

In solution protein digestion using trypsin as protease

Urea is a chaotropic agent and disrupts three dimensional structure of proteins and denatures them. *However, urea* + *heat* + *protein* = *carbamylation; urea in solution is in equilibrium with ammonium cyanate, that may decompose to ammonia and isocyanic acid* (HNCO). *Isocyanic acid attach the N*-*terminal of the protein, but also the side chains of lysine and arginine residues rendering a protein unsuitable for many enzymatic digests* ($HN=C=O + H_2N \sim \rightarrow H_2N$ -*CO*-*NH* \sim). Urea will *always degrade to isocyanic acid, so urea solutions must be made fresh, and it is recommended to add* 20mM methylamine (CH_3NH_2)*to the urea solution prior to use* (*urea can also be removed before digestion using reversed phase chromatography*)

Protein solvation/denaturation (applies for 100 µg protein or lower)

Dissolving the protein pellet;

The pellet may be difficult to dissolve. Add **<u>20µl</u> <u>urea solution</u>** (*see right panel*) and pipette gently up and down, sonicate if necessary. <u>Urea solution; 8M Urea/20mM</u> <u>methylamine:</u> Add **480 mg Urea** (art. no. 51458, Sigma-Aldrich), **1.7µl 40 wt% methylamine in H₂O** (art. no. 426466, Sigma-

Aldrich) and 630µl dH₂O.

Add <u>**20µl trypsin buffer**</u> (*see right panel*), and incubate at RT in Eppendorf mixer for 5 min (slow agitation).

<u>Trypsin buffer; 50mM</u> <u>Tris/1mM CaCl₂:</u> Add **0.61g Tris** (art. no. 252859, Sigma-Aldrich) and **15mg CaCl₂ x 2H₂O** (art. no. 21097, Sigma-Aldrich, <u>inhibits</u> <u>chymotrypsin activity</u>) to about 90ml dH₂O. Correct the pH to 7.8-8 with HCl and adjust the volume to 100ml. Store the solution at 4 °C.

Reduction and alkylation

Reduction;

Add **4µl 100 mM DTT** (*see right panel*), and incubate for 1 hour at room temperature (do NOT use 56°C as with gel pieces. That will cause carbamylation due to the presence of urea in the sample).

Alkylation;

Add **5µl 200 mM IAA** (*see right panel*) for cystein alkylation, and incubate for 1 h at room temperature (dark).

To avoid unwanted protease alkylation, add 0.8μ l 100 mM DTT, and incubate 10 min. at room temperature.

Digestion

Sample dilution;

Add 110.2 μ l Trypsin buffer (the urea concentration is now 1M).

Trypsin;

Add trypsin at a concentration about 50 times lower than the amount of protein in the sample. If the sample contains approx. 100 μ g protein, add 2 μ g of protease (*see right panel*). Measure pH using an indicator paper (litmus paper or similar), and incubate samples at 37°C overnight on a shaker

100 mM DTT in MillQ water:

Add **15.4 mg DTT** (DiThioThreitol, art. no. 171318-02, Amersham Biosciences) to $1ml dH_2O$ (may be aliquoted as a 1M solution, and kept in freezer).

200 mM IAA in MilliQ water:

Add 18.5mg IAA

(Iodoacetamide, art. no. I-6125, Sigma Aldrich) to 0.5ml dH_2O (must be freshly made and kept in the dark).

<u> 2µg Trypsin Porcine (4µl)</u>

(Promega, art. no. V 5111);

Dissolve each ampoule (20 μg trypsin porcine) in 40 μ l 50 mM acetic acid (resuspension buffer supplied from Promega with the trypsin powder). The trypsin concentration in this stock solution is then 0.5 $\mu g/\mu$ l

Acidification

In this final step, add 15 μ l 10% FA (formic acid) to quench the digestion activity. We now have approximately 0.5 mg/ml digested protein solution at pH 3. The Urea concentration in this solution (below 1M) allows analysis directly by MALDI or LC-MS. The solution should be desalted/concentrated on reversed phase_microcolumns before either MALDI-ToF or nanoflow LC-MS.