Structure and Catalytic Properties of Pt-Modified Hyper-Cross-Linked Polystyrene Exhibiting Hierarchical Porosity

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The structural transformation and catalytic properties of metal/polymer nanocomposites derived from hyper-cross-linked polystyrene (HPS) exhibiting both microporosity and macroporosity, and filled with Pt nanoparticles, are investigated in the direct oxidation of L-sorbos to 2-keto-L-gulonic acid. Transmission electron microscopy, X-ray photoelectron spectroscopy, X-ray diffraction, anomalous small-angle X-ray scattering, and catalytic studies suggest that the catalytically active species, nanoparticles of mixed composition with a mean diameter of 1.6 nm, develop after the initial induction period. At the highest selectivity (96.8%) at 100% L-sorbos conversion, the catalytic activity is measured to be $2.5 \times 10^{-3}$ mol/mol Pt-s, which corresponds to a 4.6-fold increase in activity relative to the Pt-modified microporous HPS previously reported. This substantial increase in catalytic activity is attributed to the presence of macropores, which facilitate mass transport and, consequently, accessibility of the nanoparticle surface for reactants.

Introduction

Metalated polymer nanocomposites continue to receive considerable attention since their bulk properties can be greatly, and desirably, altered relative to those of pure polymers.1–4 Nanoparticles incorporated into a polymer system may impart magnetic, semiconducting, or catalytic properties, depending on the nanoparticle species and its characteristics. In recent years, catalysis employing metal nanoparticles has become a subject of intense interest due, in major part, to the enhanced activity and selectivity of nanostructured catalysts.5–7 When catalytic nanoparticles are formed on the surface of an inorganic or carbon substrate, however, the nanoparticle size and morphology cannot be precisely regulated.8,9 One reliable means by which to overcome this challenge is to grow the nanoparticles in nanostructured polymers exhibiting well-defined buried interfaces.10–12 Such interfaces can be conveniently generated via (i) microphase separation in solution or bulk13–23 or (ii) formation of nanopores, or nanocavities, within a polymer matrix.24 Nanopores can be designed into polymers by frustrating the chain packing of complex macromolecules.25 Conversely, nanoporosity can be induced in polymers by special treatment, such as exposure to a supercritical fluid under a select set of conditions.26 When nanoporous polymers are used as matrices in which to grow nanoparticles, the size of the nanoparticles may be physically restricted by the pore size of the polymer.

This approach is widespread in the production of mesoporous solids wherein metal compounds are incorporated inside a porous inorganic medium so that nanoparticles can subsequently be grown within the pores.27,28 This general strategy is not typically employed for use with polymeric media, since most polymers are dense and do not consist of regular interpenetrating cavities.

The development of nanoporous polymers has been greatly stimulated by the need for materials with a low dielectric constant for next-generation microelectronics.29 Production of such materials with closed nanopores can be achieved, for instance, by templating over block copolymers possessing a body- or face-centered cubic spherical morphology and a thermally labile block. If the pores are openly connected through the use of other morphologies, the resultant polymer membrane can be surface-decorated by metal deposition, resulting in metal–polymer nanocomposites.24,30 Alternatively, delocalized solvent crazing can yield nanoporous polymers possessing interpenetrating pores that can serve as nanoreactors in the formation and stabilization of highly dispersed metal nanoparticles. Amorphous porous poly(ethylene terephthalate) prepared in this fashion has been successfully used as a matrix in which to grow metallic Ni nanoparticles via reduction of nickel perchlorate by sodium borohydride.31 At the reaction conditions examined, pore dimensions influence the state of the final product. Subtle control over nanoparticle size and size distribution, which are vital to ultimate property development, cannot be realized, however, with the ill-defined pores formed in a crazed polymer matrix. Nanoporous polymer microspheres containing continuous channels lined with poly(acrylic acid) have been reported by Lu et al.32 These microspheres are prepared by UV-cross-linking of poly(tert-butyl acrylate-b-2-
Hyper-cross-linked polystyrene (HPS) is another nanoporous polymeric medium that can be considered for regulated nanoparticle growth. In this polymer network with a formal cross-link density exceeding 100%, rigid nanoscale cavities measuring about 2 nm form spontaneously during synthesis. This material is obtained by incorporating methylene groups between adjacent phenyl rings in dissolved PS homopolymer or gelled poly(styrene-divinylbenzene) copolymer in the presence of ethylene dichloride, which is a good solvent for the polymer. A unique feature of HPS is its ability to swell in a wide variety of different solvents, even thermodynamically poor ones. This characteristic facilitates incorporation of various organometallic compounds into the nanostructured HPS matrix and, thus, nanoparticle formation. One of our earlier studies of metalated HPS describes the formation of Co nanoparticles in HPS. The regulated growth of Co nanoparticles measuring ~2 nm in diameter over a wide range of Co concentrations is attributed to the existence of nanoscale HPS cavities, which serve to physically restrict the size of the growing nanoparticles. More recently, we have examined the formation of Pt nanoparticles in HPS. Impregnation of HPS with a tetrahydrofuran (THF) solution containing platinic acid (H₂PtCl₆·6H₂O) results in the formation of Pt(II) complexes within the nanocavities of HPS. Subsequent reduction of the complexes by H₂ yields stable, relatively monodisperse Pt nanoparticles with a mean diameter of 1.3 nm (since the size of the Pt precursor, 2 nm, is restricted by the size of the HPS nanoparticles). This robust metal/polymer nanocomposite exhibits exceptional properties in the catalytic oxidation of L-sorbos to 2-keto-L-gulonic acid, a vitamin C precursor (see Scheme 1), without the use of any chemical modifiers normally added to attain high reaction selectivity. In general, the catalytic activity and selectivity both depend on nanoparticle size and composition.

The principal shortcoming of this system is that the microporous nature of HPS (when pores are below ~2.0–2.2 nm) likewise limits the mass transport of reactant within the metalated HPS matrix and, hence, catalytic activity. To alleviate this disadvantage, we have elected to examine Pt metalation and subsequent L-sorbose oxidation within a commercially available variant of HPS that combines both micropores and macropores (larger than 20 nm in diameter). While this strategy should certainly improve mass transport during reaction, we recognize the possibility that the presence of macropores might severely jeopardize control over nanoparticle size. In this work, we report on the structure and catalytic properties of a micro/macroporous HPS nanocomposite containing Pt nanoparticles in the direct oxidation of L-sorbose. The structural transformation of this hybrid material is followed at different reaction stages by X-ray photoelectron spectroscopy (XPS), transmission electron microscopy (TEM), X-ray diffraction (XRD), and anomalous small-angle X-ray scattering (ASAXS).

**Experimental Section**

**Materials.** The HPS was generously provided by Purosolve Int. (U.K.), as Macronet MN 270/386-101/100 (hereafter designated as MN-270), and was used without further purification. According to N₂ sorption measurements, the BET (Brunauer–Emmett–Teller) surface area of the unmodified HPS was 738 m²/g and its total pore volume was 0.59 cm³/g. About 46% of the pore population measured in the 20–100 nm range, while pores <6 nm accounted for ~13%. The mean diameter of the micropores was reported by the manufacturer to be 1.5 nm. Both H₂PtCl₆·6H₂O and sodium hydrogen carbonate (NaHCO₃) were obtained from Reakhim (Moscow, Russia). Reagent-grade THF was purchased from Aldrich, whereas the L-sorbose was provided by Fluka, and both were used as received. Water was purified with a Milli-Q water purification system.

**Synthesis.** Compared to the method described elsewhere, the procedure to grow Pt nanoparticles within HPS was slightly modified to be more commercially attractive. In a typical synthesis, 0.24 g H₂PtCl₆·6H₂O was dissolved in 6 mL of THF under air and added to 3 g of MN-270. The suspension was continuously stirred for 10 min so that the solution could be absorbed by the polymer and then dried at 70–75 °C for 24 h. The Pt content was 1.72 wt % by elemental analysis.

**Catalysis.** The oxidation reaction was conducted batchwise in a specially constructed apparatus that permits independent control over parameters such as the L-sorbose and NaHCO₃ concentrations, catalyst concentration, temperature, (pure) oxygen feed rate, and stirring rate. A suspension of catalyst and L-sorbose (100 mL) prepared at a predetermined concentration was placed in the temperature-controlled apparatus equipped with a magnetic stir bar and reflux condenser. The rate of oxygen feed was controlled by a rotameter. An equimolar quantity of alkalizing agent (NaHCO₃) was fed to the apparatus as a single injection or continuously over 180 min (to maintain a constant pH of 7.7) using an automatic dispenser. The high stirring rates employed here ensured good mixing without diffusion limitation. Samples of the reaction mixture were periodically removed for analysis. At the end of each experiment, the catalyst was separated by filtration, and the filtrate was analyzed for the presence of L-sorbose and 2-keto-L-gulonic acid. The amount of residual L-sorbose was discerned by gas chromatography (GC) through the use of a Chrom-5 chromatograph, operated isothermally with a flame-ionization detector and glass column filled with 5% SE-30 on Chromaton N-AW. Analysis of the sodium salt of 2-keto-L-gulonic acid was performed on a MILLICHROM-5 HPLC chromatograph. Normal-phase chromatography using a 10 cm tungsten column characterized by a 4000 theoretical plate number was chosen for sample analysis.

**Characterization.** The Pt content was obtained by X-ray fluorescence (XRF) measurements performed with a Zeiss Jena VRA-30 spectrometer (Mo anode, LiF crystal analyzer, and SZ.
detector). Analyses were based on the Co Kα line and a series of PS/Pt standards prepared by mixing 1 g of PS with 10–20 mg of standard Pt compounds. The time of data acquisition was constant at 10 s. Nitrogen sorption was conducted at the normal boiling point of liquid nitrogen using a Micromeritics Gemini instrument. Samples were degassed at 100°C in vacuo for 12 h prior to analysis. X-ray photoelectron spectra were obtained using Mg Kα (hv = 1253.6 eV) monochromatized radiation with a modified ES-2403 spectrometer (provided by the Institute for Analytic Instrumentation of the Russian Academy of Sciences, St. Petersburg, Russia). The analyzer was operated at a pass electron energy of 100 eV. All data were acquired at an X-ray power of 100 W and an energy step of 0.1 eV. The electron-flood gun accessory was used so that the current of the total emitted electron flux from the flood-gun filament was adjustable from 0 to 100 mA, with the optimum found to be 70 mA at an electron energy of 2 eV. Samples were allowed to outgas for 15–30 min before analysis and were sufficiently stable during examination. Data analysis was performed using a standard RFES-set with Resolver program. Identification of band components was achieved by assuming that the Pt 3d5/2 and 3d3/2 spin–orbit components (i) exhibit a Gaussian peak shape, (ii) are in 3:2 peak area ratio, and (iii) are separated by ca. 5 eV. Surface compositions were discerned by measuring the peak-area intensities of different Pt species.

X-ray diffraction patterns were collected on a Scintag θ–θ powder diffractometer. X-rays were generated with Cu Kα radiation with a wavelength of 0.154 nm. The step size of each run was maintained at 0.02°. Transmission electron microscopy was performed with a Zeiss EM902 electron spectroscopic microscope operated at 80 kV and an energy loss of 0 eV. Insoluble HPS-Pt powders were embedded in epoxy resin and subsequently microtomed at ambient temperature. Images of the resulting thin sections (ca. 50 nm thick) were collected on plate negatives, digitized at 600 dpi, and analyzed with the Adobe Photoshop software package and the Scion Image Processing Toolkit. The ASAXS analysis was performed at the JUSIFA beamline35 at HASYLAB, DESY (Hamburg, Germany), over an energy range in close proximity to the LIII-absorption edge of Pt (11.564 keV). Measurements were obtained with a two-dimensional detector at two energies (11.484 and 11.562 keV). The energy dependence of small-angle scattering near the LIII-absorption edge of Pt was used to differentiate the scattering due specifically to Pt-containing heterogeneities from that associated with, for instance, pores in the polymer matrix. Details pertaining to data analysis are provided in the following section.

Results and Discussion

Structure Analysis. As discussed elsewhere,36 incorporation of the THF solution containing platinic acid into microporous HPS results in partial reduction of the Pt(IV) species and the formation of Pt(II) complexes wherein ligands are the product of THF oxidation. Figure 1 presents XPS spectra (and their deconvoluted contributions) of Pt-containing MN-270 at different stages of the catalytic reaction. The fraction of each type of Pt species ascertained from the deconvolution analysis is listed in Table 1. In the present work, addition of the THF/platinic acid solution to MN-270 yields the unchanged Pt(IV) (25.8%), as well as Pt(II) (48.3%) and even Pt(0) (25.9%). This analysis suggests that the chosen reaction conditions facilitate

![Figure 1](http://example.com/figure1.png)

**Figure 1.** X-ray photoelectron spectra (green lines) of the following HPS nanocomposites: (a) MN-270-Pt, (b) MN-270-Pt-induction, (c) MN-270-Pt-OX, and (d) MN-270-Pt-3OX. The Pt 5p3/2 binding energy at 69 eV and satellites from MgKα and MgKα4 at 8.4 and 10 eV are shown as gray lines. The blue, violet, and magenta peaks denote the deconvolutions of Pt(0), Pt(II), and Pt(IV) in K2PtCl6. The red lines represent the calculated spectra as the sum of all the deconvoluted peaks.

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<th>material notation</th>
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<th>BET surface area (m²/g)</th>
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* The binding energy.
Pt reduction since no Pt(0) was previously detected \[^{36}\] in the microporous HPS after incorporation of platinic acid. Parts a and b of Figure 2 are TEM images of the MN-270-Pt hybrid after initial incorporation of platinic acid. The material is observed to contain electron-dense nanoparticles with a diameter exceeding 0.7 nm (the estimated resolution of the microscope). The mean particle diameter calculated from a particle size histogram (Figure 2c) is 1.6 nm, and the standard deviation is 0.6 nm. Since HPS constitutes a highly hydrophobic matrix, the Pt species deposited from the platinic acid do not distribute molecularly within the HPS matrix but instead form well-defined clusters. The light background regions in this and subsequent TEM images represent the macropores alluded to earlier (the micropores are not resolved). The corresponding XRD pattern of this material is displayed in Figure 3a and confirms the absence of scattering from crystalline Pt. It does not, however, preclude the possibility of scattering from amorphous Pt. Last, it should be noted that the BET surface area of this Pt-modified HPS (see Table 1) is similar to that of the parent MN-270 homopolymer (provided in the Experimental Section), indicating that the incorporation of Pt species into the HPS matrix does not obstruct the micropores.

When this metal–polymer nanocomposite is used as a catalyst in the oxidation of L-sorbose to 2-keto-L-gulonic acid according to Scheme 1, L-sorbose is slowly consumed for the first 60 min of the reaction but the product (2-keto-L-gulonic acid) is not detected. This stage is referred to as the induction period, and the Pt-modified MN-270 isolated after this period is denoted MN-270-Pt-induction. According to the XPS spectra presented in Figure 1b and the deconvolution results recorded in Table 1, this material contains a higher fraction of Pt(0) and a lower fraction of Pt(II) and Pt(IV). The XRD pattern of this material (cf. Figure 3b) shows peaks assigned only to the Pt(0) crystalline phase. \[^{39,40}\] The average Pt(0) crystallite size (calculated from the Scherrer equation for 2\(\theta = 39.8^\circ\) along the (111) projection) is 2.8 nm. No peaks signifying the existence of PtO or PtO\(_2\) crystals are evident in Figure 3b, in which case the presence of Pt at higher oxidative states, such as Pt(II) and Pt(IV), in the XPS spectra of this sample are attributed to either (i) incomplete reduction by L-sorbose or (ii) the modification of nanoparticle surface atoms by 2-keto-L-gulonic acid. It should be recognized, however, that high binding energy values may also reflect charging of metal nanoparticles in a nonconductive polymer environment. \[^{42,43}\] In this case, the species comprising the nanoparticles would be expected to exhibit an increased oxidation state during analysis. Moreover, according to the results listed in Table 1, the BET surface area of this sample is substantially lower (by more than a factor of 2) than that of the MN-270-Pt material after incorporation of platinic acid, strongly suggesting that the micropores in the HPS matrix are impeded by the presence of reaction species (L-sorbose and 2-keto-L-gulonic acid).

The TEM images and corresponding particle size histogram presented in Figure 4 reveal that the Pt nanoparticle size and

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**Figure 2.** TEM images (a, b) and corresponding particle size histogram (c) of MN-270-Pt containing 1.72 wt % Pt.

**Figure 3.** XRD patterns of the following HPS nanocomposites: (a) MN-270-Pt, (b) MN-270-Pt-induction, (c) MN-270-Pt-OX and (d) MN-270-Pt-3OX.
size distribution in the MN-270-Pt-induction material are surprisingly comparable to those in the initial MN-270-Pt material. Despite the formation of slightly larger nanoparticles (measuring up to ca. 4 nm in diameter) following the induction period, such similarity strongly suggests that the mixed Pt(0/II/IV) clusters serve as nuclei in the formation of Pt(0) nanoparticles and that the nanoparticle size is, for the most part, controlled by the size of the HPS micropores. The larger mean particle size deduced from the XRD pattern in Figure 3b implies that the small Pt nanoparticles in Figure 4a,b may be amorphous. Recall that this possibility could not be discounted from the pattern acquired from the MN-270-Pt material. Reduction of Pt in MN-270-Pt by L-sorbose during the induction period of the reaction seems to stabilize the nanoparticle surface. The HPS macropores, on the other hand, provide easy access of the reaction medium to the growing nanoparticles, thereby facilitating nanoparticle nucleation. In addition, oxidation products may likewise adsorb onto and help stabilize the surface of the growing Pt nanoparticles. Together, these molecular-scale events contribute to the apparent tight control over nanoparticle size. In contrast, access of L-sorbose to resident Pt species and the subsequent availability of 2-keto-L-gulonic acid (a possible nanoparticle stabilizer) are both hindered in fully microporous HPS by the small pore size. Diffusion-limiting considerations such as these are deemed responsible for the larger and more broadly distributed Pt nanoparticles observed\(^3\) after the induction period in HPS possessing only micropores.

Figures 5 and 6 are TEM images and corresponding particle size histograms of HPS/Pt nanocomposite catalysts isolated after one catalytic run (MN-270-Pt-OX, Figure 5) and three cycles (MN-270-Pt-3OX, Figure 6). With the exception that a relatively small population of nanoparticles appears to aggregate into nanoparticles measuring greater than \(\sim 3.5\) nm in diameter, the mean particle diameter calculated from both histograms remains unchanged. Thus, it follows that the catalytic oxidation examined here does not destabilize the Pt nanoparticles within the HPS nanocomposite. It is particularly noteworthy that, after three consecutive catalytic runs, no further nanoparticle aggregation is observed. In fact, the relatively large nanoparticles (>4 nm
in diameter) introduced after the first cycle are no longer detected by TEM after the third cycle. Although we recognize that this population of nanoparticles is small and may easily be overlooked during sampling, we cannot discount the possibility that these nanoparticles actually disperse in the presence of the reaction media so that the mean particle diameter decreases slightly. Results acquired from XRD are consistent with this scenario. The XRD patterns of MN-270-Pt-OX and MN-270-Pt-3OX (cf. parts c and d, respectively, of Figure 3) exhibit reflexes identifying Pt(0) crystallites with mean diameters of 3.2 and 2.8 nm, respectively. The XPS spectra collected from these nanocomposites after cycling (Figure 1c for MN-270-Pt-OX and Figure 1d for MN-270-Pt-3OX), as well as the deconvolution results (see Table 1), are virtually identical to those of the MN-270-Pt-induction material (Figure 1b), suggesting that the Pt nanoparticle composition does not change after the catalytic oxidation of L-sorbose. One property that does change upon catalytic oxidation is the BET surface area, which increases sharply from 325 m$^2$/g after induction to 600 m$^2$/g after just one catalytic run (included in Table 1). These results suggest that the HPS micropores become less obstructed relative to those following the induction period when the reaction is complete.

The enhanced stability of the Pt nanoparticles within MN-270 is indeed surprising. In the presence of macropores, it is reasonable to expect greater aggregation of the nanoparticles during catalytic cycling. Such stability can be explained in terms of the following mechanism. The Pt clusters formed in MN-270 after incorporation of platinic acid preferentially locate inside (without obstructing) the micropores, not at the macropore interfaces. If they did reside at the macropore interfaces, we suspect that the nanoparticle stability might be sorely compromised after consecutive exposure to the reaction medium. Such preferential positioning of the Pt clusters would most likely occur due to their hydrophobic shell that develops because of the presence of organic ligands in the Pt(II) complex. Subsequent Pt nanoparticle nucleation and growth occur exclusively in the micropores, which effectively serve to restrict nanoparticle size. The small fraction of Pt nanoparticles with diameters larger (2.5–4.0 nm) than the average nanopore diameter (~1.5 nm) may result from either (i) clusters or nanoparticles that reside in pore defects or junctions within the native HPS matrix or (ii) neighboring nanoparticles that aggregate as the HPS matrix swells in the reaction medium. It is possible that such aggregation may occur only upon initial exposure to the reaction medium. While it would be intuitive to expect that the extent of such aggregation would increase upon repeated exposure, the TEM and XRD results reported here indicate that consecutive exposure of the Pt nanoparticles to the reaction medium may actually serve to reduce the mean nanoparticle size.

Figure 7. Small-angle X-ray scattering curves of the following HPS nanocomposites: (a) MN-270-Pt, (b) MN-270-Pt induction, (c) MN-270-Pt-OX and (d) MN-270-Pt-3OX. The total scattering curves (Δ) are measured at 11.484 keV, while the separated scattering curves (○) are obtained from measurements acquired at 11.484 and 11.562 keV. The solid lines are regressed fits to the model provided in eq 1. The inset in (d) shows the correlation function of the model proposed by Debye et al.41 (see text).
curves is a regression of a model introduced by Debye et al. with an additional $q^{-4}$ term describing large inhomogeneities (eq 1):

$$\Delta \sigma_{\text{tot}}(q, E_1, E_2) = \frac{ba^3}{(1 + a^2 q^2)^2} + \frac{c}{q^4}$$ (1)

Here, $\Delta \sigma_{\text{tot}}$ is the separated scattering intensity obtained from the total scattering curves measured at energies $E_1$ and $E_2$, $q$ ($= 4\pi/\lambda \sin \theta$) is the scattering vector with the wavelength $\lambda$ and the scattering angle $2\theta$, $a$ is the correlation length, and $b$ and $c$ are adjustable parameters. The model represents the small-angle scattering of inhomogeneities with fully random shape, size and distribution. The correlation function with the correlation length $a$ is shown for example in the inset of Figure 7d, and the related scattering curve obtained from this function (without the $q^{-4}$ term) lies normal to the ordinate. Table 2 summarizes the results obtained from fitting eq 1 to the separated scattering curves in Figure 7. The correlation lengths approximate the Pt nanoparticle sizes and agree favorably with the sizes measured by TEM.

The volume fraction of Pt nanoparticles in each specimen can likewise be deduced from the parameter $b$ if the electron density contrast of the particles relative to the HPS matrix is known. Assuming that the nanoparticles are Pt(0) (with a mass density of 21.41 g/cm$^3$), the volume fractions have been calculated according to the methodology presented earlier. From these volume fractions, the weight percents ($w_b$) of Pt in the four materials have been calculated and compared in Table 2 to results obtained from elemental analysis ($w_e$). The Pt compositions extracted from the ASAXS model lie between 35 and 51% of the compositions acquired from elemental analysis. Two plausible explanations are offered for the “missing” Pt in the ASAXS analysis: (i) the larger structures, which require the $q^{-4}$ term in eq 1 to fit the separated scattering curves, are required to contain Pt, and (ii) the electron density contrast introduced into the calculation of the volume fractions from the correlation model may be smaller than anticipated due to the presence of Pt oxides. The first explanation is confirmed by TEM images that illustrate the presence of macropores, which are expected to be partially decorated with Pt due to Pt species located in adjacent micropores. Moreover, the oxidation state of the Pt is known from the XPS spectra displayed in Figure 1 to vary (from 0 to IV) due to chemical modification in the presence of the reaction media (or possibly charging during analysis). Since these two considerations are not mutually exclusive, a combination of both explanations is most likely applicable in resolving the apparent discrepancy in Pt compositions evident in Table 2.

**Property Analysis.** The optimal conditions for L-sorbose oxidation with MN-270-Pt have been identified by systematically varying the following reaction parameters: (i) catalyst concentration ($C_C$) from 0.001 to 0.010 M Pt, (ii) L-sorbose concentration ($C_{C_{\text{L-sorbose}}}$) from 0.05 to 0.15 M, (iii) NaHCO$_3$ concentration ($C_{NaHCO_3}$) from 0.05 to 0.15 M, (iv) reaction temperature ($T$) from 60 to 80 °C, (v) reaction time ($t$) from 180 to 220 min, (vi) oxygen flow rate ($V_O$) from 7.5 × 10$^{-6}$ to 20 × 10$^{-6}$ m$^3$/s, and (vii) stirring rate from 200 to 1000 rpm. The highest selectivity obtained at 100% L-sorbose conversion during the course of this study is 96.8%. It is achieved with the MN-270-Pt nanocomposite under the following set of reaction conditions: $C_C = 0.002$ M Pt, $C_{C_{\text{L-sorbose}}} = 0.016$ M, $C_{NaHCO_3} = 0.106$ M, $T = 70$ °C, $V_O = 14 × 10^{-6}$ m$^3$/s, stirring rate = 1000 rpm, and NaHCO$_3$ added continuously for 180 min. The kinetic characteristics of selective L-sorbose oxidation with MN-270-Pt have been ascertained by adjusting the substrate/catalyst concentration ratio ($C_{C}/C_C$) from 27.0 to 77.0 mol/mol Pt at a constant O$_2$ flow rate of 14 × 10$^{-6}$ m$^3$/s and a stirring rate of 1000 rpm at 70 °C. For this analysis, the reaction medium was charged with a single shot of the alkaline agent (in an equimolar amount with respect to the L-sorbose) before the beginning of the reaction. The dependence of L-sorbose conversion on reaction time is presented in Figure 8 and demonstrates that an increase in $C_C$ at constant $C_C$ promotes an increase in reaction time ($t$), which is typical for most catalysts. These kinetic curves also verify the existence of a 60 min induction period, during which time 2-keto-L-gulonic acid yield is negligible. Comparison of this induction period with results presented in our previous study of fully microporous HPS reveals a substantial decrease in induction time from 100 to 60 min for the micro/macroporous MN-270-Pt nanocomposite. This reduction is attributed to the presence of macropores in the HPS matrix and, hence, facilitated reactant and product transport.

The effect of temperature on the kinetics of L-sorbose oxidation has also been investigated from 60 to 80 °C at the same conditions employed above, and the resulting data (see Table 3) establish that the rate of L-sorbose oxidation increases with increasing temperature. At the highest selectivity and conversion of the MN-270-Pt nanocomposite measured in this work (96.8% selectivity at 100% conversion), the value of the catalytic activity is 2.5 × 10$^{-3}$ mol/mol Pt-s, which corresponds to a 4.6× increase compared to the fully microporous HPS/Pt analogue and an even greater increase relative to other oxidation catalysts. This enhanced catalytic activity is likewise attributed to the presence of a hierarchical pore morphology.

**Table 2:** Structure Parameters and Compositions Extracted from the ASAXS Analysis

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<th>$a$ (nm)</th>
<th>$b$ (cm$^{-1}$/nm$^3$)</th>
<th>$c$ (cm$^{-1}$/nm$^3$)</th>
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<th>$w_b$ (%)</th>
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<tr>
<td>MN-270-Pt-OX</td>
<td>0.99(2)</td>
<td>2.31(4)</td>
<td>0.035(1)</td>
<td>3.24</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MN-270-Pt-3OX</td>
<td>1.02(1)</td>
<td>3.09(4)</td>
<td>0.050(1)</td>
<td>1.62</td>
<td></td>
<td>3.16</td>
</tr>
</tbody>
</table>

$^a$ The weight percent of Pt calculated from the volume fraction. $^b$ The weight percent of Pt measured from elemental analysis. $^c$ The volume fraction and weight percent of MN-270-Pt are not given, since it contains no detectable Pt(0) phase.
fully microporous HPS to 60 min for the HPS with hierarchical porosity is attributed to the presence of macropores and facilitated L-sorbose transport. The corresponding 4.6× increase in catalytic activity relative to fully microporous catalyst is further credited to the macropores and the improved accessibility of the Pt nanoparticles to the reaction medium. The surprisingly high catalytic activity and selectivity, as well as stability after three catalytic runs, render this Pt-modified HPS nanocomposite with both micro- and macroporosity a robust hybrid material for Pt-catalyzed reactions, such as the direct oxidation of L-sorbose. The superb control over nanoparticle size and stability afforded by the micro/macroporous HPS matrix further indicate that HPS can serve as a platform for the development of novel metal/polymer nanocomposites to be used in emergent technologies requiring high accessibility to the surfaces of nearly monodisperse nanoparticles.

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