Structure-Property Relationships of Topological Insulator Nanomaterials

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The last decade has witnessed an explosion of a new frontier in condensed matter physics: topological materials with an electronic band structure belonging to a different topological class than ordinary insulators and metals [1]. The non-trivial band topology gives rise to robust, spin-polarized electronic states with linear energy-momentum dispersion at the edge or surface of the materials. For topological materials to be useful in electronic devices, precise control and accurate detection of the topological states are needed in nanostructures, which can enhance the topological states.

The current challenge in the field is to ensure that the crystal quality of topological nanomaterials is sufficiently good in order to easily reveal and control the topological properties. For example, the binary chalcogenide topological insulators such as Bi2Se3 and Bi2Te3 suffer from chalcogen vacancies and surface oxidation [2], which increase the undesirable bulk carrier densities and decrease the electron mobility, detrimental to the detection of the topological edge states. In addition to the crystal quality, understanding the crystal structure – transport property relationships of topological nanomaterials is critical. For SnTe, a topological crystalline insulator, the required cubic crystal symmetry that provides the topological protection gets lost at low temperatures because SnTe goes through a structural transition from the cubic to the rhombohedral structure [3]. Thus, transmission electron microscopy and spectroscopy is critical for characterization of topological nanomaterials. The structural characterization obtained by electron microscopy must be then correlated to transport measurements to overcome the current challenge.

In this talk, I will discuss my group’s efforts in making superconducting SnTe nanowires [4], a candidate material for 1D topological superconductivity. Superconductivity can be induced in SnTe by doping with indium (In) (Figure 1). SnTe nanostructures are fabricated by metal-catalyzed chemical vapor deposition [3-7]. Transport properties of pristine, In-doped, and alloyed SnTe nanostructures are measured and correlated to structural characterization of the nanostructures, obtained by transmission electron microscopy and spectroscopy.

I will highlight the importance of careful structural and morphology characterizations in understanding the transport properties of topological nanostructures. A few examples will be discussed. We observed a structural transition of SnSexTe1-x nanostructures at a critical concentration of Se doping (Figure 2) [6], which was introduced to reduce the carrier density. We also observe surface defects on SnTe nanostructures, arising from dislocations embedded inside the bulk of SnTe nanostructures, which have consequences for the predicted topological properties [7].

Figure 1. Superconductivity in In-doped SnTe nanostructures: (a) nanoplate, (b) nanoribbon, and (c) nanowire.

Figure 2. Structural transition observed in SnSe$_x$Te$_{1-x}$ nano-plates as a function of the Se concentration, x. (a-f) pristine SnTe nanoplate with chemical mapping and structural characterization. (g) EDX spectrum to calibrate x. (h-m) SnSe$_{0.5}$Te$_{0.5}$ nanoplate with chemical mapping and structural characterization. (n) EDX spectra of SnSe$_x$Te$_{1-x}$ nanoplates with varying x. (o) Linear correlation between the nominal ratio, x, and the measured ratio, $x_{EDX}$. 