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Structural phase transitions from first-principles calculations

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Describing atomic vibrations from first-principles accurately is of paramount importance to understand the thermodynamic and transport properties of solids, especially to understand structural phase transitions. Phonon dispersions are routinely calculated within the harmonic approximation, and transport properties can be studied by estimating the electron-phonon and phonon-phonon interactions within perturbation theory. Nevertheless, whenever the amplitude of the atomic displacements largely exceeds the range in which the harmonic potential is valid, the harmonic approximation completely fails without allowing a perturbative expansion. This clearly hinders any ab initio calculation of structural phase transitions in these situations

The stochastic self-consistent harmonic approximation (SSCHA) that we have developed [1,2,3,4] offers an efficient method to calculate vibrational properties of solids even when the harmonic approximation completely collapses. The method is variational and takes into account quantum and thermal effects rigorously. With our recent developments on the SSCHA method [3], we show how phonon frequencies should be defined from the second derivative of the free energy, which allows calculating the transition temperature of structural second-order phase transitions. Moreover, the new developments [3] allow calculating thirdorder anharmonic force-constants, which determine thermal properties, beyond the perturbative limit.

In this lecture we will present the method and several applications of it in superconducting hydrides, charge-density-wave systems, and thermoelectric materials.

- [1] I. Errea, M. Calandra, and F. Mauri, Phys. Rev. Lett. 111 (2013) 177002.
- [2] I. Errea, M. Calandra, and F. Mauri, Phys. Rev. B 89 (2014) 064302.
- [3] R. Bianco, I. Errea, L. Paulatto, M. Calandra, and F. Mauri, Phys. Rev. B 96 (2017) 014111.
- [4] L. Monacelli, I. Errea, M. Calandra, and F. Mauri, Phys. Rev. B 98 (2018) 024106.

Summary

Topic

1. Theoretical and experimental methods

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