# Introduction to DSP – Proteins and Their Structure

Your Objectives:

At the end of the lesson, you should be able to describe proteins and their structure.

Proteins are composed of of amino acids. A typical protein

contains between 200 and 300 amino acids. Each amino acid has a specific chemical formula and structure.

Cells make distinctive proteins by arranging the different amino acids in unique sequences. For each type of protein, the cell's genetic code dictates which amino acids are added and in what order.

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These carefully arranged amino acids f	orm a chain-like	held
together by	bonds. Because the chem	iical bonds are weak, they
can be easily affected by factors such as		and pH.
Proteins can do their job only if they kee	o their	. Changes in the
shape of a protein make it unable to function correctly. If the target protein (product) is		
exposed to the wrong temperature or		, the protein chain will
lose its shape and ability to function e	ntirely. The	of protein
structure is called denaturation. A	pr	otein cannot return to its
original shape.		

**Protein structure** is the three-dimensional arrangement of atoms in an amino acid-chain molecule. Proteins are polymers – specifically polypeptides – formed from sequences of amino acids, the monomers of the polymer. A single amino acid monomer may also be called a **residue**, indicating a repeating unit of a polymer. Proteins form by amino acids undergoing condensation

, whereby the amino acids lose one water molecule per reaction in

order to attach bond to one another which has a peptide bond. By convention, a chain of under 30 amino acids is often identified as a peptide rather than as a protein. To be able to perform

their

function, proteins fold into one or more specific spatial

conformations driven by a number of non-covalent , such as

hydrogen bonding, ionic interactions, Van der Waals forces and hydrophobic packing. To understand

the functions of proteins at a level, it is necessary to determine

their three-dimensional

. The scientific field of structural biology

employs techniques such as X-ray crystallography, NMR spectroscopy, cryogenic (anstatt nur: cryo) electron microscopy (cryo-EM) and dual polarisation interferometry to determine the structure of proteins.

Protein structures range in size, from tens to several thousands of amino acids. By

size, proteins are classified as nanoparticles ranging between 1-

100 nm. Very large aggregates can be formed from protein subunits. For example, many thousands of actin molecules assemble into a microfilament.

A protein generally undergoes reversible changes in performing

its biological function. The alternative structures of the same protein are referred to as different conformational isomers, or simply, conformations, and transitions between them are called conformational changes.

## **Primary structure**

The primary structure of a prote	ein refers to the		of amino acids in
the polypeptide chain. T	he primary structu	re is held togeth	er by peptide
r	made during the proces	ss of protein biosynthesis	. The two ends of
the polypeptide chain are referred to as the carboxyl terminus (C-terminus), and the amino terminus (N-terminus), based on the nature of the free group on each extremity. A counting of residues always starts at the N-terminal end (NH2-group), which is the end where the amino			
group is not involved in a pe	ptide bond. The		structure of a
protein is determined by the gene corresponding to the protein. A specific sequence of nucleotides in DNA is transcribed into mRNA, which is read by the ribosome in a process called			
translation. The sequence of	amino acids in insulir	n was	by
Frederick Sanger, who established that proteins have defining amino acid sequences. The			
sequence of a protein is		to that protein and de	fines the structure
and function of the protein. The sequence of a protein can be determined by methods such as Edman degradation or tandem mass spectrometry. Often, however, it is read directly from the			
sequence of the gene, using the	e	code. It is stric	tly recommended
to use the words "amino acid residues" when discussing proteins because, when a peptide			
bond is formed, a <u>water</u>		is lost, and therefore	proteins are made
up of amino acid		Post-translational mod	lification such as

phosphorylations and glycosylations are usually also considered a part of the primary structure, and cannot be read from the gene. For example, insulin is composed of 51 amino acids in two chains. One chain has 31 amino acids, and the other has 20 amino acids.

#### Secondary structure

Secondary structure refers to highly regular local	structures on the

actual polypeptide backbone chain. Two main types of secondary structure, the  $\alpha$ -helix and the  $\beta$ -strand or  $\beta$ -sheets, were suggested in 1951 by Linus Pauling et al. These secondary structures are defined by patterns of hydrogen bonds between the main-chain peptide groups. They have a regular geometry, being constrained to specific values of the dihedral angles  $\psi$  and  $\varphi$  on the Ramachandran plot. Both the  $\alpha$ -helix and the  $\beta$ -sheet represent a way of saturating all the

bond donors and	in the peptide

backbone. Some parts of the

are ordered but do not form any

regular structures. They should not be confused with random coil, an unfolded polypeptide chain lacking any fixed three-dimensional structure. Several sequential secondary structures may form a "supersecondary unit".

## **Tertiary structure**

Tertiary structure refers to the three	structure of monomeric and

multimeric protein molecules. The  $\alpha$ -helixes and  $\beta$ -pleated-sheets are folded into a compact globular structure. The folding is driven by the non-specific hydrophobic

, the burial of hydrophobic residues from water, except that the

structure is stable only when the parts of a protein domain lock into place by specific tertiary interactions, such as salt bridges, hydrogen bonds, and the tight packing of side chains and

disulfide bonds. The disulfide bonds are extremely

in cytosolic

proteins, since the cytosol (intracellular fluid) is generally a reducing environment.

## Quaternary structure

 Quaternary
 structure
 is
 the
 three-dimensional
 structure
 consisting
 of
 the

 of
 two
 or
 more
 individual
 polypeptide
 chains
 (subunits)
 that

operate as a single functional unit (multimer). The resulting multimer is stabilized by the same

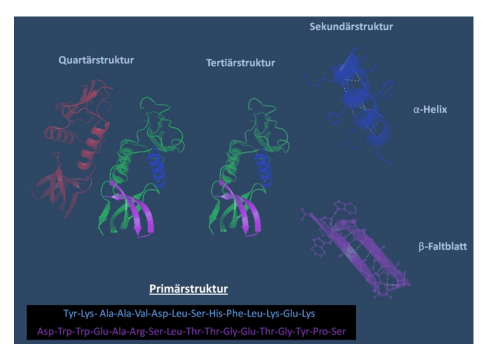
non-covalent interactions and disulfide bonds as with a tertiary structure. There are many possible quaternary structure organisations. Complexes of two or more polypeptides (i.e. multiple subunits) are called multimers. More specifically, it would be called a dimer when containing two subunits, a trimer with three subunits, a tetramer for four subunits, and a pentamer with five. The subunits are frequently related to one another by symmetry operations,

such as a twofold axis in a dimer. Multimers made up of identical

are referred to with the prefix of "homo-" whereas those made up of different subunits carry

the prefix of "\_\_\_\_\_\_-", as with a heterotetramer: two alpha and two

beta chains of hemoglobin.



Aufgabe Lückentext:

Folgende Wörter bitte in den Lückentext einfüllen. Jedes Wort kommt einmal vor. Bitte Gross- und Kleinbuchstaben beachten.

acceptors, aggregation, biological, bonds, chains, chemical, discovered, -dimensional, denatured, genetic, hydrogen, hetero, interactions, interactions, loss, molecular, molecule, primary, physical, pH, protein, rare, reactions, residues, sub-, subunits, structure, shape, structure, structural, sequence, temperature, unique,