**Weekly Colloquium**  
Tuesday, 5/22/2018, 12:30pm, Billings Building – Rosedale Conference Room

**“Reversible Redox Regulation of Cytoskeletal Dynamics to Control Neuronal Form and Function”**

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Our brains control such diverse abilities as movement, sensation, intelligence, speech, emotion, and memory only because our neurons communicate with one another and the rest of our body through highly organized networks. These neuronal networks or connections are formed when neurons send out their stalk-like axon and dendrite appendages. To accomplish this, neurons require actin and tubulin proteins to assemble together into long polymers (F-actin and microtubules, respectively) – and numerous extracellular stimuli have now been identified that alter the assembly and organization of these cytoskeletal structures. Yet, we still know little of how these extracellular cues exert their precise effects on the cytoskeleton. To better understand these mechanisms, my lab has been focusing on one of the largest families of extracellular cues, the Semaphorins (Semas) – which alter neuronal behaviors by eliciting destabilizing effects on both F-actin and microtubules. Our strategy has been to use model organisms and screening approaches to search for proteins that work in the signal transduction cascade utilized by Semas and their Plexin receptors. Among the proteins that we have identified, is a new family of intracellular proteins called the MICALs that are required for Sema/Plexin signal transduction. Now, recent work in my lab has revealed that the MICALs employ a previously unknown Redox signaling system to control the actin cytoskeleton. Namely, we have found that Mical is a novel F-actin disassembly factor – and our results reveal that Sema/Plexin-mediated reorganizations of the actin cytoskeleton can be precisely achieved in space and time through activation of Mical. We have also found that the MICALs belong to a class of oxidoreductase (Redox) enzymes and that Mical employs its Redox enzymatic activity to alter the properties of F-actin. Our work has gone on to identify that Mical uses F-actin as a direct substrate and post-translationally oxidizes conserved amino acids on actin, simultaneously dismantling F-actin and decreasing polymerization. Moreover, we find that this Sema/Plexin/Mical-mediated Redox regulation of actin is reversible (by a protein called SelR/MsrB) – and that this specific reversible Redox actin regulatory system directs multiple different biological processes in neuronal and non-neuronal tissues. Thus, MICALs and SelR/MsrBs form a reversible Redox cellular signaling system controlling actin filaments – the building blocks of neuronal form and function.

**Publications:**
