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Suppression of the Phase Coexistence of the fcc-fct Transition in Hafnium-Hydride Thin Films

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perspective, the results highlight the profound influences of the nanostructuring and nanoconfinement of metal hydrides on their structural response to hydrogen with a significant impact on their applicability in future devices.

Metal hydrides are a class of materials that may play an important role in a hydrogen-powered economy. Traditionally considered to be hydrogen storage materials,¹⁻³ other applications such as in hydrogen-purifying membranes,^{4,5} switchable mirrors,^{6,7} fuel cells,⁸ and especially hydrogen sensors^{3,9-14} have attracted attention recently. In general, the *x* (hydrogen-to-metal ratio) – *T* (temperature) phase diagrams of metal hydrides are relatively complex, involving a combination of body- or face-centered cubic, tetragonal, orthorhombic, or hexagonal metal hydride phases that may or may not coexist.¹⁵ A detailed understanding of (the transitions between) these phases is of vital importance because it determines, together with the entropy and enthalpy of formation, the performance of metal hydrides in real-life applications.

Although many applications of metal hydrides rely on nanostructered materials such as thin films and nanoparticles, most studies determining the phase diagram and nature of phase transitions focused on the bulk. Nanostructuring may significantly alter the properties of metal hydrides, changing their stability and the occurrence of phases and phase transitions (see, e.g., refs 16–24). Such effects may arise from an increased surface-to-volume ratio or, especially for thin films, geometric constraints on the possibility to expand volumetrically and clamping to the supporting substrate. In particular, for hydrogen-sensing applications, metal hydrides are typically structured as thin films or nanoparticles, 3,11-14,25 and understanding the occurrence and nature of phase

transitions is of vital importance to designing hysteresis-free hydrogen sensing materials that are stable over time with repeated exposure to hydrogen.

Hafnium (Hf) is a promising optical hydrogen-sensing material that features an extraordinary hysteresis-free sensing range spanning over 6 orders of magnitude in hydrogen pressure.^{3,26,27} In the bulk, similar to other group IV elements titanium (Ti)^{28–30} and zirconium (Zr),^{31,32} a large two-phase region separates the hexagonal-close-packed (hcp) solid solution at low x in HfH_x from the face-centered-cubic (fcc) phase ($x \gtrsim 1.5$), where hydrogen occupies the interstitial tetrahedral sites (4f).^{28,33,34} Upon further hydrogenation, a transformation to the face-centered tetragonal (fct) phase is observed, in which the fcc lattice is compressed along the *c* axis. The attractiveness of hafnium as a hydrogen-sensing material stems from the fact that hafnium thin films show a highly reproducible change in optical transmission in response to a hydrogen pressure, which can be used to accurately determine the hydrogen pressure.^{3,13,26,27} The sensing range corresponds

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Figure 1. Diffraction and NMR results on powder HfD_x at three different deuteration states: HfD_{1.66}, HfD_{1.78}, and HfD_{1.95}. (a) X-ray diffraction patterns (Cu K α , λ = 0.1542 nm). The continuous lines show the fit to the data, while the additional lines underneath the diffraction patterns indicate the corresponding residuals of the fits. (b) Neutron diffraction patterns obtained at NOVA located at the Japan Proton Accelerator Research Complex (J-PARC). The insets shows the split of the {200} reflection at HfD_{1.66} to (002) and (110) at HfD_{1.95}, while the coexistence of these three reflections is observed at HfD_{1.78}. (c) ²H magic angle spinning nuclear magnetic resonance (MAS NMR) spectra.

to a hysteresis-free hydrogenation of Hf from about HfH₁₅ to HfH_{1.98}.^{26,27} The appearance of a hysteresis-free sensing range is most remarkable, as for bulk Hf a first-order fcc \leftrightarrow fct phase transition involving hysteresis is reported between HfH_{1.78} and $HfH_{1.86}$.^{28,34} As such, it is unclear whether the phase transition is suppressed in thin films or whether nanoconfinement pushes the transition toward second order, thereby eliminating the hysteresis. Studying this and other phase transitions in thin films is typically complicated by the strong epi-texturing of the films: while powder X-ray diffraction (XRD) provides information on multiple lattice reflections, typical Bragg-Brentano XRD measurements of thin films provide limited information on the out-of-plane direction only: in the case of hafnium-hydride only the (111) and (222) reflections can be observed, preventing the observation of the fcc \leftrightarrow fct phase transition.

Here, we combine *in situ* out-of-plane, in-plane XRD and Xray reflectometry (XRR) to elucidate the nature of the fcc \leftrightarrow fct phase transition in Hf thin films. We unambiguously identify the fcc \leftrightarrow fct phase transition in Hf thin films, but at a much larger value of x than in the bulk and with a smaller degree of tetragonality. Different from bulk HfH_x no sign of phase coexistence is observed, suggesting that the confinement of the film suppresses the phase coexistence. In a more general perspective, these results highlight the profound influence of nanoconfinement on the presence and nature of phase transitions in thin films and other nanostructured metal hydrides.

Before discussing the results on thin films, we first consider the nature of powder Hf to confirm the first-order nature of the fcc \leftrightarrow fct phase transition in the bulk. Figure 1 displays X-ray and neutron diffraction results as well as ²H magic angle spinning nuclear magnetic resonance (MAS NMR) spectroscopy results of HfD_x in three different deuteration states— HfD_{1.66}, HfD_{1.78}, and HfD_{1.95}—obtained after pressure– composition isotherm measurements at 300 °C (Figure S3). As previous results indicate that the hafnium–hydrogen and hafnium–deuterium phase diagrams are very similar,³⁴ we use deuterium instead of hydrogen to suppress the incoherent background in the neutron diffraction experiments and to enable a higher resolution in NMR.

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The X-ray and neutron diffraction patterns (Figure 1(a,b)) of HfD_{1.66} and HfD_{1.95} are well described by single-phase models of fcc ($Fm\overline{3}m$) and fct (I4/mmm), respectively, while the pattern of HfD_{1.78} can be fitted using only a two-phase model where the fcc and fct phases coexist. While for fitting the XRD pattern the contribution of deuterium is assumed to be negligible, neutron diffraction confirms that deuterium atoms occupy the tetrahedral sites. The lattice parameters that are refined by neutron diffraction are a = 0.487648(6) nm for HfD_{1.66} and a = 0.487805(6) nm and c = 0.434134(10) nm (c/a = 0.89) for HfD_{1.95}. The refinement of the pattern for HfD_{1.78} using the two-phase model provides a = 0.467416(33) nm for the fcc phase and a = 0.481491(12) nm and c = 0.443678(26) nm (c/a = 0.92) for the fct phase, respectively. The obtained weight fraction of these two phases is fcc/fct = 39:61.

The ²H MAS NMR spectra confirm the single-phase behavior for HfD_{1.66} and HfD_{1.95} and the phase coexistence for HfD_{1.78}. The spectra of HfD_{1.66} and HfD_{1.95} show a single component (Figure 1(c)), which have negative Knight shifts of -9.9 and -33.9 ppm, respectively. The Knight shift is attributed to the density of states at the Fermi energy: HfD₁₆₆ is relatively more insulating than HfD₁₉₅, consistent with the reduction of the optical transmission upon increased hydrogenation (Figure 4(d,e) and refs 26 and 27). The spectrum of HfD_{1.78} consists of two contributions due to the phase coexistence. As such, it was fitted using two Gaussian functions with an area ratio of fcc/fct = 4:6. Assuming that the deuterium contents of the fcc and fct phases are D/Hf = 1.75and 2, respectively, and using the fact that the intensity of NMR signals is proportional to the molar ratio of deuterium, the weight fraction is estimated to be fcc/fct = 43:57, which is close to the one obtained by neutron diffraction. Most importantly, the phase coexistence observed for HfD₁₇₈ by both X-ray and neutron diffraction as well as NMR unambiguously confirms the first-order nature of the fcc-fct phase transition.

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Figure 2. *In situ* out-of-plane and in-plane XRD measurements (Cu K α , $\lambda = 0.1542$ nm) on the 60 nm Hf films capped with a 10 nm Pd layer and with a 3 nm Ti adhesion layer at T = 120 °C and the partial hydrogen pressures P_{H_2} indicated. During the experiment, the hydrogen pressure was increased stepwise. (a) Out-of-plane diffraction and in-plane diffraction with the sample tilted in the direction perpendicular to the X-ray beam by (b) $\chi = 35^{\circ}$, (c) $\chi = 55^{\circ}$, and (d) $\chi = 70^{\circ}$. The continuous lines show fits of pseudo-Voigt function(s) to the data from which the *d* spacing (Figure 3) has been derived.



Figure 3. Partial hydrogen pressure dependence of the *in situ* out-of-plane and in-plane *d* spacing of 60 nm Hf films capped with a 10 nm Pd layer and with a 3 nm Ti adhesion layer at T = 120 °C. During the experiment, the hydrogen pressure was stepwise increased and decreased. The *d* spacing was obtained by fitting pseudo-Voigt functions to the data of Figure 2 and normalized to the *d* spacing of the unloaded state as measured in air. *d* spacing was obtained by (a) out-of-plane diffraction and in-plane diffraction with the sample tilted in the direction perpendicular to the X-ray beam by (b) $\chi = 35^\circ$, (c) $\chi = 55^\circ$, and (d) $\chi = 70^\circ$.

Turning to the results of the thin films, Figure 2 displays the out-of-plane and in-plane diffraction measurements obtained for different partial hydrogen pressures at 120 °C. The thin films are produced by magnetron sputtering on fused quartz substrates and are composed of a 4 nm Ti adhesion, a 60 nm Hf, and a 10 nm Pd capping layer to prevent oxidation and promote hydrogen dissociation. As the films are highly textured, the out-of-plane results reveal only the (111) peak (Figures S4 Figure S5). As such, we performed in-plane diffraction by tilting the sample in the direction perpendicular to the beam by $\phi = 35$, 55, and 70° in order to monitor the hydrogen pressure dependence of the (200), (220), and in-plane (111) reflections.

Most importantly, the diffraction patterns show the hallmarks of the fcc \leftrightarrow fct transformation: comparing the

diffraction patterns in the unloaded state and the highest hydrogen pressure measured, $P_{\rm H_2} = 613$ Pa, indicates that the (111) reflection (Figure 2(a,d)) remains unsplit, both in the in- and out-of-plane directions, while the (200) (Figure 2(b)) and (220) (Figure 2(c)) reflections, both oriented in-plane, each split into two peaks. On the basis of the peak positions, we observe a substantial expansion of the *a* axes from 0.472 nm in the unloaded state to 0.479 nm in the fully hydrated state. At the same time, the *c* axis contracts to 0.457 nm, resulting in $c/a \approx 0.95$. As such, the volume of the unit cell in the fully hydrogenated state of V = 0.0524 nm³ is approximately the same as for the bulk (V = 0.0516 nm³), while the compression of the *c* axis and expansion of the *a* axes are much less pronounced in the thin films than in the bulk ($c/a \approx 0.90$).



Figure 4. Reflectometry and optical transmission measurements on the 60 nm Hf films capped with a 10 nm Pd layer and with a 3 nm Ti adhesion layer. (a) *In situ* X-ray reflectometry (XRR) measurements at T = 120 °C and the partial hydrogen pressures P_{H_2} indicated. During the experiment, the hydrogen pressure was stepwise increased. The dots indicate the measurement points, while the continuous lines indicate the fits to the data. (b) Partial hydrogen pressure dependence of the Hf layer thickness at T = 120 °C derived from fitting the XRR data that was collected by stepwise increasing and decreasing the partial hydrogen pressure. (c) Hydrogen content at T = 120 °C of 40 nm Hf films capped with a 10 nm Pd layer as deduced from neutron reflectometry measurements. Data were obtained from ref 27. (d) Changes in the white light optical transmission where the film was exposed to various increasing and decreasing pressure steps of $P_{H_2} = 0.22-9.5$ Pa at T = 120 °C. The optical transmission \mathcal{T} is measured relative to the transmission of the film after unloading in air (\mathcal{T}_{unl}). The dashed lines indicate levels of the same transmission and pressure. (e) Partial hydrogen pressure dependence of changes in the white light optical transmission at T = 90, 120, and 150 °C.

The fcc \leftrightarrow fct transition at $P_{\text{H}_2} = 10^{+1}$ Pa is accompanied by a considerable reduction in volume. The out-of-plane diffraction results indicate a reduction of the out-of-plane d_{111} spacing by about 1% (Figure 3(a)) when fcc \leftrightarrow fct occurs. Differently, the in-plane d_{111} spacing remains unaltered and thus does not change upon increased hydrogenation. Consistent with this, the XRR measurements of Figure 4(a,b) indicate that the Hf layer thickness decreases by about 1%, thus indicating that the reduction of the d_{111} spacing is translated completely in the out-of-plane direction.

Different from bulk measurements, where phase coexistence of fcc- and fct-HfH_x is observed in the region between $x \approx 1.78$ and $x \approx 1.86$,³⁴ the XRD patterns of thin film Hf do not show any indication of the simultaneous occurrence of these two phases. The absence of phase coexistence in thin films could imply that the confinement of the film suppresses the phase coexistence associated with a first order phase transition. It has the important implication that the hysteresis associated with first-order phase transition is (practically) eliminated. Indeed, both earlier^{26,27} and present (Figure 4(d,e)) optical transmission measurements of Hf indicate a completely hysteresisfree response to changing hydrogen pressures: The optical transmission at a certain $P_{\rm H_2}$ is exactly the same after increasing and decreasing the pressure, making the response of hafnium as an optical sensing material completely free of hysteresis.

In addition, we observe that the phase transition in thin films occurs at a much larger x than in the bulk. Both our data (Figure 1) and data in the literature (see, e.g., refs 28 and 34) indicate that the onset of the transition occurs at $x \approx 1.78$ in the bulk. For thin films, neutron reflectometry measurements (Figure 4(c)) indicate that the partial pressure of the transition, $P_{\rm H_2} \approx 10$ Pa, corresponds to $x \approx 1.92$. Different from bulk materials, two-dimensional films that are clamped to the substrate have to obey constraints on lateral expansion

because expansion can be realized only in the out-of-plane direction, which may result in a very high in-plane stress and affect the stability of phases.^{17,22,23,35,36} Indeed, the diffraction results indicate that the unit cell expands only in the out-of-plane direction and continuously deforms from its unloaded state: at $P_{\rm H_2} = 9.5$ Pa (HfH_{1.90}), the out-of-plane d_{111} spacing is approximately 2% longer than the in-plane spacing (Figure 2(a,d)). Moreover, the nucleation of domains, inducing locally large stresses, may also be hindered considerably by the clamping of the film to the substrate.^{22,24,35,37-40} The collective result of the clamping is that the onset of the transition is pushed to higher values of *x*, the coexistence of phases is suppressed, and the tetragonality is much smaller than for the bulk.

Most remarkable is that an abrupt increase in both the outof-plane d_{111} spacing (Figure 3(a)) and layer thickness (Figure 4(b)) of about 1% is observed at $P_{\rm H_2} = 10^{+2}$ Pa, while no changes are seen in the in-plane XRD results. Neutron reflectometry measurements (Figure 4(c)) reproduced from ref 27 indicate that this abrupt change in layer thickness and d_{111} spacing occurs when the layer is nearly fully hydrogenated, i.e., at $x \approx 1.98$. While we have no direct explanation for this remarkable observation, a direction for future research would be to determine if such a volume reduction also occurs in other group IV elements Zr and Ti.

In the literature, the root cause of changes in the optical transmission of metal hydride thin films upon exposure to hydrogen has been debated. Often, Beer-Lambert's law is taken as a staring point: $T = I/I(d) = e^{-\alpha d}$, with T the optical transmission of the film, d the thickness and α the linear attanuation coefficient. In this framework, changes in the optical transmission upon hydrogenation can be due to the volumetric expansion of the film (that needs to be completely translated in an expansion of the film thickness d) or to

differences in hydrogen concentration in the metallic thin film causing changes to the dielectric constants and thus the adsorption coeficient α . Different from a previous study of vanadium films for which it was reported that the optical transmission changes are driven by the volumetric expansion of the thin film,⁴¹ our results are consistent with optical transmission changes dictated by changes in the hydrogen concentration affecting the dielectric properties. Indeed, the fcc \leftrightarrow fct transition including the pronounced reduction in volume is not reflected in the partial pressure dependence of the optical transmission that changes monotonically with increasing partial hydrogen pressure (Figure 4(d,e)). Thus, our results suggests that the changes in optical transmission are fully dictated by the change in hydrogen concentration, causing changes to the dielectric properties, and not by the variation of the thickness of the thin film. This is consistent with neutron reflectometry and optical transmission measurements of Hf²⁶ (and Ta, 27 Pd_{1-y}Au_y²⁴) that indicate a linear relationship between the optical transmission and the hydrogen concentration of the metallic layer. The fact that the optical transmission is unaffected by the fcc \leftrightarrow fct transition has the convenient implication that Hf can be used as a hydrogen sensor in which the optical response monotonically changes with the hydrogen pressure/concentration in the environment over a large range.

In conclusion, using *in situ* in- and out-of-plane X-ray diffraction and reflectometry, we unambiguously identify that the fcc \leftrightarrow fct transition also exists in Hf thin films. The confinement of Hf and the clamping to the substrate significantly affect the transition: compared to bulk Hf, the transition is pushed to higher values of x, the tetragonality of the fct phase is reduced, and the coexistence of phases is suppressed. These results highlight the profound influences of the nanoconfinement of metal hydrides on their structural response to hydrogen. In the case of Hf, the nanoconfinement effects facilitate the extraordinary performance as an optical hydrogen-sensing material including a hysteresis-free sensing range over 6 orders of magnitude in partial hydrogen pressure.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpclett.1c03411.

Experimental details; spectrum of the LED lights used for the optical transmission measurements; photographs of the *in situ* XRD/XRR setup. Pressure–composition isotherm of bulk powder HfD_x; out-of-plane and inplane XRD measurements of the Hf thin films; and rocking curves of the Hf thin films (PDF)

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Notes

The authors declare no competing financial interest.

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