MOLECULAR RESONANCES OBSERVED IN THE PREDISSOCIATION OF Cs₂

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Very cold molecular beam of Cs₂ is generated by a high temperature supersonic jet source. State-specific photofragment yield spectrum is obtained in the orange band of Cs₂. Asymmetric line shapes are observed. The Fano line shape parameter, q, shows a gradual sign change, which is called "q-reversal". A very broad absorption band which has about 50 cm⁻¹ width is observed and explained to occur through the perturbation by the D ¹Σ⁺ state. The complex resonance line shapes are attributed to the interfering resonances.

KEY WORDS: Cesium dimer, Predissociation, Complex resonance, q-reversal

INTRODUCTION

In the last few years, we have seen increasing interest in a new field called 'transition state spectroscopy', in which spectroscopic methods are used to probe directly the transition region of molecules where chemical bonds are about to form or break.¹,² This 'transition-region species', which refers to the short-lived molecular complex in this unstable region, is a quasibound state occurring when discrete states are coupled to a continuum, and can be called 'resonance' in more general term. In the study of molecular reaction dynamics, the resonance can be a very sensitive probe of the reactive scattering processes, providing spectroscopic information on the detailed process of chemical changes.³

The resonance occurring in the predissociation of diatomic molecules is the simplest example of such transition-region species. There are close similarities between the diatomic predissociation and the atomic autoionization, with the fragmented particle being an atom in the former, and an electron in the latter. The resonances in the autoionization have been studied in much more detail than in the predissociation. Fano showed that lineshapes in the autoionization can be described by a simple formula when a discrete state in coupled to a single continuum, as

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where \( \varepsilon = (E - E_r)/(\Gamma/2) \) is the reduced energy, \( \Gamma \) is the width of the resonance, \( E_r \) is the resonance energy, \( \sigma_0 \) corresponds to the dissociative continuum cross section, and \( q \) is the line shape parameter. 4

Among alkali dimers, \( \text{Cs}_2 \) shows the most complex spectrum due to very compact potential energy curves and large spin-orbit couplings. They are, therefore, very suitable to study the predissociation occurring through multichannel interactions. When illuminated by blue or orange light, \( \text{Cs} \) is known to produce excited \( \text{Cs} \) atoms in several different internal states. 5 Detailed spectroscopic studies on the photodissociation of \( \text{Cs}_2 \) have, however, been hampered by the complex absorption spectrum caused by numerous hot bands and its very small rotational constant. We have designed a new high temperature pulsed nozzle and prepared a very cold molecular beam of \( \text{Cs}_2 \), in which all the vibrational hot bands are practically eliminated and the rotational temperature is reduced to about 1 K. 6-12 In this very cold molecular beam the number of partial cross sections for each \( J \), the angular momentum of the initial state, superimposed in the observed dissociation cross section, are greatly reduced. The dramatic cooling in a pulsed molecular beam turns out to be very important in the study of resonance, since direct comparison with theory now becomes possible.

EXPERIMENTAL

Figure 1 shows the experimental setup. \( \text{Cs}_2 \) is produced by expanding \( \text{Cs} \) vapor with \( \text{Kr} \) gas using a high temperature pulsed nozzle. The nozzle is made of a modified fuel injector of an automobile engine (Nippon Denso). An electric pulse of 1 msec duration and 50 V height is applied to open the nozzle. The diameter of the nozzle is 0.8 mm. While the pulsed nozzle set is heated to about 450°C, the electromagnet part is cooled down to room temperature by water. We fill 1 gram of 99.95% \( \text{Cs} \) (Mitsuwa Chemicals) in the sample room of the pulsed nozzle, which lasted for longer than one week when the nozzle was operated continuously at 4 Hz. The most difficult part of the experiment is keeping the nozzle in continuous operation in order to prevent the condensation of the \( \text{Cs} \), which hinders smooth operation of the pulsed beam. The pulsed supersonic jet, in which \( \text{Cs}, \text{Cs}_2 \), and the seeding gas (Kr) were the major species, was collimated by a 1.2 mm diameter skimmer (Beam Dynamics) located 5 cm from the nozzle. The skimmed molecular beam was intersected at right angles 15 cm from the skimmer by the pump and probe laser pulses that were propagating collinearly. A typical pressure was \( 2 \times 10^{-4} \) Torr for the source chamber and \( 4 \times 10^{-7} \) Torr for the detection chamber when the pulsed nozzle was operated at 4 Hz with 600 Torr of Kr backing pressure.

Ions formed by the absorption of two photons from the excitation laser were accelerated by a double electrostatic field time-of-flight lens to about 3500 eV and traveled through a 70 cm long field free region toward a dual microchannel plate detector. The ion signals were then amplified by a fast preamplifier and sent to
boxcar integrators. By collimating the laser beam with a skimmer, we could achieve very good mass selection, with the mass resolution of about 500 at the mass of $^{133}\text{Cs}_2$ (265.8 a.u.). The ion signals were monitored by using two boxcar integrators. By adjusting the time gates of the two boxcars, $\text{Cs}^+$ and $\text{Cs}_2^+$ ion signals were both recorded at the same time. The frequency of the excitation laser was measured simultaneously at the time of the experiment by sending a fraction of the laser beam through an optogalvanic device filled with Ne. The pump and probe laser pulses were provided by two dye lasers pumped by an excimer laser. The probe laser pulse was delayed 5 nanosecond with respect to the pump pulse by optical path length

**Figure 1** Schematic illustration of the molecular beam apparatus with the linear time-of-flight mass spectrometer. DG: delay generator, PND: pulsed nozzle driver, PNV: pulsed nozzle valve, SK: skimmer, MCP: microchannel plate detector, LN: liquid nitrogen, GAS: gas supply system.
difference. The nominal bandwidth of the pump and probe laser pulses are 0.2 cm\(^{-1}\). The pump and probe laser power were reduced to less than 200 \(\mu\)J per pulses by neutral density filters in order to avoid saturation and power broadening.

RESULTS AND DISCUSSION

Figure 2 shows the observed ionization spectra obtained in the supersonic molecular beam of Cs\(_2\). The upper trace is the resonance enhanced two photon ionization spectrum of Cs\(_2\), in which Cs\(^+\) ions are detected. The lower trace is the photofragment yield (PFY) spectrum of Cs\(_2\), in which Cs (6 \(^2\)P\(_{3/2}\)) atom resulted from the photodissociation of Cs\(_2\) is selectivity monitored by a single photon ionization. The ionization laser has an energy of 19720 cm\(^{-1}\), which is larger than the ionization energy for the Cs (6 \(^2\)P\(_{3/2}\)) atom (19674.36 cm\(^{-1}\)). Extra bands observed in the PFY spectrum at frequencies higher than 17100 cm\(^{-1}\) are due to the Cs\(^+\) ions fragmented from the Cs\(^+\) ions generated by two photon ionization and correspond to the vibrational progression of the \(D \, ^1\Sigma^+\) state. When the energy of the ionization laser (probe pulse) is less than the ionization energy (19674.36 cm\(^{-1}\)), all the peaks except those extra bands disappear. The rotational temperature of Cs\(_2\) in the molecular beam in estimated to be about 1.0 K by simulating the rotational contours with the known molecular parameters of the D and X states. The intensity of the hot bands are very weak, and the vibrational temperature is thus estimated to be lower than 5 K.

Figure 3 shows the ionization schemes of these spectra. The first laser prepares Cs\(_2\) molecule in its excited state. The coupling to the repulsive electronic state predissociates the excited Cs\(_2\) molecule. The second laser pulse of frequency \(\nu_2\) ionizes the dissociated Cs (6 \(^2\)P\(_{3/2}\)) atom. When the second laser pulse of frequency \(\nu_2\) is applied to the excited Cs\(_2\) molecule, two electronic processes are in competition. They are the predissociation caused by the coupling of the excited bound state to the continuum and the molecular ionization caused by the absorption of the second photon of the frequency of either \(\nu_1\) or \(\nu_2\). If the predissociation rate is fast, or in the order of picosecond, the molecular ionization process becomes much less effective, making the Cs\(^+\) ion signal too weak to observe. When the predissociation rate is in the order of nanosecond, we observe both the predissociated atomic fragments (Cs\(^+\)) and the Cs\(^+\) ions.

The experimental PFY spectrum in Fig. 2 (lower trace) shows many broad and asymmetric line profiles. At close inspection, the PFY spectrum shows a gradual sign change of the line shape parameter \(q\) passing through zero (near 16900 cm\(^{-1}\)) and through infinity (near 17050 cm\(^{-1}\)). The Beutler-Fano profile remains valid only in the isolated resonance case, i.e., when only a single level of the discrete state is coupled to the continuum. When more than a single discrete state are coupled to the continuum, second and higher order perturbations should be taken into account, and the profile cannot be described by the simple Fano formula. Note that the band near 17050 cm\(^{-1}\) cannot be simulated by a single Beutler-Fano profile and shows clear multichannel quantum interference effect.
Molecular resonances observed in the predissociation of CS$_2$

Figure 2 Ionization spectrum of CS$_2$. The seeding gas is 1 atm. of Kr. a) Upper trace: 2 photon ionization spectrum of CS$_2$. Cs$^+$ ion is monitored. b) Lower trace: photofragment yield (PFY) spectrum of CS$_2$. Cs (6$^2$P$_{3/2}$) atom is ionized and monitored.

The possibility that the asymmetric line shapes are due to rotational structures is excluded based on the following evidences. Firstly, the vibrational progressions show very regular spacing. Rotational bands for diatomic molecules are asymmetric and blue-shaded if the rotational constant of the excited state (B') is larger than that of
the ground state (B''), and red-shaded if B' is smaller than B''. The rotational constant (B'), however, would not change so widely in a region where the potential curve is close to harmonic. Secondly, if the broadening were due to rotational structures, its shape would change when the rotational temperature of the Cs$_2$ in molecular beam is changed. Figure 4 shows the ionization spectrum of the Cs$^+_2$ (upper trace) and Cs$^+$ ions (lower trace) with the coexpanding gas changed to Ar and backing pressure reduced to 100 Torr. Under these expansion conditions, the rotational temperature becomes higher than 5 K. In the spectra of Fig. 4, however, the line shapes of the broad resonances of the Cs$^+$ signal remain about the same. This observation shows that the broad and asymmetric line shapes are not due to rotational structures. Lastly, the band at 17050 cm$^{-1}$ shows a dip in the center, which cannot be explained by a rotational structure.

Figure 3  The ionization scheme of the spectra shown in Fig. 2.
MOLECULAR RESONANCES OBSERVED IN THE PREDISSOCIATION OF CS$_2$

Figure 4 Ionization spectra of Cs$_2$ obtained under the same conditions as in Fig. 2, except that the seeding gas is changed to 100 torr of Ar. The line shape of the broad resonances are not changed.

Figure 5 shows the potential energy diagram of Cs$_2$ in the interested frequency range. The potential curves are from the ab initio calculations by Spiess and Meyer obtained by the pseudo-potential method. The spectroscopic data on these electronic states have been fragmentary. Only the low vibrational levels of the C $^1\Pi_u$ (up to $v' = 13$), and (2) $^3\Pi_u$ states (up to $v' = 6$) were observed previously. Complex perturbations are observed for higher vibrational levels of the C $^1\Pi_u$ state. The $\Omega = 0$ component of the (2) $^3\Pi_u$ state is predicted to perturb the D $^1\Sigma_u^+$ state at frequencies higher than 17500 cm$^{-1}$, which was confirmed recently.

Among these three electronic states, only the C $^1\Pi_u$ state has oscillator strength associated with the ground state, which has a $^1\Sigma_g^+$ symmetry. Since the equilibrium bond lengths of the C $^1\Pi_u$ and (2) $^3\Pi_u$ states are very close to that of the ground state, the high vibrational levels of the C $^1\Pi_u$ and (2) $^3\Pi_u$ states have very small Franck-Condon overlap with the $\nu'' = 0$ level of the ground state at the observed frequency range. Only the (2) $^3\Sigma_u^+$ state, which is a continuum state, has a good Franck-Condon overlap with the ground state ($\nu'' = 0$) at the observed frequency range, and carries most of the oscillator strength, which is borrowed from the low vibrational levels ($\nu = 0$–6) of the C $^1\Pi_u$ state. The resonances occurring in this system are then described as a result of configuration interactions between dark discrete states near the resonance energy, a dark continuum, and remote bright discrete states.
The PFY spectrum in Fig. 2 shows a couple of very narrow resonance lines (at 17097 and 17020 cm\(^{-1}\)). The narrow resonances with a very long lifetime have been observed in the autoionization of Ba atom and that of Li atom in strong magnetic field.\(^{16,17}\) They are interpreted as occurring when several resonances associated with different channels interfere strongly and the coupling matrix elements to the open channel are accidentally canceled out. This produces a metastable state with an anomalously long lifetime, which is referred to as a 'bound state in a continuum'.
(BIC). The close-coupling calculations predict the existence of several very narrow resonances. Since this resonance is borrowing the oscillator strength from the continuum of \( (2) \ 3\Sigma_u^+ \), which borrows the oscillator strength from the low vibrational levels of the \( C \ 1\Pi_u \) state in turn, the radiative lifetime is much longer than that of the low vibrational levels of the \( C \ 1\Pi_u \) state and would be of the same order as the predissociation lifetime.

Compared to some broad resonances as wide as about 5 cm\(^{-1}\) (fwhm) and thus about 600 fs of lifetime, these narrow resonances are expected to have lifetimes in the order of nanoseconds, which is more than about \( 10^4 \) times enhancement of the lifetime. While the BIC’s in the autoionization can be used as possible storage states of vacuum-UV energy above the ionization threshold, those observed in the predissociation can be used as good intermediate states for double resonance experiment, since they are mixtures of different electronic states with long lifetimes.

According to the Fano’s multi-level formula for the configuration interaction, the absorption cross section vanishes once in each of successive resonances. The experimental dissociation cross section (lower trace in Fig. 2), however, does not show such zero cross sections. Although sharp dips may disappear by the overlap of the rotational bands and the laser bandwidth, broad resonances should not be affected by these broadening effects. This lack of zero cross section can be ascribed to the existence of a noninteracting, or weakly interacting dissociation channel.

Figure 6 shows a vertically expanded ionization spectrum of Cs\(^+\). It is evident that Cs\(_2\) has a continuous absorption band near 17050 cm\(^{-1}\) with a band width of 50 cm\(^{-1}\) (fwhm). The most likely candidate for the origin of this broad band is the absorption by the \( (2) \ 3\Sigma_u^+ \) state. The \( D \ 1\Sigma_u^+ \) state has no direct coupling to the continuum \([ (2) \ 3\Sigma_u^+ ] \) in the first order approximation. The \( D \ 1\Sigma_u^+ \) state is, however, coupled to the \( \Omega = 0 \) component of the \( (2) \ 3\Pi_u \) state through the spin-orbit coupling, which, in turn, interacts with the \( \Omega = 1 \) component of the \( (2) \ 3\Sigma_u^+ \) state through L-uncoupling. Thus the nominal \( (2) \ 3\Sigma_u^+ (\Omega = 1) \) state has some character of the \( D \ 1\Sigma_u^+ \) state, from which it borrows additional oscillator strength. Since the \( (2) \ 3\Sigma_u^+ (\Omega = 1) \), \( (2) \ 3\Pi_u (\Omega = 1) \), and \( C \ 1\Pi_u \) states are not directly coupled to the \( D \ 1\Sigma_u^+ \) and \( (2) \ 3\Pi_u (\Omega = 0) \) states in the first order, this additional dissociation cross section can be incoherently added to the dissociation cross section contributed by the coupling between the states with \( \Omega = 1 \). This then explains the apparent lack of the zero cross section in the experimental PFY spectrum. At frequencies higher than 17200 cm\(^{-1}\), the \( (3) \ 3\Sigma_u^+ \) state and its nonadiabatic coupling with the \( (2) \ 3\Sigma_u^+ \) state need to be included in the consideration of the dissociation cross section.

We showed that interference can occur between the continuum and discrete state as long as the continuum has oscillator strength from the initial state, even when the discrete state does not have the oscillator strength. This is because the discrete state is modified by an admixture of continuum, the contribution of which to the transition dipole matrix element \([ (\psi_i l T l \psi_f) ] \) interferes with those of the continuum, as explained by Fano’s configuration interaction theory.
Figure 6 Expanded part of the two photon ionization spectrum of Cs₂ shown in Fig. 2. Cs⁺ ion is monitored. A broad absorption band is seen centered at 17050 cm⁻¹.

The spectrum in Fig. 2 shows that the line shape parameter changes sign at least twice. This kind of the gradual sign change of q is called a ‘q-reversal’. While several examples of the q-reversals have been observed in autoionization, they are interpreted as occurring through a structured continuum which is perturbed by another discrete state from the third channel. Thus, the q-reversal in autoionization is considered as a result of 3-channel interaction. On the other hand, we showed that in predissociation the q-reversal can occur when only 2 channels are involved. This can be ascribed to the property of the dissociating continuum. While the coupling of the ionization continuum to discrete state shows monotonous variation in autoionization, that of the continuum of nuclear dissociation shows oscillatory variations in the case of outer limb crossing. It is likely that such oscillatory variations have caused q-reversal with only 2 channels involved.

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References

Integration of nanotechnology into the treatment of various diseases such as cancers represents a mainstream in the current and future research due to the limitations of traditional clinical diagnosis and therapy. The early detection of cancer has been universally accepted to be essential for treatment. However, it remains challenging to detect tumors at an early stage. For instance, the goal of molecular imaging in breast cancer is to diagnose the tumor with approximately 100–1000 cells, compared to the traditional techniques which may require more than a million cells for accurate clinical diagnosis. On another hand, anticancer drugs are designed to simply kill cancer cells, and their entrance into healthy organs or tissues is undesirable due to the severe side effects. In addition, the rapid and widespread distribution of anticancer drugs into nontargeted organs and tissues requires a lot of drugs with high cost. These difficulties have largely limited the successful therapy of cancer.

Nanomaterials are anticipated to revolutionize the cancer diagnosis and therapy. The development of multifunctional polymeric nanoparticles allows for the early detection of cancers. The construction of intelligent polymeric nanosystems can be used as controlled delivery vehicles to improve the therapy efficiency of anticancer drugs, that is, such vehicles are capable of delivering drugs to predetermined locations and then releasing them with preprogrammed rates in response to the changes of environmental conditions such as pH and temperature. Besides polymers, these nanomaterials can also be composed of supraparamagnetic iron oxide, carbon nanotube, metallic nanoshell, core-shell aggregate, or composites. These nanomaterials represent new directions for more effective drug administration in cancer.

This special issue of the Journal of Nanomaterials will cover a wide range of nanomaterials for cancer diagnosis and therapy. It will mainly focus on the preparations, characterizations, functionalizations, and properties of nanoparticles, nanostructured coatings, films, membranes, nanoporous materials, nanocomposites, and biomedical devices. Fundamental understanding of the basic mechanisms on material and biological processes related to the unique nanoscale properties of the materials will be the highlights of this special issue.

Papers are solicited in, but not limited to, the following areas:

- Synthesis and functionalization of polymer nanoparticle/nanomicelle/nanocomplex
- Polymer nanoparticle/hydrogel for drug delivery
- Synthesis of intelligent nanogel
- Hydrogel in nanoscale sensing
- Supraparamagnetic nanoparticle for magnetic resonance imaging applications
- Carbon nanotube-based devices for drug delivery
- Core-shell nanoparticle for molecular imaging
- Metallic nanoshell for drug delivery
- Nanoporous and nanoscaled materials for drug delivery
- Nanotechnologies for targeted delivery
- Controlled drug-delivery nanovehicles
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Special Issue on
Solar Energy and Nanomaterials for Clean Energy Development

Call for Papers

This special issue on solar energy and nanomaterials for clean energy development is comprised of selected, full-length versions of papers presented at the International Conference on Powering a Greener Future: Nanomaterials for solar energy conversion (solar’09). The highly successful solar conference series, started in 1991, provide a place where researchers interested in fundamental and applied aspects of photochemistry can meet and inspire one another. The solar conference recognizes that successful photochemical applications go hand in hand with advancement of fundamental understanding of photoinduced processes and excited states.

This conference was held in Luxor, Egypt, during the period 10–14 January 2009. It aimed to discuss in an interdisciplinary sense the newest trends, break-through, discoveries, as well as applications of nanomaterials in photochemistry, photocatalysis, solar energy conversion, and environmental aspects.

Topics of interest include but are not limited to:

- Nanomaterials for photovoltaics and 3rd-generation DSSCs, batteries, fuel cells, ceramics coating, solar energy conversion devices, polymers in water desalination and RO membrane production, OLED, photonics and photoluminescent materials
- Molecular/nano-photochemistry, photosciences and applications
- Environmental remediation: photocatalysis (air, soil, as well as water treatment and disinfection)
- Photosynthesis and artificial light-harvesting systems
- Photocatalytic hydrogen production and storage

Submission to this special issue is opened to participants of the International Conference on Powering a Greener Future: Nanomaterials for solar energy conversion (solar’09). Other contributions from interested scientists worldwide are also welcomed.

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Special Issue on
Polymer Nanocomposite Processing, Characterization, and Applications

Call for Papers

Polymers reinforced with nanoparticles, such as carbon nanotubes, are of great interest due to their remarkable mechanical, thermal, chemical properties as well as optical, electronic, and magnetic applications. In the general research area of polymer nanocomposites, a number of critical issues need to be addressed before the full potential of polymer nanocomposites can actually be realized. While a number of advances have recently been made in the area of polymer nanocomposites, the studies on understanding of the effects of processing parameters on the structure, morphology, and functional properties of polymer nanocomposites are deficient. There is a need for characterization techniques to quantify the concentration and distributions of nanoparticles as well as to assess the strength at the interface between the polymer and nanoparticles. Also, there is a need for the development of better models able to predict the mechanical properties of the polymer nanocomposites as functions of myriad factors including nanoparticle orientation, the type of functional groups, and the molecular weight of polymer chain. The relationships between the structural distributions and the ultimate properties of the polymer nanocomposites also need to be elucidated.

This special issue of the Journal of Nanomaterials will be devoted to emerging polymer nanocomposite processing techniques and call for new contributions in the field of characterization and applications of multifunctional nanocomposites. It intends to cover the entire range of basic and applied materials research focusing on rheological characterization, nanoparticle dispersion, and functional properties of polymer nanocomposites for sensors, actuators, and other applications. Fundamental understanding of the effects of processing and nanoparticles on the polymer structure and morphology, their optical, electrical, and mechanical properties as well as novel functions and applications of nanocomposite materials will be the highlights of this special issue.

Papers are solicited in, but not limited to, the following areas:

- Solution and melt processing of polymer nanocomposites
- Rheological and thermal characterization of nanocomposites
- Generation of nanofibers using extrusion and electrospinning of nanocomposites
- Processing-induced orientation of nanoparticles
- Quantification of nanoparticle dispersion
- Effect of nanoparticle incorporation on polymerization
- In situ nanoparticle formation in polymer matrix
- Noncovalent functionalization techniques and characterization of properties at polymer-nanoparticle interface
- Novel applications of polymer nanocomposites

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