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#### Structural phase transitions in yttrium up to 183 GPa

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Angle-dispersive x-ray powder diffraction experiments have been performed on yttrium metal up to 183 GPa. We find that the recently discovered oF16 structure observed in the high-Z trivalent lanthanides is also adopted by yttrium above 106 GPa, pressures where it has a superconducting temperature of ~20 K. We have also refined both tetragonal and rhombohedral structures against the diffraction data from the preceding "distorted-fcc" phase and we are unable to state categorically which of these is the true structure of this phase. Finally, analysis of yttrium's equation of state reveals a marked change in the compressibility upon adoption of the oF16 structure, after which the compression is that of a 'regular' metal. Electronic structure calculations of oF16-Y confirm its stability over oF8 structure seen in Nd and Sm, and provide insight into the nature of the shift of orbital character from s to d under compression.

#### I. INTRODUCTION

The trivalent transition metals yttrium (Y) and scandium (Sc) are considered members of the rare-earth elements, along with the lanthanide series (La to Lu). The unique structural and material properties exhibited by the rare-earth elements at extreme pressures has incentivized a great number of experimental and theoretical studies<sup>1-7</sup>. The regular trivalent lanthanides (La to Lu, excluding Ce, Eu and Yb) each exhibit a common phase transition sequence on compression, the structures of which differ only in the stacking arrangement of their close-packed atomic layers: hcp (space group  $P6_3/mmc$ , hP2 in Pearson notation)  $\rightarrow \alpha$ -Sm type  $(R\bar{3}m, hR9)$  $\rightarrow$  double-hcp (P6<sub>3</sub>/mmc, hP4)  $\rightarrow$  face-centred cubic  $(Fm\bar{3}m, cF4) \rightarrow \text{distorted-}cF4 \ (R\bar{3}m, hR24)^{1,8,9}$ . As the reverse sequence of structures (hP4 - hR9 - hP2) is adopted by the same elements at ambient conditions as one traverses them with increasing atomic number (Z). it was initially thought to arise from changes in the occupancy of the f-band with Z or with pressure. However, Y, which has no f electrons, was later shown to exhibit the same transition sequence on  $compression^{10,11}$ , and the subsequent reassessment of the structural changes found that they arise from an increase in the *d*-band occupancy, as a result of  $s \to d$  electron transfer<sup>9</sup>. Y has thus historically been used as a benchmark for comparing the electronic character of the structural transitions in the rare-earth metals  $^{10-13}$ .

None of the phase transitions listed above results in a measurable volume change<sup>6</sup>. However, upon further compression, the distorted-cF4 phase transforms to a

"collapsed" phase accompanied by a discontinuity in the atomic volume<sup>3,5,6</sup>. This transition occurs at  $\sim$ 95 GPa in Y with a volume change of  $-2.6\%^{13}$ . For the last 20 vears, the structure of the collapsed phases was generally perceived to be either orthorhombic (Cmcm, oC4) in Ce, Pr, Nd and Sm, or monoclinic (C2/m, mC4) in Ce, Nd, Sm, Tb, Gd, Dy, Ho, Er, Tm and Y (see Reference<sup>6</sup>, and references cited therein). The fits of the mC4 structure to the reported diffraction data were typically poor, except in the case of  $Ce^{14,15}$ , and we have recently shown that the true structure of the collapsed phase in Tb, Gd, Dy, Ho, Er and Tm is orthorhombic, space group Fddd with 16 atoms per unit cell  $(oF16)^7$ , while in Nd and Sm it is also orthorhombic with space group Fddd, but with 8 atoms per unit cell  $(oF8)^{16}$ . The oF8 structure was previously found in the actinides Cf, Am and Cm at high pressure,  $^{17-19}$  and the *oF*16 and *oF*8 structures, along with the hP3 structure observed in Nd, Sm and Yb<sup>20,21</sup>. form a new family of structures which differ only in the stacking sequence of their quasi-close packed layers<sup>7</sup>.

Remarkably, while the oF16 structure had not previously been observed experimentally in any element, although it has recently been confirmed as the stable phase of Ho above 103 GPa<sup>22</sup>, both it and the hP3 structure were calculated to be the stable phase(s) of Y above 97 GPa<sup>12,23,24</sup>, pressures at which Y has a remarkably high superconducting transition temperature ( $T_c$ ) of 19.5 K at 115 GPa<sup>25</sup>. It is then a reasonable assumption that the superconducting phase adopts one of these two structures, as both oF16 and hP3 are estimated to have a  $T_c$ in the range of 16–19 K at 97 GPa<sup>23,26</sup>. Furthermore, electronic structure calculations have found that shifts of the *d* electron states to low-energy levels, and a significant decrease in the density of states (DOS) at the Fermi level, contribute to the stabilities of both the oF16 and hP3 structures<sup>23</sup>. In addition, the accumulation of electrons into interstitial sites in both structures was also reported to play an important role in their stability<sup>23</sup>.

The diffraction patterns from the mC4 and oF16 structures are somewhat similar, which led to the incorrect assignment of the former to the collapsed phases of the high-Z trivalent lanthanides<sup>7</sup>. The previous determination of the structure of Y above 100 GPa as being  $mC4^{13}$ then suggests that the structure is more likely to be oF16rather than hP3, which has a different and much simpler diffraction pattern<sup>20</sup>. In order to determine the true structure of Y above 100 GPa, we have made diffraction studies to 183 GPa using x-ray powder diffraction and synchrotron radiation. We find that Y does indeed adopt the oF16 structure above 95 GPa, in agreement with computational studies  $^{23,24}$ . Our result confirms that the high-pressure structural behaviour of Y and the trivalent lanthanides is indeed very similar, but different to that of the other rare-earth element  $Sc^{27}$ .

The high quality of our diffraction data also enabled us to address another long-standing issue with the rareearth series, that is, the crystal structure of the distortedcF4 phase (d-cF4). Although the most widely accepted structure, including in Y, is hR24 (spacegroup  $R\bar{3}m$ )<sup>28,29</sup>, other possibilities have been suggested in order to explain peak splittings which are forbidden by the rhombohedral symmetry of this structure, for example in Dy<sup>30</sup>. In Y, recent *ab initio* density functional theory (DFT) calculations have suggested tetragonal or triclinic distortions of hR24 as being the true structure of the d-cF4 phase<sup>31</sup>, and we have tested these new models against our diffraction data.

#### **II. EXPERIMENT**

We conducted synchrotron x-ray diffraction experiments on four separate Y samples, attaining a maximum pressure of 183 GPa at 300 K. Diamond anvil cells (DACs) equipped with diamond culets ranging in diameter from 80 to 300  $\mu$ m were prepared with tungsten (W) gaskets and the sample chamber was filled with Y powder (99.9% purity, Sigma Aldrich) under a dry Ar atmosphere  $(<1 \text{ ppm } O_2 \text{ and } <1 \text{ ppm } H_2O)$  to prevent oxidation. A Cu sphere  $\sim 5 \,\mu \text{m}$  in diameter was loaded into three of the DACs as a pressure calibrant, while a small ruby sphere was used as a calibrant in the fourth. Angle-dispersive diffraction data were collected at the high pressure beamlines P2.02 at PETRA-III in Hamburg, at beamline I15 at the Diamond Light Source (DLS), and at beamline 9.5HP at the now-closed Synchrotron Radiation Source (SRS) at Daresbury Laboratory in the UK. Monochromatic x-rays of wavelength 0.34439 Å (PETRA-III), 0.42454 Å (DLS), and 0.44379 (SRS) were focused down to a FWHM of 2  $\mu m$  (PETRA), 20  $\mu m$  (DLS) and 50  $\mu m$  (SRS). Diffrac-



FIG. 1. Background-subtracted angle-dispersive diffraction profiles from Y collected on compression showing the diffraction patterns obtained from each of the different phases. For the four profiles collected below 95 GPa, the reflection positions predicted by the different phases (hP2, hP2 + hR9,hP4 and hR24, respectively) are indicated by vertical bars below each profile. The profile collected at 138 GPa is from the "collapsed" phase. The  $\checkmark$  symbols identify peaks from the W gasket,  $\bigtriangledown$  denotes peaks from the Cu calibrant and asterisks identify weak peaks from a sample impurity. The  $(2\bar{2}4)$  peak in the hR24 profile at 82 GPa is enlarged in the inset to highlight the asymmetric shape that may indicate it is a closely-spaced doublet – something that is forbidden by the rhombohedral symmetry of the hR24 structure.

tion data were recorded on a Perkin-Elmer area detector (PETRA-III) and a MAR345 image-plate detector (DLS and SRS). In one run at PETRA-III the detector was initially placed  $\sim 400$  mm from the sample, and later increased to  $\sim 600$  mm to improve the low-angle peak resolution, while in other runs at PETRA-III the detector was maintained at  $\sim 400$  mm. At DLS and SRS the detectors were maintained at  $\sim 300$  mm from the sample. The exact sample-detector distance and the detector tilts were determined using diffraction standards ( $LaB_6$ ,  $CeO_2$  and Si). The sample pressures were derived from the published Cu equation of state  $(EoS)^{32}$  and standard ruby fluorescence calibration data<sup>33,34</sup>. The 2D diffraction images were integrated to 1D profiles using  $DIOPTAS^{35}$ , and these were analysed using Rietveld<sup>36,37</sup>, Le Bail profile fitting, and least-squares fitting<sup>38</sup> to individual peak positions.



FIG. 2. Rietveld refinements of the (a) hR24 and (b) tI16structures to a diffraction profile of Y obtained at 102 GPa, showing the observed (cyan crosses) and calculated (black line) diffraction patterns. The calculated reflection positions (vertical bars), principal Miller indices, and difference profiles (lower lines) are shown. The  $\checkmark$  symbols identify a weak peak from the W gasket, and the asterisks identify a weak peaks from a sample impurity. The same diffraction peak is highlighted in both profiles illustrating how it is fitted as a singlet in hR24 and a doublet in tI16. The  $(1\overline{12})$  peak, which is predicted by the hR24 structure at  $4.2^{\circ}$  but not by the tI16structure, is highlighted in (a) The refined structural parameters are (a) a = 5.554(1) Å, c = 13.604 Å, atoms at (0.521(1), -0.521(1), 0.243(1)) and (0, 0.269(1)) and (b) a = 5.524(1) Å, c = 7.926(2) Å, atoms at (0.528(1), 0.219(1), 0.890(1)).

#### III. RESULTS AND DISCUSSION

At ambient pressure, Y crystallises in the hP2 structure and this produced sharp, well-defined diffraction peaks (Figure 1) and atomic volumes in excellent agreement with previous compression studies<sup>10,11</sup>. The transition from hP2 to the hR9 structure has been described previously as sluggish, starting at 13 GPa and becoming complete at 16 GPa<sup>13</sup>. We observed the transition to start at 14 and 16 GPa in two different samples, and observed the hR9 phase only as a minority component in mixed-phase samples along with either hP2 (at lower pressures) or hP4 (at higher pressures). Indeed, the similarities of the hP2, hR9, hP4 and cF4 structures, which differ only in the stacking of their close-packed atomic layers – e.g. AB in hP2 and ABACACBCB in hR9 – re-



FIG. 3. (a) The tI16 structure of Y at 102 GPa with a = 5.524(1) Å and c = 7.926(2) Å,  $c/a \sim \sqrt{2.06}$ , and atoms located on the 16*f* site at (0.528(1), 0.219(1), 0.890(1)). (b) The tI16 structure with  $c = \sqrt{2}a = 2a_{\rm FCC}$ , which reproduces the cF4 structure when the atoms are at (0.5, 0.25, 0.875). The cubic FCC unit cell highlighted by dotted lines.

sulted in significant overlap of diffraction peaks from the different phases, and sluggish transitions that produced mostly mixed-phase diffraction patterns.

These effects were perhaps exacerbated by the absence of any pressure transmitting medium (PTM) in our samples, which typically increases non-hydrostaticity. However, we note that neither Vohra *et al.*<sup>10</sup> nor Samudrala et al.<sup>13</sup> reported the use of a PTM in their previous studies. Using the 2  $\mu$ m diameter x-ray beam at PETRA-III we used 'grid-scans' to map out the pressure gradients within the samples at a number of pressures. These scans revealed not only pressure variations of 4 GPa within the sample volume at 89 GPa, and 7 GPa at 125 GPa, but also revealed that these variations resulted in different phase mixtures being present at different locations within the samples. We obtained accurate compressibility data, and reduced the effects of pressure gradients and mixedphase samples, by ensuring that our diffraction patterns contained diffraction peaks from both the Y sample and Cu pressure marker, and utilising the small x-ray beam size to ensure that the pressure gradients within the sampled volume were minimal. Preliminary experiments using the larger (20  $\mu$ m) x-ray beam at the DLS resulted in both more complex mixed-phase diffraction patterns, and discrepancies in the compressibility data due to the



FIG. 4. Rietveld refinement of the oF16 structure to a diffraction profile from Y at 138 GPa, showing the observed (cyan crosses) and calculated (line) diffraction patterns, the calculated reflection positions, and the difference profile. The  $\checkmark$  symbols identify peaks from the W gasket, and the asterisk identifies a weak peak from a sample impurity. Spacegroup Fddd with Y atoms on the 16e (x,0,0) Wyckoff sites with x = 3/16 (fixed), a = 16.911(3) Å, b = 4.656(1) Å, and c = 2.728(1) Å.

pressure gradients within the much larger ( $\sim 100 \times$ ) sampled volume.

While complex mixed-phase samples were often observed below 52 GPa, only single-phase diffraction patterns from the d-cF4 phase were observed above that pressure. If the d-cF4 phase of Y has the hR24 structure, then the  $cF4 \rightarrow hR24$  transition produces two changes to the diffraction pattern. Firstly, the distortion from cubic to rhombohedral symmetry results in the peaks of the cubic cF4 structure splitting at the transition. Importantly, the (200) peak in the cF4 pattern remains a singlet in hR24 (where it indexes as the  $(20\overline{4})$  – see Figure 2(a), while the most intense (111) cF4 peak splits into a doublet (which index as the (202) and (006) peaks). Secondly, additional diffraction peaks appear at the transition as a result of the enlarged unit cell of the hR24structure and the movements of the atoms at the transition to lower-symmetry sites.

Studies of single-phase d-cF4 profiles from Y between 52 and 102 GPa using the hR24 structure revealed that the (202)/(006) doublet remains unresolved at all pressures, and the same is true for the higher-angle (404)/(00,12)) doublet, which would have a larger splitting than the (202)/(006). The overlap of the peaks in these doublets implies that the c/a ratio is indistinguishable from  $\sqrt{6}$  at all pressures. This is in marked contrast to the behaviour seen in the hR24 phases of  $Pr^{39}$ , and  $Sm^{40}$ , where the (202)/(006) and (404)/(00,12) peaks are clearly resolved into doublets, and the c/a ratio increases



FIG. 5. The compressibility of Y to 183 GPa, shown as the volume per atom versus pressure (GPa). Open symbols refer to experimental data while the filled diamonds are atomic volumes of the oF16 phase determined from our DFT calculations. The inset shows an enlarged view of the region near the  $hR24 \rightarrow oF16$  transition, highlighting the -1.8% volume change ( $\Delta V/V_0$ ). The dashed line through the data points below 100 GPa is the best-fitting AP3 equation of state (EoS) with V<sub>0</sub>=33.0177 Å<sup>3</sup> (fixed),  $K_0$ =47.3(6) GPa, K' = 1.90(13), and K'' = -0.017(6). The solid line through the data from the oF16 phase is a best-fitting AP1 EoS with  $K_0 = 11.56(4)$  GPa K' = 6.02(2) and  $V_0$  is fixed at 32.42 Å<sup>3</sup>/atom, the value for oF16-Y at ambient pressure and 0 K as determined from our DFT calculations.

rapidly with pressure. Importantly, the (200) peak in the cF4 phase (which indexes as the (20 $\overline{4}$ ) in the hR24structure) shows some evidence of being a closely-spaced doublet in the d-cF4 diffraction profiles in Y - as highlighted in the 82 GPa profile in Figure 1. A similar doublet, which is forbidden by the hR24 structure, was seen in diffraction profiles from the d-cF4 phase of Dy, and this was used by Shen *et al.* to argue that d-cF4phase of Dy has orthorhombic rather than rhombohedral symmetry<sup>30</sup>.

The computational study of Y by Chen *et al.* suggested that the d-cF4 phase of Y has the hR24 structure<sup>12</sup>, and this was confirmed experimentally soon afterwards by Samudrala *et al.*<sup>13</sup>. However, more recently, Ishikawa *et al.*<sup>31</sup> have reported that two distortions of the hR24structure, with tetragonal and triclinic symmetry, also have low enthalpies over the relevant pressure range, with the former having the slightly lower enthalpy between 40 and 80 GPa. Analysis of the atomic coordinates of the tetragonal structure listed by Ishikawa *et al.*<sup>31</sup> reveal it to be body-centred rather than primitive, and to have spacegroup  $I4_1/a$  (tI16) with atoms on the



FIG. 6. Linearisation of compressibility of Y in the form of an  $\eta_{APL} - \sigma$  plot, where  $\sigma = \sigma_0 x$ ,  $x = (V/V_0)^{1/3}$  and  $\sigma_0$  is the Thomas-Fermi radius  $(3ZV_0/4\pi)^{\frac{1}{3}}$ . The data from the different phases of Y are plotted using different symbols, alongside the same plot for Sm (irregular compressibility, black filled squares) and the "regular" compressibilities of Cu, Au and Pt (grey filled circles), as determined from the compression data of Dewaele *et al.*<sup>42</sup> for comparison. The red line labelled "ideal solids" has the gradient  $\bar{\eta} = -5.67$  nm<sup>-1</sup>, and is the average gradient expected for all elements with regular compression<sup>43,44</sup>.

16f sites at (0.5295, 0.2327, 0.8918). This structure becomes cF4 when  $c = \sqrt{2}a = 2a_{\rm FCC}$  and the atoms are at (0.5, 0.25, 0.875). A key feature of the tI16 structure is that the (111) reflection in cF4 remains a singlet in tI16, while the (200) reflection becomes a doublet. In contrast, if the d-cF4 phase of Y were to adopt the oS8structure, first proposed by Porsch and Holzapfel<sup>41</sup> and reported to be the structure of the d-cF4 phase of Dy<sup>30</sup>, then both the (111) and the (200) peaks would become doublets. In the triclinic structure suggested by Ishikawa  $et al.^{31}$ , the (111) and the (200) peaks would split into 4 and 3 peaks, respectively. The tI16 structure therefore provides a high-symmetry, few-parameter solution to the structure of the d-cF4 phase when the (111) peak of cF4remains a singlet while the (200) becomes a doublet.

As highlighted in Figure 1, there is some evidence of the (200) peak of the cF4 becoming split in the d-cF4phase of Y, although this is evident only as an asymmetry on the low-angle side of the peak. A Rietveld fit of the tI16 structure to a profile obtained at 102 GPa (see Figure 2(b)) is excellent, and the refined structure is shown in Figure 3(a). The relationship of the tI16 structure to that of cF4 is illustrated in Figure 3(b).

However, despite the excellent fit of the tI16 structure, there are no clear features (apart from the possible doublet at 8.5°) not also accounted for in a fit using the hR24 structure with  $c/a = \sqrt{6}$  (see Figure 2(a)). While higher resolution profiles would make distinguishing the two structures straightforward, since they have different symmetry and predict a multitude of different peak splittings, our current data are unable to decide definitively between the two structural models. The hR24 structure also predicts an additional low-angle peak at ~ 4.2° (the  $(1\bar{1}2)$ ) not accounted for by tI16, but it is predicted to be extremely weak, and was not observed previously in either Pr or Sm<sup>39,40</sup>. We are thus unable to determine the structure of the d-cF4 phase of Y definitively, despite the quality of our diffraction data, and to obtain its compressibility we have refined it as having the hR24structure with the constraint  $c/a = \sqrt{6}$  (see Figure 2(a)).

We observe the transition to the "collapsed" phase of Y at 106 GPa (Figure 5) - which is much closer to the most recently predicted transition pressure of  $109 \text{ GPa}^{24}$ , than the 95 GPa reported by Samudrala et al.<sup>13</sup>. The onset of the transition is clearly evident from the splitting of the (202)/(006) doublet (of hR24), see Figure 1, and is complete by 117 GPa. A Rietveld refinement of the Fddd(oF16) structure to the diffraction profile obtained at 138 GPa is shown in Figure 4, and the fit is excellent. The oF16 structure comprises eight quasi-close-packed layers with a b/c axial ratio of  $\sqrt{2.9}$  – ideal packing would give an axial ratio of  $\sqrt{3}$ . In oF16 the atoms of each layer are located above the saddle point between two atoms in the preceding layer, resulting in ten-fold coordination (6 +2+2) - whereas in close-packed hP2, hP4, hR9 and cF4, each atom is located at the central point between the three closest atoms in the previous layer, resulting in twelve-fold coordination (6 + 3 + 3). The oF16 structure has an eight-layer stacking sequence - ABCADCBD - twice that of the four-layer ABCD repeat sequence of the isosymmetric oF8 structure observed in Sm. Nd. Am. Cm and  $Cf^{7,16-19}$ .

The compressibility of Y up to 183 GPa is shown in Figure 5. A volume change  $(\Delta V/V_0)$  of -1.8% (see inset to Figure 5) is observed at the transition to the "collapsed" phase, much closer to typical volume changes observed in the equivalent transition amongst the other rare-earth elements than the discontinuity of -2.6% reported by Samudrala *et al.* when using the C2/m structure to analyse the collapsed  $phase^{13}$ . The theoretical work of Chen et al. reported that the stability of the oF16 structure in Y arises from the shift in the *d*-electron energy levels, which results in the transfer of s electrons to the d-band<sup>23</sup>. Such changes in the electronic structure can result in changes in compressibility, which are most easily visualised by linearising the compressibility curve, enabling one to distinguish anomalies arising from changes within the electronic structure from the standard compressive behaviour expected of a 'normal' metal<sup>44</sup>.

If one fits the compression data using the second order (AP2) form of the Adapted Polynomial of order L (APL)  $EoS^{43,44}$ :



FIG. 7. The enthalpy difference between oF16-Y and the oF8-Y as a function of pressure, as calculated within DFT. The difference is normalized per Y atom and exhibits a minimum around P = 120 GPa.

$$P = 3K_0 \frac{(1-x)}{x^5} \exp(c_0(1-x))(1+x\sum_{k=2}^{L} c_k(1-x)^{k-1})$$
(1)

where  $K_0$  is the zero pressure bulk modulus, K' is its pressure derivative,  $x = (V/V_0)^{1/3}$ ,  $c_0 = -\ln(3K_0/p_{\rm FG})$ ,  $c_2 = (3/2)(K'-3) - c_0$ ,  $p_{\rm FG} = a_{\rm FG}(Z/V_0)^{5/3}$  is the Fermi-gas pressure, Z is the atomic number, and  $a_{\rm FG} = [(3\pi^2)/5](\hbar^2/m_e) = 0.02337$  GPa nm<sup>5</sup> is a constant, then the compression data can be linearised in a so-called  $\eta_{\rm APL} - x$  plot:

$$\eta_{\rm APL}(x) = \ln(\frac{px^5}{p_{\rm FG}}) - \ln(1-x)$$
 (2)

where  $x = (V/V_0)^{1/3}$ , the Fermi-gas pressure  $p_{\rm FG} = a_{\rm FG} (Z/V_0)^{5/3}$ , and  $a_{\rm FG} = 0.02337$  GPa nm<sup>5</sup>.

To realise differences in compression data of different materials with respect to "ideal" behaviour, it is most convenient to use an APL linearisation not with respect to x but rather to  $\sigma = \sigma_0 x$ , with  $\sigma_0$  being the Thomas-Fermi radius  $(3ZV_0/4\pi)^{\frac{1}{3}}$ , because this scaling brings out common trends more clearly<sup>44</sup>.

The linearised  $\eta_{\rm APL} - \sigma$  data for Y is shown in Figure 6, along with the similarly-linearised data we have recently reported for Sm<sup>16</sup> and the data for the "regular" metals Cu, Au and Pt<sup>42</sup>. In such a plot, materials undergoing "normal" compressive behaviour will show linear or quasi-linear behavior with an average "ideal' gradient of  $-5.67 \text{ nm}^{-1}$  and with the correct theoretical limit of  $\eta_{\rm APL}(0) = 0$ . This behaviour is clearly demonstrated by Cu, Au and Pt in Figure 6, while Sm provides an excelent example of a material whose compressibility is initially 'irregular' as a result of s - d electron transfer, but



FIG. 8. Band structure (BS) and density of states (DOS) for oF16-Y at ambient pressure (top panel) and at 140 GPa (bottom panel). The color code of the BS plot reflects the predominant orbital character: either s (red) or d (green). The color code of the 'stacked' DOS plots corresponds to the orbital character: red for s-orbital, blue for p-orbital, green for d-orbital, and cyan for f-orbital. These plots were made by using the PyProcar package<sup>45</sup>.

which becomes 'regular' after it transforms to the oF8 structure. Although the data from the lower-pressure phases of Y exhibit some scatter amongst the linearised  $\eta_{APL}$  values, a general positive gradient can be identified, similar to that shown by Sm. However, on entering the oF16 phase, there is a clear shift to linear behaviour with a negative gradient very similar to that found in Sm, Cu, Au and Pt. Although the pressure range over which we have compression data for oF16-Y (80 GPa) is somewhat smaller than the 100+ GPa pressure range over which we have compression data for oF8-Sm, the linearized data in Figure 6 suggest that oF16-Y, like oF8-Sm, is a metal with 'regular' compressibility.

The positive gradient exhibited by the lower-pressure phases of Y in Figure 6 suggests that the compression curve shown in Figure 5 would not be well fitted by an AP2 EoS. This was indeed the case, and we found that a 3rd-order AP3 form was required, as shown in Figure 5. However, the regular compression of the oF16 phase means that it could be well fitted using a 1st-order AP1 EoS (Figure 5).



FIG. 9. Relative contribution to the total DOS of the s and d states at P = 0GPa (solid red line: s-states, dashed green line: d-states) and at P = 140GPa (dotted blue line: s-states, dashed-dotted magenta line: d-states).

#### IV. ELECTRONIC STRUCTURE CALCULATIONS

As said, the transition to regular compressive behaviour in the collapsed oF16 phase may arise from a discrete change in the electronic structure of Y. Indeed the recent electronic band structure calculations of Li et al. for oF16-Y exhibited metallic features, with the majority of the DOS at the Fermi level collected in the 4d-band, transferred from the 4s-band<sup>24</sup>. This supports prior models predicting high critical Coulomb pseudopotentials for oF16, and it would be interesting to obtain a linearized plot (similar to Figure 6) from compression data collected below the predicted superconducting transition pressure, to observe if such treatment of standard compressibility data could provide additional insight into other material behaviours affected by its electronic structure.

To gain further insight, we have performed extensive DFT calculations of the oF16 and oF8 phases of Y. Structural optimization of bulk Y in the oF8 and oF16 phases was accomplished by using the DFT calculations with the help of the VASP<sup>46</sup> package using the Perdew-Burke-Ernzerhof functional<sup>47</sup>. The k-point sampling was performed using a Monkhorst-Pack mesh of  $8 \times 16 \times 2$  for the oF8 and  $8 \times 12 \times 2$  for the oF16 structures in the tetragonal unit cell, respectively, and a Gaussian smearing of 0.1 eV. During the DFT structural optimization, the convergence on internal forces and stress tensor of 0.01 eV/Åwas reached, and the energy cutoff was set to 500 eV. Scalar relativistic spin-orbit coupling was taken into account within the Koelling-Harmon approximation<sup>48</sup>.

Our zero-temperature DFT calculations for Y demonstrate that the oF16 phase is lower in enthalpy than the oF8 phase at all pressures between 40 and 240 GPa, as shown in Figure 7, with the maximum enthalpy gain being reached near 120 GPa. The agreement of our DFT zero-temperature volume calculations for oF16-Y as a function of pressure with the room-temperature experimental data is also very good, as can be seen from Figure 5. The absence of the imaginary phonon modes in the oF16 phase was previously established in the work of Chen et al.<sup>23</sup>.

The qualitative change of the valence electrons' orbital character in the oF16-Fddd can be best seen from the combined band structure and density-of-states plots at ambient and at high pressures, as shown in Figure 8. It can be seen that the major contribution to the band structure in the energy range of [-5:10] eV comes from the s and d states, while the main effect of the pressure on Y is to increase dramatically the bandwidth of the d states. Indeed, by applying pressure from 0 to 140 GPa, the d state's bandwidth increases by a factor of two. At the same time, the s states, located around 4 eV below the Fermi energy, are slightly raised by around 1 eV and are strongly hybridised with the d states, as a result of applied pressure.

In order to underline the difference in contribution of the s and d states to the total DOS at high and low pressure, in Figure 9, we plot the quantities:

$$\rho_s(E) = \frac{D_s(E)}{D_{tot}(E)}, \quad \rho_d(E) = \frac{D_d(E)}{D_{tot}(E)},$$
(3)

where  $D_s(E)$ ,  $D_d(E)$ ,  $D_{tot}(E)$  are the partial s, partial d and the total DOS respectively. It can be seen that at P = 0GPa, the total DOS is dominated by the s states, reaching 100% at -5eV, and the s states contribution gradually goes to zero as one approaches Fermi energy. At the same time, the d-states contribution gradually grows until it reaches a value of the order of 80% at  $E_F$ . Conversely, at high pressure, the s-states contribution remains approximately constant and small (below 20%), while the d-states contribution, starting at 100% at -5 eV, decreases slightly, but remains higher than 60% until few eV above Fermi energy. Thus, in our calculations, the  $s \rightarrow d$  charge transfer appears more like a d bandwidth increase rather than a real transfer of the charge between the orbitals, which is ill-defined in a metallic system.

#### V. CONCLUSIONS

The structure of Y metal above 100 GPa, and thus the likely structure of the superconducting phase, is found to be face-centered orthorhombic (oF16), isostructural with that observed in the collapsed phases of the trivalent lanthanides Tb, Gd, Dy, Ho, Er, and (probably) Tm. This result confirms the close structural relationship between the lanthanides and yttrium, which has no f electrons. High precision measurements of the compressibility reveal anomalous values at lower pressures, very similar to those seen in Sm, but that the compressibility of the oF16 phase is that of a regular metal, such as Au, Pt or Cu. DFT calculations confirm the lower enthalpy of the oF16 structure relative to that of oF8 structure seen in Sm and Nd, and reveal that the  $s \rightarrow d$  charge transfer occurs via a *d*-bandwidth increase, which enhances the relative contribution of the *d*-states to the density of states near the Fermi level.

Added note: Buhot *et al*<sup>49</sup> have recently reanalyzed the x-ray diffraction data of Samudrala *et al*<sup>13</sup> at 123 GPa using Rietveld methods and shown that it can be fitted with the oF16 structure, while their superconductivity measurements show that the oF16 phase can be stabilised without partially-occupied f states.

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