General Guide for Dissolving Peptides

※ The more charged residues on a peptide, the more soluble it is in aqueous solutions.
※ First try to dissolve a small amount of the peptide in either water or buffer.
※ Consider that any solvent that you use must be compatible with your assay.
※ Gentle warming and sonication are effective tools in aiding peptide solubilisation. Be aware that a peptide may appear to have solubilised after sonication, but if the solution is cloudy or has gelled it may merely be in suspension.
※ If a peptide does not dissolve and you wish to recover it, lyophilise to remove your solvent.

There is no general solvent for suitable for solubilising all peptides; the flowchart below provides a general guideline for peptide solubilisation. Please select the best solvent for your supplied peptide by testing a small amount first. It is advisable to first attempt to use solvents that are readily removed by lyophilisation.

The solubility characteristics of a peptide can be gauged by the amino acid composition; the ionic charges on the peptide will help towards determining its solubility. Please note that although hydrophilicity analysis will assist in the ability to solubilise your product, it is not a definitive determinant of the solubility of the peptide.
Peptides containing free cysteines should be dissolved in degassed acidic buffers. The thiols will be oxidized at pH>7 forming disulphide bonds. Solutions should be prepared immediately before use to avoid air oxidation.

Peptides containing a large percentage of hydrophobic residues, along with the peptides already described in the “Approaches to Peptide Solubilisation” flowchart, require the use of organic solvents due to their largely uncharged nature. Recommended solvents are DMSO, DMF, acetic acid, acetonitrile, methanol, propanol or isopropanol.

The peptide should be dissolved in the organic solvent completely before dilution in water with aqueous buffers (Sonication may be required). The buffer solution should be added dropwise with gentle agitation throughout the addition. More organic solvent may be added if the peptide starts to precipitate.

DMSO is a good solvent and has the advantage of being tolerated by cells. It is however, difficult to remove by drying.
DMSO should not be used if the peptides contain methionine (M) or free cysteine (C). Its use may cause oxidation of the side-chain functionalities if left in solution for long periods.
Calculating the Overall Peptide Charge

1. Count the number of **acidic** residues. Each has a value of \(-1\).
   The acidic residues are D, E and the C-terminal COOH.

2. Count the number of **basic** residues. Each has a value of \(+1\).
   The basic residues are K, R and the N-terminal NH2.

3. Count the number of **H** residues.
   If your intended solution is pH<6 then each has a value of \(+1\).
   If your intended solution is pH>6 then each has a value of 0.

4. Add together all of the assigned values.
   If the overall value is negative, the peptide is considered acidic.
   If the overall value is positive, the peptide is considered basic.
   If the overall value is zero, the peptide is considered neutral.

Calculating the Percentage of Charged & Hydrophobic Residues

1. Count the total number of charged residues on the peptide at pH 7: D, E, K,
   R, N-terminal NH2 and C-terminal COOH.

2. Count the number of hydrophobic residues on the peptide: A, F, I, L, M, P, V, W.

3. Count the total number of residues.
   Determine the number of charged residues as a percentage of total number of residues.
   Determine the number of hydrophobic residues as a percentage of total number of residues.
Hydropilicity Analysis – Example 1

MART-1 Peptide
ELAGIGLTV

For the above peptide we can carry out a hydropilicity analysis as an example and deduce the appropriate solvent.

One acidic residue and one C-terminal COOH = -2
No basic residues and one N-terminal NH2 = +1
Overall charge = -1

The peptide is considered acidic.
Percentage charged residues = (1/9) x 100
= 11%

This is very close to the 10% limit for using organic solvent (see “Approaches to peptide solubilisation” flowchart below). Also, almost half of the residues are hydrophobic, so it is best to initially use anorganic solvent for this peptide. Dissolving the peptide in the minimum amount of DMSO and then slowly adding dropwise to the desired aqueous buffer solution whilst constantly agitating would be an inappropriate method.

Hydropilicity Analysis – Example 2

OVA (323-329)
ISQAVHAAHAEINEAGR

Two acidic residues and one C-terminal COOH = -3
One basic residue (assuming neutral pH and ignoring the two H residues) and one N-terminal NH2 =+2
Overall charge = -1
Peptide is considered acidic at neutral pH.
Percentage charged residues = (3/17) x 100
= 18%
Percentage Hydrophobic residues = (3/17) x 100
= 18%

This peptide has quite a low percentage of hydrophobic molecules, evenly spaced throughout the peptide. It will be soluble in water, but if it did not dissolve and were applied to the flow diagram for the dissolving approach, the output would be:
Add a small amount of 10-25% acetic acid and dilute with water to the desired concentration.
Approaches to Peptide Solubilisation

- **Number of residues**
  - Generally soluble in water except where all residues are hydrophobic.
  - Add a small amount of CHCl₃ to mix solid peptide with water to the desired concentration.

- **Number of charged residues at pH 7.0**
  - Use of initial organic solvent required.
  - Sequences that are composed of a very high percentage (DTNB of pepsin-dissolved residue)
    - C, D, E, H, K, Q, R, S, T, V, and W are capable of forming proteinacious/elecrtrostatic hydrogen-bonding networks and tend to form gels or aggregate concentrated aqueous solutions.
  - Sequences that are composed of a very high percentage (DTNB of pepsin-dissolved residue)
    - Add 0.1 M sodium bicarbonate to dissolve the peptide and dilute to the desired concentration.

- **Number of hydrophobic residues**
  - Basic.
  - Acidic/Neutral.