NMR Studies of Clustering in Solids

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Abstract: We present a time-resolved multiple-quantum NMR experiment which allows us to determine the spatial distribution of atoms in materials lacking long-range order; in particular, we study the size and extent of atomic clustering in these materials. Multiple-quantum NMR is sensitive to the distribution of spins in a solid. At the two extremes, uniformly distributed spins absorb radio-frequency quanta continuously, whereas clustered groups of \( N \) spins can absorb only up to \( N \) quanta. Model experiments are demonstrated on the hydrogen distribution in selectively deuterated organic solids and in hydrogenated amorphous silicon.

In this paper, we describe the application of nuclear magnetic resonance to the question of clustering in materials lacking long-range order. In such materials, standard methods of structural characterization, such as X-ray crystallography, are not useful. Examples of such disordered materials range from minerals, semiconductors, polymers, and liquid crystals, to species adsorbed on surfaces and in zeolites. These materials often display important physical or electrical properties; for example, a critical level of hydrogen incorporation into amorphous silicon renders it a semiconductor used in many industrial applications.1

As an example of the type of problem we face in these disordered solids, Figure 1 illustrates two extreme atomic configurations: in the first, the atoms are distributed randomly but uniformly within the sample, and in the second they are grouped together, forming clusters. Usually, an NMR spectrum of a solid will not reveal the basic difference between these two situations. In both cases, the spectrum will normally be broad and featureless; the line width, due primarily to the dipolar couplings between spins, does not contain sufficient information to establish any statistical information on the atomic distribution.

In contrast, by capitalizing on the dipolar couplings in a different manner, a time-resolved multiple-quantum NMR experiment can be used to address the question of atomic distribution in such materials.2 For example, if the uniformly distributed material is irradiated with rf quanta in an NMR experiment, it might be expected to absorb them continuously whereas isolated clusters will absorb only a finite number.3 In this paper, we show that indeed a time-resolved multiple-quantum experiment, whose statistics are very sensitive to atomic distributions, can be used to probe the nature and extent of clustering in solids. Model systems containing a variety of hydrogen atom distributions are investigated; in particular, totally isolated clusters, various concentrations of clusters ranging from dilute to fairly dense, and uniformly distributed environments are all studied. We illustrate how these different distributions affect the multiple-quantum dynamics and how the tendency toward clustering and information on cluster sizes can be determined from the time dependence of the multiple-quantum intensities.

Time Development of Multiple-Quantum Coherences

In multiple-quantum NMR, individual spins become correlated with one another over time through their dipolar couplings.4 In this way, the usual Zeeman selection rule can be overcome, and "forbidden transitions", where the difference in magnetic quantum number \( \Delta M \) is equal to 0, 1, 2, ... , \( \pm N \), can be excited. When \( \Delta M = N, N \) spins "flip" collectively from the ground state to the highest excited state. Experimentally, such multiple-quantum coherences are formed by applying an appropriate radio-frequency pulse sequence to the system for a time \( \tau \), thereby creating a Hamiltonian which forces the spins to act collectively. For spins to become correlated, the system must evolve under this Hamiltonian for a time \( \tau \) roughly proportional to the inverse of the pairwise dipolar interactions. Because the magnitude of the dipolar interaction is inversely proportional to the cube of the distance between two spins \( i \) and \( j \),

\[
D_{ij} \propto \left( \frac{1}{r_{ij}^3} \right) P_2(\cos \theta_{ij})
\]

(1)

two spins which are far apart require more time to communicate with one another than spins which are closer together. Therefore, the rate at which multiple-quantum coherences develop is determined by the dipolar coupling distributions present in the system. Because the spatial arrangements of atoms are reflected by the magnitude of the dipolar couplings, a time-dependent

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quantum NMR. Distributions can be distinguished by time-resolved solid-state multiple-quantum orders arise in the spectrum. The system is no longer bounded and higher multiple-quantum orders will be addressed in more detail in a later section.

Schematically, the interspin correlations which occur during the development of multiple-quantum coherences are represented in Figure 3. If a solid is composed of a uniform distribution of atoms, where the dipolar couplings between spins may be roughly comparable, then the time development will be expected to look like the one depicted in Figure 3a. As the uniformly distributed spins absorb more and more quanta of radiation, the spins become correlated with one another in a continuous manner. Over time, the number of communicating spins is essentially unbounded and the effective “size” of the system grows monotonically. However, if a solid contains isolated clusters, then the variation between inter- and intracluster dipolar couplings is large enough to preclude intercluster correlations on the experimental time scale. Consequently, as shown in Figure 3b, after an initial induction period, during which multiple-quantum coherences develop between spins with large dipolar couplings, the number of correlated spins will be expected to remain roughly constant over time and will reflect the size of the isolated clusters. If this group is truly independent of any other groups, then no additional interactions can build up on the experimental time scale. On the other hand, if small but nonnegligible dipolar couplings do exist between spins of different groups, then with time the groups will communicate with one another. If the concentration of clusters is high, less time will be needed for intergroup communication to occur. Ultimately, for very long times, a large network of correlated spins will develop.

Experimental Methods
Multiple-quantum NMR in solids has been described in detail elsewhere, and we outline here only the salient points relevant to this work. The two-dimensional multiple-quantum experiment is divided into four periods, each characterized by a time variable: preparation ($t_P$), evolution ($t_E$), mixing ($t_M$), and detection ($t_D$). The pulse sequence design is shown in Figure 4. The radio-frequency pulse cycle of the preparation period is composed of eight $\pi/2$ pulses of length $t_p$ separated by delays $\Delta$ and $\Delta' = 2\Delta + t_p$. The phases of the first and last four pulses are 0 and 180°.

Multiple-quantum NMR in solids can be distinguished with one another over time, through pairwise dipolar interactions; the closer the spins, the shorter the time needed for interactions to develop. For uniformly distributed spins, correlations will be expected to develop monotonically with time. In a clustered material, however, the magnitudes of the inter- vs. intra-cluster dipolar couplings are quite different. At short times, the number of correlated spins will be limited to the number of atoms in the cluster, whereas at long times intercluster interactions will develop and all the spins will become correlated with one another. The difference between clustered and uniform distributions is observable in the time dependence of the $n$-quantum absorption seen in Figure 2.

In multiple-quantum NMR, spins, symbolically represented by hatched circles, become correlated with one another over time, through pairwise dipolar interactions; the closer the spins, the shorter the time needed for interactions to develop. For uniformly distributed spins, correlations will be expected to develop monotonically with time. In a clustered material, however, the magnitudes of the inter- vs. intra-cluster dipolar couplings are quite different. At short times, the number of correlated spins will be limited to the number of atoms in the cluster, whereas at long times intercluster interactions will develop and all the spins will become correlated with one another. The difference between clustered and uniform distributions is observable in the time dependence of the $n$-quantum absorption seen in Figure 2.

are effectively isolated from one another owing to the large perchlorinated cyclopentadiene rings. For this model clustered material, the appearance of the multiple-quantum spectra remains very similar up to preparation times of 300 ps, after which many high multiple-quantum orders begin to develop. These higher orders are attributed to intermolecular interactions. This material will be addressed in more detail in a later section.

Multiple-quantum experimen can yield information concerning the distribution of spins in a material.

Monitoring the time development of the multiple-quantum coherences in an infinite spin system reveals that correlations develop between spins in a monotonic fashion. As the time progresses, more spins can absorb more quanta of radiation and very high multiple-quantum orders can be detected, experimentally in excess of 100. However, if clusters of spins exist, then the time development of the multiple-quantum coherences will be interrupted; the number of interacting spins will be limited, to a large extent, by the size of the cluster. Shown in Figure 2 are $^1$H multiple-quantum spectra for a polycrystalline sample (shown in the inset) in which hydrogen atom clusters of different molecules are effectively isolated from one another owing to the large perchlorinated cyclopentadiene rings. For this model clustered material, the appearance of the multiple-quantum spectra remains very similar up to preparation times of 300 ps, after which many high multiple-quantum orders begin to develop. These higher orders are attributed to intermolecular interactions. This material will be addressed in more detail in a later section.

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Figure 4. Two-dimensional multiple-quantum pulse sequence for solids. The preparation and mixing periods are composed of eight π/2 pulses of length $t_p (=2.5\ \mu s)$ separated by delays $\Delta (=2.5\ \mu s)$ and $\Delta' (=7.5\ \mu s)$. The total preparation/mixing time $\tau$ is achieved by repeating the basic 60-μs cycle $m$ times. When the phases of the pulses are $x$ and $y$, the Hamiltonian is equal to $1/2(H_{yx} - H_{xy})$; during the mixing period, when the phases of the pulses are $y$ and $\dot{y}$, the Hamiltonian becomes $-1/2(H_{yx} + H_{xy})$, i.e., the negative of the preparation period Hamiltonian. To separate the multiple-quantum orders, the preparation period pulses are incremented by a phase $\phi$ for each value of $t_i$. During the detection period, a pulsed spin-locking sequence is used for signal enhancement in samples containing low $^1H$ concentrations.

respectively. The total preparation period ($\tau$) is created by repeating the 60-μs basic cycle $m$ times. This combination of pulses and delays creates a two-quantum average Hamiltonian:

$$H = \frac{1}{2}(H_{yx} - H_{xy}) = -\frac{1}{2}\sum_{i=1}^{N} D_{ij}(I_{i}J_{j} + I_{j}J_{i})$$

(2)

where $I_{i}$ and $J_{j}$ are raising and lowering angular momentum operators. This Hamiltonian allows multiple-quantum coherence of even order to develop. In addition, different multiple-quantum orders are separated from one another by using the method of time proportional phase incrementation. Here, the preparation period pulses are phase shifted by $\phi$ for each value of $t_{i}$. The total bandwidth in $t_{i}$ was 2.5 MHz, resulting in a separation between orders of 45°. The detection period was sampled after the first pulse and after each $\dot{y}$ pulse; usually 512 or 768 spin locked points were acquired, subject to the constraints of the $T_{1}$'s of the material. All these points are then averaged together to give a final point for a particular value of $t_{i}$. The entire sequence is repeated for different values of $t_{i}$ to obtain a multiple-quantum interferogram. Fourier transformation with respect to $t_{i}$ then yields the multiple-quantum spectrum for a particular preparation time. At very long preparation times, the overall integrated intensity of the signal decreases owing to accumulation of experimental error as the basic pulse cycle is repeated numerous times.

During the detection period, a pulsed spin-locking sequence of the form $(\pi/2)\{(\pi/2)\}$ is used, resulting in a large increase in signal to noise over the detection scheme used earlier. The pulsed spin locking is not a necessary component of the experiment, but is useful for signal enhancement in samples containing low $^1H$ concentrations. To obtain the maximum signal, the optimal value of $\phi$ is roughly 45° and the delay between pulses 40 μs. One point is sampled after the first pulse and after each $\dot{y}$ pulse; usually 512 or 768 spin locked points were acquired, subject to the constraints of the $T_{1}$'s of the material. All these points are then averaged together to give a final point for a particular value of $t_{i}$. The entire sequence is repeated for different values of $t_{i}$ to obtain a multiple-quantum interferogram. Fourier transformation with respect to $t_{i}$ then yields the multiple-quantum spectrum for a particular preparation time. At very long preparation times, the overall integrated intensity of the signal decreases owing to accumulation of experimental error as the basic pulse cycle is repeated numerous times.

Experiments were performed on a homebuilt spectrometer operating at a 4.7 T Larmor frequency of 180 MHz. The spectrometer was equipped with quadrature phase generation circuits that produce $\pi$ pulses with relative phases of 0, 90, 180, and 270°. Additional phase shifts needed for TPPI were performed by a digitally controlled phase shifter, in series with the quadrature generation, capable of 1.4° phase increments. For most spectra, obtaining a 16 multiple-quantum orders was sufficient. The orders were accommodated by incrementing the phases of the preparation

Figure 5. Top: 180-MHz $^1H$ multiple-quantum spectrum of the liquid crystal, p-benzyl-$p$-cyanophenyl in the nematic phase, for a preparation time $\tau = 660\ \mu s$. Bottom: The data points are measured from the spectrum above by integrating each peak corresponding to each multiple-quantum order. The resulting intensities are then fit to a Gaussian distribution whose standard deviation, $\sigma$, is associated with $(N/2)^{1/2}$. The solid curve is the plot of the best fit value for $N$. The effective system size, characterizes the number of correlated spins, in this case 20, at each preparation time.

Results and Discussion

Measurement of Cluster Sizes. The number of correlated spins at a particular preparation time dictates the intensity distribution of the signal over the multiple-quantum orders. For a finite cluster of size $N$, ignoring symmetry, the integrated spectral intensity per order can be related to the number of multiple-quantum transitions within that order. These can be calculated directly from combinatorial arguments; for example, the number of $\Delta M$ quantum transition in an $N$ spin system is

$$\binom{2N}{N - \Delta M} = 2N!(N - \Delta M)!(N + \Delta M)!$$

(3)

References


which can be approximated by a Gaussian distribution for large \( N \) and \( n \ll N \). As a result, the integrated multiple-quantum intensities fall off in a Gaussian manner, \( I(n) \sim \exp(-n^2/N) \), and are dictated by the size of the system.

The simplest assumption for a solid, which consists essentially of an infinite number of spins, is to subdivide the infinite spin system into finite spin systems which grow in the time-dependent fashion already portrayed by Figure 3. Thus we can represent the number of spins that have become correlated for a preparation time \( \tau \) by an effective system size \( N(\tau) \). This time-dependent parameter is calculated as shown in Figure 5, by fitting the integrated intensities to a Gaussian and associating the standard deviation, \( \sigma \), of the Gaussian with \( (N/2)\sigma \). The pattern and rate of growth of \( N(\tau) \) over time will reflect the distribution of atoms in the sample. In the following sections the time-dependent behavior of isolated clusters, concentrations of clusters, and uniform distributions will be examined from this point of view.

**Isolated Clusters: Liquid Crystal.** In a nematic liquid crystal sample, owing to rapid translational diffusion, intermolecular dipolar couplings are averaged to zero while intramolecular couplings remain large.\(^{13}\) The nematic phase of the 21-spin \( p \)-heptyl-\( p' \)-cyanobiphenyl (molecular formula shown in Figure 6) liquid crystal sample was thus selected to demonstrate an "ideal" case of isolated clusters on the NMR time scale, in the sense that individual molecules behave like clusters while still being completely independent of one another. Thus, this liquid crystal which contains 21 protons should be a good model for a 21-spin cluster. Figure 6a shows a plot of the \( n \)-quantum intensity, normalized to the total intensity, vs. preparation time; at short times, the number of multiple-quantum orders increases, whereas for times greater than 1000 \( \mu \)s the number of orders and their relative intensities remain unchanged. Referring to the trend of the multiple-quantum intensity plot, the effective system size \( N(\tau) \), plotted in Figure 6b, grows for times up to 1000 \( \mu \)s, after which it remains completely constant with a value of 21. The effective system size does not grow beyond the actual size of the molecule, confirming that intermolecular couplings are zero and that only spins within the molecule can become correlated with one another. This type of behavior, the leveling of \( N(\tau) \), is characteristic of isolated clusters.

**Concentration Effects: 1,8-Dimethylnaphthalene-\( d_9 \).** Solid solutions of six spin clusters were prepared to examine the effects of different degrees of cluster concentrations on the multiple-quantum dynamics. The six spin clusters are formed by intimately mixing 1,8-dimethylnaphthalene-\( d_9 \) (DMN-\( d_9 \)) with perdeuterated DMN-\( d_{12} \); three concentration levels (5, 10, and 20 mol %) were prepared in addition to the neat material. For the latter, the shortest intermolecular \( \text{H}^+ \text{H}^- \) distance is 2.0 A along the \( b \) axis, and the intramolecular methyl groups are separated by 2.93 A. The unit cell is arranged such that the methyl groups of a pair of molecules are pointing toward one another.\(^{14}\) Dilution of the protonated DMN-\( d_9 \) in a perdeuterated lattice forces the intercluster distances to increase without affecting the intracluster distances. Therefore, relative differences in the development of

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and the inter- and intramolecular couplings comparable enough to uniform environment of hydrogen atoms. In the intermediate cases, where clusters exist but are not sufficiently isolated to preclude small interactions between them, a second time-dependent parameter is introduced to ascribe a size to the clusters independently of intercluster events. A schematic drawing of a two-Gaussian model is presented in Figure 9. The simplest approximation is to attribute the multiple-quantum intensities to two independent events: the intra-cluster correlations which have already matured, and the inter-cluster correlations that continue to develop between spins of different groups. The multiple-quantum intensities due to the subgroup of clustered spins is approximated by a Gaussian whose variance is associated with $N_c$, the cluster size obtained from the two-Gaussian model, and $N_a$, the open ones values for $N_c$. The multiple-quantum intensities, or when $N_a > N_c$, can the inter- vs. intracluster interactions be addressed separately to obtain $N_a$.

Figure 9. Schematic drawing of the two-Gaussian model. The inset represents clusters interacting with one another on the experimental time scale. In order to determine the cluster size, a parameter $N_c$ is introduced to describe events occurring within a cluster, independently of those occurring between clusters. The multiple-quantum intensities (filled circles) are the sum of intra and inter-cluster correlations, both of which are associated, independently, with a Gaussian of standard deviation $(N_c/2)^{1/2}$ and $(N_a/2)^{1/2}$, respectively. Only when most of the multiple-quantum intensity is due to intra-cluster correlations is this approximation valid.

Two-Gaussian Model. In the intermediate cases, where clusters are varied entirely freely. To begin the search, the first guess is calculated by fitting the integrated intensities of orders $n > 2$ and $n = 4$ to a Gaussian of variance $N_c$ with magnitude $m_a$, and also by independently fitting the intensities of the two highest orders to a Gaussian of variance $N_a$ with magnitude $m_a$. Then the four parameters are varied to obtain the best fit of the multiple-quantum intensities to the two-Gaussian model; the mean square error is between 0.98 and 0.99, indicating that if distributions of cluster sizes exist, they occur in very low concentrations.

Now the two time-dependent parameters which will be considered are $N_c$, the cluster size obtained from the two-Gaussian model, and $N_a$, obtained by fitting the integrated intensities to a single Gaussian as before. The pattern of growth of both parameters is used to establish the size and extent of clustering.
in solid samples. In the same spirit as for \( N(\tau) \), absence of change in \( N_c(\tau) \) over time is evidence that clusters exist, although they may be near one another. The magnitudes of \( m_c, m_s, \) and \( N_c \) change to account for the increasing number of communicating clusters over time until finally \( m_s \) becomes larger than \( m_c \) and the approximation is no longer valid.

Returning to Figure 8, the values of \( N_c(\tau) \) for the four concentrations of DMN are now plotted along with the values of \( N(\tau) \) discussed earlier. Considering the two extremes once again, we note that \( N_c(\tau) \) is not necessary to determine the cluster size in the 5 mol % solution, as here \( N_c(\tau) \) remains fixed for a sizable duration of time. \( N_c(\tau) \) can be calculated at longer times and results in values close to 6. By contrast, \( N_s(\tau) \) and \( N(\tau) \) both increase in the 100% DMN sample, confirming that the density of spins is too high to assign a cluster size to this material. For the intermediate concentrations of 10 and 20 mol % where \( N(\tau) \) grew continuously, \( N_c(\tau) \) now remains constant over time with values about 6. These data show what we should expect in a sample like DMN containing predominantly clusters of six atoms which become increasingly close to one another as the \( ^1H \) concentration is raised.

1,8-Dimethyl-naphthalene-d_{10} (DMN-d_{10}) is very similar to 1,8-DMN-d_{8}, but here, in addition to ring deuteration, the methyl groups are partially deuterated as well, leaving only two hydrogen atoms on the entire molecule. The inter- and intramolecular \( ^1H\text{-}^1H \) distances are not affected by the additional methyl deuteration. Two solid solutions of DMN-d_{10}, 5 and 10 mol %, were prepared in the same manner as those of DMN-d_{8}. Plots of the integrated \( n \)-quantum intensities appear in the upper portions of Figure 10, parts a and b. In the 5 mol % solution, where the two-quantum intensity is essentially dominant at all times, the effective system size, \( N(\tau) \), is calculated directly from the binomial formula of eq 3. For the 10 mol % solution, the combinatorial formula is used up to 600 \( \mu \text{s} \), after which the intensities can be approximated by a Gaussian as usual.

A comparison of the six-spin and two-spin model clusters at comparable \( ^1H \) concentrations reveals similar trends for \( N(\tau) \) and \( N_c(\tau); \) in both 5% cases, \( N(\tau) \) remains level up to approximately 500 \( \mu \text{s} \) after which it increases slowly, whereas in the 10% cases \( N(\tau) \) rises more steeply. What distinguishes the two materials from one another are the actual values of the effective system and cluster sizes. In DMN-d_{10}, contrary to DMN-d_{8}, \( N(\tau) \) remains close to 2; in the 10% solution, \( N_c(\tau) \) can be calculated at long times and lies between 3 and 4. These results demonstrate that the multiple-quantum dynamics are sensitive enough to distinguish clearly between two-spin and six-spin clusters of very similar compounds.

Hexamethylbenzene. Protonated molecules of hexamethylbenzene (HMB) are mixed with perdeuterated molecules to create clusters of 18 atoms. Two molecular motions exist in HMB at room temperature: each methyl group reorients about its C, axis, and the entire molecule undergoes fast limit sixfold hopping about the C6 axis of the benzene ring. In neat HMB, C–C distances between molecules range upward from 3.7 to 6.6 A. Solutions

![Figure 10](image1.png)

**Figure 10.** \( n \)-Quantum intensity (top) and number of correlated spins (bottom) vs. preparation time for two solid solutions: (a) 5 mol % 1,8-dimethyl-naphthalene-d_{10} (diagrammed in the inset) in DMN-d_{12}, and (b) 10 mol % DMN-d_{10} in DMN-d_{12}. For the 5 mol % solution \( N(\tau) \) remains close to 2. In the 10 mol % solution, \( N(\tau) \) grows with time but \( N_c(\tau) \) remains essentially constant, hovering between 3 and 4. The cluster size in these samples is very small.

![Figure 11](image2.png)

**Figure 11.** Number of correlated spins vs. preparation time for hexamethylbenzene-\( h_{18} \) (shown in the inset) in hexamethylbenzene-d_{18}. In this 5 mol % solution, \( N(\tau) \) grows steadily whereas \( N_c(\tau) \) remains level at roughly 15 indicating fairly large clusters.


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Figure 12. (a) n-Quantum intensity vs. preparation time for the polycrystalline sample 1,2,3,4-tetrachloronaphthalene-bis(hexachlorocyclopentadiene) adduct. The two-quantum intensity is not plotted so that higher multiple-quantum orders can be seen more clearly. The time development of the multiple-quantum intensities is discontinuous. (b) Number of correlated spins vs. preparation time for this sample. The effective system size remains at four for times ranging up to 300 μs, indicating that only the four hydrogen atoms within the molecule have become correlated thus far. At long times, a large number of intercluster interactions can develop.

Figure 13. Schematic three-dimensional drawing of clustered monohydrides in hydrogenated amorphous silicon (a-Si:H). The silicon atoms are in a tetrahedral bonding configuration. Silicon atoms, hydrogen atoms, and covalent bonds are represented by open circles, filled circles, and solid lines, respectively. The drawing is intended to motivate questions concerning the distribution of hydrogen in a-Si:H thin films.

Figure 14. Number of correlated spins vs. preparation time for: (a) 50 atom % H and 8 atom % a-Si-H samples. For the 50 atom % sample, N(τ) grows very rapidly, whereas for the 8 atom % sample, N(τ) remains essentially constant at 6; (b) 16 atom % a-Si:H sample. N(τ) remains nearly level up to 250 μs, after which it increases slowly; N(τ) levels off at approximately 6. Both the 16 and 8 atom % samples contain clusters of approximately six atoms; as the hydrogen level is raised, the cluster concentration is increased as well.

Then shoots up rather rapidly. From these trends it is clear that correlations develop very quickly for hydrogen atoms within the molecule; at longer times, the smaller dipole couplings become more important and communication occurs between spins on different molecules as well. Because N(τ) and N(τ) both increase at longer times, this sample appears to have similar properties to the neat DMN sample shown in Figure 8d. Perhaps it can be best thought of as a borderline case where the cluster size can be determined from the plateau in N(τ) at short times.

The following equations are derived from the Hamiltonian $H_{ss} = J_{ss}(I_{ss} - I_{ss})$.

1. $N(τ) = \text{constant}$
2. $N(τ) = \text{constant}$
3. $N(τ) = \text{constant}$


to the narrow component is present as spatially isolated monohydrides and molecular H₂. The remaining hydrogen, which consists of between 4 and 16 atom % H in device quality films, is attributed to clusters of monohydrides.

We have used multiple-quantum NMR to determine the size and extent of clustering in a-Si:H thin films.¹ Shown in Figure 14 are plots of the number of correlated spins vs. time for a number of samples containing different concentrations of hydrogen. In Figure 14a, two extremes are presented: a 50 atom % sample for which the effective system size grows rapidly and continuously, and an 8 atom % sample for which N(τ) remains essentially constant over time. The trends in these plots indicate that the polymeric sample consists of a continuous network of atoms, whereas the 8 atom % sample is composed of clusters of approximately six atoms. A plot of the number of correlated spins for an a-Si:H sample containing 16 atom % H is shown in Figure 14b: values of N(τ) increase over time, whereas N(τ) levels off at 6. Similarly to the 8 atom % sample, the size of the clusters in the 16 atom % sample is approximately six, but now they exist in higher concentrations, as evidenced by the more rapid growth of N(τ). The values in these plots are very similar to those of 1,8-DMN in perdeuterated DMN shown in Figure 8.

Summary

Multiple-quantum NMR, a technique which induces spins to act collectively through their dipolar couplings, is used to determine the spatial distribution of atoms in materials lacking long-range order; in particular, the size and extent of clustering is probed. Based on the proximity of spins to one another, correlations between them will develop more or less rapidly. A time-resolved multiple-quantum experiment measures the number of correlated spins and the rate at which these develop. The key feature in the time-dependent experiments is that in clustered materials, where groups are physically isolated from one another, the effective system size grows rapidly and continuously, in higher concentrations, as evidenced by the more rapid growth in the 16 atom % sample. In the 8 atom % sample, the size of the clusters is approximately six, but now they exist in higher concentrations, as evidenced by the more rapid growth of N(τ). The values in these plots are very similar to those of 1,8-DMN in perdeuterated DMN shown in Figure 8.

Abstract: The kinetics of interconversion between the conjugated (1,3) and nonconjugated (1,5) isomers of cyclooctatetraene (cot) complexes of CpCo [Cp = C₅H₅] 1 and *CpCo [Cp = C₅Mes] 2 have been studied. Dynamic NMR studies were employed for neutral CpCo(cot) and gave a rate of 3 × 10⁻⁹ s⁻¹ at 370 K, extrapolated to 4 × 10⁻⁷ s⁻¹ at 398 K. The permethylated complex isomerizes more slowly, k = 3 × 10⁻⁴ s⁻¹ at 370 K. However, fast fourier transform ac polarography results show that the monooxamers of CpCo(cot) and *CpCo(cot) isomerize at the same fast rate, 2 × 10⁻⁷ s⁻¹ at 298 K. An explanation is offered in terms of the lifting of symmetry restrictions to the simplest molecular isomerization motion in the anion radicals. The dibenzocyclooctatetraene complex CpCo(1,5-dbcot) (3) reduces to a stable anion which shows no tendency to isomerize. This is explained by a likely change in electronic ground states of [CpCo(cot)]⁻ [and [CpCo(dbcot)]⁻], a conclusion supported by ESR results.

Interconversion of Conjugated and Nonconjugated Polyolefin–Cobalt Complexes in Two Oxidation States: An Electrochemical and NMR Study

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