Integrative structure and function of the yeast exocyst complex

Sai J. Ganesan1,2 | Michael J. Feyder3 | Ilan E. Chemmama1,2 | Fei Fang4 | Michael P. Rout5 | Brian T. Chait6 | Yi Shi4 | Mary Munson3 | Andrej Sali1,2

1Department of Bioengineering and Therapeutic Sciences, University of California, San Francisco, San Francisco, California
2Department of Pharmaceutical Chemistry, University of California, San Francisco, San Francisco, California
3Department of Biochemistry and Molecular Pharmacology, University of Massachusetts Medical School, Worcester, Massachusetts
4Department of Cell Biology, University of Pittsburgh, Pittsburgh, Pennsylvania
5Laboratory of Cellular and Structural Biology, The Rockefeller University, New York, New York
6Laboratory of Mass Spectrometry and Gaseous Ion Chemistry, The Rockefeller University, New York, New York

Correspondence
Andrej Sali, Department of Bioengineering and Therapeutic Sciences, University of California, San Francisco, Byers Hall, 1700 4th Street, Suite 503B, San Francisco, CA 94158.
Email: sali@salilab.org
Mary Munson, Department of Biochemistry and Molecular Pharmacology, University of Massachusetts Medical School, 364 Plantation Street, Worcester, MA 01605.
Email: mary.munson@umassmed.edu

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Abstract
Exocyst is an evolutionarily conserved hetero-octameric tethering complex that plays a variety of roles in membrane trafficking, including exocytosis, endocytosis, autophagy, cell polarization, cytokinesis, pathogen invasion, and metastasis. Exocyst serves as a platform for interactions between the Rab, Rho, and Ral small GTPases, SNARE proteins, and Sec1/Munc18 regulators that coordinate spatial and temporal fidelity of membrane fusion. However, its mechanism is poorly described at the molecular level. Here, we determine the molecular architecture of the yeast exocyst complex by an integrative approach, based on a 3D density map from negative-stain electron microscopy (EM) at ~16 Å resolution, 434 disuccinimidyl suberate and 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride cross-links from chemical-crosslinking mass spectrometry, and partial atomic models of the eight subunits. The integrative structure is validated by a previously determined cryo-EM structure, cross-links, and distances from in vivo fluorescence microscopy. Our subunit configuration is consistent with the cryo-EM structure, except for Sec5. While not observed in the cryo-EM map, the integrative model localizes the N-terminal half of Sec3 near the Sec6 subunit. Limited proteolysis experiments suggest that the conformation of Exo70 is dynamic, which may have functional implications for SNARE and membrane interactions. This study illustrates how integrative modeling based on varied low-resolution

Abbreviations: CATCHR, complexes associated with tethering containing helical rods; CXMS, chemical-crosslinking mass spectrometry; DSS, disuccinimidyl suberate; EDC, 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride; EM, electron microscopy; GMM, Gaussian mixture model; IMP, integrative modeling platform; LC/MS, liquid chromatography and mass spectrometry; MTC, multisubunit tethering complex; PH domain, Pleckstrin homology; PMI, python modeling interface; RMSD, root-mean-square deviation; RMSF, root-mean-square fluctuation; SNARE, soluble NSF attachment protein receptor.

Sai J. Ganesan and Michael J. Feyder authors contributed equally to this work.

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1 | INTRODUCTION

Vesicle trafficking and fusion in eukaryotic cells are accurately regulated to ensure that protein and lipid cargos are delivered to the correct destinations. At the plasma membrane, regulation is carried out by a multisubunit tethering complex (MTC) called exocyst, which is required for quality control of fusion of secretory vesicles at appropriate exocytosis sites.\textsuperscript{1–7} Exocyst and other MTCs function to recognize the correct vesicles and regulate SNARE complex assembly for membrane fusion. These functions are required for many fundamental cellular processes, such as secretion, cell growth, cell–cell communication, and cell division.\textsuperscript{1,6,8}

The exocyst complex is a hetero-octameric MTC composed of eight subunits, including Sec3, Sec5, Sec6, Sec8, Sec10, Sec15, Exo70, and Exo84, ranging in size from ~650 to 1,300 amino acid residues (Figure 1a). Most of these subunits were discovered in a yeast secretory pathway screen.\textsuperscript{2,9,10} The phenotype of the yeast mutants at the nonpermissive temperature is the accumulation of secretory vesicles that do not fuse, resulting in failed cargo delivery. High-resolution crystal structures of Exo70, Exo84, Sec3, Sec15, Sec10, and Sec6 domains indicate that the subunits are mainly elongated, consisting of helical bundles stacked on top of each other. Subunits without available crystal structures were predicted to contain similar helical bundles.\textsuperscript{11–15} These helical bundles indicate that the exocyst belongs to the complexes associated with tethering containing helical rods (CATCHRs) family of MTCs, along with Dsl1, Golgi-associated retrograde protein, endosome-associated retrograde protein, and conserved oligomeric Golgi complexes.\textsuperscript{17–20} In addition to the helical bundles, the exocyst subunits Sec3 and Exo84 (and Sec5 in mammals) have domains that bind small GTPases and/or PI(4,5)P\textsubscript{2}.\textsuperscript{21–23}

We previously obtained negative-stain EM 2D class averages of the yeast exocyst and used an auxin-inducible degron system to selectively degrade exocyst subunits,\textsuperscript{24} which showed that the yeast exocyst is stably assembled

\textbf{FIGURE 1}  Diagram of the exocyst subunit domain arrangements and high-resolution structures. (a) \textit{Saccharomyces cerevisiae} exocyst complex consists of eight subunits ranging from ~650 to 1,300 residues. Black bars show crystallized regions of each subunit, which are shown on the right. Structures were determined of exocyst domains from yeast,\textsuperscript{11,12,22,23} rat,\textsuperscript{21} zebrafish,\textsuperscript{14} and \textit{Drosophila}.\textsuperscript{13} (b) Comparison between the 16 Å negative stain electron microscopy (EM) density and the cryo-EM model of exocyst, showing similar architectures and overall structures.
from two 4-subunit modules or half-complexes. Full molecular architecture of the yeast exocyst was revealed by the cryo-EM structure determined at 4.4 Å resolution. Each 4-subunit module is stabilized by a four-helix bundle, called CorEx. Each CorEx bundle consists of one helix from the N-terminus of each of the subunits, except for Sec3 that contributes a helix from the middle of the sequence. Recently, we also determined a 3D negative-stain EM map of the entire octameric exocyst (Figure 1b). As seen by both negative-stain EM and cryo-EM, the structure appears compact and “closed.” This form is in stark contrast with structures of other CATCHR MTCs, whose subunits protrude from the complex in extended conformations, suggesting direct roles in tethering vesicles to the appropriate target membranes. Many protein–protein interactions between exocyst subunits and its partners have been determined over the years, but most have not been characterized in relation to the fully assembled complex. Our recent studies indicate that the closed conformation of exocyst needs to be activated by one or more of its partners to open into a functional conformation.

Here, we determined the molecular architecture of the yeast exocyst complex using an integrative approach that depends on multiple types of experimental data to maximize the accuracy, precision, completeness, and effi-

**FIGURE 2** Chemical cross-linking and mass spectrometry reveal connectivity within the exocyst. (a) Both EDC (left) and DSS (right) crosslinks are shown with intermolecular (black) and intramolecular (brown) crosslinks indicated. The map was generated using the Circos-CX software (http://cx-circos.net/). (b) The EDC and DSS crosslinks are mapped onto the previously published cryo-EM structure of exocyst, showing those that satisfy the distance restraint (<35 Å for DSS and <25 Å for EDC; blue) or violate it (>35 Å for DSS and >25 Å for EDC; red)
ciency of structure characterization. By bringing together varied low-resolution structural information, we aimed to illustrate the power of the integrative approach in the absence of high-resolution data. We also aimed to annotate the exocyst function by comparing our integrative structure with a previously determined cryo-EM structure in light of new biochemical data.

2 | RESULTS

2.1 | Chemical cross-linking of exocyst

To obtain structural information about the exocyst, purified complexes were subjected to extensive chemical cross-linking with both EDC (0 Å linker length) and DSS (11.4 Å linker length), followed by mass spectrometry (CXMS) identification of the cross-linked peptides (Section 4). This approach identified 434 unique intra- and intersubunit cross-linked pairs of residues (Figure 2 and Table S1).

2.2 | Integrative structure determination of the exocyst complex

To model the octameric yeast exocyst structure, we followed the previously described four-stage workflow (Figure 3, Supplementary Methods, Table S2). Input information included the 16 Å negative-stain EM density map,26 the 434 chemical cross-links (Section 2.1), as well as the X-ray structures,11–14,22 and comparative models for 51% of the residues in the 8 exocyst subunits. Stereochemically feasible exocyst structures were sampled to compute an ensemble containing 9,741 models that satisfy the experimental restraints sufficiently well (Figure S1, Section 4, and Supplementary Methods). The ensemble corresponds to a single cluster of models with the root-mean-square fluctuation (RMSF) of 38 Å (defining the precision of the integrative structure; Figure 4a,b). The centroid structure (the structure with the minimal sum of root-mean-square deviations (RMSD) from all the other structures in the ensemble) is used as the representative model.
2.3 Validation of the integrative structure

We begin by assessing how well the model ensemble satisfies the input cross-links and negative-stain EM density map. The ensemble satisfies the data well. First, 98% (255 DSS and 170 EDC) of cross-links are satisfied by at least one model in the cluster (Figure 4a,d). Most of the violations are small (<5%), and can be rationalized by experimental uncertainty, local structural fluctuations,
coarse-grained representation of the model, and/or finite structural sampling. Second, the shape of the negative-stain EM density map also overlaps well with that of the ensemble; the correlation coefficient between the map and the ensemble is 0.83 (Figure 4c).

Next, we assessed the model by cross-validation against the previously published cryo-EM density map,25 cross-links,25 and distances from in vivo fluorescence microscopy in live yeast cells.33

First, as for the negative-stain EM map, the shape of the cryo-EM density map also overlaps well with that of the ensemble; the correlation coefficient between the cryo-EM map and the ensemble is 0.78 (Figure 4f). However, the Sec15 and Sec6 subunits do not fit the map as well as other subunits, resulting in an unoccupied volume in the EM map (shaded gray region in Figure 4c,f). Both these subunits are also poorly resolved in the negative-stain EM 2D averages24 and the cryo-EM structure.25 This could be due to three reasons, (a) sample heterogeneity or the presence of multiple states in the sample, (b) dynamics or fluctuations in Sec15 and Sec6, or (c) lack of sufficient data.

Second, the integrative model satisfied 284 (211 DSS and 73 EDC, or 98%) out of the 290 previously published inter- and intrasubunit DSS and EDC cross-links25 (Figure 4e), even though these cross-links were not used to compute the model; four of the six violated cross-links are violated by less than 10 Å. Out of these 290 cross-links, 139 overlap between the datasets (119 DSS and 20 EDC cross-links).

Finally, we also validated our model based on 80 inter-subunit distance measurements obtained by in vivo fluorescence imaging experiments.33 The authors converted these distance measurements into a unique threshold for each of the distances between the two labeled subunits33 (Figure 4g). All 80 distance thresholds were satisfied by at least one model in the ensemble (Figure 4g). Each model in the ensemble satisfies at least 70 of the 80 distance thresholds (Figure 4g).

In conclusion, our confidence in the integrative model is increased by its consistency with the data used to compute it (our negative-stain EM map and cross-links) and the data not used to compute it (previously published cryo-EM map, cross-links, and in vivo fluorescence images).

### 2.4 Comparison of integrative and cryo-EM structures

We now compare the representative integrative structure (Supplementary Methods) and the cryo-EM structure (PDB: 5YFP).25 Upon a global least-squares superposition, the average Cα RMSD value for the regions represented at atomic resolution in both structures is 57 Å (Figure 4f). Using these global superpositions, we also quantified the similarity in individual subunit positions, orientations, and conformations. The subunit Cα RMSD values range from 10 to 32 Å. The subunit with one of the highest atomic coverages (74% of the atoms are represented explicitly), Sec10, has the highest degree of similarity in position, orientation, and conformation in the two structures (Cα RMSD of 10 Å). In contrast, the position, orientation, and conformation of the subunit with the lowest atomic coverage (33%), Sec15, is most different with an RMSD of 30 Å (Table S2).

We now interpret the observed differences between the two structures. In general, differences between two models can arise from a combination of two factors. First, a difference can appear because of the uncertainty of both

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**FIGURE 4** Validation of the integrative exocyst structure. (a) Localization probability densities of the exocyst subunits from the solution cluster. Left and right images represent two views at 180° rotation. (b) Atomic representation of the centroid structure of the solution cluster. Left and right images represent two distinct views at 180° rotation. (c) Fit of the localization probability density of the exocyst subunits from the solution cluster to input EM map shown in mesh representation. (d) Euclidean Cα–Cα distance distributions of our measured cross-links in the ensemble of solutions for the cluster (Table S1). The y-axis details the normalized number of cross-links that were mapped onto the model, x-axis details the distances. Dashed blue and green lines denotes the expected maximum reach of a DSS and EDC cross-link respectively. Satisfied DSS and EDC cross-links, with Cα–Cα distances that fall within the distance threshold of 35 and 25 Å in at least one exocyst structure in the solution ensemble, are shown in blue and green, respectively. (e) Euclidean Cα–Cα distance distributions of all previously reported cross-links25 in the solution ensemble. Satisfied DSS and EDC cross-links, with Cα–Cα distances that fall within the distance threshold of 35 and 25 Å in at least one exocyst structure in the solution ensemble, are shown in blue and green, respectively. (f) RMSD distance distribution between regions represented at atomic resolution (51% of the structure) in the solution ensemble and PDB ID 5YFP, after a global least-squares superposition (Section 2). The structure on the inset shows the fit of the localization probability densities of the exocyst subunits in the solution cluster to the published cryo-EM map (EMDB ID 6827). (g) Euclidean Cα–Cα distance measurements of previously published in vivo distance measurements33 in the solution ensemble. Box plot for each measurement across the solution ensemble is compared with the reported mean distance (red circles). (h–o) Comparison of subunits in the integrative structure with the cryo-EM structure (PDB: 5YFP) upon a global least-squares superposition. Figures h–o represent RMSD distributions of subunits Sec3, Sec5, Sec6, Sec8, Sec10, Sec15, Exo70, and Exo84, respectively.
models. Second, a difference can also appear because of the actual variation between the two structures, resulting from the different samples and/or conditions used to obtain the data for the two structure determinations. We attempt to deconvolute the two sources of difference as follows. The uncertainty of the integrative model (model precision is 38 Å) is most likely substantially higher than that of the cryo-EM structure (based on a cryo-EM map at resolution of 4.4 Å). Therefore, to visualize the significance of the structural differences between the integrative and cryo-EM structures, we plot the distribution of the RMSD values between the cryo-EM structure and each of the ensemble models (Figure 4h–o). The cryo-EM structure is within the integrative ensemble density except for Sec5, indicating there is no significant difference between the two structures other than for Sec5. In other words, at the precision of the integrative model, the actual structural difference between the two structures is limited to the position, orientation, and conformation of Sec5. Most of the difference is due to changes in position and orientation, as the conformations of the Sec5 models are similar (Cα RMSD of 5 Å).

### 2.5 Comparison of cross-links and negative-stain EM map with the cryo-EM structure

We now assess how well the cryo-EM structure satisfies our cross-links and negative-stain EM density map. The cryo-EM structure satisfies 78% of the mapped EDC cross-links (83 of 106) and 95% of the mapped DSS cross-links (189 of 199) (Figure 2b). While these cross-link satisfactions are less than that for the integrative structure (96 and 99%, respectively), our cross-links were not used in computing the cryo-EM structure and the degree of satisfaction is still high. The cryo-EM structure also fits the negative-stain EM density map well, with a cross correlation coefficient of 0.86, which is comparable to that for the integrative model (0.83). In conclusion, our cross-links and negative-stain EM density map are consistent with the cryo-EM structure.

### 2.6 Structural interpretations for biochemical data

While the exocyst complex has been studied for many decades, its structure and function are only beginning to be understood. There is considerable published and unpublished biochemical data on the yeast complex as well as exocyst complexes from other species. Here, we use the structures to help interpret several types of biochemical data.

#### 2.6.1 Protein–protein interactions

Exocyst subunits interact with each other as well as with a number of different protein and lipid factors.\(^1,3,6,7,12,15,34-37\) Many of these interactions are critical for the stability of exocyst, the recognition of secretory vesicles (via Rab GTPases and the type V myosin, Myo2) at specific sites on the plasma membrane (Rho GTPases, PI(4,5)P2, and Ral GTPase in mammals), and the regulation of membrane fusion through interactions with the SNARE proteins and Sec1/Munc18. The regions of the exocyst that interact with many of these partners are poorly resolved in our integrative model, the negative-stain EM map,\(^24\) and the cryo-EM structure.\(^25\) Examples include Sec6 interacting with SNARE and Sec1 binding partners,\(^38-41\) the Sec3 N-terminal region interacting with Rho GTPases, Pip2 and the Sso1/2 SNARE,\(^22,23,42\) and Sec15 interacting with the Rab GTPase Sec4 and Myo2.\(^43,44\) This observation suggests that these regions are flexible in the absence of their binding partners.

#### 2.6.2 Limited proteolysis

To reveal exposed surfaces of exocyst, the assembled complex was incubated with trypsin in solution overnight, and the digested peptides were analyzed by liquid chromatography and mass spectrometry (LC/MS). Generally, the digested regions mapped to flexible loops and short helices in the cryo-EM structure, consistent with the solvent accessibility of those regions (Figure 5a). Exo70 was the second-most cleaved subunit after Sec3 (discussed in the next section). Not surprisingly, an exposed helix that links two of the Exo70 helical bundles was digested. However, several buried regions of Exo70, including its CorEx helix, were also cleaved. This result suggests that Exo70 is sufficiently dynamic to make its CorEx bundle residues accessible. We showed that gain-of-function mutants of Exo70 led to a dynamic conformational change in which Exo70 shifted away from the rest of the complex,\(^26,45\) potentially exposing regions that were cleaved in our limited proteolysis experiment. These dynamics of Exo70 were not observed in the cryo-EM structure. However, the cryo-EM data were obtained using glutaraldehyde-crosslinked complexes,\(^25\) which likely stabilized the complex on grids at the cost of limiting configurational and conformational variability. In contrast, the integrative model reflects a substantial configurational and conformational uncertainty of Exo70 (Figure 4n), although it is not clear to what degree it originates from the lack of data or actual flexibility/heterogeneity (above). Our limited proteolysis data support a
model in which exocyst is autoinhibited and undergoes structural rearrangements as part of its tethering and SNARE regulation function.\textsuperscript{7,26,46}

### 2.6.3 Deletions and mutations

Deletion of most exocyst subunits are lethal in eukaryotes.\textsuperscript{6} In yeast, the secretory (sec) mutants\textsuperscript{9} as well as auxin-induced degradation studies provided insights into the function of the complex.\textsuperscript{2,24,47–49} The yeast sec6-4 mutant (L633P) had previously been sequenced.\textsuperscript{11,50} We sequenced several others (sec3-2 is Q772*; sec5-24 is W300*; sec8-9 is W758*; sec10-2 is E439K; sec15-1 is G836*). Surprisingly, many mutations result in premature stop codons that truncate exocyst subunits (Figure 5b). These truncated proteins appear to function reasonably well at permissive temperatures; for example, while the yeast cells grow at 20°C, the deletions are destabilizing at higher temperatures and the cells die.\textsuperscript{2} In each case, the truncation occurs in the sequence after the CorEx helical bundles, indicating that an essential function of most of the subunits is to stabilize the complex through their CorEx bundles.

### 2.6.4 Phosphorylation

To investigate possible structural and functional consequences of phosphorylation, we mapped phosphorylation sites (Table S3) from high throughput studies.\textsuperscript{51–56} and

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**FIGURE 5** Limited trypsin proteolysis, mutations, and phosphorylation of the assembled exocyst. (a) Limited proteolysis using trypsin identified flexible and exposed regions within the intact exocyst complex. Regions colored in red indicate peptides that were cleaved after an overnight trypsin digest in solution. Arrows point to regions of Exo70 that are buried in the cryo-EM structure. (b) Regions of exocyst that are truncated or mutated in the original sec mutant screen;\textsuperscript{9} sec6-4 (L633P); sec3-2 is Q772*; sec5-24 is W300*; sec8-9 is W758*; sec10-2 is E439K; sec15-1 is G836*. Truncated regions are shown as lines. (c) Phospho-sites mapped to the cryo-EM structure (Table S3)
our own MS studies, onto the current cryo-EM model (Figure 5c). Phosphorylation sites often occur in intrinsically disordered protein regions, and may control disorder–order transitions. Of the total of 49 phosphorylation sites, 40 are in regions absent from the cryo-EM structure. The remaining nine phosphorylated sites all map on interhelical loops. All 24 of the phosphorylated sites on Sec3 are in the Sec3N region, mostly clustered around the PH domain (Figure 6b). Seventeen sites are in Exo84. In particular, five of these sites lie near interaction surfaces with other subunits (including Sec5, Sec8, and Sec15). These phospho-sites have previously been implicated in the assembly of the complex, as their mutation leads to defects in secretion and cell growth. Other phospho-sites on Sec5, Sec8, and Sec10 are in the vicinity of regions critical for conformational changes that activate Sec6 for SNARE binding or near SNARE binding sites. A phosphorylated serine in the C-terminus of Sec15 is located in a region near putative Rab and Myo2 binding sites, although this region of the cryo-EM

**FIGURE 6** The N-terminal domain of Sec3 (Sec3N) is disordered and localizes near the C-terminal region of Sec6. (a) Sec3N (1–611 residues) is predicted to be disordered using PrDOS, with the exception of the PH domain (71–241). The y-axis gives the probability of each residue to be disordered, and the threshold is the false positive rate. The graph below shows the results from the limited trypsin digestion; the digestion showed that most of Sec3N is susceptible to proteolysis (red), consistent with the predicted disorder in most of this region. (b) CXMS revealed a large cluster of contacts between Sec3N and the region near the C-terminal end of Sec6 (purple), suggesting that Sec3N can occupy this area in various positions, presumably transiently. Intramolecular Sec3N crosslinks and phosphorylation sites in this domain are shown in black and green, respectively (Table S3)
structure is dynamic and poorly resolved. These phosphorylated regions may be critical for regulation of exocyst assembly, vesicle binding, and/or SNARE function.

2.7 | The Sec3 N-terminal region is flexible and proximal to the SNARE-binding region of Sec6

We now focus on Sec3—the entire Sec3 protein is included in the integrative model, while the N-terminal half of Sec3 is missing in the cryo-EM structure. Sec3 is unique among the budding yeast exocyst subunits, as it appears to be substantially longer (1,336 residues) than its homologs in other species, ranging from fission yeast to humans (~800 residues). Most of this difference in length lies in the N-terminal region (1–611 residues), which is considerably longer in *Saccharomyces cerevisiae*.60–63 The length difference may explain the structural differences among the exocyst complexes in different organisms; biochemical studies of the S. cerevisiae exocyst indicate that it functions as a single octameric complex,24 while smaller subcomplexes appear to exist in other species.5,64–70 For example, Sec3 appears to dissociate from the mammalian complex after tethering.65

In contrast, the longer budding yeast Sec3 protein may extend the physical reach of the complex to promote interactions with other factors during tethering, thus avoiding post-tethering dissociation by maintaining the complex as a stable octamer and bypassing the need for subcomplexes.

Residues 71–241 of Sec3 fold into a Pleckstrin homology (PH) domain (Figure 1a), which is a key functional region in the N-terminal half of Sec3 (Sec3N).22,23,42 Association of the polybasic region in the PH domain with PI(4,5)P2 was suggested to play an important role in localization of the complex and in tethering.23 Moreover, this domain binds Sso2 and catalyzes assembly of SNARE complexes, likely by interacting with the inhibitory N-terminal H\(_{abc}\) domain of Sso2 and relieving auto-inhibition.42,71 Interestingly, deletion of Sec3N is not lethal unless combined with phosphoinositide-binding defective mutants of Exo70, suggesting that functional redundancy exists between the two subunits.48 This functional redundancy also extends to binding of Rho GTPases, including Cdc42, by both Sec3 and Exo70,72 suggesting that binding of exocyst to both PI(4,5)P2 and Rho GTPases are essential for cell growth and secretion.

Apart from the PH domain, the rest of Sec3N is predicted by the PrDOS and PONDR algorithms73,74 to be disordered (Figure 6a). In contrast, the disordered regions in the C-terminal half of Sec3 were predicted to be only in short loop regions. The N-terminal flexible regions were confirmed by limited proteolysis experiments of exocyst in solution (Figure 6a). In contrast, in the C-terminal half of Sec3, only a single loop was cleaved (Figure 6a). The predicted flexibility of the N-terminal half of Sec3 is consistent with the lack of density for Sec3N in the cryo-EM model. Heterogeneity in Sec3N phosphorylation could also lead to structural heterogeneity (Table S3 and Figure 6b). Differential phosphorylation of Sec3N is likely to provide regulation; Pck1 phosphorylation of Sec3 upon wounding of the plasma membrane was shown to lead to Sec3 degradation, exocyst relocalization, and inhibition of exocytosis.75

Although the position of Sec3N was unresolved in the cryo-EM model, our CXMS experiments identified the region of exocyst that Sec3N interacts with, albeit transiently (Figure 6b). It had been suggested that Sec3N is located near Sec15, based on a single crosslink between Sec3 and Sec15.25 In contrast, our more extensive EDC and DSS CXMS revealed that Sec3N forms multiple crosslinks in a region at the other end of the complex, near Sec6 and its SNARE interacting domains (Figure 6b). This crosslinking data informed the integrative model, which localizes the N-terminal domain of Sec3 near Sec6 (Figure 4a), in agreement with two independent considerations as follows. First, both Exo70 and Sec3 interact with PI(4,5)P2. Therefore, we expect them to be positioned on the same face of exocyst for optimal membrane binding (Figure 4a). Second, as Sec3 and Sec6 are the SNARE-binding exocyst subunits, they are likely to be close to each other to coordinate SNARE complex assembly.

3 | DISCUSSION

3.1 | Utility of integrative structure modeling

No experimental method for structure determination is universally applicable. For example, crystals suitable for X-ray crystallography cannot always be produced, especially for insufficiently homogenous assemblies of multiple components. While cryo-EM can be used to study large assemblies, it often produces structural models at worse than atomic resolution due to sample heterogeneity and/or a limited number of single particle images. Finally, molecular biology, biochemistry, and proteomics techniques, such as yeast two-hybrid, affinity purification, and mass spectrometry, yield information about the interactions between proteins, but not the positions of these proteins within the assembly or the atomic structures of the proteins themselves.

One approach to overcome the limitations of individual methods is integrative structure modeling.27–32,76–79 Integrative modeling aims to simultaneously use all available information from any method, including from varied experiments (e.g., EM and chemical cross-linking),
physical theories (e.g., molecular mechanics force field), statistical analyses (e.g., statistical potentials), and prior models (e.g., atomic structures of complex subunits). Therefore, integrative modeling maximizes the accuracy, precision, completeness, and efficiency of structure determination. Numerous structures refractive to traditional methods have already been solved using this approach.28

This study illustrates how integrative modeling based on low-resolution structural data can be useful even in the absence of high-resolution data. In particular, we determined the molecular architecture of the yeast exocyst complex at 38 Å precision (Supplementary Methods) by an integrative approach as implemented in our open source IMP package.32 Modeling relied on a negative-stain EM density map at 16 Å resolution, 434 intra- and intermolecular cross-links, partial atomic models of the 8 exocyst subunits, sequence connectivity, and excluded volume. The model was largely consistent with the previously published cryo-EM structure,25 chemical cross-links, and intersubunit distances from in vivo fluorescence measurements in yeast. Importantly, the integrative model reproduces locations of critical functional regions, such as the C-terminal regions of Sec15 (for vesicle binding) and Sec6 (for SNARE interactions) at opposite ends of the complex, as found in the cryo-EM structure.

3.2 Biochemical experiments indicate regulatory regions in exocyst

We examined the yeast exocyst structure in relation to previously published as well as new biochemical and genetic data regarding flexible/disordered regions, mutations, and phosphorylation sites (Figures 5 and 6). The data suggests regions that are likely involved in (a) the regulation of the conformation and function of exocyst, (b) the assembly and stability of the complex, and (c) the function of exocyst controlled by phosphorylation. Our cross-linking data also places the N-terminal region of Sec3, which interacts with PI(4,5)P2, Rho GTPases, and the plasma membrane SNARE Sso1/2, nearby the region of Sec6 that has been implicated in SNARE and Sec1 binding. This information is critical for beginning to tease apart exocyst’s mechanism of action, although further structural and biochemical information will be necessary for a molecular understanding of this fundamental process.

In conclusion, our integrative exocyst model is similar to the previous structure based on a cryo-EM density map at 4.4 Å resolution. Therefore, it allows us to reach many of the same conclusions about the function of exocyst. Because chemical cross-links and negative-stain EM density map are often much easier to determine than a higher resolution cryo-EM density map, this study illustrates the efficacy of integrative structure determination. Thus, the integrative approach can facilitate rationalizing existing data and suggesting hypotheses for future experiments, even in the absence of higher resolution cryo-EM data.

4 MATERIALS AND METHODS

4.1 Cross-linking and mass spectrometry analyses

4.1.1 EDC and DSS cross-linking of the exocyst complex

Exocyst was isolated from cells based on the previously described protocol.24 Briefly, 8 L of either GFP-Exo84/Sec15-PrA (for EDC crosslinking) or Sec8-GFP (for DSS crosslinking) were grown to an approximate OD600 of 1.5, and lysed using a ball mill grinder.80 The frozen powder was resuspended in 40 ml of 40 mM Tris pH 8.0 and 200 mM sodium citrate lysis buffer with a Roche protease inhibitor tablet and spun at 13,000 rpm to remove cell debris. The soluble lysate was then incubated with either IgG-conjugated Dynabeads (EDC) or anti-GFP nanobody Dynabeads (DSS) for 1 hr at 4°C. The beads were washed with 600 μl of lysis buffer three times, then washed with 600 μl of 20 mM MES pH 6.5, 250 mM NaCl, 2 mM MgCl2, 0.1% CHAPs, and 1 mM DTT, 3x. The slurry for EDC was resuspended in 100 μl of 20 mM MES pH 6.5, 250 mM NaCl, 2 mM MgCl2, 0.1% CHAPs, and 1 mM DTT, plus PreScission protease, for 1.5 hr at 4°C. The eluted exocyst was then treated with either 2.5 mM EDC and 5 mM NHS for 30 min at 25°C. The reaction was quenched by adding Tris pH 8.0 and β-mercaptoethanol to a final concentration of 50 and 20 mM, respectively. For DSS, either 5 or 10 g of frozen yeast powder was used, the cross-linking was carried out on beads, and the peptides eluted by trypsin digest.81

4.1.2 Proteolytic digestion of EDC crosslinked exocyst complex

After EDC cross-linking, the complex was reduced by 10 mM tris-(2-carboxyethyl)-phosphine (Invitrogen, MA) at 80°C for 15–20 min, cooled to room temperature and alkylated by 50 mM iodoacetamide for 20 min in the dark to block the formation of disulfide bonds. After reduction and alkylation, the cross-linked complexes were separated by 3–12% SDS-PAGE (NuPAGE Bis-Tris, Invitrogen) to reduce the complexity of the sample. For in-gel
digestion, the high-molecular-weight-region gel bands (>220 kDa, estimated by the high-molecular-weight protein markers, Invitrogen) corresponding to the cross-linked exocyst proteins were sliced and proteolysed by trypsin as previously described. In brief, gel plugs were crushed into small pieces, 0.5 µg of sequencing-grade trypsin (Promega, Madison, WI) per ~10 µg protein was added with subsequent 6–8 hr incubation. Peptides were extracted by formic acid and acetonitrile, desalted on C18 cartridges (Sep-Pak, Waters, MA) and snap-frozen before fractionation.

4.1.3 Fractionation of cross-linked peptides by high pH chromatography

For the high pH chromatography fractionation, the tryptic peptides were separated in a home-made reverse-phase C18 column in a pipet tip. Peptides were eluted into seven fractions using a stepwise gradient of increasing acetonitrile (6, 9, 12, 15, 18, 22, and 35%) at pH 10 adjusted with ammonium hydroxide, and the 6 and 35% fractions were combined to one fraction. All of the six fractions were lyophilized in a vacuum concentrator (Thermo Fisher Scientific, MA), and analyzed by LC/MS.

4.1.4 Mass spectrometric analysis of cross-linked peptides

For cross-link identification, the fractionated peptides were dissolved in the sample loading buffer (2% ACN and 0.2% formic acid) and loaded onto a Picochip® column with an integrated electrospray ionization emitter tip (360 outer diameter, 75 inner diameter with 15 µm tip, New Objective, Woburn, MA). The column was packed with 25 cm of reverse-phase C18 material (3 µm porous silica, 120-Å pore size, Dr. Maisch GmbH, Entringen, Germany). Mobile phase A consisted of 2% acetonitrile with 0.1% acetic acid, and mobile phase B consisted of 80% acetonitrile with 0.1% acetic acid. The peptides were eluted in a 90-min LC gradient (5% B to 12% B, 0–10 min; 12% B to 32% B, 10–75 min; 32% B to 100% B, 75–80 min, and followed by 100% B until 90 min) using an easy nLC-1200 system (Thermo Fisher) and analyzed with an Q Exactive HFX or QE Plus mass spectrometer (Thermo Fisher). The flow rate was 350 nl/min. The capillary temperature was 250°C, and Funnel RF level was set at 55%. The instrument was operated in the data-dependent mode, where the top 10 or top 6 most abundant ions were fragmented by HCD (HCD energy 30) and analyzed in the Orbitrap mass analyzer. The target resolution for MS1 was 60,000, and for MS2 it was 7,500. Peptide ions with a charge state of >3 were selected for fragmentation. An auto dynamic exclusion of was used. Other instrumental parameters include the following: Isolation window of 1.8 Th, and minimal threshold of 50,000 to trigger an MS/MS event. The AGC targets were set as 1 × 10⁶ for full mass and 2 × 10⁶ for MS2. The maximal ion injection times for MS1 and MS2 were set as 100 and 200 ms, respectively.

4.1.5 Identification of cross-linked peptides using pLink

The raw data were transformed to Mascot generic format by pXtract 2.0 and searched by pLink3 using a target decoy search strategy with a concatenated FASTA protein sequence database containing the eight subunits of the exocyst complex. We also included the BSA sequence for the target-decoy database search to quickly prefilter the false positive identifications containing the BSA sequence. Other search parameters included cysteine carboxymethylation as a fixed modification, methionine oxidation, and protein N-terminal methionine cleavage as a variable modification. A maximum of two trypsin missed-cleavage sites were allowed. The data were filtered using a mass accuracy of MS1 ≤ 10 ppm (parts per million) and MS2 ≤ 20 ppm as specified in the software. We then manually applied additional filters to remove potential false positive identifications from our dataset as previously described. The cross-linking data were analyzed and plotted by an online software tool, CX-Circos (http://cx-circos.net).

4.2 Integrative structure modeling of the exocyst complex

The architecture of the yeast exocyst complex was determined using the integrative structure modeling approach described previously. All available structural information on the exocyst complex was used for computational analyses (Table S2, Figure 3, and Section 4). Representations of individual subunits relied on (a) atomic structures of three yeast exocyst subunit domains, determined by X-ray crystallography (PDB codes 2FJI, 2D2S, and 2BIE) and (b) eight comparative models built with MODELLER 9.13 based on known template structures detected by HHPred (Table S2 and Section 4). The atomic regions of the exocyst subunits were represented by 12 rigid bodies, corresponding to different domains of the eight constituent proteins (Table S2). Regions with unknown structures were modeled as flexible strings of beads. The EM density map was
approximated by a Gaussian mixture model (GMM) containing 200 components and every 50–100 residues of each subunit was approximated by a GMM component. Finally, the proximity between specific residue pairs was determined by DSS and EDC XL-MS experiments, which identified a total of 256 DSS and 178 EDC (Table S2 and Figure 4) inter- and intrasubunit interactions. The maximum Ca–Ca distance between any two residues cross-linked by DSS was estimated to be ~35 Å, and by EDC was estimated to be ~25 Å, based on spacer length, flexibility of side chains, and backbone dynamics.

Next, 2,000,000 exocyst models were computed by optimizing spatial proximities using a Monte Carlo method, as informed by cross-linking data, EM map, excluded volume, and sequence connectivity from 200 random initial models. The sampling yielded an ensemble of 9,741 models that sufficiently satisfied the input restraints; this entire ensemble is the model of the exocyst, with its uncertainty reflecting both the lack of input data and the actual heterogeneity of the samples used to collect the data. The clustering of these models revealed a single cluster containing the majority (99% or 9,669 models) of the models (Figure S1 and Supplementary Methods). The precision of the cluster defines the overall precision of the structural model (Figure S1). It is quantified by the average root-mean-RMSF with respect to the centroid model and is 38 Å for the modeled exocyst structure.

In general, an ensemble can be visualized as a localization probability density map. The map specifies the probability of any volume element being occupied by a given bead in superposed models. The probability localization density for the subunits of exocyst is sufficiently precise to define their position, but not their orientation (Figure 4a).

### 4.3 In solution limited trypsin proteolysis of native complexes

#### 4.3.1 In-solution digestion

Ten μg of eluted exocyst (purified as above) was added to 20 μl of 0.1% ProteaseMAX Surfactant (Promega) in 50 mM ammonium bicarbonate, an additional aliquot of 20 μl of 50 mM ammonium bicarbonate was then added. For reduction, a 2 μl aliquot of 45 mM DTT was added and the samples were incubated at 50°C for 30 min. For alkylation of cysteines, a 2 μl aliquot of 100 mM iodoacetamide was added and the samples were incubated at room temperature for 30 min. Tryptic digestion was initiated with a 1 μl aliquot of trypsin (0.2 μg/μl Sigma Proteomics grade). Samples were incubated at 37°C for 18 hr. Tryptic digests were acidified with 4 μl of 5% trifluoroacetic acid (TFA) and lyophilized in Speed Vac.

#### 4.3.2 LC–MS/MS analysis

A 3 μl injection was loaded by a Thermo Easy-nLC II in 5% acetonitrile (0.1% formic acid) at 4.0 μl/min for 4.0 min onto a 100 μm I.D. fused-silica precolumn packed with 2 cm of 5 μm (200 Å) Magic C18AQ (Bruker-Michrom). Peptides were eluted at 300 nl/min from a 75 μm I.D. gravity-pulled analytical column packed with 25 cm of 3 μm (100 Å) Magic C18AQ particles using a linear gradient from 5 to 35% of mobile phase B (acetonitrile + 0.1% formic acid) in mobile phase A (water + 0.1% formic acid) over 60 min. Ions were introduced by positive electrospray ionization via liquid junction at 1.4 kV into a Thermo Scientific Velos Pro mass spectrometer. Mass spectra were acquired over m/z 350–2,000 at 60,000 resolution (m/z 200) with an AGC target of 1e5, and data-dependent acquisition selected the top 10 most abundant precursor ions for tandem mass spectrometry by CID fragmentation using an isolation width of 2 Da, max fill time of 110 ms, and AGC target of 1e5. Peptides were fragmented by a normalized collisional energy of 35, and fragment spectra acquired at a resolution of 17,500 (m/z 200).

#### 4.3.3 Data analysis

Data was analyzed by Mascot Server (version 2.4, Matrix Science) against the S. cerevisiae (Swissprot) FASTA file (downloaded 06/2013). Search parameters included Trypsin/P specificity, up to two missed cleavages, a fixed modification of carbamidomethyl cysteine, and variable modifications of oxidized methionine, pyroglutamic acid for Q, and N-terminal acetylation. Assignments were made using a 15 ppm mass tolerance for the precursor and 0.05 Da mass tolerance for the fragments. All non-filtered search results were processed by Scaffold (version 4.0.5, Proteome Software, Inc.) utilizing the Trans-Proteomic Pipeline (Institute for Systems Biology) with threshold values set at 1% FDR for peptides and 1% FDR for proteins (two peptide minimum) and quantitative comparisons made using the total unique peptide count method with all samples normalized by total ion current for the run.

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AUTHOR CONTRIBUTIONS
Sai Ganesan: Conceptualization; methodology; validation; visualization; writing-original draft; writing-review and editing. Michael Feyder: Conceptualization; formal analysis; investigation; visualization; writing-original draft; writing-review and editing. Ilan Chemmama: Conceptualization; validation; visualization; writing-review and editing. Fei Fang: Data curation; formal analysis; methodology; validation; visualization; writing-review and editing. Michael Rout: Conceptualization; resources; writing-review and editing. Brian Chait: Conceptualization; formal analysis; funding acquisition; project administration; validation; writing-review and editing. Yi Shi: Conceptualization; data curation; formal analysis; funding acquisition; investigation; methodology; supervision; validation; visualization; writing-review and editing. Mary Munson: Conceptualization; funding acquisition; project administration; writing-original draft; writing-review and editing. Andrej Sali: Conceptualization; funding acquisition; project administration; supervision; writing-original draft; writing-review and editing.

CODE AVAILABILITY
The integrative structure modeling scripts are available at https://salilab.org/exocyst.

MODEL AVAILABILITY
The integrative exocyst structure has been deposited in the nascent public PDB repository, PDB-dev (https://pdb-dev.rcsb.rutgers.edu/), accession number PDBDEV-00000041. Raw data for Figure 4 and Figure S1 are available at https://salilab.org/exocyst.

DATA AVAILABILITY STATEMENT
The EM density map has been deposited in the EMDB, accession code 21226.

ORCID
Sai J. Ganesan © https://orcid.org/0000-0002-8335-2866
Michael J. Feyder © https://orcid.org/0000-0001-5233-6161
Ilan E. Chemmama © https://orcid.org/0000-0002-5657-3516
Michael P. Rout © https://orcid.org/0000-0003-2010-706X

Brian T. Chait © https://orcid.org/0000-0003-3524-557X
Mary Munson © https://orcid.org/0000-0003-2297-8053
Andrej Sali © https://orcid.org/0000-0003-0435-6197

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SUPPORTING INFORMATION
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