Conducting Polyaniline Filaments in a Mesoporous Channel Host

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Conducting filaments of polyaniline have been prepared in the 3-nanometer-wide hexagonal channel system of the aluminosilicate MCM-41. Adsorption of aniline vapor into the dehydrated host, followed by reaction with peroxydisulfate, leads to encapsulated polyaniline filaments. Spectroscopic data show that the filaments are in the protonated emeraldine salt form, and chromatography indicates chain lengths of several hundred aniline rings. The filaments have significant conductivity while encapsulated in the channels, as measured by microwave absorption at 2.6 gigahertz. This demonstration of conjugated polymers with mobile charge carriers in nanometer channels represents a step toward the design of nanometer electronic devices.

Efforts to create electronic functions and devices based on molecules instead of bulk semiconductors are inspired by the anticipated enormous increase in computing speed and storage density (1, 2). Previous studies have shown that conjugated systems with appropriate end groups (3-5) and confined aromatic systems (6) can transfer excitations upon photon absorption or after reduction or oxidation. A major challenge is to achieve charge transfer with low fields as in metallic wires and to establish communication with individual, electrically separated nanometer structures or molecules.

We have demonstrated previously the encapsulation of several different conjugated polymers such as polypyrrole and polylyzed polycrorynitrile in the well-defined channels of zeolite molecular sieves (7-10). The template synthesis of conducting polymers in the much larger, random pores (about 0.1 to 1 μm) of insulating host membranes has also been described (11). Networks of poly(3-methylthiophene) dendrites have been grown between electrodes (12). We have now achieved the stabilization of conducting polyaniline filaments in the ordered, 3-nm-wide hexagonal channels of the mesoporous aluminosilicate host (13, 14) MCM-41. We were able to demonstrate the ac conductivity of such encapsulated filaments of nanometer dimensions.

A distinctive feature of polyaniline among the conducting polymers is that its conductivity is not only controlled by the degree of chain oxidation but also by the level of protonation in \[\{(\text{B-NH-B-NH})_y(\text{B-N}=\text{N})_{1-y}\}_{1-x}\text{(HA)}_x\] (15, 16). In the conducting form [emeraldine salt or polyaniline (PANI)], y is close to 0.5, B and Q are \(\text{C}_6\text{H}_4\) rings in the benzenoid and quinoid states, respectively, and HA is a strong acid. The MCM host was synthesized with \(\text{C}_6\text{H}_5\text{N}(-\text{CH}_3)\text{OH}\) according to (13) with a Si/Al ratio of 18 and was then calcined at 580°C and evacuated at 400°C and 10⁻² torr. Vacuum-distilled aniline was adsorbed into the host from the vapor phase at 40°C for 24 hours (Fig. 1). The aniline-saturated host (AN-MCM; 0.5 g of aniline in 1.0 g of host) was then immersed in an aqueous solution of peroxydisulfate at 0°C for 4 hours (1:1 molar ratio of oxidant versus aniline, 1 g of AN-MCM in 50 ml of 0.2 M HCl), and a drastic color change to dark green was observed (17). After a thorough washing with water, the materials were dried under vacuum. A typical polymer loading was 0.28 g per 1.00 g of MCM host.

Electron micrographs of the MCM host before and after polyaniline encapsulation show the characteristic hexagonal channel pattern (unit cell size, 4.0 nm) in both cases (Fig. 2). The similarity of the ring and C-H bending modes between PANI-loaded host (PANI-MCM) and emeraldine salt, seen in their Raman spectra (Fig. 3), show that the encapsulated polymer is in the conducting, protonated state (18). In particular, the modes at 1309 and 1364 cm⁻¹ are indicative of the protonated quinone radical cation structure. Infrared spectra of PANI-MCM and of extracted polymer from PANI-MCM resemble the ring vibration pattern of bulk emeraldine salt (peaks at 1585, 1497, 1306, 1240, and 1124 cm⁻¹) (19). The electronic absorption spectrum of PANI-MCM presents major transitions at 4.6, 3.4 (shoulder), and 1.6 eV, typical for the band-gap and polaron transitions of emeraldine salt (20). The encapsulated, evacuated polymer shows a single, fairly broad (8 G) electron spin resonance line at \(g = 2.0032\), suggesting slightly lower protonation levels than in emeraldine salt (21) or dipolar interactions with the MCM channel walls.

The relative chain length of intrachannel polyaniline [versus polystyrene (PS), in N-methylpyrrolidinone with 2.5% LiCl] was determined with gel permeation chromatography after decomposition of the host in 5% HF and deprotonation. Polyaniline is stable toward HF under these conditions. A correction factor of ~0.5 is required to estimate absolute molecular weights (22). The peak molecular weight of the intrachannel polyaniline (35,000 versus PS; correction makes it equivalent to ~190 polyaniline rings) is consistently shorter than that of bulk material synthesized under comparable conditions (52,000 versus PS; corrected to ~290 rings). This difference suggests that diffusional constraints experienced by the reactants limit the chain growth in the MCM host.

Infrared spectra of the hydroxy region of MCM-41 before aniline loading show a single peak at 3650 cm⁻¹ (in Nujol), attributable to Si-OH, which disappears on contact with aniline. This indicates complete coverage of the intrachannel surface with aniline. The intrachannel polymerization reaction was probed by nitrogen sorption. The isotherm of the PANI-MCM shows a residual pore volume of 0.40 ml per gram of MCM host, compared with 0.64 ml pore volume in the empty host (Fig. 4). If we assume a polymer density of 1.3 g/cm³ (23), the loading of 0.28 g per gram of host closely corresponds to the change in poros-

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polyemeraldine salt (23), a geometric calculation shows that about 20 aligned poly-
alinene chains could fit in the 3-nm MCM channels if the latter were completely filled.
Thermogravimetric analysis in oxygen (100 ml/min, heating at 5°C/min) indicates drastic
changes in pyrolysis rate when the polymer is encapsulated. Whereas bulk emeraldine
salt decomposes rapidly between 300° and 400°C, pyrolysis of the encapsulated polymer proceeds more slowly over a much greater temperature range (350° to 600°C). This
difference indicates diffusional constraints in the channel system. The above results show that polyaniline was synthesized within the channels of the MCM host.

Detailed studies of bulk emeraldine salt led to the model of a granular metal where charge hopping in amorphous regions between metallic bundles dominates the macroscopic conductivity (24, 25). How does the polymer behave when it is encapsulated in nanometer channels? The dc conductivity of PANI-MCM is in the 10−8 S/cm range, similar to the conductivity of un-
loaded MCM host under ambient conditions and more than seven orders of magnitude lower than that for bulk PANI. When the polymer is extracted, the dc conductivity is 10−2 S/cm, a striking difference resulting from the removal of the separating MCM channel walls. This result confirms that the polyaniline is located inside the MCM channels and that no percolating conducting paths develop on the external crystal surfaces. To probe the charge transport of PANI filaments in the insulating MCM host, we used the contactless micro wave absorption technique (26). The microwave conductivity of evacuated PANI-MCM (27) obtained from the perturbation of a rectangular cavity at 2.63
GHz is 0.0014 S/cm, one-fourth of that of evacuated bulk PANI (0.0057 S/cm), after correction for the volume fraction of the polymer in the host. As synthesized, bulk polyaniline has a microwave conductivity of 0.018 S/cm and a dc conductivity of 0.2 S/cm. The significant low-field conduc-
tivity of the polymer filaments demonstrates that conjugated polymers can be encapsu-
lated in nanometer channels and still support mobile charge carriers. In a previous study, we established that oxidized polypr-
role chains in the channels of zeolite Y and mordenite (0.7-nm channels) do not exhibit significant ac conductivity up to 1 GHz (28). The charge carriers are trapped in those systems, probably because of electro-
static interactions with the channel walls and the lack of interchain contacts. In contrast, the channels in the MCM host provide more space and allow some important interchain contact to occur.

REFERENCES AND NOTES
1. C. Carter, Molecular Electronic Devices (Dekker, New York, 1982).
18. Raman spectra of polyaniline obtained with 632.8-
nm excitation have been published [Y. Furukawa, T. Hara, Y. Hyodo, J. Harada, ibid. 16, 189 (1986)]. The spectral positions observed here with excitation at 752 nm are shifted because of different resonance conditions. With excitation at 632.8 nm, a peak at ~1590 cm−1 in the polyaniline base form was assigned to a substituted benzene ring vibration, which is shifted to ~1580 cm−1 in the salt form. A peak at 1471 cm−1 was associated with the mode of the unprotonated quinone dimine structure (mixed C=N and ring mode) that shifts to much lower energy on protonation (1344 or 1334 cm−1), indicating a delocalized linkage between the rings. The latter vibrations can serve as markers for the proto-
nonated form. Another band of the resulting quinone radical cation structure was observed at 1484 cm−1. Also, C–H in-plane bending occurs at about 1160 cm−1.
23. The densities of crystalline model compounds of
Role of Deep Cloud Convection in the Ozone Budget of the Troposphere

Jos Lelieveld and Paul J. Crutzen

Convective updrafts in thunderstorms prolong the lifetime of ozone (O₃) and its anthropogenic precursor NOₓ [nitric oxide (NO) + nitrogen dioxide (NO₂)] by carrying these gases rapidly upward from the boundary layer into a regime where the O₃ production efficiency is higher, chemical destruction is slower, and surface deposition is absent. On the other hand, the upper troposphere is relatively rich in O₃ and NOₓ from natural sources such as downward transport from the stratosphere and lightning; convective overturning convays the O₃ and NOₓ toward the Earth’s surface where these components are more efficiently removed from the atmosphere. Simulations with a three-dimensional global model suggest that the net result of these counteractive processes is a 20 percent overall reduction in total tropospheric O₃. However, the net atmospheric oxidation efficiency is enhanced by 10 to 20 percent.

Deep cumulonimbus (Cb) convection is a primary mechanism for transporting heat, moisture, and momentum from the lower to the upper troposphere, thus forcing the general circulation of the atmosphere (1). In the Cb outflow region in the cold upper troposphere, cirrus clouds develop and cover areas much larger than the cores of these convective systems. The vigorous motions in the Cb cores, which tend to maximize in the mid-troposphere, are driven by the release of latent heat by condensation (2). Conversely, mesoscale descent in the cloud environment is caused by radiative and evaporative cooling and melting of ice (3). Upward transport through Cb clouds takes about 0.5 to 1 hour, whereas the mesoscale subsidence takes about 10 to 50 hours over a proportionally larger area (4). Although thunderstorms constitute a major force in the convective overturning of the troposphere, especially in the tropics, synoptic disturbances such as extratropical cyclones also cause rapid vertical mixing; vertical velocities in these systems are highest in the region of organized convection ahead of the frontal zone (5). Deep convective motions play an important role in the vertical redistribution of trace gases (6). In their absence, surface-emitted pollutants would be transported to the upper troposphere in several months, whereas through their presence large parts of the troposphere are mixed on much shorter time scales. In this study, a global three-dimensional transport–chemistry model of the troposphere was used to evaluate the effects of convection on the chemistry and distribution of O₃.

Ultraviolet radiation causes the photochemical formation of O₃ in the background troposphere; this occurs mostly through the oxidation of carbon monoxide (CO) and methane (CH₄), catalyzed by NOₓ (7). The global budgets of these gases are dominated by anthropogenic emissions in the lower troposphere, notably from combustion (Table 1). In addition, in polluted or forested areas the oxidation of reactive hydrocarbons can play a significant role. In this study, we are concerned with the background troposphere; thus, we concentrate on the effects of CO and CH₄ oxidation. The lifetime of NOₓ in the troposphere is about 1 to 2 days, and that of CO is a few months. Hence, convection strongly alters the vertical profiles of these species, particularly that of NOₓ. The photochemical O₃ yield per emitted NOₓ molecule decreases as NOₓ concentrations increase (8), so that the convective dilution of surface-emitted NOₓ from the boundary layer into the free troposphere enhances the formation of O₃. Moreover, the lifetime of NOₓ in the upper troposphere is longer by up to an order of magnitude than in the boundary layer. Chemical conversions and dry deposition act on NOₓ rather than NO. Ultraviolet radiation fluxes and thus photolysis rates are relatively high and temperatures are low in the upper troposphere, which shifts the equilibrium between NOₓ and NO toward NO. Also, in the absence of deposition processes nitric acid (HONO) accumulates so that NOₓ can partly reform through photodissociation and reaction with hydroxyl (OH). Furthermore, the lifetime of O₃ that has been formed photochemically in the boundary layer is enhanced by upward convective transport. At low altitudes, O₃ is efficiently destroyed by deposition on land surfaces (9) and through O₂ photolysis and subsequent reaction of O(1D) with water vapor; the latter reaction, yielding OH, is most efficient in the humid lower troposphere. Consequently, it has been concluded that convective transport of O₃ and its surface-emitted precursors enhances the tropospheric O₃ abundance (6, 10). We show, however, that this assessment may not be true.

In the classical theory regarding tropospheric O₃, downward transport from the stratosphere is the dominant O₃ source (11). Although there is no agreement about the relative contributions of downward O₃ mixing and in situ photochemical O₃ formation, several studies suggest that the anthropogenic release of O₃ precursors is of major importance in the contemporary troposphere.

The large model of trace-gas emissions (in teragrams per year) of which two-thirds are from anthropogenic sources in the lower troposphere.

Table 1. Model trace-gas emissions (in teragrams per year) (10), of which two-thirds are from anthropogenic sources in the lower troposphere.

<table>
<thead>
<tr>
<th>Source</th>
<th>CO</th>
<th>CH₄</th>
<th>NOₓ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fossil fuel use</td>
<td>450</td>
<td>115$</td>
<td>25</td>
</tr>
<tr>
<td>Biomass burning</td>
<td>400</td>
<td>30</td>
<td>9</td>
</tr>
<tr>
<td>Natural</td>
<td>470$</td>
<td>135$</td>
<td>11.5$</td>
</tr>
<tr>
<td>NMHC oxidation</td>
<td>230$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ruminants</td>
<td>105$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rice paddies</td>
<td>70</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Landoilis</td>
<td>25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sewage</td>
<td>25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>1550</td>
<td>520</td>
<td>45.5</td>
</tr>
</tbody>
</table>

$Includes 5 TG/yr from CH₄ hydrate destabilization.
†Three hundred teragrams per year from natural nonmethane hydrocarbon (NMHC) oxidation in the atmosphere, 100 TG/yr from vegetation, 40 TG/yr from oceans, and 30 TG/yr from wildfires.
‡Thirty teragrams per year from high-latitude wetlands and 70 TG/yr from low-latitude wetlands, 20 TG/yr from termites, 10 TG/yr from oceans, and 6 TG/yr from wild ruminants.
§Five teragrams of N per year from lightning, 6 Tg of N per year from soils, 0.5 Tg from the stratosphere.
||From anthropogenic NMHC oxidation in the atmosphere.

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27. Evacuated at 25°C and 10⁻¹⁰ torr for 12 hours.
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