

ANALYSIS OF GENE EXPRESSION

1

Objectives

- Determination of RNA level
 - Northern blot
 - Microarrays
- Analysis of proteins
 - ELISA
 - Western blot
- Gene Sequencing and application

ANALYSIS OF GENE EXPRESSION

Determination of RNA levels

Northern blots

Microarrays

Analysis of proteins

Enzyme-linked immunosorbent assays (ELISA)

Western blots

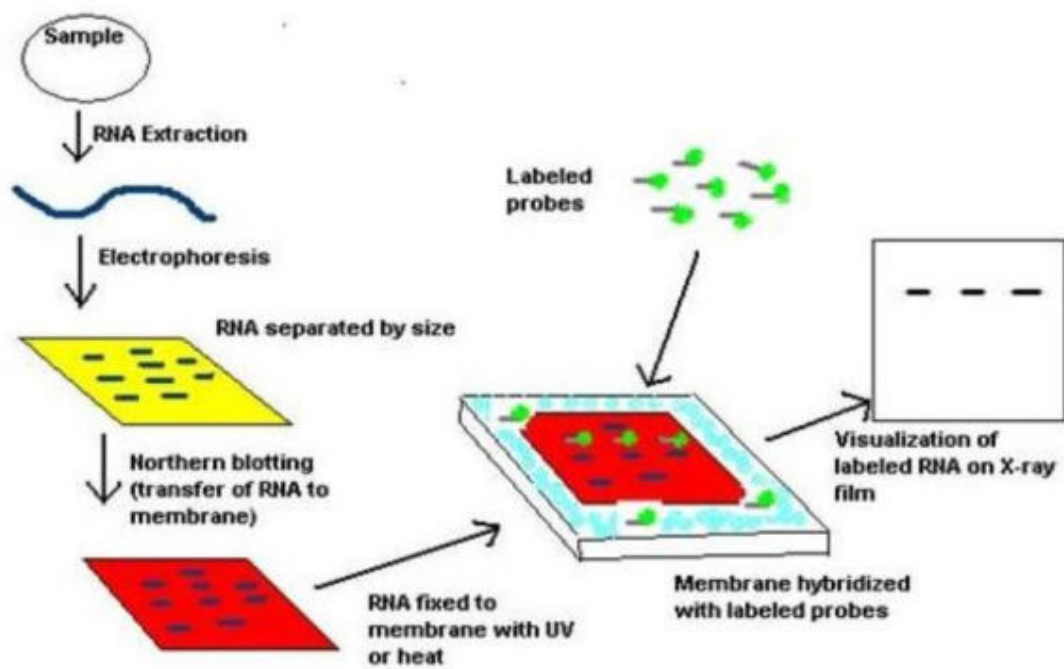
Proteomics: The study of all proteins expressed by a genome, including their relative abundance, distribution, posttranslational modifications, functions, and interactions with other macromolecules, is known as proteomics. Proteomics offer the potential of identifying new disease markers and drug targets.

3

Northern Blot

- The northern blot technique was developed in 1977 by James Alwine, David Kemp and George Stank at Stanford University
- Principle:
- Northern blotting involves the use of electrophoresis to separate RNA samples by size and detection with a hybridization probe complementary to part of or the entire target sequence.

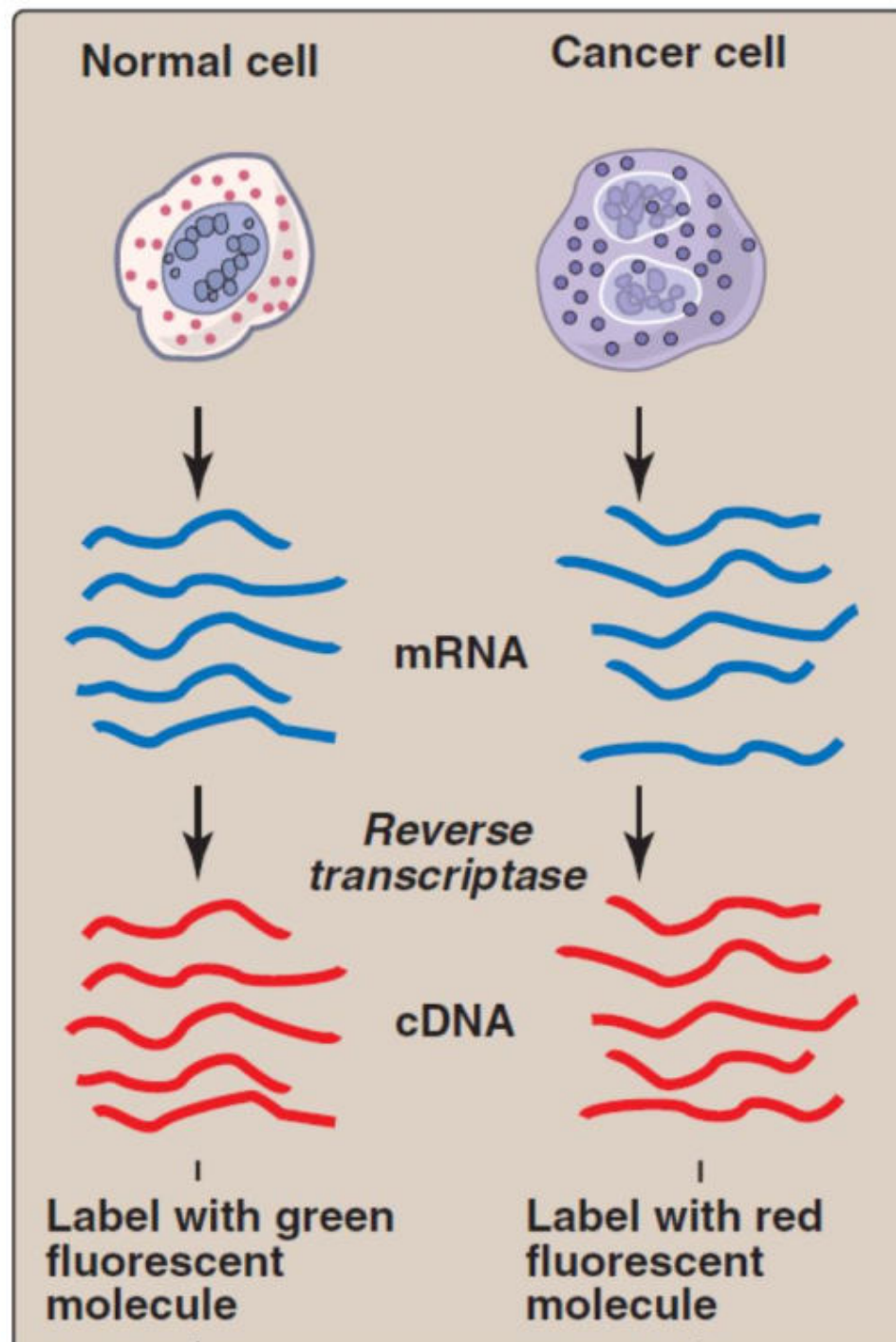
Procedure of Northern Blotting



5

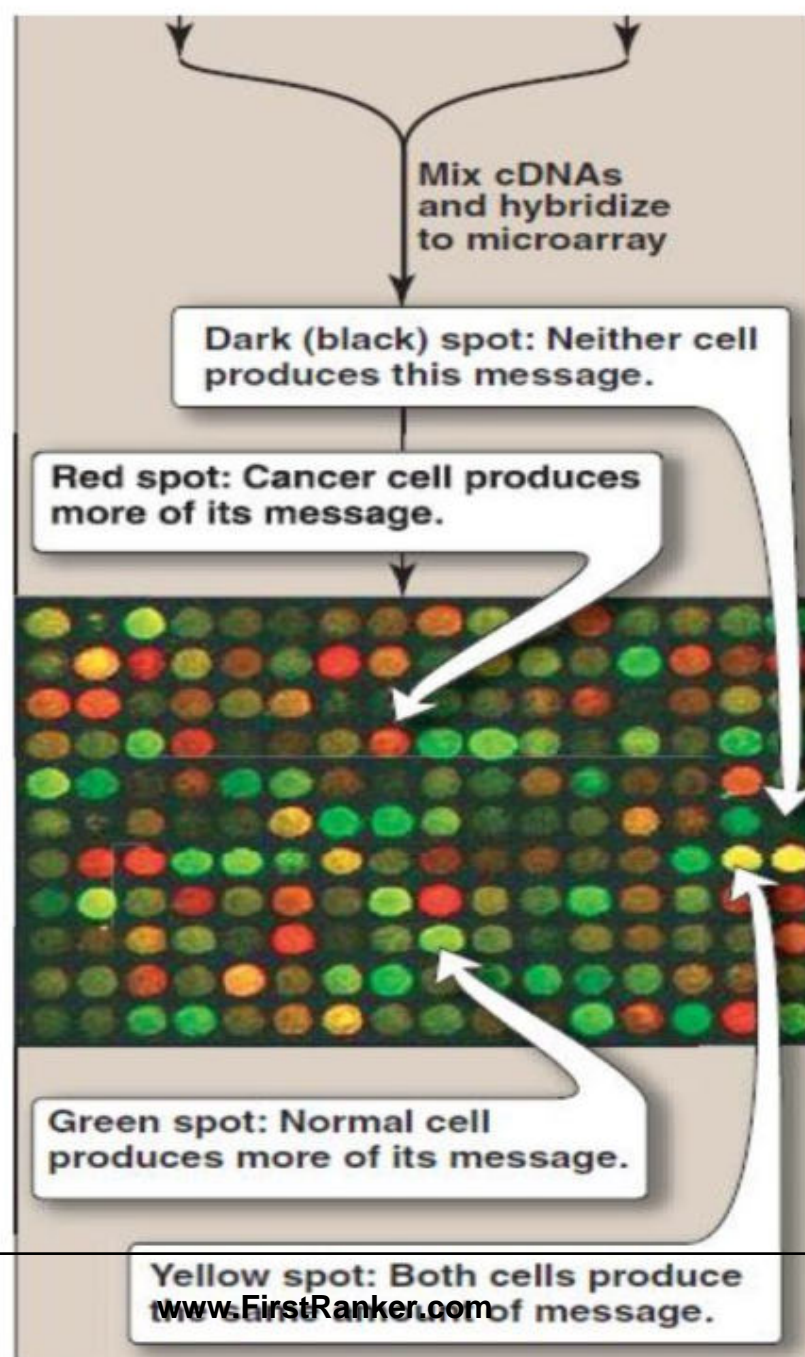
- Limitations:
- Northern Blotting using radioactive probes is very sensitive, but very **time-consuming**. Northern blotting is **not practical in large clinical studies** to detect the expression of hundreds of miRNAs and it also **requires large amounts (5–25 µg)** of total RNA from each sample

Microarray analysis of gene expression



7

Microarray analysis of gene expression



8

Gene Sequencing

9

“... [A] knowledge of sequences could contribute much to our understanding of living matter.”

Frederick Sanger

History

- The first method for determining DNA sequences involved a location-specific primer extension strategy established by Ray Wu in 1970
- The first DNA fragment to be sequenced belonged to T4 bacteriophage
- In the mid-1975, Frederick Sanger and Aln Coulson sequenced by using a **plus-minus system** for running a sequencing reaction.
- .

11

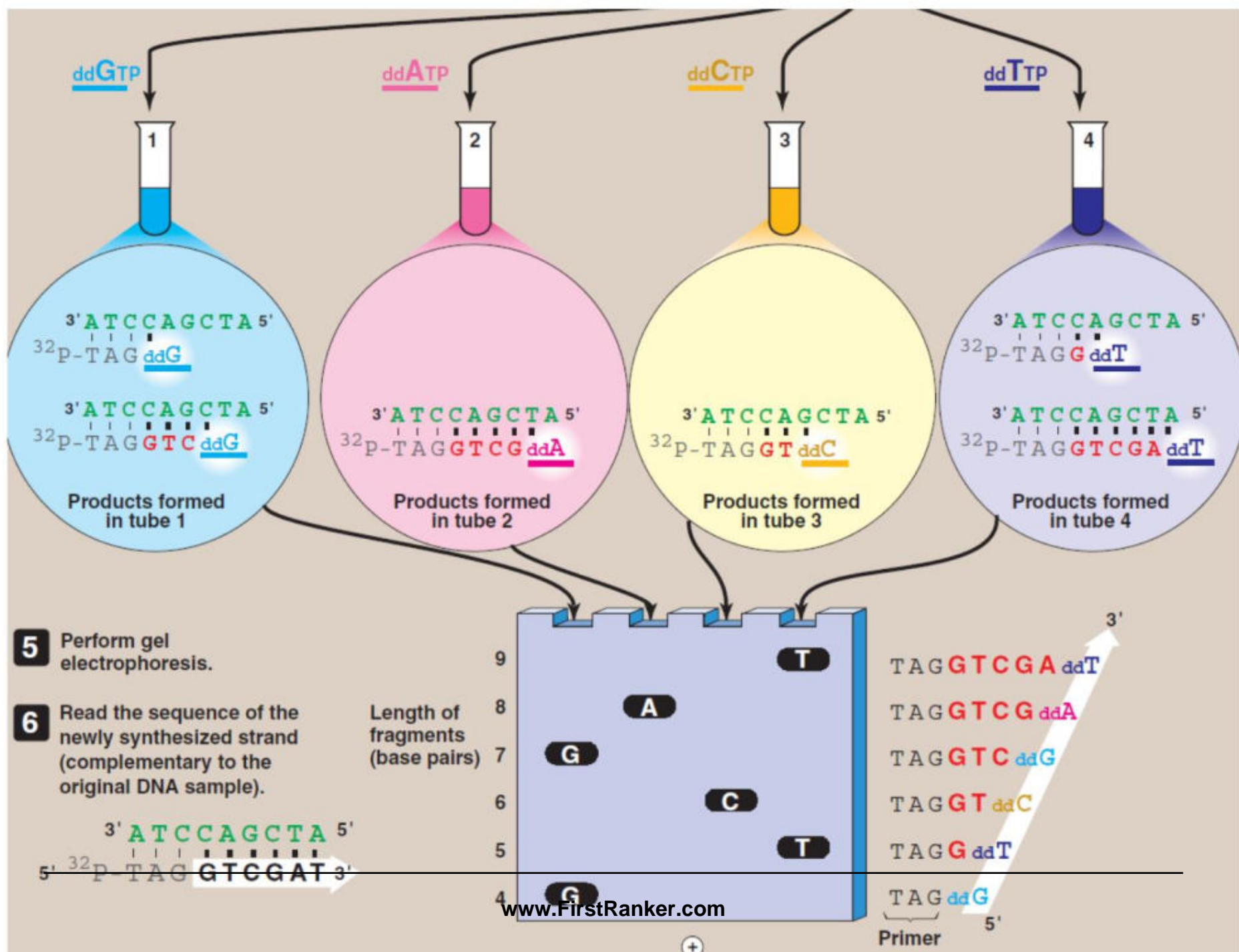
History contd

- Maxam and Gilbert further modified this method by using radiolabeled DNA and chemicals (such as hydrazine)
- One of the biggest breakthroughs in this field was the development of chain-termination technology using modified nucleotides by the Sanger lab in 1977
- In 1983, polymerase chain reaction (PCR) for amplifying stretches of DNA was discovered.

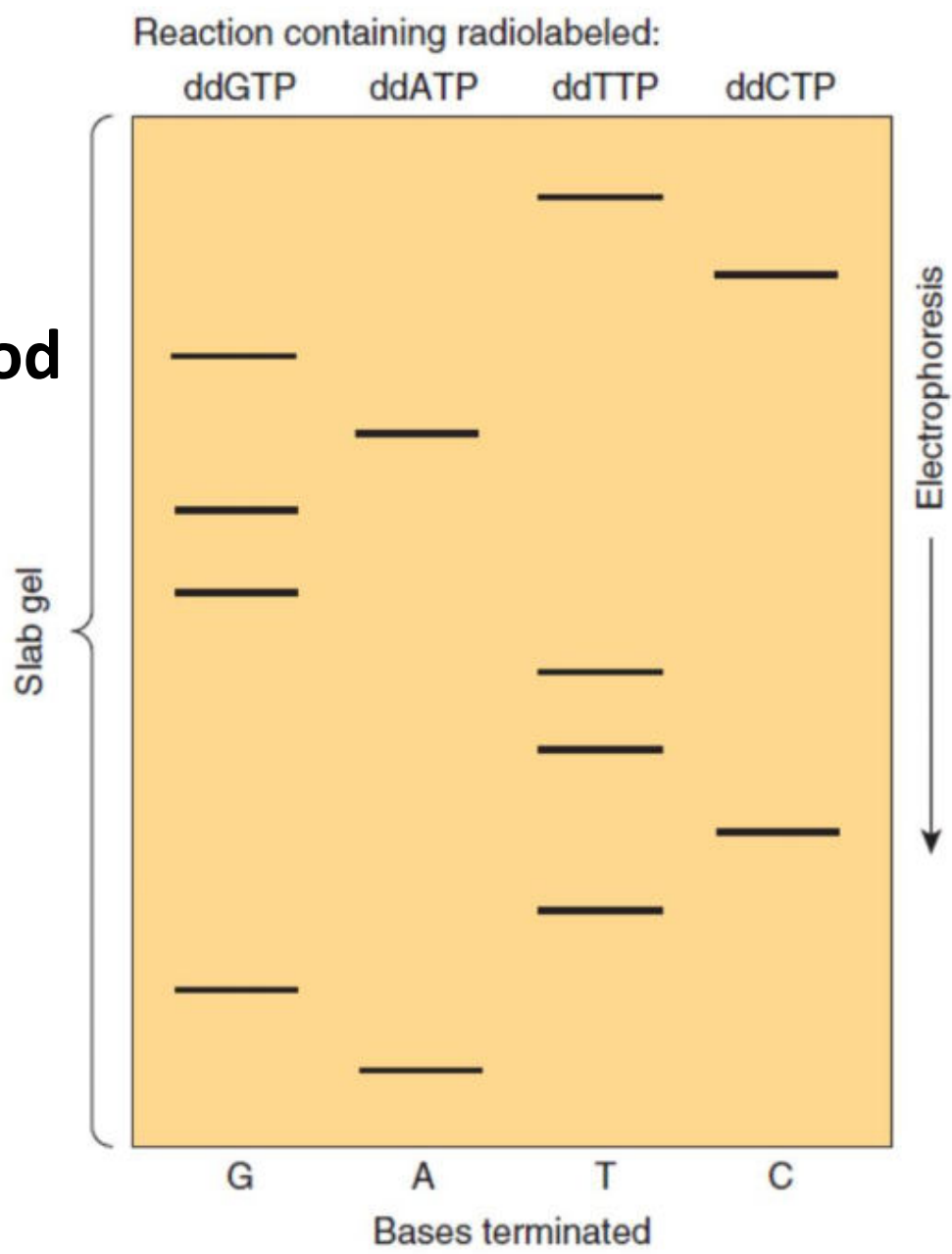
Type of gene sequencing

- **First generation DNA sequencing**
 - Sanger sequencing
 - Maxam Gilbert sequencing
 - Automated DNA sequencing
 - Shot Gun sequencing

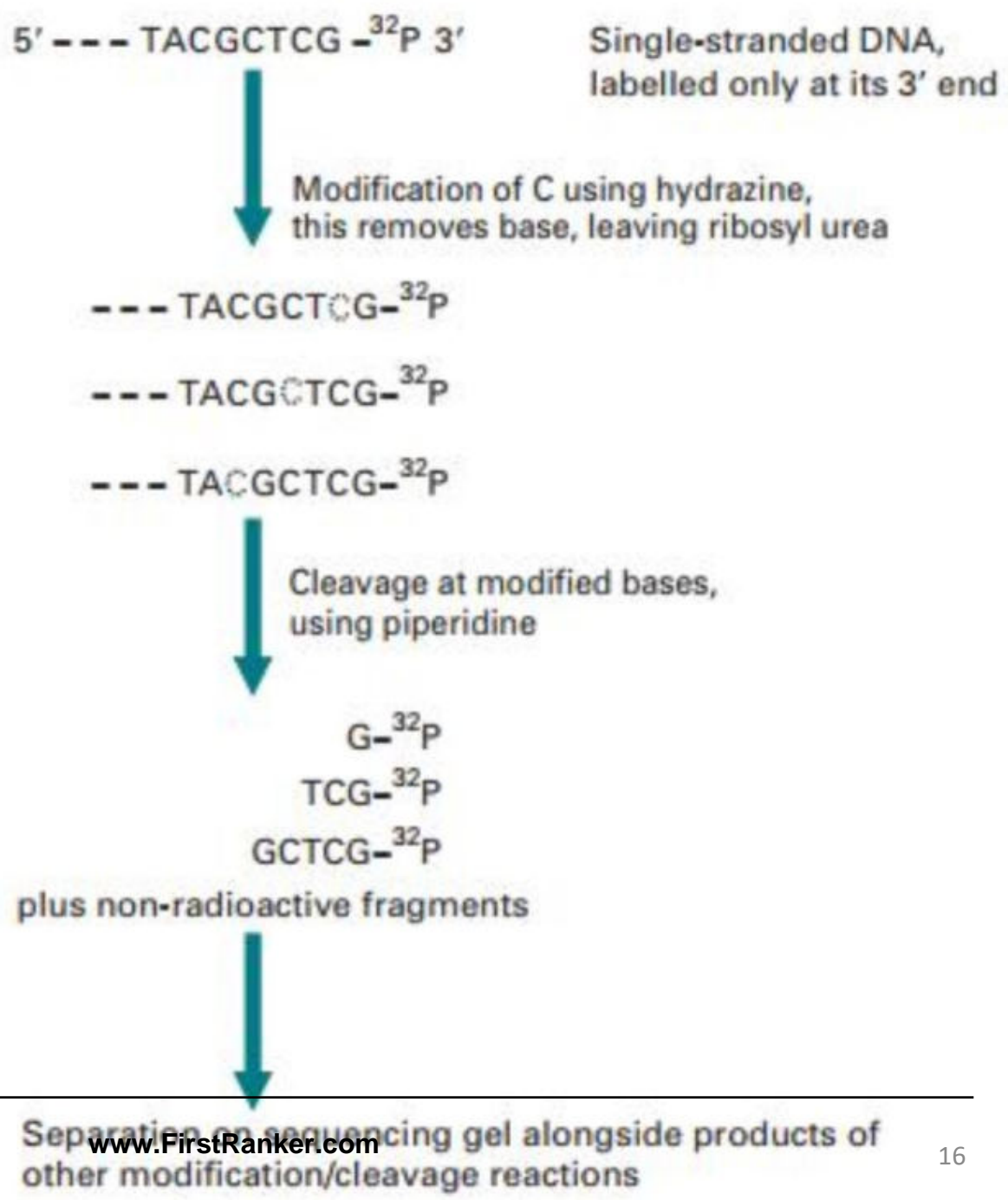
13



Sequencing of DNA by
the chain termination method
devised by Sanger



Maxam and Gilbert
sequencing of DNA

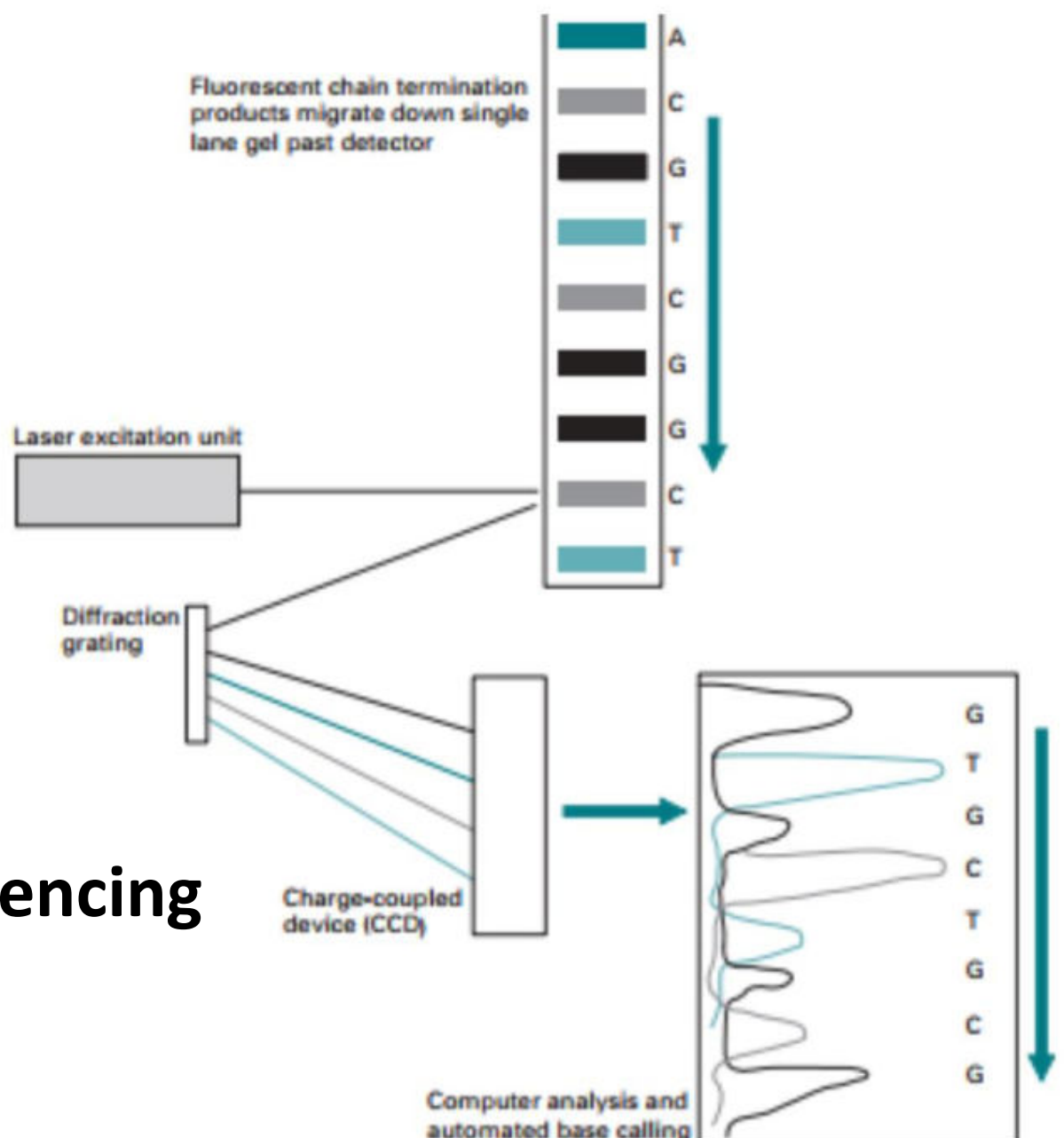


Automated DNA sequencing

- PCR used for making sequencing templates
- Fluorescently labelled ddNTPs are used
- Capillary electrophoresis

17

Automated fluorescent sequencing detection



- **Second-generation DNA sequencing**
: (Next generation sequencing)
 - Pyrosequencing
 - Sequencing by synthesis
 - Sequencing by ligation
 - Ion semiconductor sequencing

19

- **Third-generation DNA sequencing**
 - **Real-time, single-molecule sequencing**
 - capable of sequencing single molecules, negating the requirement for DNA amplification shared by all previous technologies.

Application of DNA sequencing

- Forensics:
 - To identify the individual
- Medicines
 - To detect genes associated with some hereditary or acquired diseases
 - E,g. Huntingtons disease
 - CAG codon in exon 1
 - Fragile X syndrome
 - CGG >200
 - Myotonic Dystrophy
 - CUG >100
- Agriculture
 - Mapping and sequencing of whole genome of microorganisms helps in making them useful for foods or crops

21

MCQ 1

- Which of the following is not an exclusively DNA sequencing method?
- 1. Sangers
- 2. Maxam Gillbert
- 3. Edman
- 4. LMPCR (Ligation mediated PCR)

LMPCR (Ligation mediated PCR)

- (1) primary DNA nucleotide sequences
- (2) cytosine methylation patterns
- (3) DNA lesion formation and repair, and
- (4) in vivo protein-DNA footprints

23

MCQ2

- The sample in Sangers method after reaction separated in
- 1. AGE
- 2. PAGE
- 3. PFGE (Pulse field gel electrophoresis)
- 4. 2-D gel electrophoresis

MCQ3

- If a hypothetical peptide has the sequence Phe-Tyr-Met-Pro-His.
- Calculate number of possible nucleotide sequences.
- A. 11
- B. 8
- C. 22
- D. 32