

Quaternary Structure Modeling Through Chemical Cross-Linking Mass Spectrometry: Extending TX-MS Jupyter Reports

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Materials

Name	Company	Catalog Number	Comments
Two Protein DataBank files of the proteins of interest.	N/A	N/A	Example files available on txms.org and zenodo.org, DOI 10.5281/zenodo.3361621
An mzML data file acquired on a sample where the proteins of interest were crosslinked.	N/A	N/A	Example files available on txms.org or zenodo.org, DOI 10.5281/zenodo.3361621