

Lab (5): Protein fractionation by ammonium sulphate and dialysis


Aim:

- Fractionation of animal and plant crude extracts by ammonium sulphate.
- Removing of salts ions using dialysis.

Introduction:

Purification should yield a sample of protein containing only one type of molecule, the protein in which the biochemist is interested. This protein sample may be only a fraction of 1% of the starting material, whether that starting material consists of cells in culture or a particular organ from a plant or animal. How is the biochemist able to isolate a particular protein from a complex mixture of proteins? ⁽¹⁾

Several thousand proteins have been purified in active form on the basis of such characteristics as solubility, size, charge, and specific binding affinity. Usually, protein mixtures are subjected to a series of separations, each based on a different property to yield a pure protein. At each step in the purification, the preparation is assayed, and the protein concentration is determined. ⁽¹⁾


 **PAUSE AND THINK** → What will happen for the protein concentration during the purification scale?

The purification of proteins by altering the solubility achieved by what called *salting out*. Most proteins are less soluble at high salt concentrations, an effect called salting out. The salt concentration at which a protein precipitates differs from one protein to another. Hence, salting out can be used to fractionate proteins (as proteins will precipitate at different points with increases in salt concentration). Salting out is also useful for concentrating dilute solutions of proteins, including active fractions obtained from other purification steps. ⁽¹⁾ In fact, salting out is an effective means for initial molecule purification, but lacks the ability for precise isolation of a specific protein. Ideally, the type of salt being used, and the concentration of the salt can be varied to selectively precipitate the molecule. ⁽²⁾ Ammonium sulphate is common substance used to precipitate proteins selectively since it is very soluble in water, its relative freedom from temperature effects and harmful effects of proteins like irreversible denaturation. ⁽³⁾

Proteins can be separated from small molecules (salts) by *dialysis* through a semipermeable membrane, such as a cellulose membrane with pores. Molecules having dimensions significantly greater than the pore diameter are retained inside the dialysis bag, whereas smaller molecules and ions traverse the pores of such a membrane and emerge in the dialysate outside the bag. ⁽¹⁾

Principle:

The most effective region of salting out is at the isoelectric point of the protein, because all proteins exhibit minimum solubility in solutions of constant ionic strength at their isoelectric points. Different proteins will precipitate at different salt concentration, where protein size is inversely correlated with salt concentration. A typical protocol consists of adding ammonium sulphate to give specific percentage saturation, followed by a period of time for proteins to precipitate and a centrifugation step to collect the precipitate. Precipitation of proteins is conventionally carried out at 0 °C to avoid possible denaturation of proteins. Following fractionation by ammonium sulphate, dialysis is applied to remove salts. During dialysis, the small, unwanted salts ions removed from proteins in a solution by selective and passive diffusion through a semi-permeable membrane. Sample molecules (proteins) that are larger than the membrane-pores are retained on the sample side of the membrane, but small molecules and buffer salts pass freely through the membrane, where the salt molecules move from the more concentrated solution (from inside the dialysis bag) to the less concentrated solution (e.g. buffer). The movement of the salt molecules will stop, when the solution reaches the equilibrium. At this point, the buffer is changed to drive the diffusion and salts movements. (1,4,5,6)

 PAUSE AND THINK → What if you did not change the buffer? Why?

Materials:

Chemical

Prepared crude extracts, ammonium sulphate, 0.1M Tris-HCl (pH 7.4), phosphate buffer 0.1M (pH 7.0), distilled water.

Equipment and Glassware

Beakers, measuring cylinder, centrifuge tubes, dialysis bags, electronic balance, centrifuge, magnetic stirrer.

Protocol:

A. Salting out of protein A by 60% ammonium sulphate saturation:

1. Measure the volume of your crude extraction and calculate the weight in g of ammonium sulphate needed to saturate the solution 40% using **Table 1**.
2. Add the required salt to the solution slowly and gradually with small quantities and mix well continuously using magnetic stirrer while the sample is placed in ice.
3. After the addition is completed and the salt is completely dissolved, centrifuge at 3500 rpm for 10 min.
4. Take the supernatant, measure its volume, then saturate the solution to 60% using **Table 1**.

Supporting material:

- A video shows how to dialysis a sample by using dialysis tube:
https://f1.media.brightcove.com/4/3663210762001/3663210762001_5214982793001_5214973220001.mp4?pubId=3663210762001&videoId=5214973220001
- How to make a dialysis bag:
<https://www.youtube.com/watch?v=mWN-eE6fmpM>

References:

1. Berg JM, Tymoczko JL, Stryer L. Biochemistry. 5th edition. New York: W H Freeman; 2002.
2. [https://chem.libretexts.org/Bookshelves/Physical_and_Theoretical_Chemistry_Textbook_Maps/Supplemental_Modules_\(Physical_and_Theoretical_Chemistry\)/Thermodynamics/Real_\(Non-Ideal\)_Systems/Salting_Out](https://chem.libretexts.org/Bookshelves/Physical_and_Theoretical_Chemistry_Textbook_Maps/Supplemental_Modules_(Physical_and_Theoretical_Chemistry)/Thermodynamics/Real_(Non-Ideal)_Systems/Salting_Out).
3. Chapter 9: Protein expression, purification and characterization", *Proteins: Structure and Function*, Whitford, 2005, John Wiley & Sons, Ltd.
4. <http://tools.thermofisher.com/content/sfs/brochures/D21227~.pdf>
5. https://en.wikibooks.org/wiki/Structural_Biochemistry/Proteins/Purification/Salting_Out
6. <https://www.thermofisher.com/sa/en/home/life-science/protein-biology/protein-biology-learning-center/protein-biology-resource-library/pierce-protein-methods/dialysis-methods-protein-research.html>
7. http://kuchem.kyoto-u.ac.jp/seika/shiraiishi/protocols/as_precipitation.html