The nucleus of every human cell contains the full complement of the human genome, which consists of approximately 30,000 to 70,000 named and unnamed genes and many intergenic DNA sequences. The double-helical DNA molecule in a human cell, associated with special proteins, is highly compacted into 22 pairs of autosomal chromosomes and an additional pair of sex chromosomes. The entire cellular DNA consists of approximately 3 billion base pairs, of which only 1% is thought to encode a functional protein or a polypeptide. Genetic information is expressed and regulated through a complex system of DNA transcription, RNA processing, RNA translation, and posttranslational and cotranslational modification of proteins. Advances in molecular biology techniques have allowed accurate and rapid characterization of DNA sequences as well as identification and quantification of cellular RNA and protein. Global analytic methods and human genetic mapping are expected to accelerate the process of identification and localization of disease genes.

The Human Genome Project has provided near-total sequence information about our genes and their surroundings.1,2 This monumental accomplishment has been accompanied by equally remarkable progress in biomedical research and computer science.3-5 Biologists and clinical investigators are now well positioned to perform global analysis of genetic polymorphisms—mutations,6,7 gene expression,8 protein identity and function (proteomics),9 and the 3-dimensional structure of proteins (structural genomics).10,11 The resulting information should accelerate the development of novel and targeted therapeutics that are tailored to individual genomic profiles (pharmacogenomics).12,13 The massive amount of data generated by genome-wide experiments requires the development of comprehensive genomic-proteomic databases and software that enable megadata access, transfer, and manipulation (bioinformatics).14,15

For editorial comment, see page 745.

The development of a structural and functional genomic infrastructure will undoubtedly accelerate our understanding of diseases and help refine methods of disease prediction, diagnosis, staging, and treatment.16,17 Clinicians, clinician investigators, and students of medicine must be prepared to participate effectively in the “new” genomics revolution and tackle the economical and ethical issues that are integral to all aspects of medical genomics.18-21 The intent of the current communication is to begin the process of education in this regard, with a clinician-oriented review of basic principles and methods in molecular biology that are relevant to genomics. In subsequent articles, the aforementioned components of genomics will be discussed and updated periodically.

GENES AND GENE EXPRESSION

Genes and DNA

A gene, the original Mendel “factor,”22 is the basic unit of heredity.23,24 A gene is defined as a contiguous region of
DNA that includes a defined set of exons (protein-coding regions of DNA) and introns (DNA regions interspersed between exons). The basic unit of DNA is called a deoxyribonucleotide (nucleotide for short) and is made up of 1 of 4 nitrogenous bases (adenine [A], guanine [G], thymine [T], cytosine [C]) that is attached to the 1′ carbon of a deoxyribose sugar, which in turn is attached to a phosphate group through its 3′ and 5′ carbons (Figure 1). Each DNA strand is a polymer of many nucleotides that are attached to each other through their phosphate and sugar groups (forming a sugar-phosphate backbone of DNA). There are 2 orientations of sugar-phosphate connections: 5′ to 3′ (carbon 5′ of the sugar moiety of a particular nucleotide in a strand is attached to a phosphate group that is in turn attached to carbon 3′ of the next nucleotide) and 3′ to 5′.

A DNA molecule is made up of 2 antiparallel (1 strand is oriented 5′ to 3′ and the other 3′ to 5′) polymers (strands) of nucleotides (side-by-side chains) in a helical configuration that forms a double helix (Figure 1). The 2 strands of DNA are noncovalently (loosely) connected to each other through their nitrogenous bases by hydrogen bonds (A always bonds with T and G with C). Both strands of DNA are used as templates for new complementary DNA (cDNA) synthesis during DNA replication that precedes cell division. As such, each of the daughter cells contains 1 parental and 1 newly synthesized strand of DNA (ie, DNA replication is semiconservative).

Although the accurate number of genes in the human genome remains to be clarified, the initial drafts from the Human Genome Project suggest the presence of approximately 30,000 named and unnamed genes and many intergenic DNA sequences. In other words, the human genome represents the entire DNA sequence (approximately 3 billion base pairs [bp] per haploid set of chromosomes) in a human cell and is made up of approximately 1% exons (protein-coding regions of DNA), 24% introns (DNA regions interspersed between exons), and 75% intergenic DNA (DNA regions that flank genes). The intergenic DNA consists largely (55% of the total genome) of repetitive sequences, including inverted repeats. Tandem repeats make up “satellite” DNA, a band of DNA that separates from other DNA on a cesium chloride density gradient centrifugation. Satellite DNA is usually localized near the centromere. The terms minisatellites (20-70 bp repeats) and microsatellites (2-4 bp repeats) are used to refer to tandem repeats of small size. The total length of such tandem repeats is highly variable among individuals and between cells of different clonal origin. Accordingly, this variability has been used in different investigations, including person identification and cell chimism.

Scattered DNA sequence repeats can also be short (short interspersed nuclear elements [SINEs], eg, Alu repeats of 100-300 bp long) or long (long interspersed nuclear elements [LINEs], eg, L1 [line-1; long inserted element-1] sequences of 1-5 kilobase [kb] long). An Alu element is a specific DNA sequence that derives its name from the Alu 1 endonuclease because of a single recognition site for the particular enzyme located near the center of the Alu sequence. Approximately one half million Alu repeats are scattered throughout human genome and serve no apparent function. Some DNA sequences can self-replicate, and the additional copies might migrate to different locations within the genome (mobile elements or transposons). Some mobile elements may have originated from viral...
RNA and propagate via an RNA intermediate by using reverse transcriptase (retrotransposons).48,49

Most human DNA is organized into 46 chromosomes that reside in the nucleus of every nucleated cell. However, some DNA resides in mitochondria that are located in the cytoplasm and number in the hundreds per individual cell.50 Each mitochondrion has many copies of a circular DNA molecule (the mitochondrial chromosome), which is approximately 16,500 bp long, and is replicated and expressed within the organelle.51 The mitochondrial DNA (mtDNA) may have coevolved with nuclear DNA and encodes for 2 ribosomal RNAs (rRNAs), 22 transfer RNAs (tRNAs), and 13 functional polypeptides that are involved in oxidative phosphorylation.52 Mitochondrial DNA differs from nuclear DNA in terms of not having introns and displaying high mutation rates.53,54 Because mitochondria are located in the cytoplasm, inheritance of mtDNA is exclusively maternal.55 The combination of uniparental inheritance and high mutation rates has made mtDNA important in the study of human evolutionary genetics.56-58

Each chromosome represents 1 double-stranded DNA molecule that is closely packed with special proteins called histones (DNA + histones + regulatory proteins = chromatin) (Figure 2).59,60 Chromatin condenses substantially during the metaphase stage of mitosis and becomes visible by light microscopy. Chromatin that remains condensed during interphase is called heterochromatin.61-63 Chromatin that condenses only during metaphase is called euchromatin. In general, euchromatin contains more functional genes, and heterochromatin contains more nonfunctional repetitive DNA elements.

In women, the 46 chromosomes in each cell are found in 23 homologous pairs (2 haploid sets of chromosomes): 1 set inherited from the father and the other from the mother. The same is true for men except the twenty-third chromosome pair is only partially homologous, at the point where it pairs an X to a Y chromosome.64 The smallest and largest human chromosomes contain about 50 and 250 million bp, respectively. Each human gene is usually more than 10,000 bp long.1 However, only a tenth of this actually codes for amino acids of the mature protein product.

Gene Expression

The information contained in a given gene is expressed if the DNA sequence of the gene is first transcribed into an RNA molecule that, after being processed in the nucleus, will carry the information into the cytoplasm for translation into a protein.65 Unless indicated otherwise, the mechanisms described subsequently pertain primarily to eukaryotic cells.

Transcription.—Transcription starts with formation of an RNA sequence from a DNA template representing a gene. The protein-coding gene has both transcribed and nontranscribed regions of DNA.66 The transcribed region is first copied onto a nuclear RNA called the primary transcript (pre-messenger RNA [mRNA] or heterogeneous nuclear RNA) in a process catalyzed by the enzyme RNA polymerase II.67 Part of the nontranscribed region (the promoter sequence) of the gene is the site of the initial contact with RNA polymerase II and controls transcription of the primary transcript; it is usually located upstream (20-100 bp) of the start of the transcribed region.68,69 Other regulatory elements of transcription (enhancers-silencers) can be on either side of the transcription start site and be several thousand nucleotides away from the place where transcription actually initiates.70 However, both promoters and enhancers-silencers usually reside on the same stretch of DNA (ie, are cis-acting).
Of note, neither the cis-acting sequences of DNA nor the RNA polymerase II alone could initiate transcription without the help of proteins called transcription factors.71-73 These “trans-acting” proteins recognize and bind to the cis-acting DNA sequences and together with RNA polymerase II and other coactivators of transcription factors form the transcription complex that ultimately regulates transcription either in a positive or a negative manner (Figure 3).74,75

In this regard, promoters often contain characteristic sequences such as TATA and CATT that are recognized by transcription factors.76-79 Transcription factors may be grouped into different classes based on motifs present in their 3-dimensional protein structure (eg, helix-turn-helix, helix-loop-helix, Leucine zipper, zinc finger).80,81 These unique motifs allow recognition of the aforementioned specific DNA sequences that are cis-acting regulators of transcription.82

During transcription and after completion, the primary transcript (always synthesized from 5′ to 3′ orientation) undergoes several modifications before the mature mRNA is released into the cytoplasm for protein translation.83 These include “capping,” addition of a “poly-A tail,” and “splicing” of the primary transcript.84 First, an extra nucleotide (the cap), a 7-methylguanosine (m7G-P-P-P) is placed at the 5′ end soon after the initiation of transcription.85 This modification is believed to protect RNA degradation and mark the starting point for protein translation by binding to ribosomes. Subsequently, the end of transcription is heralded when RNA polymerase II reaches a “termination sequence” (TTATT on the DNA template and AAUAAA on the primary transcript), which, together with other proteins, signals cleavage at the polyadenylation site (10-35 nucleotides downstream of the termination sequence).86,87 Finally, poly(A) polymerase adds a stretch of approximately 200 adenine nucleotides (poly-A tail) at the 3′ end of the primary transcript.88

Once synthesis of the primary transcript is completed, the introns of pre-mRNA are spliced out and the remaining segments rejoined to form the mature mRNA.89,90 The latter includes both translated and untranslated regions (UTRs) of exons, the cap, and the poly-A tail (Figure 4).86 Splicing occurs in a large complex called a spliceosome, which contains the primary transcript and its associated proteins, several protein splicing factors, and a class of small nuclear RNAs that are present in the cell as small nuclear ribonucleoproteins (snRNPs).91-95 There is increasing evidence that both snRNPs and intronic RNA may possess catalytic activity (ie, are ribozymes) and contribute to RNA splicing.96-98 The spliceosome recognizes the 5′ and 3′ splice sites on the exon-intron boundaries via conserved consensus sequences within the intron.99 The actual process involves both lariat (loop and tail) formation of the excised intron and transesterification of the 5′ and 3′ splice sites of the exons.100,102 Alternative splicing occurs not infre-
quently, resulting in different mRNA species and subsequently different proteins from the same primary transcript. Alternative splicing may occur in as many as one half or more of all human genes and sometimes results in altered translation efficiency rather than synthesis of a different polypeptide. The widespread occurrence of this phenomenon increases the coding capacity of genes and contributes to proteomic diversity. The regulation of alternative splicing is an active area of current research, and splicing regulators may include snRNPs.

In conventional terms, the final mRNA represents the sense template, whereas the cDNA strand that was used as a template for its synthesis is referred to as the antisense strand. Regardless, not all the nucleotide sequences of mRNA are ultimately translated into proteins. The very first and last transcribed regions of the primary transcript contain sequences (UTRs) that do not code for protein despite being part of the first and last exons. Nonetheless, these sequences are important for the function of the mRNA and often control the stability, subcellular localization, and translational efficiency of an individual mRNA.

Translation.—The mRNA is translated into protein in the cytoplasm of the cell with the help of ribosomes. Ribosomes are made up of 2 subunits (60S and 40S), each consisting of proteins and rRNA. The mature mRNA that is released into the cytoplasm docks onto rRNA, and its ORF is recognized by another form of cytoplasmic RNA called transfer RNA (tRNA) (Figure 5). A cloverleaf-shaped adaptor molecule, tRNA carries a specific amino acid at its 3’ end. At the opposite end of the amino acid–binding

![Figure 4](image1.png)

**Figure 4.** Steps leading from gene to protein for eucaryotes (A) and procaryotes (B). mRNA = messenger RNA. See text for further details. Copyright 1998 from Alberts et al. Reproduced by permission of Garland Publishing, Inc, part of The Taylor & Francis Group.

![Figure 5](image2.png)

**Figure 5.** Translation of messenger RNA (mRNA) into amino acids in the cytoplasm of a cell. Process requires the concerted effort of mRNA, transfer RNA (tRNA), and ribosomal RNA (rRNA). See text for further details. Reproduced with permission from Lodish et al.
site, the tRNA is equipped with a 3 nucleotide sequence (ie, the “anticodon”) that undergoes complementary base pairing with the mRNA “codon.” This codon-anticodon interaction, which is dictated by the specific nucleotide sequence on the mRNA (the genetic code), aligns anticodon-specific amino acids that subsequently form a growing polypeptide chain through covalent peptide bonding.

Polypeptide synthesis on the ribosome may be subdivided into initiation, elongation, and termination phases. First, elongation factors (eg, eIF4F) recognize the 5′ cap of mRNA, recruit other components of the initiation complex, and subsequently bind to the rRNA on the small (40S) ribosomal subunit. The initiation complex (40S ribosome, elongation factors such as eIF4, initiator tRNA, etc) rolls along the mRNA sequence until it reaches the “start codon” (AUG). Subsequently, polypeptide synthesis starts with disintegration of the initiation complex and recruitment of the 60S ribosome along with other elongation factors. As the ribosome continues moving along the mRNA genetic code, successive amino acid–attached tRNAs are recruited, and the polypeptide chain grows until a “stop codon” (UAA, UAG, or UGA) signals release of the finished product. Current evidence suggests that the enzymatic activity required to form peptide bonds between adjacent incoming amino acids may reside in the rRNA, which functions as a ribozyme.

Many newly synthesized proteins are further modified after or during polypeptide synthesis. Posttranslational or cotranslational modifications of proteins ensure proper protein folding, targeting, activation, and stability. The specific processes include protein translocation, glycosylation, phosphorylation, acylation, metal binding, and assumption of secondary and tertiary structures. Chaperones are enzymes that guide proper protein folding by binding and releasing misfolded proteins. Damaged or surplus proteins are removed through various mechanisms, including lysosomal and nonlysosomal protein degradation. Nonlysosomal protein degradation is facilitated by ubiquitination (a process of tagging proteins with ubiquitin to remove them by proteolysis). Ubiquitin is a small protein (76 amino acids long) that binds and marks damaged or excess proteins for subsequent degradation by the proteasome complex. The proteasome is a multiunit protease complex that degrades ubiquitin-tagged proteins. Ubiquitination also influences signal transduction and apoptosis by facilitating controlled degradation of pertinent regulatory molecules.

GENOMICS-RELATED LABORATORY METHODS

Polymerase Chain Reaction

Molecular cloning or DNA amplification often involves synthesis of multiple copies (clones) of a DNA sequence of interest (the target DNA). This may be accomplished by either inserting the target DNA into a bacterium that is capable of extended cell division (recombinant DNA technology) or using the polymerase chain reaction (PCR) for in vitro cloning of DNA (Figure 6). Polymerase chain reaction is an in vitro method of replicating relatively small DNA sequences into millions of copies over a short period.

A typical PCR reaction requires (1) two oligonucleotide primers (PCR primers), (2) a thermally stable DNA polymerase, (3) an ample amount of free deoxynucleotides, and (4) a small amount of DNA (the sample) that contains the sequence of interest (ie, the region to be amplified). The PCR primers (short DNA sequences of approximately 20 bp in length) are synthesized in the laboratory, and their nucleotide sequences are designed to be complementary to DNA sequences that flank (lie on either side of) the DNA fragment of interest (the target). Therefore, one has to know the nucleotide sequence of the target DNA fragment to use PCR technology. Because DNA synthesis can proceed only in the 5′ to 3′ direction, a PCR primer allows extension of DNA in only 1 direction (ie, the other side is a dead end). Therefore, to replicate both strands of the DNA molecule (the complementary strands), 2 PCR primers (forward and reverse primers) are needed, and they are placed on either side of the DNA sequence of interest on complementary strands (ie, the forward primer on 1 strand of DNA and the reverse primer on the second strand of DNA that is complementary to the first strand). This approach allows amplification of the double-stranded target sequence that is located between the 2 primers.

At the beginning of PCR, the sample DNA is first heated to a temperature greater than 90°C to separate the 2 paired DNA strands (DNA denaturation). The reaction is then cooled to approximately 40°C, in the presence of excess primers, to allow annealing (hybridization) of the specific primers to their complementary sequences on the target DNA (which is now in single strands). After the primers are bound to the DNA sequences that are immediately adjacent to the target sequence, new complementary strands of DNA, which are extensions of the bound PCR primers, are formed by recruiting the free nucleotides in the reaction. This step of primer extension occurs in an intermediate temperature of approximately 70°C with the help of a thermally stable DNA polymerase (an enzyme originally derived from the bacterium Thermus aquaticus). The newly synthesized DNA is double stranded, starting at the 5′ end with 1 of the 2 primers (Figure 6). After this first cycle of PCR, the same process is repeated 20 to 30 times (20-30 heating and cooling cycles), and each cycle results in the doubling of the target DNA region, with a final theoretical target amplification of approximately 1 million-fold or more.
Figure 6. Amplification of DNA using polymerase chain reaction technique. See text for further details. Copyright 1998 from Alberts et al. Reproduced by permission of Garland Publishing, Inc, part of The Taylor & Francis Group.
RNA species, among a mixture of RNAs, is first size-separated by gel electrophoresis. The RNA is subsequently transferred to a membrane where hybridization occurs with either radioactively or nonisotopically labeled DNA probes. A nuclease protection assay uses single-strand–specific endonucleases to remove by digestion single- but not double-stranded DNA. In the particular assay, the sample RNA is incubated with excess labeled, single-stranded DNA molecules that are complementary to the specific mRNA. The subsequent hybridization between the mRNA and the labeled DNA strands protects the specific mRNA from being digested with the single-strand–specific endonucleases. In both Northern blotting and nuclease protection assays, the amount of steady-state mRNA level is estimated from the density of the labeling molecule.

In RT-PCR, traditional PCR is combined with reverse transcription to amplify mRNA. In reverse transcription, mRNA (sense) in the test sample is converted to a single-stranded cDNA (antisense) in a reaction mediated by reverse transcriptase. Reverse transcriptase (a retroviral enzyme) is an RNA-dependent DNA polymerase that can synthesize a DNA sequence that is complementary to a specific mRNA. The synthesized cDNA represents only the exons of the corresponding genomic DNA. The subsequent PCR amplification method is called reverse transcription-polymerase chain reaction to distinguish it from the standard PCR used to amplify genomic DNA. Reverse transcription-PCR is an extremely powerful and sensitive tool to detect and quantify mRNA (ie, gene expression) from minute amounts of test samples.

In RT-PCR, mRNA quantification may be performed both in relative (relative RT-PCR) and in absolute (competitive RT-PCR) terms. In relative RT-PCR, the amount of the target mRNA is estimated in comparison with a coamplified internal control. The resultant ratio is then compared across different samples. In competitive RT-PCR, a known quantity of a synthetic RNA molecule, which is similar in sequence but shorter compared with the target mRNA, is added to the reaction. This results in the synthesis of 2 cDNA templates (target and synthetic) that will compete for the same primers during PCR amplification. As a result, the abundance of the target PCR product is inversely proportional to the amount of the synthetic RNA that is added to the reaction. Therefore, by performing a series of PCRs with addition of serial dilutions of the synthetic RNA, a concentration curve can be constructed that is used to estimate the quantity of the endogenous target product.

Recent medical uses of RT-PCR include detection of lymph node micrometastasis in prostate cancer, breast cancer tumor contamination of bone marrow, viral contamination of vaccines, and monitoring of minimal residual disease in acute leukemia.

Real-Time PCR.—In RT-PCR, the steady-state level of a specific mRNA is indirectly estimated from the level of the corresponding amplified product. The amplified product is usually quantified after the reaction is completed (ie, “end point” quantification). Alternatively, the presence and quantity of PCR products may be monitored in “real time” as the reaction progresses. Monitoring the abundance of PCR products in multiple cycles of PCR provides dynamic range values that are more informative than end point measurements. Of note, target amplification during the initial cycles of PCR is exponential because the presence of excess primers in the reaction minimizes renaturation of the complementary strands of the target DNA. As the concentration of an amplified product increases, the possibility of renaturation of its complementary strands increases. As a result, the amplification rate decreases and plateaus subsequently. The transition point from the exponential to the plateau phase of PCR may vary markedly for different samples. Therefore, it is more accurate to quantify the specific mRNA products of PCR during the exponential phase of target amplification. Real-time PCR accomplishes this task by quantifying reaction products in every cycle of PCR.

Real-time PCR depends on the detection and quantification of fluorescent molecules that are bound to the PCR product. Specific binding of fluorescent molecules to the PCR product is achieved by several methods, including TaqMan and SYBR green. The TaqMan probe is a dual-labeled oligonucleotide probe that contains a fluorescent dye at its 5′ base and another molecule (a quencher) at its 3′ base. The quencher absorbs (quenches) fluorescence when it is situated close to the fluorescent molecule. This phenomenon is called fluorescence resonance energy transfer (FRET). Therefore, the probe itself does not fluoresce. However, the fluorescent molecule is separated from the quencher molecule during primer extension through the exonuclease activity of Taq polymerase (the specific polymerase used in this particular PCR), whereby fluorescence is unmasked. Accordingly, the degree of fluorescence increases with an increase in the level of PCR product. Other FRET-dependent methods include molecular beacons and scorpions. In contrast, the SYBR green method is not FRET dependent. SYBR green is a dye that emits fluorescence only when it binds to double-stranded DNA. Therefore, the fluorescence signal is enhanced with continued synthesis of the PCR product.

A discussion on the quantification of the signals obtained from real-time PCR is beyond the scope of this review, and the reader is referred to recent overviews on this subject. Potential applications of real-time PCR
include disease diagnosis and monitoring, biosafety and genetic stability testing, mutation detection, quantification of cytokine profiles in immune cells or inflamed tissue, and in vivo imaging of cellular processes.

Electrophoresis and Blotting

DNA, RNA, or proteins may be separated by size with use of agarose or polyacrylamide gels through electrophoresis. DNA and RNA are uniformly, negatively charged, and under most circumstances their gel separation is a function of their size and topology (linear fragments move in a different manner from supercoiled fragments of the same length). In contrast, proteins must first be denatured (unfolded) and negatively coated with an ionic detergent (sodium dodecyl sulfate) to allow gel separation solely by size, avoiding disturbances from their inherent positive and negative charge. Isoelectric focusing separates molecules by their net (positive or negative) charge.

In Southern blotting (named after Dr Edward Southern, who described the technique), DNA is first cut into pieces using restriction endonucleases (enzymes that digest DNA at specific sites that are marked by a 4- to 8-member specific nucleotide sequence), and then the restriction fragments are subjected to agarose gel electrophoresis (Figure 7). The double-stranded DNA fragments on the gel are then turned into single strands by denaturation and transferred to a DNA-binding membrane, such as nitrocellulose or nylon, to make a permanent copy of single-stranded DNA. A DNA sequence of interest is then visualized by a radiolabeled reporter probe after a hybridization step followed by autoradiography. In addition to target identification and quantification, Southern blotting may allow detection of mutations that result in alterations of restriction fragment lengths (ie, detect restriction fragment length polymorphisms [RFLPs]). Larger DNA molecules are separated by pulsed-field gel electrophoresis (interruption application of voltage gradients to disorient large DNA molecules and optimize separation by size only). 

Northern blotting (named to reflect similarity to Southern blotting) is used to identify specific RNAs. Western blotting is used to identify specific proteins that have been separated by size with use of polyacrylamide gel electrophoresis.

Cutting, Pasting, and Altering the Topology of DNA

Different enzymes specialize in digesting and twisting (and untwisting) the DNA molecule. Endonucleases cleave DNA (break the sugar-phosphate ester bond between 2 deoxynucleotides) from the middle, while exonucleases sequentially remove nucleotides from the ends of DNA strands. DNA polymerases often possess exonuclease activity as part of their proofreading capability in addition to their synthetic function. For many laboratory experiments, it is useful to remove the exonuclease activity of Escherichia coli DNA polymerase I by brief proteolytic digestion, leading to a synthetically active DNA polymerase called the Klenow fragment. Restriction endonucleases act at specific DNA sites, which are marked by precise 4- to 8-nucleotide sequences. Such restriction digestion of DNA is the key to recombinant technology and

Figure 7. Southern blotting. See text for further details. Reproduced with permission from Lodish et al.
chromosome mapping. Ligases attach 2 separate DNA fragments (reestablish the sugar-phosphate bond) and are important in recombinant technology.218

DNA is naturally highly compacted but may take a relaxed (α-helical) or even a supercoiled state (a coiling and double coiling of the α-helical thread). A positive supercoil tends to tighten further and supercoil the neighboring double helix, and a negative supercoil tends to unwind the neighboring double helix. Supercoiling and relaxing of DNA are often mediated by topoisomerases.219,220 Topoisomerases alter the topology (degree of coiling) of DNA by creating transient, either single (topoisomerase I) or double (topoisomerase II) strand breaks and allowing DNA to wind or unwind through those breaks. DNA gyrase is a type of topoisomerase that relieves supercoiling.221

Clones, Libraries, and Polymorphisms

Clones.—Any DNA can be amplified (mass and strand number increased) in the laboratory by using either recombinant technology or PCR. Recombinant cloning starts with preparation of restriction fragments of DNA (eg, from human cells) that are then fused (using DNA ligase) to vector DNA (a carrier DNA molecule capable of self-replication) that has been digested with the same restriction enzymes so that they have similar split ends that allow DNA ligation.222 This “recombinant” DNA molecule is transported and amplified by the vector (a carrier molecule capable of self-replication) (Figure 8). Vectors in current use include plasmids (circular double-stranded DNA molecules that replicate extrachromosomally in bacteria), bacteriophages (bacterial virus with a larger than plasmid double-stranded DNA molecule), cosmids (large plasmids capable of carrying large DNA inserts), and bacterial artificial chromosomes (BACs) or yeast artificial chromosomes (YACs) (artificially constructed chromosome-like structures that carry very large DNA fragments to be amplified [cloned]).29,223-226 Specifically, plasmids and bacteriophages are used for the insertion of small foreign DNA (<20 kb); BACs and YACs are used for larger DNA molecules (300-1000 kb). Because YACs can carry large DNA fragments, they have been particularly important in the mapping of the human genome.227

Libraries.—Chromosomal DNA isolated directly from a cell is called genomic DNA. DNA made from reverse transcription of mRNA is called complementary DNA. A set of DNA clones from a particular source is called a DNA library (genomic or cDNA library depending on the source of the clones) (Figure 9). To prepare a human DNA library, genomic DNA or cDNA is first digested with restriction enzymes, and the restriction fragments (inserts) are recombined with a similarly digested vector DNA. A population of recombinant DNA molecules with different human DNA inserts is then cloned in a bacterial host. A collection of thousands of vector-linked human DNA inserts represents the human DNA library.228

Complementary DNA libraries have the advantage of carrying only exons (protein-encoding regions of genes) and may be modified by flanking the inserts with transcription and translation signals (expression vectors), thus allow-
Complementary DNA clones representing named and unnamed genes may be partially sequenced to generate expressed sequence tags (ESTs). Although incomplete, ESTs provide a snapshot of the expressed sequences in a given cell type and also can be used to identify gene location in genomic DNA sequences. Further sequence information on the identified gene can then be obtained by chromosome walking (described subsequently). A section of a DNA sequence that starts with an initiation codon, which is a recognition site to start protein synthesis (translation), and ends in a stop codon (a codon that signals termination of translation), is called an open reading frame, suggesting the potential to code for a protein.

Polymorphisms.—The human genome is extensively affected by sequence variation. When the rate of variation at a specific point in the DNA yields a variant sequence that is found in more than 1% of a given population, it is referred to as a polymorphism. When the incidence of a variant sequence is less than 1%, it is referred to as a mutation. However, such definitions are imprecise because what may be considered a mutation in a given population may qualify as a polymorphism in another population (eg, sickle cell mutation).

Genetic polymorphisms may be revealed by restriction enzyme digestion that results in variably sized restriction fragments, a phenomenon known as restriction fragment length polymorphism (Figure 10). Usually occurring in intergenic DNA regions, RFLPs generally have no phenotypic effect. Certain regions of DNA are highly polymorphic because of the presence of variable numbers of tandem repeats (VNTRs). The latter are short DNA sequences that are repeated in sequence multiple times. Variations among individuals in the number and size of such repeat sequences may be revealed by using RFLPs induced by restriction enzymes that cut near the boundaries that encompass the repeat sequences. The degree of polymorphism in VNTRs is sufficiently high to allow DNA fingerprinting.

Single nucleotide polymorphisms occur as often as every 200 bp in the human genome. Single nucleotide polymorphisms may be detected by microarrays and have the potential to be used in studies of genetic susceptibility to diseases. Of note, however, only about 1% of single
nucleotide polymorphisms result in an alteration in protein coding at the amino acid level.

DNA Sequencing

DNA sequencing entails spelling out the nucleotide chain of a DNA region of interest. Sufficient quantities of the target DNA region of interest are obtained from either a bacterial-cloned fragment or a PCR-amplified sequence. A short DNA primer is used to start the synthesis of a complementary strand to the original template to be sequenced. With the use of specific chemical analogues of nucleotides (dideoxynucleotides), which are capable of terminating chain extension at specific nucleotide bases, the synthesis of the complementary strand is halted (nested) at specific nucleotide bases, resulting in the synthesis of DNA fragments with varying lengths that differ by 1 nucleotide (Figure 11). The set of nested fragments that are radiolabeled or tagged with fluorescent dyes can then be separated by gel electrophoresis, resulting in a pattern that allows the determination of the specific nucleotide at the end of each successive fragment. Currently, an automated computerized laser detection system is used for rapid accrual of sequence information.

Gene Mapping

Human genetic mapping entails (1) the localization or mapping of genes to each of the 23 chromosome pairs and (2) the determination of the order and spacing of the gene on the particular chromosome. This is done in 1 of 2 ways: physical mapping (based on estimates of physical distance measured in bp) or genetic linkage mapping (based on the frequency of meiotic crossing over that is observed between 2 loci of a chromosome). Linkage distance is measured in centiMorgans (cM), named for Thomas H. Morgan, the father of Drosophila (fruit fly) genetics, who defined the concept of genetic linkage. Locating a disease gene is the first step toward cloning the gene itself.

Physical Mapping—Physical mapping of genes may be achieved by different methods with differing levels of resolution that range from an entire chromosome to a single base pair (the base sequence). Methods of low-resolution physical mapping include karyotyping, fluorescence in situ hybridization (FISH), and somatic cell hybridization.

Human chromosomes may be identified by size and by distinctive banding patterns seen under light microscopy after staining with special DNA-binding dyes (Giemsa). This method of chromosome visualization is called karyotyping and provides a metaphase picture of the 22 homologous pairs of autosomal chromosomes and the 1 additional pair of sex chromosomes lined up by size. Such a chromosome picture allows identification of gross abnormalities (chromosomal deletions, duplications, translocations) that may be associated with a particular disease phenotype (eg, chronic myeloid leukemia is associated with a gross translocation between chromosomes 9 and 22). As a result, the disease gene is physically mapped to the vicinity of the abnormal karyotypic lesion.

In FISH, a priori knowledge of a DNA sequence of the specific gene or of a marker near the gene is required. Fluorescent-tagged probes, by cloning of the specific sequence, are then prepared and used to localize the gene of interest to a particular chromosome by slide exposure of metaphase or interphase chromosomes with the labeled probe.

In somatic cell hybridization, human and nonhuman cells are fused under specific experimental conditions.
Subsequently, the hybrid cells are allowed to replicate under specific conditions whereby they start losing a variable number of the human chromosomes. Eventually, hybrid cells that have retained 1 or 2 specific human chromosomes are selected for further testing. Karyotyping allows distinction of human from nonhuman (usually mouse) chromosomes and identification of the specific human chromosome that is retained in a hybrid cell. Additional experiments are then performed to look for the presence or absence of a specific DNA sequence or protein product in a given hybrid cell, and this would allow the assignment of the gene for a particular protein product to a specific chromosome. Somatic cell hybrids may provide additional resolution when the hybrid is constructed from cells containing either structurally abnormal or radiation-damaged human chromosomes (so-called radiation hybrids). Similarly, somatic cell hybridization and FISH techniques may be combined for improved resolution.

Other physical maps include cDNA maps that localize the chromosomal positions of expressed genes, RFLP maps of restriction cleavage sites, and contig maps of overlapping sequences of YACs, BACs, or cosmids (Figure 12) (described in more detail subsequently). The resolution grade of karyotypic identification of a genetic
Genetic Linkage Mapping.—Genes that are located on different chromosomes segregate independently during meiosis (the law of independent assortment). In other words, 2 genes on 2 different chromosomes have a 50% chance of being inherited together in a daughter cell. In contrast, genes on the same chromosome have a tendency to be inherited together (ie, are genetically linked).25 However, the chance of segregating together is not 100% because of the “phenomenon” (process) of recombination (crossing over).252-256 During recombination, the naturally occurring exchange of chromosome regions between homologous pairs of chromosomes happens during meiosis.257 The chance for a recombination event to occur depends on the distance between the 2 genes on the same chromosome.258 The closer the genes are to each other, the less chance for a recombination event to occur and vice versa.

Genetic linkage is defined as the tendency for genes on the same chromosome to stick together during meiosis (Figure 13). The strength of this linkage is used as a unit of measurement (Morgan) of the distance between the 2 linked genes.259 One centiMorgan denotes a recombination event of 1% (ie, the genes stick together 99% of the time during meiosis). A genetic distance of 1 cM therefore denotes 2 genes that are relatively close to each other. Accordingly, a genetic linkage map may be perceived as a map of a chromosome with genes or DNA markers that are aligned in a linear fashion with intervening distances that reflect the frequency of recombination between the adjacent genes or markers.252 The total genetic length of the human genome is estimated at 3000 cM. As such, 1 cM corresponds to about 1 million bp.

Of note, chromosomal regions (DNA sequences) that are evaluated for linkage analysis may not be associated necessarily with a phenotypically apparent trait. For example, hair color is a phenotypically apparent trait, and variation in the DNA sequence responsible for hair color is reflected by variation in hair color. In contrast, DNA sequence variation of a phenotypically silent genetic marker is revealed by molecular techniques that exploit the presence or absence of specific restriction enzyme cleavage sites (RFLPs) and variations in the number of tandem repeats at specific chromosomal regions (VNTRs).260 Regardless of whether the study markers are phenotypically apparent (genes) or not (RFLPs, VNTRs), linkage analysis requires that heterozygosity exists at the locus of both the gene and the genetic marker (ie, alleles of the specific gene or DNA sequence must be heterozygous).261

Alleles are variants of the same gene occupying the same locus on chromosomes. Therefore, a homologous pair of chromosomes may either have the same allele on the 2 chromosomes forming the pair (ie, are homozygous for the specific allele) or have 2 different alleles at a specific locus (ie, are heterozygous). Genetic mapping methods are based on the observed segregation of alleles on a homologous pair of chromosomes. For example, if 2 separate traits are inherited together 99% of the time, then the alleles responsible for the traits are probably on the same chromosome at a distance that is 1 cM between each other.

Precise Localization and Sequencing of the Gene of Interest.—In most diseases, a causal genetic aberration is not yet known. As described previously, one can use linkage analysis to associate a putative disease gene with a...
chromosomal locale that is detectable either molecularly or phenotypically.\textsuperscript{260} Once the gross location of the putative disease gene is determined by linkage analysis, one can apply positional cloning for more precise identification and sequencing of the gene of interest.\textsuperscript{262,263}

Positional cloning starts with the selection, by using the DNA sequence of the linked marker as a probe, of vector clones from a genomic library that carry DNA sequences that overlap with those from the linked genetic marker. As mentioned previously, a genomic library is constructed by partially digesting the entire genome with restriction enzymes, followed by cloning of the resultant fragments into appropriate vectors.\textsuperscript{264} Because the DNA products of restriction digestion are relatively large, the vectors usually used are YAC and BAC vectors. Using the DNA sequence of the linked marker, one can pick YAC or BAC clones with sequences that overlap with those of the linked marker.\textsuperscript{265} Alternatively, sequence tagged sites, which are unique, relatively short DNA sequences with known chromosomal location, may be used to identify clones with overlapping sequences. Subsequently, the overlapping sequences themselves may be used as probes to select additional YAC or BAC clones with sequences that are partially overlapping with the probes used.\textsuperscript{239,266}

The stepwise process of identifying YAC or BAC clones based on overlapping DNA sequences is called \textit{chromosome walking}.\textsuperscript{267} This process of chromosome walking continues until the disease gene is identified. The linear alignment of YACs or BACs with overlapping sequences results in a continuous DNA segment called a \textit{contig map}. Such contig maps have been essential for the analysis of human and nonhuman genomes as well as for specific chromosomal regions.\textsuperscript{255,257,264,269}

Probe-identified clones can also be individually mapped for sites complementary to ESTs. An EST that cosegregates with the gene of interest is determined and then sequenced. Alternatively, the clone containing the linkage markers may have already been sequenced, and resident genes are sought by looking for ORFs. Finally, the candidate gene is identified by mutational analysis that compares ORFs from a patient-derived DNA library to those obtained from nonaffected individuals.

\section*{CONCLUSION}

This review on the basic principles of molecular biology is not intended to be comprehensive but to serve as an introduction to this subject. Additional information is easily accessible from numerous outstanding textbooks of molecular biology.\textsuperscript{30,121,270}

\section*{REFERENCES}


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**Glossary of Terms Frequently Used in Medical Genomics**

**Allele**—One of 2 or more alternative forms of a DNA sequence; eg, most persons have 2 functional (normal) alleles of the phenylalanine hydroxylase gene, whereas carriers of phenylketonuria have 1 functional allele and 1 mutant allele.

**Alternative splicing**—Process by which different messenger RNAs (mRNAs) are produced from the same primary transcript, through variations in the splicing pattern of the transcript.

**Alu element**—Short repetitive DNA sequence that is scattered throughout the genome and is identified by a restriction enzyme (Alu 1 endonuclease) recognition site near the middle of the sequence.

**Autosomes**—Chromosomes not involved with sex differentiation (1-22 in humans).

**Bacteriophage**—Virus that infects bacteria, commonly known as a phage.

**Gene bank**—Collection of recombinant DNA molecules containing inserts that together comprise the entire genome of an organism.

**CentiMorgan (cM)**—Distance between DNA loci as determined for forensic purposes. DNA from blood, semen, or other tissue sample is isolated and cleaved with a restriction endonuclease, the restriction fragments that are unique to an individual, especially in blood, semen, or other tissue.

**Centromere**—Heterochromatic-constricted portion of a chromosome where the chromatids are joined.

**Chromosome**—Self-replicating nucleic acid molecule containing a number of genes. In bacteria, the entire genome is contained within 1 double-stranded (ds), circular DNA chromosome. In eukaryotes, chromosomes are linear DNA duplexes, and most organisms have genomes divided between a number of such chromosomes, that number being characteristic for a particular species.

**Chromosome walking**—Method used to identify which clone in a gene bank contains a desired gene or sequence that cannot be selected for easily. The gene bank must contain the entire DNA sequence of the chromosome as a series of overlapping fragments. These fragments can be generated either by random shear or by partial digestion with a 4-base pair [bp] cutter such as Sau 3A. A series of colony hybridizations is then carried out, starting with same cloned gene that has already been identified and that is known to be on the same chromosome as the desired gene. This identified gene is used as a probe to pick out clones containing adjacent sequences; these are then used as probes themselves to identify clones carrying sequences adjacent to them, etc. At each round of hybridization one “walks” further along the chromosome from the identified gene.

**Clone**—Colony of cells, all descended from a single cell, therefore genetically homogeneous; also the act of preparing and propagating a genetically defined cell. This term is used in a number of senses. As a noun it may mean (1) a population of recombinant DNA molecules all carrying the same inserted sequence or (2) a population of cells or organisms of identical genotype. It is most frequently used to describe a colony of microorganisms that harbors a specific DNA fragment inserted into a vector molecule. As a verb, “to clone” means to use in vitro recombination techniques to insert a particular gene or other DNA sequence into a vector molecule.

**Coding sequence**—Portion of a gene that directly specifies the amino acid sequence of its protein product. Noncoding sequences of genes include control regions, such as promoters, operators, and terminators, as well as the intron sequences of certain eukaryotic genes.

**Complementary**—Nucleic acid strand is said to be complementary to another strand if it is able to form a perfectly hydrogen-bonded duplex with it, according to the Watson-Crick rules of base pairing, which include the antiparallel alignment of the strands 3' to 5' at each end of the duplex. Example: An mRNA molecule is complementary to one of the DNA strands of the gene that encodes it. The RNA sequence 5'-AUGGCAUUCGCCCCACUGA-3' is complementary to the DNA sequence 3'-TACCGTAAAGCAGGGGTAGCT-5'.

**Complementary DNA (cDNA)**—DNA complement of an RNA sequence; it is synthesized by the enzyme RNA-primed DNA polymerase of reverse transcriptase. The single-stranded DNA product of this enzyme (the reverse transcript) may be converted into the double-stranded form by DNA-primed DNA polymerase (DNA polymerase) and inserted into a suitable vector to make a cDNA clone; cDNA cloning is commonly used to achieve expression of mammalian genes in bacteria or yeast. Since the cDNA clone is a copy of the mature RNA molecule, it contains no introns to act as a barrier to expression.

**Digestion**—Treatment of a substrate molecule with an enzyme preparation in which covalent bonds are hydrolyzed.

**DNA fingerprinting**—Method to generate a pattern of DNA restriction fragments that are unique to an individual, especially for forensic purposes. DNA from blood, semen, or other tissue is isolated and cleaved with a restriction endonuclease, the products being separated by polyacrylamide gel electrophoresis, and fragments containing a specific nucleotide sequence are detected by hybridization with an appropriate DNA probe to produce the unique pattern.

**DNA gyrase**—A type II topoisomerase of *Escherichia coli*, which is essential for DNA replication. This enzyme can induce or relax supercoiling.

**DNA polymerase**—Any enzyme that synthesizes DNA by copying a template strand. DNA polymerases synthesize DNA in the 5’ to 3’ direction by successively adding nucleotides to the free 3’ hydroxyl group of the growing strand. The template strand determines the order of addition of nucleotides via Watson-Crick base pairing. *E. coli* has 3 DNA polymerases: polymerase I is involved in repair synthesis, polymerase III is responsible for DNA replication, and the function of polymerase II is unknown at present (see DNA polymerase I, nick translation, Okazaki fragment). Animal cells have α, β, γ, and δ polymerases, with α and δ apparently responsible for replication of nuclear DNA and γ for replication of mitochondria. All these functions with a DNA strand as a template. Retroviruses possess a unique DNA polymerase (reverse transcriptase) that uses an RNA template.

**DNA topoisomerase**—Enzyme capable of altering the linking number of DNA, usually measured by following a change in the degree of supercoiling of dsDNA molecules. Various topoisomerases can increase or relax supercoiling, convert single-stranded rings to intertwined double-stranded rings, tie and untie knots in single-stranded and duplex rings, and catenate and decatenate duplex rings. Topoisomerase II of *E. coli* = gyrase.

**Endonuclease**—One of a large group of enzymes that cleave nucleic acids at positions within the chain. Some act on both RNA and DNA (eg, S1 nuclease, EC 3.1.30.1, which is specific for single-stranded molecules). Ribonucleases such as pancreatic, T1, etc, are specific for RNA, deoxyribonucleases for DNA. Bacterial restriction endonucleases are crucial in recombinant DNA technology for their ability to cleave dsDNA at highly specific sites.
Condensed chromatin in certain chromatin-specific gene.

Euchromatin—In cytology, the lightly staining regions of chromosomes that contain less-condensed chromatin and that are the regions transcribed as RNA (see heterochromatin).

Exon—From “expressed,” a region of a eukaryotic gene that encodes protein, as opposed to an intron, ie, a region of the gene that does not.

Expressed sequence tags (ESTs)—Known DNA sequences from gene segments expressed as RNA that can be used to clone and characterize these genes.

Fluorescence in situ hybridization (FISH)—Nonisotopic method to label DNA probes for in situ hybridization. The ability to use multiple fluorochromes in the same reaction increases the utility of this procedure. The resolving power of FISH is further enhanced if interphase chromosomes are studied.

Genomic library—Collection of cloned DNA fragments from a single genome. Ideally, the bank should contain cloned representatives of all the DNA sequences in the genome. For large genomes, such banks are made by shotgun cloning.

Genetic map—Indirect measure of distance, constructed by determining how frequently 2 markers (DNA polymorphisms, physical traits, or syndromes) are inherited together. Distances in genetic maps are measured in cM (see physical map).

Genomic library—Collection of transformed cells, each of which contains DNA fragments, the entire population representing the total genome of an organism, eg, a rat library containing which contains DNA fragments, the entire population representing the total genome of an organism, eg, a rat library containing which contains DNA fragments, the entire population representing the total genome of an organism, eg, a rat library containing which contains DNA fragments, the entire population representing the total genome of an organism, eg, a rat library containing which contains DNA fragments, the entire population representing the total genome of an organism, eg, a rat library containing.

Gene bank—Collection of cloned DNA fragments from a single genome. Ideally, the bank should contain cloned representatives of all the DNA sequences in the genome. For large genomes, such banks are made by shotgun cloning.

Genetic map—Indirect measure of distance, constructed by determining how frequently 2 markers (DNA polymorphisms, physical traits, or syndromes) are inherited together. Distances in genetic maps are measured in cM (see physical map).

Histones—Proteins found in the nuclei of all eukaryotic cells where they are complexed to DNA in chromatin and chromosomes. They are of relatively low molecular weight and are basic, having a very high arginine-lysine content; highly conserved and are extensively processed to give the mRNA which is exported to the cytoplasm where protein synthesis takes place. This processing includes the addition of a 5′-5′-linked 7-methylguanylate “cap” at the 5′ end and a sequence of adenylnucleotides at the 3′ end (the poly-A tail), as well as the removal of any introns and the splicing together of exons. Only 10% of hnRNA leaves the nucleus. Eukaryote mRNAs are comparatively long lived, with half-life ranging from 30 minutes to 24 hours.

Microsatellites—Same as for minisatellites except that the polymorphism allele size is smaller, eg, <1 kilobase, and the basic core repeat unit involves a 2- to 4-nucleotide bp repeat motif; also known as SSRs (simple sequence repeats). One example is repeats of the motif CA, eg, CACACACACAC, which is also written as AC (CACACACACAC, etc). Terminology is confusing because these repeats are identical.

Minisatellites—Repetitive DNA consisting of small numbers of repeating nucleotides in a tandem configuration, used for genetic linkage and DNA fingerprinting studies.

Northern blotting—Procedure analogous to Southern blotting in which RNA, not DNA, is transferred or blotted from a gel to a suitable binding matrix, such as a nitrocellulose sheet.

Nucleic acid—DNA or RNA molecule that can be single stranded or double stranded. When very small, it is an oligonucleotide.

Nucleotide—Repeating structural unit of chromatin that consists of a complex of 8 molecules of histones and a DNA double helix wrapped twice around them.

Okazaki fragment—Short fragments of newly synthesized DNA strands produced during DNA replication. All the known DNA polymerases can only synthesize DNA in 1 direction, the 5′ to 3′
direction. However, as the strands separate, replication forks will be moving along 1 parental strand in the 3’ to 5’ direction and 5’ to 3’ on the other parental strand. On the former, the leading strand, DNA can be synthesized continuously in the 5’ to 3’ direction. On the other, the lagging strand, the DNA synthesis can occur only when a stretch of single-stranded DNA has been exposed and proceeds in the direction opposite to the movement of the replication fork (still 5’ to 3’). Thus, it is discontinuous, and the series of fragments are then covalently linked by ligases to give a continuous strand. Such fragments were first observed by Okazaki using pulse-chase labeling with radioactive thymidine. In eukaryotes, Okazaki fragments are typically a few hundred nucleotides long, whereas in procaryotes they may contain several thousands of nucleotides.

**Oligonucleotides**—Small single-stranded segments of DNA, typically 20 to 30 nucleotide bases, that are synthesized in vitro. Uses include DNA sequencing, DNA amplification, and DNA probes.

**Open reading frame (ORF)**—A reading frame uninterrupted by stop codons. The existence of ORFs is usually inferred from the DNA (rather than the RNA) sequence.

**Polyacrylamide gel electrophoresis (PAGE)**—Method for separating nucleic acid or protein molecules based on their molecular size. Molecules migrate through the inert gel matrix under the influence of an electric field. For protein PAGE, detergents such as sodium dodecyl sulfate are often added to ensure that all molecules have a uniform charge. Secondary structure often leads to anomalous migration of molecules. Thus, it is common to denature protein samples by boiling them before PAGE. For nucleic acids, denaturing agents such as formamide, urea, or methyl mercuric hydroxide are often incorporated into the gel itself, which may also be run at high temperature. PAGE is used to separate the products of DNA-sequencing reactions, and the gels used are highly denaturing because molecules differing in size by a single nucleotide must be resolved.

**Polymerase**—Enzyme that will form oligomeric molecules from monomers. A DNA polymerase will synthesize DNA from deoxyribonucleoside triphosphates by using a cDNA strand and a primer. An RNA polymerase will synthesize RNA from monoribonucleoside triphosphates and a cDNA strand.

**Polymerase chain reaction (PCR)**—Technique to amplify a specific region of dsDNA. An excess of 2 oligonucleotide primers complementary to 2 sequences that flank the region to be amplified is annealed to denatured DNA and subsequently elongated, usually by a heat-stable DNA polymerase from *Thermus aquaticus* (Taq polymerase). Each cycle involves heating to denature dsDNA and cooling to allow annealing of excess primer to template and elongation of the primers by the Taq polymerase; the number of target sequence fragments between flanking primers doubles with each cycle.

**Primary structure**—Amino acid sequence of a protein or the nucleotide sequence of a polynucleotide.

**Primer**—DNA polymerase, unlike RNA polymerase, is unable to initiate the de novo synthesis of a polynucleotide chain; it can only add nucleotides to a free 3’-hydroxyl group at the end of a preexisting chain. A short oligonucleotide, known as a primer, is needed to supply such a hydroxyl group for initiation of DNA synthesis; RNA primer is used for initiation of DNA replication in vivo.

**Probe**—As a noun, a probe is a specific DNA or RNA sequence that has been radioactively labeled to a high specific activity. Probes are used to detect complementary sequences by hybridization techniques such as Southern or Northern blotting or colony hybridization. As a verb, “to probe” is the act of hybridization to detect a specific gene or transcript, e.g., “We probed our bank with labeled rRNA to detect clones containing rDNA sequences.”

**Processing**—Posttranscriptional modification of RNA or post-translational modification of a protein.

**Promoter**—DNA region, usually upstream to the coding sequence of a gene or operon, which binds RNA polymerase and directs the enzyme to the correct transcriptional start site.

**Quaternary structure**—Arrangement in space of polypeptide subunits that make up a multicrystalline protein.

**Recombination**—Crossing over (breakage and rejoining) between 2 loci, resulting in new combinations of genetic markers—traits at those loci, e.g., imagine that 1 locus has 4 genetic markers linearly arranged, a-b-c-d, and the second locus is b-b-c-a. Recombination involving these 2 regions between the b-c markers would give new genetic combinations, i.e., a-b-c-a and b-b-c-d.

**Recombination frequency**—Number of recombinants divided by the total number of progeny. This frequency is used as a guide in assessing the relative distances between loci on a genetic map.

**Restriction endonucleases**—Class of bacterial enzymes that cut DNA at specific sites. In bacteria their function is to destroy foreign DNA, such as that of bacteriophages (host DNA is specifically modified by methylation at these sites). They have very specific recognition and cutting sites. The recognition sites are short, 4 to 8 nucleotides, and are usually palindromic sequences. Because both strands have the same sequence running in opposite directions, the enzymes make double-stranded breaks, which, if the site of cleavage is off-center, generate fragments with short single-stranded tails; these can hybridize to the tails of other fragments and are called “sticky ends.” They are generally named according to the bacterium from which they were isolated (the first letter of genus name and the first 2 letters of the specific name). The bacterial strain is identified, and multiple enzymes from the same strain are given Roman numerals. For example, the 2 enzymes isolated from the R strain of *E coli* are designated Eco RI and Eco RII.

**Restriction fragment length polymorphism (RFLP)**—DNA polymorphisms deriving from changes in nucleotides at restriction sites, yielding DNA segments of different lengths after restriction and gel electrophoresis.

**Reverse transcriptase**—RNA-directed DNA polymerase; enzyme first discovered in Retroviridae, which can construct dsDNA molecules from the single-stranded RNA templates of their genomes. Can be inhibited by the drug zidovudine. Reverse transcription now appears also to be involved in movement of certain mobile genetic elements such as the Ty plasmid in yeast, in the replication of other viruses such as hepatitis B, and possibly in the generation of mammalian pseudogenes.

**Ribonucleases**—Phosphodiesterases that degrade RNA, very widely distributed.

**Ribonucleoprotein (RNP)**—Complexes of RNA and protein involved in a wide range of cellular processes. Besides ribosomes (with which RNP was originally almost synonymous), in eukaryotic cells both initial RNA transcripts in the nucleolus (hnRNA) and cytoplasmic mRNAs exist as complexes with specific sets of proteins. Processing (splicing) of the former is carried out by small nuclear RNPs. Other examples are the signal recognition particles responsible for targeting proteins to endoplasmic reticulum and a complex involved in termination of transcription.
Ribosomal protein—Protein present within the ribosomal subunits. In prokaryotes there are 31 to 34 proteins in the large subunit and 21 in the small subunit. Eukaryotic subunits have 45 to 50 (large subunit) and 33 (small subunits) proteins.

Ribosome—Subcellular complex responsible for protein synthesis. It consists of 2 subunits that both contain 1 or more RNA molecules (rRNAs) and a large number of proteins. The small subunit is responsible for binding mRNA and is subsequently joined by the large subunit, which accepts the aminoacyl tRNA molecule and carries out the process of peptide bond formation.

RNA polymerase—Enzyme that transcribes DNA into RNA; it is able to initiate RNA synthesis on a DNA template in the absence of any primer molecule. Prokaryotes have a single RNA polymerase that synthesizes all classes of RNA molecules. Eukaryotes have 3 RNA polymerases with different transcriptional specificities: RNA polymerase A (or I) synthesizes the large rRNA precursor, B (or II) synthesizes mRNA, C (or III) synthesizes tRNA and 5s rRNA species.

Satellite chromosome—Chromosome that seems to be an addition to the normal genome.

Satellite DNA—Polynucleotides that are separable (based on their characteristic density) from the bulk of nuclear DNA and that have repetitive sequences.

Sodium dodecyl sulfate (SDS) gel electrophoresis—Technique to dissociate multimeric proteins and to separate all proteins according to their apparent molecular weights; proteins are treated with SDS, an anionic detergent, and are separated by electrophoresis on polyacrylamide gels that incorporate SDS, the detergent adsorbing to the protein in proportion to its mass and dominating its electric charge.

Secondary structure—Regular folding of a protein in repeated patterns, eg, α-helix, β-pleated sheet, β-turns; also the double helix structure of a polynucleotide.

Sequence tagged site (STS)—Method to provide unambiguous identification of DNA markers generated by the Human Genome Project. STSs comprise short single-copy DNA sequences that characterize mapping landmarks on the genome.

Single nucleotide polymorphisms—Single nucleotide substitutions detected by DNA sequencing.

Somatic cell hybrid—Hybrid formed from the fusion of different cells. These usually come from different species, eg, human and rodent hybrids are frequently used for human gene mapping.

Southern blotting—Technique that combines the resolving power of agarose gel electrophoresis with the sensitivity of nucleic acid hybridization. DNA fragments separated in agarose gel are denatured in situ and then blotted or transferred, usually by capillary action, from the gel to a nitrocellulose sheet, or other binding matrix, placed directly on top of the gel. Single-stranded DNA binds to the nitrocellulose and is then available for hybridization with labeled, 32P or biotinylated, single-stranded DNA or RNA. Labeled nucleic acid is known as the probe and, in the case of DNA, is often prepared by nick translation. The hybrids are detected by autoradiography, in the case of 32P, or a color change, in the case of a biotinylated probe. This extremely sensitive and powerful technique, often described as “blotting,” is named after its inventor, E. M. Southern.

Tandem repeats—Small sections of repetitive DNA in the genome, arranged in head-to-tail formation.

Telomeres—Two ends of a chromosome.

Termination codon—Three codons, UAA known as ochre, UAG as amber, and UGA as opal, that do not code for an amino acid but act as signals for the termination of protein synthesis. They are not represented by any tRNA, and termination is catalyzed by protein release factors. There are 2 release factors in E coli: RF1 recognizes UAA and UAG, and RF2 recognizes UAA and UGA. Eukaryotes have a single GTP-requiring factor, eRF.

Topoisomerases—Enzymes that change the degree of supercoiling in DNA by cutting 1 or both strands. Type I topoisomerases cut only 1 strand of DNA; type I topoisomerase of E coli (omega protein) relaxes negatively supercoiled DNA and does not act on positively supercoiled DNA. Type II topoisomerases cut both strands of DNA; type II topoisomerase of E coli (DNA gyrase) increases the degree of negative supercoiling in DNA and requires ATP. It is inhibited by several antibiotics, including nalidixic acid and ovobioin.

Transfer RNA (tRNA)—Low-molecular-weight RNAs that specifically bind amino acids by aminoacylation and possess a special nucleotide triplet, the anticodon, sometimes containing the base inosine, by which they recognize codons on mRNA. By this recognition the appropriate tRNAs are brought into alignment. See text for further classification.

Variable numbers of tandem repeats (VNTRs)—DNA polymorphisms deriving from genomics regions with stretches of repeating nucleotides, yielding differently sized DNA segments after restriction and gel electrophoresis. Largest category of VNTRs are minisatellites including CA repeats.

Vector—General term applied to a DNA molecule, derived from a plasmid or bacteriophage, into which fragments of DNA may be inserted or cloned. The vector should contain 1 or more unique restriction sites for this purpose and be capable of autonomous replication in a defined host or vehicle organism, such that the clone sequence is reproduced. The vector molecule should confer some well-defined phenotype on the host organism, which is either selectable, eg, drug resistance, or readily detected, eg, plaque formation.

Western blotting—Technique used to detect specific proteins, a mixture of proteins being separated by PAGE, blotted onto nitrocellulose, and then exposed to a radiolabeled immunoglobulin directed to the desired protein, which is revealed by autoradiography (see Northern blotting, Southern blotting).