

Department of Biological Sciences Faculty of Science

## VIRTUAL BIOLOGY COLLOQUIUM

Friday, 30 Oct 2020 | 10 am | Online Zoom Session

Hosted by A/P Cynthia He

# Cytoskeletal architecture and cell morphogenesis in *Trypanosoma brucei*



#### About the Speaker

My diverse training in chemical biology, cell biology, and parasitology has given me a unique skillset to study cytoskeletal biogenesis in trypanosomatids. My graduate work in the laboratory of Carolyn Bertozzi focused on the importance of the localization of Golgi-resident enzymes in their substrate specificity and the production of cell surface glycoconjugates. I pursued my interest in the Golgi in the lab of Graham Warren at Yale and the Max Perutz Lab in Vienna, where I showed that the Trypanosoma brucei Polo-like kinase homolog (TbPLK) is important for Golgi biogenesis. We performed a series of proteomic screens to identify novel TbPLK binding partners and substrates, which uncovered key components of several cytoskeletal structures that had previously only been described morphologically. Since starting at Brown University, the primary goal of my laboratory has been to use these novel protein components to develop a molecular understanding of cytoskeletal biogenesis in T. brucei.

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### By Christopher de Graffenried

Department of Molecular Microbiology and Immunology, Brown University

Trypanosoma brucei causes human African trypanosomiasis and the livestock disease known as nagana, which both cause significant health and economic burdens in sub-Saharan Africa. Key to survival within hosts is the shape of the parasite, which allows it to move rapidly through crowded, high-viscosity environments such as blood and tissue. This shape comprises a tapered posterior and a narrow, pointed anterior end, generated and maintained by a helical sheath of microtubules (MTs), which underlies the cell surface and is termed the subpellicular array (SPA). Our goal is to establish the molecular mechanisms that drive SPA assembly and which will maintenance, determine how trypanosomes shape their cells. To do this, we are studying a series of MT-associated proteins including MT-crosslinking proteins and molecular motors using cell biological and biophysical approaches. Our research will contribute to a better understanding of the plasticity of MTs and the structures they create in an early branching eukaryote, which is vital to establishing their fundamental properties and potential functions in a broader range of eukaryotes.

#### Virtual Seminar Etiquette:

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- Questions can be asked after the presentation. You are encouraged to verbally ask questions or submit your questions via chat group.
- ✓ By being present at this meeting, information presented is a privilege and you agree that you would <u>NOT UNDERTAKE</u> any forms of recording/photo-taking.