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Probing the Ab-initio Fermi Surface Efficiently using Wannier Interpolation JONATHAN YATES, IVO SOUZA, LBNL and University of California, Berkeley, XINJIE WANG, DAVID VANDERBILT, Rutgers University — Modern *ab-initio* techniques are able to provide an accurate description of the electronic structure for a wide range of materials. However, evaluation of the transport properties of metals requires an extremely detailed, and hence computationally expensive, sampling of the Fermi surface. We show that the electron group velocity and effective mass can be obtained directly from the Wannier representation of a system. This leads to an efficient and precise method for the calculations of the ordinary Hall coefficient and thermoelectric power for a variety of materials.

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