# Van Hove singularities and spectral smearing in high-temperature superconducting H<sub>3</sub>S

Yundi Quan and Warren E. Pickett\*

Department of Physics, University of California Davis, Davis, California 95616, USA (Received 3 September 2015; published 25 March 2016)

The superconducting phase of hydrogen sulfide at  $T_c = 200$  K observed by Drozdov and collaborators at pressures around 200 GPa is simple bcc  $Im\bar{3}m$  H<sub>3</sub>S from a combination of theoretical and experimental confirmation. The various "extremes" that are involved—high pressure implying extreme reduction of volume, extremely high H phonon energy scale around 1400 K, extremely high temperature for a superconductor necessitates a close look at new issues raised by these characteristics in relation to high  $T_c$  itself. First principles methods are applied to analyze the H<sub>3</sub>S electronic structure, beginning with the effect of sulfur and then focusing on the origin and implications of the two van Hove singularities (vHs) providing an impressive peak in the density of states near the Fermi energy. Implications arising from strong coupling Migdal-Eliashberg theory are studied. It becomes evident that electron spectral density smearing due to virtual phonon emission and absorption must be accounted for in a correct understanding of this unusual material and to obtain accurate theoretical predictions. Means for increasing  $T_c$  in H<sub>3</sub>S-like materials are noted.

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### I. INTRODUCTION

The recent discovery of superconducting hydrogen sulfide under high pressure by Drozdov and collaborators [1–3], and remarkably predicted a year earlier by Duan *et al.* [4], has reinvigorated the quest for room temperature superconductivity. The predicted structure has been confirmed by x-ray diffraction studies by Einaga *et al.* [5] that show that sulfur lies on a bcc sublattice; the protons cannot be seen in x-ray diffraction. The resistivity transitions were also confirmed by Einaga *et al.*. The experimental reports indicate critical temperatures up to  $T_c = 203$  K in the pressure range of 200 GPa, based on the resistivity transition, the effect of magnetic field on  $T_c$ , on a H isotope shift of the right sign and roughly the expected magnitude [1], and the Meissner effect has been demonstrated [2].

In a success of predictive theory in this area, the magnitude of  $T_c$  in the 200 GPa pressure range was obtained from first principles calculation *prior to experiment* [4] and confirmed by others [6–8], so there can be little doubt that 200 K superconductivity has been achieved in the structurally simple compound H<sub>3</sub>S, pictured in Fig. 1. The finding that H vibrations provide the mechanism seems to confirm the suggestion of Ashcroft that dense hydrogen should superconduct at high temperature [9], however, evidence is increasing that H-rich materials [10] are substantially different and more promising than pure hydrogen until TPa pressures can be reached. Early quantitative estimates [11] of  $T_c$  for metallic H were in the 250 K range; more recent values [12] for pressures of several TPa lie in the 500–750 K range. The phase diagram of this system is uncertain, however, due to the quantum nature of the proton.

Although comprehensive calculations based on density functional theory (DFT) linear response formalism and Eliashberg theory [13] have been reported and seem convincing,  $H_3S$  turns out to be more intricate than the initial reports suggest. Using their self-consistent harmonic approximation, Errea *et al.* [6] find substantial corrections due to anharmonicity: At

Other basic questions have yet to be addressed. First, why are the electron-phonon matrix elements as large as they are? While the main causative property behind the high  $T_c$  is the (understandably) high phonon frequencies that set the energy scale for  $T_c$ , substantial electron-ion matrix elements are also required. Second, Flores-Livas *et al.* [8] have investigated the energy dependence of the spectrum around the Fermi level, finding that it influences the theoretical predictions, which are overly optimistic when energy dependence is neglected. Both Akashi *et al.* [14] and Flores-Livas *et al.* have solved the gap equations, providing theoretical predictions of the gap as well as  $T_c$  without using the Allen-Dynes equation. The questions posed by intricacies in the density of states (DOS) and the role

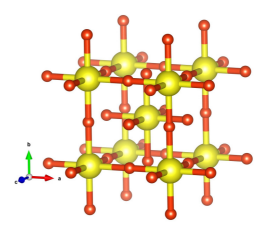


FIG. 1. Crystal structure of  $Im\bar{3}m$  H<sub>3</sub>S. Nearest neighbor S-H bonds are shown. At this nearest neighbor level, the structure consists of two interleaved ReO<sub>3</sub> sublattices, displaced relative to one another by the body-centering vector (1,1,1)a/2.

<sup>200</sup> GPa, anharmonicity increases the characteristic frequency  $\omega_{\log}$  by 3%, the electron-phonon interaction (EPI) strength  $\lambda$  is decreased by 30%, and the predicted value of  $T_c$  falls 22% from 250 K to 194 K. Potentially important for further understanding is their finding that anharmonicity shifts coupling strength to H-S bond stretch modes, from H-S bond bending (alternatively, H-H bond stretch) modes.

<sup>\*</sup>wepickett@ucdavis.edu

of zero point vibrations has stimulated work by Bianconi and Jarlborg [15].

More fundamentally there is the question "why  $H_3S$ ? why sulfur?" Several H-rich materials have been studied at high pressure (see references in Refs. [4,6–8], and Bernstein et al. [16]), and although some are predicted to superconduct up to several tens of kelvins, H<sub>3</sub>S is a singular standout. Li *et al.*, [17] for example, studied the  $H_2S$  stoichiometry for stable compounds up to similar high pressures, finding a maximum  $T_c$  of "only" 80 K. There is little understanding so far of the microscopic cause of very high  $T_c$ , beyond the obvious expectation of higher phonon frequencies at high pressure; the origin of the large matrix elements remain obscure. Papaconstantopoulos and collaborators [7] calculated the pressure dependence of matrix elements, finding increasing H scattering with increasing pressure. More basically one can ask, is there something special about sulfur, and the underlying electronic structure, that provides the platform for such high  $T_c$ ?

It is the last of these questions we address initially in this paper. An obvious feature for study is the strikingly sharp peak in the density of states N(E) due to two van Hove singularities (vHs) separated by 300 meV very near the Fermi level  $E_F$ . There is a large literature on the connection between peaks in N(E) and high  $T_c$  in the A15 class of materials [18] and later in the high temperature superconducting cuprates [19,20], but their importance for H<sub>3</sub>S is unclear. van Hove singularities near the Fermi level can enhance  $N(E_F)$  and thus the electronphonon coupling strength  $\lambda$  due to increased number of available states to participate, but there are additional questions to address.

The paper is organized as follows. Methods are described in a brief Sec. II. In Sec. III the general electronic structure and the charge density near  $E_F$  are presented and discussed. The two van-Hove singularities are identified, quantified, and analyzed, and the relation between them is identified. In Sec. IV we address the peak in N(E) in the light of strong electron-phonon coupling (EPC), high frequencies, and thermal and EPC-induced smearing. Section V presents scenarios for further increase in coupling strength  $\lambda$ , and raising of  $T_c$  toward room temperature, in this and similar systems. A short summary is provided in Sec. VI.

### **II. METHODS**

Density functional calculations have been carried out using both the linearized augmented plane wave (LAPW) method based Wien2k code [21] and the linear combination of atomic orbitals based FPLO code [22]. The PBE implementation [23] of the generalized gradient approximation (GGA) is used as the exchange correlation functional. Except where noted, the results will be those from Wien2k. The crystal structure of H<sub>3</sub>S is  $Im\bar{3}m$  with a lattice constant of 5.6 a.u. corresponding [7] to a pressure of 210 GPa. All atoms lie at high symmetry sites.

To study the vHs points near the Fermi level, a very fine k mesh containing 8094 points in the irreducible Brillouin zone is used. Sphere radii R for H and S are 0.97  $a_0$  and 1.81  $a_0$  respectively, with basis set cutoff determined by  $R_H K_{max} = 6$ . The results we discuss are insensitive to these choices. The tight-binding parameters we present

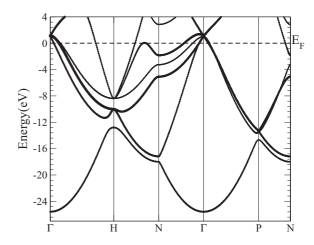


FIG. 2. DFT band structure of  $H_3S$  at a = 5.6 a.u., with the Fermi level set to zero. The darker bands in the -10 eV to +4 eV region have more H 1s character than the others. The position of the van Hove singularity discussed in the text is not visible along these symmetry lines.

were obtained from projection to symmetry-adapted Wannier functions as implemented in FPLO.

#### **III. ELECTRONIC STRUCTURE**

The calculated lattice constant [7] at 210 GPa, a = 5.6 a.u. corresponding to a volume 58% of the zero pressure volume in the same structure, is used in all calculations. The rather simple band structure of H<sub>3</sub>S is shown in Fig. 2 and, up to some differences near the Fermi level, is consistent with previous work [4,16,24]. The very large occupied bandwidth of 26 eV reflects both the nearly free electron aspects of the the lower part of the valence bands, with this aspect being enhanced by the reduced volume that increases electron density and hopping amplitudes. Both effects increase the bandwidth: The reduced volume increases the density and hence elevates the Fermi energy, also the increased bandwidth reflects increased hopping amplitudes. Several bands cross the Fermi level, with varying velocities.

The DOS N(E) on a broad scale is presented in Fig. 3. The occupied bandwidth is 26 eV. Over the lowest 20 eV of this range, the DOS has a remarkably free-electron-like  $\sqrt{E}$  shape, without significant structure. Over the lower end of this region, the DOS is dominated by S 3s character, above which H 1s and S 3p character enter and mix. Then, at -4 eV and +5 eV two substantial and rather narrow peaks emerge, indicative of very strong hybridization, presumably being bonding and antibonding signatures. Double valleys lie at -2 eV and +2 eV, between which a very sharp peak, related to two van Hove singularities (vHs) 300 meV apart in energy, juts upward. The Fermi energy  $E_F$  (set to zero throughout) lies very near the upper vHs. The vHs are discussed further in Sec. III D.

Papaconstantopoulos *et al.* provided [7] the orbital projected DOS for H<sub>3</sub>S. The lower peak, at -4 eV, is very strongly S 3*p* in character, while the sharp peak at E<sub>F</sub> is a strong mixture of S 3*p* with H 1*s*. The upper peak at +5 eV is a mixture of these two characters with states of more planewave-like character. We note that the shape of the DOS in Fig. 3 is

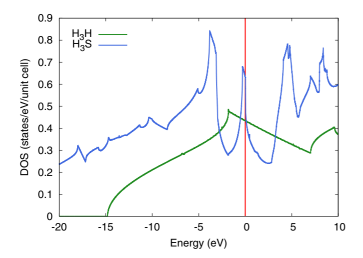


FIG. 3. Density of states of  $H_3S$  compared to that of  $H_3H$  at the same volume, from Wien2k calculation. The bandwidth is extremely large due to the broadening caused by short interatomic distances. The peak in N(E) arises from two van Hove singularities within 300 meV of each other and occurring in the region of the Fermi energy.  $N(E_F) = 0.63$  states/eV for  $H_3S$ . The impact of H bonding with the S 3*p* orbitals is evident.

significantly different than that of Duan *et al.*, [4] in spite of the similarity of the bands along symmetry directions.

We demonstrate in Sec. IV that for physical superconducting properties, thermal and, more importantly, dynamical broadening makes details of N(E) fine structure unimportant in some respects. This unimportance *does not however apply* for the underlying theory based on the static lattice, where it has serious consequences as discussed in Sec. V. Briefly, the issue is that so much in conventional EPC theory is formulated and evaluated in terms of the specific value of N( $E_F$ ) assuming N(E) is slowly varying, which is not the case in H<sub>3</sub>S.

### A. Importance of sulfur

As one clear means to assess the effect of H-S orbital mixing, Fig. 3 displays N(E) also for a hypothetical material, H<sub>3</sub>H, in which the S atom is replaced by H in the same structure. The spectrum is very different in a wide range around the Fermi level. Unusually, an almost perfectly linear region of N(E) extends between two vHs 9 eV apart, on opposite sides and far from the Fermi level. The point however is that the strong structure in H<sub>3</sub>S in the -4 eV to +5 eV region is missing in H<sub>3</sub>H. The hypothetical nature of H<sub>3</sub>H, which is in fact a simple cubic lattice of hydrogen, is corroborated by our calculation of the phonon spectrum, which results in many imaginary frequencies signaling dynamic instability.

Returning to the H<sub>3</sub>S DOS, such strong structure in N(E) reflects strong mixing between orbitals lying in this energy range, which are the H 1*s* and S 3*p* valence orbitals. The orbital projected DOS (PDOS) presented by Papaconstantopoulos *et al.* [7] shows that S 3*s* participation is becoming small around  $E_F$ . Their PDOS helps to understand the strong DOS structure. The peak at -4 eV is largely S 3*p* character with some H 1*s* contribution. The peak at +5 eV has, surprisingly, a large contribution from Bloch orbitals with *d* symmetry around

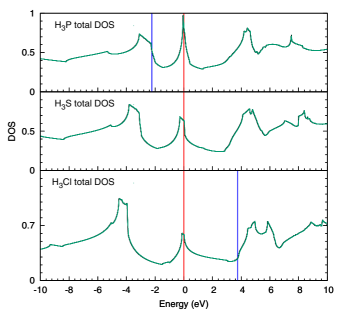


FIG. 4. Density of states (per eV-f.u.) of the sequence  $H_3X$ , X = P, S, Cl, aligned at the band filling of  $H_3S$ . The Fermi levels of  $H_3P$  and  $H_3Cl$  (using the  $H_3S$  structure, hence hypothetical) are shown by the vertical lines at appropriate band fillings. The similarity is striking; a rigid band (or virtual crystal) picture of alloys near  $H_3S$  holds qualitatively.

the S site, with some participation of all of the orbitals besides S *s*. The peak at  $E_F$ —the important one bounded by two closely spaced vHs—is a strong mixture of H 1*s* with S 3*p*, whose corresponding tight binding hopping parameters will have a correspondingly large hopping amplitude.

We also compare in Fig. 4 N(E) for H<sub>3</sub>X, X = P and Cl, to that of H<sub>3</sub>S, aligned at the H<sub>3</sub>S band filling. Despite some variation in the on-site 3*p* energy and electronegativity, N(E) is quite similar among the three. The peak in N(E) at E<sub>F</sub> is narrower for both P and Cl, i.e. the vHs are closer together. The strength of the peak at E<sub>F</sub> appears to decrease somewhat from  $P \rightarrow Cl$ . The distinction of S is that it places the band filling at the peak in N(E).

To quantify H-S hybridization relative to that of H-H, symmetry projected Wannier functions (WFs) were calculated using the FPLO code, projecting on S 3s and 3p, and H 1s atomic orbitals, for a total of seven WFs per primitive cell. The WF basis provides a tight-binding representation of the DFT bands over a 30+ eV region, although requiring hopping parameters out to several neighbors. We denote the sulfur states by S and P and the H orbital by s in the subscripts of the hopping integrals. The WFs are pictured in Fig. 5, with both S 3p and H 1s WFs showing anisotropy and extension, reflecting the strong H-S mixing.

The on-site energies and largest Slater-Koster parameters are listed in Table I. Relative to  $E_F = 0$ , the on-site energies (compared to those reported by Bernstein *et al.* [16], provided here in parentheses) are:  $\varepsilon_S = -8.0$  (-8.6) eV;  $\varepsilon_s = -5.5$ (-5.0) eV;  $\varepsilon_P = 0.03$  (-1.3) eV. The procedures used by Bernstein *et al.* are not exactly the same as ours, with the difference indicating the level of confidence one should assign to these site energies considering the nonuniqueness of

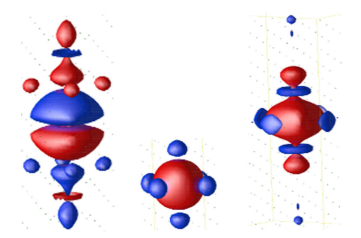


FIG. 5. Left: sulfur  $3p_z$  Wannier function, showing strong mixing with H 1s contributions above and below. Center: sulfur 3s Wannier function, with minor hybridization to neighboring 1s orbitals. Right: hydrogen centered Wannier function, revealing strong hybridization of H 1s with S  $3p\sigma$  above and below,, as well as mixing with neighboring H 1s orbitals along the other two axes.

tight-binding representations. It is eye catching that our sulfur P on-site energy is indistinguishable from  $E_F$ .

The largest hopping parameters (Table I) are nearest neighbor (n.n.)  $sP\sigma$  (-5.4 eV) and  $sS\sigma$  (-4.4 eV), the former confirming that H-S n.n. hopping is dominant in creating the DOS structure. We have calculated the DOS for a single sublattice, which is ReO<sub>3</sub>-structure H<sub>3</sub>S, and it is nothing like the full DOS. The n.n. H-H hopping  $ss\sigma$ , at -2.8 eV, couples the sublattices strongly, and S-S second neighbor hoppings, one lattice constant apart, are 1.3–1.8 eV in magnitude. Interesting are the 4th neighbor H-H hoppings, between atoms one lattice constant apart. The hopping through the S atom  $(ss\sigma)_4$  is -1.1 eV, while through an intervening H atom  $(ss\sigma)'_4$  is half that size with opposite sign.

As mentioned, the peak in N(E) reveals that the band filling in H<sub>3</sub>S happens to be almost perfect for large response to perturbations, with  $E_F$  lying within the sharp peak. Retaining the same band filling suggests substituting Se or Te for S. Flores-Livas *et al.* [8] have done parallel calculations for H<sub>3</sub>S and H<sub>3</sub>Se. The H<sub>3</sub>Se frequency  $\omega_{log}$  is 10% *higher* but the

TABLE I. Selected tight-binding hopping amplitudes (in eV) from the transformation to Wannier functions. The subscripts indicate the index of the neighbor:  $1 \equiv 1$ st neighbor,  $2 \equiv 2$ nd neighbor, etc. For H there are two 4th neighbors one lattice constant apart: one through the S atom, denoted "4," one through another H atom, denoted "4'."  $(\widetilde{PP})_1$  indicates  $\frac{1}{4}(PP\sigma)_1 + \frac{3}{4}(PP\pi)_1$ , which only occurs in this combination.

$\epsilon_{S}$	-7.98	$(sP\sigma)_1$	-5.42
$\epsilon_s$	-5.46	$(PP\sigma)_2$	-1.83
$\epsilon_P$	-0.03	$(SP\sigma)_2$	1.29
$(sS\sigma)_1$	-4.37	$(SS\sigma)_2$	0.94
$(ss\sigma)_1$	-2.80	$(sP\sigma)_2$	-0.93
$(ss\sigma)_4$	-1.14	$(\widetilde{PP})_1$	0.60
$(ss\sigma)'_4$	0.55	$(SS\sigma)_1$	0.30

calculated value of  $\lambda$  is lower by 40%, with the resulting  $T_c$  being lower by 27%. The changes of  $\omega_{\log}$  and  $\lambda$  indicate that the product  $\eta = N(E_F)\langle I^2 \rangle$  (not reported) is lower by 20% for the Se compound. Here  $\langle I^2 \rangle$  is the Fermi surface average of the square of the electron-H ion scattering matrix element. With H so dominant in the EPC and H modes separated from S (or Se) modes, electron-phonon coupling and superconductivity is dominated [7] by the H contribution

$$\lambda_H = \frac{N(E_F)\langle I_H^2 \rangle}{M_H \omega_H^2},\tag{1}$$

where the matrix element refers to scattering from the displaced H potential and  $\omega_H$  is a characteristic frequency from H modes.

The other isovalent "chalogenide" is oxygen, which is quite different from S chemically with H. We have found that the DOS of H<sub>3</sub>O in the H<sub>3</sub>S structure differs substantially from that of H<sub>3</sub>S. It may be relevant that H<sub>2</sub>O does not metalize until *much higher* pressures than are being considered here. Heil and Boeri [24] have considered bonding, EPC, and  $T_c$  where sulfur is alloyed with other group VI atoms. With alloying treated in the virtual crystal approximation (averaging pseudopotentials), they have suggested that a more electronegative ion will help. This leaves only oxygen in that column, and they calculated that a strong increase in matrix elements compensates a considerable decrease in N(E<sub>*F*</sub>), so that  $\lambda$  might increase somewhat. Ge *et al.* have also suggested partial replacement [25] of S, with P being the most encouraging, due to the increase in  $N(E_F)$ .

While S changes the electronic system very substantially from that of H<sub>3</sub>H and its  $T_c$  is spectacular, it may not be so special. A variety of calculations have predicted (see Durajski *et al.* [26] for references) high values of  $T_c$  (in parentheses) for H-rich solids: SiH<sub>4</sub>(H<sub>2</sub>)<sub>2</sub> (107 K at 250 GPa), B<sub>2</sub>H<sub>6</sub>H (147 K at 360 GPa), Si<sub>2</sub>H<sub>6</sub> (174 K at 275 GPa), CaH<sub>6</sub> (240 K at 150 GPa). Whether any general principles can be extracted from these results remains to be determined.

### B. Charge density within 1 eV of Fermi level

The charge density from states within 1 eV of  $E_F$  is shown in Fig. 6; it is this density whose coupling to H vibrations gives strong coupling and the very high value of  $T_c$ . Results from smaller energy slices are no different, indicating that the states in this range have the same character. The density around S is strongly distorted from spherical symmetry, having substantial maxima in the direction of neighboring H atoms. The H density is strongly elongated toward the two neighboring S atoms. These shapes reflect strong covalent H 1s-S  $3p_{\sigma}$  interaction discussed above. There remains a density minimum in the bond center rather than a bond charge maximum. This character is typical of strong directional bonding in metallic compounds.

#### C. van Hove singularities of H<sub>3</sub>S

N(E) in the energy range -1.0 eV to 0.2 eV was fit to the following piecewise expression [27] for 3D vHs near two

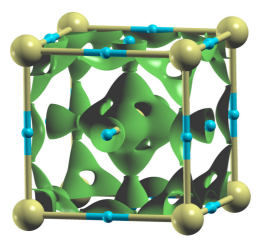


FIG. 6. Isosurface of the charge density obtained from states in the energy range  $E_F \pm 1$  eV. Sulfur atoms (yellow spheres) lie at the corners and at the body center; H atoms are denoted by small blue spheres.

singularities at energies  $\epsilon_{lo} < \epsilon_{hi}$ 

$$N(E) = \begin{cases} a_1 \sqrt{|b_1 - \epsilon|} + c_1 \epsilon + d_1 & \epsilon < \epsilon_{lo} \\ a_2 \epsilon + b_2 & \epsilon_{lo} < \epsilon < \epsilon_{hi}. \end{cases}$$
(2)  
$$a_3 \sqrt{|\epsilon + b_3|} + c_3 \epsilon + d_3 & \epsilon_{hi} < \epsilon \end{cases}$$

Energy isosurfaces at the two vHs energies are presented in Fig. 7. The vHs points lie at either end of a line where low velocity regions of two sheets of Fermi surface break apart at the zone boundary and then "unzip" until they separate into disjoint sheets. The vHs (from the FPLO bands) at -0.43 eV occurs at  $(-0.42, 0.21, 0)\pi/a$  and symmetric points. In the local principal axis coordinate system the effective masses are  $-0.15m_e$ ,  $1.36m_e$ ,  $0.14m_e$ , giving a thermal (or DOS) mass  $m_{\rm th} \equiv |m_1m_2m_3|^{1/3} = 0.31m_e$ . The thermal mass is one measure of the strength of the vHs. For the one at -0.11 eV, the masses are  $-0.83m_e$ ,  $-0.16m_e$ ,  $0.56m_e$ , and  $m_{\rm th} = 0.42m_e$ . We return to the importance of vHs effective masses in Sec. V.

# IV. IMPACT OF DYNAMICAL ELECTRON-PHONON INTERACTION

Thermal broadening as described by the Fermi-Dirac distribution function has well understood consequences and is a factor in determining (in limiting)  $T_c$ , though it is usually not thermal broadening of the DOS, which is normally constant over an energy range of several  $k_BT_c$ . The purely thermal aspect is formalized in the gap to thermal broadening scale ratio  $2\Delta/\pi k_BT_c \approx 1$ , and is central to, but standard in, Eliashberg theory. Specifically, it is thermally excited electron and hole quasiparticles that overcome pairing and restores the normal state above  $T_c$ .

However, the sharp structure in N(E) on the scale of relevant phonon energies requires an extension of conventional implementations of Eliashberg theory [13]. Conventionally, a constant N(E) on the phonon energy scale is assumed, so all scattering processes can be considered as confined to the Fermi surface  $E = E_F$ . This issue was confronted long

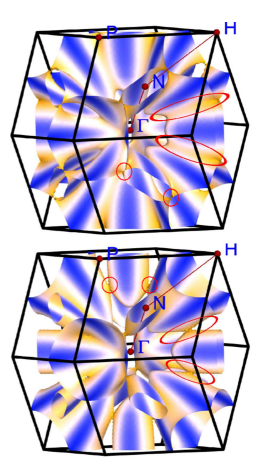


FIG. 7. Top: isoenergy contour from FPLO for the vHs at E = -0.43 eV. Bottom: similarly for the vHs at E = -0.11 eV. The red circles pinpoint two of the symmetry related vHs in each case. The red ellipse outlines the region joined by the two van Hove singularities; two energy surfaces are "unzipped" as the energy increases between the two vHs. Color denotes the velocity, which ranges from zero (darkest tan, at the vHs) to  $2.5 \times 10^8$  cm/s (deep blue).

ago [28,29], because of the sharp structure in N(E) in several of the then-high- $T_c$  A15 structure compounds, viz. Nb<sub>3</sub>Sn, V<sub>3</sub>Si, Nb<sub>3</sub>Ge, with  $T_c \sim 20$  K, and has been followed up in related applications [30,31].

In H<sub>3</sub>S around 200 GPa, the representative frequency is  $\Omega \sim 1300 \text{ K} = 112 \text{ meV}$ . We have chosen this value based on the DFT-based calculations of (harmonic)  $\omega_{log}$  frequencies ranging from 1125 K to 1450 K. [4,6,8,14,24,25], A single frequency moment is sufficient, since the logarithmic, first, and second moments differ only at the 2% level; we denote this frequency  $\Omega$ . Using a somewhat different value of  $\Omega$  below would not change our conclusions.

From visual examination, the sheets of the constant energy surfaces at  $E_F \pm \Omega$  do not differ much from the Fermi surfaces at  $E_F$  (see Fig. 7), the differences occurring in small pockets around  $\Gamma$  (not visible) and along a line connecting the two vHs. Inter-vHs scattering could be interesting: Although it involves a small amount of phase (*k*) space, it incorporates a disproportionate fraction of states with low to vanishing velocity. Possible complication from inter-vHs scattering and nonadiabatic processes lie beyond the scope of our discussion.

### A. Formalism

It has been known since the work of Engelsberg and Schrieffer [32] and Shimojima and Ichimura [33] fifty years ago that, for a characteristic phonon frequency  $\Omega$  in an interacting electron-phonon system, electron spectral density is spread from its noninteracting  $\delta$ -function spike at  $E_k$  up to a few  $\Omega$ . The spectral density arises from the electronic self-energy that is treated for superconducting systems by Eliashberg theory. When N(E) is slowly varying over a scale of a few  $\Omega$  it is rare to notice the effects of such broadening except possibly in direct measurements where phonon sidebands may be observed in photoemission spectra. For situations as in H<sub>3</sub>S where Bloch state character is slowly varying in energy but N(E) varies rapidly, the normally simple electron-phonon formalism becomes more challenging. Drozhov studied the effects of EPC in the vicinity of a vHs where Migdal's theorem is violated, and found severe renormalizations [34]. If these are confined to a very small phase space, however, the effects on most properties may be minor.

For the case of rapidly varying N(E) but neglecting violations of Migdal's theorem, the generalization of Eliashberg theory has been formulated and applied to the A15 compounds [29,35]. One feature that is distinctive in H<sub>3</sub>S compared to most other EPC superconductors is that  $T_c$  is an order of magnitude higher, because the frequencies are comparably higher, and simple thermal broadening is correspondingly larger and requires attention. The second factor in common, and the important one, is that strong EPC causes an effective smearing of the electronic spectral density due to exchange of virtual phonons appearing in Migdal-Eliashberg theory excitations described by the electron Green's function are part electron, part phonon. This broadening is given by the imaginary part of the interacting electronic Green's function

$$G_k^{-1}(\omega) = \omega - [E_k - \mu(T)] - M_k(\omega; T) - i\Gamma_k(\omega; T).$$
(3)

Here  $E_k$  is the DFT band energy,  $\mu(T)$  is the chemical potential, and M and  $\Gamma$  are the real and imaginary parts of the phonon-induced self-energy.

The spectral density  $A(\omega)$  is the interacting analog of the band DOS N(E):

$$A(\omega) = \sum_{k} A_{k}(\omega) = \frac{1}{\pi} \sum_{k} |ImG_{k}(\omega)|$$
(4)  
$$= \frac{1}{\pi} \sum_{k} \frac{\Gamma_{k}(\omega)}{[\omega - (E_{k} - \mu) - M_{k}(\omega)]^{2} + \Gamma_{k}(\omega)^{2}}$$
$$\rightarrow \int d\xi \frac{\Gamma/\pi}{(\omega - \xi)^{2} + \Gamma^{2}} N(\xi).$$

In the last expression the Brillouin sum has been converted into an energy integral by inserting  $\int dE\delta(E - E_k)=1$ , assuming that only  $E_k$  (and not wave function character, hence not Mor  $\Gamma$ ) depends on k near  $E_F$ , and  $\xi_k = \varepsilon_k - \mu + M(k,\xi_k)$  is the quasiparticle energy. This simplification is usually fine for electron-phonon coupling in a standard Fermi liquid, as wide-band H<sub>3</sub>S appears to be.

There is strong rearrangement of spectral density even before this smearing effect of electron damping  $\Gamma$ . For temperatures and frequencies  $\omega$  up to the order of the characteristic phonon energy  $\Omega$  or more, the behavior of the real part  $M_k$  is linear  $dM_k/d\omega = -\lambda_k$ , where  $\lambda_k$  is the EPC strength at k whose average over the Fermi surface is  $\lambda$ . The equation for  $\xi_k$  in the previous paragraph then gives for the quasiparticle energy

$$\xi_k = \frac{E_k - \mu}{1 + \lambda_k}.\tag{5}$$

This equation expresses the phonon-induced Fermi surface mass enhancement  $1 + \lambda_k$ , and  $(1 + \lambda_k)^{-1}$  is the quasiparticle strength, i.e., the fraction of the electron's  $\delta$ -function spectral density at  $\xi_k$  and whose average in H<sub>3</sub>S is 1/3 ( $\lambda \approx 2$ ). Twothirds of the spectral weight is spread from the quasiparticle energy  $\xi_k$  by up to a few times  $\Omega$ . This is a serious redistribution of weight that we cannot treat in any detail without explicit solution for the Eliashberg self-energy on the real axis.

Notwithstanding the complications, in an interacting system the thermal distribution function containing all complexities can be handled formally to provide insight into this "varying N(E)" kind of system. The interacting thermal distribution (state occupation) function  $f(E_k)$  is defined as the thermal expectation of the number operator  $n_k$ 

$$f(E_k) = T \sum_{-\infty}^{\infty} G_k(i\omega_n) e^{i\omega_n \eta}$$
$$= \int_{-\infty}^{\infty} d\omega f_{\circ}(\omega) A(E_k, \omega), \qquad (6)$$

where the Matsubara sum with positive infinitesimal  $\eta$  has been converted into an integral in the last expression and  $f_{\circ}(E)$  is the (noninteracting) Fermi-Dirac distribution. The *k* dependence of  $M_k$  and  $\Gamma_k$  are considered to be weak and replaced by a Fermi surface average. The interacting distribution function can be expressed as the noninteracting one broadened by [35]  $\Gamma_k(\omega)$  as N(E) is broadened in Eq. (4).

Several thermal properties can be formulated [35] in terms of the interacting (broadened and in principle mass renormalized) density of states  $\mathcal{N}(E)$ . Returning to single particle language, the spectral density at  $E_F$  is approximately

$$\mathcal{N}(E_F) = \int dE \frac{\Gamma/\pi}{(E - E_F)^2 + \Gamma^2} N(E).$$
(7)

For energies a few  $\Omega$  around  $E_F$  the extension  $E_F \rightarrow E$  to give  $\mathcal{N}(E)$  will be reasonable. Then, returning to the distribution function, the total electron number can be written [35] in two ways

$$N_{el} = \int dE f(E) N(E) = \int d\omega f_{\circ}(\omega) \mathcal{N}(\omega), \qquad (8)$$

illustrating that interaction effects can be exchanged between the distribution function and, in this instance, the interacting and noninteracting density of states. Thus in a region around  $E_F$  the spectral density is the band density of states broadened by a Lorentzian of halfwidth  $\Gamma$ .

#### B. Thermal and phonon smearing in H<sub>3</sub>S

We now estimate the impact of this spectral density smearing for  $H_3S$  using the Wien2k result for N(E). The mass renormalization effects (from the real part of the self-energy) in Migdal theory will be disregarded here, leaving our estimate as an *underestimate* of the effect of smearing. Investigation of the (Migdal) self-energy equations (i.e., in the normal state) gives the quasiparticle inverse decay rate via phonons over most of the relevant energy range, [28,29,32] for an Einstein model, as

$$\Gamma \approx \pi \lambda \Omega \left[ n_B(\Omega) + \frac{1}{2} \right]. \tag{9}$$

For H<sub>3</sub>S the characteristic frequency  $\Omega$  is in the ballpark of  $\Omega \approx 1300$  K (see above). Since we are interested only in relative values of quantities affected by smearing, we will not distinguish  $\omega_{\log}$  from  $\omega_2$ , etc.

With the choice of  $\mu^* = 0.15$ , a value of  $\lambda = 2.17$  is necessary to give the observed  $T_c = 200$  K, which also is in the range that has been quoted as resulting from DFT calculations. At 200 K, the Bose-Einstein thermal distribution  $n_B$  gives a negligible fraction of phonons excited:  $n_B(\Omega) \sim 10^{-4}$ . Thus

$$\Gamma = \frac{\pi}{2}\lambda\Omega = 5 \times 10^3 K = 0.38 \text{ eV},$$
 (10)

the halfwidth is proportional to the product  $\lambda\Omega$ . This smearing in Eq. (6) arises from the virtual excitation of phonons that provides the coupling, even at low  $T_c$  where phonons are not excited. We note that zero-point vibrations do not scatter electrons; sharp Fermi surfaces survive strong electron-phonon coupling, with de Haas-van Alphen oscillations remaining visible, and the resistivity  $\rho(T) \rightarrow 0$  at  $T \rightarrow 0$ .

Figure 8 shows three curves: the calculated (Wien2k) static lattice N(E) from Fig. 2, the thermally broadened version at 200 K, and the virtual phonon broadened DOS with halfwidth  $\Gamma = 0.38$  eV from above. The thermal broadening function is the derivative  $-df_{\circ}(E - E')/dE$  with half width of about  $\pi k_B T = 625$  K = 55 meV. This thermal broadening at 200 K is minor on the scale of interest, reducing the effective value of N(E<sub>F</sub>) slightly. The phonon broadening however is severe, with the peak value of 0.70/eV-f.u. for N(E) dropping by 37%.

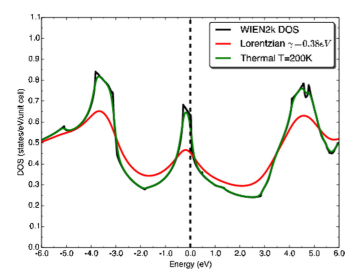


FIG. 8.  $H_3S$  density of states without broadening (black line, Wien2k DOS), with thermal broadening at 200 K (green line, which is hard to distinguish from the unbroadened one), and the virtual phonon broadened effective DOS N(*E*) using a Lorentzian halfwidth of 0.38 eV (red line). Note the large drop in the value at the Fermi energy (dashed vertical line).

The unbroadened value  $N(E_F)$  of 0.64 states/eV-f.u. is lowered by nearly 1/3, to 0.45 states/eV-f.u. The effect of the shift of the chemical potential is secondary when broadening is so large.

#### C. Implications for the theory of H<sub>3</sub>S

The Eliashberg equations including the energy dependence [28,29] of N(E) indicate that it is this 1/3 reduced value of N(E<sub>F</sub>) that should be used with the standard implementation to get a good estimate of  $\lambda$ , the superconducting gap, and  $T_c$ . The impact of experiment/theory agreement for H<sub>3</sub>S is substantial and negative: the spectral function  $\alpha^2 F(\omega)$ , proportional to  $N(E_F)$ , is reduced by 1/3 by EPC. The naive value of  $\lambda \approx 2.17$  becomes, after reducing by 1/3,  $\lambda = 1.45$ . The phonon frequency moments, which involve  $\alpha^2 F/\lambda$ , seem at this stage to remain unchanged; we return to this point below.

We have evaluated the magnitude of this phonon broadening correction on  $T_c$  using the Allen-Dynes equation [36], taking the representative values for H<sub>3</sub>S of the phonon moments to be  $\Omega = 1300$  K and  $\mu^* = 0.15$ . For  $\lambda = 2.17$ ,  $T_c = 200$  K; for the 1/3 reduced value  $\lambda = 1.45$ ,  $T_c = 130$  K. The agreement between theory and experiment is strongly degraded. It is worthy of note that H<sub>3</sub>S happens to be in a nearly linear regime of T<sub>c</sub>( $\lambda$ ), where a reduction of  $\lambda$  by 1/3 results in a decrease of  $T_c$  by nearly 1/3. We note that the corresponding strong coupling factor  $f_1$  in the Allen-Dynes parametrization is 1.13 (1.07);  $f_1$  is the crucial improvement of the Allen-Dynes equation over the McMillan equation and seems sometimes for hydrogen sulfides to have been neglected.

What this comparison implies is that theory-experiment agreement for  $T_c$  is not as good as has seemed, since taking phonon smearing into account, theory would only be predicting of the order of 2/3 of the "constant  $N(E_F)$ " value. It should be noted that this 1/3 reduction factor depends on the accurate calculation of N(E), for which we have used the Wien2k spectrum. With the FPLO result for the DOS,  $N(E_F)$  is lower and thus the effect of smearing will be smaller.

A few papers have reported calculations of the band structure and of N(E), and some studies have reported  $\lambda$ , but little attention has been given to the value of N(E<sub>F</sub>), which is sensitive to method and computational procedures. At 200 GPa (a = 5.6 a.u.) Papaconstantopoulos *et al.* quote 0.51/eV-f.u.; Duan *et al.* obtain [37] 0.41/eV-f.u.; Bianconi and Jarlborg report 0.50/eV-f.u. (their table numbers must be per cubic cell). Our Wien2k and FPLO values are 0.64 (0.42)/eV-f.u. respectively, indicating that even all-electron full potential methods can differ.

The point is that in conventional Eliashberg theory constant N(E) on the phonon scale— $\lambda$  is proportional to N(E<sub>F</sub>), and the values that have been used are sensitive to methods and cutoffs (depending on method, see above), but more seriously they are obtained from unbroadened N(E). Because of this, the reported values of  $\lambda$  and hence  $T_c$  are quantitatively uncertain, assuming they are converged BZ integrals. And on this point, McMahon and Ceperley [12] and Akashi *et al.* [14] have discussed the various challenges in reaching convergence, before even confronting the energy variation question. The (unsmeared) prediction of  $T_c \approx 200$  K indicates that improved theory, by taking into account phonon broadening, would give a substantially reduced critical temperature.

Flores-Livas *et al.* have recognized the issue of the variation of N(E) on the scale of the phonon frequencies. In their implementation of density functional theory for superconductors (DFTSC), this variation is accounted for. They reported, for their calculation of N(E) (details were not reported), taking into account the variation resulted in a 16% decrease in  $T_c$ , from 338 K to 284 K in their calculation. Their methods also involve calculation of  $\mu^*$  that is not treated in the Eliashberg form as well as other methodological differences, and these differences make direct comparison with other reports difficult. Still, the relative effects of energy variation of N(E) are clear.

This DOS variation issue extends to the calculation of phonon frequencies. The phonon self-energy involves electron-phonon scattering in which a phonon is absorbed, scattering an electron from  $E_k < E_F$  to  $E_k + \omega_q > E_F$ . Most methods of calculating phonon frequencies do not include effects of the density of available initial or final states in this energy region being variable. Thus calculation of phonon spectra will need to be re-evaluated for situations such as that imposed by H<sub>3</sub>S, and the associated nonadiabatic corrections considered.

# V. ROUTES TO HIGHER CRITICAL TEMPERATURE

The foregoing section indicates that 200 K superconductivity has been achieved with an effective DOS around  $\mathcal{N}(E_F) \approx$ 0.44/eV-f.u. compared to a peak value between the two vHs around 0.7/eV-f.u. This means as mentioned above that the theory needs refining, as already noted by Flores-Livas *et al.*, to determine just how much is understood quantitatively and what features may require more attention.

This difference in N(E) versus  $\mathcal{N}(E)$  has larger and much more positive implications. The fact that a reduced effective value of  $\mathcal{N}(E_F)$  should be used for H<sub>3</sub>S also provides the glass-half-full viewpoint: a much larger value of  $\mathcal{N}(E_F)$  and therefore  $T_c$  may be achievable in this or similar systems. Suppose that the two vHs can be moved apart, each by (say) 0.5 eV, leaving a value N(E<sub>F</sub>)  $\approx 0.7/eV$ -f.u. between. Then the Lorentzian smearing will have much less effect. There is also the question of increasing the *magnitude* of N(E) at the peak, i.e., N(E<sub>vhs</sub>). This value depends on the effective masses at the vHs, but also on the volume of the region in which the quadratic effective mass representation holds. If it holds in an ellipsoidal region defined by

$$\sum_{j} \frac{\hbar^2 k_j^2}{2m_j} < G_c^2,\tag{11}$$

the DOS from this region is

$$N(E) \propto m_{\rm th}^{3/2} \bigg[ \alpha G_c - \beta \frac{|E - E_{\rm vhs}|}{G_c} + \dots \bigg],$$
 (12)

where  $\alpha$  and  $\beta$  are numbers of order unity. The value N(E<sub>vhs</sub>) at the vHs is proportional to the thermal mass and to the radius  $G_c$  of the region of quadratic dispersion, and the decrease away from the vHs (the second term) is inversely proportional to  $G_c$ . There will be additional smooth contributions from outside this region, of course. However, increasing  $m_{th}$  and the region of

quadratic dispersion is favorable for increasing  $\mathcal{N}(E)$  in the vHs region, and hence increasing  $T_c$ . These observations seems to implicate the *topology* of the Fermi surfaces, rather than more conventional electronic structure characteristics such as relative site energies and hybridization strengths.

Numerical examples are illuminating. Suppose that the DOS peak can be widened so that  $\mathcal{N}(E_F) \approx \mathrm{N}(\mathrm{E}_F) = 0.70/\mathrm{ev}$ -f.u. as outlined just above, rather than the reduced effective value of 0.45 eV/f.u. that gives, experimentally,  $T_c = 200$  K. With  $\Omega = 1300$  K and  $\mu^* = 0.15$  as above,  $\lambda_{\mathrm{exp}} = 2.17$  is required to account for  $T_c = 200$  K. For a 0.70/0.44 = 1.55 larger value of N(E\_F),  $\lambda = 3.38$  and we find  $T_c = 277$  K—room temperature in a cool room. The increase in effective  $\mathcal{N}(E_F)$  we have assumed is ambitious but not outlandish, given the calculated spectrum of  $Im\bar{3}m$  H<sub>3</sub>S. It is clearly worthwhile to explore other H-rich compounds for higher critical temperatures.

Of course, the increase in  $\mathcal{N}(E_F)$  will give additional renormalization (softening) of the phonons. However, the modes are very stiff even with  $\lambda = 2.1$ , so this may not be a major effect. Note that decreasing  $\Omega$  increases  $\lambda$  but decreases the energy scale prefactor in  $T_c$ , one reason why increasing  $\lambda$  by decreasing frequencies is rarely a profitable means of increasing  $T_c$ . If  $\Omega$ , the prefactor in the Allen-Dynes  $T_c$  equation, is softened by 10% without change in matrix elements,  $\lambda$  increases by 20% while  $T_c$  increases by only 4%. Evidently softening of hard phonons is a minor issue when looking for higher  $T_c$  in this range of  $\lambda$ . This behavior was formalized by Allen and Dynes [36], who obtained the rigorous strong coupling limit

$$T_c \to 0.18 \sqrt{\lambda \langle \omega^2 \rangle} = 0.18 \sqrt{N(E_F) \langle I^2 \rangle / M},$$
 (13)

where  $\omega$  is expressed in kelvins. The last expression is strictly true only for an elemental superconductor, think of the electron-ion matrix element *I* and mass *M* as those of H for H<sub>3</sub>S.

#### VI. SUMMARY

In this paper we first addressed the electronic structure and especially the delicate van Hove singularity induced spectrum, bonding characteristics and effect of S, and the charge density of states near the Fermi level. The occurrence of two closely spaced van Hove singularities is definitely a central issue for the properties of  $H_3S$ . We list some of the main points.

(i) At the most basic level, why is  $H_3S$  superconducting at 200 K? It is because both  $\lambda$  is large but, more importantly, the characteristic phonon frequency  $\Omega$  is very high. This reminds one of the Allen-Dynes limit for strong coupling,

$$T_c \to 0.18\sqrt{\lambda\Omega^2} \to 0.18\sqrt{N(E_F)\langle I^2 \rangle/M}.$$
 (14)

Though not yet in this limit, this provides the right picture one can check that keeping all fixed except for  $\Omega$  and then varying it, the change in  $T_c$  is minor because the change in prefactor  $T_c \propto \Omega$  is compensated by  $\lambda \propto \Omega^{-2}$ .

(ii) Sulfur 3*p* states hybridizing with hydrogen 1*s* is crucial in producing the strong large scale structure in N(E) within  $\pm 5$  eV of the Fermi level, and in leaving E<sub>F</sub> at the top of a peculiarly sharp peak between two vHs. The van Hove points on the constant energy surfaces that define the peak in N(E) were identified, finding they lie on opposite ends of a line of Fermi surface "ripping apart" with energy varying between the two van Hove singularities. This region of very low velocity electrons affects a significant fraction of the zone. It is unclear how replacing S with other elements will affect the electronic structure near  $E_F$ , but small changes may have large effects. Ge *et al.* have noted that alloying 7–10% of P with S moves  $E_F$  to the peak in N(E), within the virtual crystal approximation which does not account for alloy disorder broadening. In any case, strong coupling smearing as discussed here will nullify this apparent gain.

(iii) The fine structure and energy variation of N(E) near the Fermi level must be taken into account to obtain quantitative results for  $\alpha^2 F$ ,  $\lambda$ , and  $T_c$ . The energy dependence of N(E) may even affect calculation of phonon frequencies, though this is untested so far.

(iv) The closely spaced van Hove singularities very near the Fermi level have been shown to have significance, both on the detailed theory of H<sub>3</sub>S but, as importantly, on the question of whether  $T_c$  can be increased in related materials. Sulfur and the specific  $Im\bar{3}m$  structure are important for high  $T_c$  though other elements will need to be studied to learn more about precisely why.

(v) The prospect for increased  $T_c$  is affirmative—it will require only evolutionary changes of the electronic structure to achieve room temperature superconductivity, though the road to this goal requires study, and additional insight into the origins of van Hove singularities may be important. Increasing the vHs effective masses, or increasing the volume within

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which quadratic dispersion holds, will increase N(E) at the vHs energy. Structural or chemical changes that affect the electronic structure rather modestly may lead to significant increase in the effective (broadened) density of states at  $E_F$ . Other studies have suggested that substitution of some sulfur with chemically related elements may increase  $T_c$ . Altogether, the prospects of achieving increased critical temperatures are encouraging.

(vi) An issue that is almost untouched is a deeper understanding, or rather an understanding at all, of electron-ion matrix elements  $\langle I_H^2 \rangle$ —what contributes to strong electron-H atom scattering, and what degrades this scattering. These matrix elements are the same that determine resistivity in the normal state; notably most of the best superconductors have high resistivities. Further study should address the EPC matrix elements.

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