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THE ELECTRONIC HAMILTONIAN FOR CUPRATES

James F. Annett
The Pennsylvania State University
University Park, PA

A.K. McMahan
Lawrence Livermore National Laboratory
Livermore, CA

Richard M. Martin
University of Illinois at Urbana-Champaign
Champaign, IL

A realistic many-body Hamiltonian for the cuprate superconductors should include both copper d and oxygen p states, hopping matrix elements between them, and Coulomb energies, both on-site and inter-site. We have developed a novel computational scheme for deriving the relevant parameters *ab initio* from a constrained occupation local density functional. The scheme includes numerical calculation of appropriate Wannier functions for the copper and oxygen states. Explicit parameter values are given for La_2CuO_4 . These parameters are generally consistent with other estimates, and with the observed superexchange energy.

Secondly, we address the question: can this complicated multi-band Hamiltonian be reduced to a simpler one with fewer basis states per unit cell? We propose a mapping onto a new two-band effective Hamiltonian with one copper d and one oxygen p derived state per unit cell. This mapping takes into account the large oxygen-oxygen hopping given by the *ab initio* calculations.

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