

anchored to the air/solution interface. From this result is easily inferred a folding of the alkanediyl chain and its penetration into the inner micelle. The folding can also be examined by the following approach. The partial molar volume  $v$  of tetradecane-sulfonate anion in the micellar state is estimated to be  $254.4 \text{ cm}^3 \text{ mol}^{-1}$ <sup>20</sup> by applying the Stokes' radius of  $1.84 \text{ \AA}$  to the sodium ion. Then, the micellar surface area ( $S$ ) made only of surfactant ions is given by

$$S = (4\pi)^{1/3}(3\bar{n}v)^{2/3} \quad (7)$$

where the micelles are assumed to be spherical due to relatively small aggregation numbers. On the other hand, the distance ( $d$ ) between anionic head groups is related with the above surface area in the form

$$S = \left(\sqrt{3}/2\right)\bar{n}d^2 \quad (8)$$

where the hexagonal packing is assumed as for the anionic groups.

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From the  $v$  and  $\bar{n}$  values obtained above, the distance  $d$  can be evaluated for each counterion. At the same time, the distance between two cationic charges in the counterions is also evaluated from the CPK model. These two distances are plotted against the carbon number of the alkanediyl group of the counterions (Figure 6). From this figure, the distance between the two cationic charges of the counterions becomes longer than that between anionic head groups of micellized surfactant ions above an eight  $\text{CH}_2$  groups separation of counterions, which also strongly indicates the folding of the alkanediyl chain of the counterions.

The above two simplified calculations substantiate the assumption made from the cmc change in the previous paper<sup>12</sup> that the divalent cationic charges whose separation is more than eight methylene groups become anchored to the micellar surface due to electrostatic attraction by anionic head groups of surfactant ions, while the alkanediyl chain folds and penetrates into a hydrophobic inner micelle, leading to a decrease in cmc.

Registry No. C<sub>2</sub>BP, 46347-63-1; C<sub>4</sub>BP, 33884-65-0; C<sub>6</sub>BP, 80038-42-2; C<sub>8</sub>BP, 80038-44-4; C<sub>10</sub>BP, 47244-14-4; C<sub>12</sub>BP, 47377-62-8; C<sub>14</sub>BP, 47510-98-5.

## Reduction and Cluster Growth of Palladium in Zeolite Y Containing Transition Metal Ions. X-ray Absorption Studies

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The effect of a second transition metal on the reducibility and agglomeration behavior of palladium in zeolite Y is studied by EXAFS spectroscopy. Special attention is given to the potential bonding interactions between both metal constituents that could result in "chemical anchoring" of the noble metal to the support via the unreduced cocation. Zeolite Y was coexchanged with  $\text{Fe}^{2+}$  or  $\text{Co}^{3+}$  and  $\text{Pd}^{2+}$  and with each ion alone. Dehydration in oxygen atmosphere and reduction with hydrogen was performed at different temperatures up to 623 K. The structure of the samples during various stages of pretreatment and reduction was studied with EXAFS on both absorption edges. Palladium is partially reduced to  $\text{Pd}^0$  at room temperature and forms small metal clusters at higher temperatures. Iron is oxidized under oxygen atmosphere and is partially present in the form of iron oxide particles, which are redispersed under reducing conditions. A small enhancement of the reducibility of palladium is observed if iron is present in the zeolite, but neither a chemical anchoring nor an effect on the final agglomeration process of palladium is detected by the presence of iron cations. In contrast, coexchange with cobalt results in a substantially higher dispersion of the  $\text{Pd}(0)$  phase after high-temperature reduction. Apparently, the change in cation distribution induced by the Co ions influences the reduction process of palladium. When palladium ions are diffused into the zeolite containing aqueous  $\text{Fe(II)}$  ions, a  $\text{Pd}(0)$  phase consisting of larger crystallites is formed at room temperature by a redox reaction with  $\text{Fe(II)}$ .

### Introduction

The preparation of supported small metal particles continues to attract wide interest due to its importance in catalytic applications. Numerous studies have shown the sometimes dramatic effect of a second metal on the catalytic behavior of a metal catalyst.<sup>1-7</sup> Structural studies on the formation of metal/support catalyst systems under different conditions are a prerequisite for

their rational design. X-ray diffraction (XRD) and transmission electron microscopy (TEM) allow determination of the presence of different phases and particle sizes but have limited utility when amorphous metals and/or clusters of dimensions smaller than  $20 \text{ \AA}$  are present. In this context, extended X-ray absorption spectroscopy (EXAFS) is complementary to X-ray diffraction studies since it is most sensitive for the local structure in a range smaller than about  $5 \text{ \AA}$  around the absorber atom.

EXAFS spectroscopy has evolved as a powerful technique to analyze materials lacking long-range crystalline order, such as amorphous materials, thin films, liquids, etc.<sup>8,9</sup> Element-specific structural information can be obtained from the modulations of the X-ray absorption coefficient: Photoelectrons excited by X-ray absorption are backscattered from atoms in the first few neigh-

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boring shells. The resulting interference pattern of the absorption coefficient contains information about the number and nature of the surrounding neighbors as well as their distances to the absorber atom. A radial distribution function is obtained through Fourier transformation of the normalized EXAFS modulation from momentum space ( $k$ , Å<sup>-1</sup>) into the distance space ( $r$ , Å).

Supported bimetallic catalysts containing a variety of transition metals have been studied with EXAFS, e.g., Cu-Ru, -Os, -Rh, -Pd; Ir-Pt, -Rh;<sup>10</sup> Rh-Co;<sup>11,12</sup> Ni-Cu;<sup>13</sup> and Pt-Mo.<sup>14</sup> In many cases, the formation of alloys was observed, and frequently the bulk composition differed from that on the surface. A common problem encountered with supported metal particles is the migration and agglomeration of the metal phase at high-reduction and/or -reaction temperatures. It was proposed by Yermakov and co-workers that the introduction of a second nonreducible transition metal could limit undesired agglomeration by Coulomb interactions with the noble catalyst metal. The reduced transition metal would be "chemically anchored" to the support through an  $M^{n+}-M^0$  interaction. Sachtler and co-workers suggested that such an effect was responsible for their observation of reduced particle sizes when Fe was added to Pt supported in zeolite Y.<sup>15</sup> They found a similar increase in metal dispersion when chromium was coexchanged into rhodium zeolites, and EXAFS measurements at the Rh K-Edge indicated an interaction between chromium ions and Rh particles.<sup>16</sup>

There are only few EXAFS studies where a potential anchoring effect by a second metal was examined directly by probing the local environment around *both* metal constituents. Tin(II) ions in an Sn/Pt-alumina system were reported to be located at the interface between the support and Pt(0) clusters.<sup>17</sup> A similar interaction was found for Fe<sup>3+</sup> and Rh<sup>0</sup> on silica at Fe/Rh ratios up to 0.3.<sup>18</sup> Higher ratios resulted in the reduction of iron to Fe<sup>0</sup> which covered the surface of the rhodium particles. Both reports describe systems reduced at high temperatures. Our recent EXFAS studies of a related Co/Pd coexchanged zeolite system showed that improved dispersion is not caused by chemical anchoring but by a modified cation distribution that affects the reducibility of the palladium ions.<sup>19</sup> In order to clarify the origin of the effects of a nonreducible second metal, we report an investigation of the interaction between iron and palladium cations coexchanged into zeolite Y. The preparation of small palladium particles in a zeolite matrix has been subject of many studies. TPD-MS and thermogravimetric analysis (TGA) of the reduction mechanism of [Pd(NH<sub>3</sub>)<sub>4</sub>]<sup>2+</sup> in zeolite X and Y showed that an autoreduction process, caused by the decomposing ammine ligands, can be suppressed with oxygen atmosphere during the dehydration step.<sup>20,21</sup> Bergeret et al. observed that a pretreatment between 570 and 870 K in oxygen and subsequent reduction at low temperature leads to atomic dispersion of a portion of the palladium.<sup>22</sup> EXAFS studies demonstrated the formation of small metal particles when the Pd(II)-exchanged zeolite was dehydrated under

vacuum, while oxygen treatment resulted in the coordination of palladium cations to the zeolite surface.<sup>23</sup> It was found that palladium clusters of 2-4 atoms were stabilized by the zeolite X matrix.<sup>24</sup> In the present study, we compare the reduction and Pd particle formation in Pd-Y zeolite with the corresponding bimetallic Fe/Pd-Y systems to explore a possible influence of the iron phase on the reduction and agglomeration behavior of Pd. We follow the changes in oxidation state and coordination environment of *both* constituents during reduction by monitoring the iron and palladium K-edge absorption spectra. This approach provides the maximal information about any possible interaction of both constituents.

### Experimental Section

Five different zeolite sample sets based upon zeolite Y (Linde LZ-Y52) were used in this study. Single-metal-exchanged samples were prepared as follows: Zeolite Pd-Y was obtained by aqueous cation exchange with [Pd(NH<sub>3</sub>)<sub>4</sub>]<sup>2+</sup> (0.01 M). Sample Fe-Y was prepared by an anaerobic exchange with FeSO<sub>4</sub> (0.01 M) in degassed, slightly acidified water (pH = 4). The samples were washed, filtered, and subsequently dehydrated in a flow of oxygen at 623 K, at a heating rate of 1 K/min. After evacuation, small portions of the samples were reduced with hydrogen at room temperature (600 Torr) and at 423 and 623 K in a flow reactor.

Zeolites containing Pd and Fe were prepared according to three different sequences: Sample FePd-Y resulted from hydrated Fe-Y that was subsequently exchanged with [Pd(NH<sub>3</sub>)<sub>4</sub>]<sup>2+</sup>. Sample Fe<sup>III</sup>Pd-Y was derived from Fe-Y, dehydrated at 623 K under oxygen, and subsequently ion-exchanged with [Pd(NH<sub>3</sub>)<sub>4</sub>]<sup>2+</sup>. Sample PdFe-Y was prepared in the reversed sequence, i.e., after exchanging with Pd<sup>2+</sup> the zeolite was dehydrated in an oxygen flow at 475 K and subsequently exchanged with FeSO<sub>4</sub> under anaerobic conditions. All three sample sets were degassed under oxygen at 623 K, evacuated, and reduced with hydrogen at 295, 423, and 623 K. Subsequent evacuation was performed at the reduction temperatures to pressures smaller than 10<sup>-3</sup> Torr. Sample Fe<sup>III</sup>Pd-Y was also reduced at 823 K.

Chemical analysis (Galbraith Laboratories) gave the following zeolite unit cell compositions:

Fe-Y	Na <sub>45</sub> Fe <sub>6</sub> (AlO <sub>2</sub> ) <sub>57</sub> (SiO <sub>2</sub> ) <sub>135</sub>
Pd-Y	Na <sub>33</sub> Pd <sub>12</sub> (AlO <sub>2</sub> ) <sub>57</sub> (SiO <sub>2</sub> ) <sub>135</sub>
Fe <sup>III</sup> Pd-Y	Na <sub>18</sub> Fe <sub>8.7</sub> Pd <sub>10</sub> (AlO <sub>2</sub> ) <sub>57</sub> (SiO <sub>2</sub> ) <sub>135</sub>
PdFe-Y	Na <sub>29</sub> Fe <sub>3.8</sub> Pd <sub>10</sub> (AlO <sub>2</sub> ) <sub>57</sub> (SiO <sub>2</sub> ) <sub>135</sub>

Zeolite powders were embedded in an octadecane/eicosane mixture and kept under nitrogen atmosphere until the EXAFS measurements were carried out. EXAFS experiments were performed at the X-11A beam line at NSLS (Brookhaven National Laboratory) with an electron energy of 2.5 GeV and ring currents between 40 and 120 mA. Data were collected with a Si(400) crystal pair monochromator at the Pd K-edge (24 350 eV) and with a Si(111) monochromator at the Fe K-edge (7112 eV) at 100 K. Samples Fe<sup>III</sup>Pd-Y (reduced at 623 and 873 K) and PdFe-Y (reduced at 623 K) were measured at the Pd K-edge at a later time at X-11A with a Si(111) monochromator. Data analysis was performed following standard procedures.<sup>25</sup> Backscattering amplitude and phase functions used in the least-squares fitting routine were extracted from the following reference materials and used as listed (see text): Pd foil (fcc), 1st Pd-Pd shell,  $N = 12$ ,  $R = 2.744$  Å; 3rd Pd-Pd shell,  $N = 24$ ,  $R = 4.76$  Å; 4th Pd-Pd shell,  $N = 12$ ,  $R = 5.50$  Å; 7th Pd-Pd shell,  $N = 8$ ,  $R = 7.26$  Å; PdO powder (tetragonal), Pd-O,  $N = 4$ ,  $R = 2.017$  Å; Fe<sup>III</sup>acac (acac = acetylacetonate), Fe-O,  $N = 6$ ,  $R = 1.992$  Å; FeO powder (NaCl structure), Fe-Fe,  $N = 12$ ,  $R = 3.063$  Å; MnS powder (NaCl structure), Mn-S for Fe-Si,  $N = 6$ ,  $R = 2.61$  Å; Co powder (hcp), Co-Co for Pd-Fe,  $N = 12$ ,  $R = 2.514$  Å.

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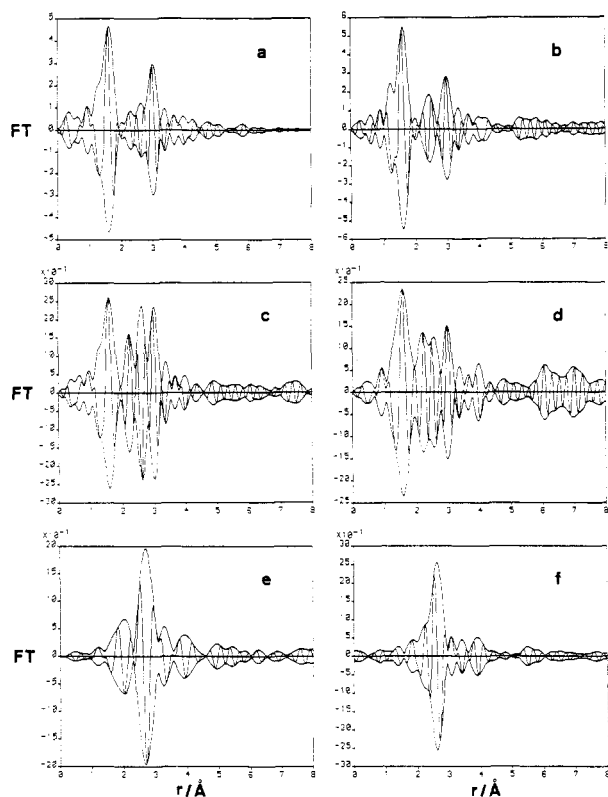
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**Figure 1.**  $k^3$ -Weighted Fourier transforms of the Pd-edge of sample Pd-Y (a,c,e) and  $\text{Fe}^{\text{III}}\text{Pd-Y}$  (b,d,f) after dehydration under oxygen atmosphere at 623 K (a,b) and hydrogen reduction at room temperature (c,d) and 423 K (e,f).

## Results and Discussion

**Pd-Edge EXAFS.** In Figure 1, the EXAFS spectra of palladium-exchanged zeolites Pd-Y (parts a, c, and e) are compared to those of  $\text{Fe}^{\text{III}}\text{Pd-Y}$ , which were iron-exchanged, dehydrated in oxygen, and subsequently exchanged with palladium (parts b, d, and f). Spectra of PdFe-Y are very similar to those of  $\text{Fe}^{\text{III}}\text{Pd-Y}$  and are therefore not included. Structural parameters obtained from the data analysis are listed in Table I. The Fourier transforms were  $k^3$ -weighted over a range of 2.8–14  $\text{\AA}^{-1}$  and show the samples after treatment at 632 K in oxygen atmosphere, reduction with hydrogen at room temperature, and reduction at 423 K. Peak positions are not corrected for phase shift effects (0.3–0.5  $\text{\AA}$ ), but true bond distances are obtained from the fitting process with model compounds.

**Dehydration under Oxygen Atmosphere.** Samples measured directly after ion exchange show the same ligand environment of Pd as the aqueous precursor ion  $[\text{Pd}(\text{NH}_3)_4]^{2+}$ , as described earlier<sup>23</sup> (spectra not shown). Upon dehydration under oxygen atmosphere, the ammine ligands are split off and the palladium cations coordinate to the zeolite lattice. This is demonstrated in parts a and b of Figure 1 for samples Pd-Y and  $\text{Fe}^{\text{III}}\text{Pd-Y}$ , where the first strong peak represents the oxygen atoms of Pd-O atom pairs. The palladium ions are coordinated on average to 2.8 nearest oxygens in both samples with a bond length of 1.99  $\text{\AA}$ . Concomitant to the oxygen peak, an outer-shell contribution appears at about 3  $\text{\AA}$  (uncorrected). Back-transformation and inspection of the amplitude function reveals that it originates from a light element, such as Si or Al atoms from the zeolite framework. Similar observations have been made upon dehydration of other cation-exchanged zeolites, and this feature can be considered as an indication for the migration of cations into defined coordination sites in the zeolite framework.<sup>26–28</sup>

**TABLE I: Structural Data As Derived from EXAFS Analysis, Pd Absorption Edge**

sample	atom pair	$R^a/\text{\AA}$	$N^b$	$\Delta\sigma^{2c}/\text{\AA}^2$	$\Delta E^d/\text{eV}$
623 K Oxygen					
Pd-Y	Pd-OZ	1.99	2.8	0.000	-0.1
$\text{Fe}^{\text{III}}\text{Pd-Y}$	Pd-OZ	1.99	2.9	0.000	-0.1
	Pd-Pd	2.78	0.5	0.000	+11.9
PdFe-Y	Pd-OZ	1.98	3.2	0.000	+1.8
CoPd-Y	Pd-OZ	1.98	2.8	0.000	+1.7
295 K Hydrogen					
Pd-Y	Pd-OZ	1.98	1.9	0.002	-1.6
$\text{Fe}^{\text{III}}\text{Pd-Y}$	Pd-OZ	1.99	1.4	0.000	-0.1
PdFe-Y	Pd-OZ	1.99	1.2	0.000	-0.4
CoPd-Y	Pd-OZ	2.00	1.2	0.000	-1.6
	3. Pd-Pd	4.77	3.2	0.001	-5.2
	4. Pd-Pd	5.47	1.2	0.000	+0.6
	7. Pd-Pd	7.27	0.8	0.000	-2.2
423 K Hydrogen					
Pd-Y <sup>e</sup>	1. Pd-Pd	2.80	2.2	0.009	-1.0
$\text{Fe}^{\text{III}}\text{Pd-Y}$	1. Pd-Pd	2.83	2.6	0.007	+2.2
PdFe-Y	1. Pd-Pd	2.83	3.1	0.008	+3.0
CoPd-Y	1. Pd-Pd	2.74	2.3	0.001	-0.2
	3. Pd-Pd	4.76	3.3	0.001	-3.7
	4. Pd-Pd	5.45	1.5	0.001	-4.7
	7. Pd-Pd	7.28	0.8	0.001	-6.2
623 K Hydrogen					
Pd-Y <sup>e</sup>	1. Pd-Pd	2.74	8.0	0.001	-5.9
	3. Pd-Pd	4.75	9.5	0.001	-3.4
	4. Pd-Pd	5.50	4.1	0.001	-6.4
	7. Pd-Pd	7.26	1.6	0.004	-3.7
$\text{Fe}^{\text{III}}\text{Pd-Y}$	1. Pd-Pd	2.73	8.8	0.004	-5.2
	3. Pd-Pd	4.76	8.0	0.003	-7.7
PdFe-Y	1. Pd-Pd	2.72	7.7	0.004	-2.2
CoPd-Y	1. Pd-Pd	2.75	5.0	0.001	-7.0
	3. Pd-Pd	4.77	7.6	0.002	-7.4
	4. Pd-Pd	5.50	3.7	0.002	-4.4
	7. Pd-Pd	7.29	1.2	0.000	-7.1
723 K Hydrogen					
$\text{Fe}^{\text{III}}\text{Pd-Y}$	1. Pd-Pd	2.73	8.6	0.003	-4.3

<sup>a</sup> Bond distances: accuracy,  $\pm 0.01$   $\text{\AA}$ . <sup>b</sup> Coordination numbers: accuracy,  $\pm 10$ –20%. <sup>c</sup> Deviations of the Debye-Waller factor: accuracy,  $\pm 10$ –20%. <sup>d</sup> Deviations from the inner potential. <sup>e</sup> The accuracy for the Pd-Pd atom pairs in sample Pd-Y is 10% for  $N$  and  $\Delta\sigma^2$ .

The oxygen neighbors of the palladium cations in sample Pd-Y and  $\text{Fe}^{\text{III}}\text{Pd-Y}$  are labeled Pd-OZ, where OZ stands for zeolite oxygen. The fit results suggest that the palladium ions are coordinated to three framework oxygens. This corresponds to cation siting in SI' (or SII, SII') positions in six-ring Si/Al windows of the zeolite lattice. X-ray diffraction studies of similarly treated Pd-Y zeolites revealed  $\text{Pd}^{2+}$ -OZ bond distances between 2.01 and 2.07  $\text{\AA}$  with all located cations in SI' positions.<sup>29,30</sup> We observe a slight difference between  $\text{Fe}^{\text{III}}\text{Pd-Y}$  and the Pd-exchanged sample Pd-Y. A small peak at about 2.5  $\text{\AA}$  can be isolated in  $\text{Fe}^{\text{III}}\text{Pd-Y}$  and fitted with the palladium foil reference. It indicates the onset of palladium reduction even under an oxidizing atmosphere. A Pd-Pd scatterer pair with a bond distance of 2.78  $\text{\AA}$  and average coordination number of 0.5 is found (Table I).

**Reduction at Room Temperature.** After reduction of Pd-Y and  $\text{Fe}^{\text{III}}\text{Pd-Y}$  at room temperature (Figure 1c,d), a more complex picture arises. Both samples still show the peaks corresponding to Pd-OZ and Pd-Si/Al contributions (note the smaller scale), but these are now obscured by other features with intermediate distances. The average Pd-OZ coordination number is reduced to 1.9 in Pd-Y and to 1.4 in  $\text{Fe}^{\text{III}}\text{Pd-Y}$ , while the bond distance remains at 1.99  $\text{\AA}$ . This is readily interpreted with a partial reduction of the  $\text{Pd}^{2+}$  ions to  $\text{Pd}^0$ . Provided that the unreduced  $\text{Pd}^{2+}$  ions remain bonded to these oxygen atoms, the degree of reduction can be calculated. Hence, about 35% of all Pd ions have

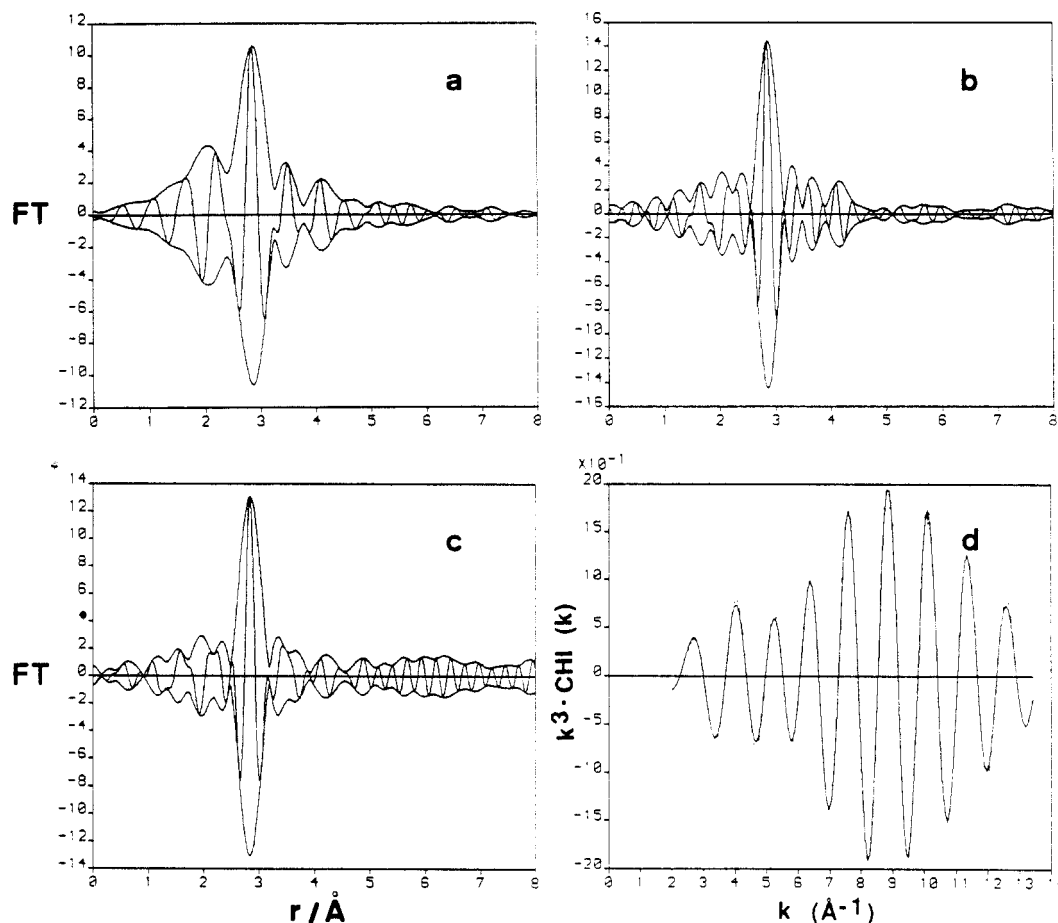
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**Figure 2.** Phase- and amplitude-corrected Fourier transforms ( $k^3$ -weighted; corrected with Pd foil) of Pd-edge spectra after reduction at room temperature of (a) Pd-Y, (b) Fe<sup>III</sup>Pd-Y, (c) PdFe-Y, and (d)  $k^3$ -weighted, isolated Pd-Pd EXAFS contribution of PdFe-Y (solid line) and fit (broken line).

been reduced in sample Pd-Y. The presence of iron increases the degree of reduction to about 55% in sample Fe<sup>III</sup>Pd-Y (60% in PdFe-Y). The additional peaks in the distribution junction can be explained by interactions of these reduced metal atoms with the zeolite.

**Reduction at 423 K.** Drastic changes occur when the samples Pd-Y and Fe<sup>III</sup>Pd-Y are treated at 423 K under hydrogen (Figure 1e,f). The oxygen contribution at 1.99 Å has totally vanished, indicating complete reduction of the palladium phase. New Pd-Pd interactions are visible in a peak at about 2.7 Å (uncorrected). The magnitude of this interaction is extremely small compared to bulk metal, indicating the formation of molecule-sized palladium clusters. No outer coordination shells associated with bulk palladium are detectable. However, small features on both sides of the Pd peak are observed in Pd-Y. An almost identical distribution function was obtained for Pd-X zeolites when reduced at room temperature, which has been described in detail.<sup>24</sup> The additional small peaks were found to arise from interactions of the palladium clusters with the stabilizing zeolite matrix in the form of a long Pd-OZ bond with a distance of 2.76 Å and a Pd-Si distance of 3.44 Å. Model calculations of the cluster arrangements show that two to four Pd atoms are arranged in the sodalite subunits in SI\* and SII\* positions, having a bond length Pd-OZ > 2.65 Å while detecting each other at a distance of 2.75 Å. A similar arrangement of reduced palladium atoms is apparent in sample Pd-Y with an average Pd-Pd coordination number of 2.2. The bond distance of 2.80 Å exceeds the value for bulk palladium by 0.05 Å, which is in contrast to the bond contraction usually found for small metal particles of ca. 10–20 Å diameter.<sup>31,32</sup> Lattice expansions can be found in bulk metal hydrides or borides.<sup>33,34</sup> We can exclude the formation of bulk

palladium hydride since clearly no bulk palladium is present (*no* outer shells, very small coordination numbers in the first shell). Furthermore, under the chosen experimental conditions (vacuum, elevated temperatures), a hydride phase would thermodynamically not be stable. As described above, the sodalite unit represents an ideal matrix for stabilizing palladium atoms in six-ring positions, giving rise to the formation of small ensembles with unusual bond distances.

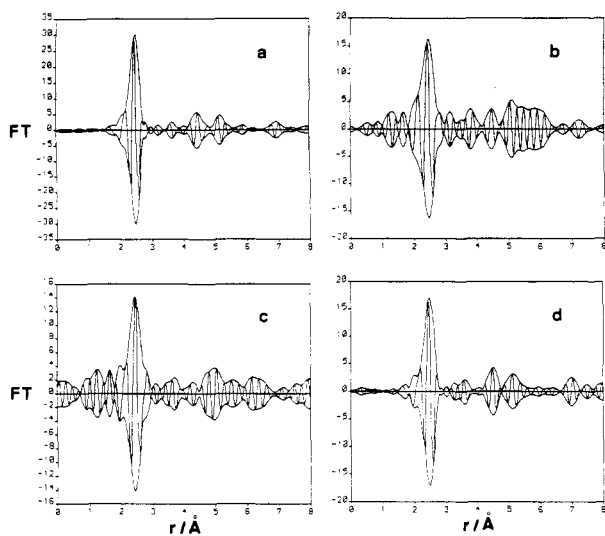
In Fe<sup>III</sup>Pd-Y (and PdFe-Y) the oxygen interface interactions are less pronounced. The absence of the "satellite peaks" in iron-exchanged samples is more apparent in Figure 2. Here, amplitude- and phase-corrected Fourier transformations (with palladium foil as reference) are shown for Pd-Y, Fe<sup>III</sup>Pd-Y and PdFe-Y. A shift of 5 eV was sufficient to position the imaginary part in the maximum of the magnitude. Totally symmetric peaks are obtained for the Pd-Pd atom pairs, while the influence of the zeolite interface is best visible in Pd-Y (Figure 2a). A back-transformation was applied on sample Fe<sup>III</sup>Pd-Y and PdFe-Y between 2.5 and 3.2 Å, dividing out the applied corrections. Excellent fits were thus obtained for the Pd-Pd interaction of which one example is shown for PdFe-Y in Figure 2d (note the almost complete match between fit and experimental data). Coordination numbers of 2.6 (Fe<sup>III</sup>Pd-Y) and 3.1 (PdFe-Y) are slightly increased with respect to 2.2 for Pd-Y, as are the bond distances (2.83 Å for Fe<sup>III</sup>Pd-Y and PdFe-Y, 2.80 Å for Pd-Y). These fit results emphasize that no additional phase besides Pd-Pd is present in the bimetal-exchanged zeolite samples when reduced at 423 K. It further suggests that, when bimetal-exchanged, on the average less Pd atoms are present that interact with the zeolite matrix. Small metal clusters do exist that are less densely packed than bulk palladium.

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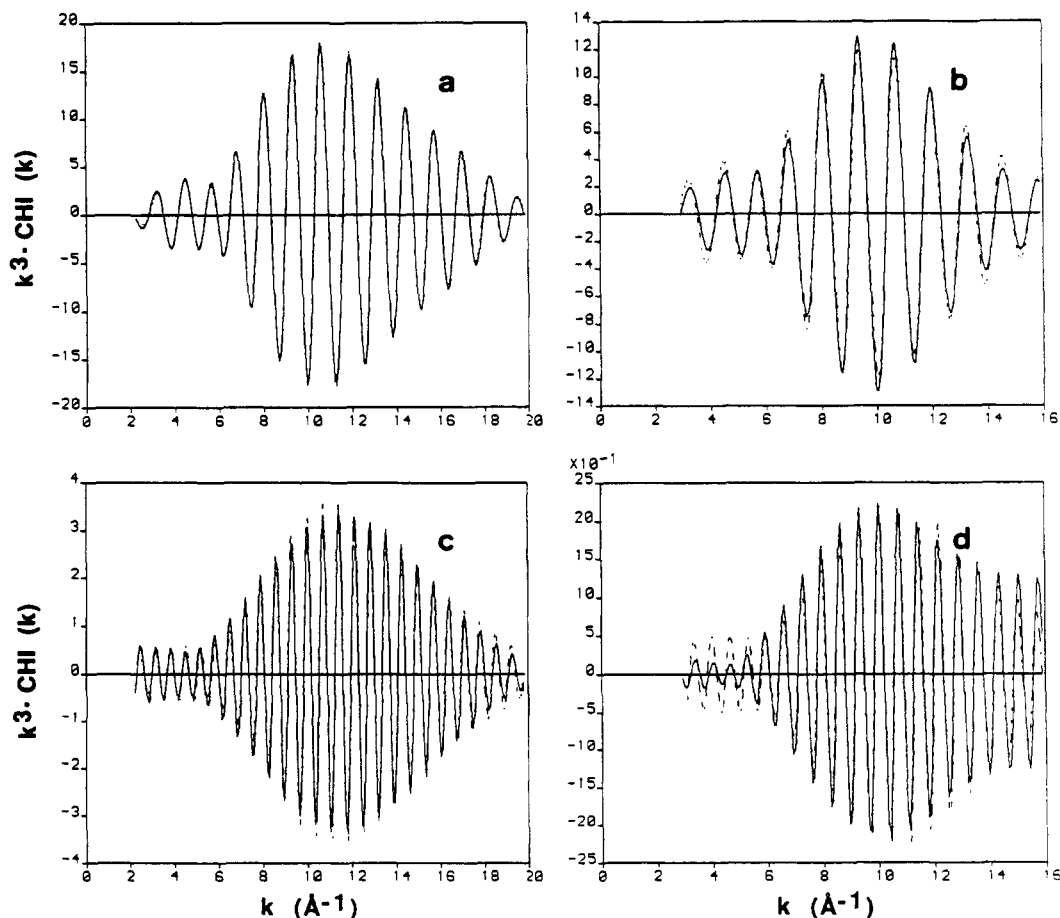


**Figure 3.**  $k^3$ -Weighted Fourier transform of the Pd-edge of Pd-Y (a),  $\text{Fe}^{III}\text{Pd-Y}$  (b), PdFe-Y (c) and CoPd-Y (d) reduced at 623 K.

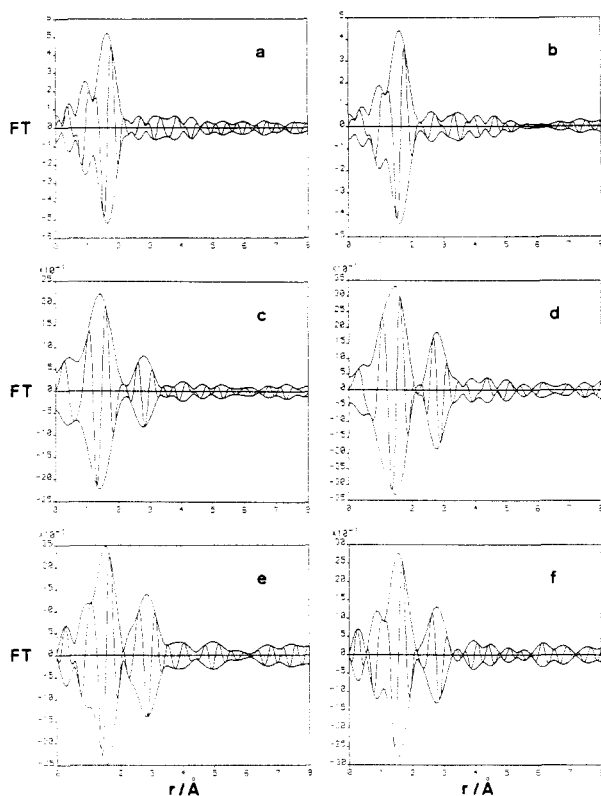
**Reduction at 623 K.** Hydrogen reduction of the Pd-zeolite samples at 623 K causes metal aggregation and formation of larger particles as demonstrated in Figure 3. The first Pd shell is strongly increased in Pd-Y (note scale) and outer-shell scattering up to the 7th coordination shell is now visible (Figure 3a). Fit results of the 1st, 3rd, 4th and 7th shells are listed in Table I. All bond distances are now identical with those of bulk palladium within the limits of accuracy. Both iron-containing samples (Figure 3b,  $\text{Fe}^{III}\text{Pd-Y}$  and 3c, PdFe-Y) apparently show a smaller first-shell scattering, but data fitting gives similar coordination numbers  $N$  of about 8 in all samples. The smaller peak height is caused by a higher static disorder in these samples as compared to Pd-Y.

Outer-shell features are also visible but are much less resolved. A fit of the third shell in  $\text{Fe}^{III}\text{Pd-Y}$  nonetheless shows again coordination numbers comparable to Pd-Y. The situation is changed in a sample coexchanged with Co(III) cations, which is shown in Figure 3d for comparison ( $\text{Na}_{16}\text{Co}_7\text{Pd}_{10}(\text{AlO}_2)_{57}(\text{SiO}_2)_{135}$ ). Small metal particles are also formed in this sample, but their dispersion seems to be higher based on the coordination numbers of the respective shells (see Table I and ref 19 for detailed description).

The higher static disorder in the reduced iron-containing samples could indicate the incorporation of iron in the palladium particles. This consideration was tested by fitting the first coordination shell with a second contribution. A standard for a possible Pd-Fe scatter pair is not available. It could be approximated with an Fe-Fe reference, which contains the (most important) correct backscatterer atom but differs for the absorber atom. Iron foil would furnish such an atom pair, but in the *bcc* crystal structure the two closest coordination spheres are only 0.4 Å apart and are therefore not easily separated. As a substitute, we chose cobalt powder that, as a neighbor in the periodic table, has phase and amplitude functions transferable to iron. Two-shell fits with Pd-Pd and Co-Co were performed on sample  $\text{Fe}^{III}\text{Pd-Y}$ , but a reasonable fit of higher quality than a single-shell fit with palladium was never achieved. Fits for the first and third shells in Pd-Y and  $\text{Fe}^{III}\text{Pd-Y}$  are compared in Figure 4, all obtained exclusively with palladium foil (first and third shell) as reference. It can be seen that the results are slightly better for Pd-Y (Figure 4a,c) than for  $\text{Fe}^{III}\text{Pd-Y}$  (Figure 4b,d), but the latter are still of high quality (nearly complete match between fit and experimental data). Furthermore, all bond distances in the bimetal-exchanged samples are now almost identical with values characteristic for bulk Pd, which would not be expected if iron ions or atoms were part of the Pd cluster. A sample of  $\text{Fe}^{III}\text{Pd-Y}$  reduced at 723 K under hydrogen showed no increase in agglomeration nor an indication for a Pd-Fe interaction. It must be concluded that an



**Figure 4.**  $k^3$ -Weighted isolated contribution of the 1st Pd shell (a,b; solid line) and 3rd Pd shell (c,d; solid line) of sample Pd-Y (a,c) and sample  $\text{Fe}^{III}\text{Pd-Y}$  (b,d) reduced at 623 K and fit (broken line, performed with Pd foil as reference).



**Figure 5.**  $k^3$ -Weighted Fourier transform of the Fe-edge of sample Fe-Y (a,c,e), sample PdFe-Y (b), and sample Fe<sup>III</sup>Pd-Y (d,f) after ion exchange (a,b,d), dehydration under oxygen atmosphere at 623 K (c), and hydrogen reduction at 623 K (e) and 423 K (f).

interaction between palladium and iron is not detectable in the bimetallic zeolites at the Pd edge.

**Iron-Edge EXAFS.** The above conclusion is best verified by examination of the same samples at the iron absorption edge. Fewer iron ions than palladium atoms are present in the samples and an interaction of these two constituents should therefore be more readily probed at the iron edge. Any Pd-Fe interaction should be detected at similar distances as observed at the palladium edge. In Figure 5, the  $k^3$ -weighted Fourier transformations (3.5–11 Å) of the bimetal-containing samples are shown and compared to monometallic, iron-exchanged zeolites treated similarly.

**Ion-Exchanged Fe-Y and PdFe-Y.** EXAFS data of sample Fe-Y and PdFe-Y, obtained after ion exchange, are shown in Figure 5a,b. Only one contribution due to oxygen coordination is visible. When fitted with Fe<sup>III</sup>acac, coordination numbers of 6.5 and a bond length of 2.11 Å (see Table II) suggest the presence of the hexaquo complex [Fe(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup> in the large cavities of the zeolite framework. This is also supported by the greenish color of the zeolite powder.

**Dehydration under Oxygen Atmosphere.** When sample Fe-Y is dehydrated in oxygen at 623 K (Figure 5c), the appearance of an outer shell indicates migration of the ions to cation positions in the zeolite framework, as observed for the Pd<sup>2+</sup> ions in Pd-containing zeolites at the Pd absorption edge. A back-transformation of this contribution, however, results in an amplitude function that differs from a pure Fe-Si/Al backscattering pair. Since no ideal reference for the analysis of this atom pair exists, an EXAFS reference file was prepared from MnS. The first-shell Mn-S coordination with a bond distance of 2.61 Å was used to calculate a scatterer pair Mn-S with a bond length of 3.4 Å, which is close to the Fe-Si distance. Fits with this model compound were only satisfying if an Fe-Fe atom pair from the second-coordination sphere of FeO was included. Best fits were obtained with two iron nearest-neighbors with a bond distance of 3.01 Å and 1.3 Si/Al atoms at 3.47 Å (see Table II). Thus, only a portion of the iron cations is isolated in cation positions, while the remainder forms very small iron oxide clusters (no additional outer

**TABLE II: Structural Data As Derived from EXAFS Analysis, Fe Absorption Edge**

sample	atom pair	$R/\text{Å}$	$N$	$\Delta\sigma^2/\text{Å}^2$	$\Delta E/\text{eV}$
295 K Hydrated					
Fe-Y	Fe-OZ	2.11	6.5	0.004	+0.5
PdFe-Y	Fe-OZ	2.11	6.4	0.007	+1.9
623 K Oxygen					
Fe-Y	Fe-OZ	1.92	3.1	0.007	+1.8
	Fe-Si/Al	3.47	1.3	0.000	-0.7
	Fe-Fe	3.01	2.0	0.000	-9.0
Fe <sup>III</sup> Pd-Y <sup>a</sup>	Fe-OZ	1.97	4.8	0.004	+2.3
	Fe-Si/Al	3.46	3.7	0.000	-2.7
	Fe-Fe	3.01	2.1	0.000	+9.9
PdFe-Y	Fe-OZ	1.95	2.2	0.004	-9.8
	Fe-OZ	2.11	1.5	0.000	-0.4
295 K Hydrogen					
Fe <sup>III</sup> Pd-Y	Fe-OZ	1.95	2.8	0.004	+4.5
PdFe-Y	Fe-OZ	2.06	3.2	0.006	-1.6
423 K Hydrogen					
Fe <sup>III</sup> Pd-Y	Fe-OZ	2.04	3.6	0.006	-0.8
	Fe-Si/Al	3.45	4.3	0.001	-1.2
PdFe-Y	Fe-OZ	2.05	4.2	0.005	-1.3
623 K Hydrogen					
Fe-Y	Fe-OZ	2.03	4.0	0.009	+0.0
	Fe-Si/Al	3.46	5.7	0.003	-3.6

<sup>a</sup>This sample was not dehydrated after the second ion exchange (Pd<sup>2+</sup>).

shells are visible). The first shell fit gave an Fe-O coordination number of 3.1 as compared to 6.5 of the hexaquo complex and showed a bond contraction from 2.11 to 1.92 Å. Bond distances of Fe-O = 2.16 and 2.36 Å were found with X-ray diffraction in Fe-Y zeolites dehydrated under vacuum at 673 K and exposed to 100 Torr of oxygen.<sup>35</sup> However, the observed bond lengths in our samples are comparable with those of hematite: Fe<sup>3+</sup>-O = 1.945 Å, Fe<sup>2+</sup>-O = 2.116 Å.<sup>36</sup> Thus, we suggest that the bond contraction is a strong indication for the oxidation of Fe<sup>2+</sup> to Fe<sup>3+</sup> (see also discussion of the corresponding XANES data).

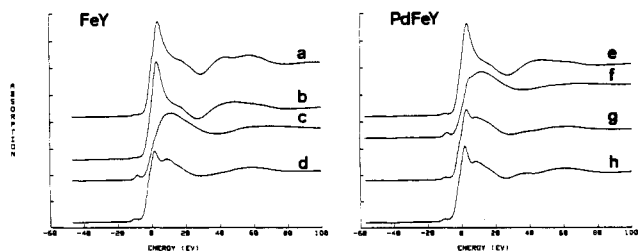
The spectrum of sample Fe<sup>III</sup>Pd-Y as shown in Figure 5d was taken after Fe(II) ion exchange, oxidative dehydration, and subsequent Pd exchange. Comparison with the Fe-Y analogue (Figure 5c) reveals a close resemblance to the latter sample. Here, a higher coordination number  $N = 4.8$  is obtained for the Fe-O shell. This sample was not yet dehydrated for a second time after palladium exchange; thus, the additional oxygens have to be interpreted as water coordinated to the anchored iron cations. This results in a slight increase of the bond distance to 1.97 Å (1.92 Å for Fe-Y). Probably some more iron ions reside in cation positions as indicated by a larger Fe-Si coordination number ( $N = 3.7$ ; Fe-Y,  $N = 1.3$ ).

The sequence of cation exchange has some influence on the homogeneity of the Fe phase: Sample PdFe-Y, first exchanged with Pd<sup>2+</sup>, dehydrated, and subsequently exchanged with Fe<sup>2+</sup> shows two different oxygen coordination distances after the second dehydration (see Table II).

**Reduction with Hydrogen.** The EXAFS data of sample Fe-Y after hydrogen reduction at 623 K (Figure 5e) reveal that intrazeolite Fe<sup>3+</sup> can be reduced to Fe<sup>2+</sup> under simultaneous re-dispersion of the small amount of FeO<sub>x</sub>. A longer Fe-O bond of 2.03 Å is obtained, and a fit of the outer shell with the Fe-Si calculated reference results in Fe-Si:  $N = 5.7$  and  $R = 3.46$  Å (a fit with an Fe-Fe shell was not possible). These values are consistent with a coordination of iron to the six-ring windows of the zeolite framework. A similar re-reduction is observed in samples Fe<sup>III</sup>Pd-Y and PdFe-Y when reduced at 423 K. Bond relaxation to 2.04 Å and the appearance of solely Fe-Si/Al in

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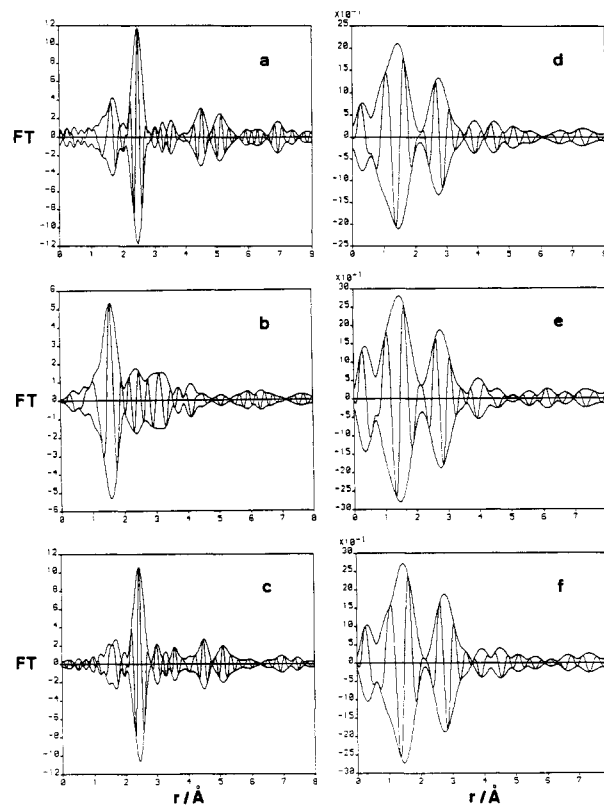
**Figure 6.** Preedge and XANES region of Fe-edges of sample Fe-Y: (a)  $\text{FeSO}_4$  precursor, (b) Fe-Y after ion exchange, (c) Fe-Y after dehydration under oxygen at 623 K, and (d) Fe-Y after hydrogen reduction at 623 K. Sample PdFe-Y: (e) after ion exchange with  $\text{FeSO}_4$ , (f) after dehydration under oxygen at 623 K, (g) after hydrogen reduction at room temperature, and (h) 423 K.

the outer shell takes place. Similar results are also obtained with sample PdFe-Y.

**Iron-Absorption Edges (XANES).** The oxidation and reduction processes of the intrazeolite iron ions under different conditions are clearly reflected in the signatures and positions of the respective iron absorption edges. In Figure 6 (left), the normalized edge spectra of the precursor compound  $\text{FeSO}_4$  (a) are compared with those of sample Fe-Y after ion exchange (b), dehydration at 623 K in oxygen (c), and hydrogen reduction at 623 K (d). Almost no change is visible between the precursor and aqueous Fe-Y (6b). Both samples show very weak  $1s \rightarrow 3d$  transitions and a distinctive "white line" as found for a number of divalent, 6-fold oxygen coordinated iron compounds.<sup>37</sup> The edge features of these samples confirm the above EXAFS results, indicating that neither the oxidation state nor the ligand sphere (symmetry) has changed upon diffusing the iron cation into the zeolite. However, oxygen treatment causes drastic changes in the form of an edge shift of about 2 eV to higher energies (c) and an increase of the preedge feature. A similar edge shift was reported for the oxidation of  $\text{Fe}^{2+}$  to  $\text{Fe}^{3+}$ , and it appears that the change in the preedge is related to a decrease in the Fe-O coordination number.<sup>37</sup> The edge shape itself is similar to that of  $\text{Fe}^{\text{III}}\text{acac}$ . Changes in the edge shape indicate a different ligand sphere after oxidation/dehydration as found in the corresponding EXAFS data (Fe-Si, Fe-Fe, and different oxygen coordination). In Figure 6d, the reduction to a lower oxidation state is apparent by the reverse shift of the absorption edge. Together with the EXAFS data of this sample, it can be interpreted with the coordination of most of the iron ions to the zeolite framework in the form of  $\text{Fe}^{2+}$ .

The absorption edges of sample PdFe-Y (Figure 6, right) show similar trends:  $\text{Fe}^{2+}$  is in an octahedral environment comparable to the precursor  $\text{FeSO}_4$  after ion exchange (e). Oxidation to  $\text{Fe}^{3+}$  occurs after oxygen exposure (f). From g and h, it is apparent that hydrogen treatment at room temperature (g) and 423 K (h) is already sufficient to reduce  $\text{Fe}^{3+}$  to  $\text{Fe}^{2+}$ .

**Potential Fe-Pd Interactions.** From the analysis of the Fe EXAFS and XANES data, it can be concluded that the intrazeolite iron ions are always at least partially coordinated to the zeolite framework. Only minor differences in the degree of iron oxide formation could be found between Fe-Y and the bi-metal-exchanged samples. An Fe-Pd interaction does not occur, since no peak in the Fe EXAFS spectra corresponding to any peak in the palladium EXAFS spectra is observed in the Fourier transforms of the bimetallic samples. If present at all, such interaction should result in a peak of a distance at about 2.6 Å, since this is the only overlap of peaks in the spectra of each absorption edge. Compare for example Figure 1f to Figure 5f for  $\text{Fe}^{\text{III}}\text{Pd-Y}$  after 423 K hydrogen reduction. Fits of the peaks at about 2.6 Å (uncorrected) on either absorption edge were perfect when only a single scatterer pair (Pd-Pd or Fe-Si) was used. Furthermore, the sample characteristics are very similar to their respective single-metal-exchanged counterparts (Pd-Y or Fe-Y). An EXAFS study about  $\text{Pd}_3\text{Fe}$  alloy formation between Fe and Pd was reported of a sample prepared by coimpregnation



**Figure 7.**  $k^3$ -Weighted Fourier transforms of sample FePd-Y of the Pd-edges (a,b,c) and Fe-edges (d,e,f) after ion exchange (a,d), after dehydration under oxygen at 623 K (b,e), and after hydrogen reduction at room temperature (c,f).

of the respective chlorides on an amorphous support and hydrogen reduction.<sup>38</sup> We can definitively rule out the presence of such a composite in the zeolite since the iron is always in an oxidized form.

**Reversed Sequence of Pd/Fe Ion Exchange.** It is of interest to compare the above results with those obtained when Fe/Pd bimetallic zeolites are prepared in the reversed sequence of ion exchange.  $\text{Fe}^{2+}$  ions were introduced first into the zeolite framework under anaerobic conditions and washed, and without any intermediate dehydration, palladium tetraammine chloride was added to the dispersion (sample FePd-Y). Immediately upon introduction of the palladium ions, a drastic color change of the zeolite dispersion from light green to black was observed. The sample was filtered and partitions were dehydrated similar to the previously discussed samples under oxygen atmosphere at 623 K and subsequently reduced under hydrogen atmosphere at room temperature.

**Pd-Edge EXAFS.** Figure 7 shows the  $k^3$ -weighted Fourier transforms of EXAFS data of these samples taken at the Pd ( $3-17 \text{ \AA}^{-1}$ ) and Fe ( $3.5-9.5 \text{ \AA}^{-1}$ ) absorption edges. Data of sample FePd-Y after ion exchange are depicted in Figure 7a. The formation of fairly large palladium metal particles is obvious, as Pd-Pd scattering can be identified up to the 7th shell. All Pd-Pd atom pairs have bond distances identical with those of bulk palladium and do not deviate even in the static disorder. This indicates that a large part of the  $\text{Pd}^{2+}$  ions are reduced to  $\text{Pd}^0$  by the presence of iron(II) ions. Corresponding electron micrographs show Pd particles with diameters greater than 50 Å clustered at the surface of the zeolite crystals. However, the reduction is incomplete indicated by the first-shell Pd-O contribution at about 1.8 Å with a coordination number of 2.5. This number is considerably smaller than  $N = 6$  for  $\text{Pd}^{2+}$  in solution and can be interpreted with the reduction of about 50% of the total palladium ions present. Fit results including the Pd-Pd outer shells are listed in Table III.

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**TABLE III: Structural Data for Sample FePd-Y As Derived from EXAFS Data**

atom pair	$R/\text{\AA}$	$N$	$\Delta\sigma^2/\text{\AA}^2$	$\Delta E/\text{eV}$
Pd-Edge				
295 K Hydrated				
Pd-O	1.99	2.5	0.000	-1.5
1. Pd-Pd	2.75	2.5	0.000	-2.6
3. Pd-Pd	4.76	5.0	0.001	-1.2
4. Pd-Pd	5.49	2.8	0.001	+0.0
623 K Oxygen				
Pd-O	1.98	3.3	0.000	+1.7
295 K Hydrogen				
Pd-O	1.99	1.5	0.000	-2.8
1. Pd-Pd	2.74	2.5	0.000	-2.9
3. Pd-Pd	4.77	4.0	0.000	-6.0
4. Pd-Pd	5.49	2.0	0.001	-4.3
Fe-Edge				
295 K Hydrated				
Fe-O	1.97	3.0	0.005	+1.4
Fe-Fe	3.06	4.7	0.002	-2.1
623 K Oxygen				
Fe-O	1.95	3.4	0.005	+0.5
Fe-Fe	3.05	7.2	0.002	-3.8
295 K Hydrogen				
Fe-O	1.98	3.4	0.004	+0.0
Fe-Fe	3.05	7.9	0.004	-3.7

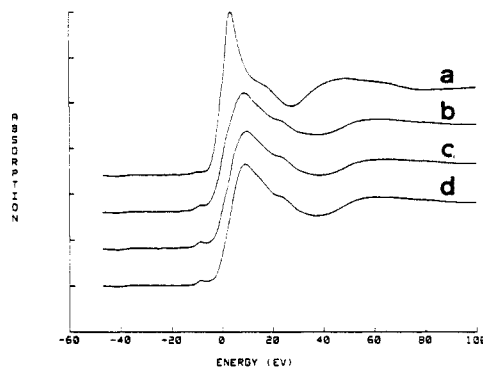
Figure 7b shows that heating of this sample up to 623 K in oxygen atmosphere causes nearly complete redispersion of the metal phase to Pd<sup>2+</sup> cations and/or Pd<sup>0</sup> atoms. Only about 1/10th of the previous Pd-Pd peak is still visible. Bergeret et al.<sup>22</sup> observed a similar redispersion with 15-Å Pd particles heated in oxygen. Subsequent hydrogen reduction at room temperature results again in agglomeration to larger particles, as shown in Figure 7c. The degree of reduction appears to be higher than after the initial reduction, since only 1.5 Pd-oxygen neighbors are found on average.

**Fe-Edge EXAFS.** The above observations demonstrate a redox reaction between intrazeolite Fe<sup>2+</sup> and Pd<sup>2+</sup>. Analysis of the hydrated sample FePd-Y after ion exchange at the iron absorption edge shows a short bond length of  $R = 1.97 \text{ \AA}$  for Fe-O (Figure 7d). This is considerably shorter than that of Fe-Y (2.11 Å) and is usually only found after dehydrating and oxidizing the iron to Fe<sup>3+</sup>. Furthermore, the coordination number  $N = 3$  is very different compared to  $N = 6$  of the hexaquo complex. Fitting of the outer shell was obtained with a single-shell contribution of Fe-Fe with a bond length identical with that of FeO ( $R = 3.06 \text{ \AA}$ ). All these findings point to a redox reaction between palladium and iron and the *simultaneous precipitation of iron oxide*. The iron oxide is stable against subsequent treatments: when the sample is heated in oxygen, only a slight increase of the first and second shell is observed (Figure 7e), which are almost unchanged after hydrogen exposure at room temperature (Figure 7f; see Table III).

The oxidation to Fe<sup>3+</sup> is also visible in the absorption edge spectra (Figure 8). It can be seen that the absorption edge of Fe-Y (a) is shifted to higher energies as soon as palladium is introduced to the aqueous dispersion to give sample FePd-Y (b). The new edge shape is comparable to that of Fe-Y after oxidation at 623 K (Figure 6c) where a substantial contribution of iron oxide particles was found. The edge features do not change when treating FePd-Y with oxygen or hydrogen (c,d). The chemistry of iron and palladium in aqueous intrazeolite "solutions" therefore favors a redox reaction under formation of an iron oxide phase and palladium particles.

### Conclusion

This study addresses the question of whether a less reducible transition metal has an influence on the reduction and agglomeration behavior of a noble intrazeolite metal, i.e., palladium. A



**Figure 8.** (a) Absorption-edge features of sample Fe-Y after ion exchange, (b) absorption edge of sample FePd-Y after ion exchange, (c) FePd-Y after dehydration under oxygen at 623 K, and (d) FePd-Y after hydrogen reduction at room temperature.

chemical anchoring mechanism was proposed by Sachtler et al. to explain for their observation that the addition of iron to Pt-exchanged zeolites resulted in an enhanced dispersion as concluded from X-ray diffraction and adsorption measurements.<sup>15</sup> This effect was only detected at an Fe/Pt ratio greater than 0.5 and at high reduction temperatures of 923 K under hydrogen. Since X-ray powder diffraction is not very sensitive for changes in particle size smaller than 50 Å or any M-M' interaction, other techniques such as EXAFS are expected to give more insight as to the structure and confinement of metal particles. A related EXAFS study of the Rh-edge of Rh/CrNa-Y zeolite showed a decrease in the Rh-Rh peak under high-temperature reduction and was explained by the presence of an additional Rh-Cr contribution.<sup>16</sup> However, a subsequent Pt-edge EXAFS study of Ca, La, or Fe coexchanged Pt-Y zeolites by Sachtler et al. did not confirm interactions between Fe<sup>2+</sup> ions and the reduced Pt phase.<sup>39</sup>

In a recent EXAFS study on Co/Pd-exchanged zeolites where spectra were taken at both absorption edges,<sup>19</sup> we could show that the addition of cobalt *increased* the reducibility of the palladium phase. Similar trends were reported when Ni, Cr, or Mn ions were added to Pt in zeolite Y.<sup>40</sup> In the Co/Pd system, palladium particles formed at room temperature were larger than in the parent Pd-Y zeolite (EXAFS results are included in Table I). However, reduction at higher temperatures resulted in *higher* dispersion as compared to Pd-Y under similar conditions. This effect could not be associated with a chemical anchoring process, since the cobalt absorption edge spectra did not differ from those of the Co-Y sample. It was concluded that the reduced particle size is associated with a change in the cation distribution: Co cations are preferentially coordinated in the sodalite cages and force the palladium cations into the supercages. The latter are thus more accessible to the reducing atmosphere and form Pd clusters of about supercage dimensions at low temperatures. The resulting clusters are locked into the zeolite cages and are *less mobile* at higher reduction temperatures; thus, they do not agglomerate further (compare samples Pd-Y and CoPd-Y at 623 K hydrogen reduction in Table I).

The present study shows that the situation is different in the Fe/Pd-Y systems. The enhanced reducibility of palladium upon addition of a second metal is still noticeable at room temperature but to a much smaller extent (about 55–60% of all Pd ions are reduced compared to 35% in Pd-Y). High-temperature reduction shows only a slightly higher dispersion when iron is present, but the effect is much less drastic compared to the cobalt/Palladium system. Thus, iron does not influence the Pd particle growth at the microscopic scale, and no chemical interaction between the two intrazeolite metals is detected by analysis of both the Pd and Fe X-ray absorption edge.

A possible explanation for the difference between iron and cobalt systems may be based on the formation of iron oxide clusters

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in the iron-containing samples. In all samples, some oxide formation was detected by the appearance of an Fe-Fe shell at 3.01 Å. The particle size of these clusters is estimated to be at least 6 Å; thus, it is very likely that they are located in the supercages rather than sodalite units. This could eliminate the site-directing effect of the second transition metal on the position of the Pd phases, and the effect on the reducibility is expected to be minimal.

This analysis shows that the interplay of metals in bimetallic zeolite catalyst is not easily predictable and is strongly dependent on the chemistry of each constituent. EXAFS is a valuable technique for structure determination of these complex systems by probing the ligand spheres of each component separately. On

the basis of subtle changes in zeolite oxygen coordination of the metal, it is demonstrated that EXAFS is also very sensitive for the analysis of intrazeolite reduction degrees.

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**Registry No.** Pd, 7440-05-3; Fe, 7439-89-6; Co, 7440-48-4.

## Electrophoretic Light Scattering of Surfactant Micelle Colloids

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Aqueous micellar solutions were investigated by the electrophoretic light scattering method. The electrophoretic mobilities of  $(2-4) \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$  were obtained for charged micelles of tetradecyl- and hexadecyltrimethylammonium salicylates ( $\text{C}_{14}\text{TASal}$ ,  $\text{C}_{16}\text{TASal}$ ) and partly protonated tetradecyl- and oleyldimethylamine oxides ( $\text{C}_{14}\text{DAO}$  and  $\text{ODAO}$ ).

### Introduction

Electrophoretic light scattering is a beneficial method in which the electrophoretic mobility and the mutual diffusion coefficient can be measured simultaneously. Since the initial development of theory and experimental procedures by Ware and Flygare,<sup>1</sup> the technique has been applied to solutions of biomolecules, biocolloids, and polyelectrolytes and to suspensions of polymer latex particles and mineral colloidal particles.<sup>1-6</sup> However, applications to surfactant micelle colloidal systems have not been achieved as yet.

In the present paper, electrophoretic light scattering measurements were carried out for aqueous micellar solutions of  $\text{C}_{14}\text{TASal}$ ,  $\text{C}_{16}\text{TASal}$  and partly protonated  $\text{C}_{14}\text{DAO}$  and  $\text{ODAO}$ . The electrophoretic mobilities of charged micelles are obtained together with their mutual diffusion coefficients.

### Experimental Section

Samples of  $\text{C}_{14}\text{TASal}$  and  $\text{C}_{16}\text{TASal}$  were prepared by reacting the corresponding alkyltrimethylammonium halides with sodium salicylate in water. The preparation will be described in detail in a subsequent paper.<sup>7</sup> Samples of  $\text{C}_{14}\text{DAO}$  and  $\text{ODAO}$  are the same as those previously prepared and used.<sup>8,9</sup>  $\text{C}_{14}\text{DAO}$  and

$\text{ODAO}$  are weak bases and are partly protonated in the presence of HCl.

The electrophoretic mobility and the mutual diffusion coefficient were measured at 25 °C on Otsuka Electronics electrophoretic light scattering spectrophotometer, ELS-800. He-Ne laser light at 632.8 nm was used. The electrophoretic rectangular cell of  $2 \times 10 \times 17$  mm is separated by cellulose semipermeable membranes from the electrode chambers, which hold the Pt electrodes coated with Pt-black.

The electric field, which is perpendicular to the incident radiation, is applied in 1-s pulses that are reversed on each successive application. The sign of the electrophoretic mobility was determined by modulating the frequency of the reference light. The autocorrelation function of the dynamic light scattering was analyzed by a fast Fourier transform routine.

### Results

When the electric field,  $E$ , is applied to aqueous solutions of charged particles, the particles are moved by an electrophoretic drift velocity of  $v$ . Then, the electrophoretic mobility,  $U$ , is described as the velocity per unit electric field, that is

$$U = v/E \quad (1)$$

On the basis of the photon correlation method by the heterodyne mixing mode, the first-order autocorrelation power spectrum,  $F_1(\mu, \omega)$ , of the light scattered from the moving particles in solutions under the external electric field applied perpendicularly to the incident radiation is written by

$$F_1(\mu, \omega) = \frac{\langle N \rangle}{\pi} \frac{\mu^2 D}{(\omega + \mu \cdot v)^2 + (\mu^2 D)^2} \quad (2)$$

and

$$\mu \cdot v = \mu v \cos(\theta/2) \quad (3)$$

where  $\mu$  and  $\mu$  are the scattering vector and its magnitude, respectively;  $\omega$  is the angular frequency;  $\langle N \rangle$  is the average number of scatterers;  $D$  is the mutual diffusion coefficient; and  $\theta$  is a scattering angle. The power spectrum exhibits a Lorentzian peak

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