Anomalous lattice dynamics and thermal properties of supported size- and shape-selected Pt nanoparticles

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Anomalous lattice dynamics and thermal behavior have been observed for ligand-free, size-, and shape-selected Pt nanoparticles (NPs) supported on nanocrystalline γ-Al2O3 via extended x-ray absorption fine-structure spectroscopy. Several major differences were observed for the NPs with respect to bulk Pt: (i) a contraction in the interatomic distances, (ii) a reduction in the dynamic (temperature-dependent) bond-length disorder and associated increase in the Debye temperature (θD), and (iii) an overall decrease in the bond-length expansion coefficient coupled with NP stiffening. The increase in the Debye temperature is explained in terms of the NP size, shape, support interactions, and adsorbate effects. For a similar average size, we observe a striking correlation between the shapes of the NPs and their θD values.

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I. INTRODUCTION

Metal nanoparticles (NPs) display anomalous vibrational properties such as phonon confinement effects and strong modifications of the phonon density of states as compared to bulk materials.1–6 Although some of the factors contributing to this unusual behavior are understood (e.g., NP interaction with the support, encapsulating ligands, and/or adsorbates), they are normally discussed in isolation. Furthermore, while it is recognized that environmental effects do contribute to the thermodynamics of small systems,7 very little is known on how to sort out and control such complex interactions at the nanoscale. The resolution of these challenges requires the combination of highly reproducible NP synthesis and sensitive methods for determining thermodynamic properties in different environments.

Recent extended x-ray absorption fine-structure (EXAFS) experiments revealed two peculiarities in the thermal properties of certain supported metal NPs: (i) a contraction in their nearest-neighbor (NN) bond lengths at elevated temperatures and (ii) an increase in their Einstein (and, hence, Debye, θD) temperature with respect to bulk.8,9 The former effects were tentatively attributed to thermally induced dynamic changes in the structure of the NPs and to the strong interaction of the NPs with defects (vacancies) in the support.10 An increased θD relative to the bulk has been reported for a variety of supported NP systems9,11–17 and unsupported nanocrystalline agglomerates.14–16 By contrast, reduced values of θD have also been observed for seemingly analogous systems.17,18 The wide discrepancy observed for θD suggests that many factors are at play. In free clusters, enhanced surface energy and the concomitant increase in the surface stress causes a lattice contraction at the NP surface, softening of interatomic force constants, and a suppression of θD and melting temperatures.1 On the other hand, the increased θD observed for certain systems has been attributed to structural stiffening and inhomogeneous internal stress correlated with the NP size, surroundings (e.g., adsorbates/surface ligands,15,16,19 a matrix encapsulating the NPs or a support12,13), and to the presence of structural defects and multiple grains in large NPs.14 It is evident that the understanding of thermal, structural, and electronic properties of these systems is hindered by their complexity, due to the multiple competing factors that define their behavior, and any effort in sorting out these influences will be helpful for developing first-principles theories. In the present study we investigate the relationship between thermodynamic quantities (θD and thermal-expansion coefficient) and geometrical properties of ligand-free, homogeneous, size-, and shape-selected Pt NPs supported on γ-Al2O3 using EXAFS.

II. EXPERIMENTAL AND THEORETICAL METHODS

A. Sample preparation

Size- and shape-selected Pt NPs were prepared by inverse micelle encapsulation methods using poly(styrene)-block-poly(2vinylpyridine) (PS-P2VP) diblock copolymers. Briefly, commercially available PS-P2VP diblock copolymers were dissolved in toluene to form inverse micelles with the P2VP head groups constituting the micelle core. Shape-selected Pt NPs are created by dissolving H2PtCl6 into the P2VP/Pt block to form a polymeric solution. Subsequently, the nanocrystalline γ-Al2O3 support (average size ~40 nm) is added. After drying, the encapsulating ligands are eliminated by annealing in O2 at 375 °C for 24 h. Different NP shapes can be obtained by tuning the molecular weight of the different blocks of the encapsulating polymer, the metal/P2VP ratio (micelle loading), and the post preparation annealing treatment and atmosphere.20 Further details on the sample preparation method and synthesis parameters can be found in Refs. 20–22 and Table I. The NP diameters obtained by high-angle annular dark-field scanning transmission electron microscopy (TEM) are: 1.0 ± 0.2 nm for S1, S2, and S4, and 0.8 ± 0.2 nm for S3.20

B. Structural characterization (EXAFS)

EXAFS data were acquired at beamline X18B of the NSLS at Brookhaven National Laboratory in transmission mode using the Pt L3 edge. The EXAFS samples were pre-
pared by pressing the Pt/γ-Al2O3 powders into thin pellets which were mounted in a sample cell that permitted sample heating via an external proportional-integral-derivative controller, liquid-nitrogen cooling, as well as the continuous flow of gases during data acquisition and online mass spectrometry analysis. A bulk Pt foil was measured simultaneously with all samples (in reference mode) for energy alignment and calibration purposes. Multiple scans (up to 6) were collected at each temperature of interest and averaged in order to improve the signal-to-noise ratio. Measurements were done at different temperatures under H2 (50% H2 balanced with He for a total flow rate of 50 ml/min) or He (100% He flow) atmospheres. Samples S1, S2, S4, and a Pt foil were measured in H2 while S3 was measured in H2 as well as in He.

Data processing and analysis with the IFEFFIT package was conducted by analyzing the first shell Pt-Pt contribution for all temperature data concurrently, as described in Refs. 9 and 24, with the exception that the EXAFS Debye-Waller factors were obtained following the correlated Debye model (CDM) (Ref. 25) instead of the Einstein model. The CDM was used to extract the dynamic mean-square relative bond-length disorder, σs2, from the analysis of temperature-dependent EXAFS data, and through that θp. The total bond-length disorder σs2 also includes a configurational, or static, σc2 contribution: σs2 = σc2 + σp2. Due to the linear dependence of the total disorder on temperature in the classical regime (i.e., when the temperature of the measurement is greater than the Einstein temperature, ~190 K), σp2 can be estimated either by extrapolating the total disorder factor to zero temperature, or from the fit using the CDM (the present approach). The following parameters were varied in the EXAFS analysis: the corrections to the photoelectron energy origin, the nearest-neighbor bond lengths, the third cumulant of the first-NN (1NN) pair distribution function, and the values of θp and σc2.24 We carefully investigated the effect of the energy origin (E0) on the values of the structural parameters (distances and third cumulants) extracted from the fits since they correlate. In most of the samples the x-ray absorption near-edge structure portion of the absorption coefficient exhibits a small shift (approximately 1–1.5 eV, depending on the sample) toward lower energies with increasing temperature. The same effect was observed earlier by Sanchez et al.9 To compensate the effect of this shift on the change in the phase of the EXAFS signals we compared two different fitting procedures: (1) the correction to the energy origin was varied independently for each temperature and (2) all the data were aligned in energy before extracting the EXAFS signals, and subsequently the energy origin correction, same for all data that were analyzed concurrently, was varied during the fitting routine. The best-fit results for these two approaches were very close (within the uncertainties). Supplementary Tables I and II (Ref. 26) contain details of the fitting parameters used for the analysis of EXAFS spectra acquired at different temperatures presented in our paper, in particular, the number of relevant independent data points, the number of variables, factors reflecting the fit quality (reduced χ2 and the r factor), and values of the third cumulant.

Quantitative determination of the NP shape was carried out by analyzing low-temperature EXAFS data up to the fourth-NN (4NN) contribution, including multiple scattering, as described in Ref. 20, and references therein. Degeneracy of single- and multiple-scattering paths in EXAFS analysis is simply related to the coordination number of the nearest neighbors, and can be obtained from the nonlinear least-squares fit of the theoretical EXAFS equation to the experimental data.23 Physically reasonable constraints between the fitting variables were applied to maximize the number of degrees of freedom in the fits. The passive electron reduction factor was fixed to that found for the Pt foil used as reference: 0.861. Only the most important single- and collinear multiple-scattering paths were used in the fits through the fourth coordination shell. Among the multiple-scattering paths we used double- and triple-scattering paths to the 4NN through the intervening atom (first neighbor, or 1NN) along the same line. The total Debye-Waller values (σs2) for these paths were constrained to be the same as that of the 4NN Pt-Pt pair. In addition, a double-scattering path connecting the central atom, its 1NN, and another 1NN in the opposite direction was used in the fit. The disorder parameter of this path was varied independently from the others. Distance corrections were constrained to vary in accordance with the isotropic lattice expansion or contraction for all paths except for the first one, for which the third cumulant of the pair distribution function accounting for its asymmetry was varied as well. This approximation is justified by the apparent strong structural order in all particles, evident by comparing raw data in r space for all the samples and the reference bulk Pt.

### Table I: Description of synthesis parameters and size (TEM) information of micellar Pt NPs supported on nanocrystalline γ-Al2O3.

The TEM values correspond to NP diameters (D) and the parameter “L” represents the metal-salt to polymer head (P2VP) ratio used in the NP synthesis.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Polymer</th>
<th>L</th>
<th>Annealing T (°C) [% O2]</th>
<th>TEM analysis after annealing</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>PS(16000)-P2VP(3500)</td>
<td>0.05</td>
<td>375 [50]</td>
<td>1.0 (2)</td>
</tr>
<tr>
<td>S2</td>
<td>PS(16000)-P2VP(3500)</td>
<td>0.1</td>
<td>425 [70]</td>
<td>1.0 (2)</td>
</tr>
<tr>
<td>S3</td>
<td>PS(27700)-P2VP(4300)</td>
<td>0.1</td>
<td>375 [50]</td>
<td>0.8 (2)</td>
</tr>
<tr>
<td>S4</td>
<td>PS(27700)-P2VP(4300)</td>
<td>0.2</td>
<td>375 [50]</td>
<td>1.0 (2)</td>
</tr>
</tbody>
</table>
TABLE II. Model polyhedron shapes for fcc-Pt NPs providing the best fit to the experimental coordination numbers (EXAFS) and volume-weighted NP diameters (TEM). The volume-weighted TEM diameters (D), the total number of atoms in each NP (Nt), the ratio of the number of Pt atoms at the NP surface and perimeter to Nt (Ns/Nt), and the ratio of the number of Pt atoms in contact with the support to Nt (Nc/Nt) are given. A factor (Q) representing the deviations of the coordination numbers and diameter of each of the model shapes with respect to the experimental samples are also included.

<table>
<thead>
<tr>
<th></th>
<th>S1 - weighted D: 1.1 (2) nm</th>
<th></th>
<th>S2 - weighted D: 1.2 (3) nm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nt</td>
<td>Ns/Nt</td>
<td>Nc/Nt</td>
</tr>
<tr>
<td>33</td>
<td>0.82</td>
<td>0.55</td>
<td>0.8</td>
</tr>
<tr>
<td>46</td>
<td>0.78</td>
<td>0.54</td>
<td>1.2</td>
</tr>
<tr>
<td>43</td>
<td>0.76</td>
<td>0.58</td>
<td>1.2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>S3 - weighted D: 0.9 (2) nm</th>
<th></th>
<th>S4 - weighted D: 1.2 (2) nm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nt</td>
<td>Ns/Nt</td>
<td>Nc/Nt</td>
</tr>
<tr>
<td>44</td>
<td>0.84</td>
<td>0.23</td>
<td>0.3</td>
</tr>
<tr>
<td>70</td>
<td>0.72</td>
<td>0.26</td>
<td>1.1</td>
</tr>
<tr>
<td>67</td>
<td>0.72</td>
<td>0.27</td>
<td>1.1</td>
</tr>
</tbody>
</table>

foil [Fig. 5(b) of Ref. 20 and Fig. 1 of this work]. Coordination numbers of all single-scattering paths were varied independently; those of the multiple-scattering paths were constrained to those of the single-scattering paths to the 4NN or 1NN, depending on the geometry of a particular path. Finally, the same energy correction (\(E_0\)) was used for all paths. Supplementary Table III displays coordination numbers and distances of each coordination shell to the absorbing atom obtained from the multiple-scattering analysis of EXAFS data acquired for sample S3.20 The data were obtained at 173 K in \(H_2\) after NP reduction. Reference 20 contains analogous information for the rest of the samples discussed in the present paper.

The possibility of morphological changes in our samples as a function of temperature has not been accounted for in our analysis. This approach decreases the number of fitting parameters and is justified by the following arguments: (i) the behavior of our raw EXAFS data throughout the entire temperature range is not consistent with a size or shape change, (ii) our samples were measured under a hydrogen atmosphere that passivates the NPs and stabilizes their shapes, and (iii) the analysis of the vibrational properties of our NPs was conducted up to a maximum annealing temperature of 375 °C, which is the same temperature used for sample calcination (for 24 h) prior to the EXAFS measurements. If any changes in the NP morphology (size and/or shape) were to occur at 375 °C, they should have already taken place before the EXAFS measurements. Evidence of the lack of NP sintering at this annealing temperature can be
C. Nanoparticle shape modeling

The shapes of our Pt NPs have been resolved by matching structural information obtained experimentally via EXAFS (coordination numbers up to the fourth-nearest neighbor, \(N_1-N_4\)) and TEM (NP diameter, \(D\)) to analogous data extracted from a database containing ~4000 model fcc NP shapes. Only closed shell clusters were considered in the models. In our analysis, after taking into consideration the error bars in the EXAFS coordination numbers as well as the TEM diameters, we have found on average only three-model fcc cluster shapes consistent with all five experimental parameters (\(N_1, N_2, N_3, N_4, \) and \(D\)). Table II shows the best three-model shapes for each of the samples investigated together with the total number of atoms, the ratio of the number of Pt atoms at the NP surface, and perimeter to \(N_1(N_c/N_1)\), the ratio of the number of Pt atoms in contact with the support to \(N_1(N_c/N_1)\), and a factor \(Q\) indicating how much each of the specific model shapes deviates from the experimentally measured EXAFS coordination numbers and volume-weighted TEM NP diameters. The shapes with the best \(Q\) factors are depicted in a darker color in Table II and appear as inset in Fig. 3. The \(Q\) factor has been obtained following Eq. (1):

\[
Q = \sqrt{\frac{NE_1^2 + NE_2^2 + NE_3^2 + DE^2}{4}},
\]

where \(NE_1, NE_2,\) and \(NE_3\) are normalized errors of the experimental first, second, and third coordination numbers, and \(DE\) is the normalized error of the TEM diameter. The latter errors are calculated following Eqs. (2) and (3):

\[
NE = \frac{\text{Experimental } CN - \text{Model } CN}{\text{Experimental error of } CN},
\]

\[
DE = \frac{\text{Experimental } D - \text{Model } D}{\text{Experimental error of } D},
\]

where \(D\) is the NP diameter and \(CN\) are the respective coordination numbers (\(N_1, N_2,\) or \(N_3\)). In the calculation of the \(Q\) factor we have not considered the fourth coordination number extracted from EXAFS. As shown by Jentys,\(^27\) either \(N_3\) or \(N_4\) can be used interchangeably to estimate the particle shape, but since \(N_3\) is generally larger than \(N_4\) for most shapes of fcc particles (e.g., 24 vs 12 in the bulk), the former number has greater impact on the shape determination.

A low-\(Q\) factor indicates a good fit between experimental and model \(CN\) and \(D\). As can be seen in the Table II, only one shape provides a \(Q\) factor \(\leq 1\) for samples S1 and S3, while as few as three shapes satisfy that constraint for S2 and S4, indicating the low degeneracy in the determination of the cluster shape for samples containing NPs in our size range (~0.8–1 nm). For S4, two shapes with identical \(Q\) factor were obtained. The final shape selected to be displayed as inset in Fig. 3 is a shape (85-atom NP) that we have also experimentally resolved on identically prepared Pt NPs supported on TiO\(_2\) (110) via scanning tunneling microscopy (STM, not shown here). For the rest of the samples, all shapes characterized by the lowest \(Q\) factors in Table II have also been observed by our group via STM on analogous micellar Pt NPs. Examples of the NP shapes resolved by STM can be found in Refs. 20 and 28.

III. RESULTS

EXAFS spectra in (a) \(k\) space and (b) \(r\) space from a ligand-free micellar Pt NP sample supported on \(\gamma\)-Al\(_2\)O\(_3\) (S3) acquired in \(H_2\) at temperatures ranging from 173 to 648 K are shown in Fig. 1. A decrease in the intensity of the EXAFS signals with increasing temperature is evident. Such effect can be attributed to an increase in the bond-length disorder in the framework of Pt atoms, to a decrease in the Pt-Pt first-nearest-neighbor coordination number due to changes in the NP size and/or shape, or to both. We can single out the disorder as the dominant factor responsible for this behavior when we examine the \(k\)-space data, Fig. 1(a). If
the NP size and/or shape were changing with temperature, the coordination number effect would be visible for all \( k \) values, since it affects the EXAFS signal as a constant multiplicative factor. However, the decrease in the intensity of our \( k \)-space data is more prominent at the end of the \( k \) range than at the beginning. Such observation is in agreement with the disorder effect since it is expected to increase with increasing \( k \).

A representative multiple-scattering fit of the 173 K spectrum is included as inset in Fig. 1. Supplementary Figs. 1 and 2 show EXAFS data for the remaining samples at all temperatures in \( k \) space, and representative fits in \( r \) space to temperature-dependent data from S2. Supplementary Fig. 3 displays the imaginary part of the \( k^2 \)-weighted EXAFS data, \( s \), and residual contribution obtained for S2 and S3 at 173–183 K.

Figure 2 displays (a) Pt-Pt distances (\( R \)) and (b) the dynamic bond-length disorders (\( \sigma^2_c \)) in the temperature range of 150–700 K for samples S1–S4 and for a reference bulk like Pt foil. All measurements were carried out under a continuous H \(_2\) flow, with the exception of sample S3, for which a He and H \(_2\) adsorbate effect was clearly observed [inset, Fig. 2(a)]. The data in Fig. 2(a) correspond to the best-fit results of the experimental EXAFS spectra obtained for the INN Pt-Pt bond lengths (\( R \)) at different temperatures.

The dynamic bond-length disorders (\( \sigma^2_c \)) obtained from the fits of the experimental EXAFS data following the CDM (Ref. 25) are shown in Fig. 2(b). In the high-temperature approximation \( \sigma^2_c = k_b T / k \sim T / \theta_D \), with \( k \) being the effective force constant. Hence, the slope of the temperature dependence of \( \sigma^2_c \) depends on \( \theta_D \), with larger Debye temperature giving smaller slopes. The data in Fig. 2(b) reveal a striking behavior of our NPs in this regard, which will be discussed in greater detail below. The best-fit values of \( \sigma^2_c \) are shown in Table III together with the first-nearest-neighbor coordination numbers (\( N_1 \)) and \( \theta_D \).

Figure 3 displays the Debye temperatures obtained from the CDM analysis of our EXAFS data [Fig. 2(b)] as a function of \( N_1 \). Samples S1, S2, and S4 have the same TEM diameter (1.0 ± 0.2 nm) but different shapes. The data correspond to samples passivated by H \(_2\) under identical conditions.

### IV. DISCUSSION

The thermal-expansion coefficients (\( \alpha \)) of the Pt NP samples were estimated from the slope of the linear interpolation of the data in Fig. 2(a), Table III. Changes in the magnitude and sign of \( \alpha \) are observed when comparing the different samples. As expected, the Pt foil displays a positive thermal expansion. Surprisingly, only two of our samples (S2 and S4) follow the same trend, although a smaller \( \alpha \) than for bulk Pt is obtained, suggesting NP stiffening. The remaining samples (S1 and S3) show negative thermal expansion with a more prominent effect observed for the sample containing the NPs with the smallest TEM diameter (~0.8 nm, S3). The negative slope of the \( R \) vs \( T \) curves in supported Pt NPs represents markedly nonbulk behavior, which can be attributed to the competing interactions of cluster-support and cluster-adsorbate (H \(_2\) or He in our case) effects, both depending on the NP size and shape. The effect of H adsorbates on the Pt-Pt spacing is expected to cause relief of both the surface tension and compressive stress which is always present in NPs, and thus increase their average bond lengths. We postulate that the extent of adsorbate-induced relief can be correlated with the surface-to-volume ratios of different NPs. This effect is demonstrated in the inset of Fig. 2(a), where larger \( R \) distances were measured for S3 under H \(_2\) as compared to He. A more prominent negative thermal expansion was observed for S3 under He. Sample S3 has the largest surface-to-volume ratio (Table III) among all samples and consequently is expected to be most affected by the presence of adsorbates. An alternative mechanism proposed to explain negative thermal expansion for materials characterized by strong bonding anisotropy relates to enhanced atomic displacements in the transverse direction to the bond.

Analysis of the EXAFS data in Fig. 2(b) obtained for the Pt foil via the CDM lead to a \( \theta_D \) of 244 ± 3 K, which is in...
agreement with literature reports. Surprisingly, significantly higher Debye temperatures (262–284 K) were observed for samples S1, S2, and S4 (~0.8 nm in diameter). However, a similar \( \theta_D \) value to that of the bulk Pt was observed for the samples with the smallest NPs (S3, \( \sim 0.8 \) nm, \( \theta_D = 250 \pm 5 \) K). If we compare the Pt-Pt bond lengths in Fig. 2(a) for different NP samples at the lowest temperatures, i.e., when the hydrogen coverages, and hence the ordering of NP facets, are the greatest, we can see that samples S1 and S2, which are characterized by the highest \( \theta_D \) (Table III and Fig. 3), have the smallest values of Pt-Pt bond lengths. This suggests some correlation between increased compressive strains and enhanced \( \theta_D \). In fact, due to the asymmetry of the effective pair potential, where the short distance branch is steeper than the long distance branch (due to the hard-core repulsion), the Pt-Pt bond stiffness should be large in the systems having bonds with large compressive strain, namely S1 and S2. It should also be mentioned that the compressive strain present in supported NP systems might have different origins as, for example, the presence of ligands on the NP surface (not the present case), adsorbates, or a support (\( \gamma \)-Al\(_2\)O\(_3\)). However, the explanation of the enhanced \( \theta_D \) observed appears to be more complex than a direct one-to-one correlation with a contraction of Pt-Pt distances, \( R \). Such contraction is theoretically predicted for free (unsupported) clusters due to surface tension arguments while the same systems show \( \theta_D \) smaller than the corresponding bulk metals. In fact, based on EXAFS measurements, Yokoyama et al. reported metal-metal bond contractions for small Ag and Pd NPs (1.3–1.5 nm) accompanied by reduced \( \theta_D \) with respect to the bulk. Also, the above explanation for the enhanced \( \theta_D \) values in terms of a contraction of \( R \) does not take into account the distinct differences (outside of the error bars) observed in \( \theta_D \) for the three samples (S1, S2, and S4) characterized by the same TEM diameter (~1.0 nm).

In order to gain further insight into the origin of the anomalous behavior of the Debye temperature of our NP samples we should consider their different geometrical structures. As was described in detail in the Results section, the shapes of our Pt NPs have been resolved by matching structural information obtained experimentally via EXAFS (\( N_1 \), \( N_2 \), \( N_3 \)) and TEM (NP diameter, \( D \), obtained from Z-contrast images) to analogous data extracted from a database containing model fcc NP shapes. The most representative shapes for each of the samples are a truncated octahedron (S1), cuboctahedron (S2), and octahedron (S3 and S4). As can be seen in Fig. 3, all samples but sample S1 contain Pt NPs with a three-dimensional (3D) shape. This conclusion is in agreement with the static disorder values displayed in Table III, since the largest \( \sigma^2 \) was obtained for S1 (~0.0028 A\(^2\)), which according to our model-shape analysis is the sample with the highest contact area with the support [two-dimensional (2D) shape]. The low \( \sigma^2 \) value of ~0.0010 A\(^2\) obtained for our smallest NPs (S3) is consistent with their 3D shape, with a relatively large surface area available for hydrogen adsorption, leading to relaxation and NP faceting.

If one compares S3 and S4, with identical octahedron shapes but different sizes, we can see that the smallest NPs (S3) displayed also the smallest Debye temperature. We at-
ANOMALOUS LATTICE DYNAMICS AND THERMAL PROPERTIES OF SUPPORTED Pt NPs

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V. CONCLUSIONS

Our study reveals an overall significant shortening of the Pt-Pt bond lengths and a drastic increase in the Debye temperatures of ligand-free, size-, and shape-selected Pt NPs synthesized by inverse micelle encapsulation and supported on nanocrystalline γ-Al₂O₃. We attribute the origin of the enhanced θₖ of our NPs as compared to bulk Pt to compressive stress from the NP/support interface as well as to inhomogeneous strain caused by competing contributions from atoms at the NP surface (leading to relaxation and reduced θₖ) and in the interior of the NPs (experiencing enhanced stiffening leading to increased θₖ).

In this work we have demonstrated that the lattice dynamics and thermal behavior of NPs is affected by their geometric properties. Our experimental data provide important validation for nanothermodynamics theories that, as we have shown qualitatively in this work, should incorporate geometrical effects (size and shape) and environmental interactions (e.g., adsorbate and support) to adequately describe thermal properties at the nanoscale.

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26 See supplementary material at http://link.aps.org/supplemental/10.1103/PhysRevB.82.155450 for further details on the EXAFS parameters used for the fit of the experimental EXAFS data, best fit results, fit quality, as well as additional EXAFS spectra in \( \kappa \)- and \( r \)-space and corresponding fits.