

**Summary of integrative structure determination of Integrative structure of the yeast gammaTuSC-Spc110 tetramer complex (PDB ID: 9A17, PDB-Dev ID: PDBDEV\_00000079)**

<b>1. Model Composition</b>	
<u>Entry composition</u>	<ul style="list-style-type: none"> <li>- Spc97: Chain B (823 residues)</li> <li>- Spc110: Chain I (222 residues)</li> <li>- Spc97: Chain A (823 residues)</li> <li>- Spc98: Chain C (846 residues)</li> <li>- Spc110: Chain K (222 residues)</li> <li>- Tub4: Chain E (473 residues)</li> <li>- Spc110: Chain J (222 residues)</li> <li>- Tub4: Chain H (473 residues)</li> <li>- Spc110: Chain L (222 residues)</li> <li>- Tub4: Chain F (473 residues)</li> <li>- Tub4: Chain G (473 residues)</li> <li>- Spc98: Chain D (846 residues)</li> </ul>
<u>Datasets used for modeling</u>	<ul style="list-style-type: none"> <li>- Experimental model, PDB ID: 5FLZ</li> <li>- Crosslinking-MS data, Linker name and number of cross-links: EDC, 44 cross-links</li> <li>- Crosslinking-MS data, Linker name and number of cross-links: DSS, 42 cross-links</li> </ul>
<b>2. Representation</b>	
<u>Resolution</u>	Coarse-grained: 1, 2, 3, 4, 9, 33 residue(s) per bead
<u>Number of rigid bodies, flexible units</u>	0, 74
<u>Flexible units</u>	<ul style="list-style-type: none"> <li>- A: 1-54, 55-80, 81-89, 90-205, 206-242, 243-304, 305-319, 320-490, 491-553, 554-614, 615-622, 623-714, 715-753, 754-800, 801-823</li> <li>- B: 1-54, 55-80, 81-89, 90-205, 206-242, 243-304, 305-319, 320-490, 491-553, 554-614, 615-622, 623-714, 715-753, 754-800, 801-823</li> <li>- C: 1-179, 180-367, 368-378, 379-612, 613-627, 628-671, 672-718, 719-743, 744-755, 756-780, 781-797, 798-846</li> <li>- D: 1-179, 180-367, 368-378, 379-612, 613-627, 628-671, 672-718, 719-743, 744-755, 756-780, 781-797, 798-846</li> <li>- E: 1-445, 446-473</li> <li>- F: 1-445, 446-473</li> <li>- G: 1-445, 446-473</li> <li>- H: 1-445, 446-473</li> <li>- I: 1-165, 166-205, 206-222</li> <li>- J: 1-165, 166-205, 206-222</li> <li>- K: 1-165, 166-205, 206-222</li> <li>- L: 1-165, 166-205, 206-222</li> </ul>
<u>Structural coverage (rigid bodies)</u>	0%
<b>3. Restraints</b>	
<u>Physical principles</u>	Information about physical principles was not provided

<u>Experimental data</u>	- 1 unique CrossLinkRestraint: EDC, 44 cross-links - 1 unique CrossLinkRestraint: DSS, 42 cross-links
<b>4. Validation</b>	
<u>Number of ensembles</u>	1
<u>Number of models in ensembles</u>	2069
<u>Number of deposited models</u>	1
<u>Model precision (uncertainty of models)</u>	28.3, Å
<u>Data quality</u>	Data quality has not been assessed
<u>Model quality: assessment of excluded volume</u>	Satisfaction: 99.88-99.88%
<u>Fit to data used for modeling</u>	Fit of model to information used to compute it has not been determined
<u>Fit to data used for validation</u>	Fit of model to information not used to compute it has not been determined
<b>5. Methodology and Software</b>	
<u>1. Method</u>	Sampling
<u>Name</u>	Replica exchange monte carlo
<u>Number of computed models</u>	1000000
<u>Software</u>	- <a href="#">IMP PMI module</a> (version 2.14.0) - <a href="#">Integrative Modeling Platform (IMP)</a> (version 2.14.0)