

View info regarding our last PhD Proteomics **course** (including sample prep) that took place Februari 2023 in Wageningen, the Netherlands.

<https://www.vlaggraduatschool.nl/en/courses/course/PRO23.htm>

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## 1. General information

### 1.1 pH stuff and more

You can check the pH of your sample by putting 0.1 ul of sample (or less) on a piece of pH paper.

#### pH UP from pH 2 to pH 8

1 ml/l HCOOH = formic acid (FA):     pH = 2.4

For 1 ml:                     Make pH 8 – 8.5 by adding 15 – 20 ul 10\* diluted conc. NH3 (max 37%).

0.5 ml/l TFA = TriFluoroAcetic acid:     pH = 2.1

For 1 ml:                     Make pH 8 – 8.5 by adding 6- 9 ul 10\* diluted conc. NH3 (max 37%).

#### pH DOWN from pH 8 to pH 2 - 4

50 mM ABC pH 8.0:

For 1 ml:                     Make pH 3 by adding 35 ul 10\* diluted conc. TFA.

1% Sodium DeoxyCholate (DC) can be used to **extract hydrophobic proteins** to replace 4% SDS:

Pasing, Y., S. Colnoe and T. Hansen (2017). "Proteomics of hydrophobic samples: Fast, robust and low-cost workflows for clinical approaches." *Proteomics* **17**(6).

Schmidt, A., K. Kochanowski, S. Vedelaar, E. Ahrne, B. Volkmer, L. Callipo, K. Knoop, M. Bauer, R. Aebersold and M. Heinemann (2016). "The quantitative and condition-dependent Escherichia coli proteome." *Nature Biotechnology* **34**(1): 104-110.

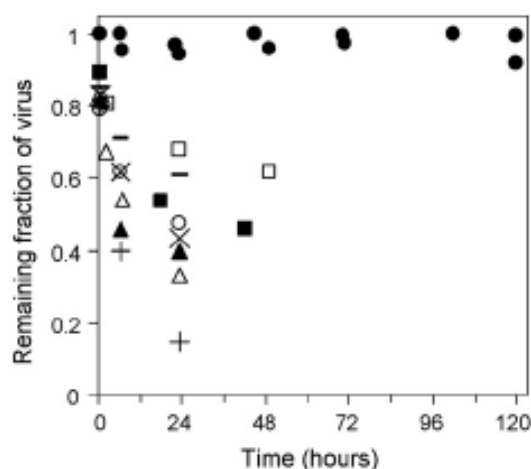
## 1.2 Abbreviations and solutions

		Find it in:	MW			in
<b>AA</b>	<b>AcrylAmide (Wear gloves)</b>	lab 3030	71	20 mM = 1.4 mg/ml	200 mM = 14 mg/ml	water
AcNi	Acetonitril	lab 3030				
ABC	Ammonium BiCarbonate	lab 3030	79	50 mM = 0.2 g /50 ml	2 M = 0.16g + 0.84ml	water
AmAc	Ammonium Acetate	lab 3030	77	10 mM = 38.5 mg	/ 50 ml	water
Cystein	Fluka 30090, >99%	lab 3030	121	125 mM = 15 mg/ml	200 mM = 24 mg/ml	water
<b>DTT</b>	<b>Dithiotreitol (4°C)</b>	FRESH! (refrig)	154	20 mM = 3.1 mg/ml	150 mM = c	water
IAA	Iodoacetamide (4°C)	FRESH! (refrig)	185	20 mM = 3.7 mg/ml	200 mM = 37 mg/ml	water
TFA	TriFluoro-Acetic acid	lab 3030		Safety cupboard MS stuff	dilute in fume cupboard only	
TCEP	Tris(CarboxyEthyl)Phosphine	-20°C	287		100 mM = 29 mg/ml	water
Tris	Stock 10*. pH 8 with HCl	weighing room	121	1 M = 121 g/l	6.0 g / 50 ml	water
Trypsin	we have a stock solution of Bovine Sequencing grade Trypsin (Roche 11 047 841 001) of 0.5 ug/ul (500 ng/ul) in 1 mM HCl. Generally dilute 100* in ABC before use unless stated otherwise					
Urea	Make FRESH	weighing room		Do not warm up		Tris
UT	8 M urea (Sigma, U0631)	weighing room	60	<b>100 ul 1M Tris + 0.48 g urea made up to 1.0 ml pH will increase to 8.2 due to the addition of urea</b>		

- 0.5 ml protein LoBind tube: order# eppe0030108.094
- 2.0 ml protein LoBind tube: order# eppe0030108.132
- Pall 3K or 10K omega filter (20kDa / 50kDa cutoff) order# Pall OD003C34 = Sigma-Aldrich Z722049-100EA

## 1.3 Tips and ways to reduce the amount of Keratins in your samples.

1. **DO:** Use commercial electrophoresis gels. They contain less keratin than your own ones.  
Use Eppendorf low binding tubes to minimize protein losses (see figure below).  
Keep gels covered as much as possible. Put them into a new square large petri dish for staining.
2. Use Nitril gloves, not Latex ones.
3. Wash your hands under running tap water before you start and as often as possible in between handlings.
4. Do not work in a standard flow cabinet intended for microbiological work. Due to the large air flow, they increase the amount of keratin passing your vials resulting in more keratin in your samples.
5. Do not wear clothes of wool while doing proteomics sample prep.
6. Try not to lean over the samples too much.
7. Never ever use glass vials for proteins. You will lose a lot of protein.
8. Use new (or only used for proteomics) throw away (polypropylene) plastics.
8. Do not use hand creams when preparing samples for MS.
9. Do not use any glassware that has been cleaned with detergent (e.g. in a washing machine).
10. Cheap (non-Eppendorf) micro tubes may contain polymers, mold release agents, plasticizers, etc.
11. Do not use pipet tips that have been sterilized by heating them. The heat may release plasticizer compounds.
12. Use Eppendorf LowBind tubes, not siliconized tubes.
13. Clean your pipets before use. Especially when you have pipetted detergent (Triton X100, Tween 80 etc) with them.



Eppendorf LoBind tubes bind much less viruses than other brands of LoBind tubes.

From:  
E.I. Trilisky, A.M. Lenhoff: Sorption processes in ion-exchange chromatography of viruses. J. Chromatogr. A 1142 (2007) 2 - 12.

Fig. 5. Virus loss in different containers (20mM HEPES, 150 mM NaCl, pH 7.8; initial Ad5 concentration was approximately  $6 \times 10^{10}$  p/mL  $\approx 0.02$  g/L): (●) LoBind™ protein tube (Eppendorf), (□) Fisherbrand 0.5 mL tube, (■) Glass (National Scientific), (○) Fisherbrand PCR tube, (×) polycarbonate tube (Eppendorf), (▲) prelubricated tube (Costar), (△) polypropylene tube (ULP), (Δ) Maxym Recovery (Axygen), (+) silanized glass (National Scientific).

## 1.4 Recommended procedures

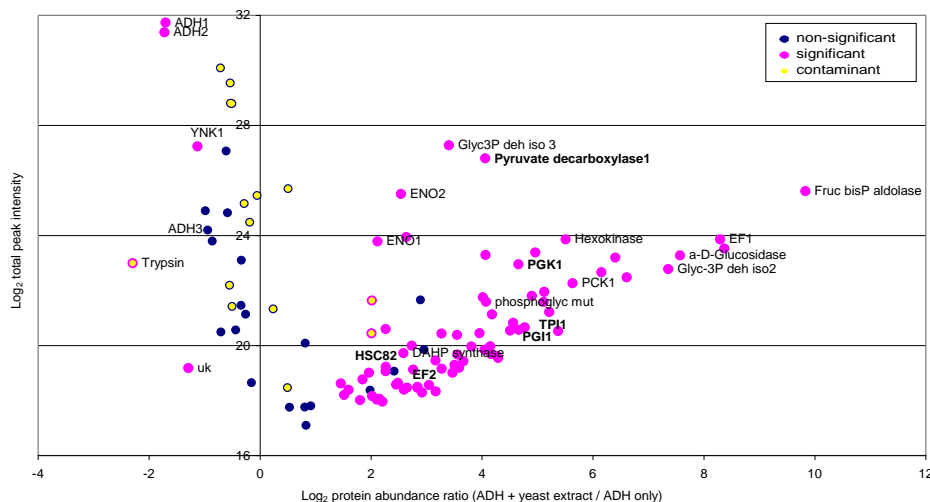
	Protein Identification	Label free relative Quantitation	Silac labeled Relative Quantitation	Dimethyl labeled Relative Quantitation	Absolute Quantitation
PAC	+	+	+	+	+
FASP	+	+	+	+	+
In Gel Digestion (IGD)	+	+	+		
On column Dimethyl labeling	-	-	-	+	-
Ovalbumin Standard addition	-	Possible but not advised	-	-	-
Peptide (labeled) Standard addition	-	-	-	-	+
Peptide fractionation	+ (always possible e.g. with High pH RP on $\mu$ Columns)				
Measurement	LC-MS/MS by Easy nLC1000 $\rightarrow$ Orbitrap Exploris 480				
Raw Data handling	MaxQuant (Protein identification and relative quantitation of peptides larger than 7 AA) MaxNOVO can be activated in the MaxQuant search to include smaller peptides of 3-7 AA.			QualBrowser + MaxQuant	
MQ data handling	Perseus (extra filtering of MQ result, statistics and intensity or ratio based clustering of proteins)				
Bioinformatics	<p>Overview of GO tools (but not complete): <a href="http://geneontology.org/">http://geneontology.org/</a> --&gt; Tools.                      First <b>pathway analysis</b> can be done with eg: Biocyc (WUR has a free subscription now): <a href="http://Biocyc.org">Biocyc.org</a>.                      Reactome (<a href="http://www.reactome.org">www.reactome.org</a>), PathVisio (<a href="http://www.pathvisio.org/">http://www.pathvisio.org/</a>), KEGG (<a href="http://www.genome.jp/kegg/tool/map_pathway2.html">http://www.genome.jp/kegg/tool/map_pathway2.html</a>).  <b>GO enrichment analysis</b> can be done within:                      - Internet resources like DAVID (Database for Annotation, Visualization and Integrated Discovery, <a href="https://david.ncifcrf.gov/">https://david.ncifcrf.gov/</a>, very easy), or (InterProScan)/PloGO (see below) + WeGO.                      - Cytoscape (<a href="http://www.cytoscape.org">www.cytoscape.org</a>) with plugins like BinGO or ClueGO (or EnrichmentMap).                      - Use the "R project" environment e.g. PloGO (includes abundance information = more advanced).  <b>Interactome</b> studies may benefit from database contained information e.g. in <a href="http://STRING-db.org/">STRING-db.org/</a> but also from Cytoscape plugins like Bionetbuilder or Bisogenet.  <b>Alternatives:</b> commercial software like ProteinCenter, Ingenuity or Metacore (these commercial softwares are not available at WUR Biochemistry).</p>				
(Not all software mentioned has been tested by me)					

Silac: Stable Isotope Labeling by Amino acids in Cell culture.

FASP: Filter Aided Sample Preparation

MQ: MaxQuant

MaxQuant label free relative quantitation result after analysis with Perseus statistical software. The Figure was prepared in Excel.



**Table 1 |** Comparison of spin filter, short SDS-PAGE and TFE methods

Method	Protein load	Peptide identifications	Protein identifications
Spin filter	50 $\mu$ g	5,369	642
Short SDS-PAGE		4,176	593
TFE		4,663	593
Spin filter	150 ng	86	46
Short SDS-PAGE		298	106
TFE		626	150

Samples of human RKO colon carcinoma cells containing the indicated amounts of protein were prepared in triplicate by the indicated methods and analyzed by reverse phase LC-MS/MS. Peptide identifications are total MS/MS spectrum-to-sequence database matches at 5% false discovery rate; protein identifications are nonredundant identifications with at least two identified peptides and parsimonious protein assembly. Reported values are the means of three technical replicate analyses.

## 1.5 Sample losses, methods comparison.

Table 1 was taken from: Liebler, D. C.; Ham, A. J. L.: Spin filter-based sample preparation for shotgun proteomics. Nature Methods (2009), Volume: 6, Issue: 11, p785-785.

	Yield (%)	Remark
FASP/PAC	50/60	The yield can be somewhat increased by using multiple digestion steps. (Wisniewski, J.R. and Mann, M. (2012) Consecutive Proteolytic Digestion in an Enzyme Reactor Increases Depth of Proteomic and Phosphoproteomic Analysis. <i>Analytical Chemistry</i> 84(6), 2631-2637)
In-gel digestion (IDG)	20	Purified protein: load a few ug of protein on the gel. Protein extract: more proteins will be identified when you load more protein. Load 50 – 75 ug for a complete proteome.

## 1.6 nLC-MSMS sample necessities.

Hand in minimally 25 ul acidic peptide sample (pH 2 – 4) with a concentration of approximately 50 – 500 ng/ul.

Peptide sample solutions to be measured should not contain detergents (SDS, Tween, Triton, NP etc) or other charged nonvolatile ions like TRIS or phosphate or particles of any kind.

Proteins can be transferred into the appropriate (ABC) solution by treatment with a Spin Filter (3 or 5 kDa) filter before doing the Trypsin digestion. Another good and proven method to get rid of impurities is to do an acetone precipitation, FASP or an SDS gel-electrophoresis step (see section 4) or at least a C18 uColumn cleanup (see section 1.7).

Peptide samples can be desalted/concentrated/transferred by performing the C18 uColumn cleanup as well (see section 1.7).

## 1.7 General peptide sample cleanup procedures with $\mu$ Columns.

Peptide solutions can be concentrated or desalted, and beads (from PAC/IP/chromatography/SP3) can be removed using  $\mu$ Columns (= C18 Stage tip (Affinisep SPE-Disks-Bio-C18-100.47.20) + LichroPrep RP-18 (Merck 1.09303) column material). With this  $\mu$ Column, there will be no loss of hydrophilic peptides (in contrast to using commercially available micro tips like Zip Tips or StageTips)

1. Prepare your own  $\mu$ column by:

Use the “cookie cutter” method to cut a small (1.6mm = Gauge 14) piece of a C18 Empore disk (= frit). Do this a second time. Transfer the 2 frits to the 200 ul tip with a good fitting plunger and tap it mildly. Add 200 ul methanol to the tip with frit.

Prepare a 50% slurry of LichroPrepC18 column material in methanol and add 4 ul of the 50% slurry into the methanol in the tip + frit.

The prepared  $\mu$ Column can be eluted by hand with a plastic 10 ml syringe, or with the vacuum manifold (connected to a vacuum pump) in lab3030. Whatever method you use, do not let the  $\mu$ Columns run dry.

2. Wash the  $\mu$ Column ones more with 100 ul MeOH.

3. Equilibrate the  $\mu$ Column ones with 100 ul 1 ml/l HCOOH in water.

4. Dissolve the sample preferentially in 15 – 200 ul of 1 ml/l HCOOH in water (not containing any AcNi) or in any other aqueous buffer.

5. Add sample(s) to the uColumn(s) (wash gel pieces with an extra 50 or 100 ul 1 ml/l HCOOH in water and add that too) and elute through.

6. Wash the uColumn ones with 100 ul 1 ml/l HCOOH in water.

7. Transfer the uColumn to a new 0.5 ml Eppendorf low binding tube.

8. Manually elute peptides from the C18 Stage tip+ uColumn by adding and eluting with 50 ul 50%AcNi + 50% 1 ml/l HCOOH in water directly into the 0.5 ml Eppendorf low binding tube.

9. The sample is now MalDI-tof ready.

10. For LCMS analysis, reduce the AcNi content by putting the samples in a Concentrator (with open cap) at 45 °C for 2 hours or longer when necessary. The final volume should be below 15 ul. Adjust the sample volume with 1 ml/l HCOOH in water to exactly 50.0  $\mu$ l. Sonicate (water bath sonicator) for 5 sec in the hot spot when the sample had been dried completely by accident.

The sample now has <5% AcNi, is pH 3, is particle free and therefore is nLCMS ready. Peptide concentrations can be measured e.g. using Pierce Quantitative Colorimetric Peptide Assay (23275) when the concentration is high enough.

Remark 1: The  $\mu$ Column cleanup step can also be used when you want to concentrate or combine samples. An alternative option to concentrate or combine peptide samples is the Eppendorf concentrator.

Remark 2:  $\mu$ Columns can be step eluted at pH 10 with 50 ul of a step gradient of AcNi [e.g 5, 8, 11, 18 and 25% AcNi with 20 mM NH<sub>4</sub>Formate pH10] to **fractionate** peaks according to their high pH affinities for the C18 column.

Fractions 1 + 4 (5 + 18%) and 2 + 5 (8 + 25%) can be mixed to decrease the number of samples to be injected to 3.

Better: fractionate extensively and mix fractions concatenated: 1+6+11, 2+7+12 etc. (Yang, F., Y. F. Shen, D. G. Camp and R. D. Smith (2012). "High-pH reversed-phase chromatography with fraction concatenation for 2D proteomic analysis." *Expert Review of Proteomics* 9(2): 129-134).

## 1.8 Methionine oxidation

(incubate with 0.1v/v% or 10 $\mu$ M H<sub>2</sub>O<sub>2</sub> by incubating at 4 °C overnight) can be used to confirm a Methionine within a peptide. Oxidized methionines shift the molecular weight up by 16 Da for each methionine (Koudelka, T et.al. (2012) Methionine Oxidation Enhances kappa-Casein Amyloid Fibril Formation. *Journal of Agricultural and Food Chemistry* 60(16), 4144-4155).

**1.9 Acetone precipitation as a general protein clean-up method** (100 ul protein extract in 100 mM Tris pH8: add 400 ul acetone and incubate at room temperature for 30 min. Centrifuge and remove supernatant) can be used as a general low loss sample cleanup method when necessary. Ref: Nickerson, J. L. and A. A. Doucette (2020). "Rapid and Quantitative Protein Precipitation for Proteome Analysis by Mass Spectrometry." *Journal of Proteome Research* 19(5): 2035-2042.

## 2. Protein determination (BCA).

by the Bicinchoninic Acid (BCA) method (Pierce:#23225):  
(linear sensitivity: 2 – 20 ug protein as measured at 562 nm in a 96 well microplate)

The BCA protein concentration determination method is almost independent on the amino acid composition but cannot be used for samples containing reducing agents (like DTT or TCEP) or chelators (like EDTA).

Alternative: Pierce 660 nm Protein Assay (#22660) with the Ionic detergent Compatibility reagent (#22663) can be used in the presence of maximally 5% SDS and/or 500 mM DTT and is linear between 2.5 and 600 ug BSA or 5 to 1200 ug of Ovalbumin. Compared to BSA, the Ovalbumin response is 54%. This strong protein dependence is the main disadvantage of the Pierce 660 nm Protein Assay.

### BCA Solutions:

BCA working solution (BCA ws): 12.5 ml reagent A + 0.25 ml reagent B  
Standard protein BSA: 2.0 mg/ml (ThermoFisher 23210)

When you made it yourself, measure the exact concentration BSA by measuring the absorbance at 280 nm (blanc is water):

$$C_{BSA} = A_{280} * 1.50 \text{ (mg/ml)}$$

$$C_{Ovalbumin} = A_{280} * 1.30 \text{ (mg/ml)}$$

This measurement is necessary since purified proteins always contain some salts.

### Method:

1. Pipet indicated volumes shown in bold in the table below into 0.5 ml low binding eppendorf micro tubes. Mix.
2. **Pipet 50 ul of the BSA standard or sample directly in a 96 well plate.**
3. **Add 200 ul of the BCA working solution.** Make sure samples are mixed well in the plate and seal the plate tightly with parafilm. Incubate (at 21 °C for 60 min or) at 37 °C for 30 min or more.
4. Measure the  $A_{562}$  in the micro plate reader you can find in room 3072.

	Dilute Cal curve into a 0.5 ml LB tube				
	Volume Water (ul)	Volume BSA (ul)	BSA amount (ug/50ul)	Measured A562 (1)	Measured A562 (2)
<b>Cal 1</b>	<b>500</b>	<b>0</b>	0	0.086	0.082
<b>Cal 2</b>	<b>490</b>	<b>10</b>	2	0.18	0.19
<b>Cal 3</b>	<b>190</b>	<b>10</b>	5	0.302	0.332
<b>Cal 4</b>	<b>180</b>	<b>20</b>	10	0.473	0.514
<b>Cal 5</b>	<b>170</b>	<b>30</b>	15	0.695	0.693
<b>Cal 6</b>	<b>160</b>	<b>40</b>	20	0.837	0.886
<b>Cal 7</b>	<b>150</b>	<b>50</b>	25	0.957	0.989
<b>Sample</b>	<b>40</b>	<b>10</b>		0.626	0.633

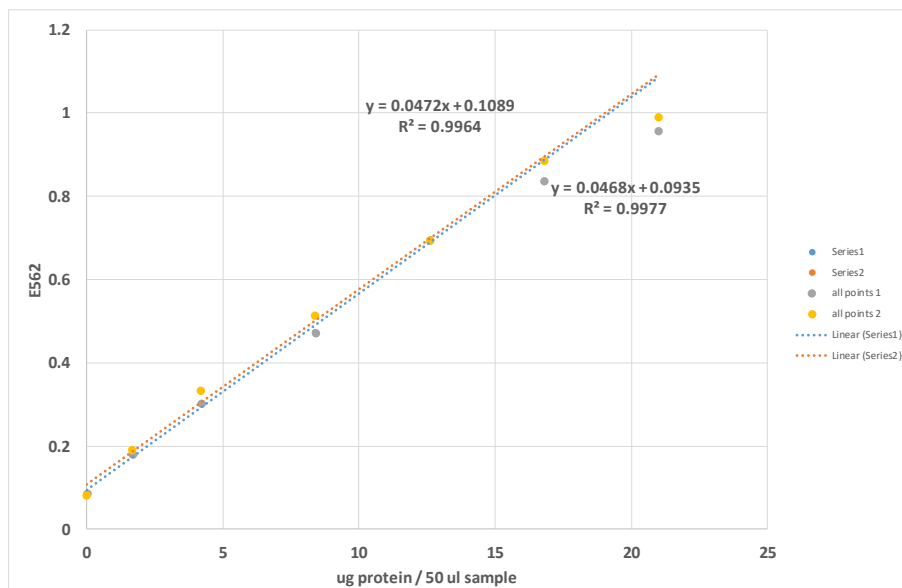
### Example measurement:

When the BSA was prepared and dissolved at 2 mg/ml. The absorbance at 280 nm was measured to be 1.118

Therefore the real concentration was:

$$1.118 * 1.50 = 1.64 \text{ mg/ml}$$

Adjust x-axis with factor 1.64 / 2



$$\text{ug protein in the 50 ul well} = \frac{[E_{562} - 0.10]}{0.047} =$$

$$\frac{([0.626 + 0.633] / 2 - 0.10)}{0.047}$$

$$\Rightarrow 1.3 \text{ ug} / 50 \text{ ul} * 50 / 10$$

$$= 1.13 \text{ ug/ul}$$

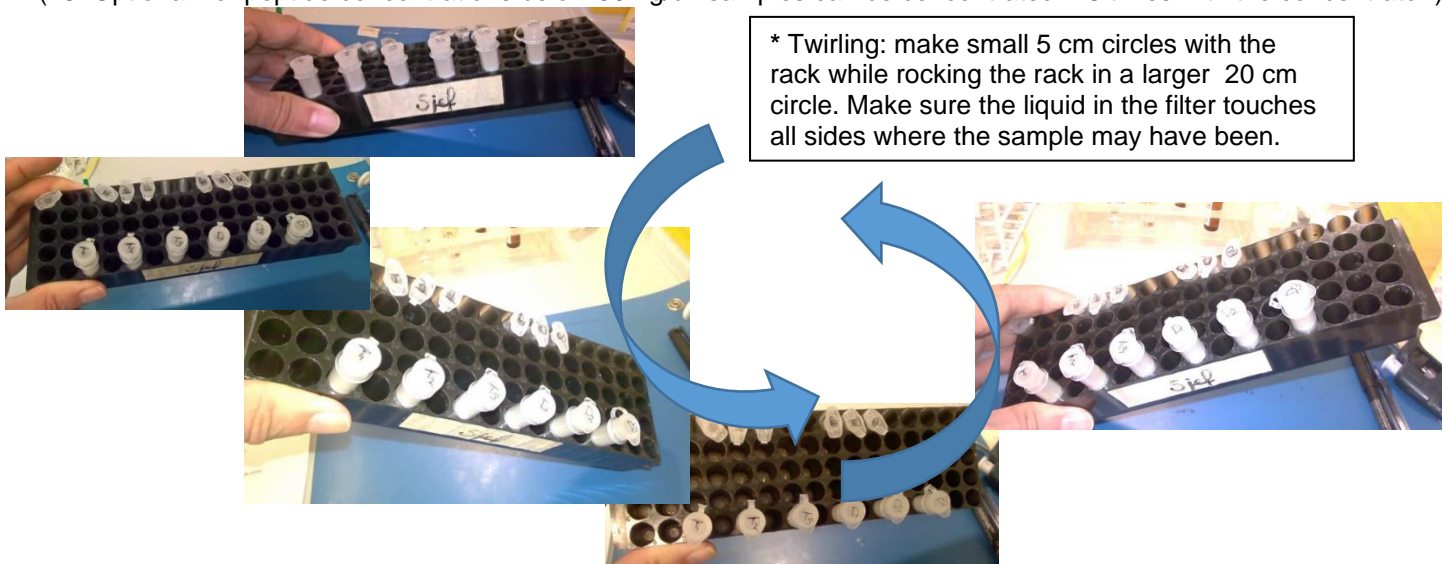
## 3. Gel free protein digestion methods.

### 3.1 Filter aided sample preparation (FASP, 100 ug protein, easy and reliable)

Modified from ref: Wisniewski, J. R.; Zougman, A.; Nagaraj, N.; Mann, M.: Universal sample preparation method for proteome analysis. Nature Methods (2009) Volume: 6, Issue: 5, Pages: 359-360

Remark 1: All centrifugation steps should be done at 12 kRPM in the Eppendorf 5424 centrifuge in lab 3030 ( $\leq 14000 \times g$ )  
Remark 2: Do check whether the filter is liquid free after each centrifugation step. When not, continue centrifugation until the filter is liquid free.

1. Before use, **clean the Pall nanosep filters** (Pall 3K omega (10–20 kDa cut-off, OD003C34) by adding 650  $\mu$ l ethanol and centrifuge at 12 kRPM for 45'. This will remove most Triton X100 present on the filter.
2. **Prepare BCA solutions** as shown in Chapter 2 Protein determination on p5.
3. **Sample lysis: prepare a concentrated protein sample** e.g. by sonicating 10 mg wet weight (washed twice) cells (~100  $\mu$ g protein, from 10 ml bacterial culture with an  $OD_{600} \geq 0.3$ ) or more in 100  $\mu$ l 100 mM Tris pH8 + protease inhibitor cocktail in a 0.5 ml low binding Eppendorf tube for 3 \* 15 sec or more with the Q-sonic cup sonicator (e.g. 15 min 30s on, 30s off at 100% = 30 min total time at 4 °C). (V=100  $\mu$ l)
4. Perform the **acetone precipitation** as a general protein clean-up method (section 1.9) when necessary.
5. **Measure the protein content *in duplo*** using the BCA method (See chapter 2) with e.g. 2 \* 10  $\mu$ l sample. Do the BCA incubation (at 37 °C) and measurement during the reduction and alkylation incubation times of the next 2 steps. (V=80  $\mu$ l)
6. **Reduction:** Add 10% of the volume of 150 mM dithiothreitol (M=154: 150 mM=23 mg/ml water) here: + 8  $\mu$ l --> 14 mM (V=88  $\mu$ l)  
Incubate at 45 °C for 30 min in a thermomixer. Immediately cool to 20 °C afterwards.  
(When using Deoxycholate, DTT reduction will make the solution slightly turbid because of some DC precipitation.)
7. **Unfold:** Add 2\* the volume of fresh prepared and cool 100 mM Tris/HCl pH 8.0 + 8 M urea (0.48g/ml) in a low binding Eppendorf tube. here: + 176  $\mu$ l (V=264  $\mu$ l)
8. **Alkylation:** Add 10% of the volume of 200 mM Acrylamide (M= 71: 14 mg/ml acrylamide in water) here: + 26.4  $\mu$ l --> 18 mM (V=290  $\mu$ l)  
Incubate for 10 min while mildly shaking at room temperature.  
In the meantime, measure the BCA protein determination 96 well plate.
9. **Transfer** 100  $\mu$ g alkylated protein sample as found in the BCA protein determination (e.g. 145  $\mu$ l) to an ethanol washed Pall filter cup (see step 1).  
**Pipet directly on the membrane in the middle without touching the filters poly-propylene side.**  
Centrifuge at 12 kRPM for at least 45 min and carefully check whether all liquid is out of the filter cup.
10. **Wash 1:** Take all samples out of the centrifuge. Check filter content. Remove the liquid from the ep and add 150  $\mu$ l 2 M  $NH_4HCO_3$  (0.32 g / 2 ml) in water to the filter unit. **Twirl\*** the 2M ABC around over the filter. Centrifuge: 12 kRPM for 45'.
11. **Wash 2:** Take all samples out of the centrifuge. Check filter content. Remove the liquid from the ep and add 100  $\mu$ l 70% ethanol/30% water to the filter unit. **Twirl\*** the liquid around over the filter. Centrifuge at 12 kRPM for 30 min.
12. Remove the Pall filter cup from its original micro tube and put it into a clean **2 ml low binding Eppendorf tube**.
13. **Digestion:** Add 100  $\mu$ l 100\* diluted Trypsin/50 mM ABC (5 ng/ $\mu$ l) to the filter and incubate overnight while mildly shaking at room temperature on the shaking platform in the fume hood of lab 3030.
14. Add 10% trifluoroacetic acid (ca 3  $\mu$ l) to adjust the pH of the sample to around pH 3. Mix/twirl immediately.
15. **Centrifuge** the peptides through the membrane at 12 kRPM for 30min.
16. Add 100  $\mu$ l 1 ml/l HCOOH in water, twirl\* over the filter. Centrifuge at 12 kRPM for 30 min.
17. Store samples in a freezer until they will be measured.
- (18: Optional: for peptide concentrations below 50 ng/ $\mu$ l: samples can be concentrated 4-8 times with the concentrator.)



# Sample preparation for proteomics by MS

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## 3.2 Protein aggregation capture (PAC, 100 ug protein)

PAC generally yields more hydrophobic peptides than FASP. This results in more quantified proteins.

Batth, T. S., M. A. X. Tollenaere, P. Ruther, A. Gonzalez-Franquesa, B. S. Prabhakar, S. Bekker-Jensen, A. S. Deshmukh and J. V. Olsen (2019). "Protein Aggregation Capture on Microparticles Enables Multipurpose Proteomics Sample Preparation." *Molecular & Cellular Proteomics* 18(5): 1027-1035.

Dagley, L. F., G. Infusini, R. H. Larsen, J. J. Sandow and A. I. Webb (2019). "Universal Solid-Phase Protein Preparation (USP3) for Bottom-up and Top-down Proteomics." *Journal of Proteome Research* 18(7): 2915-2924.

Modified SP3 from: Hughes, C. S., S. Foehr, D. A. Garfield, E. E. Furlong, L. M. Steinmetz and J. Krijgsveld (2014). "Ultrasensitive proteome analysis using paramagnetic bead technology." *Molecular Systems Biology* 10(10).

Tip: Perform all washes by removing the eppendorf tubes from the magnetic rack and short mixing on a whirlmixer (originally not to be removed). Pulse centrifuge the eps to remove liquid from the cap before removing the liquid from the bottom of the ep.

SpeedBeads (magnetic carboxylate modified particles) = GE Healthcare 45152105050250 + 65152105050250 of 50 ug/uL. For e.g.12 samples, prepare 120 ul taking 60 ul of each type, mix and wash 2\* with 1 ml water and re-suspend into 120 ul water.

Step					Total volume
1 Prepare	Pipet into 2 ml low binding ep	5 ul of each speedbead type per sample			
2	<b>Sample lysis: Prepare a concentrated protein sample</b> e.g. by sonicating 10 mg wet weight (washed twice) cells (~100 ug protein, from 10 ml bacterial culture with an OD <sub>600</sub> ≥ 0.3) or more in 100 ul 100 mM Tris pH8 + protease inhibitor cocktail in a 0.5 ml low binding Eppendorf tube for 3 * 15 sec or more with the Q-sonic cup sonicator (e.g. 15 min 30s on, 30s off at 100% = 30 min total time at 4 °C).				
3	Perform the <b>acetone precipitation</b> as a general protein clean-up method (section 1.9) when necessary.				
4	Pipet 100 ug protein sample in a 2 ml LB tube		µl	60 ul (max 110 ul)	
5	100 mM Tris pH 8		µl	0	Vt = 60 ul
6 <b>Reduction</b>	150 mM DTT (C=14 mM)	M=154: 23 mg/ml	µl	10% of the Volume (here 6 ul)	Vt = 66 ul
7	Incubate at 45 °C for 30', mix	In the meantime: prepare fresh 8M urea: 0.48 g/ml in 100 mM Tris pH8			
8 <b>Unfold</b>	8 M urea / 100 mM Tris	M=60: 0.48 g/ml	µl	3 * total Volume (here 198 ul)	Fresh, cool Vt=264 ul
9 <b>Alkylation</b>	200 mM AcrylAmide (18 mM)	M= 71: 14 mg/ml	µl	10% of the Volume (here 27 ul)	Vt=291 ul
10	Incubate at 21 °C for 30'	In the meantime: wash SpeedBeads 2* in water (see top)			
11	<b>Check pH, adjust to 7</b>	+ 10% TFA	µl	ca 1.25 - 1.5% (here 4 ul)	Vt=295 ul
12 <b>PAC</b>	SpeedBeads	50 ug/ul	µl	8	Vt=303 ul
13	AcNi		µl	2.5 * total Volume (here 750 ul)	
14	Incubate shaking for 20' at rT				
15	Place tubes on a magnetic rack and allow to separate for 30 seconds. Remove all liquid.				
16 <b>Wash</b>	Wash 1 70% ethanol			1000	
17	Wash 2 100% acetonitril			1000	
18 <b>Digestion</b>	100* diluted Trypsin / ABC	5 ng/ul in ABC	µl	100	
19	Incubate overnight while shaking at room temperature				
20	Add 10% TFA	(pH = 3)	µl	3 - 4 ul	
21					
22 <b>Separate</b>	Pulse centrifuge the samples and put them onto a magnet. Tilt the samples with the liquid towards the magnet for at least 20 s. Without taking any beads, transfer the liquid to a 0.5 ml low binding ep.				
23	Wash the beads with 100 ul 1ml/l HCOOH in water. Again, tilt the samples with the liquid towards the magnet for at least 20 s. Without taking any beads, transfer the liquid to the same 0.5 ml low binding ep.				
24	Centrifuge the 0.5 ml low binding eps at 10 kRPM for at least 10 min.				
25	Transfer the liquid to new 0.5 ml low binding eps WITHOUT ANY BEADS.				
26	Concentrate the samples in the concentrator until a low volume.				
27 <b>Reconstitute</b>	Adjust the final volume to exactly 50.0 µl with 1 ml/l HCOOH (or at least to 25.0 µl when your protein starting amount was low)				
	<b>All samples should be 100% particle free and have &lt;4% acetonitrile.</b>				
28	Inject onto nLC-MSMS		µl	Minimally 0.5 - Maximally 5	

## 3.3 On beads digestion.

Ref: Wendrich, J. R., S. Boeren, B. K. Möller, D. Weijers and B. D. Rybel (2017). "In Vivo Identification of Plant Protein Complexes Using IP-MS/MS." *Methods in Molecular Biology* 1497 (Plant Hormones: Methods and Protocols): 147-158.

Beads: can be e.g. Chromotek GFP-Trap Magnetic Agarose for IP's of GFP tagged proteins or Pierce™ Streptavidin Magnetic Beads to pull-down biotinylated proteins.

1. Wash the beads 2\* with 20\* the beads volume of extraction buffer, e.g. 100 mM Tris pH8 + 1% Nonidet P-40;
2. Wash the beads 2\* with 20\* the beads volume of extraction buffer without detergent e.g. 100 mM Tris pH8.
3. Wash the beads 2\* with 20\* the beads volume with 50 mM ABC.
4. Add 90 ul 50 mM ABC
5. Add 10 µL 150 mM DTT (23 mg/ml) in water and incubate gently shaking for 0.5 h at 45 °C.
6. Add 10 µL 200 mM IAA (37 mg/ml) in water to the samples. Mix and incubate gently shaking for 0.5 h at room temperature.
7. Add 10 µL 200 mM cysteine (24 mg/ml) in water to the samples. Mix.
8. Add 10 µL 10\* diluted in ABC sequence-grade trypsin (0.05 µg/ul), mix well, and incubate gently shaking overnight at 20 °C.
9. Stop the digestion by adding (ca 3.5 ul) 10% TFA to decrease the pH to 3.
- 10a. Large µm sized Beads: Without taking any beads, filter all sample through a double C8-Filter into a 0.5 ml Low Binding ep.
- 10b. Small nm sized Miltenyii beads: do the C18 µColumn peptide purification described in section 1.7.
11. Wash the beads with 100 ul 50% AcNi/50% 1m/l HCOOH in water and filter/elute the sup into the same ep.
12. Dry of most of the AcNi/water in the concentrator until a volume of 10 to 15 ul
13. Adjust the final volume to exactly 50.0 µl with 1 ml/l HCOOH (or at least to 25.0 µl when your protein starting amount was low)

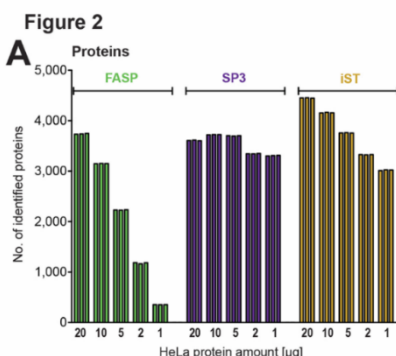
## 3.4 Methanol and TriFluoroEthanol (TFE) sample preparation method

Ref. Wang, H. X, Qian, W. J.; Mottaz, H. M.; ..... , Smith, R. D.: Development and evaluation of a micro- and nanoscale proteomic sample preparation method. *Journal of Proteome Research* (2005) 4 (6) p2397-2403

- 3.4.1. Sonicate in a 2 ml low binding ep 10 mg of wet cell sample in 100 ul of either:
  - a. 50 mM ABC (pH 8),
  - b. MeOH/50 mM ABC (60:40 v/v),
  - c. TFE/50 mM ABC (50:50 v/v)
- 3.4.2. Continue with the reduction and alkylation as above under 3.3.4 – 3.3.9.

## 3.5 Other proteomics sample prep methods:

**SP3:** Hughes, C. S., S. Foehr, D. A. Garfield, E. E. Furlong, L. M. Steinmetz and J. Krijgsveld (2014). "Ultrasensitive proteome analysis using paramagnetic bead technology." *Molecular Systems Biology* 10(10).  
 Improved SP3: Moggridge, S., P. H. Sorensen, G. B. Morin and C. S. Hughes (2018). "Extending the Compatibility of the SP3 Paramagnetic Bead Processing Approach for Proteomics." *J Proteome Res* 17(4): 1730-1740.  
 Improved SP3: Sielaff (2017). "Evaluation of FASP, SP3, and iST Protocols for Proteomic Sample Preparation in the Low Microgram Range." *J Proteome Res* 16(11): 4060-4072 with the figure (2A) shown below:



**iST:** Kulak, N. A., G. Pichler, I. Paron, N. Nagaraj and M. Mann (2014). "Minimal, encapsulated proteomic-sample processing applied to copy-number estimation in eukaryotic cells." *Nature Methods* 11(3): 319-324.

**OmSET:** (modified iST) Kostas, J. C., M. Gregus, J. Schejbal, S. Ray and A. R. Ivanov (2021). "Simple and Efficient Microsolid-Phase Extraction Tip-Based Sample Preparation Workflow to Enable Sensitive Proteomic Profiling of Limited Samples (200 to 10,000 Cells)." *Journal of Proteome Research* 20(3): 1676-1688.

**GASP:** Fischer, R. and B. M. Kessler (2015). "Gel-aided sample preparation (GASP)-A simplified method for gel-assisted proteomic sample generation from protein extracts and intact cells." *Proteomics* 15(7): 1224-1229.

SB: SP3, iST and GASP gave poorer results than PAC in my hands.

**STrap:** Ludwig, K. R., M. M. Schroll and A. B. Hummon (2018). "Comparison of In-Solution, FASP, and S-Trap Based Digestion Methods for Bottom-Up Proteomic Studies." *J Proteome Res* 17(7): 2480-2490.

Zougman, A., P. J. Selby and R. E. Banks (2014). "Suspension trapping (STrap) sample preparation method for bottom-up proteomics analysis." *Proteomics* 14(9): 1006-1010.

SB: STrap uses four C18 plugs + five MK360 quartz plugs = fiber glass. Fibre glass gives a small protein loss (ca 10 ng/disc).

**PROTRAP XG:** Crowell, A. M. J., D. L. MacLellan and A. A. Doucette (2015). "A two-stage spin cartridge for integrated protein precipitation, digestion and SDS removal in a comparative bottom-up proteomics workflow." *Journal of Proteomics* 118: 140-150.

SB: not convenient + can go wrong. Better use acetone precipitation + PAC.



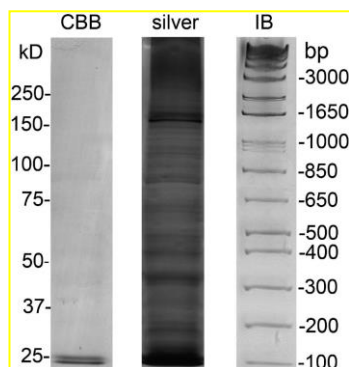
# Sample preparation for proteomics by MS

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## 4. In-Gel Digestion method (IGD, 2 µg for 1 purified protein to 60 µg for a complex mix)

### 4.1 General info

Standard protein gel: 12% bisacrylamide (MW 15 – 200 kDa): e.g. Thermo Bolt 4-12 Bis-Tris gel: NW04120BOX (<https://www.thermofisher.com/order/catalog/product/NW04120BOX>) or from Invitrogen.



Possible internal markers: 2 µg DNA ladder (1 kb plus, Invitrogen 10488-085) added to the sample. Stain after electrophoresis with “Indo-1 blue” DNA stain (Sigma R325147) as described by:

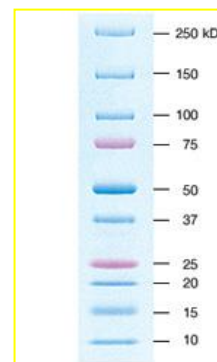
Guoan Zhang, David Fenyö, and Thomas A. Neubert: Use of DNA Ladders for Reproducible Protein Fractionation by Sodium Dodecyl Sulfate-Polyacrylamide Gel Electrophoresis (SDS-PAGE) for Quantitative Proteomics.

Journal of Proteome Research 7 (2008) 2, p678-686

Suggested external markers to get an indication of the size of a sample protein:

Protein dual color markers (Bio-Rad 161-0374) or  
Prestained protein MW marker (ThermoFisher 26619).

Use 10 µl per well.

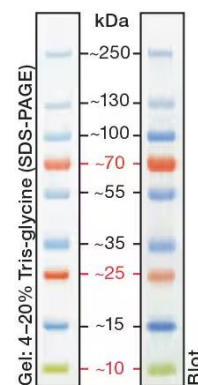


### Comments (by Andrej Shevchenko):

For highest sensitivity, rinse for 60 minutes or more after the gel has been run and fixed. This helps to keep the background transparent during development.

Do not use glutaraldehyde as the sensitizing agent - it is also a protein cross linking agent!

Ref: Shevchenko A., Wilm, M., Vorm, O. and Mann, M. Anal. Chem. T68T, 850-858 (1996). Mass spectrometric sequencing of proteins from silver stained polyacrylamide gels.



### 4.2 Recommended procedure for CCB or Oriole protein gel staining:

1. With the procedure described on the next page, fixation is not necessary. When you still want to fix the proteins, then do so by incubating the entire gel in: 10% acetic acid / 50% water / 40% methanol. Definitely do NOT use cross-linking agents like glutaraldehyde or formaldehyde!
2. Stain either according to the Colloidal Coomassie Staining method (we use the Colloidal Blue Staining Kit which contains Coomassie G-250 (854.02 g/mol), from Invitrogen (ordering# LC6025) described below for visible staining or use the Oriole (Bio-Rad 161-0496 ready for use stain) fluorescent stain to observe bands by UV light. Colloidal Coomassie Staining can be used to see more than 50 ng of protein, Oriole can go down to the low ng range.

#### 4.2.1 Colloidal Coomassie Preparing Staining Solution

Shake the Stainer B solution before using it. Prepare the solutions fresh (from top to bottom = keep this order and mix after each addition) as described in the table below in a 50 ml (Greiner) tube. Then directly transfer it to the gel in a new square petri dish.

Solution*	ml per gel
Deionized Water	13.75
Methanol	5
Stainer B	1.25
Stainer A	5

\*When Stainer A and Stainer B are combined a precipitate may form which will dissolve within 30 seconds.

#### 4.2.2 Colloidal Coomassie Gel staining procedure

1. Shake gel in 25 ml staining solution in a new square petri dish for 1 - 2 hours.  
Note: Staining intensity does not vary significantly if left in stain for 3 hours or 12 hours.
2. Decant staining solution and replace with a minimum of 200 ml of deionized water per gel. Shake gel in water for at least 7 hours. The gel will have a clear background after 7 hours in water.  
Note: Gels can be left in water for up to 3 days without significant change in band intensity and background clarity.
3. For long-term storage (over 3 days), keep the gel in a 10% Methanol solution at 4°C.

## 4.3 In Gel Digestion protocol

### 4.3.1 Remarks

Proteins visible in a SDS gel after Colloidal Coomassie Staining can be measured by Proxeon nLC-LTQ-Orbitrap XL MS-MSMS provided you read and follow the remarks concerning Keratin in section 1.3 Reduce the amount of Keratins in your samples as much as possible by...

### 4.3.2 Procedure

#### 1. Sample preparation.

a. Make a concentrated protein sample in a suitable buffer e.g. by sonication.

E.g. 50 mg washed cells in 450  $\mu$ l SDT (= 4% SDS + 20 mM DTT + 100 mM Tris pH 8 lysis buffer pH 8)  
(= maximally 5  $\mu$ g protein/ $\mu$ l) or 2  $\mu$ l 2.0  $\mu$ g/ $\mu$ l BSA in 398  $\mu$ l SDT-lysis buffer pH 8 (= 10 ng BSA/ $\mu$ l)

b. Sonicate and incubate at 95°C (heating block) for 10 min. Cool down to room temperature.

c. Pipet 80  $\mu$ l of sample in an ep and add 20  $\mu$ l of Sample loading buffer (e.g. Pierce 39000 or [50% glycerol/50% water + 1 g/l bromophenol blue]). Incubate shaking at 95°C for 10 min. Centrifuge 10.000\*g for 2 min.

The sample will now have circa 1 - 4  $\mu$ g protein/ $\mu$ l from 50 mg cells or 8 ng BSA/ $\mu$ l from 2  $\mu$ g BSA

f. Apply **2  $\mu$ g (one purified protein) to 60  $\mu$ g (protein mix)** or even **more** protein per sample well.

Each sample well holds maximally 40  $\mu$ L (but then you have to pipet very, very carefully, 30  $\mu$ l is more safe).

As a protocol control, use 100 ng of BSA exactly (= 12.5  $\mu$ l as prepared above).

**2. Run the gel** as described by the manufacturer. For 12% gels, run at 120V for about 55 min = ca 5.5 cm. This is fine for 8 gel slices. If you want to prepare less slices from one sample, then run the gel shorter (ca 5 min per gel slice).

Use gloves. Open the gel cassette with a thin spatula and put it into a large clean square petri dish. Keep the gel covered with the lid as much as possible to prevent extra keratin contamination.

Stain for 1 - 2 hours (see 5.2) and then de-stain with water for 1 to 20 hours. Refresh the water a few times.

Remark: Some (most) in-gel digestion protocols use a 50% acetonitril to wash away the Coomassie but this is not necessary when the sample is measured by LC-MSMS. The Coomassie stain will be released from the protein during the digestion procedure and will elute from the RP column after the peptides at acetonitril percentages higher than 40% but generally is poorly soluble in 1 ml/l HCOOH in water which is used to dissolve the peptides before injection onto the nLC.

#### 3. Cysteines reduction and alkylation.

a. Add 25 ml 50 mM  $\text{NH}_4\text{HCO}_3$  + 15 mM reductor: 0.057 g DTT.

Gently shake for 1 h at room temperature or at 45 °C for 0.5 h to reduce all disulfide bridges.

b. Wash with water.

c. Add 22.5 ml water + 2.5 ml 1M Tris pH 8 + 0.078 g IodoAcetamide (= 20 mM IAA pH 8).

Incubate at room temperature in the dark while gently shaking for 0.5 hour.

d. Wash with water thoroughly (5\*). DTT-IAA is difficult to remove and gives a large peak in the MS.

**4. Gel cutting.** (If the gel gets a bit dry and starts jumping around, than add a small drop of water on top of the gel).

Cut out the gel bands or slices (1 – 8) and cut them into small pieces of ca. 1 mm<sup>2</sup>. Use a sharp clean scalpel from lab 3030 on a clean piece of parafilm. Transfer the gel pieces to clean 0.5 ml low binding micro centrifuge tubes.

#### 5. Enzymatic digestion.

At this point you may store the samples in a freezer or freeze + de-freeze the gel pieces to further increase the Trypsin accessible area.

Add 50  $\mu$ l cold freshly prepared Trypsin solution (5 ng/ $\mu$ l = 100\* diluted into ABC). When there is still some gel piece sticking out of the solution, then add extra ABC (but NO Trypsin) to completely cover the gel pieces.

Preferentially incubate overnight while shaking at room temperature (20 °C).

#### 6. Extraction of peptides.

a. Add 10% TFA up to a pH between 2 and 4 (measure the pH with pH paper, ca. 3.5  $\mu$ l is needed per 50  $\mu$ l of ABC). Mix.

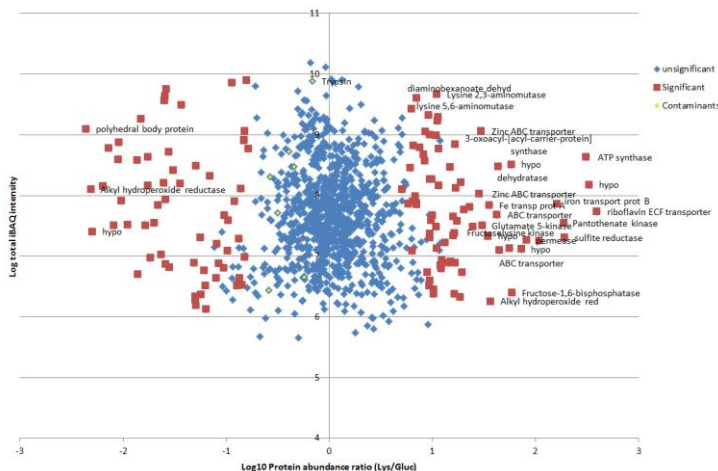
b. Put all sample on top of a double C8 filter mounted into a 200  $\mu$ l pipet tip. Elute all sample into a 0.5 ml low binding ep. Add 100  $\mu$ l 50% acetonitrile/1 ml/l HCOOH in water to the remaining gel pieces, mix, and add the liquid to the C8 filter and elute all sample into the same 0.5 ml low binding ep.

Concentrate the sample in the Eppendorf concentrator to a volume of 10 - 15  $\mu$ l and add 1 ml/l HCOOH in water up to 50.0  $\mu$ l.

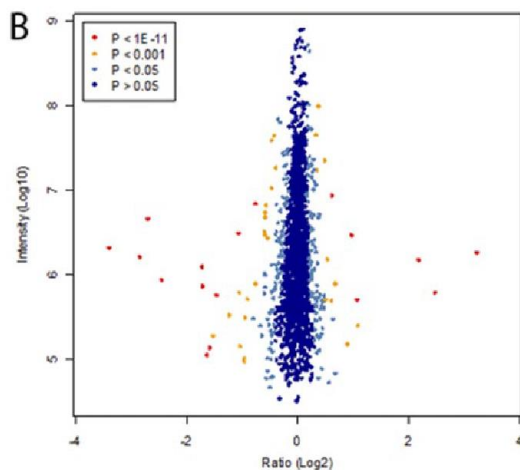
## 5. Quantitation

### 5.1 Relative quantitation (= Sample compared to Control)

Three ways of relative quantitation can be used. The easiest to do is **label free relative quantitation**. In this case, samples (to be measured at least in triplo, better in 4 fold or more) and Controls (the same amounts) are measured separately. Relative quantitation takes place after calculation of all peak intensities (by MaxQuant) in each chromatogram. This tricky method became feasible due to the high accuracy and low noise of the Orbitrap's MS measurement and release of the MaxQuant software. The statistical program Perseus can be used to find the really significant differences between samples and controls in the MaxQuant result table. The accuracy strongly depends on the sample preparation reproducibility but generally starts from a factor 10 to find significantly different protein concentrations between sample and control.



Bui, T. P., J. Ritari, S. Boeren, P. de Waard, C. M. Plugge and W. M. de Vos (2015). "Production of butyrate from lysine and the Amadori product fructoselysine by a human gut commensal." Nat Commun 6: 10062.



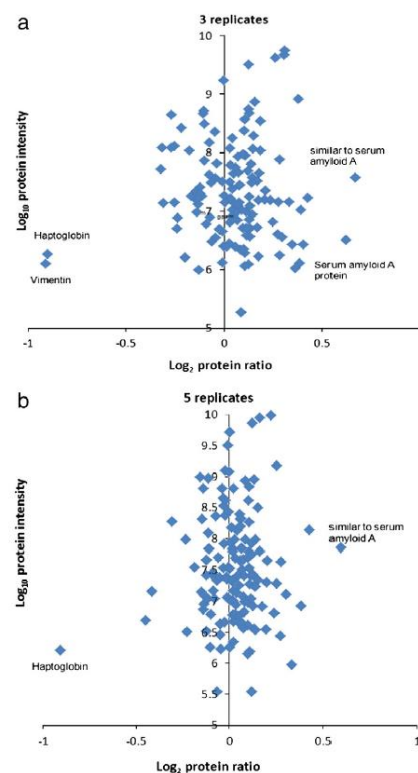
**SILAC labeling.** The essential amino acids arginin and lysin are added in Light, Intermediate and Heavy form, and these labeled amino acids become incorporated during cell growth. Different samples can therefore be mixed in a very early stage, that is directly after cell lysis and protein determination. Here, protein losses occurring during sample preparation will happen for every labeled protein (light/intermediate and heavy), and therefore will not affect the final result. Silac labeling can be regarded as the most accurate relative quantitation method (Accuracy +/- 30%, you may find significantly different protein concentrations between sample and control from a factor 1.5).

Sotoca, A. M., M. D. S. Gelpke, S. Boeren, A. Strom, J. A. Gustafsson, A. J. Murk, I. M. C. M. Rietjens and J. Vervoort (2011). "Quantitative Proteomics and Transcriptomics

Addressing the Estrogen Receptor Subtype-mediated Effects in T47D Breast Cancer Cells Exposed to the Phytoestrogen Genistein." Molecular & Cellular Proteomics 10(1).

Sometimes Silac labeling is impossible though labeling may be desired. Then, samples can also be labeled on the peptide level. So far, we have obtained best results with **Dimethyl labeling** of N-terminal amines and lysines. This is a reductive alkylation method that uses formaldehyde ( $\text{CH}_2\text{O}$ ,  $\text{CD}_2\text{O}$  or  $^{13}\text{CD}_2\text{O}$ ) and cyanoborohydride ( $\text{NaBH}_3\text{CN}$  or  $\text{NaBD}_3\text{CN}$ ) and is described in detail below. Disadvantage of labeling at the peptide level is that almost the complete sample preparation has to be done for each sample separately. Different samples are mixed after completion of the sample preparation which may result in a large error. Fortunately, not all proteins will be up- or down regulated due to the stimulus so an internal control should normally be possible (Accuracy: +/- 50%, you may find significantly different protein concentrations between sample and control from a factor 2).

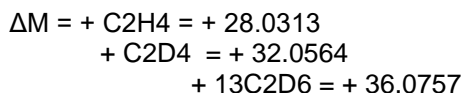
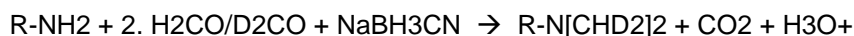
Lu, J., S. Boeren, S. C. de Vries, H. J. F. van Valenberg, J. Vervoort and K. Hettinga (2011). "Filter-aided sample preparation with dimethyl labeling to identify and quantify milk fat globule membrane proteins." Journal of Proteomics 75: 340.



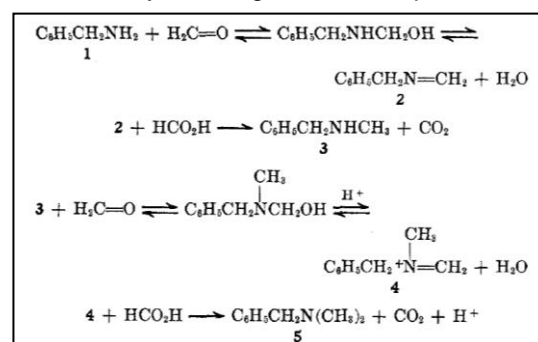
## 5.1.1 Relative quantitation by on column peptide dimethyl labelling protocol

Ref. Boersema, P.J. et al., Nature Protocols (2009) 4, 4, p484-494: Protocol: Dimethyl labeling for relative quantitation.

Reductive amination of NH<sub>2</sub> on N-terminus and lysines with aldehyde:



dΔM = 4 / 8 Da per group



### Stock solutions

500 mM NaH <sub>2</sub> PO <sub>4</sub> ·1 H <sub>2</sub> O	(M= 138)	6.9 g/l	3.45 g / 50 ml
500 mM Na <sub>2</sub> HPO <sub>4</sub>	(M= 142)	7.1 g/l	3.55 g / 50 ml
35% or 20% (vol/vol) formaldehyde in water (CH <sub>2</sub> O, CD <sub>2</sub> O or <sup>13</sup> CD <sub>2</sub> O)			
0.6 M cyanoborohydride in water (NaBH <sub>3</sub> CN or NaBD <sub>3</sub> CN): (M= 64.8)			3.9 mg / 100 ul

### Labeling reagent

Prepare per sample/label:

10 ul 500 mM NaH<sub>2</sub>PO<sub>4</sub>·1 H<sub>2</sub>O + 35 ul 500 mM Na<sub>2</sub>HPO<sub>4</sub> + 425 ul H<sub>2</sub>O (= 50 mM pH 7.5)

+ 2.9 ul 35% or 5 ul of 20% (vol/vol) formaldehyde in water (CH<sub>2</sub>O, CD<sub>2</sub>O or <sup>13</sup>CD<sub>2</sub>O).

+ 25 ul of 0.6 M cyanoborohydride in water (NaBH<sub>3</sub>CN or NaBD<sub>3</sub>CN) or 1 mg.

CAUTION Formaldehyde solutions and formaldehyde vapors are toxic, prepare solutions in a fume hood.

CRITICAL Labeling reagent mixtures should be kept at 4 °C and not stored longer than 24 h to ensure labeling efficiency.

Light: CH<sub>2</sub>O + NaBH<sub>3</sub>CN

Intermediate: CD<sub>2</sub>O + NaBH<sub>3</sub>CN

Heavy: <sup>13</sup>CD<sub>2</sub>O + NaBD<sub>3</sub>CN

### On-column stable isotope dimethyl labeling (TIMING: 1 h):

(i) Make C18+ Stage tips by: Use the "cookie cutter" method to put a small (1.6mm) piece of a C18 Empore disk into a plastic 200 ul tip. Do this by cutting the Empore disk with a large metal needle and transfer it to the 200 ul tip with a good fitting plunger from a syringe. Add 200 ul of methanol to this Stage tip. Add 5 ul of a 50% LichroprepC18 slurry in methanol to the Stage tip to create a C18+ Stage tip.

(ii) Wash the C18+ Stage tip column with 200 ul of methanol.

(iii) Condition the C18+ Stage tip column with 100 ul of 1 ml/l HCOOH in water.

(iv) Load the acidified peptide sample in water (maximally 10 ug peptide, detergent free) on a C18+ Stage tip column.

(v) Wash the C18+ Stage tip column with 100 ul of 1 ml/l HCOOH in water.

(vi) Put 2 ml Eppendorf tubes under the columns to catch reagent that runs through the column.

(vii) In 10 min. time (not faster), flush each of the C18+ Stage tip columns with 100 ul of the respective labeling reagent (light, intermediate or heavy).

CRITICAL STEP To allow for complete labeling, make sure that Step vii takes at least 10 min.

(viii) Wash the C18+ Stage tip columns with 200 ul of 1 ml/l HCOOH in water.

(ix) Add 10 ul 1 M Tris to the eps under the columns to destroy remaining reagent.

### Peptide recovery

Manually (= with a syringe) elute and collect the labeled samples in new 0.5 ml low binding eps from the C18 Stage tip columns with 50 ul of 50% AcNi/50% 1 ml/l HCOOH in water.

CRITICAL STEP When performing the protocol for the first time or with a new sample, it is advised to check the labeling efficiency and sample amounts by measuring a fraction of the sample by LC-MS before mixing differentially labeled samples.

For LCMS analysis, reduce the AcNi content by putting the samples in a Concentrator (with open cap) at 45 °C for 2 hours or longer when necessary. The final volume should be below 20 ul. Adjust the sample volume with 1 ml/l HCOOH in water to exactly 100.0 ul. Sonicate (water bath sonicator) for 5 sec in the hot spot when the sample had been dried completely by accident.

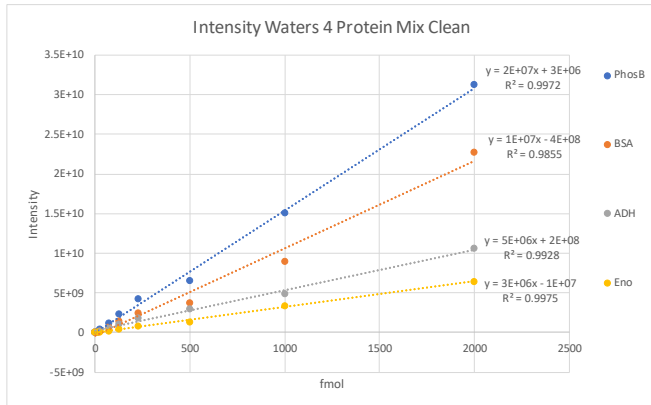
The sample now has <5% AcNi, is pH 3 and is particle free. Mix the samples in the appropriate ratio (light/intermediate/heavy) to make them nLCMS ready.

# Sample preparation for proteomics by MS

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## 5.2 Absolute quantitation

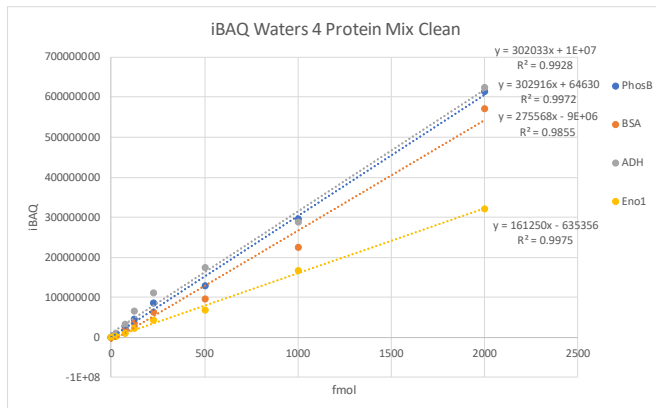
a. Absolute quantitation on a limited number of proteins can be done by making **calibration curves with synthetic peptides that preferentially contain 1 <sup>13</sup>C labeled amino acid**, e.g. the C-terminal K or R in case of a tryptic peptide. To do this, the (HPLC or NMR quantified) labeled peptide has to be added to the sample to generate the calibration curve under exactly the same conditions as the sample measurements. Later, the labeled peptide can also serve as an internal standard. This way of absolute quantitation generally gives a good accuracy (+/- 30%).



b. Less stringent but more practical when more proteins have to be quantitated, is the method of quantitation with respect to a single added internal standard like BSA (e.g. Pierce [PI23210](#) 2.0 mg/ml) or Ovalbumin. This rough absolute quantitation **“relative to an internal standard”** is accurate within a factor of 4.

BSA can be replaced by Waters MassPREP Digestion Standard Mix 1 [186002865] (a 4 protein mix, €200/set) or Sigma UPS2, a quantified set of 48 human proteins at different concentrations (€900/set).

Lahtvee, P.J.: Absolute quantification of protein and mRNA abundances demonstrate variability in gene-specific translation efficiency in yeast. *Cell Syst.* 4, 495–504.e5 (2017).

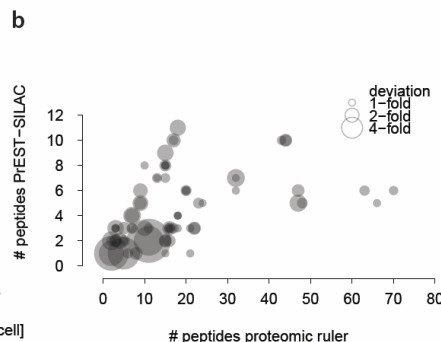
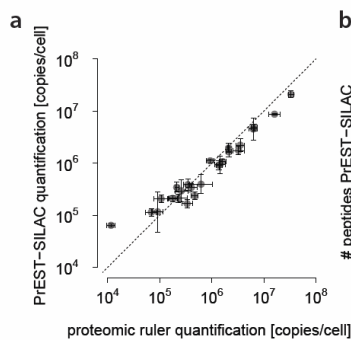
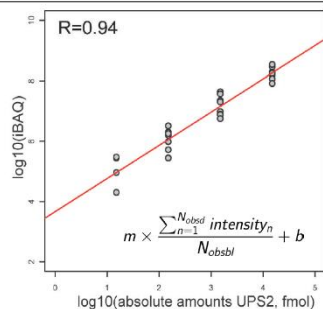


c. Somewhat less accurate but very easy to implement is the **“intensity based absolute quantification (iBAQ)”** method (Schwanhauser et al. *Nature* 2011, 473, 7347, P337-342) that uses the Total peak intensity as determined by MaxQuant for each protein and corrects that for the number of measurable peptides (=number of tryptic peptides of 7-30 amino acids long without missed cleavages).

The error in the iBAQ method is between a factor 2 (as shown to the left) and 10 as shown in Log<sub>10</sub> iBAQ vs Log [Protein] plot.

The Log-Log figure comes from a presentation by Selbach during the MaxQuant summerschool 2011. It has been published as Fig. S8 in Schwanhauser 2011 mentioned above.

Absolute protein levels: intensity-based absolute quantification (iBAQ)



d. For very large datasets with >12.000 peptides identified, the absolute amount of a protein per cell can be calculated by relating the MS signal intensity from a protein to the total MS signal intensities of all histones observed. Error = a factor 2 only (?).

Wisniewski (2014): A ‘proteomic ruler’ for protein copy number and concentration estimation without spike-in standards. *Molecular & Cellular Proteomics* 13(12): 3497-3506.

# Sample preparation for proteomics by MS

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e. Addition of labeled peptides produced by either Cell free expression systems (Takemori: MEERCAT Multiplexed Efficient Cell Free Expression of Recombinant QconCATs For Large Scale Absolute Proteome Quantification. *Mol Cell Proteomics*. 2017, 16(12):2169-2183. SB ca €5000) or Ecoli (Beynon, R. J., Doherty, M. K., Pratt, J. M., and Gaskell, S. J. (2005) Multiplexed absolute quantification in proteomics using artificial QCAT proteins of concatenated signature peptides. *Nat. Methods* 2, 587–589 SB ca €3000).