Growth and dynamics of C\textsubscript{60} molecules on Ag(111) RENEE DIEHL, HSIN LI, GUILHERME ABREU, HEEKEUN SHIN, Pennsylvania State Univ, A.K. SHUKLA, JULIAN LEDIEU, VINCENT FOURNÉE, LAURA SERKOVIC LOLI, CNRS-Université de Lorraine, KRISTIN MARINO, STEPHANIE SU, MICHAEL SNYDER, Pennsylvania State Univ — C\textsubscript{60} monolayers on Ag(111) form various structures that depend on the surface preparation. For films deposited at 300 K, the monolayer comprises close-packed domains that are incommensurate with the substrate. These domains progressively convert into a commensurate \( (2\sqrt{3} \times 2\sqrt{3})R30^\circ \) structure as the annealing temperature is increased. The annealing process activates the formation of vacancies beneath some of the C\textsubscript{60} molecules, resulting in a temperature-dependent equilibrium population of about 50\% of the molecules sitting on vacancies and 50\% sitting on top of Ag atoms in the commensurate phase. In addition to molecules in these two sites, about 0.5\% of the molecules image very brightly, and these molecules are in dynamical equilibrium with the other two types. This talk will focus on scanning tunneling microscopy and low-energy diffraction studies of the structure and dynamics of the monolayer phases of C\textsubscript{60} molecules on Ag(111).