Potential Energy Surfaces and Transition Moments of Cl Atom in Xe Solid Matrix

by

Isidore Last, Thomas F. George, Mario E. Fajardo and V. A. Apkarian

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Departments of Chemistry and Physics
State University of New York at Buffalo
Buffalo, New York 14260

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Xe solid containing a Cl atom is considered by the semiempirical diatomics-in-ionic-systems method which takes into account the charge delocalization in an ionic state and the coupling between the neutral and ionic states. The calculation shows that the Cl atom has motional freedom in substitutional trapping sites which leads to the broadening of absorption bands. The vertically accessed ionic states are treated as \( \text{Xe}^{+6}\text{Cl}_4^- \) molecules with a nonuniform positive charge distribution. The most stable ionic complex has the \( \text{Xe}_2^+\text{Cl}_4^- \) molecular configuration. Cl atoms at interstitial sites are treated as \( \text{Xe}^{+6}\text{Cl}_4^- \) molecules with \( \text{Xe}^6\text{Cl}_4^* \) excited states. The results of the calculation are in general agreement with recent experiments.
Potential Energy Surfaces and Transition Moments of Cl Atom in Xe Solid Matrix

Isidore Last* and Thomas F. George
Departments of Chemistry and Physics & Astronomy
239 Fronczak Hall
State University of New York at Buffalo
Buffalo, New York 14260

and

Mario E. Fajardo and V. A. Apkarian
Department of Chemistry
University of California
Irvine, California 92717

Abstract

Xe solid containing a Cl atom is considered by the semiempirical diatomics-in-ionic-systems method which takes into account the charge delocalization in an ionic state and the coupling between the neutral and ionic states. The calculation shows that the Cl atom has motional freedom in substitutional trapping sites which leads to the broadening of absorption bands. The vertically accessed ionic states are treated as Xe₁₂Cl⁻ molecules with a nonuniform positive charge distribution. The most stable ionic complex has the Xe₂⁺Cl⁻ molecular configuration. Cl atoms at interstitial sites are treated as Xe₆Cl molecules with Xe₆⁺Cl⁻ excited states. The results of the calculation are in general agreement with recent experiments.

*Permanent address: Soreq Nuclear Research Center, Yavne 70600, Israel
1. Introduction

Rare-gas solids containing halogen molecules demonstrate interesting spectroscopic properties when exposed to laser radiation.\textsuperscript{1-5} These properties are associated with the formation of quasistable ionic rare-gas-halogen molecules. The same processes can also be observed in gas\textsuperscript{6-11} and liquid phase\textsuperscript{12,13} rare-gas-halogen mixtures.

The formation of the ionic rare-gas-halogen molecules was investigated in detail in Ref. 1 for solid xenon initially doped with Cl\textsubscript{2} and HCl molecules. Near-UV laser irradiation of the doped solid results in emission from the quasistable ionic molecules. In pure xenon hosts, Xe\textsuperscript{+}Cl\textsuperscript{-} is responsible for this emission, whereas in mixed rare gas hosts, such as HCl:Xe:Ar or HCl:Xe:Kr (1:4:100), both Xe\textsuperscript{+}Cl\textsuperscript{-} and Xe\textsubscript{2}\textsuperscript{+}Cl\textsuperscript{-} are observed. The identity of the Xe\textsubscript{2}\textsuperscript{+}Cl\textsuperscript{-} emitter in condensed media was further verified in experiments that continuously spanned the three phases -- gas, liquid and solid.\textsuperscript{13} The theoretical consideration of the ionic xenon-chlorine molecules in Xe solid also confirmed that the observed emission is associated with the Xe\textsubscript{2}\textsuperscript{+}Cl\textsuperscript{-} molecule.\textsuperscript{14} In Xe solid doped with HCl or Cl\textsubscript{2} molecules, there are several processes which lead to the formation of the Xe\textsubscript{2}\textsuperscript{+}Cl\textsuperscript{-} molecule.\textsuperscript{1} Two-photon cooperative absorption was established as the main mechanism in the case of HCl-doped solids.\textsuperscript{1} The quantum electrodynamic formalism of two-photon cooperative absorption and its efficiency in condensed media has previously been discussed.\textsuperscript{15} In the case of Cl\textsubscript{2}-doped solids, sequential two-photon absorption could not be ruled out.\textsuperscript{1} In this mechanism the first photon dissociates the Cl\textsubscript{2} molecule and forms two Cl atoms, and the second photon transfers an electron from the Xe matrix to the Cl atom to form an activated complex which relaxes to the quasistable Xe\textsubscript{2}\textsuperscript{+}Cl\textsuperscript{-} molecule. In either scheme, the end result is the permanent dissociation of
Cl$_2$ and the creation of solids doped with atomic chlorine. The subsequent spectroscopy is conducted at low intensities such that two-photon processes are diminished. The states probed in this weak-field limit then refer to those of Cl atoms isolated in rare-gas matrices.

The processes in the rare-gas solids doped by halogen molecules and exposed to laser radiation are complicated and involve several problems which can be separately studied theoretically. We shall mention here the most important of these problems: a) motion and equilibrium localizations of halogen molecules in the host matrix; b) excitation by photons of the electronic states of the halogen molecules inside the host matrix; c) excitation by photons of the electronic states with the transfer of an electron from the host matrix to the halogen molecule; d) cage effect on the dissociation of the electronically excited halogen molecules and ions (Cl$_2^+$ and Cl$_2^-$, for example); e) motion and equilibrium localizations of halogen atoms in the host matrix; f) excitation by photons of the electronic states with a transfer of an electron from the host matrix to the halogen atom. g) relaxation of atoms after the electron transfer and the formation of quasistable ionic rare-gas-halogen molecules; h) spectroscopic properties of the ionic rare gas-halogen molecules in the rare-gas matrix.

The unambiguous interpretation of experimental data, in particular excitation spectra, requires an understanding of the initial Cl atom site distributions. These are determined by the photodissociation dynamics and subsequent atomic motions as alluded to in (d) and (f) above. Reliable potential energy surfaces are crucial for such studies. The last problem (h) for the molecule Xe$_2^+$Cl$^-$ was considered in Ref. 14 within the framework of a relatively simple model. In this model, any exchange interactions between the Xe$_2^+$Cl$^-$ atoms and the matrix Xe atoms were neglected as well as
the coupling of the ionic $\text{Xe}_2^+\text{Cl}^-$ state with neutral $\text{Xe}_2\text{Cl}$ electronic states. The geometry of the ionic $\text{Xe}_2^+\text{Cl}^-$ molecule in the solid was assumed to be the same as in the gas phase. The shift of the $\text{Xe}_2^+\text{Cl}^-$ emission wavelength in the solid matrix in comparison with that in the gas phase was found to be equal to $-0.26 \text{ eV}$, not far from the experimental value of $-0.38 \text{ eV}$. In Ref. 14 the problems (e) and (f) were also considered in order to estimate the activation energy of the ionic state formation.

The consideration of most of the problems cited above needs potential energy surfaces (PES) of the system, which can be calculated by the semiempirical diatomics-in-ionic-systems (DIIS) method. This method will be used in this paper for the PES and transition moment calculations of a Cl atom inside an Xe cage (problems (e) and (f)) and the calculation of the structure of the quasistable rare-gas-halogen ionic molecules and spectroscopic transitions in these molecules (problems (g) and (h)). We shall then report some of the experimental findings and compare them to the theoretical results to establish the validity of the present approach.

2. The Diatomic-in-Ionic Systems (DIIS) Method

The semiempirical DIIS method is designed for the calculation of polyatomic rare-gas-halogen systems $R_nX$ consisting of an arbitrary number of rare-gas atoms $R$ and one halogen atom $X$. The method takes into account the coupling between ionic and neutral electronic states and the positive charge delocalization between the rare gas atoms. The wave function $\phi$ of the $R_nX$ system is given as the linear combination

$$
\phi = \sum_{i=1}^{J} \sum_{m=1}^{3} C_{im} \phi_{im}.
$$

(1)
where \( J = n + 1 \) is the number of atoms in the system, \( \phi_{im} \) are polyatomic wave functions which describe diabatic states with fixed electronic configuration of the atoms, and \( m \) is the index of the orientation of a P-symmetry atomic wave function. The main feature of the \( R_nX \) system is that in any electronic configuration one atom only has the electron shell of P-symmetry whereas all other atoms have closed electron shells of spherical (S) symmetry. In neutral configurations the halogen atom has a P-symmetry wave function. In ionic configurations a positive rare-gas ion \( R_i^+ \) has a P-symmetry wave function; all other (neutral) rare-gas atoms \( R_j (j \neq i) \) and the negative halogen ion \( X^- \) have S-symmetry electron shells. Describing the electronic configurations by the localization of the positive charge and by the orientation of the P-symmetry atomic wave function, we do not take into account, at least directly, the spin-orbit coupling. However, the shift of the levels due to the spin-orbit coupling can be accounted for indirectly by the proper choice of diatomic potentials.

The zero overlap of atomic orbitals (ZOA0) approximation, which is common in semiempirical methods, allows us to write any diabatic polyatomic function as a product of atomic group functions,

\[
\phi_{im} = \chi_{im} \prod_{j \neq i} \chi_j, \quad i = 1,2,...J, \quad m = 1,2,3 \tag{2}
\]

where the P-symmetry group function \( \chi_{im} \) describes either the neutral \( X \) atom or an ion \( R_i^+ \), and the group functions of S-symmetry \( \chi_j (j \neq i) \) describe neutral rare-gas atoms \( R_j \) and a halogen ion \( X^- \). The 3J diabatic wave functions (2) form a polyatomic basis and produce the \( 3J \times 3J \) Hamiltonian matrix. This matrix determines the eigenstates and eigenvalues of the
adiabatic wave function (1), which describes an electronic state as a mixture of the neutral and ionic configurations.

The DIIS matrix elements are expressed in terms of the potentials of diatomic fragments of the system, except for the off-diagonal matrix elements which are responsible for the coupling between the ionic and neutral electronic configurations. These matrix elements (coupling terms) are considered as semiempirical parameters which fit known empirical or \textit{ab initio} rare-gas-halogen diatomic data, other than potentials. In Ref. 16, where the Xe\textsubscript{n}Cl systems were calculated, the coupling terms were determined by fitting the transition moments between the ionic and neutral XeCl states. These coupling terms will be used in the present calculation. The diatomic potentials of the Xe\textsubscript{n}Cl fragments, namely Xe-Cl, Xe-Cl\textsuperscript{-}, Xe-Xe and (Xe-Xe)\textsuperscript{+}, will also be taken from Ref. 16, where they were obtained mostly by fitting empirical potentials.

3. Cl Atom in a Substitutional Site

Let us consider a Cl atom which replaces an Xe atom in the Xe solid matrix and is located inside a cage formed by 12 Xe atoms. These 12 Xe atoms are considered together with the Cl atom as a Xe\textsubscript{12}Cl molecule, whereas all other Xe atoms of the solid form a rigid matrix. The Xe\textsubscript{12}Cl molecule is calculated by the DIIS method, which takes into account the electron delocalization between the atoms of the molecule but neglects any exchange interaction and electron transfer between the Xe\textsubscript{12}Cl atoms and the Xe atoms of the matrix. Consequently, the energy of the system can be presented as a sum of the Xe\textsubscript{12}Cl energy (E\textsubscript{A}), the matrix energy (E\textsubscript{M}) and the energy of molecule-matrix interaction. In an ionic state the Xe\textsubscript{12}Cl molecule has, generally speaking, a dipole moment which polarizes the Xe atoms of the matrix. Consequently, the energy of the molecule-matrix interaction is
expressed as a sum of the polarization energy $\varepsilon_p$ and the energy of the non-polarization components of the pairwise interactions $\varepsilon_{ij}$ between all atoms of the $\text{Xe}_{12}\text{Cl}$ molecule and the Xe atoms of the matrix,$^{14}$

$$E = E_A + E_M + \sum_i \sum_j \varepsilon_{ij} + \varepsilon_p,$$  \hspace{1cm} (3)

where $i$ and $j$ are atomic indices of the molecule and the matrix, respectively. In the dipole approximation the polarization component of the interaction between the $i$-th ion ($\text{Xe}^+$ or $\text{Cl}^-$) and the $j$-th atom of the matrix is

$$\varepsilon_{ij} = -\frac{e^2 \alpha_j}{2R_{ij}^4},$$  \hspace{1cm} (4)

where $\alpha_j$ is the Xe atom polarizability and $R_{ij}$ is the interatomic distance. Excluding $\varepsilon_{ij}$ from the empirical of \textit{ab initio} potential $U_{ij}$, one obtains the non-polarization component

$$0_{ij} = U_{ij} - \varepsilon_{ij}.$$  \hspace{1cm} (5)

The higher-order multiple interactions are incorporated in the $U_{ij}$ term. If the $i$-th atom of the $\text{Xe}_{12}\text{Cl}$ molecule is neutral, then the polarization is zero ($\varepsilon_{ij} = 0$) and $0_{ij}$ coincides with $U_{ij}$. In the neutral $\text{Xe}_{12}\text{Cl}$ configuration all $0_{ij}$ coincide with $U_{ij}$ and $\varepsilon_p = 0$ in Eq. (3). The diatomic potentials $0_{ij}$ or $U_{ij}$ are taken from Ref. 16 where they are expressed analytically. In the present calculation, 66 matrix atoms are taken into account in the pairwise interactions in Eq. (3). In an ionic $\text{Xe}_{12}\text{Cl}^-$
configuration the polarization energy $\epsilon_p$ of the system is determined as a sum of the polarization energy of these 66 atoms, including their dipole-dipole interaction, and the polarization energy of the matrix which is considered as a continuum.\textsuperscript{14}

Since the Xe$_{12}$Cl molecule consists of 13 atoms, the order of the Hamiltonian matrix is $39 \times 39$ and the number of electronic states is 39. While any given state includes, generally speaking, both ionic and neutral electronic configurations, one of these two configurations is always predominant. We shall designate the states according to their predominant configuration. For example, a largely ionic state will be called an ionic state. In the Xe$_{12}$Cl molecule, only three states are neutral and 36 states are ionic.

A. Ground State PES

As shown by simple calculations in Refs. 1 and 14, the Cl atom has some freedom of motion inside the Xe cage. This finding is confirmed by the present more complicated DIIS calculation. When the Cl atom is located in the center of the cage and the Xe$_{12}$Cl molecule is symmetric, the minimum energy is obtained for the cage slightly shrinked, with the Xe-Cl distance of 4.33 Å, as compared to 4.4 Å in the undeformed matrix. The ground state PES is obtained for a fixed location of the Xe atoms, with the distance to the center of the cage of 4.33 Å regardless of the Cl location. The shift of the Cl atom from the center of the cage is described by spherical coordinates $(R, \theta, \phi)$ shown in Fig. 1. The angular and Cartesian coordinates of the Xe atoms are presented in Table 1.

The PES are presented in Fig. 2 for four different planes, each with a fixed $\phi$ angle. The energy in the center ($R = 0$) is -3.4485 eV. The shape of the PES is different for different values of $\phi$, but the general feature
is the presence of the relatively broad region of almost constant potential with the variations within the range of 1 meV. Twelve very shallow potential wells are shifted by 0.5-0.8 Å from the cage center toward the Xe atoms. The energy minimum of -3.4495 eV is located at the point $R = 0.65$ Å. The potential wells are separated one from another by low barriers of 0.5 meV.

Even at absolute zero temperature, the Cl atom cannot be located in a well due to the quantal motion. The uncertainty principle gives approximately 0.4 meV for the energy of the Cl atom in a well, which is enough for Cl to tunnel between the wells. The thermal energy exceeds the quantal energy and consequently increases the attainable region at -5 K. This expansion of the region of the Cl motion leads first to a decrease of the average distance $R$ to the cage center since the Cl atom at $T > 5$ K can reach the central part of the cage. Further increase in the temperature leads to a slight increase of the average distance of the Cl atom from the cage center (Fig. 3). The $R$ minimum is located at 10 K. If we take into account the quantal motion, then the average radius $R$ will be smaller for very low temperatures $T < 5$ K and the minimum will not be as well defined as in Fig. 3.

B. Influence of the Xe Atoms Shift on the Ground State Energy

The previous consideration was restricted to the case of a rigid Xe cage with Cl atom moving inside it. However, the Xe atoms can also be shifted, first because of their vibrational motion and second because of their interaction with the Cl atom. In order to check how the shift of the cage atoms affects the energy, the minimum energy geometry was calculated for a fixed location of the Cl atom and for a cage with one Xe atom shifted from the symmetrical position ($R_{Xe} = 4.33$ Å) along the radius. All other 11
cage Xe atoms are fixed in the symmetrical position. According to the results of the calculation at low temperatures T < 20 K, when the accessible region is approximately restricted to energies less than -3.447 eV (Fig. 2), the shift of a Xe atom can lower the energy by not more than 1 meV. At the minimum energy points (R = 0.65 Å, θ = 45°, ϕ = 0°, see Fig. 2a), the shift of a Xe atom stabilizes the system by lowering the energy to -3.4505 eV. However, it is unlikely that the Xe atoms can relax to trap the Cl atom in a well, as the phonon energy is higher than 1 meV. At higher temperatures the Cl atom reaches the regions associated with a stronger perturbation of the nearest Xe atom location. However, the energy of this perturbation is still smaller than the thermal energy. The changes of the transition energies and transition moments due to the vibrational motion of Xe atoms are small and can be neglected.

C. Ionic States

We consider now the PES for ionic states. As for the ground state case, the PES are expressed as functions of the Cl spherical coordinates, suggesting the Xe atoms to be fixed. The potential energy curves of all 36 ionic states along the line running from the cage center to one of the nearest Xe atoms (i = 2, θ = 45°, ϕ = 0°, for example) are presented in Fig. 4. This line is of primary interest since it crosses the minimum energy well in the ground state (Fig. 2a). The energy of the ionic states is given in Fig. 4 relative to the ground state energy for the central (R = 0) position of the Cl atom. The potential curves form a complicated system of intersecting lines with only the lowest ionic state, k = 4, being separated from the others. Most of the intersections in Fig. 4 are real (not avoided) crossings, since the states with crossings have different symmetry with respect to the considered direction which is the symmetry axis of the
crystal. Most of the diabatic curves each have a shallow minimum in the interval 1-1.2 Å.

The potential curves for a fixed radius $R = 1.1$ Å and changing angles $\theta$ and $\phi$ are presented in Fig. 5 for the twelve lowest ionic states $k = 4-15$. The potential curves in Fig. 5a are located in a plane which includes the line $\theta = 45^\circ$, $\phi = 0^\circ$ passing through the $i = 2$ Xe atom (the potentials along this line are presented in Fig. 4) and the line $\theta = 65.905^\circ$, $\phi = 26.565^\circ$ passing between two Xe atoms, $i = 1$ and 2 (Table 1, Fig. 1). Just the last line leads to the formation of the quasistable $\text{Xe}_2^+\text{Cl}^-$ molecule, which will be considered later. The potential curves in Fig. 5b are presented for a fixed angle $\theta = 90^\circ$ and the $\phi$ angle changing from $0^\circ$ to $45^\circ$. The point $\phi = 45^\circ$ on the abscissa coincides with the direction to the Xe atom $i = 1$ (Table 1, Fig. 1). In contrast to the potential curves in Fig. 4, the potential curves presented in Fig. 5 demonstrate mostly avoided crossings. Some of the avoided crossings have very small splitting of the order of $0.01 - 0.02$ eV which are difficult to detect in the figures. The lowest potential curve $k = 4$ in Fig. 5 is also separated from other curves, as in the case presented in Fig. 4. The adiabatic ionic states are denoted here by numbers, according to their energy, as it is difficult to denote them by symmetry, mainly because of the numerous crossings of the diabatic levels. A diabatic symmetry can be associated with the $k = 4$ state, which corresponds to the $4^2\Gamma$ state in the asymptotic $\text{Xe}_2\text{Cl}$ case.

The PES of the two lowest ionic states $k = 4$ and 5, are presented in Fig. 6 for the plane $\phi = 0^\circ$. The ground state PES for this plane is presented in Fig. 2a. The PES of the $k = 4$ and $k = 5$ states are similar to each other. Each of these PES has a minimum located on the x-axis which is equidistant to four Xe atoms ($i = 1, 2, 7, 8$, Table 1). The location of these
minima differ significantly from those of the ground state which are located on a line passing through a Xe atom (Fig. 2a).

D. Excitation of the System to Ionic States

In order to study the excitation of the system by photons, one needs the energy of transitions and the transition moments between the ground and excited states as a function of the Cl atom location \((R, \theta, \phi)\). We are interested in the excitation from the neutral \((k = 1, 2, 3)\) to the ionic \((k > 4)\) states. The three lowest (neutral) states are degenerate at the center of the cage \((R = 0)\) and nearly degenerate in the vicinity of it \((R < 0.4 \text{ Å})\), so that the population of the first two excited states is equal or comparable to the ground state population. Taking into account this possibility, we have to consider the excitation to ionic states from all three neutral states, at least in the region around the cage center.

The excitation energy from a neutral \(k_1\) to an ionic \(k_2\) state,

\[
V_{k_1k_2} = E_{k_2} - E_{k_1};
\]  

depends on the location of the Cl atom inside the cage and therefore is a function of the coordinates \(R, \theta, \phi\). The probability of the \(k_1 \rightarrow k_2\) excitation for the Cl atom located in a small volume \(\Delta v\) about the point \(R, \theta, \phi\) is proportional to the probability \(\frac{dW_{k_1}}{dv} \Delta v\) of the Cl location in this volume when the system is in the initial \(k_1\) state, and to the square of the transition moment \(\mathcal{M}^2_{k_1k_2}\):

\[
\Delta Q_{k_1k_2} = \mathcal{M}^2_{k_1k_2} \frac{dW_{k_1}}{dv} \Delta v. \]  

(7)
The value $Q$ can be presented in arbitrary units as it shows only the relative contribution of different transitions and different Cl atom locations to the light absorption. If the quantal motion of the Cl atom is neglected and the thermal motion only is taken into account, then the probability density is expressed in terms of the Boltzmann distribution,

$$\frac{dW_{k_1}}{dv} = A \exp\left[-\frac{(E_{k_1} - E_0)/kT}{A}\right],$$

where $kT$ is the thermal energy, $E_0 = -3.4495\text{eV}$ is the minimal energy (Fig. 2), and $A$ is the normalization coefficient. The transition energy distribution is calculated here by a boxing procedure, whereby the accessible space for the Cl atom inside the cage is divided into small volumes $\Delta v_j$ whose centers form a grid of points $R_j \theta_j$ with transition energies $V_j$ (for certain $k_1 \rightarrow k_2$). The energy scale is divided on small equal intervals $\Delta V$ ($\Delta V = 0.1\ \text{eV}$ in this calculation) with energies $V_i$ in the centers of these intervals. The relative probability $Q_i$ of the excitation with the transition energy $V_i$ is found by summing $\Delta Q_{k_1 \rightarrow k_2}$ values of all volumes $\Delta v_j$ whose transition energies $V_j$ lie in the interval

$$V_i - \Delta V/2 \leq V_j \leq V_i + \Delta V/2.$$  

The transition energy distributions were obtained for the temperatures $T = 5\ \text{K}, 10\ \text{K}, 20\ \text{K}, 40\ \text{K}, 80\ \text{K}, 160\ \text{K}$. The distribution for $T = 10\ \text{K}$ and $T = 80\ \text{K}$ are presented in Figs. 7a and 7b, respectively. The figures demonstrate the total distributions and the distributions to three individual ionic states, namely to the lowest one $k_2 = 4$ and to two higher states, $k_2 = 5$ and 20, with a strong absorption. The distributions for 10 K and 80 K do not
differ significantly from one another, leading us to the conclusion that the thermal motion does not affect the absorption to any significant extent. Both distributions have two maxima which are separated by a deep minimum. The maxima and the lower limit of the absorption are shifted slightly to smaller energies when the temperature increases. This shift is due to the expansion of the accessible area to the regions where the transition energies are smaller than in the central region.

The strongest transition at both temperatures is $1 \rightarrow 5$, i.e., from the ground state to the second ionic state. The $1 \rightarrow 5$ transition moments fall mainly within the limits $0.9 - 1.7$ D. The excitation to the first ionic state, $k = 4$, is relatively insignificant. The excitation to the third ionic state, $k = 6$ (not shown in Fig. 7), becomes important only at the higher temperature of 80 K. This increase in absorption is due to the regions with $R > 1$ Å which are accessible for higher temperatures only. In these regions the $1 \rightarrow 6$ transition moment is large, about 2.1 D, but the transition energy is relatively low so that the $k = 6$ absorption maximum has lower energy than the maximum of the absorption to the $k = 5$ state (Fig. 7b). In all other cases the energy of the absorption maximum increases with the state index $k$.

The results presented in Fig. 7 show that due to inhomogeneous broadening, the absorption forms a wide band in the range 3.5-4.7 eV, roughly, which grows weak in the middle. The total absorption cross section (in arbitrary units) is shown in Fig. 3, which is seen to increase slightly with temperature.

E. Formation of Quasistable Ionic Molecules

After the excitation of the system to an ionic state and the formation of an activated complex, the atoms begin their shift to a new equilibrium
geometry. This relaxation of the system is associated with the redistribution of the positive charge of the Xe atoms. The charge distribution in the activated complex depends significantly on the location of the Cl atom. When the Cl atom is located in the central region of the cage, then the charge is generally shared by several Xe atoms in such a way that none of the atoms can individually carry an important part of the charge. The statistical consideration of the activated complexes shows that inside the region $R < 0.7 \, \text{Å}$ there are no cases with one atom carrying more than half of the charge, and only in 30% of the cases does one atom carry more than 0.4 of the charge. When the Cl atom is located far from the cage center ($R > 0.7 \, \text{Å}$), then the charge is transferred by the excitation mostly to one of the Xe atoms, but even in these cases more than 30% of the charge is always distributed among several other Xe atoms. These distant locations do not contribute much to the absorption, at least at low temperatures. At 10 K, for example, only in 12% of the activated complexes are there Xe atoms which carry more than half of the charge. Furthermore, there are not many cases where the charge is distributed mainly between two atoms. These statistics demonstrate that the activated complexes are neither $\text{Xe}^+\text{Cl}^-$ nor $\text{Xe}_2^+\text{Cl}^-$ molecules. They have to be considered as $\text{Xe}_{12}^+\text{Cl}^-$ molecules with various positive charge distributions among the twelve Xe atoms.

In contrast to the activated complexes, the positive charge of the most stable ionic system is found to be localized mostly on two Xe atoms forming a $\text{Xe}_2^+\text{Cl}^-$ molecule inside the Xe matrix. This finding substantiates the suggestions made in Refs. 1 and 14. The relaxation from activated complexes in different ionic states to the equilibrium geometry of the lowest ($k = 4$) ionic state is possible due to the crossings of the PES demonstrated in Figs. 4 and 5 for fixed Xe atoms. The relaxation of the
system involves obviously also the charged Xe atoms. The shift of Xe atoms increases the possibility of crossings between the states including, probably, the lowest \((k = 4)\) state which is separated from other states when the Xe atoms are fixed.

In order to find the equilibrium geometry of the lowest ionic \((k = 4)\) state, the calculations were performed for different \(\text{Xe}^{+}_2\text{Cl}^-\) configurations. Varying the locations of all 13 atoms in these configurations, we selected only those directions of the atomic shifts which lead to the energy decrease. The calculations of additional configurations show that there is one only minimum energy configuration which gives the equilibrium geometry. At this equilibrium geometry the \(\text{Cl}^-\) ion is shifted from the center of the cage by 1.12 Å, forming with two Xe atoms \((i = 1, 2)\) the \(\text{Xe}^{+}_2\text{Cl}^-\) molecule. The molecule has the shape of an isosceles triangle with the Xe-Cl-Xe angle as 60.2°, each Xe-Cl distance as 3.30 Å, and the Xe-Xe distance as 3.31 Å. This geometry does not differ much from the gas-phase \(\text{Xe}^{+}_2\text{Cl}^-\) geometry in the \(4^2\Gamma\) state.\textsuperscript{16-18} The comparison with the results obtained in Ref. 16 by the same method and with the same semiempirical parameters as in the present calculation shows that the solid matrix extends slightly, by 0.06-0.07 Å, the interatomic distances in the \(\text{Xe}^{+}_2\text{Cl}^-\) molecule. The ionic charge of the \(\text{Cl}\) atom is -0.97e, almost the same as in the gas-phase molecule (-0.964e).\textsuperscript{16} The positive charge is partly delocalized so that the common charge of the two Xe atoms of the \(\text{Xe}^{+}_2\text{Cl}^-\) molecule is +0.932e, whereas +0.038e of the charge is distributed between the other ten Xe atoms. One of these atoms \((i = 3)\) carries most of this charge, +0.021e. The symmetry axis of the \(\text{Xe}^{+}_2\text{Cl}^-\) molecule is located in a plane formed by the x-axis and the diagonal line of the yz-plane. The coordinates of the \(\text{Cl}\) atom and two Xe atoms \((i = 1, 2)\) of the \(\text{Xe}^{+}_2\text{Cl}^-\) molecule are shown in Table 2. The Xe atom \(i = 3\) which carries
some charge (0.021e) is shifted further from the Cl atom by 0.1 Å. Its coordinates are also shown in Table 2. All other Xe atoms are located at the same points as in the neutral state (Table 1).

As in the gas phase, the dominant transition from the ionic $4^2\Gamma$ state of the $\text{Xe}_2^+\text{Cl}^-$ molecule is to the $1^2\Gamma$ ground state with the large transition moment of 2.41 D. The transition energy from the $\text{Xe}_2^+\text{Cl}^-$ equilibrium geometry is $V_{4+1} = 2.35$ eV. This value is smaller by 0.21 eV than the gas-phase transition energy, calculated in Ref. 16 with the same semiempirical parameters as here. The experimental value for the shift of the $\text{Xe}_2^+\text{Cl}^-$ transition energy in the solid is larger, 0.38 eV. The simple model calculation of Ref. 14 gave 0.26 eV for the shift of the transition energy in the solid, which is closer to the experimental value. It is not surprising that the present, more sophisticated calculation gives for the transition energy shift a worse result than the simple model calculation. In the model calculation the shift was estimated directly, whereas in the present calculation it is determined as the difference between the transition energies of two different systems, namely $\text{Xe}_2^+\text{Cl}$ in the gas phase and $\text{Xe}_{12}\text{Cl}$ in the solid.

The potential curves for the inner deformation of the $\text{Xe}_2^+\text{Cl}^-$ molecule are presented in Fig. 8. These potential curves allow us to calculate the vibrational energy of the molecule. In the harmonic approximation the energy of the vibrational excitation is found to be equal to 0.034 eV along the molecular symmetry axis (Fig. 8a) and 0.012 eV along the Xe-Xe line (Fig. 8b). The vibrational ground state ($v = 0$) energies are 0.017 and 0.006 eV, respectively. If the system is in thermal equilibrium and the temperature is not higher than -80 K, then only the ground state is of importance. Its vibrational motion broadens significantly the transition
1.8 energy to the repulsive electronic ground state $k = 1$, whose potential curves are shown in Fig. 8, as well. The estimations yield for the transition band width the values $\Delta V = 0.28$ eV and $0.20$ eV for the cases presented in Fig. 8a and 8b, respectively.

The potential curves of the translational motion of the $\text{Xe}_2^+\text{Cl}^-$ molecule inside the solid matrix are presented in Fig. 9. The potential wells are relatively shallow for all three mutual perpendicular directions of the translation shown in Fig. 9. The ground state ($k = 1$) potential curves are presented in Fig. 9 for the $\text{Xe}_2\text{Cl}$ geometry coinciding with the equilibrium $\text{Xe}_2^+\text{Cl}^-$ geometry. In common with the potential curves for the $\text{Xe}_2^+\text{Cl}^- (k = 4)$ molecule, the $k = 1$ curves demonstrate minima whose locations coincide exactly (Fig. 9c) or roughly (Fig. 9a, b) with the $k = 4$ minima locations. The $k = 1$ minima are slightly deeper than the ionic $k = 4$ minima. It is important, however, to note that the ground state minima in Fig. 9 do not indicate any equilibrium configuration since the ground state potentials, in contrast to the ionic $k = 4$ potentials, are repulsive along the inner $\text{Xe}_2\text{Cl}$ coordinates (Fig. 8). Because the $k = 4$ and $k = 1$ curves are nearly parallel one to another in Fig. 9, the translational motion of the $\text{Xe}_2^+\text{Cl}^-$ molecule does not change significantly the $4 \rightarrow 1$ transition energy.

4. Cl Atom at an Interstitial Position

At the interstitial position the Cl atom has 6 nearest-neighbor Xe atoms, so that the system is considered as an $\text{Xe}_6\text{Cl}$ molecule inside the Xe matrix. The Cartesian coordinates of the atoms and the Xe-Cl distances are presented in Table 3 for the minimum energy configuration. The predominant transition is to the lowest ionic state, $k = 4$, with a transition moment of 3.72 D and transition energy of 3.37 eV. This energy is lower than the
lowest limit of the absorption band obtained in the previous case for the Cl atom inside a cage (Fig. 7).

The energy to place the Cl atom at the minimum energy interstitial position is 0.104 eV, which is lower than the energy for replacing a Xe atom by a Cl atom, found to be equal to 0.156 eV. However, the difference, 0.05 eV, is too small to make the conclusion that the interstitial position is more stable for the Cl atom than the substitutional site. The energy of the interstitial position is sensitive to the diatomic potentials for relatively short distances (Table 2), where the accuracy of the diatomic potential presentation is low.16

In the lowest ionic state, k = 4, the Xe\textsuperscript{+}Cl\textsuperscript{-} geometry is almost symmetrical with a slight shift of 0.007-0.12 Å from the lattice points. The distance from the Cl atom to one of the Xe atoms is 3.16 Å and 3.78 Å to the other Xe atoms. The state k = 4 is purely ionic with approximately uniform delocalization of the positive charge among the six Xe atoms.

The Xe\textsuperscript{+}Cl\textsuperscript{-} transition energy to the ground state is 2.32 eV, which is close to the center of the experimental emission band (2.16 eV).1 However, it is difficult to identify the experimental emission as produced by the Xe\textsuperscript{+}Cl\textsuperscript{-} molecules. The emission band of the Xe\textsuperscript{+}Cl\textsuperscript{-} molecule, as well as the absorption band of the Cl atom in the interstitial position, is expected to be relatively narrow, which contradicts the experimental data (see below). It is important also to note that the quasistable Xe\textsuperscript{+}Cl\textsuperscript{-} molecule is a configuration with a local energy minimum, which may relax to the Xe\textsuperscript{+}Cl\textsuperscript{-} configuration. However, the possibility of such relaxation is not studied here.
5. **Experimental Spectra**

The emission spectrum of Xe\(_2^+\)Cl\(^-\) in solid xenon is shown in Fig. 10. The assignment of this spectrum to the molecular triatomic exciplex and its simulation by gas-phase pair potentials was presented in Ref. 1. The emission is centered at 2.16 eV and has a width of 0.22 eV (FWHM). These values are in acceptable agreement with those calculated in the present model, namely a line center of 2.35 eV and a linewidth of 0.2-0.28 eV for the 4 \(\rightarrow\) 1 transition (see Sec. 3).

Excitation spectra of Cl atoms trapped in solid xenon, xenon:argon and xenon:krypton were reported in Ref. 1. The spectra are broad and power dependent. The weak-field excitation spectra are of concern here, which were obtained by monitoring the triatomic emission while scanning the monochromatized output of a xenon-arc lamp. Two such spectra are shown in Fig. 11, where 11a was recorded immediately after photogeneration of the Cl atoms by laser irradiation at 308 nm, while 11b was recorded after annealing the solid at 55 K. The preannealed spectrum shows a single broad peak at 3.5 eV. Upon annealing, the spectrum undergoes a permanent change: several secondary peaks develop in the range 3.5-4.5 eV, and the preannealed peak diminishes to a shoulder near the absorption threshold. Once annealed, the spectrum only undergoes reversible changes in linewidth with thermal cycling. It is worth noting that while a well-defined absorption threshold is observed in these spectra, the short-wavelength limit of 260 nm is strictly experimental.

Permanent change of absorption spectra upon annealing is diagnostic of intersite conversion. Reversible line-broadening with temperature cycling is diagnostic of population equilibrium among thermally accessible initial states. The single-peaked preannealed spectrum is then indicative of a
tight trapping site with little variation in the Cl-Xe nearest-neighbor distances. The broad multiple-peaked spectrum after annealing is indicative of a looser trapping site in which a variety of Cl-Xe distances coexist and are accessible thermally. The likely assignment is then that the preannealed spectrum is due to interstitially trapped Cl, while the annealed spectrum represent substitutionally trapped Cl atoms. The spectra are then in general agreement with calculations, namely that absorption at interstitial sites leads to a single narrow spectral distribution while absorption at substitutional sites leads to a spectral distribution that shows an energy range of ~1 eV (compare Figs. 7 and 11). The calculated absorption maxima, 3.37 eV for the interstitial position and ~3.65 eV (Fig. 7b) at the substitutional site, are located close to the corresponding experimental values of 3.5 eV and ~3.75 eV.

Independent of the exact assignment of the different sites, if different ionic states are favored in the different sites, then the emission spectra would depend on the excitation wavelength, temperature and annealing history of the solid. Emission spectra identical to that shown in Fig. 10 are obtained for excitation wavelengths ranging from 375 to 280 nm, and for temperatures ranging from 12 to 40 K. The result of annealing is a permanent line-narrowing of ~200 cm$^{-1}$. The latter observation is verification of the inhomogeneity of observed lineshapes; yet in all cases the emission lineshapes are indistinguishable. The experiments then clearly establish that independent of the initial isolation site of the dopants, and therefore independent of the initial configuration of the charge-transfer complex accessed optically, the system always relaxes to the lowest energy configuration, namely Xe$^+_2$Cl$^-$. 
6. **Conclusions**

1. A Cl atom in a substitutional site can move inside the Xe cage almost freely over distances $\leq 1$ Å at the temperatures 10-20 K. The Cl atom is mostly shifted toward the cage Xe atoms.

2. Due to the Cl motion inside the Xe cage, the energy spectrum of the excitation to ionic activated complexes is broadened and has two wide maxima in the regions 320-340 nm and 270-290 nm.

3. The positive charge in the ionic activated complexes is distributed most often between several Xe atoms forming, in general, a $\text{Xe}_{12}^+\text{Cl}^{-}$ molecule.

4. In the lowest ionic state, a quasistable $\text{Xe}_{2}^+\text{Cl}^{-}$ molecule is formed in the configuration similar to that of the gas-phase $\text{Xe}_{2}^+\text{Cl}^{-}$ molecule. The calculated decrease of the emission photon energy by the matrix is 0.21 eV, in comparison with -0.36 eV in the experiment. The inner vibrational motion of the $\text{Xe}_{2}^+\text{Cl}^{-}$ molecule increases the width of the emission band to 0.3 eV, approximately.

5. In the interstitial position of the Cl atom the quasistable $\text{Xe}_{6}^+\text{Cl}^{-}$ ionic molecule can be formed. The calculated emission frequency of this molecule is close to the experimental value; however the emission band can not be as wide as in the experiment. It follows that most probably not the $\text{Xe}_{6}^+\text{Cl}^{-}$ but the $\text{Xe}_{2}^+\text{Cl}^{-}$ molecule, formed in the substitutional site, is responsible for the experimentally detected emission.

**Acknowledgments**

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notation hereon. I. L. thanks SUNY-Buffalo for its hospitality during a visit in 1986-87 when this work was performed.
References

Table 1. Angular and Cartesian coordinates of the cage Xe atoms.

\[ R = 3.062 \text{ Å} \]

| \( \theta \) | 90 45 45 90 45 45 90 135 135 90 135 135 |
| \( \phi \) | 45 0 90 135 180 270 315 0 90 225 180 270 |
| \( \chi \) | R R 0 -R -R 0 R R 0 -R -R 0 |
| \( \gamma \) | R 0 R R 0 -R -R 0 R -R 0 -R |
| \( \zeta \) | 0 R R 0 R R 0 -R -R 0 -R |

Table 2. Cartesian coordinates (in Å) of atoms for the \( \text{Xe}^+\text{Cl}^- \) equilibrium configuration. The coordinates of the \( i = 4-12 \) Xe atoms are the same as in Table 1.

\[
\begin{array}{c|cccc}
\text{Xe} & \text{Cl} \setminus i & 1 & 2 & 3 \\
\hline
x & 1.114 & 2.887 & 2.887 & 0 \\
y & -0.083 & 2.670 & 0.329 & 3.132 \\
z & 0.038 & 0.329 & 2.670 & 3.132 \\
R_{\text{XeCl}} & - & 3.30 & 3.30 & 4.68 \\
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\]
Table 3. **Cartesian coordinates (in Å) of the Cl atom at an interstitial position and six neighboring Xe atoms for the ground state.**

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Figure Captions

Fig. 1. Angular Coordinates and the location the three Xe atoms (i = 1,2,3) of the cage.

Fig. 2. Ground state potential energy surfaces of the Cl atom in the Xe cage for the planes passing through the z-axis (the ordinate). The numbers next to the equipotential lines indicate their energy in eV. The pointers indicate the direction to the nearest Xe atom. The planes for the following values of the angle coordinates $\phi$ are presented: a) $\phi = 0^\circ$, b) $\phi = 15^\circ$, c) $\phi = 30^\circ$, d) $\phi = 45^\circ$. For the location of the planes relatively to the Xe atoms see Table 1.

Fig. 3. Average distance $R$ of the Cl atom from the cage center (the left scale) and light absorption cross section $Q$ (the right scale) as functions of temperature.

Fig. 4. Potential curves of all ionic states ($k = 4-39$) along the line running from the cage center to the nearest Xe atom. The distance of the Cl atom from the cage center is plotted against the x-axis.

Fig. 5. Potential curves of the lowest ionic states $k = 4-15$ along arcs with a fixed radius $R = 1.1$ Å. The angle shift of the Cl atom on an arc is plotted against the x-axis.
(a) The arc is located in a plane which includes the cage center and two Xe atoms. The curves start at the point on the line running to one of these atoms and end at the point on the line running between these two Xe atoms.
(b) The arc is located in the $\theta = 90^\circ$ (the xy-plane). The curves start at the point on the x-axis ($\phi = 0^\circ$) which is equidistant to
four Xe atoms and end at the point on the line running to one of these atoms ($\phi = 45^\circ$).

Fig. 6. Potential energy surfaces of the Cl$^-$ ion in the Xe$_{12}^+$ cage for the lowest ionic states $k = 4$ (a) and $k = 5$ (b) in the plane $\phi = 0^\circ$ which is the same as in Fig. 1a for the ground state. The numbers next to the equipotential lines show their energy in eV.

Fig. 7. The energy spectrum of the excitation of the Xe$_{12}$Cl system from neutral ($k = 1-3$) to ionic ($k = 4-39$) states (see the text). The solid line represents the total excitation. The dashed lines represent the excitations to the particular ionic states $k = 4, 5, 20$. The spectrum is given for the temperatures 10 K (a) and 80 K (b).

Fig. 8. The Xe$_2$Cl potential curves of the first ionic state $k = 4$ (the left scale) and the ground state $k = 1$ (the right scale) in the vicinity of the Xe$_2^+$Cl$^-$ ($k = 4$) equilibrium configuration (Table 2) for the fixed molecular plane and fixed Cl atom location:
(a) Potential curves as functions of the Xe-Cl molecular distance.
(b) Potential curves as functions of the Xe-Cl-Xe molecular angle.

Fig. 9. Xe$_2$Cl potential curves of the first ionic state $k = 4$ (the left scale) and the ground state $k = 1$ (the right scale) in the Xe$_2^+$Cl$^-$ ($k = 4$) equilibrium configuration as functions of the molecule translation shift along following mutually perpendicular lines:
(a) $x$-axis ($\theta = 90^\circ$, $\phi = 0^\circ$).
(b) The (+yz) diagonal line ($\theta = 45^\circ$, $\phi = 90^\circ$)
(c) The (-yz) diagonal line ($\theta = 45^\circ$, $\phi = 270^\circ$)
The lines (a) and (b) lie in the system symmetry plane which is perpendicular to the molecular plane; the line (c) lies in the molecular plane.

**Fig. 10.** Emission spectrum of Xe$_2^+\text{Cl}^-$ in solid xenon at 12 K. Original composition, 1:100 HCl:Xe; excitation wavelength, 308 nm.

**Fig. 11.** Excitation spectra of Cl:Xe solids at 12 K. Original composition, 1:100 HCl:Xe. The spectra were recorded with weak-field excitation after extensive irradiation of the solid with a 308 nm laser: (a) spectrum recorded prior to annealing the solid; (b) spectrum recorded after extensive annealing at 55 K.
Fig. 2(c)
Fig. 10

Intensity vs. Wavelength (nm)

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<td>800 N. Quincy Street, Arlington, Virginia 22227-5000</td>
<td>1</td>
<td>Dr. David Young</td>
<td>Code 334, NORDA, NSTL, Mississippi 39529</td>
</tr>
<tr>
<td>1</td>
<td>Dr. Bernard Dauda</td>
<td>Naval Weapons Support Center, Code 50C, Crane, Indiana 47522-5050</td>
<td>1</td>
<td>Naval Weapons Center</td>
<td>Attn: Dr. Ron Atkins, Chemistry Division, China Lake, California 93555</td>
</tr>
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<td>Attn: Dr. R. W. Drisko, Code L52, Port Hueneme, California 93401</td>
<td>1</td>
<td>Scientific Advisor</td>
<td>Commandant of the Marine Corps, Code RD-1, Washington, D.C. 20380</td>
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<td>Building 5, Cameron Station, Alexandria, Virginia 22314</td>
<td>1</td>
<td>U.S. Army Research Office</td>
<td>Attn: CRD-AA-IP, P.O. Box 12211, Research Triangle Park, NC 27709</td>
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<td>DTNSRDC</td>
<td>Attn: Dr. H. Singerman, Applied Chemistry Division, Annapolis, Maryland 21401</td>
<td>1</td>
<td>Mr. John Boyle</td>
<td>Materials Branch, Naval Ship Engineering Center, Philadelphia, Pennsylvania 19112</td>
</tr>
<tr>
<td>1</td>
<td>Dr. William Tolles</td>
<td>Superintendent, Chemistry Division, Code 6100, Naval Research Laboratory, Washington, D.C. 20375-5000</td>
<td>1</td>
<td>Naval Ocean Systems Center</td>
<td>Attn: Dr. S. Yamamoto, Marine Sciences Division, San Diego, California 91232</td>
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<tr>
<td>1</td>
<td>Dr. David L. Nelson</td>
<td>Chemistry Division, Office of Naval Research, 800 North Quincy Street, Arlington, Virginia 22217</td>
<td>1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
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Department of Chemistry
University of California
Berkeley, California 94720

Dr. J. H. Weaver
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Box 5055
Tennessee Technological University
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Troy, New York 12181

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New Haven, CT 06511-8118

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Dr. Theodore E. Madey
Surface Chemistry Section
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National Bureau of Standards
Washington, D.C. 20234

Dr. J. E. Demuth
IBM Corporation
Thomas J. Watson Research Center
P.O. Box 218
Yorktown Heights, New York 10598

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Evanston, Illinois 60637

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Dr. Keith H. Johnson
Department of Metallurgy and Materials Science
Massachusetts Institute of Technology
Cambridge, Massachusetts 02139

Dr. S. Sibener
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James Franck Institute
5640 Ellis Avenue
Chicago, Illinois 60637

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Quantum Surface Dynamics Branch
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General Electric Company
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Department of Chemistry
James Franck Institute
5640 Ellis Avenue
Chicago, Illinois 60637

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R301
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White Oak
Silver Spring, Maryland 20910

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Code 5230
Naval Research Laboratory
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Dr. L. Kesmodel
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Indiana University
Bloomington, Indiana 47403

Dr. K. C. Janda
University of Pittsburg
Chemistry Building
Pittsburg, PA 15260

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Department of Chemistry
University of North Carolina
Chapel Hill, North Carolina 27514

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Bell Laboratories
Murray Hill, New Jersey 07974

Dr. Martin Fleischmann
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Southampton 509 SNH
UNITED KINGDOM

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Rochester, New York 14627

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University of California
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Dr. G.H. Morrison
Department of Chemistry
Cornell University
Ithaca, New York 14853

Dr. J. Baldeschwieler
Department of Chemistry and Chemical Engineering
California Institute of Technology
Pasadena, California 91125