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High-Temperature Superconductivity in Hydrides: Experimental Evidence and Details

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Abstract

Since the discovery of superconductivity at ~ 200 K in H_3S [1], similar or higher transition temperatures, T_cs , have been reported for various hydrogen-rich compounds under ultra-high pressures [2]. Superconductivity was experimentally proved by different methods, including electrical resistance, magnetic susceptibility, optical infrared, and nuclear resonant scattering measurements. The crystal structures of superconducting phases were determined by X-ray diffraction. Numerous electrical transport measurements demonstrate the typical behavior of a conventional phonon-mediated superconductor: zero resistance below T_c , shift of T_c to lower temperatures under external magnetic fields, and pronounced isotope effect. Remarkably, the results are in good agreement with the theoretical predictions, which describe superconductivity in hydrides within the framework of the conventional BCS theory. However, despite this acknowledgement, experimental evidences for the superconducting state in these compounds have recently been treated with criticism [3–7], which apparently stems from misunderstanding and misinterpretation of complicated experiments performed under very high pressures. Here, we describe in greater detail the experiments revealing high-temperature superconductivity in hydrides under high pressures. We show that the arguments against superconductivity [3–7] can be either refuted or explained. The experiments on the high-temperature superconductivity [8] and eliminate it [3].

Keywords High pressure · Superconductivity · Hydrides

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1 Introduction

The discovery of new superconductors demands strict, unambiguous proof. Obviously, this is applicable to the nearly room-temperature superconductivity (RTSC) discovered in hydrogen sulfide at 203 K at high, megabar-level pressures of ~ 150 GPa [1, 9] (1 GPa ~ 10⁴ atmospheres). High pressures put significant constraints on the experimental study of superconductivity. At such pressures the typical size of samples residing in a diamond anvil cell (DAC) is necessarily small - of the order of ~ 50 µm in lateral dimension. Nevertheless, even for DACs there is a number of available experimental methods allowing these tiny samples to be probed using various electrical, magnetic, optical, and X-ray diffraction techniques [2, 10]. Electrical transport measurements demonstrated a zero resistance state in H₂S below its T_{c} [1]. The application of an external magnetic field provided further evidence for superconductivity: T_c shifts to lower temperatures. The isotope effect — a strong decrease of T_c resulting from the replacement of hydrogen atoms by

deuterium — clearly indicates the phonon-mediated BCS mechanism of superconductivity [1].

The Meissner effect, or expulsion of magnetic field from a superconductor, is a crucial and independent test for superconductivity. Therefore, samples with H_3S were subsequently probed by high-sensitivity measurements of magnetic susceptibility using a superconducting quantum interference device (SQUID) magnetometer [1] under megabar-pressure conditions. A clear diamagnetic signal (screening of magnetic field) below T_c was observed in excellent agreement with the electrical transport measurements. This result was supported by nuclear resonant scattering measurements [11]. Later the superconducting gap was estimated through infrared spectroscopy [12]. X-ray diffraction gave information on the crystal structure of superconducting phases [13, 14].

An essential and crucial step for any newly discovered superconductor is reproducibility. In contrast to the discoveries of MgB_2 and the cuprates, where superconductivity was immediately reproduced by many laboratories, the superconductivity in H₃S could not quickly be replicated by other groups because of the complicated synthesis protocol associated with the condensing and loading of H₂S at low temperatures, and the difficulties inherent to measurements at the megabar-pressure range. An important step for the independent confirmation of superconductivity in H₃S was electrical transport measurements of the prepared samples in the laboratory of Prof. K. Shimizu at Osaka University [13]. Shortly afterwards, H₃S was independently synthesized and superconductivity was fully confirmed [15–19]. Later, new hydrides with higher T_c s and crystal structures distinct from that of H₃S were discovered by different groups, including $T_c \sim 250$ K in LaH₁₀ [20–22], $T_c \sim 243$ K in YH₉ [23], and lower T_c s in many other metal superhydrides [2].

More recently, room-temperature superconductivity was announced first at 288 K under a pressure of 276 GPa [24] and then at 294 K at 21 GPa [25] in the same sulfur-carbonhydrogen system. However, these contradictory measurements with no access to the raw data have found serious criticism [5, 26–29]. After more than a year, room-temperature superconductivity in the sulfur-carbon-hydrogen system was neither confirmed experimentally nor supported by theoretical calculations. Despite the significant efforts, we also failed to reproduce these observations. Our study on the S-C-H system will be described in detail elsewhere [30].

The experimental evidence for superconductivity in hydrides is consistent with a large number of theoretical predictions and calculations [2] and the general theory of conventional superconductivity [31, 32]. Meanwhile J. Hirsch and F. Marsiglio (H&M) have been trying to explain superconductivity in hydrides through their theory, where superconductivity results from "a few holes conducting through a network of closely spaced negatively charged anions, in conducting substructures with excess negative charge" [3, 8, 33]. In contrast to BCS theory, in the proposed theory [8], superconductivity is driven by a universal mechanism that is not connected with electron-phonon interactions. H&M first tried to explain superconductivity in H_3S [33] by holes, which conduct through the direct overlap of planar s-p orbitals of sulfur atoms. Sulfur atoms, instead of hydrogen atoms, were stated to be essential for the "hole superconductivity." Explanation for superconductivity by the hopping holes between negatively charged anions (e.g., sulfur in H₃S) obviously fails for metal superhydrides such as LaH_{10} where the rare-earth atoms are cations [20, 21, 34]. The contribution of hydrogen in superconductivity is impossible to ignore because of the isotope effect - substitution of hydrogen atoms by deuterium atoms leads to shift of T_c to lower temperatures — a *negative* shift. The shift of T_c is enormous: $\Delta T_c \sim 70$ K for the $P6_3/mmc$ phase of YH_0/YD_0 [23] and the Fm-3 m phase of LaH_{10}/LaD_{10} [20]. For comparison with the previous studies, the largest shift was found in MgB₂ with $\Delta T_c \sim 1.0$ K between Mg¹¹B₂ and $Mg^{10}B_2$ [35]. The values of the isotope coefficient in hydrides are close to the value of $\alpha \approx 0.5$ for conventional superconductivity: $\alpha = 0.46$ for the *Fm-3 m* phase of LaH₁₀ [20], $\alpha = 0.39$ and 0.50 for the *Im-3 m* phase of YH_6 and $P6_3/mmc$ phase of YH_9 , respectively [23].

An inability to explain the isotope effect clearly shows that the hole superconductivity is invalid. According to this theory, the isotope effect should be small, if any, and with a *positive* shift $-T_c$ should increase for a crystal with heavier atoms. Failed to explain conventional hightemperature superconductivity in hydrides by the hole superconductivity, H&M [3-5, 8, 33] put into doubt the whole evidence of conventional superconductivity: "the experimental observations rule out conventional superconductivity but it could be either unconventional superconductors of a novel kind, or alternatively, that they are not superconductors" [4]. For several years, H&M have been trying to find inconsistencies, contradictions in the published works, demonstrating an apparent misunderstanding of the high-pressure experiments. The arguments against superconductivity raised in Refs. [3-7] can be summarized as follows: (i) observation of thermal hysteresis in resistance measurements; (ii) absence of broadening of the superconducting transition under external magnetic fields; (iii) absence of reliable magnetic measurements.

We address these issues by considering the related experiments in greater detail and including some new results in comparison with the published papers. We hope that this comprehensive analysis of the high-pressure experiments on high-temperature superconductors will be useful to the community.

2 Results

2.1 Thermal Hysteresis in Temperature-Dependent Measurements

The transformation between the normal and superconducting states under zero magnetic field in type-II superconductors is a second-order phase transition, and therefore, there should be no hysteresis in the temperature-dependent resistance measurements between cooling and warming cycles. However, typically there is a significant hysteresis in R(T) measurements of hydrides under high pressures. Based on this fact, H&M concluded that, most likely, hydrides are not superconductors [36].

Besides the sample property, an imperfect thermal equilibrium between thermometer and sample can also result in extrinsic hysteresis. Upon cooling, the DAC is usually in a cold gas flow (nitrogen or helium), and the temperature of the sample lags with respect to that of the thermometer located in the exterior of the DAC. Obviously, this difference depends on the cooling rate (Fig. 1a–c). Upon slow warming without using a heater (typical rate ~ 0.3 K min⁻¹), the DAC body thermalizes with the cryostat, and consequently



Fig. 1 The superconducting transitions in LaH₁₀ and H₃S compounds detected through the temperature dependence of the electrical resistance upon cooling (blue curves) and warming (red curves). **a**, **b** R(T) measurements of the *Fm-3 m*-LaH₁₀ phase residing in a large DAC with an outer diameter of 33 mm. The temperature sweep dT/dt is ~1 K min⁻¹ upon cooling and ~0.3 K min⁻¹ upon warming. **a** The temperature sensor is attached to the body of DAC, leading to a hysteresis of ~14 K. **b** The sensor is attached to the body of DAC and carefully insulated from the cooling gas. In this case, the hysteresis is significantly lower ~3.5 K. **c**, **d** R(T) measurements of the *C2/m*-LaH₁₀ phase in the miniature DAC with an outer diameter of

8.8 mm. **c** The sensor is attached to the body of the DAC and dT/dt is ~3 K min⁻¹ upon cooling and ~0.3 K min⁻¹ upon warming, leading to a hysteresis of ~25 K. **d** The same DAC but the sensor is attached to the gasket and dT/dt is ~0.15 K min⁻¹ upon cooling and warming, leading to a hysteresis of ~0.3 K. **e**, **f** Superconducting transitions in the *Im-3 m*-H₃S phase residing in the miniature DAC with dT/dt of 0.15–1.5 K min⁻¹. **e** The sensor is attached to the gasket and the hysteresis is ~0.3 K; **d** the sensor is attached to the gasket and the hysteresis is ~0.01 K. Insets: photos of the miniature DAC showing the location of the temperature sensors

the DAC and the thermometer are nearly in equilibrium. Therefore, we and other groups use the R(T) data taken upon warming cycle. We checked this procedure with the DACs whose hysteresis was significantly reduced through a careful insulation of the thermometer from the cooling gas (Fig. 1b). The thermal hysteresis decreased from a value of ~ 10–20 K to ~ 3 K, but the R(T) curve upon warming cycle remained nearly the same.

Various cooling/warming protocols are used by the different groups. For instance, in the experiments on LaH₁₀ [21] its higher $T_c = 260$ K was determined from the R(T)data upon cooling. This value could be inaccurate because the cooling rate was not reported. In contrast, in Ref. [22] the transition at $T_c = 250$ K was determined upon warming cycle and it is consistent with Ref. [20]. Another uncertainty in Ref. [21] is the evaluation of a pressure value, at which the superconducting transition occurred, since it was not determined directly.

The most accurate low-temperature measurements were collected with a miniature DAC (Fig. 1c–f). Since such DACs are diminutive in a size and thermal mass (an outer diameter is 8.8 mm and weight ~9 g) and have relatively high thermal conductivity (the DAC body is made of a copper alloy), the thermal gradients are minimized. Still, one observes thermal hysteresis when the temperature sensor is attached to the body of DAC in a typical experimental arrangement (Fig. 1c, e). However, the measurements are much more accurate, if the sensor is attached to the metal gasket, which surrounds the sample compressed between both diamond anvils and, thus, is in a direct thermal contact with it. In this arrangement, the difference in R(T) between cooling and warming cycles is negligibly small (Fig. 1d, f).

2.2 Electrical Measurements

H&M also falsely claim the absence of a broadening of the superconducting transition under external magnetic fields in the R(T) measurements of hydrides, raising doubts about superconductivity in these compounds. Before beginning the discussion on this issue, we present details of electrical transport measurements, which are essential for the interpretation of superconductivity.

A desired superconducting sample is synthesized within the pressure range of its stability — most high- T_c hydrides require pressures over one megabar. For that, the DAC containing a piece of a precursor compound (e.g., chemical elements (S, La, Y) or their stable hydrides (LaH₃, YH₃)) is placed in a gas loader where it is clamped under hydrogen atmosphere at elevated pressures of 0.10–0.15 GPa. The sample is further pressurized to the required pressure value, at which it transforms into hydrogen-rich product via the diffusion of hydrogen into the precursor compound. The diffusion is a long process at room temperature, but it can be greatly accelerated by heating a sample in an oven (up to hundreds of degrees), or by a pulsed laser (up to thousands of degrees). This method has two important advantages: it yields solely binary products without impurities of third elements, and allows for a good control of the hydrogenation. Excess hydrogen can be easily monitored both optically and spectroscopically when hydrogen surrounds the final product. A much easier but less controlled method is the use of hydrogen containing materials such as BH₃NH₃ [21], AlH₃, or paraffins [37] as an alternative source of hydrogen. Upon heating, these compounds decompose and release free hydrogen. This technique was recently successfully introduced for the synthesis and studies of lanthanum superhydride by Somayazulu et al. [21]. We note that a long time ago this method was used by Ponyatovskii for the synthesis of hydrides in large-volume high-pressure cells [10, 38]. However, this method has its flaws. For instance, the decomposition products may contaminate the superconducting phase or even react with it. It is also difficult to control the amount of the released hydrogen, which is crucial for the reliable synthesis of the stoichiometric hydrogen-enriched phases that require excess hydrogen.

The electrical resistance is typically measured using a four-probe arrangement. This guarantees that the resistance of the electrical leads is excluded from the measurements — only the sample is examined (Fig. 2). The simplest four-probe arrangement for micrometer-size samples clamped between diamond anvils is the van der Pauw geometry [39, 40] (Fig. 2a, b). By alternating the combination of current and voltage contacts, one can measure resistance from different parts of the sample that provides information about its uniformity (Fig. 2c, d). In addition, by passing electrical current through opposite contacts under an applied magnetic field, one can pick up the Hall signal from another pair of contacts [40, 41].

The interpretation in terms of the emergence of superconductivity when the R(T) dependence sharply drops to zero below T_c is obvious for perfect samples. However, many real samples are contaminated by unreacted precursor compounds, contain impurity by-product phases (e.g., unsaturated lower hydrides), or are poorly crystallized. These imperfections are often unavoidable, since some tiny areas of a sample are not thoroughly heated in order to prevent the electrical leads from damage by a pulsed laser. In addition, samples of a larger size, which are not surrounded by a quasi-hydrostatic medium (e.g., excess hydrogen), have considerable pressure gradients. In such samples, the superconducting transition broadens and displays additional steps, indicating that the R(T) dependence is affected by non-uniform current flow (Fig. 2c-e). These distortions of the superconducting transition were also observed in other superconductors [42]. These interesting multiple resistance steps around the main transition were



Fig. 2 Typical four-probe electrical transport measurements in DACs. **a** Photo of the sample of Fm-3 $m-LaH_{10}$ phase with excess H_2 at 138 GPa. **b** SEM photo of four Ta/Au electrical leads spattered on the diamond culet before loading the sample. As an example, two out of four possible combinations of current and voltage contacts, which can be realized in the typical R(T) measurements are marked by blue and red. **c**-**e** R(T) measurements displaying the distortions of superconducting transitions (broadening, steps, and peaks) in the non-uniform samples. **c** Superconducting transition in Fm-3 $m-LaH_{10}$

not studied theoretically. An empirical model for MgB₂ [42] attributes these transitions to inhomogeneous doping (resulting in parts of the sample with different T_c values). The shape of R(T) is explained by a core–shell model: the outer layers or the shell of grains would correspond to a doped MgB₂ phase, which displays the lower T_c while the core of the grains still contains pure MgB₂ phase exhibiting the superconducting transition at higher temperatures.

Perhaps even more complicated is another kind of distortions of the superconducting transition: a peak in R(T)[43-45] — an anomalous increase of the resistance that precedes the sharp drop to zero (see Fig. 2c, d). This peak has been observed in many disordered superconductors [44] and has been the subject of many studies [46]. It was systematically studied, for instance, in boron-doped diamond [45, 47]. It was suggested to be due to the granular character of superconductivity: Cooper pairs first condense, and then localize into "bosonic islands," causing a sharp increase in the resistance as the islands are separated from each other [47]. At lower temperatures, percolation through these sizeincreasing islands leads to the drop in resistance down to zero. This model describes well the peak that is observed in hydrides (see Fig. 2c, d and, for instance, Ref. [43]). We repeatedly observed a peak in R(T) with a particular combination of van der Pauw contacts, whereas a normal sharp step is observed with another combination (Fig. 2c, d). Most likely, this reflects the non-uniformity of the

at 136 GPa: two combinations (green and black) show the broadened transitions with a T_{onset} of ~241 K, whereas other two combinations (red and blue) show a peak in R(T) that precedes the sharp drop to zero-resistance. **d** The broadening of the superconducting transition (red and blue) and additional steps in R(T) (black and green), indicating that the R(T) is affected by non-uniform current flow through the sample of $Im-3 m-D_3S$ at 154 GPa. **e** The suppression of distortions and bumps at the superconducting transition under external magnetic fields in the $Im-3 m-H_3S$ phase at 155 GPa (taken from Ref. [41])

superconducting phase in samples. It is clear that T_c should be defined as the temperature where the peak arises upon cooling (Fig. 2c and Ref. [43]), not as the temperature associated to the maximum of the peak. This anomalous peak in R(T) loses amplitude, broadening into a bump under applied magnetic fields [45, 46].

Importantly, not only the observation of zero resistance strongly supports superconductivity in hydrogen-rich compounds, but also the transition imperfections (broadening, steps, and peaks) discussed above, since these features are common among inhomogeneous superconductors.

2.3 Superconducting Transitions at High Magnetic Fields

Measurements of the superconducting transition in external magnetic fields independently verify the presence of superconductivity and additionally provide valuable information on the properties of superconductors. Whereas the magnetic field has often a small effect on the resistance of normal metals, a T_c is strongly suppressed as the magnetic field increases, and superconductivity can be completely suppressed at fields above an upper critical field, which is defined as H_{c2} for type-II superconductors. This suppression is difficult or impossible to achieve in high- T_c hydrides that have very high values of $H_{c2} \ge 100$ T [20, 41, 48]. Nonetheless, available laboratory magnetic fields in the order of ~ 10 T allow one to define the slope of $H_{c2}(T)$ near T_c and estimate the value of $H_{c2}(0)$ by extrapolating the values of the $H_{c2}(T)$ to lower temperatures. An accurate evaluation of H_{c2} and a study of $T_c(H)$ up to the record static fields up to 45 T was done at the NHMFL in Tallahassee using the hybrid magnet and up to 65 T in pulsed magnetic fields at the NHMFL in Los Alamos through the use of miniature nonmagnetic DACs. Experiments in pulsed magnetic fields are especially demanding due to the potential heating of the DAC and the sample due to eddy currents associated to fast varying magnetic fields. It turned out that the miniature DAC, which was specially designed for SQUID measurements (Fig. 3), is also suitable for magnetic fields [41, 48].

We often observe that the steps in R(T), which accompany the superconducting transition in imperfect samples, are strongly affected by the external magnetic field; e.g., they disappear above a modest field of just 3 T in H₃S [41] (Fig. SM1 in Ref [41]). Similar behavior was also observed in LaH₁₀ [20]. The distortions of the superconducting transition are suppressed under high magnetic fields in hydrogen-rich compounds as well as MgB₂ [20, 41, 42] (see Fig. 2e and also Fig. 2a in Ref. [20]). There are no systematic studies of these phenomena; however, we attributed them to inhomogeneities of the superconducting phase in samples, where superconductivity first sets across weak inter-grain links but is easily suppressed by magnetic field [41].

We note that despite the complicated shape of the R(T) dependence in non-uniform samples (bumps, steps, and broadening), the value of a T_c — the temperature corresponding to the emergence of superconductivity — remains nearly unchanged and is close to the value measured in perfect samples.



Fig. 3 The design of the miniature nonmagnetic DAC. A photo of the assembled miniature DAC (right) and its parts (left)

2.4 Broadening of the Superconducting Transition Under Applied Magnetic Fields

Previously, we discussed the different reasons for the broadening of superconducting transition: impurities, poor crystallinity, pressure gradients, inhomogeneity of the superconducting phase in samples, etc. A special non-trivial case is the broadening of a superconducting transition in perfect type-II superconductors under strong magnetic fields. Magnetic field penetrates the type-II superconductor in the form of vortices, and vortex motion associated to the electrical currents produces dissipation. As a result, the width of the resistive transition ΔT is expected to increase with the field as ~ $H^{2/3}$ (Ref. [49]). Indeed, we observed the broadening of the superconducting transitions under magnetic fields (Fig. 4). Naturally, the broadening can be observed only if the superconducting transition is sharp enough in zero-field. Also, one should keep in mind that the vortex dissipation that leads to the broadening of the transition depends on vortex pinning mechanisms and vortex phase diagram. For instance, in cuprates the existence of a vortex liquid phase leads to a dramatic broadening of the superconducting transition under an external magnetic field [50] whereas the smaller broadening was observed for iron-based superconductors [51].

However, in Refs. [4, 52] it is argued that the high-temperature superconducting hydrides display no broadening of the superconducting transition under magnetic fields. Unfortunately, the authors manipulated the published data and cherry-picked the references that include samples exhibiting broad superconducting transitions already at zero-field. For these samples, it is difficult to detect the magnetic field-induced broadening. H&M ignored the publications, which contradict their beliefs, i.e., the initially sharper superconducting transitions that broaden in applied fields, e.g., Ref. [41]. Therefore, the arguments of H&M in Refs. [4, 52] against superconductivity in hydrides, which are largely based on their claims on nonstandard absence of the broadening of the superconducting transitions in external magnetic fields, are false. The broadening is indeed observed and is consistent with the well-established scenario where the hydrides are type-II superconductors [1].

There is an apparent exception: Snider et al. [24] reported no broadening of the superconducting transition under magnetic fields in their experiments (Fig. 4). This is particularly puzzling because the transitions are extremely sharp $\Delta T < 0.5 \text{ K} (\Delta T/T_c < 0.2\%)$, even narrower than those in MgB₂ where the broadening is pronounced [42, 53]. Unfortunately, H&M [4] groundlessly extrapolated this controversial result to all other hydrides.





Fig. 4 Broadening of the superconducting transition under external magnetic fields in different superconducting compounds: **a** SnH_{12} [68], MgB₂ [53], CeH_x [69], H₃S [41], C-S-H [24]. The related coordinates are chosen for easier comparison. The solid lines are a guide

2.5 Magnetic Susceptibility Measurements: Observation of the Magnetic Field Screening

A major claim of H&M is the "absence of magnetic evidence for superconductivity in hydrides under high pressure." The authors question the measurements of the magnetic moment of H₃S samples as a function of both temperature M(T) and magnetic field M(H) [1]. Here, we will elucidate these experiments.

Magnetic susceptibility measurements under very high pressures are very challenging, and in fact much more difficult than the electrical transport measurements. The diameter of a superconducting sample at megabar pressures is smaller than $\sim 100 \,\mu\text{m}$ and its thickness is just a few μm . Such small superconducting disk has a magnetic moment of just ~ 10^{-6} - 10^{-7} emu that is quite small, and in fact comparable to the sensitivity threshold of a SQUID magnetometer (~ 10^{-8} emu). Nevertheless, it was possible to perform such measurements even at megabar pressures using the miniature DAC specially designed for this purpose (Fig. 3). The body of the DAC is made of a high-purity Cu-Ti alloy with 3 wt % Ti in order to minimize the magnetic signal over a wide temperature range and provide a high mechanical strength. The piston and diamond seats, which are subjected to the highest load, were made of the harder Cu-Be alloy with 1.8-2.0 wt % Be. Such combination of materials allows us to construct the miniature DAC with an outer diameter of 8.8 mm, which is capable of reaching pressures as high as 220 GPa retaining low overall magnetic response.

to the eye for $\Delta T(H) \sim H^{2/3}$ broadening expected at high fields [49]. **b** Broadening of the superconducting transition derived in Ref. [4] is compared with the present study (blue points)

So far, measurements of the magnetic susceptibility in a SQUID magnetometer were reported only in the original work [1], for both H₃S and D₃S (see Fig. 5a, b and Supplementary Fig. 6d in Ref. [1]). A sharp step was observed in the magnetization curve M(T) under zero-field cooled, ZFC, condition at 203 K for H₃S and at 136 K for D₃S counterpart. It coincides with the superconducting step observed in the R(T) data for both H₃S and D₃S, and thus provides clear evidence for superconductivity in both compounds. In H₃S ZFC and field-cooled, FC, M(T) plots are different — the ZFC curve displays a sharp step, but practically no kink is observed under FC conditions, as the magnetic flux penetrates the material (Fig. 5a). However, some samples still display a small signature for the superconducting transition when measured under FC conditions (Fig. 5b).

The major obstacle for proper magnetic susceptibility measurements is a background from the DAC, which is still much larger than the signal from the sample. It was not possible to subtract the background of the DAC in the usual way by measuring the DAC with and without the sample. Instead, the magnetic susceptibility was measured just above T_c where the sample is in its normal metallic state [1]. This magnetic signal was used as a background for subsequent measurements in SQUID magnetometry. Fortunately, at high temperatures in the vicinity of 200 K, the magnetic susceptibility of the materials of the DAC body displays a weak dependence on temperature. Apparently, subtraction of a background measured at a single temperature point should not be universal in the whole temperature range below T_c ,



Fig. 5 Magnetization measurements under high pressures using the miniature DAC. **a** M(T) measurements of H₃S at 155 GPa (taken from Ref. [1]). **b** More typical, the smaller magnetic signal is registered from the H₃S sample at a pressure of 135 GPa. The measurements of the same sample in another run are shown in the inset. **c**–**h** The recent improved measurements of M(T) under high pressure with accurate background subtraction (taken from Ref. [55]). **c**, **e**, **g** M(T) data of the *Fm*-3 *m*-LaH₁₀ at 130 ± 8 GPa under magnetic field H=10

mT: **c** M(T) of the sandwiched sample with LaH₃ and NH₃BH₃ before laser heating; **e** the heated sample with the superconducting *Fm-3 m-*LaH₁₀ phase; and **g** the difference plot. **d**, **f**, **h** M(T) data of the *Im-3 m-*H₃S at 155±5 GPa under magnetic field H=4 mT: **d** M(T) of the sandwiched sample with S and NH₃BH₃ before laser heating; **f** the heated sample with the superconducting *Im-3 m-*H₃S phase; and **h** the difference plot. The black and red circles correspond to ZFC and FC measurements. Blue lines represent smoothed FC data

especially at low temperatures. Nevertheless, this approach allowed us to observe the superconducting transition in the M(T) measurements. The absolute value of the diamagnetic transition of ~ 10⁻⁶ emu (Fig. 5a) corresponds to the signal from a thin diamagnetic disk with a diameter of ~ 100 µm which is comprable to the sample size estimated by optical microscopy. Nevertheless, the subtraction of the background was not perfect and therefore the M(T) curve is shifted vertically due to a paramagnetic component (Fig. 5a), and this was criticized in Ref. [54]. However, this superconducting transition fully agrees with the recent and more accurate measurements [55] (Fig. 5c–h).

Measurements of hysteresis in M(H) reported in Ref. [1] are even more difficult because the magnetic susceptibility of the materials of the DAC body, as well as copper wires, solder and electrical leads mounted for the corroborative resistance measurements, depends on a magnetic field as well. A lower critical magnetic field H_{cl} of ~ 30 mT was only roughly estimated as the point associated to the bending of the M(H) hysteretic loops without considering the geometry of the sample (demagnetization factor) (Fig. 4c, d in Ref. [1]). However, the correct value of the inflection point in the M(H) virgin curves was determined in the recent work [55] and its value is higher ~96 mT at 0 K for H_3S . This is not a large disagreement keeping in mind the absence of M(H)virgin curves in Ref. [1] and related uncertainties associated with the criterion of the deviation of M(H) from linearity in hysteretic loops. Even for the less experimentally challenging magnetic measurements at ambient pressure conditions, one can see factor 2–3 difference among different publications for the penetration depth (that results in even larger spread of the H_{cl} values) evaluated for the same superconductors [56].

The original magnetic measurements [1] were criticized in Ref. [4] and even qualified as a "myth" [5] given that they had not been reproduced for 6 years. In fact, we have a number of unpublished results, which agree with the Ref. [1] (Fig. 5b). Here, we present one of such results in Fig. 6, which was obtained right after the publication of Ref. [1]. We attempted to evaluate the lower critical field H_{c1} using the virgin curve of M(H). The sample was prepared according to the original recipe, i.e., through the pressure-induced disproportionation of compressed H₂S, and subsequently pressurized to $P \sim 140$ GPa. We did not characterize the sample by X-ray diffraction; however, the T_c value of ~ 140 K, which was observed in the ZFC M(T) measurements





Fig. 6 Magnetization measurements of superconducting H₃S at pressure $P \sim 140$ GPa. **a** M(T) data measured at H=3 mT after subtraction of the background collected at 250 K. **b** M(H) data measured at 100 K showing the initial virgin curve and hysteretic loop on alternat-

(Fig. 6a) suggests that the sample had the rhombohedral distorted crystal lattice or the R3m phase of H₃S, according to the later established dependence $T_c(P)$ (Fig. 3 in Ref. [14]). Figure 6b shows magnetization measurements of the H₃S sample at T = 100 K. The measurements were performed in the background subtraction mode using the reference data measured from the same DAC at T=210 K. Original hysteretic loop shown in the inset in Fig. 6b reveals a strong paramagnetic component originating mainly from the DAC. After subtraction of this paramagnetic background, which is approximated by linear dependence M(H), the superconducting hysteresis has the typical shape for a type-II superconductor (Fig. 6b). At H = 30 mT the M(H) virgin curve starts to deviate from a straight line allowing the determination of a lower critical field H_{cl} . Taking into account a demagnetization correction $\frac{1}{1-N} \sim 9$ for a thin disk shaped sample, which was estimated from the measured value of ΔM , we obtain H_{c1} (100 K) = 270 mT. The extrapolation of H_{c1} to lower temperatures using the equation $H_{c1}(T) = H_{c1}(0K)$ • $(1 - (\frac{T}{T})^2)$ yields the value $H_{cl}(0 \text{ K}) \sim 550 \text{ mT}$. The ZFC M(H) dependence coincides with the full magnetization loop at $H \approx 0.2$ T. Considering also the demagnetization factor, one can estimate the full penetration field $H_p(100)$ K \approx 1.8 T. Note that the hysteretic loop has an asymmetric shape and is broader than that in Ref. [1]. The hysteresis asymmetry results from the polycrystalline nature of the sample. According to Refs. [57, 58], the pronounced asymmetry is observed when the grain size $R < 10\lambda_L$ (λ_L is the London

ing the magnetic field. Inset shows the original hysteretic loop before subtraction of the background signal originating from the DAC at 210 $\rm K$

penetration depth). One can expect more symmetric hysteretic curves for samples of better quality with a larger grain size. The dependence of the superconducting hysteresis on the grain size was systematically studied in MgB_2 in Refs. [53, 59].

For a long time we did not clearly appreciate the role played by the quality and integrity of the samples. Even when the sample has a large enough size and clearly shows superconductivity in electrical transport measurements, the magnetic susceptibility signal can turn out to be elusive or below the sensitivity of the SQUID magnetometer. The reason for that can be the granular or non-uniform distribution of the superconducting phase in samples. The electrical current finds a continuous path through superconducting grains and metallic grain boundaries in the transport measurements whereas much smaller thin superconducting grains have a relatively small superconducting volume leading to a smaller signal due to the demagnetization factor.

Only recently, we succeeded in improving our measurements of the magnetic susceptibility significantly [55] (see Fig. 5c–h). For that we used a solid BH_3NH_3 as an alternative source of hydrogen instead of pure hydrogen. This allowed us to produce large samples with a diameter close to that of the culet of the diamond anvils, and to obtain a pronounced diamagnetic signal from superconducting phases under high pressures. We believe that this approach can be adopted by other laboratories.

We were able to extract the small diamagnetic signal of a superconductor from the measured overall magnetic moment, which includes magnetic moments of the DAC body, diamonds, and rhenium gasket, using a different approach. First, we measured the magnetic signal of DACs with the starting pressurized precursors (S or LaH₃ and BH₃NH₃) before laser heating. Then, we subtracted these reference data from the magnetic moment collected from the same DACs after laser heating and chemical synthesis and obtained the superconducting response of the H₃S and LaH₁₀ compounds. The temperature dependence of the magnetic moment of both samples changed dramatically after laser heating, indicating that they became superconducting (Fig. 5c-h). We note that Ref. [6] incorrectly states that ZFC and FC curves for H₃S coincide at 200 K before and after heating, which is clearly not the case for Fig. 5d, f. Abrupt steps in the ZFC M(T) curves yield values of $T_c \sim 231$ K for *Fm-3 m*-LaH₁₀ at 130(8) GPa and ~ 196 K for *Im-3 m*-H₃S at 155(5) GPa. These values are in excellent agreement with the previously reported values from four-probe electrical transport measurements under the same pressures [1, 14, 18, 41, 48]. FC curves show the small step for the Im-3 m-H₃S sample (Fig. 5h) and the subtle step for the Fm-3 m-LaH₁₀ samples. The subtle steps observed on FC curves or their apparent absence are common for superconductors with strong pinning of vortices [60].

In addition, we determined lower critical fields and consequently the London penetration depth for both H₃S and LaH₁₀ [55]. For H₃S, a lower critical field $H_{cl}(0 \text{ K})$ is estimated to be ~ 0.74-1.09 T, leading to the London penetration depth $\lambda_L(0K) \sim 18-23$ nm. For LaH₁₀, the lower critical field $H_{cl}(0 \text{ K})$ is estimated to be ~ 0.42–1.75 T and $\lambda_L(0K)$ ~14–35 nm. The coherence length, $\xi(0K)$, was estimated from our previous measurements of the dependence of T_c on external magnetic fields: it is 1.84 nm for H_3S [41] and 1.51 nm for La H_{10} [48]. These values yield the Ginzburg–Landau parameter $\kappa = \frac{\lambda}{\epsilon} \sim 10-13$ for H₃S and ~ 10-23 for LaH₁₀. These parameters indicate that both H₃S and LaH₁₀ compounds belong to the category of "moderate" type-II superconductors, rather than being hard superconductors as would be intuitively expected from their high T_c s. One can see that the coherence length is comparable to those of other high-temperature superconductors, e.g., the cuprates, but the London penetration depth is significantly shorter [56].

2.6 Critical Current Density

Using the values of the thermodynamic critical field, $H_c(0 \text{ K}) = 5.6 \text{ T}$ for H₃S and 5.1 T for LaH₁₀, and the London penetration depth, λ_L (0 K), we can evaluate the depairing critical current density as $j_d \sim H_c/\mu_0 \lambda$. This yields values of $j_d \sim 1.4 \times 10^{10} \text{ A cm}^{-2}$ and $\sim 7.4 \times 10^9 \text{ A cm}^{-2}$ for H₃S and LaH₁₀, respectively, which are approximately two orders of magnitude higher than the $j_d(0 \text{ K})$ extracted for YBa₂Cu₃O₇₋₈ [61] and three orders of magnitude higher than those of MgB₂ [62] or V₃Si [63]. It has to be mentioned that the depairing critical current density represents the theoretical maximum or the critical current density and is usually found to be of nearly one order of magnitude higher than the actual depinning critical current density in a number of materials.

The experimentally evaluated critical current densities from magnetization measurements are significantly lower values ~ 7×10^6 A cm⁻² at 100 K for H₃S and LaH₁₀ [55]. We note that the recent criticism of the estimate incorrectly ignores substantial demagnetization factors [6]. The electrical transport current–voltage measurements yield similar order of magnitude values, viz, ~ 2.6×10^7 A cm⁻² for YH₆ [43] and ~ 2.7×10^6 A cm⁻² at 4.2 K for La(Y)H₁₀ [64]. This could suggest that there is room for improvement of the critical currents in the hydrides.

2.7 Magnetic Field Screening in the Nuclear Resonance Scattering Experiments [11]. Numerical Analysis

Further evidence for superconductivity was demonstrated on the screening of the magnetic field by H₂S compressed under 153 GPa in Ref. [11]. In this work, a thin ¹¹⁹Sn film was inserted inside the H₃S sample, and the magnetic field in the film was monitored via nuclear resonance scattering of synchrotron radiation. In that experiment, an external magnetic field of about 0.7 T was shielded by a H₃S superconductor at least up to 140 K. Importantly, the same field was attenuated only to half of this value when it was applied along the sample. Considering nearly zero demagnetization factor in this configuration, it cannot be understood if the sample around Sn foil was not superconducting. This experiment was questioned in Refs. [4, 5, 7, 54] where the authors claim that the magnetic field should not have been expelled. We note that magnetic field expulsion (i.e., in a FC experiment) is very different from the magnetic field screening (i.e., a ZFC experiment when magnetic field is applied at low temperature after cooling in zero field).

The ability to shield a magnetic field by a type-II superconductor involves Meissner-London screening at the sample edge [65] and penetration of Abrikosov's vortices forming a Bean critical state [66]. Unlike any other magnetic material, the latter is non-local on the scale of the whole sample. We therefore considered two orientations of an applied field with respect to the sample assembly. A careful analysis of both phenomena [67] in the experimental geometry of [11] suggests that the average magnetic field measured in the interior cavity of a superconductor in two orientations is fully consistent with the conventional behavior of a type-II superconductor with a lower critical field, $0.3 \le H_{cl} \le 0.6$ T and critical current density larger than $j_c \sim 4 \times 10^6$ A cm⁻². Considering large uncertainties of the estimates, these values are in a good agreement with the estimates obtained from the magnetization measurements discussed above. On the other hand, the difference found in two orientations cannot be understood assuming any other non-superconducting state of the studied sample.

Based on a detailed analysis of the experiment discussed in Ref. [11], it appears that the nuclear resonance scattering is a new, non-trivial, and sophisticated technique to detect superconductivity. Hopefully it will find further use to study novel near room-temperature superconductors in difficult conditions, such as ultra-high pressure, and perhaps in compounds exhibiting superconductivity even above room temperature. A full analysis of this experiment is given elsewhere [67].

3 Conclusions

To conclude, we presented a detailed description of experiments such as the observation of zero resistance that is concomitant with the onset of the pronounced diamagnetic response, as well as the broadening of the resistive transition under external applied fields. The observed temperature–magnetic-field phase diagram is akin to those observed in orbitally limited superconductors. These observations represent standard as well as very solid evidence for hightemperature superconductivity in hydrogen-rich compounds under high pressures and disavow the doubts raised in Refs. [3–7]. The pronounced isotope effect in hydrides is in favor of conventional superconductivity, but contradicts the hole superconducting mechanism [8], and in the essence eliminate this theory, as this declared in Ref. [3].

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Author Contribution The group in Mainz (M.I.E., V.S.M., A.P.D., P.P.K.) synthesized samples and performed electrical transport measurements; F.F.B., D.S., S.M., and L.B. performed measurements of electrical resistance under high magnetic fields; V.S.M, V.K., and S.I.S. performed magnetic measurements in SQUID; F.F.B., S.L.B., V.S.M., and V.K. analyzed magnetization data; R.P. contributed finite element modelling. All authors collaborated in the discussion of the manuscript. M.I.E. wrote the manuscript along with V.S.M., F.F.B., S.L.B., V.K., and R.P.

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