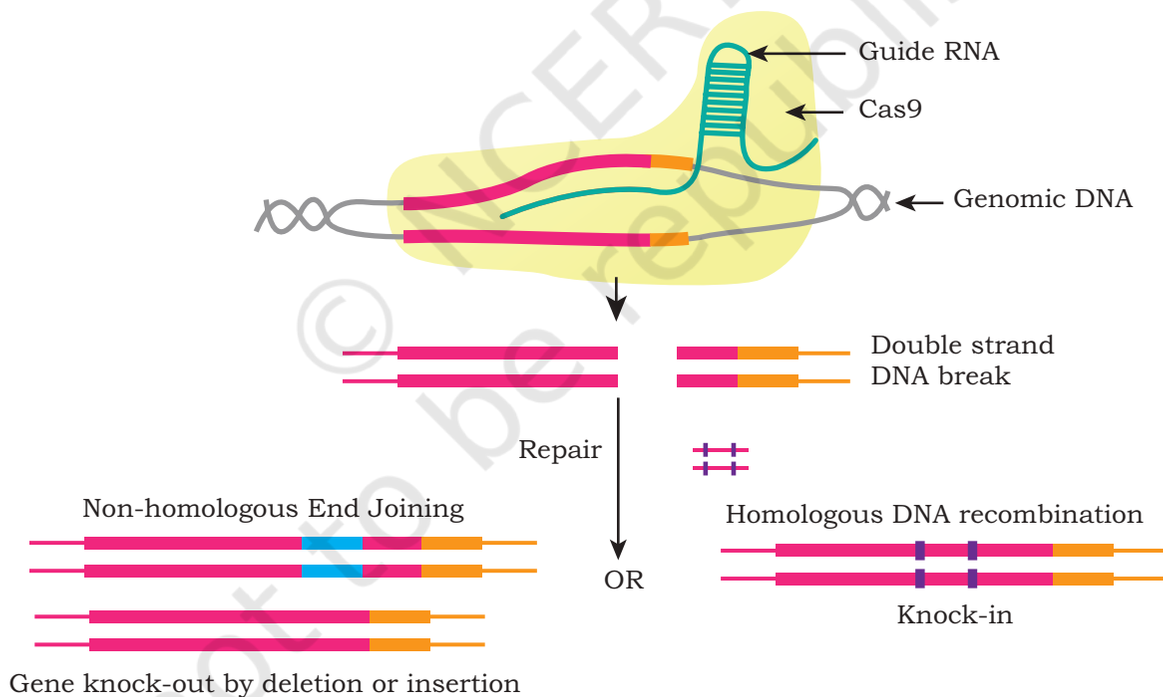


Fig. 5.8: Gene editing using CRISPR-Cas9 technology. Guide RNA position the specific region of genomic DNA with Cas9 protein complex for inducing double strand break. After double strand break, gene knock-out can be created using deletion or insertion of DNA sequence. For gene editing, a desired mutant copy (for correction/replacement around the break site) can be introduced to this region to undergo homologous recombination in the presence of desired mutant template DNA, thereby introducing these mutations in genomic DNA at a specific place.

In this approach, a guide RNA is designed and produced in the cell along with Cas9 endonuclease. The guide RNA are required for positioning of Cas9 endonuclease at a location, where DNA are required to be edited or engineered. Part of the guide RNA is complementary to the DNA sequence, where the sequence needs to be edited. The other part of guide RNA attracts and binds to Cas9 endonuclease. Once this whole complex (the guide RNA bound to target DNA and Cas9) is assembled, a double strand break is introduced in the DNA by the endonuclease (Fig. 5.8). The broken DNA undergoes a template based homologous DNA repair where a modified or edited DNA template is provided that has the desired changes (edited sequence). As a consequence of such double strand break and template guided DNA repair, the repaired genomic DNA now possess an edited DNA sequence.

Box 1 Nobel Prize in Chemistry 2020 for discovering the CRISPR-Cas9 genetic scissors



Emmanuelle Charpentier



Jennifer Doudna

Emmanuelle Charpentier, a researcher at the Max Planck Unit for the Science of Pathogens, in Berlin, Germany; and Jennifer A. Doudna, a biochemist at the University of California, Berkeley were awarded the Nobel Prize in chemistry 2020 for discovering the CRISPR-Cas9 genetic scissors, a gene-editing method.

The concept of CRISPR-Cas9 was modified from naturally occurring ancient bacterial immune systems. Bacteria use CRISPR-Cas systems to recognise and cut the DNA of invading viruses into inactive segments called CRISPR arrays. This effectively disables the virus and contributes to immune ‘memory’ of the bacteria. While studying *Streptococcus pyogenes* in 2011, Dr. Charpentier discovered a new molecule in the CRISPR-Cas system, called tracrRNA, and started a collaboration with Dr. Doudna to reprogram the CRISPR system. Together, they discovered that CRISPR could be directed to cut DNA at the target sequences using Cas9 protein. This allowed the genetic material to be subsequently added, deleted or changed as desired, using the cell’s own DNA repair machinery. Researchers can use this technique to modify the DNA of living organism with extremely high precision. This specialised gene editing tool allows very small and precise changes to be made to the genetic code, it can halt a disease that causes premature ageing in a mouse model and double the lifespan of the treated animals.

5.5 STRUCTURAL, FUNCTIONAL AND COMPARATIVE GENOMICS

Next generation sequencing projects generate large volume of data that require extensive computing for data analysis. Computational genomics is a field that uses high performance computing clusters and workstations to analyse genomics data including the data generated by the sequencing technology. Computational genomics helps in understanding the functions of the genome based on computational and statistical analysis to decipher biology from genome sequences and related data based on DNA and RNA sequences, as well as other experimental data. With the increasing number of genome sequence of different organisms, one major step is the identification of common genes through homology search and gene annotation (assigning functional value). Secondly, sequence comparisons across multiple sequences can be carried out by advanced computational genomics tools, such as BLAST, ClustalW, Phylip to understand the evolutionary relationship. Rapid progress in the field has led to advancements in structural genomics, which deals with the three dimensional information of the proteins encoded by the genome.

Advancements in the area of powerful computer graphics processing units (GPU) and graphic cards have been utilised [General purpose (GP-GPU) and Computer Unified Device Architecture (CUDA) programming] for scientific computing including computational genomics to bring high performance computing for accelerated data analysis.

Genomics work is capital intensive. Since genomics has become a major tool to study biology and has a profound effect on biotechnology, the Department of Biotechnology (under Ministry of Science and Technology, Government of India) has taken an initiative to establish three National Genomics Core facilities in India. These facilities would help Indian scientists and entrepreneurs to execute their genomic projects. These facilities are located within two national laboratories (Centre for DNA Fingerprinting and Diagnostics, Hyderabad and National Institute

of Biomedical Genomics, Kalyani) and one university (University of Allahabad, Prayagraj).

5.5.1 Structural genomics

Structural genomics means two different things to two different sets of people. For those who are interested in protein structure, it refers to 3D structure of every protein encoded by a particular genome. For those who are interested in chromosome organisation, it refers to physical nature of the whole genome.

Earlier, the focus of structural genomics (as defined by structural genomics consortium) was to search for new protein folds or structure in newly sequenced genomic DNA. Through this approach, scientists address the need for solving more protein structures, especially of proteins with sequence that are of unknown fold types.

An expanded view of structural genomics includes the study of structural organisation of DNA region in chromosome and nucleosome status of the genome. This includes the study of large structural changes in genome organisation of related species. An interesting example is to see as to how genes that are common between mice and human are redistributed across different chromosomes of two species.

5.5.2 Functional genomics

Functional genomics aims to understand as to how the physiological function information encoded in genome is executed. A set of genomics based tools and techniques, such as RNAseq, ChipSeq, Metagenomics (of microbial community), etc., are used to understand the physiological and pathological function associated with the state of a cell.

5.5.3 Comparative genomics

Comparative genomics deals with the comparison of genes and genomes of multiple species or within multiple individuals of the same species. It can help in genome annotation of the newly sequenced genome based on its comparison with related well-known genome. It can also help identify the set of common genes that form core



by which point mutation can be introduced at a specific site), the engineered protein becomes stable even in the presence of bleach (Fig. 5.9).

2) Recombinant protein for protein purification: A peptide with six consecutive His amino acids called 6-His-Tag, can bind to metal ions, like Nickel, Zinc and Cobalt with great affinity at mildly alkaline pH. This property of 6-His-tag is exploited for the purification of recombinant protein. DNA encoding for 6-His-tag is fused with the gene sequence of recombinant protein of interest. During protein purification by affinity chromatography, the 6-His-tagged recombinant protein will be immobilised by the presence of metal ions in the agarose gel. All other proteins will be removed by washing the gel. 6-His-tagged recombinant protein can then be eluted by using an elution buffer (Fig. 5.10).

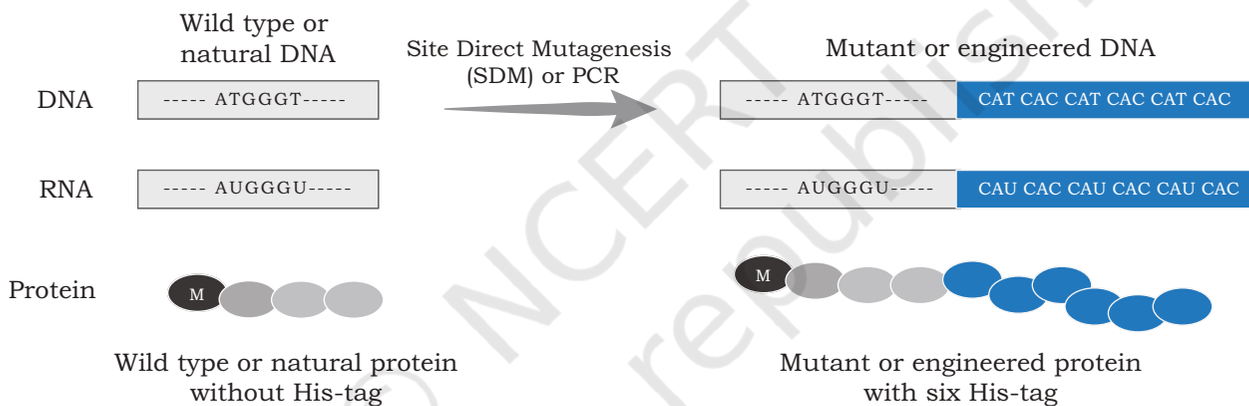


Fig. 5.10: Protein engineering for protein purification. The engineered protein with 6-His-tag are easily and efficiently purified on Nickel ion columns

3) Recombinant protein for tracking the location of the protein in a cell: The gene encoding for green fluorescent protein (GFP) obtained from jelly fish, *Aequorea victoria*, is fused with the DNA sequence of interest. This chimeric DNA now encodes for an engineered protein that appears as a green fluorescent molecule when exposed to UV light, thus revealing its location inside the cell (Fig. 5.11).

4) Recombinant protein as immunotoxin: Recombinant immunotoxins are rationally engineered protein agents (often anti-cancer agents) that combine the selectivity of antibody with cell-killing potency of toxin.

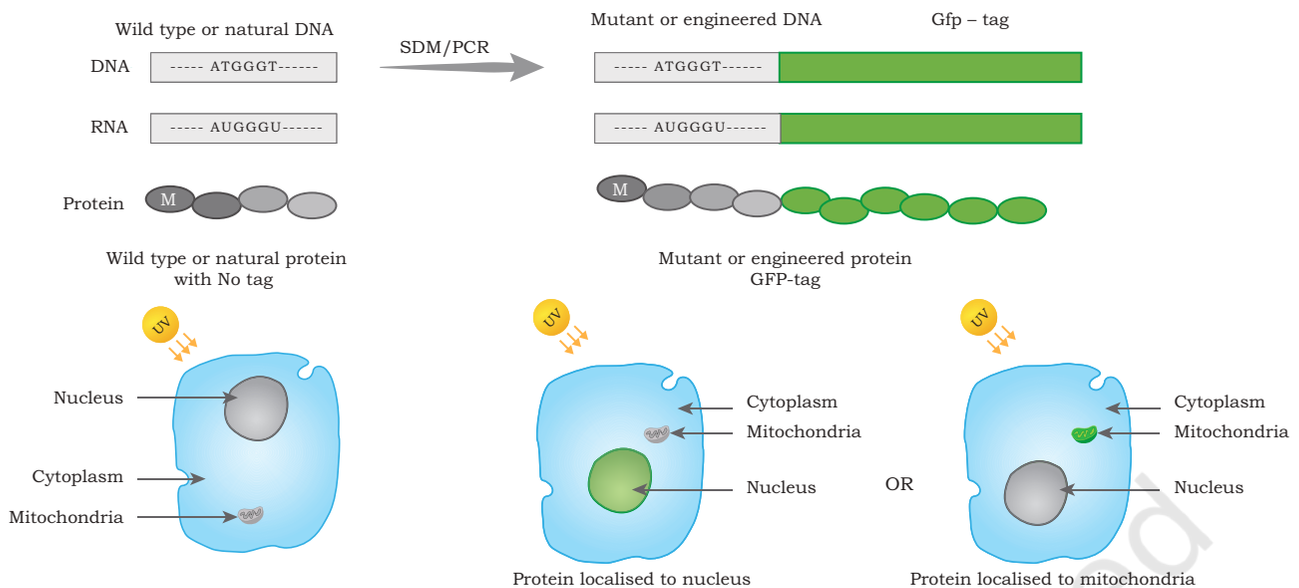


Fig. 5.11: Protein engineering for protein localisation. The engineered protein with Green Fluorescent protein (GFP) tags are easily visualised and localised in the specific regions of cell using epifluorescence or confocal microscopy. The nucleus localised GFP tagged protein appears green in the presence of UV light.

Box 2 Recombinant Immunotoxin

Recombinant immunotoxins are engineered proteins in which the selectivity of antibody is combined with the cell killing potency of toxins. Antibodies are in fact multi-chain proteins having two heavy and two light chains. The function of each fragment of antibody is well defined i.e., the antigen binding activity (present in Fab fragment) and the effector function (in the Fc fragment). The Fab fragment is further divided into the Fv fragment, the smallest fragment which is responsible for antigen binding *via* contact with both the heavy and light chains. For the production of recombinant immunotoxins, recombinant single-chain antibodies, also known as single-chain variable fragments (scFv) are produced through rDNA technology. These antibodies consist of the variable regions of the heavy (VH) and light (VL) chains of immunoglobulins, linked by a short flexible peptide linker. Such scFv can be conveniently produced in bacteria as it can be encoded by a single gene with genetic manipulation and is expressed as a single-function polypeptide chain. During production of recombinant immunotoxins, scFv that specifically binds to the target antigen on the surface of cancer cells are designed. Then toxin fragment that can effectively kill the target cells are selected (such as *Pseudomonas* exotoxin A (PE), *Diphtheria* toxin (DT), or ricin). The designed scFv with the toxin is finally fused with a linker, which allows for proper folding and activity of both the components (Fig. 5.12). This engineered protein uses the antibody part for selectivity targeting the cancer cells displaying their antigens and gets internalized. The toxin part gets released from the endosome and effectively kills the cells.

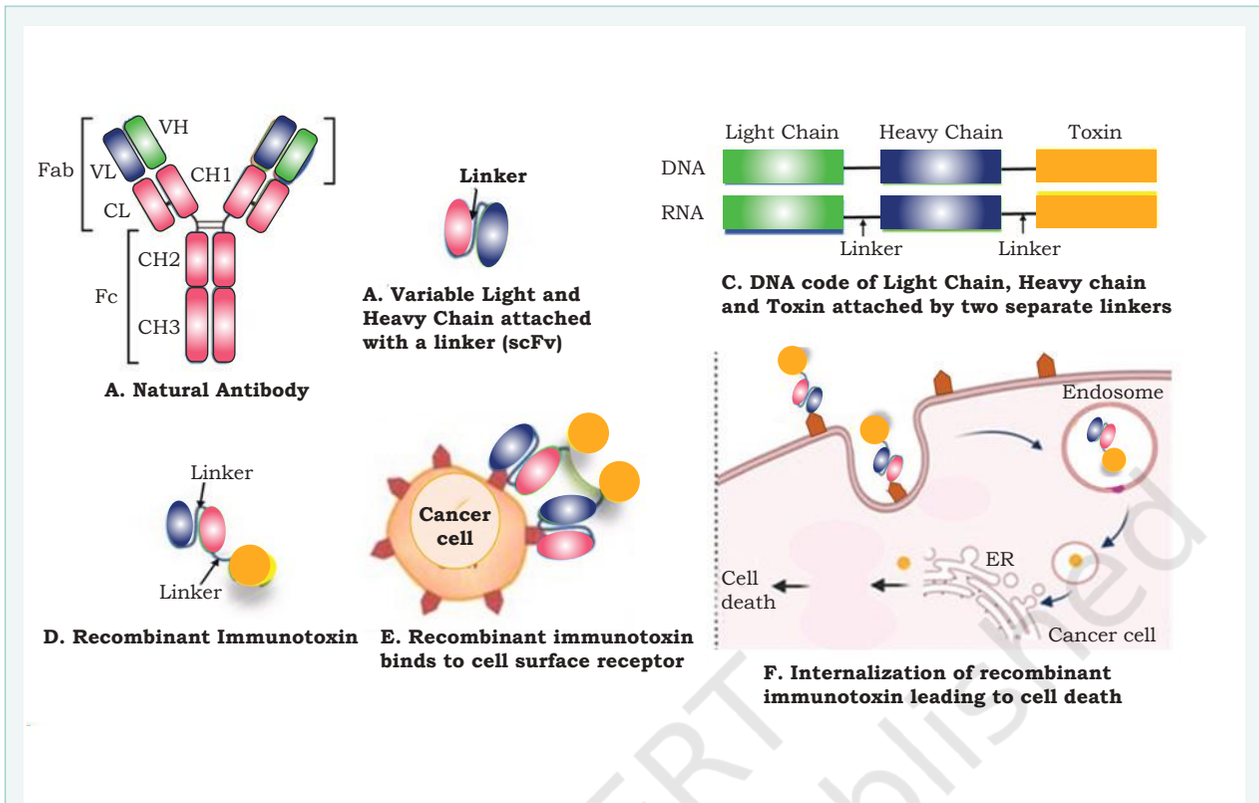
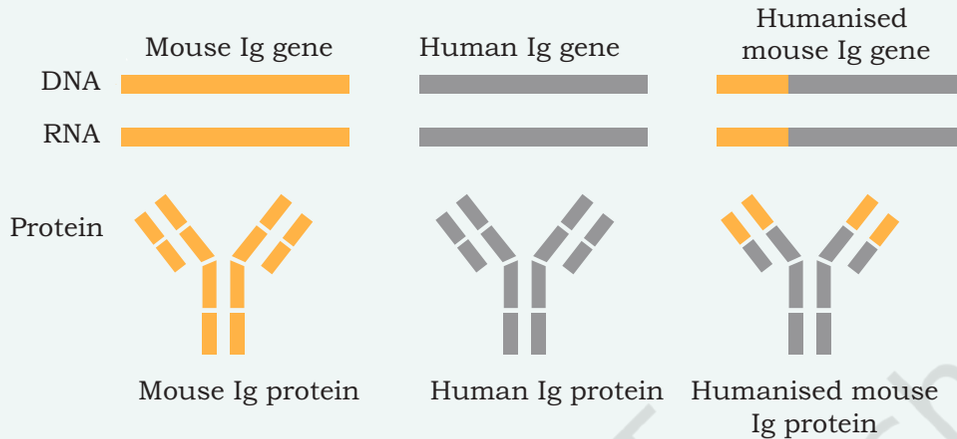


Fig. 5.12: Production of recombinant immunotoxin (A) Natural Antibody (B) Single chain variable fragment joined with a linker (C) Variable part of light chain coding DNA (blue) is fused with the equivalent part of heavy chain coding DNA (green) and toxin coding DNA (yellow) through a flexible linker DNA code (black) giving rise to an engineered single chain recombinant immunotoxin; [D] Recombinant immunotoxin; [E] The antigens expressed on the cancer cell are the antigen binding sites on which the recombinant immunotoxin binds. [F] After binding it gets internalized by endocytosis and the toxin is released in the cytosol from the endosome that kills the cancer cells.

Box 3 Humanised monoclonal antibody

Monoclonal antibodies are very useful reagents that bind their targets in a well-defined way. These antibodies are easily produced using Mouse hybridoma technology (refer Section 4.5, Chapter 4), a biotechnological tool that exploits normal mouse immune response to develop a specific antibody producing immortalised mouse plasma cells by fusing it to mouse myeloma cells. Such immortalised hybrid cells become cell factories for producing specific antibodies. While these mouse monoclonal antibodies are useful as reagents but the same cannot be used as immune-therapeutics in humans. This restriction is due to the ability of human immune system to recognise the mouse antibody as a foreign antigen leading to anti-mouse monoclonal antibody immune response. Raising monoclonal antibodies is not practical in humans. As a result, a biotechnological strategy of protein engineering is applied to humanise the mouse monoclonal antibody. In this strategy, the part of mouse antibody involved in antigen binding is identified and transferred to a human antibody,

which again is done at the DNA level. The DNA coding sequence that codes part of mouse antibody involved in binding the antigen is transferred to the equivalent position in a human antibody coding genes using recombinant DNA techniques, thereby producing a humanised version of mouse monoclonal antibody (Figure). The humanised antibody is often considered self and therefore, has good immune acceptance in human beings when administered for therapeutic purposes.



Humanised mouse Ig using protein engineering

SUMMARY

- The genome of an organism is the complete DNA content, which includes DNA present in the nucleoid in prokaryotes and the DNA present in the chromosomes of the nucleus, mitochondria and plastids in eukaryotes.
- There are two different approaches of genome mapping, physical mapping and genetic mapping.
- A genetic map estimates genetic distances between genetic loci, which are responsible for a set of well-known phenotypes. Genetic distances are measured in centimorgan or map unit (a unit to measure genetic linkage of two phenotypes).
- Physical mapping is based on map features, such as restriction enzyme sites (restriction mapping) and sequence tagged sites (STS).
- In restriction mapping, restriction enzymes cut DNA at a specific base sequence, resulting in fragments of DNA that can be visualised by agarose gel electrophoresis in which DNA fragments are separated based on their sizes.
- STS are unique DNA sequences (200–500–bp) with well-known location, these serve as a useful landmark in the creation of a physical map of a genome. STS can be demonstrated by PCR.

- First generation DNA sequencing technology was a multistep procedure involving separation of chromosomes, restriction digestion and ligation to high capacity cloning vectors.
- First generation DNA sequencing technique used a method of sequencing by chain termination, where a ladder of single stranded DNA with specific fluorescence tagged terminating bases are separated and detected by a fluorescence detector. By this method, base position and type of DNA bases are identified.
- Next generation DNA sequencing technology depends on advance computing algorithms to take millions of short sequence outputs, assembling in large contigs, representing DNA sequences of large genomes.
- Next generation sequencing includes techniques, like whole genome sequencing (WGS), Targeted resequencing, clinical exome sequencing, etc., and ChIPseq, RNAseq are some of the advance applications of sequencing technologies.
- Nanopore sequencing technology uses two proteins: DNA helicase and porins like molecules to bind, unwind double stranded DNA and push single stranded DNA through porins like molecules to identify base pattern. This technique is simple, rapid, cost efficient, and displays results in real-time, and useful for genotyping and high mobility testing.
- Metagenomics involves sequencing of DNA or cDNA present in a microbial community.
- Computational genomics involves usage of high performance computing clusters and workstations to analyse genomics data.
- Genome engineering is a technology to modify a genome (such as inactivate, delete, integrate, transduce and edit the genome).
- Transposons are DNA elements with the ability to move from one position of DNA to another, and can be exploited to engineer a knockout of the existing genes.
- Genome editing using CRISPR-Cas9 (use guide RNA along with Cas9 endonuclease) involves double strand break, homologous DNA repair to generate the edited DNA sequence.
- Structural genomics includes the study of structural organisation of DNA region in chromosome and nucleosome status of genome.
- Comparative genomics identify a set of common genes that form core genome and other genes that are unique to a species. So, it is a starting point for genome based taxonomy and phylogenetic lineage study.
- Functional genomics aims to study the physiological and pathological function associated with the state of a cell.



- Protein engineering is applied to develop novel reagents, diagnostics and therapeutics. It is used to develop recombinant protein with 6 histidine-tag (6-His-tag) for the ease of purification and to develop fluorescent proteins to track the cellular localisation of proteins.
- Protein engineering technology is also involved in the production of humanised monoclonal antibody, single chain antibody and a recombinant immunotoxin which is an engineered antibody that delivers toxin to specific cellular targets.

EXERCISES

1. What is a genome? How is the genome of prokaryotes different from eukaryotes?
2. What are the types of genome mapping? Explain each with a comparative approach.
3. Suppose a DNA is extracted, purified and digested with enzyme BamH1. What type of mapping can be achieved? Mention the application of this type of mapping.
4. What is STS and its role in genome mapping?
5. Give a brief insight about the development of DNA sequencing technology and genomic workflow.
6. Discuss how next generation DNA sequencing technology has overcome the drawbacks of the first-generation DNA sequencing technology. Elaborate the methodology.
7. What is a unit of physical mapping? Discuss in detail.
8. Discuss the methodology and applications of third generation sequencing technology.
9. What are the different variations of Next Generation Sequencing? Explain in brief.
10. What are the applications of advanced sequencing technologies? Elaborate.
11. Discuss the applications of metagenomics.
12. What are the goals of genome engineering? Explain each in detail.
13. One of the applications of genome engineering is to achieve high level recombinant protein production. How can we achieve this? Explain in detail.

14. What is a genome editing? Why is there a need to edit a genome? Explain the methodology of editing using CRISPR-Cas9.
15. What are the structural, functional and comparative genomics?
16. What are the applications of protein engineering? Discuss in brief.
17. If you are given a recombinant protein with 6-His-tags, how are you going to use it? Explain the methodology and application in detail.
18. How do you apply protein engineering to track cellular localisation of proteins?
19. How to generate a recombinant immunotoxin? Why do you think there is a need for generation of an immunotoxin? Mention its mechanism of action.
20. **Assertion:** RFLP is variation in length of DNA fragments when cleaved by same restriction endonuclease.
Reason: Genome of every individual has different position of restriction sites.
 - (a) Both assertion and reason are true and the reason is the correct explanation of the assertion.
 - (b) Both assertion and reason are true but the reason is not the correct explanation of the assertion.
 - (c) Assertion is true but reason is false.
 - (d) Both assertion and reason are false.
21. **Assertion:** Recombinant immunotoxins are rationally engineered protein agents.
Reason: It is prepared by fusing DNA-coding region of antibody with the DNA code of toxin.
 - (a) Both assertion and reason are true and the reason is the correct explanation of the assertion.
 - (b) Both assertion and reason are true but the reason is not the correct explanation of the assertion.
 - (c) Assertion is true but reason is false.
 - (d) Both assertion and reason are false.

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