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Retraction: Carbon content drives high temperature superconductivity in a carbonaceous sulfur hydride below 100 GPa

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Retraction of 'Carbon content drives high temperature superconductivity in a carbonaceous sulfur hydride below 100 GPa' by G. Alexander Smith et al., Chem. Commun., 2022, 58, 9064-9067, https:// doi.org/10.1039/D2CC03170A

We, the named authors, hereby wholly retract this Chemical Communications article based on our concerns over the origins of the electrical transport measurements presented. While the validity of the X-ray crystallographic study and structure calculations of carbonaceous sulfur hydride (C-S-H) are maintained, we have lost confidence in the origin of the electrical transport measurements, and therefore all conclusions deduced from the electric measurements, including the superconductivity properties are uncertain. Therefore, this article is being retracted to avoid misleading readers and to protect the accuracy and integrity of the scientific record. We regret any confusion or inconvenience caused to the scientific community.

Ranga P. Dias was contacted but did not respond.

Signed: G. Alexander Smith, Ines E. Collings, Elliot Snider, Dean Smith, Sylvain Petitgirard, Jesse S. Smith, Melanie White, Elyse Jones, Paul Ellison, Keith V. Lawler and Ashkan Salamat, 22nd December 2023.

Retraction endorsed by Richard Kelly, Executive Editor, Chemical Communications.

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Carbon content drives high temperature superconductivity in a carbonaceous sulfur hydride below 100 GPa†

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We report a previously unobserved superconducting state of the photosynthesized carbonaceous sulfur hydride (C-S-H) system with a maximum $T_{\rm C}$ of 191(1) K below 100 GPa. The properties of C-S-H are dependent on carbon content, and X-ray diffraction and simulations reveal the system remains molecular-like up to 100 GPa.

The superhydride superconductor is envisioned as a hydrogen dominant alloy which lowers the pressure required to achieve the favorable properties and high- T_c predicted for dense met hydrogen. Hydrogen within these alloys takes part in an extend bonding network, be it the purely hydrogenic clathrate a metal superhydride, or a covalent network with other in H₃S.²⁻⁴ There have been record breaking miles covalent superhydrides, including a 203 K T_c r hydride and a 288 K T_c at 267 GPa in carbonaceous C-S-H was first synthesized from elemental plantsors at 4 GPa, and then compressed without the all annealing ato its final athy likely leading to metareported superconducting state, nthes d by reacting elestable states. C-S-H has since be nciple, this method mental S and CH₄-H₂ flui tures permits greater control C co centration, although the reported

C-H Raman modere parably weak, and whether it leads to oe st ed. From either synthetic route, high- T_c states a rich ph agram below 100 GPa where evidence C-S-H disp ntion of molecular-like packing as well as points to a ion.^{8,9}

√hile the exact identity of the record-breaking C-S-H material vet to be iscerned, candidate structures have been proposed ructure prediction (CSP) and virtual crystal approximation simulations. 10-12 Many of the CSP candidates for C-S-H are lar or exhibit a molecular sub-unit, including the leading candidates with CH4 intercalating or replacing an H3S unit within the H₃S perovskite-like lattice. 10-13 While these low-dimensional sub-units seem contrary to the extended bonding network, our recent simulations showed that dispersion interactions can potentially be important in covalent superhydrides with such sub-units. 14 Along these lines, it was recently discovered that a metal superhydride with a relatively low hydrogen concentration, YH6, exhibits an anomalously high T_c at lower pressures than its more studied higher hydrogen content counterparts.¹⁵

Building on the high T_c reported at 100 s of GPa for the superhydrides, the next goal towards achieving ambient superconductivity is to lower the critical pressure required to form superconducting phases. 16 Herein, we investigate C-S-H below 100 GPa to probe for lower-pressure superconducting states predicted by CSP, and to further understand the consequences of the thermodynamic pathway for synthesizing C-S-H from elemental precursors. We present electrical transport measurements in this previously unexplored pressure regime that reveal a remarkably high T_c in some crystals, raising the question as to how these macroscopic quantum states emerge over such dramatically different *P–T* ranges. Synchrotron single crystal X-ray diffraction (SC-XRD) identifies structural evolution of C-S-H up to 100 GPa and Raman spectroscopy shows that the C content in C-S-H produced by photochemistry varies in each crystal synthesised. That variation directly affects the material properties with subtle differences in packing densities. Density functional theory (DFT) assists in understanding the H positions of the determined phases.

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All crystals of C-S-H here are synthesized using the procedure of Snider et al.8 (full details in ESI†). Ball-milled mixtures of elemental C and S with dimensions about 15% of the diamond culet (typically 100-250 µm) are placed into the sample chamber of a diamond anvil cell, along with a ruby sphere. Gas phase H₂ is loaded at 0.3 GPa. Samples are then pressurized to 3.7-4.0 GPa and excited for several hours using light from a 514 nm laser with power ranging from 10 and 150 mW depending on sample response. Crystal growth is monitored in situ by visual observation, and Raman spectroscopy confirms the transformation into C–S–H by the presence of characteristic C-H, S-H, and H-H Raman modes. Samples are compressed to 10 GPa after transformation and characterization by Raman spectroscopy to avoid decomposition.

We performed electrical transport measurements on 3 crystals of C-S-H - Runs T1, T2, and TN - following the methods described in Snider et al.8 (Fig. 1). In 2 separate runs, we observe maximum T_c of 191(1) K at 97(5) GPa (Run T1, Fig. 1a and b) and 188(1) K at 98(5) GPa (Run T2, Fig. 1b). These transitions occur at roughly half the pressure required to achieve a similar T_c in either C-S-H or S-H/S-D. 8,17 Runs T1 and T2 are contrasted with Run TN, which exhibits no superconducting transition at 90(5) GPa on cooling to 10(1) K, despite exhibiting metallic character (Fig. 1a inset). The shape of the T_c

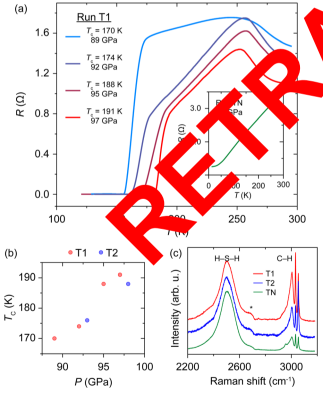


Fig. 1 (a) Resistance response of C-S-H (Run T1) on cooling, displaying a superconducting transition at 191 K at 97 GPa. (Inset) R response from Run TN at 90 GPa showing metallic behavior. (b) Evolution of T_c with P for Runs T1 and T2. (c) Comparative Raman spectra of Runs T1 and T2, and Run TN at 4.0 GPa and 300 K. The feature marked with an asterisk (*) is secondorder Raman scattering from diamond

vs. pressure (Fig. 1b) implies this superconductivity comes from a distinct phase than that at 267 GPa. Also observed in Run T1 is the previously noted behavior of C-S-H to exhibit narrowing $\Delta T/T_{\rm c}$ as a function of increasing pressure and $T_{\rm c}$, exhibiting a minimum $\Delta T/T_c$ of 0.0373 at 97 GPa (data in ESI†).

By virtue of our focus on the lower pressure phases of the C-S-H ternary, the samples used in this study are significantly larger than those in Snider et al., by a factor of 3-10, and these larger crystals have a heterogeneous C concentration compared with crystals from our previous work. This inhomogeneity is evidenced by variations in the relative intensities of Raman modes originating from C-H stretches around 3000 cm⁻¹ and H-S-H bends around 2500 cm⁻¹, i.e. I_{C-H}/I_{H-S-H} . Fig. 1c shows representative Raman spectra of C-S-H crystals from each of the three runs following their initial synthesis at 4 GPa, with variations in I_{C-H}/I_{H-S-H} evident. Run TN, which did not exhibit a superconducting kion at 90(5) GPa, has an intensity ratio I_{C-H}/I_{H-S-H} of $\sqrt{7}$. Mean ile, Runs T1 and T2 have $I_{\text{C-H}}/I_{\text{H-S-H}}$ of 1.16 and 93, pective It is important to note that even our Run Z nas a hig I_{H}/I_{H-S-H} than the samples et which become superconducting at reported in Snid room tempera com $\epsilon_{\text{CSSion}} (I_{\text{C-H}}/I_{\text{H-S-H}} = 0.08)$. Thus, centratio. e C-S-H ternary system is linked to a increased @ on in the pressure required to reach the supersignificant redu g regime

ach of the R(T) responses at the different pressures measured Run Timature a turning point around 250 K (Fig. 1a). At these –H exhibits the temperature response of a finite gap con ystem, whereas below 250 K the temperature response is metallic. havior likely results from either a structural or electronic phase transition. An electronic transition would not likely be accompanied by a change in symmetry, and a structural transition in a hydride material might also be indistinguishable using XRD if the heavy atom sublattice does not re-order, as is the case for the R3m to $Im\bar{3}m$ transition in H_3S .⁷ Resistance continues to decrease with lowering temperature before a sharp drop to zero resistance as the critical temperature is crossed. Such a difference in T_c to that of Snider et al.8 could be expected, as their thermodynamic approach to a superconducting state begins from cooling in the recently confirmed $Im\bar{3}m$ phase emerging above 159 GPa⁹ rather than the previously reported phase IV.8

SC-XRD measurements on other crystals were conducted at HPCAT with $\lambda = 0.34453$ Å. Conical diamonds with 80° apertures were used for greater completeness in SC-XRD. Fig. 2 shows the P-V response of 8 C-S-H crystals from 3 separate runs, with all data on phase III/IV collected during Run X2. 2nd -order Birch-Murnaghan equations of state are fit to each crystal and phase (values in Table SI, ESI†). We observe subtle systematic differences in V-P relations across the different crystals measured at the same thermodynamic conditions. The largest difference in V is 2.9% at 28.9(5) GPa in Run X2 between crystals C1 and C4. K0 was found to range between 7.32 and 14.50 GPa for Runs X1 C3 and X2 C3. V trends for all of the C-S-H crystals measured are equal or lower than that of our own measurements on pure H2S + H2, which in turn is noticeably lower than that reported for C-S-H prepared from mixtures of molecular gases.¹⁸ This, along with differences in ChemComm Communication

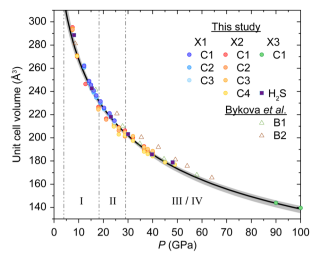


Fig. 2 P-V relations of C-S-H at 300 K compared with values from H₂S and d Bykova et al. 18 A 2nd order Birch-Murnaghan equation of state was fit with initial volume $V_0 = 400.573 \text{ Å}^3$ and bulk modulus $K_0 = 11.028 \text{ GPa}$ (black line), and the gray area denotes uncertainty derived from high and low bands for Runs X1 and X2. Phase division for I $(I4/mcm) \rightarrow II (C2/c) \rightarrow II$ III/IV (I4/mcm) are taken from Snider et al.8

the electronic response between crystals measured here and in Snider et al., suggests a large variability in C-S-H stoichiometry generated by photochemistry under pressure.

Leading up to 18 GPa, SC-XRD measurements confirm phase I⁸ as the Al₂ Cu-type structure (I4/mcm) previously identified in CH₄-H₂ and H₂S-H₂ mixtures. The I4/mcm phase inferred between 4 to 9 GPa as no change is observed by Ram spectroscopy. Due to insufficient C concentration crystallographic placements, SC-XRD measureme nable to resolve between C and S on the 8 h Wyckoff Fig. 3a displays only H₂S units on the 8 ying the Bernal-Fowler "ice rules" to determine I positio. 14/mcm of the H₂S molecular units results in tially occupied 16k Wyckoff positions, and this strains the units to be planar within {002} in sobel et al. 20

A CSP study on the H-S syste. entifica a P1 modification dons owing to out-ofwhich mostly varies from sub-units.6 Comparing several plane rotation of the olecv (keeping the lattice and S planar arrangements of

positions fixed at their experimental values) versus the arrangement of the P1 structure with DFT and the vdW-DF2 functional shows a ~0.44 eV preference for a non-planar H arrangement. 22 This indicates C-S-H will have non-planar arrangements of H₂S molecular units to facilitate interactions between the shorter 3.30 Å interplane nearest neighbor S atoms. The magnitude of the enthalpic differences shows weak packing forces that could enable the molecular sub-units to behave as weakly constrained rotors within their respective molecular volume when thermalized. Given the orientational preference in the interplane direction and the S-S nearest neighbor distances being within the van der Waals and H₂S dimer H-bonding distances, ²³ there is at least some weak H-bonding contributing to the cohesion of the lattice along with the primary van der Waals forces.

Above 18 GPa, C-S-H transforms into a C2/c phase (Fig. 3b). This transition was observed crystals of Run X1 and in $H_2S + H_2$, but was not perfect in Cand C4 of Run X2. The absence of C2/c-type C-H some stals is consistent with a et al. 18 ncharov et al., where the observations in Byk phase is observe only a crystals with low C content, and varion in stoichiometry in C–S–H further exem It is worth noting the similarities otochem. formed by tochem. It is worth noting the similarities structure of C–S–H and previously documenbetween The C. tures of The Cccm H–S structure from Duan et al. 6 referred by Pace et al.²⁴ owing to its H–S–H network additional distinct environment for molecular p. H_2 ch is reflected in the Raman vibron. This and the I222 structure reported by Strobel et al. 20 differ from the C2/c re only in the orientation of H₂S sub-units and apparent directionality of the H bonding network. The C2/c phase resembles a monoclinically-distorted version of the I4/mcm phase where the [101] direction of the C2/c structure roughly corresponds to the [001] direction of the I4/mcm structure. In both cases, that direction resembles a 2-dimensional pore formed by S atoms interconnected by inter-plane H-bonding that encapsulates the H2 molecules, and the views shown in Fig. 3 are oriented to look along these pores. The H positions determined by SC-XRD are reminiscent of the 9 GPa structural optimizations.

C-S-H transforms back into an I4/mcm structure around 29 GPa (Fig. 3c) which persists to our highest measurements at

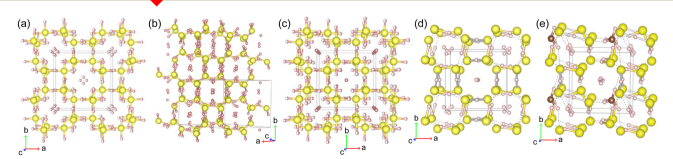


Fig. 3 SC-XRD determined structure at (a) 9 GPa /4/mcm (b) 29 GPa C2/c and (c) 50 GPa /4/mcm C-S-H. (d) DFT derived structure at 90 GPa – bicolor cylinders represent bonds (≤1.43 Å), silver cylinders represent H atoms shared between two heavy atoms (1.43-1.53 Å), and dashed lines represent H bonds (1.53–2.0 Å). (e) Lowest enthalpy structure found here when substituting a CH_4 for an H_2S in the 90 GPa structure shown in (d). Yellow spheres represent S throughout, brown spheres C, and pink spheres H.

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100 GPa. Our measured phase transitions by SC-XRD agree well with those reported in Raman studies.8 The H positions of the H₂S units are again best modeled crystallographically within the constraint of the I4/mcm group to be in a planar configuration. However, DFT dictates that orientations with out-ofplane H positions are 5-7 eV more enthalpically favorable, and the lowest enthalpy configuration found here (structure in ESI,† but like Fig. 3d) shows a H-bonding network creating 2 dimensional channels along [001].

The previous reported transformation from phase III to IV around 45 GPa or metallization above 60 GPa are not distinguished by SC-XRD as the structural solution remains 14/mcm up to our highest measurement at 100 GPa. Optimizing the lowest enthalpy 50 GPa configuration using the lattice and S positions determined by SC-XRD at 90 GPa shows a H-bond symmetrization along [001] as in Im3m H₃S (Fig. 3d).⁷ Other configurations were evaluated confirming the structure with zig-zag H-bonding along [001] is the most enthalpically favorable at 90 GPa. This marks a transition from a double- to a single-well potential for those H atoms, and is accompanied by a significant drop in band gap (of the S and H only system) from 1.99 eV at 50 GPa to 0.25 eV at 90 GPa. Thus, the transition from phase III to IV is a transition from H-bonding to covalency which drives metallization. It should be noted that planar configurations considered at 50 and 90 GPa are metallic, so any H₂S molecules metastably trapped in planar orientations could drive metallization sooner than the double- to single-well transition.

A prototypical carbonaceous model can be created by substituting one of the H2S molecules of Fig. 3d with a CH4 mole Optimizing the H positions of that model shows a disruption to zig-zag S-H-S network along [001] in the vicinity (structure in ESI†), coupled with a reduction of metal to the S-H system. The lowest enthalpy structure the band gap to 1.36 eV but does orient the 4 to 1 linkages reminiscent of those seen in R3m CSI higher (structure in ESI†) structure rotates the CM₄ such that the adjacent H₂S molecules are more like Fig. 30 ccompanied a ~0.27 eV lower bad gap. While a metallic reffice of this model was not identified here, these results sugge the ning points of the rient ordering and H-bond R(T) curves in Fig. 1a arise symmetrization within C-S sample

In conclusion, new surements on C-S-H with greater C content show a sition to a superconducting state with maximum T_c of 191 K at GPa – significantly lower than previously observed. SC-XRD confirms a phase evolution of I4/ mcm to C2/c to I4/mcm in crystals with lower C content, while more carbonated crystals bypass the monoclinic phase. The absence of an measurable transition from phase III to IV seen in earlier Raman studies indicates that the transition is likely a reordering of the H which leaves the S sublattice unchanged, which is supported by DFT simulations. That greater C content inihibits the formation of monoclinic C-S-H, but also promotes a transition to a superconducting state at significantly lower pressures is worthy of further study, and a major challenge for the study of C-S-H is to ensure control of the product and controllable concentration of the constituent elements during the photo-induced reaction.

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Conflicts of interest

There are no conflicts to declare

Notes and references

- 1 N. W. Ashcroft, Phys ev. Leti , 187002.
- 2 C. J. Pickard, I. E a and M. I. ts, Annu. Rev. Conden. Mater. 76. Phys., 2020, 11,
- brock mmon, R. McBride, X. Wang, 3 E. Snider, . V. Law rek, A. Salamat and R. P. Dias, Phys. N. Meyers Rev. Let 1, 126, 11
- a, J. Contreras-García and I. Errea, Nat. Commun., 4 F. Bell, T. N
- Drozdov, M. . Eremets, I. A. Troyan, V. Ksenofontov and . I. Shylip *Nature*, 2015, **525**, 73–76.
- Liu, F. Tian, D. Li, X. Huang, Z. Zhao, H. Yu, B. Liu, . Duan, Tian 🛭 T. Cui, *Sci. Rep.*, 2014, **4**, 6968.
- . Calandra, C. J. Pickard, J. R. Nelson, R. J. Needs, Y. Li, H. Liu, Y. Zhang, Y. Ma and F. Mauri, *Nature*, 2016, **532**, 81–84.
- nider, N. Dasenbrock-Gammon, R. McBride, M. Debessai, H. Vindana, K. Vencatasamy, K. Lawler, A. Salamat and R. Dias, Nature, 2020, 586, 373-377.
- A. F. Goncharov, E. Bykova, M. Bykov, X. Zhang, Y. Wang, S. Chariton, V. B. Prakapenka and J. S. Smith, J. Appl. Phys., 2022, 131, 025902.
- 10 W. Cui, T. Bi, J. Shi, Y. Li, H. Liu, E. Zurek and R. J. Hemley, Phys. Rev. B, 2020, 101, 134504.
- 11 Y. Sun, Y. Tian, B. Jiang, X. Li, H. Li, T. Iitaka, X. Zhong and Y. Xie, Phys. Rev. B, 2020, 101, 174102.
- 12 T. Wang, M. Hirayama, T. Nomoto, T. Koretsune, R. Arita and J. A. Flores-Livas, Phys. Rev. B, 2021, 104, 064510.
- 13 D. R. Harshman and A. T. Fiory, J. Appl. Phys., 2022, 131, 015105.
- 14 L. Novakovic, D. Sayre, D. Schacher, R. P. Dias, A. Salamat and K. V. Lawler, Phys. Rev. B, 2022, 105, 024512.
- 15 I. A. Troyan, D. V. Semenok, A. G. Kvashnin, A. V. Sadakov, O. A. Sobolevskiy, V. M. Pudalov, A. G. Ivanova, V. B. Prakapenka, E. Greenberg, A. G. Gavriliuk, I. S. Lyubutin, V. V. Struzhkin, A. Bergara, I. Errea, R. Bianco, M. Calandra, F. Mauri, L. Monacelli, R. Akashi and A. R. Oganov, Adv. Mater., 2021, 33, 2006832.
- 16 X. Zhang, Y. Zhao, F. Li and G. Yang, Matter Radiat. Extremes, 2021, 6,068201.
- 17 M. Einaga, M. Sakata, T. Ishikawa, K. Shimizu, M. Eremets, A. Drozdov, I. Troyan, N. Hirao and Y. Ohishi, Nat. Phys., 2016, 12, 835–838.
- 18 E. Bykova, M. Bykov, S. Chariton, V. B. Prakapenka, K. Glazyrin, A. Aslandukov, A. Aslandukova, G. Criniti, A. Kurnosov and A. F. Goncharov, Phys. Rev. B, 2021, 103, L140105.
- 19 M. S. Somayazulu, L. W. Finger, R. J. Hemley and H. K. Mao, Science, 1996, 271, 1400-1402.
- 20 T. A. Strobel, P. Ganesh, M. Somayazulu, P. R. C. Kent and R. J. Hemley, Phys. Rev. Lett., 2011, 107, 255503.
- 21 J. D. Bernal and R. H. Fowler, J. Chem. Phys., 1933, 1, 515-548.
- 22 K. Lee, E. D. Murray, L. Kong, B. I. Lundqvist and D. C. Langreth, Phys. Rev. B: Condens. Matter Mater. Phys., 2010, 82, 081101.
- 23 A. Das, P. K. Mandal, F. J. Lovas, C. Medcraft, N. R. Walker and E. Arunan, Angew. Chem., Int. Ed., 2018, 57, 15199-15203.
- 24 E. J. Pace, X.-D. Liu, P. Dalladay-Simpson, J. Binns, M. Peña Alvarez, J. P. Attfield, R. T. Howie and E. Gregoryanz, Phys. Rev. B, 2020, 101, 174511.