**Structural Insight into the Regulation of the SNARE Assembly by the Cell Polarity Protein Sro7**

The cells that comprise many tissues are polarized, meaning that they have distinct ‘sides’ with different membrane identities. For example, in the cells that line the intestine, the membrane that faces the space in the gut has special proteins responsible for uptake of nutrients, whereas the sides that contact neighboring cells have different proteins on their surfaces. Cell polarity is fundamental to many aspects of cell and developmental biology and it is implicated in differentiation, proliferation and morphogenesis in both unicellular and multi-cellular organisms. Loss of cell polarity can lead to uncontrolled tissue growth and cancers. To generate and maintain this polarized structure, specific proteins and lipids must be delivered to particular locations on the cell membrane. This process involves active transport of membrane-enclosed vesicles containing specific cargo to a target site, where the vesicle and target membranes then fuse to deliver the cargo. Soluble N-ethylmaleimide-sensitive factor activating protein receptor (SNARE) proteins in the vesicle membrane and target membrane form a complex that mediates membrane fusion. The process by which SNARE-mediated membrane fusion is coordinated with the machinery that transports the vesicle to the correct location is poorly understood.

In this work, Hattendorf *et al.* studied a yeast protein called Sro7 that is essential for delivery of vesicles from a mother to a budding daughter cell, which is another example of polarized cell growth. Sro7 and its relatives in higher organisms bind to SNARE proteins and are known to be essential for cell polarity, but their mechanism is unknown. The crystal structure of Sro7, determined with data measured at SSRL (beamline 11-1) and the Advanced Light Source, revealed a double-domain structure that is followed by a “tail” that binds to the surface of one of the domains (Figure 1). It was shown that removal of the tail promotes binding to a yeast SNARE protein and thereby blocks formation of SNARE complexes. Thus, the structure led to the discovery of an unanticipated mechanism for regulating SNARE complex assembly (Figure 2), and provided the first mechanistic data on this essential family of proteins.

**Figure 1.** Structure of Sro7. The N-terminal barrel is light green, the C-terminal barrel is light blue, and the tail is shown in dark blue.
Figure 2. Model for how Sro7 may coordinate release of the SNARE Sec9 with arrival of a secretory vesicle. Sro7 is associated with the plasma membrane at the site of budding. It is proposed that membrane association of Sro7 is coupled to displacement of the tail from its binding site. In this state, Sro7 binds to the Sec 9 SNARE regions (marked Qb and Qc) and prevents Sec9 from binding to its partner SNAREs (shown in red and blue) while localizing it to the eventual site of membrane fusion. Factors associated with the arriving vesicle would stimulate rebinding of the tail to the body of Sro7, releasing the Sec9 SNARE regions and allowing SNARE complex formation and membrane fusion to proceed, thereby coordinating arrival of the vesicle with membrane fusion.

This work was supported by grants from the NIH and the American Cancer Society.

Primary Citation

References