Mass Spectrometry-Based Absolute Quantification of Single Xenopus Embryo Proteomes

Rik G.H. Lindeboom,1,4 Arne H. Smits,2,4 Matteo Perino,3 Gert Jan C. Veenstra,3 and Michiel Vermeulen1,5

1Department of Molecular Biology, Faculty of Science, Radboud Institute for Molecular Life Sciences, Radboud University, Nijmegen 6500 HB, The Netherlands; 2Genome Biology Unit, European Molecular Biology Laboratory, 69117 Heidelberg, Germany; 3Department of Molecular Developmental Biology, Faculty of Science, Radboud Institute for Molecular Life Sciences, Radboud University, Nijmegen 6500 HB, The Netherlands

Early Xenopus development is characterized by a poor correlation between global mRNA and protein abundances due to maternal mRNA and protein loading. Therefore, proteome profiling is necessary to study gene expression dynamics during early Xenopus development. In contrast to mammals, single Xenopus eggs and embryos contain enough protein to allow identification and quantification of thousands of proteins using mass spectrometry-based proteomics. In addition to investigating developmental processes, single egg or blastomere proteomes can be used to study cell-to-cell variability at an unprecedented depth. In this protocol, we describe a mass spectrometry-based proteomics approach for the identification and absolute quantification of Xenopus laevis egg or embryo proteomes, including sample preparation, peptide fractionation and separation, and data analysis.

MATERIALS

It is essential that you consult the appropriate Material Safety Data Sheets and your institution’s Environmental Health and Safety Office for proper handling of equipment and hazardous materials used in this protocol.

RECIPES: Please see the end of this protocol for recipes indicated by <R>. Additional recipes can be found online at http://cshprotocols.cshlp.org/site/recipes.

Reagents

ABC buffer (50 mM ammonium bicarbonate, freshly prepared)
Britton-Robinson (B&R) buffer (pH 11/pH 8/pH 2) <R>
Buffer A (0.1% formic acid)
Buffer B (80% acetonitrile, 0.1% formic acid)
C18 disks (Empore) and reagents for preparation of C18 StageTips (see Steps 11–14 of Protocol: Enrichment of Phosphopeptides via Immobilized Metal Affinity Chromatography [Swaney and Villen 2016])
Cysteine (3% in 0.25× MMR)
IAA buffer <R> (freshly prepared, kept in the dark)
MMR buffer (0.25×, pH 7.4) <R>
NaCl (0.5 M)

1These authors contributed equally to this work.
5Correspondence: michiel.vermeulen@science.ru.nl
From the Xenopus collection, edited by Hazel L. Sive.
© 2019 Cold Spring Harbor Laboratory Press
Advanced Online Article. Cite this protocol as Cold Spring Harb Protoc; doi:10.1101/pdb.prot098376
METHOD

This method is based on the work of Smits et al. (2014) (Fig. 1). Yolk-deprived egg or embryo extracts are first digested with trypsin using the so-called Filter-Aided Sample Preparation (FASP) protocol (Wisniewski et al. 2009a) followed by SAX fractionation (Wisniewski et al. 2009b). A protein spike-in is used to facilitate quantification of the protein molecules (Schwanhausser et al. 2011). After sample preparation, peptides are measured in a mass spectrometer. Finally, data analysis is performed using the MaxQuant software suite (Cox and Mann 2008).

Embryo Collection and Lysis

1. Dejelly fertilized eggs by adding 3% cysteine in 0.25× Marc’s modified Ringer’s (MMR). Wash thoroughly four times with 0.25× MMR as soon as the embryos start touching each other.
2. Stage the embryos according to Nieuwkoop and Faber (1994), and collect single embryos in separate tubes. Remove any carryover MMR.

3. Add 20 µL of WCE-LS buffer per sample and homogenize the embryos by gentle pipetting.

4. Centrifuge the samples at 3500g for 5 min at 4°C.

5. Transfer the supernatant to a fresh tube without touching the yolk/pigment pellet.

Filter-Aided Sample Preparation

All of the following steps are performed at 20°C.

6. Add UA buffer to the supernatant from Step 5 to a total volume of 220 µL. Incubate with gentle shaking for 5 min in a thermostaker.

7. Transfer 20 µL of the lysate in UA buffer to a new tube. This is the FASP-iBAQ sample, which is not fractionated and is used for absolute quantification. The residual 200 µL is the FASP-SAX sample, which is subjected to fractionation to obtain deep proteome coverage.

8. Add 2 µL of UPS2 standard (0.5 µg/µL) and 178 µL of UA buffer to the FASP-iBAQ sample and mix.

9. Transfer both samples (FASP-iBAQ and FASP-SAX) to two 30-kDa filters. Centrifuge at 11,600g for 15 min. Discard the flowthrough (FT).

10. Add 100 µL of IAA buffer to each filter. Incubate for 1 min in a thermostaker at 600 rpm.

11. Incubate the filters for an additional 20 min protected from light and without shaking. Centrifuge at 11,600g for 15 min.

12. Wash the samples by adding 100 µL of UA buffer to each filter. Centrifuge at 11,600g for 15 min.

13. Repeat Step 12 two times. Discard the FT.

14. Wash the samples by adding 100 µL of ABC buffer to each filter. Centrifuge at 11,600g for 10 min.

15. Repeat Step 14 two times. Discard the FT.

16. Add trypsin to each filter as follows.

   i. To the FASP-iBAQ sample, add 0.1 µg of trypsin in ABC buffer to a total volume of 40 µL.
   ii. To the FASP-SAX sample, add 1.0 µg of trypsin in ABC buffer to a total volume of 40 µL.

17. Mix the samples at 600 rpm for 1 min and then incubate in a sealed plastic box containing moist paper towels overnight at 37°C.

18. Transfer each filter to a new tube. Centrifuge at 11,600g for 15 min. Retain the FT. Steps 18–20 are performed in the same tube. The FT from these steps are combined during centrifugation.

19. Process each filter as follows.

   i. To the FASP-iBAQ sample, add 50 µL of 0.5 M NaCl. Centrifuge at 11,600g for 10 min.
   ii. To the FASP-SAX sample, add 50 µL of ABC buffer. Centrifuge at 11,600g for 10 min.

20. Repeat Step 19.

   The FT of ~140 µL from each filter represents the tryptic digest of each sample.

Strong Anion eXchange Fractionation (FASP-SAX Sample Only)

21. Prepare a 20 plug SAX StageTip according to Rappsilber et al. (2007).

22. Wash the StageTip by adding 200 µL of B&R buffer (pH 11). Centrifuge for 24 min at 1000g.

23. Add 140 µL of B&R buffer (pH 11) and 10 µL of 1 M NaOH to the FASP-SAX sample and mix.

   The pH should be between 11 and 12.
24. Transfer the FASP-SAX sample to the StageTip.

25. Centrifuge the sample for 45 min at 500g. Retain the FT.
   
   The FT represents the FASP-SAX FT fraction.
   
   See Troubleshooting.

26. Transfer the StageTip to a new tube. Add 200 µL of B&R buffer (pH 11) to the StageTip. Centrifuge for 24 min at 1000g. Retain the FT.

   The FT represents the FASP-SAX (pH 11) fraction.

27. Repeat Step 26 with 200 µL of B&R buffer (pH 8) to obtain the FASP-SAX (pH 8) fraction.

28. Repeat Step 26 with 200 µL of B&R buffer (pH 2) to obtain the FASP-SAX (pH 2) fraction.

**Peptide Desalting and Elution**

*Use all samples (1 FASP-iBAQ + 4 SAX fractions).*

29. Prepare, wash and activate five C18 StageTips (or filtration tips) as described in Protocol: Enrichment of Phosphopeptides via Immobilized Metal Affinity Chromatography (Swaney and Villen 2016) (Steps 11–14).

30. Add 10 µL of 10% TFA to each sample. Apply each sample to a separate C18 StageTip.

31. Wash each StageTip by adding 30 µL of buffer A. Push the buffer slowly through the StageTips with an airtight, fitted Combitip.

32. Elute the peptides of each sample in a separate 0.2-mL tube by adding 30 µL of buffer B to each StageTip and pushing the buffer slowly through with a Combitip.

33. Concentrate the samples to ~5 µL with a speed vacuum concentrator. Add 7 µL of buffer A to each sample (final volume ~12 µL).

**Mass Spectrometry**

34. Inject 5 µL of each sample into the nano-HPLC coupled to a C18 column of 25–30 cm.

   The nano-HPLC acetonitrile gradient varies between the different fractions (Table 1).

35. Acquire mass spectra on a high-resolution mass spectrometer, e.g., time-of-flight or Orbitrap mass analyzer.

   For examples of optimized nano-HPLC and mass analyzer settings, see Richards et al. (2015).

**Data Analysis**

36. Perform data analysis using the MaxQuant software suite (www.maxquant.org). Specify for each FASP-SAX sample the same experiment name but different fraction numbers (assign uneven numbers only). Assign the FASP-iBAQ sample its own experiment name.

37. Upload the FASTA files with protein sequences of *Xenopus* and UPS2 proteins.

   Protein databases from Smits et al. (2014), Wahr et al. (2014) or UniProt can also be used as a reference for the *Xenopus* proteome. A FASTA file containing the sequences of the UPS2 proteins can be downloaded from the supplier’s webpage.

| TABLE 1. Linear nano-HPLC acetonitrile (ACN) gradients for different samples |
|-----------------------------|---------------|--------------|--------------|
| Sample                     | % ACN at 0 min | % ACN at 214 min | % ACN at 240 min |
| FASP-iBAQ                  | 4             | 26           | 76           |
| FASP-SAX FT fraction       | 4             | 14           | 76           |
| FASP-SAX pH11 fraction     | 6             | 17           | 76           |
| FASP-SAX pH8 fraction      | 7             | 18           | 76           |
| FASP-SAX pH2 fraction      | 9             | 21           | 76           |

Advanced Online Article. Cite this protocol as Cold Spring Harb Protoc; doi:10.1101/pdb.prot098376
38. Empty the contents of the “ibaq.txt” file in the `1.X.X.X_MaxQuant\bin\conf\` folder.

39. Make sure to enable the “iBAQ” option to facilitate absolute quantification and enable the “match-between-run” option to share peptide identification information between runs.

40. After the MaxQuant analysis, find the quantified protein abundances in the `proteinGroups.txt` file.

   The relative iBAQ quantification of the UPS2 proteins in the FASP-iBAQ sample can be used for absolute quantification of all identified proteins in the fractionated sample.

   - A linear regression between the supplied concentrations of the spike-in proteins and the measured iBAQ values can be used to calculate the abundance in femtomoles for all proteins identified in the FASP-iBAQ sample.
   - A second linear regression between the absolute abundances of the quantified proteins in the FASP-iBAQ sample and the iBAQ values of same proteins in the FASP-SAX sample can then be used to extrapolate the absolute amounts of all identified proteins in the FASP-SAX sample.

   See Troubleshooting.

**TROUBLESHOOTING**

**Problem (Steps 9 and 25):** The 30-kDa filters or SAX StageTips are not completely cleared after centrifugation.

**Solution:** It is essential that the 30-kDa filters and StageTips are completely cleared before continuing to the next step. Increase the centrifugation time accordingly.

**Problem (Step 40):** The peptide identification rates are low after MaxQuant analysis.

**Solution:** To boost low peptide identification rates, keep temperature stable in Steps 6–13 to prevent peptide carbamylation by urea buffers.

**DISCUSSION**

While RNA-sequencing is often used to study genome-wide gene expression, it is not sufficient when studying early *Xenopus* development due to decoupled proteome and transcriptome dynamics (Smits et al. 2014; Peshkin et al. 2015). Sample preparation of early-stage *Xenopus* embryos for mass spectrometry requires proper removal of yolk and protein solubilization with strong detergents (Peuchen et al. 2016). Recent advances in the field of mass spectrometry-based proteomics allowed the study of early *Xenopus* proteome dynamics at an unprecedented depth (Sun et al. 2014; Wuhr et al. 2014). The relatively large protein content also enabled the first single egg and embryo deep-proteome studies (Smits et al. 2014; Lombard-Banek et al. 2016; Sun et al. 2016).

**RECIPES**

**Britton-Robinson (B&R) Buffer (pH 11/pH 8/pH 2)**

40 mM H₃PO₄
40 mM CH₃COOH
40 mM H₃BO₃

Titrated buffer to pH 11, pH 8, or pH 2 with 1 M NaOH. Store for up to 2 yr at room temperature.
**IAA Buffer**

UA buffer <R> (freshly prepared)
50 mM iodoacetamide (IAA)
Prepare fresh before use and keep in the dark.

**MMR Buffer (0.25×, pH 7.4)**

22 mM NaCl
0.5 mM KCl
0.5 mM CaCl₂
0.25 mM MgCl₂
1.25 mM HEPES
Store for up to 2 yr at room temperature.

**UA Buffer**

8 M urea
0.1 M Tris-HCl (pH 8.5)
50 mM dithiothreitol
Prepare fresh before use.

**UPS2 Standard Solution (0.5 µg/µL)**

UA buffer <R> (freshly prepared)
UPS2 standard (Sigma-Aldrich) (0.5 µg/µL)
Dissolve the UPS2 standard (0.5 µg/µL) in UA buffer for >30 min with gentle shaking at 20°C.
Prepare fresh before use.

**WCE-LS (Whole Cell Extract-Low Salt) Buffer**

20 mM Tris-HCl (pH 8.0)
70 mM KCl
1 mM EDTA
10% glycerol
0.1% IGEPAL CA-630
5 mM dithiothreitol (DTT)
1× cOmplete EDTA-Free Protease Inhibitors (Roche)
Prepare fresh before use.

**REFERENCES**


Mass Spectrometry-Based Absolute Quantification of Single *Xenopus* Embryo Proteomes

Rik G.H. Lindeboom, Arne H. Smits, Matteo Perino, Gert Jan C. Veenstra and Michiel Vermeulen

*Cold Spring Harb Protoc*; doi: 10.1101/pdb.prot098376; published online August 13, 2018

To subscribe to *Cold Spring Harbor Protocols* go to: http://cshprotocols.cshlp.org/subscriptions