SEC-MALS: CMI Guide to Protein Conjugate Analysis

dn/dc Sample Preparation dn/dc Data Collection Protein Conjugate Analysis

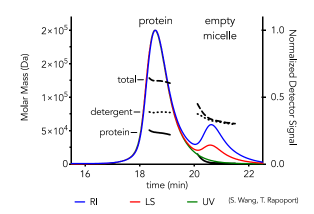
Introduction

MALS calculations are dependent on a good estimate of the refractive index increment (dn/dc) of the sample. All proteins have a dn/dc near to 0.185. However, common modifiers, such as glycans or detergent micelles, have different dn/dc values. To calculate accurate masses from MALS data for modified proteins, the fractional mass contribution of the protein and the modifier must be deconvoluted. A method called Protein Conjugate Analysis is applied to SEC-MALS data to perform these calculations. Protein Conjugate Analysis requires that two concentration detectors (RI and UV) are used simultaneously during the MALS data collection and that the dn/dc, and UV extinction coefficient of each component are known. With sufficient pure material (e.g. detergent), dn/dc values can be determined using a batch refractive index experiment.

$$\frac{\left(\frac{dn}{dc}\right)_{conjugate}}{\left(\frac{dn}{dc}\right)_{p} \cdot X_{p}} + \frac{\left(\frac{dn}{dc}\right)_{m} \cdot (1-X_{p})}{\left(\frac{dn}{dc}\right)_{m} \cdot (1-X_{p})}$$

$$\epsilon_{conjugate} = \epsilon_{p} \cdot X_{p} + \epsilon_{m} \cdot (1-X_{p})$$

 X_p = weight fraction protein dn/dc = refractive index increment (ml/g) ϵ = UV extinction coefficient (mg/ml•cm)



Standard dn/dc values

Protein 0.185 ml/g
 Glycan 0.145 ml/g
 DNA 0.17 ml/g

Detergents Variable (check supplier for value or measure it yourself, see below)

References

Strop, P., & Brunger, A. T. (2005). Refractive index-based determination of detergent concentration and its application to the study of membrane proteins. *Protein Science: a Publication of the Protein Society*, 14(8), 2207-2211.

Barer, R., Ross, K. F. A., & Tkaczyk, S. (1953). Refractometry of Living Cells, 171(4356), 720 EP —724. Pasternack, S. G., Veis, A., & Breen, M. (1974). Solvent-dependent changes in proteoglycan subunit conformation in aqueous guanidine hydrochloride solutions. *The Journal of Biological Chemistry*, 249(7), 2206–2211.

Instrument Overview

The CMI has a Wyatt WISH module, a high pressure manual injection system with 1 ml injection loop. This module is attached downstream of the chromatography pump and upstream of the refractive index detector to control batch measurements of differential refractive index.

Required Supplies

• Syringe(s) for filling the injection loop.

dn/dc Sample Preparation

Assay Buffers

- Water or Buffer is needed for the pump and to dissolve and dilute sample.
- Running buffer and Dilution buffer are typically the same (but can differ).
 - o Water is often used for both the pump and to dissolve sample.
 - o Prepare at least 200 ml of Running buffer.
 - o If different from Running buffer, prepare sufficient Dilution buffer to dissolve and dilute your samples and for two blank injections (to normalize the RI signal).

Samples

- Typically detergent, but could be performed for any sample type (even protein) provided you have sufficient quantity of pure sample.
- Accurate concentration measurements are critical for dn/dc measurements
 - o Collect replicate data (2-3 replicates).
 - Weigh your sample out independently for each replicate.
 - Prepare least 2 ml of 6-7 concentrations of your sample, in replicate.
 - o Use a high concentration of ~5 mg/ml and a low concentration of ~0.1 mg/ml.
 - o Prepare a concentration series (e.g. 5, 2.5, 1.25, 0.625, 0.3125, ... mg/ml).

dn/dc Data Collection

- 1. Book time on the PPMS calendar before you start.
- 2. Login to the computer using your PPMS credentials (eCommons ID and password).
- 3. Attach the Wyatt WISH module to the Agilent chromatography system and to the Optilab TrEX RI detector.
 - a. Make sure there is NO COLUMN inline on the HPLC system
 - b. Attach the outlet on from HPLC system UV detector (or other convenient location downstream of the pump) to the inlet on WISH.
 - c. Attach the outlet on WISH to the inlet on the Optilab TrEx.
- 4. Equilibrate the RI detector:
 - a. Purge pump on the Agilent chromatography system with water (or buffer) for 5 min at 5 ml/min flow rate.
 - b. Turn on the flow at 0.5 ml/min.
 - c. Rinse the WISH module in both inject and load positions for several minutes.
 - d. Rinse the RI detector while the purge valve is open for at least 30 min.
 - e. Close the purge valve.

- f. Zero the RI detector.
- 5. Open Astra software.
- 6. Write a Batch method.
 - a. File \rightarrow New \rightarrow Experiment from Method.
 - i) Go to System \rightarrow Method \rightarrow RI Measurement \rightarrow Batch (determine dn/dn).
 - ii) Or Find a batch dn/dc method in the user methods folder.
 - b. Open Basic Collection (double click).
 - c. Set calculated duration to 60 min (or longer).
- 7. Start Method in Astra:
 - a. Click Run button to start method.
 - b. Click OK when prompted (after pump has started).
- 8. Inject the buffer for several minutes to establish a baseline.
- 9. Inject samples (from low to high concentration):
 - a. In the load position, inject the sample into WISH module (must use over 1 ml to ensure that the loop is filled).
 - b. Turn valve to the inject position for sample injection.
 - c. After injection is complete (or nearly complete, turn valve to load position.
 - d. Repeat injections for each concentration.
 - e. Run duplicate measurements, either in two experiments, or in one continuous experiment.
- 10. Inject the buffer again for 5-10 minutes.
- 11. Stop the Method once finished.
- 12. Draw Baseline:
 - a. Open Baselines.
 - b. Draw a baseline between the two regions of buffer injection.
 - c. Click OK.
- 13. Select Peaks:
 - a. Open Peaks.
 - b. Click and drag through a flat region to define each peak (usually the 2nd half of the injection peak).
 - c. Enter the concentration in mg/ml for each injection.
 - d. Click OK.
- 14. Extract dn/dc:
 - a. Open dn/dc from RI.
 - b. Examine the linear fit.
 - c. Get the dn/dc value.
- 15. Wash system
- 16. Return system to Storage buffer.
 - a. Turn off RI LED.
 - b. Purge the RI detector.
 - i) Turn Purge "ON" on the RI detector LCD screen to open the reference cell so it is washed during equilibration.
 - ii) Leave the purge valve ON at all times, except during data collection
 - c. Stop flow in HPLC Manager.

- d. Purge the Pump:
 - i) Change the solvent to storage buffer (Filtered water with 0.02 % NaN₃).
 - ii) Open the purge valve on the Agilent Infinity 1260 isocratic pump.
 - iii) Go to HPLC manager.
 - iv) Set the maximum flow gradient to 100 ml/min².
 - v) Set the flow rate to 5 ml/min.
 - vi) Run at least 25 ml buffer through degasser and pump (5 min).
 - vii) Close the purge valve on the Agilent pump.
- e. Equilibrate system in storage buffer:
 - i) Set equilibration flow rate (0.5 ml/min for most shared columns).
 - ii) Set flow to 0 ml/min.
 - iii) Set maximum flow gradient to 0.1ml/min².

17. Shutdown

- a. Clean up in and around the instrument.
- b. Close ASTRA.
- c. Logoff from PPMS

Protein Conjugate Analysis

- 1. Open Protein Conjugate Analysis Method.
 - a. Open a SEC-MALS data file in ASTRA
 - b. Right click on the file name of the open file in ASTRA.
 - c. Select Apply Method.
 - d. Go to System \rightarrow Method \rightarrow Light Scattering \rightarrow With QELS \rightarrow Protein. Conjugate.
 - e. Click Create.
- 2. Edit Peaks.
 - a. Open Peaks.
 - b. Expand Protein Conjugate Analysis.
 - c. Enter modifier dn/dc value for each peak (and UV if non-zero).
 - d. Click OK.
- 3. View data.
 - a. Open EASI Graph.
 - b. Display Protein Conjugate.
 - c. Open EASI Table to view summary data for all open files.
- 4. View Report.
 - a. Open Report (detailed).
 - b. Protein mass, modifier mass and conjugate (the total) mass are reported.

Book time and Report Problems through the PPMS system: https://ppms.us/hms-cmi

• rates are based on <u>booked and real time usage</u>

Contact cmi@hms.harvard.edu with questions.

last edited: 2021-04-21