Room Temperature Superconductivity: 
the Roles of Theory and Materials Design

Warren E. Pickett

Department of Physics and Astronomy,
University of California Davis,
Davis, California 95616

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For half a century after the discovery of superconductivity, materials exploration for better superconductors proceeded without knowledge of the underlying mechanism. The 1957 BCS theory cleared that up: the superconducting state occurs due to strong correlation in the electronic system: pairing of electrons over the Fermi surface. Over the following half century higher critical temperature $T_c$ was achieved only serendipitously as new materials were synthesized. Meanwhile the formal theory of phonon-coupled superconductivity at the material-dependent level became progressively more highly developed: by 2000, given a known compound, its value of $T_c$, the corresponding superconducting gap function, and several other properties of the superconducting state became available independent of further experimental input. In this century, density functional theory based computational materials design has progressed to a predictive level – new materials can be predicted from free energy functionals on the basis of various numerical algorithms. Taken together, these capabilities enable theoretical prediction of new superconductors, justified by application to superconductors ranging from very weak to quite strong coupling. Limitations of the current procedures are discussed briefly; most of them can be handled with additional procedures. Here we recount the process that has resulted in the three new highest temperature superconductors, with compressed structures predicted computationally and values of $T_c$ obtained numerically, that have been subsequently confirmed experimentally: the designed superconductors SH$_3$, LaH$_{10}$, YH$_9$. These hydrides have $T_c$ in the 200-260K range at megabar pressures; the experimental results and confirmations are discussed. While the small mass of hydrogen provides the anticipated strong coupling at high frequency, it is shown that it also enables identification of the atom-specific contributions to coupling, in the manner that has been possible previously only for elemental superconductors. The challenge is posed: that progress in understanding of higher $T_c$ is limited by the lack of understanding of screening of H displacements. Ongoing activities are mentioned and current challenges are suggested, together with regularities that are observed in compressed hydrides that may be useful to guide further exploration.
2. An alternative paradigm: activating bonding states

D. Increasing accessibility; metastable structures
1. Lowering the required pressure
2. More complex hydrides; speeding searches
3. Exploring higher pressures

VII. Regularities in Compressed Hydrides

VIII. Epilogue

IX. Acknowledgments

References

I. PROLOGUE

Room temperature superconductivity (RTS) has been one of the grand challenges of condensed matter physics since the BCS theory of pairing (see Sec. II A) was proposed and its predictions verified. The remarkable electronic and magnetic properties of the superconducting state readily suggest revolutionary applications, both in the laboratory and in the public sector. The slow progress in increase in the maximum critical temperature $T_c$ from 4K in 1911 to 23K in 1973 – 3K/decade – followed by a 15 year plateau, moved this grand challenge well into the background. The discovery of high $T_c$ cuprates (up to 134K, or 164K under pressure [Gao et al. 1994]), involving magnetic interactions, introduced a new type of superconductivity (SC) and generated renewed excitement. After eight years of advancement of $T_c$, another plateau in cuprate superconducting $T_c$ has lasted for (so far) 28 years. This recent advancement of the maximum $T_c$, revealing a breakthrough increase toward room temperature superconductivity that prompted this article, is shown in the upper right corner of Fig. 1. After preliminary information, in Secs. V and VI these advancements and some of their microscopic origins will be discussed.

Much attention has centered on guidelines (‘rules,’ or ‘roadmaps’) necessary for high $T_c$, thereby presumably pointing the way to higher $T_c$. Such rules have been based primarily on known superconductors, and have subsequently been set aside as entirely new classes of superconductors were discovered, almost entirely by serendipity. In the 1960s empirical trends led to “Matthias’s Rules.” These stated that for high $T_c$ one should search for (1) cubic materials, (2) $d$ electrons at the Fermi level, and (3) specific electron/atom ratios. This latter rule was soon understood to mean a high density of electron states at the Fermi level ($\varepsilon_F=0$) $N(0)$, i.e. a high density of superconducting pairs. Bernd Matthias’ group had been the leader in the discovery of new superconductors in the 1950s and early 1960s, after which the search extended worldwide.

Though unwritten, additional rules were advertised: (4) stay away from oxygen, which produces unpredictable behavior including insulation, and (5) stay away from magnetism, which at the time competed too strongly with superconducting pairing. Matthias had an additional personal rule: (6) stay away from theorists, they are no help. (Matthias did write [Matthias 1972], without elaboration ‘I never realized how many of my friends are theorists.’) This last rule was based on his observation that knowing the theory of SC, at least in a broad model BCS way, had been useless in helping discovery; what worked (but ploddingly so) was just to “follow the simple roadmap.” However, the rules did not produce high $T_c$ materials. It continued to be true through cuprate days – after the initial high $T_c$ – that theory played no part in the advance of the maximum $T_c$ from 30K to 134K in cuprates.

Over the intervening decades theory and numerical implementation have advanced. While overt activity toward higher BCS superconductivity waned, intellectual interest in the goal of RTS persisted, with evidence given by various international workshops indicating continued emphasis on room temperature explorations [1]. With the 2014-2015 discovery of extreme high $T_c$ SC in compressed metal hydrides under pressure discussed in this paper, the roles of experiment and theory evolved and were reversed. Theory has assumed a prominent role as predictor beyond the maximum known $T_c$ ($T_c^{max}$) for phonon-coupled SCs as they jumped from 40K to 200K, then rapidly marched toward room temperature.

After recounting the sequence of necessary theoretical advances in Sec. II, Sec. III gives a brief indication of the computational innovation and implementations that were required to design real, heretofore unknown, materials. An overview of the rising research area in this century of crystal structure prediction is given in Sec. IV. Section V provides a concise description of the first three revolutionary discoveries of critical temperatures in the 200-260K range. So far these (and a few others) all require megabar pressures. The discussion in Sec. VI reveals how the light H atom restores the sort of analysis that was developed for elemental metals, which leads to demonstration that H dominates the metal component in promoting very high $T_c$; in fact, the contribution of the

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1 This list provides six of the meetings.

- 1994 The Road to Room Temperature Superconductivity, Bodega Bay, California, 1992.
- 2005 The Possibility of Room Temperature Superconductivity, Notre Dame, June 10-11, 2005. https://www3.nd.edu/~rts/index.html
- 2014 (Toward) Room Temperature Superconductivity, Leiden, The Netherlands, 2014.
- 2016 SUPERHYDRIDES Towards Room Temperature Superconductivity, Hydrides and More, Rome, Italy, 2016.
- 2017 Towards Room Temperature Superconductivity: Hydrides and More. Chapman College, California, 2017.
- 2022 Challenges in Designing Room Temperature Superconductors. L’Aquila, Italy, 2022.
metal atom provides confusion by contributing in opposing ways to the properties that promote high $T_c$. This leads to Sec. VI, which gives an overview of near-term challenges and opportunities that have been identified in the very high $T_c$ arena. These are provided in terms of several regularities of high $T_c$ compressed hydrides that may guide the next level of searches, as compiled in Sec. VII. Section VIII provides a brief Epilogue.

II. THEORETICAL DEVELOPMENTS THROUGH THE DECADES

The impetus for this stunning breakthrough has been a sequence of advances in theory, numerical implementation, and computational design together with mastery of high pressure techniques. Since this advance in computational theory has extended over six decades (i.e. three generations of physicists), we provide in this section an overview of the fertile path of breakthroughs in theory and numerical implementation that have enabled current capabilities and their paradigm-revising results.

A. BCS Theory

In 1957 Bardeen, Cooper, and Schrieffer published their 30-page *magnus opus* “Theory of Superconductivity,” (Bardeen et al., 1957) that includes around 275 numbered equations. This theory introduced a correlated many-body wavefunction based on Fermi surface pairing of electrons demonstrated a year earlier by Cooper. This wavefunction described a thermodynamic condensate of correlated pairs below a critical temperature $T_c$, assuming an attractive effective pairing potential arising from exchange of phonons and screened electron-electron scattering. The primary result of their theory was the $T_c$ equation (units $\hbar = 1$, $k_B = 1$ will be used in this article)

$$T_{c}^{BCS} = 1.14 \ \Omega \ e^{-1/\lambda^*}$$

(1)

where $\Omega$ is the characteristic phonon frequency. $\lambda^* = N(0)V - \lambda - \mu^*$ in terms of terminology developed later (see below). The Fermi level density of states $N(0)$ is a measure of the density of electrons interacting through the phenomenological coupling $V$, which BCS noted is the *net interaction* of phonon attraction plus Coulomb repulsion.

The electron-phonon coupling strength $\lambda$ is the measure of the attractive pair coupling strength by exchange
of phonons. Out of interest for the following content of this article we note that the strong coupling limit $\lambda^* \to \infty$ leads in this expression to a maximum SC temperature of 1.14 $\Omega$, which can be well above room temperature. The theory is however only valid for weak coupling, leaving open the question of high $T_c$ regime. “High $T_c$” would only be confronted three decades later with the discovery of cuprate superconductors, for which there is no predictive theory and hereafter decades later with the discovery of cuprate superconductors, for which there is no predictive theory and hereafter not be considered.

The second property of note, provided by BCS theory and its extensions, is the superconducting gap $[\Delta(\omega, T)]$ equation that gives $T_c$ when linearized ($\Delta \to 0$) and at finite temperature provides the frequency $\omega$ and temperature $T$ dependence of the energy gap that is responsible for many of the fascinating properties of superconductors. The novelty is that a perfect conductor can have a gap for excitations; this gap leads to a vanishing magnetic susceptibility in its interior; it is a basic player in a gap for excitations; this gap leads to a vanishing magnetic susceptibility in its interior; it is a basic player in the energy gap, in excellent agreement with experimental data.

FIG. 2 Figure from the BCS paper [Bardeen et al. (1957)] paper illustrating the predicted temperature dependence of the ratio of the specific heat in the superconducting state to that in the normal state value (equal to 1.0 on the abscissa). The behavior is exponentially small at low temperature due to the energy gap, in excellent agreement with experimental data.

B. Migdal-Eliashberg Theory

The microscopic Migdal-Eliashberg theory of 1960 (henceforward called Eliashberg theory) of electron-phonon (EP) coupled superconductivity, weak or strong, followed very quickly after the model BCS theory (1957) based on a model pairing Hamiltonian and a variational treatment. Recall that the full, exact (non-relativistic) Hamiltonian $H$ for a system of ions $\{\vec{R}\}$ and electrons $\{\vec{r}\}$ is given by

$$H = [T^{el}(\{\vec{r}\}) + V^{el-el}(\{\vec{r}\})] + V^{el-ion}(\{\vec{r}\}, \{\vec{R}\}) + T^{ion}(\{\vec{R}\}) + \gamma^{ion-ion}(\{\vec{R}\})$$ (2)

in terms of the various kinetic $T$ and potential $V$ energies. The middle term contains the (bare) electron-ion interaction. With roughly $10^{20}$ dynamical coordinates per mm$^3$, this all-encompassing operator is the fundamental, intractable feature of materials theory that requires innovative approaches and reliable approximations.

Migdal formalized the observation [Migdal 1960] that the great differences in energy scales (or velocities, or masses) of electrons and ions leads to negligible contributions beyond second order perturbation theory in EP scattering. Eliashberg made the formidable step of placing the pairing theory within the newly developed many-body theory (thermal Green’s function) that made it applicable to the superconducting state, [Eliashberg 1960 1961] leaving generalizations to materials with crystal structure and lattice effects as a later step (see below). This Eliashberg formalism provided the fundamental equations for calculating the gap function (superconducting order parameter), given the Eliashberg spectral function $\alpha^2 F(\omega)$ defined in Sec. IIE that provides the essential input for the pairing process. Calculation of this function necessitated several theoretical and algorithmic advances.

C. Density Functional Theory (DFT)

Hohenberg, Kohn, and Sham followed immediately (1965-1967) with density functional theory, [Hohenberg and Kohn 1964; Kohn and Sham 1965] which makes the full crystalline Hamiltonian in Eq. 2 treatable (for static nuclei) for many properties of materials for electrons in their ground state, given a reasonable approximation for many body effects (through an exchange and correlation functional). (The many following extensions of DFT to other properties are not relevant here, except as noted.)

An extremely important aspect of DFT is that it provides a highly reliable one-electron (‘mean field’) set of Kohn-Sham one-electron band energies and wavefunctions (Kohn and Sham 1965) for use in themselves, and also for applications in treating dynamic behavior that
is averaged over in DFT. This DFT breakthrough accelerated activity in band theory, already in progress since the late 1930s based on more phenomenological grounds. The late 1970s saw the accomplishment of achieving self-consistent electronic charge densities, bands, and wavefunctions. Many applications of (electronic) ground state energies were explored in the 1980s and following.

Calculation of energies for any configuration of atoms enabled the evaluation of interatomic force constants between displaced atoms, and thereby phonon spectra. Initially this was accomplished for each phonon independently, but capabilities were rapidly extended, especially by applying density functional perturbation theory and Wannier function techniques. (Cockayne, 2005; Mostofian et al., 2008; Rabe and Wagmare, 1995). The outcome was true harmonic phonon frequencies through formalism making use of infinitesimal atomic displacements. The change in the electronic potential due to an atomic displacement is also the root factor in EP coupling, and calculation of EP matrix elements was achieved only around 2000.

D. Extension of Eliashberg Theory to Real Materials

Also in the mid-1960s, the challenge of addressing real superconductors versus simplified models, a prerequisite for materials design and discovery, was accomplished by Scalapino et al. in 1964. (Scalapino, 1969; Scalapino et al., 1966). Starting from the full Hamiltonian of a solid (Eq. 3) they derived a remarkably complete formalism for the superconducting Green’s function and the full frequency and temperature dependence of the complex gap function. Their formalism awaited a viable description of the underlying electronic energy bands and phonon frequencies, and their coupling. The DFT capabilities discussed in Sec. II.C would provide the underlying electronic bands and wavefunctions, phonon dispersion curves and polarizations, and EP matrix elements.

The validity of their formalism, which underlies today’s numerical implementation, relies on a few approximations. One is Migdal’s theorem mentioned above: the vast differences in masses of electrons and ions (more precisely, differences in the frequencies of their dynamic responses) specifies that second order perturbation theory is sufficient for the electron and phonon self-energies. Second, electron-electron Coulomb repulsion effects leave the metal as a conventional Fermi liquid; specifically, the possibility of magnetic order and magnetic fluctuations is not included at the level we discuss here.

Thirdly, the symmetry that is broken at the superconducting phase transition is so-called gauge symmetry; in picturesque language, two electrons can form a Cooper pair and disappear into the condensate, or the inverse process can happen. In this language, electron number conservation is broken. [C. N. Yang liked to emphasize that electrons do not actually disappear, and that the fundamental signature of the superconducting state is appearance of long-range order in the two-particle density matrix.] Finally, for EP-coupled superconductors it is nearly always the case that the superconducting order parameter is proportional to the gap function, i.e. a complex scalar and not a vector or tensor quantity, and other symmetries are not broken at $T_c$. The resulting theory has passed numerous tests as being accurate for EP-coupled superconductors.

E. Analysis of $T_c$; discussion of a maximum

1. McMillan’s analysis

In 1967 McMillan advanced the advanced analysis of the origins of $T_c$ substantially. (McMillan, 1967) He defined the electron-phonon spectral function $\alpha^2 F$, also called the Eliashberg function, in terms of the various quantities appearing in the Scalapino et al. expressions:

$$\alpha^2 F(\omega) = N_f(0) \sum_{kk'} |M_{kk'}|^2 \delta(\omega - \omega_{kk'}) \delta(\varepsilon_k) \delta(\varepsilon_{k'}),$$

where the Fermi energy $\varepsilon_F = 0$, and $kk'$ is the wavenumber of the phonon scattering an electron from state $k$ to $k'$, each confined to the Fermi surface. Necessary sums over bands and phonon branches are not displayed explicitly. The sums that are shown are over the three dimensional Brillouin zone, each confined by the pair of $\delta$-functions to the Fermi surface. The frequency $\delta$-function provides the frequency resolution of coupling. For clarity, the density of states factor is for a single spin, designated by the arrow on $N_f(0)$.

The EP coupling strength $\lambda$, and the two frequency moments $\omega_{2\log}$ and $\omega_2$ used prominently by Allen and Dynes (see the next subsection) are given in terms of moments of $2\alpha^2 F(\omega)/\omega$ by

$$\lambda = \int d\omega \frac{2\alpha^2 F(\omega)}{\omega},$$

$$\omega_2^2 = \int d\omega \omega^2 \frac{2\alpha^2 F(\omega)}{\lambda} = \int d\omega \omega^2 \log g(\omega),$$

$$\omega_{2\log} = \exp \left[ \int d\omega \log \omega g(\omega) \right].$$

Here $g(\omega)$, defined by the second line, is the normalized shape function of $\alpha^2 F\omega/\omega$. Aside from normalization, these are all different moments of $2\alpha^2 F(\omega)/\omega$. The calculation of $\alpha^2 F$, necessary for the computational program to advance, is discussed further in Sec. III.B.

For elemental metals, McMillan obtained initially two fundamental expressions from $\alpha^2 F$. First is that its first frequency moment is independent of vibrational proper-
ties aside from an overall inverse mass factor:

$$\int \omega^2 F(\omega) d\omega = \frac{N(0) I^2}{2M}, \quad (7)$$

where $I^2$ is the average atomic scattering strength by phonons from the Fermi surface to the Fermi surface:

$$I_{k,k'} = k|dV(r; \{R\})/dR|_{k'} >, \quad (8)$$

$$M_{k,k'} = \frac{h}{2M \omega_{k-k'}} I_{k,k'}, \quad (8)$$

$$I^2 = \langle |I_{k,k'}|^2 >_{FS}. \quad (8)$$

Here $V(r; \{R\})$ is the total electronic potential which depends on all ionic coordinates $\{R\}, R_i$ is the coordinate of the displaced atom under consideration, and $I_{k,k'}$ is the electron-ion matrix element between electron states $k$ and $k'$. $M_{k,k'}$ is the electron-phonon matrix element that contains the atom mass $M$ and phonon frequency $\omega_{k-k'}$, and the outside brackets indicate that an average is taken over all $k$ and $k'$ both on the Fermi surface to obtain $I^2$. This first-order change in potential $dV/dR$ has a long history (Pickett, 1979) and much progress in numerical evaluation, without much progress in understanding. This integral expression holds only for elements with mass $M$, a restriction we manage to relax for hydrides in Sec. VII.B.

McMillan’s second observation was that (again for elemental metals) $\lambda$ can be expressed simply in terms of physical quantities

$$\lambda = \frac{N(0) I^2}{M \omega^2} = \eta \kappa, \quad (9)$$

The numerator $\eta = N(0)I^2$ is the McMillan-Hopfield factor that reflects an “electronic stiffness,” divided by $\kappa = M \omega^2$ which is the standard harmonic oscillator form of lattice stiffness (mean force constant), with $\omega$ representing the entire spectrum of frequencies. All are normal state properties. This relation indicates that increasing the frequency scale $\omega$, which helps $T_c$, will decrease $\lambda$ according to its square, hurting $T_c$, with net outcome yet to be understood.

An aside, for now. The simple expression Eq. 9 does not apply to compounds, as the quantities $N(0)$ and $\omega$ have contributions from all atoms. and $I^2$ and $M$ are distinct atomic quantities. It will be seen in Sec. VI.B that this simple ‘elemental’ expression for $\lambda$ can be extended to compressed metal hydrides to obtain these quantities separately for H and the metal atom, thereby allowing deeper analysis into the origins of high $T_c$.

Most prominently McMillan provided, with justification for chosen algebraic forms, a seminal analysis (McMillan, 1967) from Eliashberg theory of $T_c$ and its dependence on $\alpha^2 F(\omega)$, presenting his iconic “McMillan equation” expressing $T_c$ via two materials quantities, the phonon (Debye) frequency $\theta$ and $\lambda$. A phenomenological retarded Coulomb repulsion $\mu^* = 0.13$ was included in the Eliashberg equation that was solved to relate $T_c$ to $\alpha^2 F$. Solutions for various $\alpha^2 F$ functions were fit to a generalization of the BCS equation of the form

$$T_c^{McM} = \frac{\theta}{1.45} \exp \left[-1.04 \frac{1 + \lambda}{\lambda - \mu^* - 0.62 \lambda \mu^*} \right]$$

$$= \frac{\theta}{1.45} \exp \left[-\frac{1 + \lambda}{\lambda - \mu^*_{eff}} \right]^{1.04} \quad (10)$$

where the second expression emphasizes the generalization of the BCS relation. The constants provided best fits to the computational data: the ‘exponent’ 1.04 seems clearly to be a parameter giving best fit without any physical significance. Note that the McMillan equation is not appropriate for very weak coupling; when the denominator of the argument of the $exp$ function becomes negative, the result is non-sensical. What the McMillan equation does is to replace the very involved functional dependence $T_c(\alpha^2 F, \mu^*)$ by a simple equation for $T_c$ involving two quantities from $\alpha^2 F$ plus $\mu^*$.

The “effective Coulomb repulsion” $\mu^*_{eff} = \mu^*(1 + 0.62\lambda)$ is defined here (without physical justification) to emphasize the similarity to the BCS equation (1). The $1 + \lambda$ factor in the numerator is often assigned to the strong-coupling electron mass renormalization by EP coupling. The mass enhancement is well known to increase the density of active states at the Fermi level by $1 + \lambda$, but in this expression the effect in the numerator is to decrease $T_c$, compensated somewhat by the $\lambda \mu^*$ term in the denominator. Flores-Livas et al. have reproduced the model that gives a similar result, for $\mu^*(\lambda = 0)$ and a different expression for the prefactor. (Flores-Livas et al., 2020)
In McMillan’s time the experimental Debye frequency \( \theta \) for elemental metals was the most accessible measure of the frequency spectrum, however McMillan recognized that using moments from \( \alpha^2F \) (such as the second moment \( \omega_2 \)) would be more representative.

Only values of \( \lambda \) to around unity were included in McMillan’s fit; extrapolation of \( T_c^{\text{McM}} \) shows that it saturates for large \( \lambda \), however the equation is justified only for moderately strongly coupled superconductors. The limited range of input has been missed or forgotten by many readers, and until the discoveries discussed in the next section one could still find statements in the literature and in presentations that “theory shows that the maximum \( T_c \) is limited, perhaps to around 40 K.” This point will be clarified below.

2. Bergmann and Rainer’s functional derivative

In 1973 Bergmann and Rainer [Bergmann and Rainer, 1973] raised a straightforward question: to increase \( T_c \), one needs to understand the effect on \( T_c \) due to an added increment of coupling \( \Delta\alpha^2F(\omega) \) at frequency \( \omega \).

It seemed clear from the BCS equation that adding coupling at high frequency, seemingly increasing both \( \omega \) (or \( \Omega \), or \( \theta \), depending on treatment of the phonon system) and \( \lambda \) (adding an increment), would raise \( T_c \). Study by Allen [Allen, 1973, 1974] soon after McMillan gave suggestions that, due to this complex interdependence of materials properties, including \( N(0) \) and \( T^2 \), coupling at low frequency might even be harmful for \( T_c \). We will see in Sec. VLB that compressed hydrides provide an unexpected field of study of this unsettled question.

Bergmann and Rainer addressed the question by calculating the differential quantity in the kernel of

\[
\Delta T_c = \int d\omega \frac{\delta T_c[\alpha^2F, \mu^*]}{\delta \alpha^2F(\omega)} \Delta \alpha^2F(\omega),
\]

(11)

this being the functional derivative of \( T_c \) with respect to \( \alpha^2F(\omega) \): given an increase in coupling by an increment \( \Delta \alpha^2F \) at frequency \( \omega \), what is the change \( \Delta T_c \)?

They established that this functional derivative is smooth (\( \alpha^2F \) is not) and non-negative, thus every phonon helps, as anticipated. At least for conventional shapes and strengths of \( \alpha^2F \) it has a broad peak around \( \omega_{BR} \sim 2\pi T_c \) followed by a slow decrease at high frequency, shown in Fig. 4. Conversely, below \( \omega_{BR} \), the usefulness of coupling decreases linearly to zero. Low frequencies can and do contribute strongly to \( \lambda \); viz. the corresponding functional derivative is \( \delta \lambda/\delta \alpha^2F(\omega) = \omega/2\omega \).

This result, viz. that coupling at high frequencies is most important, does not quite resolve all issues. While adding some differential coupling \( \Delta \alpha^2F(\omega) \) does affect both \( \lambda \) and \( \omega \) calculated from \( \alpha^2F \), within the larger picture it also will affect the EP coupling that causes

changes in \( \alpha^2F \) at frequencies besides \( \omega \). This effect is not settled by simply including a single (extrinsic) increment in \( \alpha^2F \). It will be seen in Sec. VLB that hydrides provide counterexamples to the simple interpretation offered by the functional derivative of Bergmann and Rainer.

3. Allen and Dynes’ reanalysis

Building on the work of McMillan, Allen and Dynes in 1975 presented a re-analysis [Allen and Dynes, 1975] of \( T_c \), using more than two hundred solutions of the Eliashberg equation for \( \lambda \) up to 10 and \( \mu^* \) up to 0.20, including experimental determinations of \( \alpha^2F \) (by numerical inversions of tunneling data) with associated measured \( T_c \). Allen and Dynes chose to generalize the McMillan equation. Most obvious is that the frequency prefactor is updated from \( \omega_2 \) (altered earlier from McMillan’s \( \theta \)) to the logarithmic moment \( \omega_{log} \) with an adjusted constant, as the latter produced a more consistent fit to their data.

More importantly, they observed that corrections were needed for both the strength of coupling – they considered much larger values of \( \lambda \) than previously studied – and for and unusual shapes of the \( \alpha^2F/\omega \) spectrum. For example, the textbook shape of the Nb phonon spectrum is much different from that of (say) PdH, which consists of low frequency Pd acoustic modes separated from high frequency H optic modes. We return to the implications of this two-peak shape in Sec. VI.B.

The shape dependence was framed by Allen and Dynes in terms of the ratio \( \omega_2/\omega_{log} \), which is always greater than unity. In their fit to extensive data, the argument
of the exponent was not changed, rather each change (strong coupling and shape) was incorporated into its own prefactor $f_1$ and $f_2$ respectively. The Allen-Dynes equation can be written, considering the McMillan form given above,

$$T_{c}^{AD} = \frac{\omega_{log} 1.45}{1.20} f_1(\lambda, \mu^*) f_2(\omega_{log}, \omega_2) T_{c}^{McM}(\lambda, \mu^*)$$

$$= \frac{\omega_{log}}{1.20} f_1 f_2 \exp \left[-\frac{1 + \lambda}{\lambda - \mu^*} \right]^{1.04}$$

$$= \frac{\omega_{log}}{1.20} f_1 f_2 \exp \left[-1.04 \frac{1 + \lambda}{\lambda - \mu^* - 0.62\lambda\mu^*} \right], \quad (12)$$

the latter form being the more common one and the one presented by Allen and Dynes. The simple expressions for $f_1$ and $f_2$ can be found in the original paper. [Allen and Dynes 1975] Note that $\theta$ cancels in the first expression, and is replaced by the logarithmic frequency moment, which Allen and Dynes found to be the most useful moment for the prefactor of $T_c$.

Due to the $f_1$ and $f_2$ factors, $T_{c}^{AD}$ is no longer exponential in the argument involving $\lambda$ except in the weak-coupling region where $\lambda$ begins to approach $\mu^*$. In this regime a solution of the Eliashberg equation, as well as $T_{c}^{AD}$, is highly sensitive to each quantity. For this reason, their equation is not a meaningful fit to any data in this weak coupling region. For the high $T_c$ compressed hydrides discussed in Sec. V, each prefactor can approach a 10-20% enhancement of $T_c$ (or increasingly more as $\lambda$ is increased still further).

A crucial finding of Allen and Dynes was that in the large $\lambda$ regime, where analytic results could be extracted, the asymptotic behavior is

$$T_c \rightarrow 0.18 \sqrt{\lambda\omega^2} = 0.18\omega_2\sqrt{\lambda} = 0.18\sqrt{\eta/M} \quad (13)$$

for $\mu^* = 0.10$ (the prefactor depends somewhat on $\mu^*$). Here $\lambda = \eta/(M\omega^2)$ for elements has been used to display different viewpoints. Note that the asymptotic limit involves only the same frequency independent constant identified by McMillan. A primary implication is that

FIG. 5 (a) Allen and Dynes’ plots of the ratio $T_c/\langle \omega \rangle$ versus $\lambda$ for the McMillan equation, and for truncated and exact numerical results from the Eliashberg equation. (b) Similar plots (also from Allen and Dynes (1975)) using data for several known strong coupled superconductors. Also shown is the exact solution for an Einstein model with frequency $\langle \omega \rangle = \Omega$, with the deviation from the McMillan equation beginning at $\lambda \sim 1$. Here $\langle \omega \rangle$ is the conventional first moment of $\alpha^2 F(\omega)/\omega$. Figures are from Allen and Dynes (1975) with permission.
Eliashberg theory poses no limit on $T_c$. Figure 5(a) shows some of the numerical data that establishes the ever-increasing (not plateauing) of $T_c(\lambda)$, and panel (b) of this figure provides numerical data using experimental shapes of $\alpha^2 F$.

An informative observation is that, as far as the limiting value goes, neither $\lambda$ nor $\omega_2$ are separately relevant; physically, increasing $\lambda$ decreases $\omega_2$ and there are numerous examples of this. It should be recognized that performing this limit assumes nothing else happens except that $\lambda$ increases. This procedure is non-physical; making changes that increase $\lambda$ in turn decrease frequencies due to the increased coupling from electronic states at higher energy. Increasing $\lambda$ often creates lattice instabilities, so the more strongly coupled material cannot exist. On the positive side, nothing prohibits finding superconductors with increasingly larger $\lambda$ with appropriate frequencies and higher $T_c$. Further analysis of $T_c$ can be found in extended descriptions by Allen (Allen, 1979, 1980) and Allen and Mitrović. (Allen and Mitrović, 1982)

F. Solidifying the formal theory

1. General remarks

Eliashberg theory providing $T_c$, the frequency and temperature dependent gap $\Delta(\omega, T)$, and more, as described above and as commonly applied, is based on the common but somewhat ad hoc choices of (i) using the DFT mean-field eigenvalues and eigenfunctions as including the essential electronic interaction effects in a static lattice (this choice has no rivals), (ii) from that, the self-consistently determined phonon spectrum including static electronic screening, as obtained from DFT, and (iii) the simple bubble diagrams giving the EP coupling contribution to the electron and phonon self-energies (always consistent with Migdal’s theorem). The Hamiltonian then consists of band (bare) electrons (in the static lattice), bare harmonic phonons oscillating in the frozen electron density which are non-physical and ill-defined), and (iv) the bare electron – bare phonon coupling due to first-order change in the lattice potential due to atomic motion. This step-by-step ansatz, though seemingly first principles, leaves questions about the full self-consistent treatment of the Coulomb interaction.

A general formulation of the electron-phonon problem based on the full crystal Hamiltonian, provided by Allen, Cohen, and Penn. (Allen et al., 1988) reveals that the combined electron + lattice polarization (and resulting dynamic dielectric screening function) is the central quantity to be addressed by many-body theory. While treating the polarizations separately has become intuitive and works well for conventional metals, a full treatment might reveal novel processes and possibly new phases of matter in complex materials.

A rigorous underlying formalism for a complex problem often guides progress even when rigor must be relaxed. DFT provides an excellent example. DFT gives a rigorous foundation for treating the full crystal Hamiltonian in Eq. 3 with the restriction to static nuclei. Beyond the ground state energy and magnetic properties, it is established that the Kohn-Sham band structures are reasonable to excellent representations of single particle excitations in conventional (Fermi liquid) metals, as soon as a sufficiently sophisticated exchange-correlation functional is incorporated. This characterization is especially relevant near the Fermi surface of metals, where necessary corrections can be incorporated when desired.

2. Superconducting density functional theory

Such a formal underpinning for superconductors was devised by Gross and a sequence of collaborators. For this article we emphasize the density functional theory for superconductors (SCDFT) formulated for material
specific studies, which was achieving implementation by 2005. The formulation includes a number of innovations, with the basic ones being the extension of ground state DFT to thermodynamic DFT, followed by generalization to description of quantum nuclear degrees of freedom, then incorporating an allowance for a superconducting order parameter.

Accounting for superconductivity involves a functional $F[n, \chi, \Gamma]$ of the electron density $n(\mathbf{r})$, the superconducting order parameter $\chi(\mathbf{r}, \mathbf{r}')$, and the diagonal of the nuclear density matrix $\Gamma(\{R\})$. The resulting three Euler-Lagrange equations involve one for the nuclear coordinates, of generalized Born-Oppenheimer type, and two Kohn-Sham Bogoliubov-de Gennes-like equations for the electronic states. Each of the equations involves its own exchange-correlation potential that requires approximation.

In the standard approximation invoking Migdal’s theorem the first equation describes fully dressed harmonic phonons. The formalism is extended to non-zero temperature thermal equilibrium, finally providing properties of the superconductor below $T_c$. Above the calculated $T_c$, the order parameter and corresponding potential vanish, and the solution reverts to the usual DFT description of the normal state. The schematic flow chart in Fig. 6 provides an outline of the computational program.

Another central feature of SCDFT is that the empirical Coulomb parameter $\mu^*$ no longer appears. Instead, the screened Coulomb interaction is calculated from a functional involving the electron dielectric function $\epsilon(q, \omega)$ as the central quantity. A few choices for this functional have been tested, analogous to the DFT exchange-correlation functional that likewise has undergone refinements over several decades.

3. Tests for elemental metals

This formulation with its computational implementation (see the flow chart in Fig. 6) has been shown to be remarkably accurate for elemental superconductors, especially considering the lack of any empirical input. In the upper panel of Fig. 7 the calculated value of $T_c$ is compared with the experimental value for five elements with very different strengths $\lambda$ and shapes of $\alpha^2 F(\omega)$, all within the 1-10K range of $T_c$. For the “TF-FE” choice of screening functional (for details see the original papers Lüders et al. 2005; Marques et al. 2005; Sanna, 2017), there is negligible error (the maximum is 0.7 K for Ta). This accuracy for low to modest $T_c$ superconductors is impossible when $\mu^*$ is uncertain (i.e. is used as an empirical parameter) because the difference $\lambda - \mu^*$ involves both the uncertainty in $\lambda$ (from approximate functionals and lack of full numerical convergence) and the fundamental uncertainty in $\mu^*$, while the value of $T_c$ is approaching zero because $\lambda$ is approaching the vicinity of $\mu^*$.

The lower panel Fig. 7(b) illustrates a different test of the theory. It displays the calculated $T_c=0$ static gap $\Delta(0,0) \equiv \Delta_0$ compared to the experimental value. The correspondence is somewhat off for the transition metals Ta and Nb, depending on the approximate screening functional that is used, but the theory can be improved with better screening functions, likely at a cost in computational effort. The report Marques et al. 2005 of these results indicates how the shapes and magnitudes of the electron-phonon spectral function are very different for Pb than for Al, Nb, or Ta, yet the theory is impressively
accurate for all.

4. Application to intermetallic compounds

In the design of and search for new superconductors, improved numerical efficiency is much desired. With such alterations made to the functionals (sometimes with concepts borrowed from models), the procedure was applied by Sanna, Pellegrini, and Gross to elemental metals [Sanna et al. 2020] including those mentioned above, to the transition metal carbide TaC and nitride ZrN, to intercalated graphite CaC$_6$, to the A15 compound V$_3$Si, to the ‘high T$_c$’ boride MgB$_2$, and to the compressed hydride SH$_3$. For these compounds the results are much improved, sometimes dramatically (except for ZrN, with T$_c$ predicted 3 K too high) compared to the 2005 implementation applied to the same compounds. The variation of T$_c$ in this set of materials ranges from 1 K for Al to 203 K for SH$_3$, and a range of calculated $\lambda$ from 0.4 to 2.8. A likely area of needed improvement was identified as the functional related to Coulomb repulsion – what one might heuristically associate with $\mu^*$.

Due to the stated ‘low computational cost’ of the changes allowing substantially improved numerical efficiency, additional capabilities have become available. The full $k$-dependence of the gap over the Fermi surface, a property of increasing interest especially after the discovery of the strong multigap character of MgB$_2$, is one such capability.

The point of this subsection is to emphasize that a fully ab initio and unusually accurate calculation of superconducting state properties and T$_c$ has become available for electron-phonon superconductors. The existence proofs of the underlying formalism allows for extended functionals in which the electron-electron interaction plays a more active role in the pairing mechanism.

G. Limitations of SCDFT/DFT-Eliashberg theory

SCDFT is formally exact for a condensed matter system in thermal equilibrium, analogous to the formal exactness of diagrammatic perturbation theory. In practice one relies on (1) three functionals, most obviously that of DFT for the static lattice, then those necessary for the superconducting state, and (2) the Migdal theorem, which states that electron and phonon propagators can be treated in the single bubble approximation – vertex corrections (more involved diagrams) are negligible. SCDFT involves approximations of the various functionals that appear when treating pairing.

1. Anharmonicity; non-linear EP coupling; quantum proton

For metallic EP superconductors, the theory outlined above has reached a level comparable to that for metals in the normal state (though not yet widely applied), whose phenomenology was formalized in Landau Fermi liquid theory. If the metal is not a Fermi liquid, which can be due to low dimensionality, disorder, or strong electronic interactions, these effects must be built into the functionals. Complications include anharmonicity of the phonons and non-linear EP coupling, both of which arise diagrammatically as vertex corrections but are not often approached strictly strictly diagrammatically. An example is anharmonicity, which can be folded into a quasi-harmonic treatment. Quantum behavior of the nuclei is more prevalent for hydrides due to the light proton mass. Effects on T$_c$ are material dependent; while corrections to T$_c$ from each of these effect are at the ±5-8% level, and further can be either additive or competing, their impact can be more far-reaching in materials with a complex unit cell.

One extreme example of the impact of quantum nature of the proton on structure is provided by LaH$_{10}$ with its clathrate-like H cages around La. Using a classical description of the proton the Fm$\bar{3}$m structure becomes dynamically unstable below 220 GPa, while it is observed to be a 250-260 K superconductor around 200 GPa and somewhat lower (see Sec. VI for more details and references). Errea et al. found that the enthalpy surface contained several atomic configurations [Errea et al. 2020] with a lower enthalpy than the Fm$\bar{3}$m structure, incon-
sistent with the observed structural data. Including the zero-point energy of quantum fluctuations of the proton along with anharmonic corrections, the enthalpy surface reverted to a single minimum at the $Fm\bar{3}m$ structure down to 130 GPa, restoring agreement with experiment.

The fully corrected SCDFT results are shown in Fig. 8 along with those from more conventional Eliashberg theory but with the quantum proton and anharmonic corrections included so LaH$_{10}$ remains stable down to 130 GPa, as found experimentally. The two theory curves differ by about 10% for the hydride and 5% for the deuteride, where anharmonic and quantum fluctuation effects would be smaller for the deuteride. For this case, it seems that current SCDFT is not as accurate in reproducing the experimental results,\(^\text{[Drozdov et al., 2019]}\) which might indicate needed improvement in the SCDFT functionals or might even be accidental given possible complexities of sample preparation. The comparison in Fig. 8 is with experimental results from two groups.\(^\text{[Drozdov et al., 2019; Somayazulu et al., 2019]}\)

2. Eliashberg theory at strong coupling

Over many years there have been studies following the expectation that strong EP coupling alone provides a breakdown of Eliashberg theory for the normal state. The question needs to be specified more specifically, since the DFT-based Eliashberg theory is a given formalism. In the absence of divergences, it makes specific predictions for the system, and these are what current and past solutions describe.

The implicit question is more likely this: at strong coupling, does Migdal-Eliashberg theory give correct results for known or realistically predicted materials? Conclusions (see for example \(^\text{[Bauer et al., 2011; Chubukov et al., 2020; Schrodi et al., 2020]}\)) for treatments and references) have often been based on the Holstein model, originally put forward as a minimal model treatment of EP coupling, applicable to very low density carrier systems, \textit{i.e.} semimetals and doped semiconductors.

In the Holstein model, the EP interaction is entirely on-site: electron charge on a site is coupled to a non-specific scalar displacement on that site. Intersite coupling is due only to indirect coupling through the conserved charge density; there is no modulation of an intersite hopping parameter. As mentioned in Sec. II.E.3, Allen and Dynes provided study of $T_c$ in Eliashberg theory to very large values of coupling, finding a smooth $T_c(\lambda)$ relation with no evidence of a phase transition or even a crossover. The question is: at very large coupling, does Migdal-based Eliashberg theory cease to provide the behavior of real systems, and if so, how does this happen?

DFT-based Eliashberg theory treats the force on an atom due to its displacement as a collective effect, involving changes in potential that induce forces from neighboring shells that impact, sometimes strongly, dispersion of the renormalized (physical) phonons. An electron mass enhancement fixed to the Fermi energy results in an enhancement of the fermion excitation density of states by a factor $1 + \lambda$. The resulting phonon dispersion determines the possible instabilities of the lattice. Numerically exact (Monte Carlo) and other non-perturbative techniques cannot handle the resulting complications of real materials in currently available computational time. Put another way, the strong EP coupling limit based on the full crystal Hamiltonian of Eq. 3 has yet to be attempted.

Experience up to the present, both experimental and computational, indicates that the primary limitation of DFT-Eliashberg theory derives not from strong coupling \textit{per se}, \textit{i.e.} electron mass enhancement, but rather from lattice instability sometimes already occurring at moderate coupling. There are numerous examples indicating that the strong coupling related lattice instability, which is strongly material-dependent and often Fermi surface related, and can occur already at moderate coupling. Phonon branches are renormalized to lower energies in a momentum- and branch-dependent fashion, followed by zero and (calculated) imaginary frequencies as coupling strength increases. Calculations up to $\lambda=4$ (see the following sections) indicate that it is possible to have standard EP coupled systems at that coupling strength (though lattice instability is imminent).

3. Inapplicability of Migdal’s Theorem

We return briefly to the more formal limitations of Migdal-based Eliashberg theory. At the most basic level, the regular decrease in importance of successive terms in the perturbation expansion relies (in a self-consistent treatment at a chosen number of terms) on the smallness of a ratio of phonon velocity to Fermi velocity

$$\frac{\omega_q/q}{v_k^*} = \frac{\omega_q/q}{v_k/(1 + \lambda_k)} << 1,$$

often stated heuristically in terms of the electron-to-ion mass ratio $\sqrt{m/M}$, around 1/300 for medium mass ions. The ratio of phonon phase or group velocity to Fermi velocity is typically $10^{-2}$ or smaller in Fermi liquid metals, providing an excellent expansion parameter. The renormalized (downward, by the EP mass enhancement $1 + \lambda_k$) Fermi surface velocity $v_k^*$ allows for a self-consistent treatment at second order perturbation theory.

In a given band this velocity might approach zero near $E_F$ due to a van Hove singularity, but typically in a small region of the zone and for a single band out of several, perhaps giving rise to unusual effects but not an invalidation of the theory. The platform of flatband materials provides possible realizations for violations of this condition, but the small phase space in a conventional
metal, viz. a wideband compressed hydride (compressed hydrides have occupied conduction bandwidths or 25 eV or more), argues that such occurrences will be rare and, in a many-band background, of minor impact.

Fermi surface nesting, which has an extensive literature, can exaggerate a related condition of this sort, providing a larger phase space in which inter-Fermi-surface scattering acquires a significant phase space. This possibility is real and the resulting behavior of the system (several possible instabilities involving broken symmetries) is material- (and model-) dependent.

4. Thermal fluctuations at high \( T_c \)

Superconductivity theory, especially in the strong coupling regime, posits a complex-valued order parameter (related to the energy gap function). In bulk EP superconductors fluctuation of the magnitude of the gap is rarely regarded as a limiting factor for high \( T_c \). In the quasi-2D high \( T_c \) cuprates, fluctuations of the phase of the order parameter has received a great deal of attention, with one viewpoint being that fluctuations are especially large, and possibly limiting, in the pseudogap region of the doping-temperature phase diagram. While fluctuations are much less of a factor in three dimensions than in two, the elevated values of \( T_c \) in compressed hydrides invites consideration of the question.

An energy scale for the phase stiffness was given by Emery and Kivelson as

\[
V_o = \frac{(\hbar e)^2 a}{16\pi e^2 \Lambda_L^2 (0)}
\]

in terms of the London penetration depth \( \Lambda_L (T) \) at zero temperature and with characteristic length scale \( a \) which would be comparable to the superconducting coherence length. If \( V_o \) (in temperature units) begins to approach \( T_c \), fluctuations would begin to arise, requiring a generalization of Eliashberg theory where no fluctuations appear (it is a mean field theory in this respect).

Eremets and collaborators \( \text{[Eremets et al., 2022] and Minkov et al., 2022a,b]} \) have provided, from upper and lower critical field measurements, the necessary data for \( \text{SH}_3 \) and \( \text{LaH}_{10} \), with \( T_c \) around 200 K and 260 K respectively. The values are, respectively, \( a \approx \xi = 1.84 \) (1.51) nm, \( \Lambda_L (0) = 18.2 \) (14.4) nm. A more recent design for measurement of flux pinning, for different samples of \( \text{SH}_3 \), gave \( \Lambda_L (0) = 27 \) nm. In each compound the ratio \( V_o/T_c \) on the order of \( 10^3 - 10^4 \), so order parameter phase fluctuations are negligible in compressed hydride superconductors.

5. Broader comments

The conclusion based on the full DFT-Eliashberg theory and behavior of real materials is that there is currently no established limit on EP-mediated superconducting \( T_c \), if the aforementioned vertex corrections are taken into account. A few calculations are predicting above room temperature superconductivity in certain compressed hydrides where the strongly coupled phonons have very high frequency and \( \lambda \) exceeds four (see below). While there is no known fundamental limit, it does not follow that prospects are rosy. However, the high \( T_c \) regime (versus the more specific but elusive large \( \lambda \) regime) remains open to new discoveries.

The challenge is (i) to retain lattice stability and (ii) achieve strong coupling to high frequency phonons: both high \( \Omega \) (an appropriate phonon frequency scale) and large \( \lambda \). Very high \( T_c \) seems to require a \( T_c \) expression that is increasing in both \( \Omega \) and \( \lambda \). The theory gives the Allen-Dynes conclusion: the regime of very high \( \lambda \) is

\[
T_c \propto \sqrt{\lambda \omega_0^2} = \sqrt{\eta/M}.
\]

Somewhat peculiarly, \( \eta = N(0) \xi^2 \) is the purely electronic quantity shown by McMillan to be the vibration frequency independent moment of \( a^2 F \). This result is subject to vibrational behavior by only requiring no unstable modes, but an otherwise arbitrary vibrational spectrum. The inverse square root of mass factor is in line with Ashcroft’s original proposition \( \text{[Ashcroft, 1968]} \) – hydrogen systems might have an order of magnitude times larger limit than Nb compounds (with other differences depending on \( \eta \)). Worth repeating: the regime of very high \( T_c \) may not be the same as that of very large \( \lambda \). Hydrides suggest that very high \( T_c \) is the regime of very high frequencies. Unhappily, it is difficult to imagine higher frequencies except at even higher pressures.

III. APPLYING THE THEORY

With DFT-Eliashberg theory in hand, the periodic table and compilations of known superconductors provide an imposing number of possible applications posing the question: which materials systems are the most rewarding for study. This question is discussed briefly in this section, with the question of how to sample the many possibilities giving some discussion in the following section.

A. Choice of Favorable Materials Platforms

Regarding ever higher \( T_c \), transition metal based materials had been sampled and studied, experimentally and theoretically, by 2010, and promising directions were few, but included organic superconductors and interfacial (excitonic) superconductivity. Considering possibilities more generally, the favored palette of atomic constituents for higher, possibly room temperature, superconductors had already been presented by Ashcroft. In 1968, while
research focused on transition metal compounds, he proposed elemental metallic hydrogen as a very high $T_c$ material. \cite{Ashcroft1968} based on (1) its small mass favoring high frequencies, and (2) the vibrating proton without core electrons, which if not too strongly screened should provide the strong scattering that would be required for pairing at high frequencies.

These properties were supported by the BCS Eq. 4 although little of a material-specific nature was understood relating to higher $T_c$. Ashcroft understood that it would require high pressure to metaize hydrogen, specifically to break the $H_2$ molecular bond to create an “atomic hydrogen metal” and bring strong scattering processes to the Fermi energy. That it would require of the order of 1 TPa (ten megabar) or more \cite{McMahon2011} to break the $H_2$ bond may not have been anticipated.

The theoretical and computational capabilities to address this question would not be available until the next century. Skipping ahead: in 2004 Ashcroft refocused his concept, arguing \cite{Ashcroft2004} that hydrogen-rich molecules, viz. $H_2$, $CH_4$, $NH_3$, etc., could circumvent the challenge of breaking the strong $H_2$ bond by replacing it with a weaker bond, and also provide “precompression” (higher H density) in the experiment, thereby lowering the required pressure to produce what would be essentially metallic hydrogen. Section V reveals how remarkably successful this path has been.

B. Computational Implementation

The formalism and some important analysis was complete by the mid-1980s, where self-consistent DFT calculations could provide the electron wavefunctions and band structures giving Fermi surfaces in excellent agreement with experiment, and phonons were becoming available but were still challenging computationally. The expression for the Eliashberg function, re-expressed from Eq. 3, illustrates one of the computational bottlenecks:

$$\alpha^2 F(\omega) = N_\uparrow(0) \frac{\sum_{k,k'} |M_{kk'}|^2 \delta(\omega - \omega_Q) \delta(\epsilon_k) \delta(\epsilon_{k+Q})}{\sum_{k,Q} \delta(\epsilon_k) \delta(\epsilon_{k+Q})}$$

where the Fermi energy $\epsilon_F=0$, and $Q = k' - k$ is the wavevector of the phonon scattering an electron from state $k$ to $k + Q$, both on the Fermi surface. Necessary sums over bands and phonon branches are implicit.

The sums that are shown are each over the three dimensional Brillouin zone, confined by the pair of $\delta$-functions to lines of intersection of Fermi surfaces, one displaced by $Q$ from the other and requiring fine meshes for convergence. As before, the density of states factor is for a single spin, designated by the arrow on $N_\uparrow(0)$. This expression makes evident the geometrical interpretation of a double average of $|M|^2$ over the line of intersection of two Fermi surfaces with relative displacement $Q = k' - k$, all done in a frequency $\omega$-resolved fashion.

To emphasize the numerical challenge, we note that the EP matrix element $M_{kk'}$ of Eq. 8 requires the computation of the self-consistently screened potential due to each phonon displacement, then requiring identification and evaluation of the matrix element between electron states $k$ and $k + Q$ on the Fermi surface, with necessary band and phonon branch indices. These are included within the six dimensional integral. This extensive computation has, through innovative algorithms, been brought to viable although time-consuming level. The electron states are expressed in terms of localized Wannier functions, \cite{Mostofia2008} and the phonons are expressed in terms of localized lattice Wannier functions. \cite{Cockayne2005, Rabe1995} This combination considerably speeds the various required zone samplings.

Linear response algorithms have been implemented to enable the phonon-induced change in screened potential to be calculated from the formal infinitesimal-displacement limit. Beyond the basic DFT formalism and codes, evaluating $\alpha^2 F$ to convergence required a sequence of advancements of formalism and construction of codes, many of which were adapted to parallel computation. In lieu of attempting to describe them, we refer the reader to a modern treatment of anisotropy by Margine and Giustino, \cite{Margine2013}, an extensive review article on the EP formalism with several of the algorithms and references, \cite{Giustino2017}, and a monograph on materials modeling \cite{Giustino2014}.

IV. CRYSTAL STRUCTURE PREDICTION

A. General recent activities

The previous section described the developments that have led to the current capability: given a dynamically stable specific compound, the EP coupling strength $\lambda$, superconducting $T_c$, and several properties of the superconducting state can be calculated reliably. Design of new superconductors requires a separate capability: the prediction of new stable crystalline materials. Design and discovery of new materials was an occasional occurrence until the several agency-wide U.S. program Designing Materials to Revolutionize and Engineer our Future (DMREF) that was initiated in 2011. Related programs have emerged in other countries.

The idea was to push the ever expanding computational power, and theoretical and algorithmic development, to design new materials and properties in many classes, to accelerate experimental discovery, then to speed time to market of new products. Many new programs have supported this initiative, which emphasized computational theory-experiment partnership and research, development, and industry synergy.

Materials design, even restricting oneself to crystalline
materials with modest sized unit cells, is a challenging process. First is the choice of number of elements (we discuss binaries; ternaries and beyond require thoughtful choices) and their stoichiometry, with some chosen property in mind but initially in the background. A great deal is known about structural phases of elemental materials, and about the concentration-temperature phase diagrams of several binary compounds, and certain ternary materials, and about the concentration-temperature phase diagram is known about structural phases of elemental property in mind but initially in the background. A great choices) and their stoichiometry, with some chosen prop-

lenging task (here the formula is normalized to one M atom). The focus here is compressed hydrides, adding the essential pressure variable to the space to be explored. The challenge is to identify candidates that are metallic and thermodynamically stable (metastable phases that are not far from stability may be of interest), then check whether they are dynamically stable. Finally, calculation of electron bands and wavefunctions, and the phonon spectrum, followed by $\alpha^2 F$. Then calculation of $T_c$, $\Delta(\omega,T)$, and a few other properties of the superconducting state can be carried out, using the algorithms and codes mentioned above.

1. Free energy functional

To identify thermodynamic stability (compounds that will not decompose into two or more phases with lower energy) for any chosen pressure P and temperature T, an efficient numerical scheme is required due to the computational complexity of the exercise. Fundamentally, the goal is to identify, for a given stoichiometry $\text{M} \text{H}_n$, the minimum enthalpy $H(P,T)$ over the possible crystal structures. Somewhat more precisely, one really needs for the most precise prediction the minimum of the Gibbs free energy

$$F(P,T) = [E(P,T) + PV(P,T)] - TS(P,T) \quad (17)$$

in terms of the internal energy $E$, enthalpy $H$, volume $V$, and entropy $S$ (mostly lattice vibrational at temperatures of interest).

It is found that the lattice zero-point energy in $E(0,0)$ is more important for hydrogen than for heavier atoms, and the entropy term can shift phase boundaries. The zero point energy and entropy requires calculation of the phonon spectrum, but much screening of candidates can be done without this step. The volume $V$ and internal structural parameters are relaxed (at $T = 0$) to get the enthalpy $H(P,0)$, and the vibrational entropy is calculated when that level of precision is desired. When $H(P,0)$ is plotted versus concentration of H the resulting curve is called the convex hull. The minimum gives the predicted most stable stoichiometry, which is calculated for each pressure of interest.

In the area of compressed hydrides, it has become fairly standard to calculate the complex hull for each pressure of interest, and very often to check the most favorably case(s) for dynamic stability. That information makes it worthwhile for the experimenter to attempt synthesis and characterization, and this symbiosis is evident in several productive collaborations.

B. Evolutionary prediction

An enabling capability in the structural search has been the development, in this century, of evolutionary crystal structure prediction. There are a few methods in use, but the concept is to choose a few candidate structures, relax them, and compare properties, especially the total energy, or under pressure, the formation enthalpy. The most favorable candidates are chosen to guide the construction of new candidates – typically, derived “evolutionarily” by some algorithm – until the most favorable candidate is obtained (for a given pair and a given cell or supercell size, as a practical limitation).

Crystal structure prediction progressed from early random sampling, basin-hopping, and force-field molecular dynamics to first principles DFT-based enthalpy comparisons, often outlining the convex hull of thermodynamic crystal stability. Modern methods, with some prominent ones (with clues to methods contained in their acronyms) being CALYPSO (Yang et al., 2012) (Crystal structure AnalyLysis by Particle Swarm Optimization), USPEX (Glass et al., 2006) (Universal Structure Predictor: Evolutionary Xtallography), AIRSS (Pickard and Needs, 2011) (Ab Initio Random Structure Searching), and XtalOpt (Lonie and Zurek, 2011) (crystal structure prediction and optimization). These codes incorporate various algorithms from simulated annealing, evolutionary/genetic algorithms, minima or basin hopping, particle swarm optimization, metadynamics, and (quasi)random searches, to search the necessarily broad configuration space. A monograph can be consulted for further information (Oganov, 2010).

Actual procedures differ, and the likelihood of finding the true stable formation enthalpy minima is constrained by a few factors, such as unit cell size (faced by all methods) and the effort spent to sample the full phase space, which as mentioned is a daunting task. A typical pro-
FIG. 9 Top row of panels. For the LaH\textsubscript{n} system, the computed convex hull, with specific calculated formation enthalpies with respect to elemental La and H\textsubscript{2}. Shown in the three top panels are pressures of 100, 150, and 300 GPa for the stoichiometries that were obtained in the search. LaH\textsubscript{10} is the compound of most interest, marked with light green background (or gray in grayscale print). Convex hulls often result in several compounds near the convex hull, and may be obtained from hundreds or thousands of enthalpy calculations. Adapted from Errea \textit{et al.} with permission. Bottom row of panels. A schematic diagram of the progression of the high T\textsubscript{c} search program. First, the thermodynamic stability is obtained from compounds on the convex hull. Second, dynamic stability is studied, and phonon spectra are obtained. Third, the DFT-Eliashberg calculation: EP coupling and the phonon spectrum to obtain \(\alpha^2F(\omega)\), and \(T_c\) is obtained from the Eliashberg equation or from one of the fit expressions \(T_c(\lambda, \mu^*, \omega_{log}, \omega_2)\). DFTB, DFPT, and SCDFT indicate separate DFT-based capabilities. Adapted from Flores-Livas \textit{et al} with permission.

Procedure might proceed as follows. After choices of metal atom, H concentration \textit{n}, and some candidate structures (or space groups), the volume and atomic positions are relaxed subject to the chosen pressure. In early steps faster methods with somewhat lower precision can be used. Evolutionary steps based on the most favorable candidates produce new candidate structures. The candidate structures are combined, using various algorithms, to produce the next level candidates.

The process is continued (a computationally taxing procedure) until negligible improvement is occurring, signaling convergence. The H concentration \textit{n} is changed and the process repeated; sometimes this is automated. Varying \textit{n}, which for a given system can involve hundreds of thousands of structures, [Hilleke and Zurek 2022] the “convex hull” delineating the stable compounds in this binary system can be plotted. Examples of the convex hull of LaH\textsubscript{10} at three pressures are shown in Fig. 9(a). This method, involving enthalpy calculated from DFT, has proven successful in predicting stable candidates. Various chosen pressures must be computed independently.

Thermodynamically stable candidate structures then must be checked for dynamic stability (absence of imaginary frequencies) to be valid predictions. Stability can be determined either by phonon spectrum calculations or by \textit{ab initio} molecular dynamics. Only then can DFT-Eliashberg theory be applied to obtain \(T_c\) and a selection of other desired properties obtained from the gap equation. The procedure for full search for high \(T_c\) is outlined in schematic form in Fig. 9(b).
C. Machine learning, data mining

As in many of the sciences and elsewhere, machine learning techniques are being applied to materials design, but few yet with applications to compressed hydrides. The basic idea is pattern recognition: provide a large database to give the neural net the opportunity to identify certain characteristics (“training”) that are (statistically) related to given descriptors. Application of the trained apparatus to new possibilities produces likely candidates for the desired characteristics (viz. $T_c(P)$) with rectitude estimated by various statistical measures. While earlier applied to address other material properties, applications to general superconducting materials have been implemented only more recently. A large database of known superconducting materials exists, consisting of the compound formula, crystal structure, and $T_c$. However, no experiment-derived database for compressed hydrides exists due to the dearth of hydride examples, so training must be done on computationally predicted cases.

In 2017 an example of machine learning related to superconducting $T_c$ was provided by Stanev et al. [Stanev et al., 2018] Their study covered a large range of values of $T_c$ and the SuperCon database of over 12,000 superconductors to provide training. With additional guidance of materials properties from the AFLOW repository, Curtarolo et al. [2012] the procedure was applied to the entire Inorganic Crystal Structure Database. Bergerhoff et al. [1983] More than 30 non-copper and non-iron materials, with all being multicomponent oxides, were identified as the most promising candidates.

The machine learning study of hydrides by Hutcheon et al. [2020] provides a more specific instructive example. For descriptors for candidates $MH_n$, they chose H content ($n$), and the metal element ($M$) size (van der Waals radius), atomic number, mass, and electronic configuration (number of $s,p,d,f$ electrons). Notice that these descriptors have little direct relation to the quantities that determine $\lambda$ or phonon frequencies or interaction strength. Their result identified the first three columns of the periodic table as best candidates for $M$, a feature also noted in various hydride overviews.

These low electron affinity elements give up much or even all of their valence electrons to the H sublattice. This added charge raises the H 1s occupation above the half-filled level, promoting metallicity. In studies of atoms $M$ across the entire periodic table, a few outliers exist: atoms with open $d$ or even $f$ shells. Interestingly, the first discovery, SH$_3$, is a different sort of outlier, with sulfur’s open, roughly half-filled $p$ shell. Also unusual for SH$_3$ is the sharp van Hove singularity at the Fermi level, which could contribute to it being an outlier. [Quan et al., 2019]

V. THE BREAKTHROUGH DISCOVERIES: THEORY THEN EXPERIMENT

A. High pressure experimentation

According to Ashcroft’s original concept, metallic hydrogen should provide the acme of $T_c$, since the atom has the smallest mass, with highest frequencies if force constants are strong, and the potential for strong scattering of electrons by proton displacements that seemed likely to Ashcroft. Calculations sometimes including the quantum nature of the proton have predicted the stable structures versus pressure. The predictions are $T_c$ of 500K or higher, [McMahon and Ceperley, 2011] requiring pressures of 500 GPa or higher. Few attempts have reached a static pressure this high, so Ashcroft’s second suggestion [Ashcroft, 2004] has been the avenue of choice: use H-rich molecules as the ambient pressure sample to avoid strong H$_2$ bonding and antibonding states being pulled away from the Fermi level, and to exploit precompression (higher H concentration and density).

Another point merits mention. Over past decades there have been reports of signals of possible room temperature superconductivity, usually in resistance or susceptibility measurements, which are the most straightforward evidences of superconductivity. The samples were invariably polycrystalline, multiphase, or disordered to the point of amorphous. Transport and magnetic measurements often show anomalies in such samples. When such signals are not reproducible, they have made the community skeptical to the point that “USO” is a recognizable acronym – unidentified superconducting object. It is possible that some of them could be evidence of interface superconductivity or some other unusual type, but if not reproducible a report does not receive extended notice.

For this reason the discoveries below focus on reproducible results, noting confirmations. It must however be recognized that the samples in diamond anvil cells are far from the ideal single crystals that are often available at ambient pressure. The compounds are synthesized within the tiny pressure cell at megabar pressures, at temperatures up to 2000K are varied. Resulting samples will typically be polycrystalline, strained, and possibly multiphase, and with hard-to-determine stoichiometry. For first order phase transitions the free energy barriers may be high, making it challenging to reach certain Gibbs free energy minima. For these reasons various confirmations of these first three discoveries of approaching or near

2 DICE: a data platform for materials science. National Institute of Materials Science, Materials Information Station, SuperCon, http://supercon.nims.go.jp/index_en.html (2011).
room temperature superconductivity will be noted. Also, given the complexities of samples (mentioned above) at a given P and T, experimental data will not be uniform across groups, nor even across a given laboratory’s runs. Thus “reproducibility” and comparison with theoretical predictions should be interpreted accordingly.

B. SH₃: the initial breakthrough

1. Theory

In 2014 Li et al. (Li et al., 2014) employed the CALYPSO structure prediction code (Yang et al., 2012) to identify candidate structures with SH₂ stoichiometry at high pressure. Note that this formula is isovalent with H₂O, and the molecules are isostructural. The much more strongly bonded H₂O is known to remain insulating up to the current highest static pressure available. Their study, finding insulating structures at lower pressures, focused on a transition between two predicted metallic structures in the 130-160 GPa region. The lower pressure compound has a low symmetry PI space group, polymeric structure, with an atomic H Cmca phase with a layering of H and of S atoms occurring at higher pressure. The calculated Tc was maximum at the transition (∼160 GPa), around 60K in the PI structure and 82K for Cmca, and decreasing with pressure in this latter phase. The maximum coupling strength, occurring at the structural transition, was λ=1.25, with 40% attributed to S modes (which comprise 1/3 of the phonon branches).

Appearing immediately after, also in 2014, was a prediction by Duan et al. (Duan et al., 2014) that for a SH₃ stoichiometry [labeled at the time after the initial constituents, (H₂S)₂H₂], and later as H₂S], a high symmetry, atomic H structure pictured in Fig. 10(a) was predicted to be the most stable one in the 150-200 GPa range, and the calculated Tc peaked just above 200 k near the lower pressure range of the 160-200 GPa region. Coupling strength λ=2.2 was calculated, with 25% arising from the low energy S modes. The cubic structure has one of the highest symmetries possible for this stoichiometry: SH₃ octahedra in a body-centered arrangement are connected on two sublattices by H atoms which are coordinated only with two S atoms, see Fig. 10(a). Such a remarkably high prediction must have seemed to most readers as beyond reproducibility and comparison with theoretical calculations.

In 2015 Akashi et al. (Akashi et al., 2015) and Errea et al., (Errea et al., 2015) which promoted the understanding of the relation of electronic structure and microscopic processes to promote high Tc.

2. Experiment

Independently, experimentalists were working on S-H samples. Even in 2014, Drozdov and collaborators (Drozdov et al., 2014) posted a notice of pressure-induced Tc up to 190 K in sulfur hydrate samples, and in 2015 the published announcement (Drozdov et al., 2015) was Tc as high as 203 K around 160 GPa in the H-S system. The superconductor responded to magnetic field in the anticipated way, revealing the expected critical field Hc(T) behavior versus temperature.

The sample was later determined by Einaga et al. (Einaga et al., 2016) to have the bcc S sublattice predicted by Duan et al., and it was concluded, based also on computational input and experimental volume, to be bcc SH₃ as shown in Fig. 10(a). The hydrogen→deuterium isotope shift of Tc was large, as theory predicted, and Tc versus pressure was reproduced. This discovery opened the field to the realistic possibility of room temperature superconductivity. High Tc in this T-P regime has been confirmed, for example by Huang et al. who reported from susceptibility measurements Tc as high as 183K around 150 GPa. (Huang et al., 2019)

Magnetic measurements by Erements and collaborators (Erements et al., 2022; Minkov et al., 2022a,b) led to the superconducting material parameters λGL ∼ 18−27 nm, ξ =18.4. The resulting Ginzburg-Landau parameter ΛGL=η/ξ=10−15 indicates strong Type II superconductivity.

In retrospect not surprising, there was significant skepticism about SC at 200K until the following two higher Tc discoveries were made and reproduced. However, for those (relatively few) who understood the degree and accuracy of the computational theory, the fact that experiment agreed with theory would have been convincing in itself, as it was for this author.

C. LaH₁₀: approaching room temperature

1. Theory

The remarkable success of design and discovery for SH₃ emboldened the superconducting materials design community. Binary hydrides MHₙ were the focus. Varying the metal M = valence, size, chemistry and moving toward superhydrides (n > 6, say), would seem to approach the optimal combination to move toward the idealistic case of Ashcroft’s metallic hydrogen.

In 2017 two theoretical groups, Liu et al., (Liu et al., 2018) and Peng et al., (Peng et al., 2017a) nearly simultaneously predicted MH₁₀, with isovalent M=La and Y, to have high Tc, in the 275-325K range depending on element and pressure (always above 200 GPa). Liu...
FIG. 10 (a–c). The structures of SH₃, YH₆, and LaH₁₀, respectively, illustrating the high symmetry and the progressions toward sodalite or clathrate-type structures with increased H content. Small circles indicate H toms, larger circles denote metal atoms. (d) A schematic ω₂–Tc phase diagram of calculated results for five compressed hydrides in three crystal structure classes shown above. Note that pressure is increasing in the upper left direction, the blue region (lower right region) denotes the low pressure region of instability of the structures that are considered. Solid lines connect values of a compound changing with pressure. Pressure lowers Tc; as pressure increases, ω₂ increases but λ and Tc decrease. Lowering pressure increases Tc (and λ) but leads to dynamic instability. The trends suggest an “island of instability” for compressed hydrides. (e) Pickard’s scatterplot in the P-Tc plane of experimental (hexagons) and theoretical (circles) positions for several superconductors. The contours indicate values of the ad hoc figure of merit $S = T_c(K)/[T_{c,MgB_2}(K)^2 + P(GPa)^2]^{1/2}$, which focuses attention on the desirability of low pressure P for higher $T_c$ (see Boeri et al., Sec. 12.)

et al. calculated $\lambda = 2.2$, with $T_c$ around 265K. The structure again was the highest symmetry possible for this stoichiometry, cubic with a clathrate-like shell of 32 H atoms surrounding the metal atom on its fcc sublattice, pictured in Fig. 10(c). Values for $\lambda$ were in the 2.2-2.6 range, similar to SH₂ but also similar to Pb-Bi-Tl alloys from the 1970s with maximum $T_c$=9K that is 35 times lower. (Allen and Dynes, 1975) The differences in $T_c$ are due to the very high H vibrational frequencies (Ashcroft’s primary point) while retaining strong coupling to Fermi surface electrons.

Further studies of this superconductivity were provided by Liu et al. (Liu et al., 2019) and the quantum (zero point motion) nature of the structure by Errea et al. (Errea et al., 2020) as discussed in Sec. II.G.1. Theoretical work by Ge et al. (Ge et al., 2021) indicated that doping LaH₁₀ by B or N on either the La or H site at the few percent level might raise $T_c$ by 30K, i.e. to $T_c \approx 290K$. 
in the 240 GPa regime. This doping also strongly tends to drive the alloys toward lattice instability, a common occurrence when coupling is increased.

2. Experiment

The superconducting materials discovery (experimental) community was also stimulated by the developments on SH$_3$. Synthesis and evidence of superconductivity in lanthanum superhydride around 260 K was announced in two publications (Geballe et al., 2018; Somayazulu et al., 2019) in 2018-2019. Resistivity drops occurred in the 180-200 GPa range for various runs upon cooling and heating, and x-ray diffraction established the fcc sublattice of La, as in the LaH$_{10}$ structural prediction. Superconductivity was confirmed by Drozdov and collaborators, initially at 215 K (Drozdov et al., 2018) but soon thereafter up to 250 K (Drozdov et al., 2019). The latter paper reported vanishing resistivity around 170 GPa, a H isotope effect, $T_c$ decreasing with applied magnetic field, and evidence of the predicted crystal structure.

Magnetic measurements by Erements et al. (Erements et al., 2022; Minkov et al., 2022a,b) were noted in Sec. V.C.2, the latter introducing a new measurement technique probing flux pinning in the very small samples. For LaH$_{10}$, the behavior was characteristic of conventional Type II behavior, with the derived values of $\Lambda(0)_{\perp} = 14.4\, \text{nm}$, $\xi = 14.4\, \text{nm}$, and Ginzburg-Landau parameter $\kappa_{GL} \approx 9$.

D. Yttrium superhydrides: the third discovery

1. Theory

The Y-H system has a more extensive history than S-H or La-H systems, partly because some of the design of La-H materials included the isovalent Y-H system. Li et al. reported (Li et al., 2015) materials design for this system soon after their 2014 work on SH$_2$ discussed in Sec. V.B. Their 2015 structure search at high pressure identified YH$_3$, a bct YH$_3$ lattice with both atomic H sites and H$_2$ units, and bcc YH$_6$ with a sodalite structure, [Fig. 10(b)], Y surrounded by 24 H atoms] as promising candidates. The latter two had predicted $T_c$ around 90K and 260K, respectively, in the (encouragingly low) 120 GPa pressure range.

The YH$_6$ prediction was starting on two counts: predicted $T_c$ was 30% above the already remarkable SH$_3$ value of the previous year, and the required pressure was somewhat lower. YH$_6$ contains some very strongly coupled H modes at comparatively low frequency (not far from instability), accounting for $\lambda \approx 3$ and high $T_c$ in spite of the significantly lowered phonon energy scale $\omega_{\text{log}}$ (the logarithmic frequency moment (Allen and Dynes, 1975)). However, anharmonicity and non-linear EP coupling can change predictions, especially when there are nearly unstable modes. These stoichiometries have not been reported in experimental studies as of 2022.

Peng et al. proposed (Peng et al., 2017b) a focus on hydrogen clathrate structures as a route to RTS. Results for $\lambda$H$_n$ structures with $\lambda$ = Y and La, for $n = 6, 9, 10$, and $\lambda$ = Sc for $n$ = 6 and 9 were presented. The results pertinent for this discussion are for two yttrium hydrides. YH$_9$ at 150 GPa has the largest calculated coupling: $\lambda \approx 4, T_c \approx 250$K. However, YH$_9$ at 400 GPa has an even larger predicted $T_c$: $T_c \approx 290$-300K with a smaller $\lambda \approx 2.3$. Considering the differences in structures and in optimum pressures, even their substantial amount of data only begins to provide guidelines for just what factors are most important in promoting RTS.

In 2019 Heil et al. predicted (Heil et al., 2019) clathrate-like structures from their structural search, focusing on YH$_6$ and YH$_{10}$. The results for YH$_6$ were similar to those of Li et al. (Li et al., 2015), with calculated anharmonic effects accounting for some of the differences. $T_c$ was predicted to be similar, 275K, for YH$_6$ at 100 GPa, an encouraging result for the efforts to produce and retain high $T_c$ hydrides at more accessible pressures. For YH$_{10}$ predicted $T_c=300$K around 300 GPa was obtained, similar to the results of Peng et al. (Peng et al., 2017b).

For this compound a remarkably large coupling $\lambda \approx 4.5$ was reported, reproducing reasonably well results of Peng et al. (Peng et al., 2017b). This value is among the largest values from DFT-Elishashberg theory for a real (if still only predicted) compound. This large value of $\lambda$ ‘benefits’ from very soft phonons, that is, being very close to a dynamical instability, which is a typical occurrence in several crystal classes. (Quan et al., 2019). When this occurs, corrections for anharmonicity, quantum fluctuations of H, and nonlinear EP coupling become necessary to pin down the critical pressure for instability as well as for the most complete prediction of $T_c$. Generally but especially in hydrides, low frequency modes do not promote $T_c$ as much as their contribution to $\lambda$ would suggest (Boeri, 2021; Quan et al., 2019) (see Sec. VI.B).

2. Experiment

Experimental verification of the prediction of high $T_c$ in the yttrium hydride YH$_9$ was announced (Kong et al., 2019) in 2019, and published in 2021: Kong et al. (Kong et al., 2021). $T_c=243$K at 200 GPa in space group $P6_3/mmc$, with the expected structure being clathrate-like. The compound YH$_{10}$ predicted to have higher $T_c$ was not observed in their experiments, which covered certain regions of phase space up to 410 GPa and 2250K. Snider et al. in 2021 provided data (Snider et al., 2021).
indicating $T_c$ up to 262 K for a sample with superconducting phase of stoichiometry likely close to YH$_6$ based on Raman data. This maximum $T_c$ occurred around 180 GPa. Extension and some degree of confirmation was provided when $T_c=253K$ was obtained in (La,Y)H$_6$ mixtures by Semenok et al. \cite{Semenok2021}.

As mentioned, other regions of the Y-H phase diagram have been predicted to display high temperature superconductivity. In 2021 Troyan et al. \cite{Troyan2021} reported $T_c$ up to 224K at 166 GPa in cubic $Im3m$ YH$_6$. This compound is an example of strong effects of anharmonicity due to the structure and small proton mass. The calculated values are $\lambda=2.4$ using harmonic quantities, reduced to $\lambda = 1.7$ with anharmonic corrections. Anharmonicity considerably hardens the lower frequency H phonons, giving a calculated value of $\omega_{\log}=115$ meV and $T_c$ in the range 180-230K depending on some choices. Nonlinear coupling corrections may be important to obtain the best predictions. Items to be aware of when comparing predictions with data have been noted in Sec. V.A.

VI. CURRENT CHALLENGES

A. Further theoretical guidance

The progress in raising $T_c$ over eleven decades of time is illustrated in Fig. 1. The lesson of the past was that increases in $T_c$ cannot be foretold. On the other hand, the discovery of new superconductors with $T_c$ approaching room temperature in compressed hydrides has been enabled by material-specific theory and computational materials design, after which near room temperature superconductivity was predicted, then verified by experiment. Intense effort continues toward discovering higher temperature or more accessible superconductors. Given the rapid progress, the way forward suggests optimism, with some understanding of the microscopic processes being partnered by computational power in the search. Considering that the full DFT-Eliashberg results can be dug into at any level of detail that one wishes, one can expect that helpful understanding will soon begin to emerge. Section VI.B indicates the current level of understanding and identifies the direction needed for further analysis.

In the previous section, some of the calculated values of EP coupling strength $\lambda$ for the verified hydrides (and some others) have been mentioned. The superconducting hydride sample is small, but it has already been clear that $\lambda$ in itself is a unreliable indicator of $T_c$; increasing $\lambda$ should not be the primary goal. Past examples show that, within a given structure, $\lambda$ is increased by lowering frequencies, which (often quickly) leads to lattice instability. Indications from Allen-Dynes analysis \cite{AllenDynes1975} through more recent studies \cite{Quan2019} indicate that increasing the electron-proton scattering matrix element $I_H$ should become the focus of attention, always assisted by large $N(0)$ of course.

Practically nothing is understood at present about what keeps $T^2$ large in the RTS materials when the electron gas is being compressed to higher density, and (in the simplest picture) should be screening more strongly, thereby reducing $T^2$. $N(0)$ itself is usually normal in size, which may help to promote stability. Several groups have noted that the H-atom DOS $N_H(0)$ per unit volume is likely to be the relevant quantity, but it does not seem to correlate strongly with $T_c$.

With leadership in materials design by the theoretical and computational modeling communities, and materials discovery by high pressure experimenters with increasingly advanced techniques, the six decade old hope for room temperature has essentially been achieved: $T_c$ up to the 250-260K range has been reproduced and accepted by 2020, in substantial measure due to agreement with prediction. The hurdle for closer study and application is that 150-250 GPa pressure is required. This observation directly suggests two future primary goals: (1) yet higher $T_c$ at high pressure, and ultimately superconducting atomic hydrogen, for the advancement of scientific achievement and for knowledge base, and (2) producing or retaining HTS to much lower, or preferably even ambient, pressure, for applications. The following comments provide items that could lead to advancement toward these goals. Some progress in these areas are noted in following subsections.

B. Analysis of H coupling

Analysis of strong EP coupling strength, especially at high frequency, is crucial in understanding how to increase $T_c$, as long as it contributes to both the frequency scale and to $\lambda$, and also helps to avoid structural instability. However, accomplishing this in compounds is more involved and less transparent than in elements because the relevant quantities – $N(0)$, matrix elements, masses, and phonon frequencies – are mixtures of the constituent atoms and their interplay.

Hydrides are special, besides their high $T_c$, because the large difference in the metal mass and the proton mass separates the phonon spectrum $F(\omega)$ and hence the Eliashberg spectral function $\alpha^2 F(\omega)$ into separate frequency ranges: \cite{Quan2019} low frequency metal acoustic modes separated by a gap from high frequency H optic modes. This separation of frequency regions allows the isolation of contribution from each atom

$$\lambda = \lambda_M + \lambda_H; \lambda_X = \frac{N_H(0)I^2_H}{M_X\omega^2_{I,X}}$$

for $X$ equal to metal $M$ or hydrogen $H$. This procedure extends the productive analysis of elemental superconductors to allow identification of the individual atomic
contribution for compressed metal hydrides. The contribution of the metal atom, discussed below, is enlightening. The comparison of hydrogen contributions, and the individual contributions contributing to $\lambda$, for the various hydrides becomes possible, with an example being given in Fig. 12.

This separation is shown for the spectral function of $\text{SH}_3$ in Fig. 11. The separation of the degree of participation is almost complete. Whereas it has been common to quote the separate metal and H contributions to $\lambda$, as quoted for a few examples above, the capability of separating all of the atomic information is available from $\alpha^2 F$. This deeper analysis is important because $\lambda$ alone is a poor indicator of $T_c$. What is of prime interest is the contribution to $T_c$ (versus $\lambda$, energy gap, or other properties) from each atom.

Analysis based on the frequency separation of metal and H modes was reported by Quan et al. (Quan et al. 2019) for five binary hydrides from three crystal structure classes. A phase diagram illustrating some aspects of the analysis is shown in Fig. 10 (4). A primary indication is of rough phase boundary identifying an “island of instability” in the high $T_c$, but lower $\omega_2$ regime. Within a high $T_c$ phase, this instability is encountered as pressure is lowered, with $\lambda$ and $T_c$ increasing until instability is reached. For brevity, only a few other results of this trend study will be discussed here.

The first observation in this study is startling. Although it is an appreciable fraction of $\lambda$, the metal atom contribution to $\alpha^2 F$ affects $T_c$ very little, and sometimes lowers it. This occurs because coupling to low frequency vibrations lowers the phonon frequency scale—the prefactor in $T_c$ in Eqs. 1, 10, and 12—off-setting the increase in $\lambda$. Quan et al. explained why this does not violate the Bergmann-Rainer ‘theorem’: additional coupling at one frequency affects the phonon spectrum and coupling at other frequencies. This conclusion can be confusing, because the metal contribution to $\lambda$ is evident and often emphasized, while the effect on the frequency scale is never calculated, therefore remaining invisible. This result focuses attention for analyzing and understanding $T_c$ on hydrogen alone—neglect the metal—thereby providing a focus for a more transparent understanding of $T_c$ in terms of hydrogen properties.

Second, and following from the first: $N_H(0)$ and $\omega_{2,\text{H}}^2$ are calculated and the mass $M_H$ is fixed. This makes it possible to extract and analyze the least understood and very important quantity: the matrix element $\mathcal{I}_H^2$ factor in Eq. 19 for scattering electrons from the moving H atom. Quan et al. provide a discussion of this and several other aspects of EP coupling in their five chosen compounds, arguing that understanding of $\mathcal{I}_H^2$ is the main missing link in the understanding of EP coupling in compressed hydrides. A synopsis of some of their analysis is shown in Fig. 12. In the $(\eta_H, \lambda_H)$ plane of Fig. 12 (a) and the pressure dependencies of $\eta_H, P_H^2$ in Fig. 12 (c), (f), respectively, the example hydrides seem to self-organize into groups. In the $(\kappa_H, \lambda_H)$ plane and the other two pressure dependencies, the magnitudes and pressure behaviors of the five phases show considerable overlap but clear trends. Note that $\mathcal{I}^2$ tends to increase with pressure, but (for example) the magnitudes at 300 GPa differ by a factor of 3.

As mentioned earlier, the strong tendency for $\mathcal{I}^2$ to increase with pressure is counterintuitive, since the increased electronic density under volume reduction would
FIG. 12 Atomic H quantities in the indicated superconducting compressed hydrides, obtained by taking advantage of the spectral separation of metal acoustic modes and H optic modes. Corresponding calculated values of $T_c$ can be seen in Fig. 10(d).

(a,b) $(\eta, \lambda)$ and $(\kappa, \lambda)$ phase planes, respectively, indicating small (resp. large) relative variations. (c-f) Pressure variations of the indicated quantities, showing the magnitudes and rates of variation with pressure. Units: $\lambda$, unitless; $\eta = N(0)I^2$ and $\kappa = M_H\omega^2$, eV/Å$^2$; $I^2$, eV$^2$/Å$^2$. The background color (or different shadings in grayscale figures) indicate the different types of plots, see the axis labels. Adapted with permission from Quan et al. 2019.

seem to provide increased screening of H motion. This should reduce the $I_H^2$ factor given in Eq. 8. The change in potential due to H displacement can be evaluated within linear response theory, either from the dielectric constant with all local field terms (highly intensive numerically) or numerically from linear response, which is the method of choice and is still computationally intensive enough to sometimes limit $Q$ (phonon) point grids.

The pressure increase in $I_H^2$ indicates there is more physics to be understood. One can compare with a simple and efficient approximation from Gaspari-Györffy theory,\cite{GaspariGyorffy1972} which uses a rigid atomic potential displacement model. The result requires negligible computation, but involves phase shifts of the potential that provide only a limited physical understanding. The Gaspari-Györffy model was applied to SH$_3$ by Papaconstantopoulos et al.\cite{Papaconstantopoulos2015}, finding indeed that $\eta_{GG}^H$ also rises steeply and nearly linearly from 18 eV/Å$^2$ at 200 GPa to 25 eV/Å$^2$ at 300 GPa. The underlying mechanisms remain unclear.

Fig. 12(f) indicates the extracted value (without approximation) of $I_H^2$ is 10-11 eV$^2$/Å$^2$ around 200 GPa, indicating a 70% overestimate by Gaspari-Györffy theory, with its neglect of screening. The calculations of Quan et al. were not extended to higher pressure, so a more complete comparison is not available. Hutcheon et al. have used Gaspari-Györffy theory to give a quick estimate of $\eta$ in their machine learning study.\cite{Hutcheone2020}.

C. How to produce strong H coupling

1. The metal-induced atomic hydrogen paradigm

The leading paradigm in compressed hydrides, from Ashcroft, is the need to break, or deter, the strong H$_2$ molecular bond or other molecular bonds (viz., CH$_4$, NH$_3$), which will move the bonding band below $E_F$ and push the antibonding bands above $E_F$, leaving little or no H contribution at the Fermi level. Breaking up molecules, leaving open 1s shell atomic hydrogen rather than molecular hydrogen, is the paradigm followed by the three first discoveries, and by nearly all of the predicted hydrides with $T_c > 100$ K. Being the leading concept for 60 years, little more needs to be said about this paradigm, but this being a new field of investigation and discovery, it is important to look for other paradigms.

2. An alternative paradigm: activating bonding states

The RTS examples introduced in Sec. V are from the anticipated class, in which hydrogen becomes atomic (no overt covalent bonding) and predominantly H 1s bands lie at the Fermi level and are primarily, almost overwhelmingly, responsible for high $T_c$. Computational explorations of ternary hydrides (see below) have found that at too low pressures, cells with molecular H$_2$ or H-rich molecules provide the stable structures. In such compounds, bonding and antibonding H 1s levels are split away from the Fermi level (below and above, respectively) and H vibrations provide little or no EP coupling. It seems that a guiding principle is that atomic H dominance leads over other productive possibilities, and its coupling strength requires further attention. Yet another
possibility has arisen.

$Li_2MgH_{16}$. Sun et al. reported calculation [Sun et al. 2019] of $T_c$, around 475K at 250 GPa for $Li_2MgH_{16}$, which is best pictured as a lattice of $MgH_{16}$ clathrate-like units intercalated by Li. The large unit is rather stable, while the Li donor adds electrons, lends the H sites a more metallic character, and doubles the value of N(0). The maximum H frequency was $2400 \text{ cm}^{-1}$, in the same range as other HTS hydrides at similar or somewhat lower pressure. As in other hydrides, $T_c$ decreases with increasing pressure while the frequency spectrum increases.

The distinction that makes $T_c$ higher than in other compressed hydrides is unclear. The small mass of Li results in the overlap of its phonon projected density of states with that of H: Mg is lower and nearly separate. For $\lambda=3.35$ in this P-$T_c$ regime, roughly 1.75 can be ascribed to the low frequency metal atoms ($\omega < 20 \text{ THz}$), with 1.60 arising from H modes extending up to 70 THz. The two H sites contribute very differently to the bands crossing the Fermi level (Y. Quan and W. E. Pickett, unpublished). Comparative analysis with other HTS compressed hydrides is needed to obtain insight into the origins of high $T_c$.

$LiB_2H_8$. As another example, Gao et al. in 2021 reported a designed (predicted) H-rich system [Gao et al. 2021] in which high $T_c$ (though not RTS) arises in a distinctive manner. The material is one in which BH$_4$ units (identifiable molecules) lie in interstitial positions within a bcc potassium sublattice, comprising KB$_2$H$_8$ = K(BH$_4$)$_2$, with identifiable BH$_4$ molecules. Extrapolating from results on other ternaries, such a compound having only molecular hydrogen should be unpromising. However, the chemistry (more specifically, the Madelung potential) is such that each molecular BH$_4$ radical (likely unstable in itself, lacking the extra electron that stabilizes methane CH$_4$) obtains $\frac{1}{2}$ electron from the K ion, leaving the uppermost (least strongly bound) molecular orbital half-empty. The resulting radicals are stable within the sublattice of positive K ions, and the compound is predicted to be dynamically stable.

KB$_2$H$_8$ is calculated to be metallic, but with the character of a heavily hole-doped wide-gap insulator. This leaves covalently bonded bands that are strongly coupled to B-H bond-stretch modes at the Fermi level, a close analog [An and Pickett 2001] of MgB$_2$ with its $T_c=40K$. The calculation of Gao et al. gave $T_c\approx140 \text{ K}$ at the modest pressure of only 12 GPa, arising from very large $\lambda=3$ but an unusually low frequency scale $\omega_{log}=33 \text{ meV}$ (100+ meV is more typical of RTS hydrides, but at pressures of 150 GPa and higher). This is a three dimensional extension of the argument that such MgB$_2$-like systems can be optimized to produce much higher EP-coupled superconductivity. [Pickett 2008] Further improvements in this direction seem possible.

However, with such a large hole density that covalent bonds may be unstable, LiB$_2$H$_8$ may not be a thermody-

namically stable composition. This scenario played out in Li$_{1-x}$BC, where for $x \sim 0.2 - 0.3$, $T_c$ up to 75 K was inferred. [Rosner et al. 2002] Substantial experimental effort [Fogg et al. 2003] could not produce the desired structure at the target doping levels, obtaining instead distorted and disordered materials. However, Sr$^{2+}$ doping on the K$^{1+}$ sites, lowering the hole doping level, and broader synthesis routes may provide pathways to desired materials. [Nakamori and Orimo, 2004]

D. Increasing accessibility; metastable structures

1. Lowering the required pressure

Of growing concern is to find, perhaps by design and discovery or perhaps by serendipity, materials that will retain their high pressure high values of $T_c$ to lower pressures, with the intention of finding applications. Most of the binary hydrides have been explored with computational means. [Bi et al. 2017, Flores-Livas et al. 2020] Analysis of the results remains to be done, and unfortunately published results often do not provide much of the information that is required, including the atom-specific quantities in $\lambda$ in Eq. 9. The study of Quan et al. has initiated such analysis, [Quan et al. 2019] but was limited to a few binary hydrides for which their recalculation provided the data required for the analysis of the electronic properties.

Separately but equally valuable is an improved understanding of the stability, or lack thereof, of high $T_c$ materials composition and structures. Considering broadly, there are several examples of this scenario: a high $T_c$ material is discovered (either computationally or experimentally) and its structure understood; pressure is lowered and $T_c$ (and calculated $\lambda$) increases but a phonon branch is lowered; a structural phase transition occurs at a critical pressure $P_{cr}$; in the low pressure phase $T_c$ is much lower or perhaps vanishing; the structure of the new phase includes H-rich molecular units, including possibly H$_2$, without much or any atomic hydrogen. One question being addressed is: how can this process be pushed to lower $P_{cr}$, or even (ideally) to ambient pressure.

2. More complex hydrides; speeding searches

After the design and discovery reported in the sections above, emphasis has broadened. Higher values of $T_c$ are of course still of great interest; after all, applications at room temperature will require $T_c$ around 30% higher (375-400K), or even higher for high current density applications. Given the considerable number of binary hydrides that have been modeled and mined for high
$T_c$ (here meaning roughly, $T_c > 100$K), useful for applications), searches are being extended to ternary hydrides.

The palette of ternary hydrides is much larger than that of binaries, thereby opening new candidates and new computational challenges. So far the emphasis has been on the more H-rich possibilities, viz. $\text{A}_i\text{B}_j\text{H}_n$, with small $i$ and $j$ and larger $n$. With atoms A and B selected from the (say) 60 most reasonable choices of elements, and with concentration $n$ ranging up to 12, this class has of the order of $10^4$-$10^5$ formulae, and for each of these, many crystal structures are possible. Given this complexity, techniques in high-throughput computing, data set construction, data mining, coupled with machine learning, are being applied to the search for promising candidates, but a full search is not in sight. Background on these activities can be found in the 2021 Roadmap compilation. (Boeri, 2021) There are too many reports already on ternary hydrides with too little analysis to attempt to provide an overview. Several candidates have predicted $T_c > 100$K, however explorations of the generalized convex hull to find the most stable stoichiometries have been limited.

3. Exploring higher pressures

While discussion of advancing high-pressure techniques is well beyond the scope of this article, it should be mentioned that experimental extensions to achieve higher pressures more readily, and to adapt measurement techniques to obtain more general data on the samples, are continually pursued in the high pressure laboratories that have contributed to high $T_c$ hydrides at high pressure.

VII. REGULARITIES IN COMPRESSED HYDRIDES

While this article is not intended as a topical review of compressed hydrides, it should have raised questions, and addressed some of them, about the properties that provide close approach to room temperature superconductivity. Several features that appear to be important clues, i.e. to have some generality, have been identified.

- A fundamental question is: what structure types of compressed hydrides are favored, and in what pressure ranges? Some guidelines seem to have arisen. In the lower pressure range, hydrogen molecule phases arise and are poor superconductors or, often not reported, insulators. At higher pressure (say, 150-300 GPa, atomic hydrogen structures are frequently favored, with H-caged metal structures (e.g. clathrate) being common. At still higher pressures (with fewer published examples) less common structures are predicted; for example, one has several layers of hydrogen followed by a few layers of metal, suggestive of incipient phase separation. This area is a complex one, and the reviews and overviews mentioned below should be consulted.
- An overriding question – what properties enable room temperature superconductivity – remains open to clarification. It is not large $\lambda$ per se; $\lambda \approx 2$-3 is similar to that in Pb-Th-Bi alloys, with their $T_c \approx 10$K. These low $T_c$ materials have very soft phonons, due to heavy masses and nondescript $s-\sigma$ bonding. Compressed hydrides have frequency scales up to 150-200 meV (1750K-2300K), compared to those heavy atoms with frequency scales of 50-75K. The enabling feature is, as Ashcroft foretold, producing H-derived modes at high frequency while retaining strong coupling to Fermi surface carriers. This is, unfortunately, more an observation than an answer.
- Strong coupling. Barring innovations, larger $\lambda$ should not be the overriding aspiration. There are numerous cases, including compressed hydrides (mentioned above), where increasing $\lambda$ increases $T_c$, but rapidly encounters soft phonons and lattice instability. $\lambda$ is an unreliable descriptor for a high $T_c$ search. Clue: $\eta/M$, independent of frequencies, is much better. (Allen and Dynes, 1975; Quan et al, 2019) Since higher frequency scales will likely require ever higher pressures, the alternative seems to be pushing strong coupling to the high frequency region. In compressed hydrides ‘bond stretch’ modes seems not to be a dominant consideration. Generally H-H ‘bonding’ is not a clear feature; H-metal bonding is more often a topic of consideration.
- Various groups (Belli et al, 2021; Semenok et al, 2020) have observed that atoms in columns I, II, and III provide, with occasional exceptions, the binary hydrides with high $T_c$. These atoms have low electronegativity, readily donating electrons to the H sublattice(s). The resulting negatively charged H atoms (versus neutral, half filled entities such as the H$_2$ molecule) promote breaking of H bonds and producing metallic ground states. This factor might also be related to the retention of strong coupling to high energy vibrations.
- Is a higher concentration of H the key? The indications are that a large fraction of H states at the Fermi surface, i.e. reflected in a large ratio $N_H(0)/N(0)$, is not a clear determining factor, or at least is not essential. This is a straightforward band structure quantity, and cannot be estimated before the band calculation is done, because band structure effects cause structure in $N(\varepsilon)$. An example for this item: SH$_3$ is somewhat exceptional, with a strong van Hove peak at the Fermi level. (Ghosh et al, 2019), (Pickard et al, 2020), (Flores-Livas et al, 2020) have highlighted the $N(0)$ factor in compressed hydrides.
- Producing “atomic hydrogen,” as opposed to molecular-bonded hydrogen, has been the overriding objective, and so far the productive one. Pressure will eventually decompose hydrogen-rich molecules, but other methods
(viz. doping) should be kept in mind. Doping holes into bonding states may be promising. (Gao et al., 2021) but too much doping will make a stable phase prone to structural instabilities.

- Lattice instabilities. In several cases of high $T_c$ hydrides, it has been found that within a given phase $T_c$ decreases with increasing pressure. Conversely, $T_c$ increases as pressure is decreased, $\lambda$ increases, and the modes giving the increase in $\lambda$ are renormalized to lower frequency and then become dynamically unstable. For the five systems illustrating this self-limiting process in Fig. 10(d), there is an indication of a phase boundary for binary hydrides in the $\omega_2$-$T_c$ plane.

- Naturally, the lattice stiffness $\kappa_H = M_H\omega_2^2$ increases with pressure. However, strong coupling is far from uniform throughout the H-derived optic modes. Pressure does increase the frequency scale, but appearing squared in the denominator it decreases $\lambda$. This trade-off has long been a persistent issue when pursuing higher $T_c$ superconductivity. More focus needs to be aimed at increasing $\eta_H$.

- The scattering efficacy of the vibrating H atom, $\mathcal{T}_H^2$, increases with pressure, according to current information. (Papaconstantopoulos et al., 2015; Quan et al., 2019) The origin of this simple fact is unclear, but the theoretical and computational means to understand it is available within DFT codes (but requiring proper extraction and analysis).

- For the small set of examples that has been studied, $\mathcal{T}_H^2$ varies from one to another over the pressure range of interest, by a factor of three, and $\eta$ by a factor of two, ([Quan et al., 2019] see Fig. 12). This difference can be attributed partially to broadening of the (largely H) bandwidth with increasing pressure, hence tending to decrease $N_H(0)$. Again, the origin is unclear but the detailed computational theory exists to analyze this fact in detail.

A careful study of the “regularities” listed above will reveal repetition and apparent inconsistencies and contradictions. Example: strong coupling at high frequency is what is really important, but also $\eta = N(0)\mathcal{T}^2$ (which is independent of frequency) is what really matters at strong coupling. Such various viewpoints are what must be confronted in the quest for higher $T_c$ at lower pressure. Also, nearly universally high $T_c$ has been couched separately in terms of $\lambda$ and one characteristic phonon frequency, viz. $\omega_\log$. This approach may be misguided (i.e. not the most profitable) for progress. It has been noted that, for the five compressed hydrides mentioned in previous sections, the simple relation $T_c \propto \sqrt{\text{area \ under \ } \alpha^2F}$, works quite well in spite of its simplicity. (Quan et al., 2019) Improvements in the $T_c$ equation and in understanding by a generalization to another 1-3 more characteristics obtained from $\alpha^2F$.

VIII. EPILOGUE

A number of overviews (Boeri and Bachelet, 2019; Pickard et al., 2020; Pickett and Eremets, 2019; Shimizu 2020) and more extensive collections (Bi et al., 2017; Boeri, 2021; Flores-Livas et al., 2020; Hilleke et al., 2022; Hilleke and Zurek, 2022; Semenok et al., 2020; Zurek and Bi, 2019) on predictions of hydride superconductivity are available. The achievement of (near) room temperature superconductivity has stimulated extended high pressure techniques and the enabling of additional measurements, in step with improved analysis and interpretation of data. (Guan et al., 2021; Hemley et al., 2019)

The first point of this perspective was provided in Secs. II and III, which summarizes the sequence of theoretical and algorithmic advances, followed by numerical implementation, that have produced an accurate, material-specific theory of EP superconducting $T_c$ as well as several superconducting properties not discussed here, mostly stemming from the complex superconducting gap $\Delta(\omega, T)$. The three initial advances predicted by the theory and then confirmed by experiment are discussed in Sec. V:

- $\text{SH}_3$, 200K at 100 GPa
- $\text{LaH}_3$, 260K at 200 GPa
- $\text{YH}_9$, 240-260K around 250 GPa.

This article is intended to provide an overview of the theory-driven forces behind the design and discovery of room temperature superconductivity. The experimental effort on hydrides has been impressive as well. Room temperature superconductivity was a much discussed but distant goal in the 1970s, but expectations faded after 13 years with no increase in $T_c$ to date. The discovery of high $T_c$ cuprates revived the dream to some extent, but the focus of research soon reverted to an intense study of the properties and mechanism(s) (versus magnitude of $T_c$) of superconducting quantum materials, a topic that remains a leading paradigm of condensed matter physics that is being broadened to other classes, properties, and applications. In terms of temperature, HTS has been superseded only by compressed hydrides – the long-sought room temperature superconductors.

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