

Combined First-Principles and Experimental Investigation into the Reactivity of Codeposited Chromium–Carbon under Pressure

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principles methods, revealing a large stabilizing effect tied to substoichiometry at the carbon site. These results offer an important case study into the current limitations of crystal structure prediction methods with regard to phase complexity and bolster the growing need for advanced theoretical approaches that can more completely survey experimentally unexplored phase space. **KEYWORDS:** *high pressure, diamond anvil cell, solid-state synthesis, density functional theory, transition metal carbides*

INTRODUCTION

Transition metal carbides (TMCs) find widespread use throughout industry due to their impressive high-resilience properties, which often include a combination of high melting points, microhardness, high thermal and electrical conductivity, and corrosion resistance.¹⁻⁵ These properties are especially pronounced in chromium carbides, which are used in a broad range of demanding applications. For example, Cr_3C_2 (*Pnma*) embedded within a NiCr binder is used for high-temperature and oxidation-resistant coatings in steam and gas turbines;^{6,7} Cr_7C_3 (Pnma) excels as a wear- and heat-resistant coating in bearing shafts, seals, and furnaces;⁸⁻¹¹ self-lubricating nanocomposites of amorphous CrC_y and the nanocrystalline NaCltype $\operatorname{CrC}_{(1-x)}(Fm\overline{3}m)$ thin films are promising materials for replacing noble metals in sliding electrical contacts;¹² and Cr_3C_2 (Pnma) embedded within carbon nanofibers shows improved cycling stability compared with conventional lithium/sulfur batteries.¹³

The chromium–carbon binary system contains three thermodynamically stable compounds (Figure 1): Cr_3C_2 (*Pnma*),^{14,15} $Cr_{23}C_6$ (*Fm* $\overline{3}m$),^{16,17} and Cr_7C_3 (*Pnma*).^{18,19} It also contains a number of metastable phases that cannot be accessed with traditional solid-state methods, including cementite Cr_3C (*Pnma*), which has been observed in small quantities following splat quenching;^{20,21} Cr_2C (*P6₃/mmc*),

which has been detected within iron alloys enriched with chromium;^{22,2,23} the Hägg carbide isomorph, Cr_5C_2 (C2/c), which has been prepared by reactive sputtering;^{24,25} the "filled Re₃B-type" Cr_3C_2 (*Cmcm*), which has also been observed with reactive sputtering;^{26,27} and finally the NaCl-type $CrC_{(1-x)}$ (*Fm*3*m*), which has been detected in reactive sputtering, ion-implantation experiments, and salt flux reactions.^{28–30} The large number of phases within the binary Cr–C system is remarkable and provides a solid foundation for chemists to study how the impressive high-resilience bulk properties are tied to crystal structures within this class of materials. Although many of the metastable chromium carbide phases have been structurally characterized, the recovery of samples suitable for bulk property characterization has remained a significant challenge.

To address this challenge, we turned to a high-pressure synthesis. This approach has proven to be one of the most productive methods for the bulk synthesis of metastable

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Figure 1. Crystal structures of selected chromium carbides. Polyhedra highlight the local coordination geometry around the carbon atoms (dark gray) bonded to chromium atoms (dark green). Cr_3C_2 (*Pnma*), Cr_7C_3 (*Pnma*), and $Cr_{23}C_6$ (*Fm* $\overline{3}m$) are thermodynamically stable, while Cr_3C (*Pnma*) and CrC (*Fm* $\overline{3}m$) are both metastable phases.

compounds that cannot be accessed with traditional heating methods, and in fact, it has already had demonstrable success in the discovery of bulk transition metal carbide phases.³¹⁻⁴⁰

We report here a study into the high-pressure phase space of the Cr–C binary system between pressures of 0 and 30 GPa. We begin with a survey of stable Cr–C phases using crystal structure prediction methods, which are commonly used to guide experiments by identifying potential synthesis targets. We then introduce a novel precursor sample preparation method that uses magnetron cosputtering of the elements to precisely control composition and mixing within diamond anvil cell experiments. We then return to computational approaches to gain deeper insight and detail on the stability of chromium carbides that we observe in our high-pressure synthesis experiments.

METHODS

Computational Details

In order to gauge the stability of a number of potential phases in the chromium carbide system, we performed first-principles density functional theory calculations within CASTEP v22.11.⁴¹ The calculations for the convex hull were performed with CASTEP's default ultrasoft pseudopotential library (C19), a maximum Monkhorst-Pack grid spacing of 0.02 Å^{-1,42} a plane-wave cutoff energy of 680 eV, and finite basis set corrections. Structures were individually relaxed at each pressure.

Calculations of the phonon modes of Cr_3C (*Pnma*) and CrC (*Fm* $\overline{3}m$) were performed under the harmonic approximation using the finite displacement method and a supercell, also known as the "direct method."^{43–45} These calculations used CASTEP's norm-conserving pseudopotential library (NCP), a maximum Monkhorst-Pack grid spacing of 0.03 Å⁻¹, a cutoff energy of 1600 eV, an acoustic sum rule correction applied in reciprocal space, ^{46,47} and spin-polarization. Prior to the phonon calculations, the Cr₃C (*Pnma*) structure was relaxed to a force tolerance of 0.0001 eV Å⁻¹. Forces on CrC (*Fm* $\overline{3}m$) are zero by symmetry.

Additional calculations on the stability of the substoichiometric $\operatorname{CrC}_{(1-x)}(Fm\overline{3}m)$ system were performed with CASTEP v19.11. These calculations used CASTEP's high-throughput pseudopotential library (QCS), a cutoff energy of 326.5 eV, a maximum Monkhorst-Pack grid spacing of 0.05 Å⁻¹, and finite basis set corrections. We sampled and geometry optimized around 400 vacancy configurations of a 32-Cr atom $2 \times 2 \times 2$ conventional supercell at each pressure of 0 and 20 GPa.

The Perdew–Burke–Ernzerhof (PBE) approach to the generalized gradient approximation of the exchange-correlation functional was used for all density functional theory calculations.⁴⁸ The c2x program,⁴⁹ atomic simulation environment (ASE) python package,⁵⁰ and pymatgen python package⁵¹ were used to convert, manage, and process the inputs and results of calculations. Most initial structures were acquired from the AFLOW Prototype Encyclopedia or web database.^{52–55} See the Supporting Information for more in-depth details of the calculations.

Preparation of Cosputtered Precursor Films

Chromium–carbon amorphous films were codeposited by magnetron sputtering (AJA International Orion 8 Sputtering System) onto Si (100) substrates using metallic chromium (99.95%) and carbon graphite (99.999%) targets, both of which were 5.08 cm in diameter and 0.635 cm thick. The chromium and carbon targets were powered by direct current (dc) and radio frequency (RF) magnetron sputtering, respectively. The power of the dc and RF supplies were set at 69 and 240 W, respectively, for the duration of the 10 h deposition. The base pressure of the sputtering chamber was below 3.0×10^{-6} Torr.

Throughout the deposition, the substrate was rotated at 100 rpm and was held 30 cm above the targets in an argon atmosphere with a sputtering pressure of 0.52 Pa. A glovebag filled with N_2 was used to prevent the oxidation of the samples while transferring from the load lock to our argon-filled glovebox. Flakes of these samples were characterized via scanning electron microscopy and energy dispersive spectroscopy using an FEI Magellan 400 XHR-SEM instrument with an Oxford X-MAX 80 mm² energy dispersive X-ray spectrometer attachment (Figure S13).

Laser-Heated Diamond Anvil Cell Synthesis

Flakes isolated from the cosputtered samples measuring approximately 90 μ m × 90 μ m were sandwiched between MgO disks (125 μ m in diameter).³¹ The MgO disks were loaded into the sample space of rhenium gaskets (originally 250 μ m thick) that were preindented to a thickness of 35–45 μ m and EDM-drilled to host a 127 μ m sample space in the center of the indent. DACs were taken to pressure in our argon-filled glovebox to ensure that the sample space was sealed with an inert environment.

In situ laser heating ($\lambda = 1064$ nm, fwhm ~ 30 μ m) and X-ray diffraction were performed at beamline 16-ID-B at the Advanced Photon Source, Argonne National Laboratory.⁵⁶ Samples were heated to 1700–2100 K. The incident X-rays ($\lambda = 0.48593$ Å) were aligned with the heating spot for in situ characterization during heating. MgO acted both as a thermal insulator and pressure calibrant.⁵⁷ Diffraction patterns were collected every 5 s with a 4.9 s exposure time on a Pilatus 1M-F detector. Temperature was determined by fitting the Planck radiation function to the blackbody radiation (600-800 nm) of the heated sample.⁵⁶ The temperature was measured through both sides of the cell, and the average was used for reporting to the nearest 100 K. Dioptas was used for real-time integration of the diffraction patterns.⁵⁸ TOPAS-Academic v6 was used to fit the patterns using Rietveld methods (see the Supporting Information for further details). Strongly textured MgO reflections and diamond Bragg reflections were masked before the integrations. Pressures were observed to differ slightly before and after the heating runs, but in all cases, the differences were below ~1 GPa.

RESULTS AND DISCUSSION

Enthalpy Calculations on the Cr-C System

We began our investigation of the Cr-C system by surveying stable phases between 0 and 30 GPa, using an approach that combined (1) a list of known stable phases in the Cr-Csystem along with simple and common phases in other binary



Figure 2. Calculated formation enthalpy of selected chromium carbide phases between 0 and 30 GPa, plotted in steps of 1 GPa. The allotropes of Cr and C with the lowest energy at each pressure were used as reference states. Cr was bcc for all pressures, while C was graphite for 0-6 GPa and diamond for 7-30 GPa. Phases that lie on the convex hull are plotted as circles, while metastable phases that lie above the convex hull are plotted as squares. The convex hull is plotted as a dashed line for 0, 10, 20, and 30 GPa. Data are colored according to pressure, as indicated by the color scale on the right of the figure. See the Supporting Information for details on all structures considered. The green points at x = 1 are the different carbon allotropes as they leave/approach the convex hull.



Figure 3. Plot of the calculated distance from the convex hull for selected Cr-C phases between 0 and 30 GPa. The data points for each phase are plotted in unique colors, with lines joining the points as a guide. Data points are not plotted for Cr_7C_3 (*Pnma*) and Cr_3C_2 (*Pnma*), which both remained on the convex hull over the entire pressure range plotted.

transition metal boride, carbide, and nitride systems (see the Supporting Information) and (2) a pool of randomly generated structures found using the AIRSS software package.^{59,60} The random structure searches were carried out at pressures of 0, 10, 20, and 30 GPa, and although they found a large number of Cr-C phases—consistent with the known stability of chromium carbides in general—they did not find any low-energy structures that were not already included in the seeded list of phases. The low-energy structures were then recalculated

over 1 GPa steps between 0 and 30 GPa using higher precision calculations (see the Methods section). Figure 2 plots the formation enthalpies of these selected phases between 0 and 30 GPa, with the convex hull being constructed at pressures of 0, 10, 20, and 30 GPa.

At 0 GPa, the calculations are consistent with the known phases reported in the literature, with $Cr_{23}C_6$ (*Fm* $\overline{3}m$), Cr_7C_3 (*Pnma*), and Cr_3C_2 (*Pnma*) all appearing on the convex hull. Notably, the η -Fe₂C-type Cr_2C (*Pnnm*) also appears on the

hull at 0 GPa, indicating that it should be synthetically accessible. The cementite analog, Cr_3C (*Pnma*), and the chromium Hägg carbide, Cr_5C_2 (*C*2/*c*), both lie just above the hull, with energy differences of 12 meV atom⁻¹ and 10 meV atom⁻¹, respectively.

Figure 3 plots the distance from the hull as a function of pressure for selected compounds. The three experimentally known phases remain on the hull up to 17 GPa, at which point $Cr_{23}C_6$ ($Fm\overline{3}m$) lifts from the hull. Cr_7C_3 (Pnma) and Cr_3C_2 (Pnma) remain on the hull over the entire range studied here. The η -Fe₂C Cr_2C (Pnnm) phase, which is not seen experimentally at ambient pressure, begins on the convex hull but then lifts sharply from the hull above 0 GPa. The metastable Cr_3C (Pnma) begins at 12 meV atom⁻¹ above the hull at 0 GPa and then gradually falls closer to the hull as the pressure is increased. Cr_5C_2 (C2/c) starts at 10 meV atom⁻¹ above the hull and increases its distance from the hull as the pressure increases. See the Supporting Information for complete details of distances from the hull.

 $CrC (Fm\overline{3}m)$ has been observed as a metastable phase in experiments,^{28–30} but it is not a low-energy structure in our searches. It is at least 248 meV atom⁻¹ above the hull over the pressure range calculated. $Cr_2C (Fd\overline{3}m)$ is an ordered supercell of the NaCl-type structure that has been observed experimentally in the metal-rich side of the Zr–C and Ti–C systems.^{61–64} This phase begins 96 meV atom⁻¹ above the hull at 0 GPa, which is lower energy than stoichiometric CrC (*Fm* $\overline{3}m$), but it increases to 184 meV atom⁻¹ above the hull at 30 GPa. This behavior is consistent with the stability of the Zr₂C (*Fd* $\overline{3}m$) phase in the substoichiometric NaCl-type Zr–C system, where Zr₂C is initially stable relative to fully stoichiometric ZrC and decreases in stability with increasing pressure.⁶⁵ Thus, our calculations are seemingly at odds with the experimental observation of this phase.

Taken together, the results of our preliminary crystal structure prediction calculations indicate that the metastable cementite Cr_3C (*Pnma*) phase could become synthetically accessible at high pressures, especially since its proximity to the convex hull is small compared to the additional energy expected from a finite temperature (i.e., these calculations are performed at zero kelvin and therefore may not accurately reflect the true potential energy surface that is relevant at higher temperatures). Based upon these results, we set out to synthesize Cr_3C (*Pnma*) in bulk form under high pressure. We did not expect to see any other phases, except for the known stable phases that exist on the hull at ambient pressure.

Laser-Heated Synthesis in the Diamond Anvil Cell

The crowded nature of the convex hull in the Cr–C system presents a technical challenge to synthesis and enforces the need for a precise control of chemical composition to avoid the formation of competing phases. This challenge is especially pronounced for high-pressure synthesis in the diamond anvil cell, where the overall composition and degree of element mixing is usually poorly constrained compared to traditional solid-state synthesis methods carried out on much larger scales (e.g., mm³ versus μ m³).^{66,67} In the present case, the targeted synthesis of the Cr₃C (*Pnma*) phase in the diamond anvil cell required us to target a precise 3:1 Cr:C precursor composition to avoid synthesizing Cr₂₃C₆ (*Fm*3*m*) and Cr₇C₃ (*Pnma*), which have similar Cr:C ratios. This motivated us to develop a new approach to precursor preparation—one that we hope will also provide a platform for a more precise and repeatable exploration of high-pressure phase space in general.

Inspired by work carried out by geophysicists,⁶⁸ we used magnetron sputtering to co-deposit chromium and carbon onto silicon, selecting deposition rates that led to a 3:1 Cr:C composition in the deposited film (see the Methods section and the Supporting Information). We deposited films with a total thickness of ~2.5 μ m. Individual flakes were lifted from the substrate and loaded into diamond anvil cells, where they were sandwiched between two laser-cut disks of (100)oriented magnesium oxide, which acted as both a thermal insulator and a pressure calibrant. The reactivity of these sputtered flakes under high pressures was studied using the laser-heated diamond anvil cell method at HPCAT beamline 16-ID-B.⁵⁶

We report here four separate heating experiments carried out within a single diamond anvil cell across different regions of the same sputtered sample flake. We performed each experiment at different pressures (13.5, 15.5, 20.2, and 24.3 GPa), adjusting the pressure of the cell between heating runs. In all runs, maximum temperatures of 1700-2100 K were reached, with total heating times on the order of 10-20 min. Temperature was quenched by turning off the lasers at the end of each heating run. Changes in the diffraction patterns were generally not observed below ~1000 K, regardless of the starting pressure. At or above ~1000 K, bcc-Cr was observed crystallizing from the amorphous signal, as evidenced by the growth and sharpening of bcc-Cr reflections. This acted somewhat as a fiducial in all of our experiments and defined the starting point of the synthesis.

The first heating run was carried out at ~13.3 GPa. Within a few seconds of chromium annealing, we observed the steady growth of peaks that could be readily matched to the cementite Cr_3C (*Pnma*) phase (Figure S25). The temperature was measured as ~1200-1400 K at the time that this phase crystallized. After holding for ~ 2 min, we then continued to increase the laser power, and as we reached ~1600-1800 K, we observed the growth of a second set of peaks, with a simultaneous decrease in the intensity of the Cr₃C (Pnma) peaks. The new peaks were not a match to any of the experimentally known phases or to any of the low-energy phases examined in our calculations. However, we could readily index the new family of peaks as belonging to a monoclinic crystal system. The lasers were then turned off after a few minutes. After thermally quenching the reaction, both the Cr₃C (*Pnma*) and monoclinic phase peaks remained, and new peaks appeared that could be readily matched to Cr₃C₂ (Pnma)-one of the three phases calculated to be on the convex hull at this pressure.

Our second heating run was carried out on a different region of the same sputtered flake at a pressure of ~16.3 GPa. As with the first heating run, we initially observed the crystallization of bcc-Cr, followed by the growth of peaks from Cr₃C (*Pnma*) above ~1200 to 1400 K, and then by peaks consistent with the monoclinic phase observed in the first heating run at higher temperatures (~1600–1800 K). The reaction was thermally quenched after a few minutes had passed with no further changes to the diffraction patterns. After thermally quenching the reaction, we again observed the growth of the peaks from Cr₃C₂ (*Pnma*). The results of the first two heating runs were thus qualitatively the same, and although this is perhaps not surprising given how close the pressures were, the repeatability of the complex phase transformations is an encouraging indication that composition was well constrained in our experiments. It is also worth noting here that each new region was unaffected by previous heating runs, with all sites yielding identical preheating diffraction patterns. This provides further support to the homogeneity of the sputtered sample flakes.

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We next increased the pressure in the cell to \sim 20.0 GPa. In this run, we once again observed the annealing of bcc-Cr, followed by the appearance of Cr_3C (Pnma) above ~1200-1400 K and then by the monoclinic phase at \sim 1600–1800 K. However, we then observed the growth of peaks matching the cubic CrC $(Fm\overline{3}m)$ phase immediately following the growth of the monoclinic phase (i.e., without any further increase in the temperature). This phase was completely unexpected from our crystal structure predictions, which calculated CrC $(Fm\overline{3}m)$ to have a formation enthalpy of no less than 248 meV above the hull over the studied pressures. The peaks from all three of the phases that grew in were retained after thermally quenching the reaction, and Cr_3C_2 (Pnma) once again appeared in the postheating diffraction pattern.

Our final heating experiment was carried out at ~24.4 GPa and produced qualitatively similar results to the reaction carried out at 20.2 GPa, with Cr₃C (Pnma) appearing first above ~1200-1400 K (albeit with noticeably weaker intensities than at lower pressures), followed by the monoclinic phase above ~1600-1800 K, and then immediately by the CrC (Fm3m). All of the phases were retained after thermal quenching. In this run, we did not observe peaks from Cr₃C₂ (Pnma) after cooling.

Figure 4 plots the X-ray diffraction data collected at each heating spot after the reaction was thermally quenched, and the laser table was removed from the X-ray path. These patterns are generally higher quality than those measured in situ, since absorption and scattering from the laser optics are removed, and the cell is able to rotate freely to achieve better orientation statistics and signal-to-noise ratio. Each of the four patterns were modeled using Rietveld methods, which allowed us to extract fitted lattice parameters for all of the identified phases.

To summarize the experimental LH-DAC results, we observed a total of four chromium carbide phases over the pressure range of 13.5 to 24.3 GPa, three of which are metastable phases that cannot be accessed under ambient pressures. Of the three thermodynamically stable phases already known from experiment, which our calculations predicted would remain on the hull over at least some of this range, only one of these, $(Cr_3C_2 (Pnma))$, was observed, and only after thermally quenching the reactions. The Cr₃C (Pnma) phase that was predicted to be accessible at higher pressure was observed experimentally at temperatures above \sim 1200–1300 K, which is in good agreement with calculations. However, the monoclinic phase and CrC (*Fm3m*), both of which grew above ~1600-1700 K, could not have been expected from our calculations. To investigate this discrepancy, and to reconcile our experimental observations with our crystal structure predictions, we returned to theoretical methods.

Assessing the Stability of NaCl-Type CrC

While the CrC ($Fm\overline{3}m$) phase would not be expected from our initial convex hull calculations, the experimental results unambiguously show that it crystallizes under high temperatures at pressures above ~ 20 GPa. One explanation is that a large substoichiometry at the carbon site could have a significant influence on the enthalpy of the phase. Indeed,



Figure 4. X-ray diffraction measured after thermal quenching at each heating site. Background subtracted experimental data are plotted in black. Rietveld methods are used to fit the multiple phases present and to extract unit cell parameters, with the modeled intensities plotted in red. Residual intensity is plotted in gray. Ticks are plotted beneath each trace to indicate the peak positions of the refined phases and are color-coded to each phase as denoted in the inset legend. Asterisks mark reflections from MgO (Fm3m).

2θ (°)

10

15

25

30

35

this is a common observation in many other TMCs that have been studied experimentally and computationally, with vacancy concentrations of up to 50% being typical in ZrC and TiC.^{61–65,69,70}

We adopted a simple approach wherein $2 \times 2 \times 2$ supercell structures of $CrC_{(1-x)}$ (Fm $\overline{3}m$) were generated with random concentrations and configurations of vacancies at the carbon site, before being relaxed using high-throughput DFT at pressures of 0 and 20 GPa (see the Methods section for more



Figure 5. Formation enthalpy of random configurations of substoichiometric $CrC_{(1-x)}$ ($Fm\overline{3}m$) plotted against the composition at 0 and 20 GPa. The enthalpy reaches a minimum for both pressures at around x = 0.35, where the formation enthalpy of the substoichiometric carbide becomes negative. This corresponds to about x = 0.39 in the domain of $Cr_{1-x}C_x$ as plotted in Figure 2 and about x = 0.65 in the notation CrC_x .



Figure 6. Gibbs free energy of formation of the CrC ($Fm\overline{3}m$) and Cr₃C (Pnma) phases at 0, 10, and 20 GPa plotted against temperature. The energies are calculated from phonon density of states using Cr (bcc) and C (diamond) as reference phases. The 0 GPa data is adjusted for reference against C (graphite) using standard reference values of the enthalpy and entropy for diamond and graphite.⁷¹ The data for CrC ($Fm\overline{3}m$) includes a term for an ideal mixing entropy with an assumed substoichiometry of $x = \frac{1}{2}$ in CrC_(1-x) which accounts for ~ 40 meV at 1300 K.

details). The results of these calculations are plotted in Figure 5. The calculations reveal that the formation enthalpy has a significant dependence on the vacancy concentration, with a prominent dip at around x = 0.35 (i.e., $CrC_{0.65}$) which is a clear indication of the energetic preference for substoichiometry in this system. We note here that Rietveld refinements of the CrC phase in our XRD data using a free occupancy at the carbon site led to a value of ~1/2. However, follow-up experiments will be required before we can be confident in reporting this

parameter accurately. For both pressures, the most stable structures appear around the same vacancy concentration and, in both cases, achieve a negative formation enthalpy, with the structures at 20 GPa having a higher magnitude formation enthalpy compared to those calculated at 0 GPa. This still however does not explain why the phase is seen in experiment given that the formation enthalpy would still be around 180 meV atom⁻¹ above the hull at 20 GPa even when considering the enthalpy contribution of substoichiometry.



Figure 7. Plot of the experimentally determined volumes of $\operatorname{Cr}_{(1-x)}(Fm\overline{3}m)$ and $\operatorname{Cr}_3C(Pnma)$ alongside volumes calculated with DFT methods. For $\operatorname{Cr}_{(1-x)}$, we plot volumes calculated for x = 0 and for the ordered $\operatorname{Cr}_2C(Fd\overline{3}m)$ phase, which we use as a proxy for the disordered phase with x = 0.5. The mean absolute percentage error between the experimental and calculated volumes is printed, with the shifted curves plotted as dashed lines.

In order to assess the dynamical stability of the observed metastable cementite Cr_3C (*Pnma*) and NaCl-type CrC phases, we calculated the phonon dispersion relations and the phonon density of states (PDOS) at 0, 10, and 20 GPa. Both the cementite Cr_3C (*Pnma*) and NaCl-type CrC (*Fm3m*) phases exhibit no imaginary modes, indicating that both phases are dynamically stable. The phonon calculations were conducted for the nonmagnetic phase of Cr_3C (*Pnma*) and both the nonmagnetic and the ferromagnetic phases of CrC (*Fm3m*). The ferromagnetic phase of NaCl-type CrC (*Fm3m*) is calculated to have lower energy and shows a noticeably larger bandwidth. All of the phonon results described above are plotted in the Supporting Information.

The phonon density of states enables an investigation into the effect that entropy, zero-point energy, and temperature have on energy, which can offer a more realistic assessment of stability as compared to calculations that use only the internal enthalpy (i.e., the zero kelvin calculations presented above). We used our PDOS to calculate the Gibbs free energy of Cr₃C (Pnma) and CrC (Fm3m) as a function of temperature at pressures of 0, 10, and 20 GPa (see the Methods section for details), and we plot the free energy of formation for both in Figure 6. We see that the free energy of CrC decreases with increasing temperature, while the free energy of Cr₃C increases with increased temperature. Although a truly quantitative comparison would rely on a consideration of the other phases as well as a more complete treatment of substoichiometry, we can nevertheless infer from our calculations that substoichiometry both lowers the enthalpy and raises the entropy of the CrC phase, thus supporting its observation in our experiments.

Finally, Figure 7 plots the experimental lattice parameters of Cr_3C (*Pnma*) and $CrC_{(1-x)}$ (*Fm* $\overline{3}m$) determined from X-ray diffraction data alongside calculated pressure-volume curves from geometry optimization calculations (see the Methodssection). For Cr_3C (*Pnma*), we see an underestimation of the calculated volumes with a mean absolute percentage error (MAPE) of 6.4%. For $CrC_{(1-x)}$ (*Fm* $\overline{3}m$), we instead see an overestimation of the calculated volume for fully stoichiometric CrC (Fm3m) (i.e., x = 0), with a MAPE of 12%. For comparison, we also plot the calculated volumes of the ordered Cr_2C (*Fd3m*) phase. This phase has an MAPE of 4.0% compared with the experimental volumes. The experimental volumes also agree well with the volumes of the calculated substoichiometric $CrC_{(1-x)}$ structures for around x = 0.5 which are plotted in Figure S11 in the Supporting Information. The low volume determined experimentally for $CrC_{(1-x)}$ could thus further support the presence of significant vacancies at the carbon site.

CONCLUSIONS

We have reported a combined first-principles and experimental investigation into the reactivity of chromium and carbon under high pressures. Crystal structure prediction methods were first employed to assess whether any novel Cr-C phases could be expected to stabilize under pressures up to 30 GPa. These calculations suggested that Cr_3C (*Pnma*) approaches the zero-kelvin convex hull under high pressures and could therefore become synthetically accessible; no other metastable phases were deemed to be promising candidates for synthesis.

We then enlisted a novel precursor preparation method using magnetron cosputtering of chromium and carbon to prepare amorphous well-mixed films with a tightly controlled 3:1 composition so that we could target the synthesis of Cr_3C (*Pnma*) while avoiding the other competing phases on the convex hull. Double-sided laser heating experiments carried out in the diamond anvil cell between pressures of 13.5–25.2 GPa on a flake of the cosputtered precursor led to the synthesis of the predicted Cr_3C (*Pnma*) phase but also to two other metastable phases: a novel monoclininc Cr–C phase and a NaCl-type CrC (*Fm*3*m*).

To rationalize the unexpected appearance of the NaCl-type CrC ($Fm\overline{3}m$), which we had calculated to be very far from the convex hull, we carried out first-principles studies into the stabilizing effect of substoichiometry at the carbon site, considering both the enthalpic effects as well as the entropy effect at higher temperatures. These calculations clearly showed that the effect of vacancies is sufficient to bring this phase close to the convex hull under our synthesis conditions. The newly discovered monoclinic phase has yet to be conclusively identified, but is likely to be either a lowsymmetry phase with a large number of atoms in its primitive cell (i.e., more than we can include in our random structure searching approach); a phase that becomes competitive under high temperatures; or a highly substoichiometric form of a simple phase that had been generated by our searches but which was simulated without carbon vacancies. It may also exhibit some combination of all of these factors.

In summary, we have shown that even in complicated systems such as Cr-C, where a large number of competing thermodynamic phases are already known, high pressures still offer a broad space for discovery. Our novel codeposition method for the preparation of stoichiometrically precise precursors is a promising tool for such systems, and we hope that it will enable the selective and high-yield synthesis of other phases being targeted with diamond anvil cell methods. This new synthetic handle on composition could also open the door to studying complicated ternary and quaternary systems under pressure or the use of site doping as a method for tuning the intrinsic properties and stabilities of metastable high-pressure phases.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsmaterialsau.3c00086.

Calculation and experimental details; Rietveld refinement details; random structure searches; convex hull plots; formation enthalpy plots; volume plots with substoichiometry; phonon dispersion and density of states plots; sputtering deposition rates; experimental setup illustrations; X-ray diffraction waterfall plots with difference traces; structure details with structure prototypes; tables of formation enthalpy distances from hull; and tables of Rietveld refinements (PDF)

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