

Abstract Submitted
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Quantum Monte Carlo calculations of defects in ZnO WILL TIP-
TON, RICHARD HENNIG, Cornell University — The semiconductor ZnO holds
much promise for many applications due to its ability to display a wide variety of
properties. However, an understanding of the structure and electronic structure of
the defects which lead to the material's properties has proven difficult. Previous
density functional theory (DFT) calculations by Van De Walle et. al. [1] using
the local density approximation, suggest that hydrogen interstitials and oxygen va-
cancies may be the dominant defect present in the as-grown material leading to its
n-type conductivity. However, the corrections accounting for DFT's band gap prob-
lem lead to a wide variance in defect formation energies reported and uncertainty in
the qualitative results. More recent calculations by Oba et. al. [2] using the HSE
hybrid functional confirm the presence of these defect structures and their electronic
transition levels. We are performing Quantum Monte Carlo calculations for the de-
fects in ZnO to determine the defect formation energies and transition levels and to
verify the previous DFT calculations. [1] C. G. van de Walle, Phys. Rev. Lett. 85,
1012 (2000). [2] F. Oba, Phys. Rev. B 77, 245202 (2008).

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