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ATOMS, MOLECULES AND CLUSTERS IN MOTION

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Bound and unbound rovibrational states of the methane-argon dimer

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ABSTRACT

Peculiarities of the intermolecular rovibrational quantum dynamics of the methane-argon complex are studied using a new, *ab initio* potential energy surface [Y.N. Kalugina, S.E. Lokshtanov, V.N. Cherepanov, and A.A. Vigasin, J. Chem. Phys. **144**, 054304 (2016)], variational rovibrational computations, and detailed symmetry considerations within the molecular symmetry group of this floppy complex as well as within the point groups corresponding to the local minimum structures. The computed (ro)vibrational states up to and beyond the dissociation asymptote are characterised using two limiting models: the rigidly rotating molecule's model and the coupled-rotor model of the rigidly rotating methane and an argon atom orbiting around it.

j^M R fixed

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Large-amplitude motion; rovibrational computation; coupled quantum rotors; methane-argon

1. Introduction

Methane is an abundant molecule not only on Earth, but in the Universe [1,2]. On Earth it has the third largest contribution to the greenhouse effect, after water vapour and carbon-dioxide, meanwhile on Titan, the moon of Saturn it is the most important greenhouse gas [3]. Therefore, the interaction of methane with other simple molecules is of interest for astrochemistry and for atmospheric chemistry. There are several experimental and theoretical results for such systems including methanewater, $CH_4 \cdot H_2O$ [4,5], methane-fluoride, $CH_4 \cdot F^-$, [6], and the mehane dimer, $CH_4 \cdot CH_4$ [7].

The first measurements for the methane-argon gas mixture at several temperatures were reported by Dore and Filabozzi [8]. The infrared spectrum was reported first by McKellar *et al.* in 1994 [9], then Pak *et al.* measured the high-resolution spectra of the CH_4 ·Ar and CH_4 ·Kr complexes in the 7 μ m region. Later on, further high-resolution measurements and *ab initio* computations were performed for the CH_4 ·Ar system [2,10].

In spite of repeated attempts, the spectral lines within the bound region of the complex have remained elusive to experimentalists due to the very low induced dipole moment of the complex dominated by dispersion interactions (both isolated methane and isolated argon are apolar). This region is however well accessible to theoretical investigations due to the low dimensionality (3D) of the intermolecular degrees of freedom and the excellent rigid-monomer approximation. On the contrary, the experimentally well-accessible transitions to predissociative states of the complex are extremely challenging for theory due to the high-dimensionality (12D)

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of the six-atomic methane-argon system. A rigorous fulldimensional rovibrational study has not been completed, and full-dimensional, *ab initio* potential energy surface have not been developed yet. At the same time, the energy and lifetime of predissociative states carry ample information on the fine details of the inter- and intramolecular quantum dynamics taking place under the influence of weak (here: dispersion) interactions. Due to the very weakly bound nature of this complex and the high-order symmetry of the methane molecule, it has been anticipated that kinetic couplings play an important role in its intermolecular and predissociative dynamics.

In the present work we take the opportunity to use a new, *ab initio* atom-molecule interaction potential energy surface of this dimer [1] to study peculiarities of the rovibrational dynamics and related symmetry considerations.

2. Computational details

a. Intermolecular potential energy surface. We use the three-dimensional potential energy surface (PES) determined by Kalugina et al. by ab initio computations at the CCSD(T)/CBS level of theory, and by fitting an analytic functional form to the data points [1]. The ab initio computations were carried out with a regular tetrahedral methane structure with $r_{\rm CH} = 2.06735$ bohr. There are two non-equivalent minima on this PES: the argon atom is in face configuration at the global minimum (GM) structure corresponding to an R = 6.95 bohr carbon-argon distance, and in edge configuration with a larger, R = 7.34 bohr value at the secondary minimum (SM) structure. The saddle point connecting the two non-equivalent minima is located at the vertex position of the argon atom (Figure 1). The dissociation energy (D_e) of the complex from the global minimum is 141.47 cm⁻¹ and it is 115.06 cm⁻¹ from the secondary minimum.

b. Rovibrational computations. We carried out rovibrational computation on this intermolecular PES using the GENIUSH programme [11,12]. GENIUSH stands for GENeral Internal-coordinate, USer-define Hamiltonians, and it has been successfully used for the computation of bound and resonance states of floppy molecular systems, including $ArNO^+$ [13], HeH_2^+ [14], NH_3 [11,12], H_5^+ [15], CH_5^+ [16], $CH_4 \cdot H_2O$ [4,5], with various internal coordinate and frame definitions. An interesting feature, the existence of formally negative-energy rotational excitations [4,5,15,17–19] had been observed in a growing number of floppy systems and led to the introduction of the notion of astructural molecules in Ref. [15]. Ref. [15] also presented a one-dimensional torsional model to better highlight the phenomenon. Ref. [4]

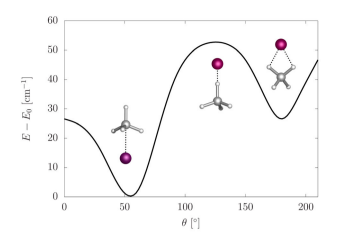


Figure 1. Angular dependence of the potential energy surface of Ref. [1] represented in terms of (R, θ, ϕ) spherical polar coordinates. For each θ value in the plot the *R* coordinate was chosen to minimise the interaction energy, while ϕ was set to 0.0°.

reported an elaborate study on the quantum dynamical background of these special features for the example of the methane-water dimer using a quantum mechanical, coupled-rotor model of the subsystems.

The general rovibrational Hamiltonian implemented in GENIUSH is

$$\begin{aligned} \hat{H} &= \frac{1}{2} \sum_{kl=1}^{D} \tilde{g}^{-1/4} \hat{p}_k G_{kl} \tilde{g}^{1/2} \hat{p}_l \tilde{g}^{-1/4} \\ &+ \frac{1}{2} \sum_{k}^{D} \sum_{\alpha}^{3} \left(\hat{p}_k G_{k,D+\alpha} + G_{k,D+\alpha} \hat{p}_k \right) \hat{J}_{\alpha} \\ &+ \frac{1}{2} \sum_{\alpha}^{3} G_{D+\alpha,D+\alpha} \hat{J}_{\alpha}^2 \\ &+ \frac{1}{2} \sum_{\alpha}^{3} \sum_{\beta > \alpha}^{3} G_{D+\alpha,D+\beta} \left(\hat{J}_{\alpha} \hat{J}_{\beta} + \hat{J}_{\beta} \hat{J}_{\alpha} \right) + V, \quad (1) \end{aligned}$$

where $\hat{p}_k = -i\partial/\partial q_k$ is a differential operator corresponding to the q_k internal coordinate, and \hat{J}_{α} ($\alpha = x, y, z$) labels the body-fixed angular momentum operators. The quantities $G_{kl} = (\mathbf{g}^{-1})_{kl}$ and $\tilde{g} = \det(\mathbf{g})$ are obtained from the g_{kl} mass-weighted metric tensor, defined as

$$g_{kl} = \sum_{i=1}^{N} m_i \mathbf{t}_{ik}^{\mathrm{T}} \mathbf{t}_{il}, \quad k, l \in \{1, 2, \dots, D+3\}$$
 (2)

$$\mathbf{t}_{ik} = \frac{\partial \mathbf{r}_i}{\partial q_k},\tag{3}$$

$$\mathbf{t}_{D+\alpha} = \mathbf{e}_{\alpha} \times \mathbf{r}_i,\tag{4}$$

Table 1. Internal coordinate, basis function, number of DVR points (*N*), and grid intervals used in the computations.

Coordinate	Basis function	Ν	Grid interval
R [Å]	Laguerre	51	[2.65–15]
$\cos \theta$	Legendre	81	Unscaled
ϕ [°]	Fourier	21	[0–360)

which are evaluated over a direct product grid in the programme using the user-defined body-fixed Cartesian coordinates in terms of the internal coordinates.

Geometrical constraints (in the present case for the methane molecule) are accounted for in an automated fashion by choosing D being equal to the number of active vibrational dimensions (D=3 for the present case) in Equation (1). This choice is equivalent to fixing the structural parameter (and a corresponding zero velocity) in the classical Lagrangian [11]. The Hamiltonian matrix is obtained as the matrix representation of Equation (1) constructed with the discrete variable representation (DVR) for the vibrational degrees of freedom, and using the Wang functions for rotations. The kinetic energy operator is evaluated numerically over a direct product DVR grid. The eigenvalues and eigenfunctions of the Hamiltonian matrix are computed with an iterative Lanczos eigensolver.

We define the body-fixed Cartesian coordinates for the methane-argon complex, as follows: we place the carbon atom in the origin, two hydrogen atoms are in the *y*-*z* plane and the other two are in the *x*-*y* plane with the same methane geometry as the PES [1]. The position of the argon atom is defined by the *R*, $\cos \theta$, and ϕ spherical polar coordinates, which are the active vibrational coordinates (for the basis functions, detailed intervals and DVR parameters see Table 1). The coordinate definition is completed by shifting the system to the centre of mass of the complex. We used atomic masses [20], $m_{\rm H} = 1.007$ 825 032 23 u, $m_{\rm D} =$ 2.014 101 778 12 u, $m_{\rm T} = 3.016$ 049 277 9 u, $m_{\rm C} =$ 12 u, and $m_{\rm Ar} = 39.962$ 383 123 7 u, throughout the computations.

3. Symmetry considerations and analysis tools

This section is dedicated to a general description of the symmetry analysis and symmetry properties of the limiting (rigid-rotor and coupled-rotor) models, which will be used to understand the numerical results in Section 4.

3.1. Point groups and the molecular symmetry group of the methane-argon dimer

The PES of Ref. [1] features two non-equivalent minima. The global minimum has a C_{3v} point-group (PG)

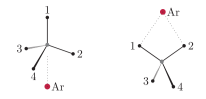


Figure 2. The global minimum has C_{3v} point-group symmetry, whereas the local minimum has C_{2v} point-group symmetry (this particular numbering of the nuclei is used in the derivation in the text).

symmetry, while the secondary minimum has C_{2v} PG symmetry (Figures 1 and 2). With the low barriers and the almost freely rotating methane unit in the dimer, there is an interchange among the four GM and the six SM structures that gives rise to a rich splitting pattern.

The methane-argon dimer belongs to the $T_d(M)$ molecular symmetry (MS) group and all symmetry species of the MS group are physically existing [21].

3.2. General considerations about the tunnelling splitting of vibrational states

The splitting of vibrational states, classified by PG irreducible representation (irrep) labels, will be derived by starting out from a simple picture. Let us imagine that the system exhibits only small (ε) amplitude vibrations about the minimum structures (of one of the PG symmetries). This system is almost 'frozen' at the near proximity of these equilibrium structures, and thus its molecular symmetry group is the permutation-inversion group isomorphic to the PG (with a selected numbering of the nuclei). Hence, these ε -vibrations can be characterised by irreps of the PG, C_{3v} (PG) or C_{2v} (PG), but they can also be characterised by the C_{3v} (M) and C_{2v} (M) *permutationinversion* (PI) groups. These PI groups share elements with the MS group (here $T_d(M)$), which will be used later.

For a general derivation of the vibrational splittings due to the large-amplitude motions, we proceed as follows. First, we construct the symmetry-adapted basis functions for the irreps of the PI group (isomorphic with the PG) of the minimum structure. Then, we consider the transformation properties of these functions under the symmetry operations of the MS group, determine the characters for each class of the MS group, and reduce the representation. We carry out the derivation for a general pair of MS and PI (\sim PG) groups, labelled with *G* and *g*, respectively. At the end of the derivation, we calculate the splittings for the case of methane-argon.

(1) Let us consider a function f_0 localised at an arbitrarily selected version of the molecule.

- (2) Map the f_0 basis function onto all feasible versions, $f_i = \hat{O}_i f_0$, by using all $\hat{O}_i \ i \in \{1, 2, ..., N\}$ MS group operations (N = |G| is the order of G).
- (3) It is always possible to choose f_0 in such way that $\langle f_i | f_j \rangle = \delta_{ij} \ (i, j = 1, ..., N).$
- (4) Construct a linear combination of the f_i (i = 1,...,N) functions which transform according to the Γ irrep of g:

$$\phi_i^{(\Gamma)} = \frac{1}{n} \sum_{k=1}^n c_k^{(\Gamma)} \hat{P}_k f_i, \quad i = 1, \dots, N$$
 (5)

where $\hat{P}_k \in g$ and n = |g| is the order of g.

(5) In order to determine the characters in *G*, the trace of the matrix representation of Ô_R ∈ *G* is calculated corresponding to the {φ^(Γ)_i; 1 = 1,..., N} set of functions:

$$\chi^{(\Gamma)}(\hat{O}_{R}) = \sum_{i=1}^{N} \langle \phi_{i}^{(\Gamma)} | \hat{O}_{R} | \phi_{i}^{(\Gamma)} \rangle$$
$$= \frac{1}{n^{2}} \sum_{i=1}^{N} \sum_{k=1}^{n} \sum_{l=1}^{n} c_{k}^{(\Gamma)} c_{l}^{(\Gamma)*} \langle \hat{P}_{k} f_{i} | \hat{O}_{R} | \hat{P}_{l} f_{i} \rangle$$
(6)

(6) In order to evaluate the trace in Equation (6), we first identify the non-vanishing integrals by reordering the operators as

$$\langle \hat{P}_k f_i | \hat{O}_R | \hat{P}_l f_i \rangle = \langle f_i | \hat{P}_k^{-1} \hat{O}_R \hat{P}_l | f_i \rangle.$$
(7)

Since $g \subset G$ and due to the closure property of a group, $\hat{P}_k^{-1}\hat{O}_R\hat{P}_l \in G$. Since the set of (localized) f_i functions was created by mapping f_0 with all possible operations of G, and the functions are orthogonal, a non-vanishing integral is obtained only if the product of the three operators, \hat{P}_k^{-1} , \hat{O}_R , and \hat{P}_l equal the identity:

$$\hat{P}_k^{-1}\hat{O}_R\hat{P}_l = \hat{I} \tag{8}$$

$$\hat{O}_R = \hat{P}_k \hat{P}_l^{-1}.$$
(9)

(7) The integral is non-vanishing for a selected $\hat{O}_R \in G$, if $\exists \hat{P}_r \in g: \hat{P}_r = \hat{O}_R$:

$$\sum_{k=1}^{n} \sum_{l=1}^{n} \langle f_{i} | \hat{P}_{k}^{-1} \hat{O}_{R} \hat{P}_{l} | f_{i} \rangle = n.$$
 (10)

Then, if Γ is the totally symmetric irrep (assume that it is labelled with A₁) of *g*, all $c_k^{(A_1)}$ characters equal

1, and we obtain:

$$\chi^{\Gamma}(\hat{O}_{R}) = \frac{1}{n^{2}} \sum_{i=1}^{N} \sum_{k=1}^{n} \sum_{l=1}^{n} c_{k}^{A_{1}} c_{l}^{A_{1}} \langle \hat{P}_{k} f_{l} | \hat{O}_{R} | \hat{P}_{l} f_{l} \rangle$$
(11)

$$=\begin{cases} \frac{1}{n^2}Nn = \frac{N}{n}, & \text{if } \exists \ \hat{P}_r \in g : \hat{P}_r = \hat{O}_R \\ 0, & \text{otherwise} \end{cases}$$
(12)

For a different labelling of the atoms the \hat{O}_R operators in the same class have different characters, thus we average over the $\chi(\hat{O}_R)$ values within a class of the group. (Equivalently, we could have considered all possible labelling of the molecule at the beginning.) Let us denote the class of \hat{O}_R as \mathcal{R} and $K_{\mathcal{R}}$ is the number of elements in this class of G. Then, we obtain

$$\chi^{\Gamma}(\mathcal{R}) = \frac{1}{K_{\mathcal{R}}} \sum_{\hat{O}_R \in \mathcal{R}} \chi(\hat{O}_R)$$
(13)

$$=\begin{cases} \frac{k_r}{K_{\mathcal{R}}} \frac{N}{n}, & \text{if } \exists \hat{P}_r \in g : \hat{P}_r = \hat{O}_R \in \mathcal{R} \\ 0, & \text{otherwise,} \end{cases}$$
(14)

and k_r is the number of operations in the \mathcal{R} class of G for which $\exists \hat{P}_r \in g : \hat{P}_r = \hat{O}_R$. It can be seen that this k_r number of operations equals the number of elements in the class of g which contains \hat{P}_r .

(8) In general for a Γ irrep of *g*, the characters for the \mathcal{R} class of *G* are

$$\chi^{(\Gamma)}(\mathcal{R}) = \begin{cases} c_r^{(\Gamma)} \frac{k_r}{K_{\mathcal{R}}} \frac{N}{n}, & \text{if } \exists \hat{P}_r \in g: \\ & \hat{P}_r = \hat{O}_R \in \mathcal{R} \\ 0, & \text{otherwise.} \end{cases}$$
(15)

3.3. Mathematical structure of the tunnelling splittings in methane-argon

First, we calculate the tunnelling splitting of the vibrational states for the global minimum structure. The order of $T_d(M)$ is N = 24. The global minimum (GM) has C_{3v} PG symmetry, and the order of C_{3v} (PG or PI) is n = 6. The equivalent operations and the orders of classes in the two groups are collected in Table 2.

Table 2. Equivalent operations in the $T_d(M)$ and $C_{3v}(M)$ symmetry groups for the labelling shown in Figure 2.

$T_{\rm d}({\rm M})$	Ε	(123)	(14)(23)	[(1423)]*	[(23)]*
K _R	1	8	3	6	6
$C_{3v}(M)$	Ε	(123)			[(23)]*
k _r	1	2			3

Table 3. Equivalent operations in the $T_d(M)$ and $C_{2v}(M)$ symmetry groups for the labelling shown in Figure 2.

$T_{\rm d}({\rm M})$	Ε	(123)	(12)(34)	[(1234)]*	[(12)]*
K _R	1	8	3	6	6
$C_{3v}(M)$	Ε		(12)(34)		[(12)]*
k _r	1		1		1

The $T_d(M)$ irreps for the vibrational splittings of states corresponding to the global minimum and the C_{3v} PG (and its isomorphic PI) group are obtained from Equation (15) and Table 2:

$$\Gamma_{A_1}^{C_{3v}} = A_1 \oplus F_2$$

$$\Gamma_{A_2}^{C_{3v}} = A_2 \oplus F_1$$

$$\Gamma_E^{C_{3v}} = E \oplus F_1 \oplus F_2,$$
(16)

which defines the mathematical structure of the splitting pattern for the vibrational states assignable to the GM.

We proceed similarly for the secondary minimum, use Equation (15) and Table 3 for the C_{2v} group (of order n = 4), and obtain the splitting pattern for the vibrational states assignable to the SM as:

$$\begin{split} \Gamma_{A_1}^{C_{2v}} &= A_1 \oplus E \oplus F_2 \\ \Gamma_{A_2}^{C_{2v}} &= A_2 \oplus E \oplus F_1 \\ \Gamma_{B_1}^{C_{2v}} &= F_1 \oplus F_2 \\ \Gamma_{B_2}^{C_{2v}} &= F_1 \oplus F_2. \end{split} \tag{17}$$

A difference in the mathematical structure of the GM and SM splittings is that the totally symmetric vibrations (such as the zero-point vibration) of the SM contain a doubly-degenerate species.

3.4. Coupled-rotor decomposition

Molecular systems which can be separated into two rigidly rotating distinct parts can be characterised by a model which includes only the coupling of the subsystems' angular momenta (and neglects any PES interactions). This scheme was introduced by Sarka *et al.* in relation with the general, numerical kineticenergy operator approach of GENIUSH and is called the coupled-rotor decomposition (CRD) [5].

The coupled-rotor (CR) ansatz for methane-argon (Figure 3) is constructed from the rotational wave functions of methane, $\phi_{km}^{j^{M}}$, and of the effective diatomic rotor, Y_{Λ}^{m} , coupled to a total rotational angular momentum

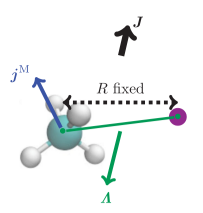


Figure 3. Coupling of the subsystems' angular momenta used to define the coupled-rotor model in CH₄.Ar.

Table 4. Characters and irrep decomposition of the J = 0 rotational functions of the methane-argon dimer in the $T_d(M)$ molecular symmetry group. n_{cl} is the number of operations in the class.

n _{cl}	Ε	(123)	(14)(23)	[(1423)]*	[(23)]*	
Г	1	8	3	6	6	Irreps
[0,0] ₀₀	1	1	1	1	1	A ₁
[1, 1] ₀₀	3	0	1	-1	1	F ₂
[2, 2] ₀₀	5	-1	1	-1	1	$E \oplus F_2$
[3, 3] ₀₀	7	1	-1	1	1	$A_1 \oplus F_1 \oplus F_2$
[4, 4] ₀₀	9	0	1	1	1	$A_1 \oplus E \oplus F_1 \oplus F_2$
[5, 5] ₀₀	11	-1	-1	-1	1	$E \oplus F_1 \oplus F_2$
[6,6] ₀₀	13	1	1	-1	1	$A_1 \oplus A_2 \oplus E \oplus F_1 \oplus 2 F_2$

state, $|JM\rangle$. Hence, the CR functions are defined as

$$[j^{M}, \Lambda]_{JM} = \sum_{\mu=-j^{M}}^{j^{M}} \sum_{M-\mu=-\Lambda}^{\Lambda} \langle j^{M}, \mu, \Lambda, (M-\mu) | JM \rangle \phi_{\mu}^{j^{M}} Y_{\Lambda}^{M-\mu},$$
(18)

where $\langle j^{\rm M}, \mu, \Lambda, (M - \mu) | JM \rangle$ are the Clebsch–Gordan coefficients. Based on this definition, the symmetry properties of the rotational functions, and the equivalent (monomer) rotations of the MS group operators, closed analytic expression can be derived for the symmetry labels of the CR functions. The characters and irrep decomposition in $T_{\rm d}({\rm M})$ are given in Tables 4 and 5 for the J = 0 and J = 1 CR functions of methane-argon.

In order to measure the similarity of the CR model and the 'exact' intermolecular wave functions, we compute CRD overlaps defined as [5]

$$\operatorname{CRD}_{nm}^{J} = \sum_{r=1}^{N_{R}} \left| \sum_{k=-J}^{J} \sum_{i=1}^{N_{\Omega}} \tilde{\Psi}_{nk}^{J}(\rho_{r},\omega_{i}) \tilde{\varphi}_{mk}^{J}(\omega_{i}) \right|^{2}.$$
 (19)

where $\tilde{\Psi}_{nk}^{J}(\rho_r, \omega_i)$ is the exact variational (ro)vibrational wave function in DVR, ρ and ω denote the radial and (the collection of) angular coordinates respectively, and

Table 5. Characters and irrep decomposition of the J = 1 rotational functions of the methane-argon dimer in the $T_d(M)$ molecular symmetry group.

n _{cl}	Ε	(123)	(14)(23)	[(1423)]*	[(23)]*	
Г	1	8	3	6	6	Irreps
[0, 1] ₁₀	1	1	1	-1	-1	A ₂
[1,0] ₁₀	3	0	-1	1	-1	F ₁
$[1, 1]_{10}$	3	0	-1	-1	1	F ₂
[1, 2] ₁₀	3	0	-1	1	-1	F ₁
[2, 1] ₁₀	5	-1	1	1	-1	$E \oplus F_1$
[2, 2] ₁₀	5	-1	1	-1	1	$E \oplus F_2$
[2, 3] ₁₀	5	-1	1	1	-1	$E \oplus F_1$
[3, 2] ₁₀	7	1	-1	-1	-1	$A_2\oplusF_1\oplusF_2$
[3, 3] ₁₀	7	1	-1	1	1	$A_1 \oplus F_1 \oplus F_2$
[3, 4] ₁₀	7	1	-1	-1	-1	$A_2 \oplus F_1 \oplus F_2$
[4, 3] ₁₀	9	0	1	-1	-1	$A_2 \oplus E \oplus F_1 \oplus F_2$
[4, 4] ₁₀	9	0	1	1	1	$A_1 \oplus E \oplus F_1 \oplus F_2$
[4, 5] ₁₀	9	0	1	-1	-1	$A_2 \oplus E \oplus F_1 \oplus F_2$
[5, 4] ₁₀	11	-1	-1	1	-1	$E\oplus 2F_1\oplusF_2$
[5, 5] ₁₀	11	-1	-1	-1	1	$E \oplus F_1 \oplus 2 F_2$

 $\tilde{\varphi}_{mk}^{J}(\omega_{i})$ are the CR functions depending only on the angular coordinates. By identifying the dominant values in the CRD matrix the assignment of the (ro)vibrational states is straightforward.

In order to be able to numerically evaluate this integral in a 'black-box fashion' of the GENIUSH protocol [11], we compute the CR functions in a reduceddimensionality GENIUSH computation with the atommolecule distance fixed at its equilibrium value, the PES switched off, and the same angular grid representation as for the 'full' problem (Table 1). The $T_d(M)$ irrep labels are assigned to the (ro)vibrational states in an almost automated fashion by inspecting the numerical CRD matrix and using the symmetry properties of the CR functions (see Tables 4–5).

3.5. Rigid-rotor decomposition

In the traditional picture of molecular rotations and vibrations, molecular rotations [21,22] are superimposed on the (higher-energy) molecular vibrations. The rigid-rotor decomposition (RRD) scheme was introduced [23] to assign parent vibrational states to rovibrational wave functions. The RRD assignment is based on the overlap of a direct product basis of pure vibrational functions, $\psi_n^{J=0}$ and the $\phi^{\text{RR},J}$ rigid-rotor functions with the full rovibrational wave function, $\psi_n^{J>0}$ (computed with a selected body-fixed frame):

$$|\psi_n^{J=0}\rangle \otimes |J,\phi_l^{\rm RR}\rangle = |\psi_n^{J=0} \cdot \phi_l^{\rm RR,J}\rangle \tag{20}$$

$$S_{nl,m} = \langle \psi_n^{J=0} \cdot \phi_l^{\text{RR},J} | \psi_m^{J>0} \rangle.$$
 (21)

By computing the $S_{nl,m}$ overlap values, the dominant contribution of a vibrational state to a rovibrational state can be determined.

Table 6. Characters and irrep decomposition of the $\phi^{\text{RR},J}$ rigid-rotor functions in the $T_d(M)$ MS group.

n _{cl}	Ε	(123)	(14)(23)	[(1423)]*	[(23)]*	
J	1	8	3	6	6	Irreps
0	1	1	1	1	1	A ₁
1	3	0	-1	1	-1	F ₁
2	5	-1	1	-1	1	$E \oplus F_2$
3	7	1	-1	-1	-1	$A_2 \oplus F_1 \oplus F_2$
4	9	0	1	1	1	$A_1 \oplus E \oplus F_1 \oplus F_2$

In the present work, besides the analysis of the numerical RRD values, we would like to better understand the regularities in the RRD matrix introduced by symmetry relations. First of all, we determine the symmetry assignment of the $\phi^{\text{RR},J}$ functions in the $T_d(M)$ MS group. The equivalent rotations $(\alpha_0, \beta_0, \gamma_0)$ for each class in the $T_d(M)$ are (123): $(0, 0, 2\pi/3)$; (14)(23): $(\pi/3, \pi, 0)$; $[(1423)]^*: (\pi/6, \pi/2, -\pi/6); [(23)]^*: (0, \pi, 0)$. The characters of this reducible representation are the traces of Wigner *D* matrices.

$$\chi^{j} = \sum_{m=-j}^{j} D^{j}_{m,m}(\alpha_{0}, \beta_{0}, \gamma_{0})$$
(22)

The irrep decompositions for several *J* quantum numbers are collected in Table 6.

Since the symmetry of the RRD basis functions, Equation (20), is determined by the direct product of the $\Gamma(\psi^{J=0})$ and $\Gamma(\phi^{\text{RR},J>0})$ representations, the overlap in Equation (21) can only be non-zero if the direct product contains the irrep of the rovibrational state. The irrep decomposition in T_d (M) of the direct product of the $\psi_n^{J=0}$ vibrational (A₁, A₂, E, F₁, F₂) and $\phi^{\text{RR},J>0}$, which is F₁ for J = 1:

$$A_1 \otimes F_1 = F_1$$

$$A_2 \otimes F_1 = F_2$$

$$E \otimes F_1 = F_1 \oplus F_2$$

$$F_1 \otimes F_1 = A_1 \oplus E \oplus F_1 \oplus F_2$$

$$F_2 \otimes F_1 = A_2 \oplus E \oplus F_1 \oplus F_2.$$
(23)

This result tells us, for example, that a rovibrational state of A_2 symmetry can only originate from an F_2 vibrational state.

4. Numerical results

4.1. Vibrational states

The depth of the PES valley of the global (secondary) minimum measured from the dissociation asymptote is $-D_e(GM) = -141.47 \text{ cm}^{-1} (-D_e(SM) =$ $-115.06 \text{ cm}^{-1})$. By considering also the zero-point vibration (ZPV) of the GM, the lowest-energy state

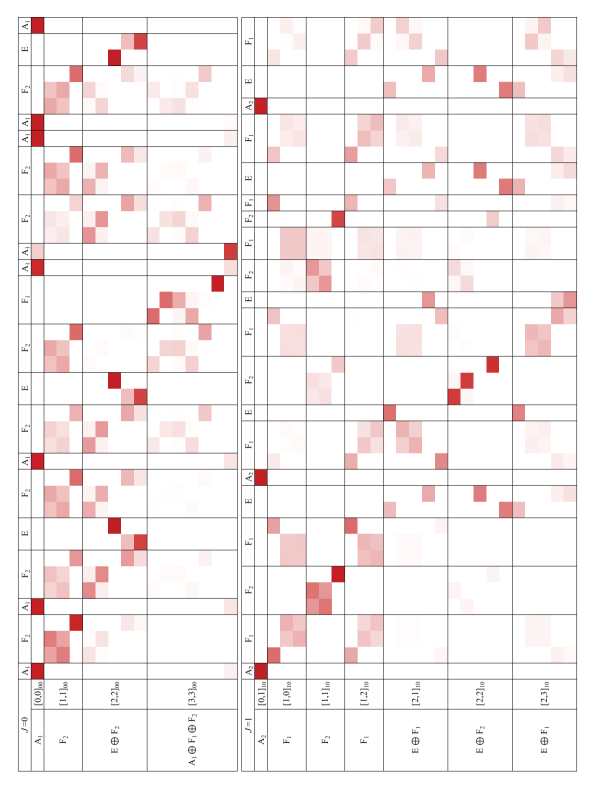


Figure 4. CRD overlap coefficients, Equation (19), computed for the vibrational (J = 0) and rovibrational (J = 1) states of CH₄·Ar (a darker colour corresponds to a larger value of the overlap). The rows correspond to the coupled-rotor functions φ_n^J , which are used to assign the full intermolecular vibrational states, Ψ_n^J , listed in the columns.

accessible to the system has an energy $-D_0 = -D_e + ZPVE = -88.60 \text{ cm}^{-1}$, measured from the dissociation asymptote. The PES bounds 20 vibrational states (if we count degenerate states only once), which have

been assigned using the limiting models described in Section 3. The computed energy levels, together with their CRD and T_d (M) irrep assignments, are collected in Table 7 (see also Figure 4).

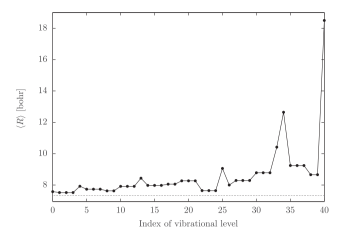


Figure 5. Expectation value of the carbon-argon distance, R, for the bound vibrational states of CH₄·Ar.

First of all, by looking at the computed and assigned list of vibrational states, we have attempted to identify complete splitting patterns, which (would) correspond to the traditional picture of molecular vibrations split up by feasible interchanges between the permutation-inversion versions. Having in mind the mathematical structure of the splittings for GM and SM vibrations, Equations (16) and (17), we have inspected the expectation value of the atom-molecule separation over the bound vibrational states (Figure 5). Due to the structural differences between GM and SM ($R_{eq}(GM) = 6.95$ bohr and $R_{eq}(SM) = 7.34$, bohr respectively), an increased atommolecule separation can be an indication of a secondaryminimum character. Alternatively, an increased $\langle R \rangle$ value can also indicate vibrational excitation along the atom-molecule distance. Stretching vibrational excitations and overtones can be recognised by nodes in the wave function along the R coordinate. Figure 6 collects cuts of A1-symmetry wave functions which highlight their nodal structure along R (the Supplementary Material contains similar cuts for all bound vibrational states).

In addition, we repeated the vibrational computation also for the two symmetrically substituted isotopologues, CD_4 ·Ar and CT_4 ·Ar (Figure 7). According to chemicalphysical intuition, we expected that the tunnelling splittings should get smaller by increasing the mass of the nuclei, which then could help the identification of the vibrational splittings.

Based on all these considerations and the numerical results, the zero-point vibration of the global minimum, ZPV(GM), could be assigned to J0.0 (0.00 cm^{-1} , A₁) and J0.1–3 (9.01 cm⁻¹, F₂). The next six states, J0.4 (28.74 cm⁻¹, A₁), J0.5–7 (31.24 cm⁻¹, F₂), and J0.8–9 (32.08 cm⁻¹, E) could be assigned to the zero-point vibration of the secondary minimum, ZPV(SM), but due to a node along the *R* coordinate it would be morea appropriate to assign J0.4 (28.74 cm^{-1} , A_1) and J0.5–7 (31.24 cm^{-1} , F_2) to the intermolecular vibrational fundamental of the global minimum, Stre(GM). In general, the overlap of the energy range of the ZPV(SM) and Stre(GM) (and other vibrations), and the appearance of the same symmetry species in the different splittings, Equations (16) and (17), an unambiguous identification of vibrational states (in the traditional sense, split up by feasible exchanges) is hardly possible.

Furthermore, the higher the energy, node counting along *R* is less obvious (Figure 6). While, we can trace (the A₁ species of) the first, the second, and perhaps also the third stretching excitations, beyond J0.25 (68.90 cm⁻¹, A₁), the wave functions along *R* become more and more oscillatory indicating that we approach the dissociation limit.

It is interesting to note that there is not any bound vibrational state of A_2 symmetry, which would be an 'indicator' of a vibrational state of A_2 (GM) or an A_2 (SM) PG symmetry, Equations (16) and (17). Table 4 shows that the lowest-energy CR-functions, which generate an A_2 -symmetry state belong to $[6, 6]_{00}$. These CR functions have $\sim 210 \text{ cm}^{-1}$ energy, much larger than the dissociation energy, which explains the absence of bound states of this symmetry species.

While it is difficult to use the traditional picture of molecular vibrations, the CR model appears to be useful and we could identify dominant CR functions for all vibrational states of methane-argon (see Figure 4 and Table 7).

Upon isotopic substitution, we could not really observe the closure of the splittings (since there are not any clear splittings apart from ZPV(GM)), but we observed another regularity with respect to the increase of the hydrogenic mass. Based on this regularity, we will explain the change in the energy ordering of the symmetry species along the CH_4 ·Ar- CD_4 ·Ar- CT_4 ·Ar sequence (Figure 7).

The rotational constant, and thus the rotational energy of methane is inversely proportional to the mass of the H (D/T) atom, and thus the energy of the $[j^{M}, \Lambda]_{00}$ CR functions is also proportional to $m_{\rm H}^{-1}$ to a good approximation (we neglect the energetic contribution from the diatom rotation, which is small). If we check the energy separation of the lowest-energy vibrational states dominated by the $[0,0]_{00}$ (A₁), $[1,1]_{00}$ (F₂), $[2,2]_{00}$ (E \oplus F₂), and $[3,3]_{00}$ (A₁ \oplus F₁ \oplus F₂) CR functions, we observe a ca. 1/2 and 1/3 reduction in this separation (indicated with double-headed arrows in Figure 7) upon the H–D and H–T replacement, respectively. Of course, the changes in

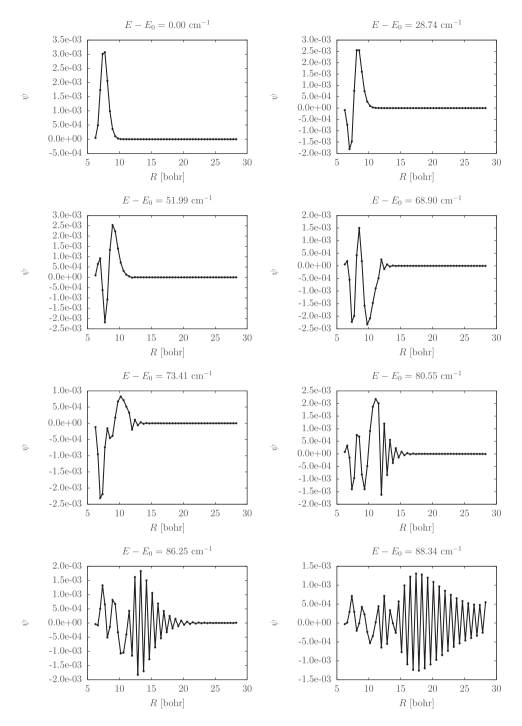


Figure 6. Wave function cuts along the *R* coordinate for the A₁-symmetry vibrational states (the energy shown above each plot is measured from the ground vibrational state).

the vibrational energy intervals only approximately follow the CR model, because the PES also plays a role in the vibrational dynamics.

In comparison with the $m_{\rm H}^{-1}$ relation of the CR splittings, the energy of the atom-molecule stretching excitations (excitation along *R*) changes approximately with $1/\sqrt{\mu}$, the square root of the reduced mass of the

effective diatom (methane and argon), which brings in a less than a $1/\sqrt{2}$ and $1/\sqrt{3}$ factor upon the H-D and H-T replacements, respectively.

Hence, the energy separation of vibrational states corresponding to angular (CR) excitations (with similar radial parts) is decreased by a larger extent than the energy separation of radial (*R*-stretching) excitations

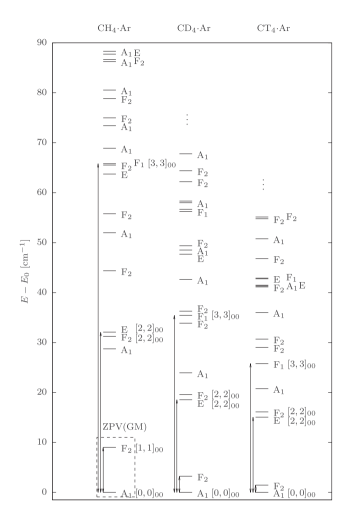


Figure 7. Comparison of the bound-state vibrational energies of CH_4 ·Ar, CD_4 ·Ar, and CT_4 ·Ar.

upon isotopic substitution. Since the radial excitation does not change the symmetry, while the angular (CR) excitation does, we observe a rearrangement of the energy ordering of the various symmetry species in the deuterated and tritiated isotopologues.

4.2. Rovibrational states

The computed J = 1 rovibrational results are summarised in Table 8. The corresponding CRD matrix is shown in Figure 4, while the RRD coefficients (obtained with the RR functions corresponding to the symmetric-top GM structure) are visualised in Figure 8. A clear assignment, of the vibrational parent (with dominant overlaps) is possible only for the lower-energy range. Nevertheless, we find it remarkable that it is possible to use the RR picture at all for this very floppy complex, in the light of the difficulties of identifying complete vibrational splitting manifolds (see previous section).

Surprisingly, the very first J=1 rotationally excited state, J1.0 (0.12 cm⁻¹, A₂), has a lower energy than

Table 7. Bound-state vibrational energies given with respect to the 52.86 cm⁻¹ zero-point vibrational energy of CH₄ · Ar obtained with GENIUSH and the PES of Ref. [1]. Each vibrational state is characterised with its $T_d(M)$ irrep label (Γ) and the dominant (and minor) CR function.

			Coupled-rotor states $(J = 0)$		
n	<i>E</i> [cm ⁻¹]	Г	dominant	minor	
J0.0	52.86	A ₁	[0,0] ₀₀		
J0.1–3	9.01	F ₂	[1,1]00		
J0.4	28.74	A ₁	[0,0]00		
J0.5–7	31.24	F ₂	[2,2]00	[1,1]00	
J0.8–9	32.08	E	[2,2] ₀₀		
J0.10–12	44.37	F_2	[1,1] ₀₀	[2,2] ₀₀	
J0.13	51.98	A ₁	[0,0] ₀₀		
J0.14–16	55.78	F ₂	[2,2]00	[1,1]00 & [3,3]00	
J0.17–18	63.73	Ē	[2,2]00		
J0.19–21	65.53	F ₂	[1,1]00	[3,3] ₀₀	
J0.22–24	65.81	F ₁	[3,3] ₀₀		
J0.25	68.90	A ₁	[0,0] ₀₀	[3,3] ₀₀	
J0.26	73.41	A ₁	[3,3] ₀₀	[0,0] ₀₀	
J0.27–29	74.97	F_2	[2,2] ₀₀	[1,1] ₀₀ & [3,3] ₀₀	
J0.30-32	78.86	F_2	[1,1]00	[2,2]00	
J0.33	80.55	A ₁	[0,0]00		
J0.34	86.25	A ₁	[0,0]00		
J0.35–37	86.70	F ₂	[1,1]00	[2,2] ₀₀ & [1,1] ₀₀	
J0.38–39	87.74	Ē	[2,2]00		
J0.40	88.33	A ₁	[0,0] ₀₀		

its vibrational parent state (9.01 cm⁻¹, F₂) highlighted in Figure 9. The rotational excitation of the ZPV(GM) (0.00 cm^{-1} , A₁) appears only as the second rotational state (9.07 cm⁻¹, F₁). This energy ordering of the rotational excitation of vibrational states is surprising if we have the traditional picture of rotating-vibrating molecules in mind. The traditional picture fails if the energy scale of rotations and intermolecular vibrations overlap which is the case for the present system.

Similarly to the vibrational states, the CR picture comes useful. The dominant CR functions in the lowest energy rotational state and in its parent vibration are the $[0, 1]_{10}$ and $[1, 1]_{00}$ functions, respectively. From this CR assignment, it can be seen that the unusual energy ordering is due to the rotational excitation of the methane unit which increases the energy but decreases the total rotational angular momentum, in the present case, due to its coupling with the rotating diatom. The opposite situation is observed for the second-lowest rotational state and its parent vibration, in which the $[1,0]_{10}$ and $[0,0]_{00}$ are the dominant CR functions, respectively. In this case, the rotational de-excitation of the methane molecule decreases the energy and (in this case) also the total rotational angular momentum is reduced. The symmetry pattern of the states follows the formal regularities derived for the RR functions, Equation (23) (see also Figure 8).

					pled-rotor tes $(J = 1)$
n	$E[cm^{-1}]$	Vib. parent	Г	dominant	minor
J1.0	0.12	F ₂ [9.01]	A ₂	[0,1] ₁₀	
J1.1–3	9.07	A ₁ [0.00]	$\overline{F_1}$	[1,0] ₁₀	[1,2] ₁₀
J1.4–6	9.91	F ₂ [9.01] & E [32.08]	F ₂	[1,1] ₁₀	.,
J1.7–9	9.98	A ₁ [0.00]	F_1	[1,2] ₁₀	[1,0] ₁₀
J1.10–11	22.57	F ₂ [9.01] & F ₂ [31.24]	É	[2,2] ₁₀	[2,1] ₁₀ & [2,3] ₁₀
J1.12	28.86	F ₂ [31.24] & F ₂ [44.37]	A ₂	[0,1] ₁₀	2, 10, 2, 10
J1.13–15	31.28	F ₂ [9.01]	F ₁	[2,1] ₁₀	[1,2] ₁₀
J1.16–17	31.93	F ₂ [9.01]	É	[2,1] ₁₀	[2,3] ₁₀
J1.18-20	31.94	E [32.08] & F ₂ [31.24]	F ₂	[2,2] ₁₀	$[1,1]_{10}$
J1.21–23	32.01	A ₁ [28.74]	F_1	[2,3] ₁₀	[1,0] ₁₀ & [2,1] ₁₀
J1.24–26	40.45	F ₂ [44.37]	F ₂	[1,1] ₁₀	[2,2] ₁₀
J1.27–29	40.46	A ₁ [28.74]	F ₁	[1,0] ₁₀	$[1,2]_{10}$
J1.30–31	42.82	F_2 [31.24]	E	[2,2] ₁₀	[2,1] ₁₀ & [2,3] ₁₀
J1.32–34	44.48	E [63.73]	- F ₁	[1,2] ₁₀	[2,3] ₁₀ & [1,0] ₁₀
J1.35	52.08	F ₂ [55.78] & F ₂ [65.53]	A ₂	[0,1] ₁₀	[2/0110 00 [1/0110
J1.36–37	55.69	F_{2} [44.37]	E	[2,2] ₁₀	$[2,1]_{10}$
J1.38–40	55.84	F_2 [31.24]	- F ₁	[2,3] ₁₀	[1,2] ₁₀ & [2,1] ₁₀
J1.41–43	58.32	F_2 [55.78]	F_2	[1,1] ₁₀	[2,2] ₁₀
J1.44–46	58.34	F_1 [51.98]	F ₁	[1,0] ₁₀	[3,2] ₁₀ & [1,2] ₁₀
J1.47–48	63.58	F ₂ [44.37]	E	[2,3] ₁₀	[2,1] ₁₀
J1.49–51	65.55	12[11.37]	F ₁	[1,2] ₁₀	[3,2] ₁₀
J1.52	65.78	F ₂ [31.24] & F ₂ [44.37]	A ₂	[3,2] ₁₀	[3,4] ₁₀
J1.53	65.83	F_1 [65.81]	A ₁	[3,3] ₁₀	[37,1][0
J1.54–56	65.91	F_2 [65.53] & F_1 [65.81]	F ₂	[1,1] ₁₀	[3,2] ₁₀
J1.57–59	66.03	A ₁ [51.98]	F ₁	[1,0] ₁₀	[3,4] ₁₀
J1.60–62	66.08	F_2 [65.53]	F ₂	[3,4] ₁₀	[1,1] ₁₀
J1.63	68.99	F ₂ [78.86]	A ₂	[0,1] ₁₀	[1,1]0
J1.64–65	71.49	F ₂ [55.78] & F ₂ [74.97]	E	[2,2] ₁₀	[2,3] ₁₀ & [2,1] ₁₀
J1.66	73.58	F_2 [44.37]	A ₂	[3,4] ₁₀	[3,2] ₁₀ & [0,1] ₁₀
J1.67–69	74.33	12[11.57]	F ₁	[3,3] ₁₀	$[3,2]_{10} \& [2,1]_{10}$
J1.70–72	74.42		F ₂	[3,2] ₁₀	$[3,3]_{10} \& [3,4]_{10}$
J1.73–75	75.12		F ₁	[2,3] ₁₀	[3,4] ₁₀
J1.76–78	78.95	A ₁ [68.90]	F ₁	[1,2] ₁₀	[1,0] ₁₀ & [2,3] ₁₀
J1.79–81	79.54	F_2 [78.86]	F_2	[1,2]10 [1,1] ₁₀	[1,0]]0 & [2,0]]0
J1.82–84	79.55	A ₁ [68.90]	F ₁	[1,0] ₁₀	[1,2] ₁₀
J1.85	80.60	A] [00.90]		[0,1] ₁₀	[1,2]10
J1.86–87	81.61	Several F ₂ states	A ₂ E	[0,1] ₁₀ [2,2] ₁₀	[2,1] ₁₀ & [2,3] ₁₀
J1.88	86.26	Several F2 states	Е А ₂	[2,2] ₁₀ [0,1] ₁₀	[2,1]10 & [2,3]10
J1.88–91	86.69		F ₁	[0,1] ₁₀ [1,2] ₁₀	[2,1] ₁₀ & [1,0] ₁₀
J1.92–94		F ₂ [86.70] & E [87.74]			
J1.92–94 J1.95–97	87.31		F ₂	[2,2] ₁₀	[1,1] ₁₀
J1.98–99	87.34 87.57	A ₁ [80.55]	F1 E	[2,3] ₁₀	[2,1] ₁₀ & [1,0] ₁₀
		F ₂ [65.53]		[2,3] ₁₀	[2,1] ₁₀
J1.100	88.31		A ₂	[0,1] ₁₀	

Table 8. Bound-state rovibrational energies (J = 1), measured from the zero-point vibration energy, of CH₄·Ar obtained with GENIUSH and the PES of Ref. [1]. The assigned vibrational parent state, $T_d(M)$ irrep label (Γ), and the dominant (minor) coupled-rotor functions are also shown for each rovibrational state.

4.3. Resonance states

Twenty vibrational bound states (not counting degeneracies) have been identified up to the dissociation asymptote (Table 7), and we continued the computations beyond the dissociation limit. Repeated computations with slightly different grid intervals of the molecule-atom distance allowed us to identify long-lived resonance states using the stabilisation method [24] and a histogram analysis technique [13]. Twenty GENIUSH computations were carried out, and 1000 eigenvalues were computed in each run with different box sizes. In every computation the maximal value for the *R* coordinate was slightly modified, from 14.05 Å to 15 Å (the PES is defined up to

R = 15 Å), incremented by 0.05 Å in each computation. The computed eigenvalues are collected and sorted from the lowest to the largest, then a histogram is made from the data. The eigenvalues corresponding to the resonance states will appear in every computation, meanwhile the continuous eigenvalues are shifted by changing the size of the box. The obtained histogram (bin size 0.01 cm⁻¹) is shown in Figure 10 and the energy positions and assignment for the peaks are collected in Table 9.

It is interesting to note that we find only E and F₂symmetry states among the identified long-lived resonances. The dominant CR functions in these states are $[1, 1]_{00}$, $[2, 2]_{00}$, and $[3, 3]_{00}$. The lack of A₁ states may be

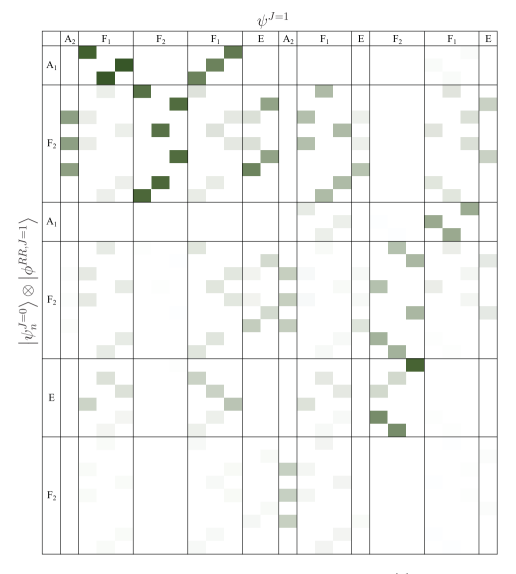


Figure 8. RRD overlap matrix, Equation (21), for the J = 1 rovibrational states of CH₄·Ar. The $\psi_n^{J=1}$ rovibrational functions listed in the columns are assigned with the product of the vibrational and rigid-rotor functions defined in Equation (20).

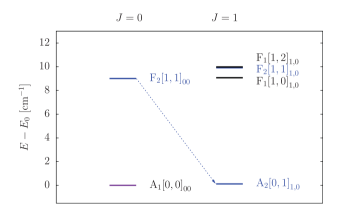


Figure 9. The lowest-energy J = 1 rovibrational level of CH₄·Ar (0.12 cm⁻¹, A₂) has lower energy than its vibrational parent state, (9.01 cm⁻¹, F₂). The unusual energy ordering can be understood by inspecting the methane's rotational excitation in the dominant CR functions, [0, 1]_{1,0} and [1, 1]_{0,0}, respectively.

explained by the fact that highly excited vibrations along the R (dissociation) coordinate also have A₁ symmetry, and due to the coupling of states with the same symmetry, we do not observe any long-lived resonance of this species.

The absence of states dominated by the $[0,0]_{00}$ CR function tells us that a fair portion of the energy in these long-lived resonances is carried by the angular degrees of freedom (the excitation of which is measured by the CRD). The presumably slow energy transfer from the angular degrees of freedom to the radial (dissociative) degree of freedom gives rise to a long lifetime of the identified states.

It is also insightful to compare the energy of the identified resonance states with the dissociation energy $D_0(\text{GM}) \approx 90 \text{ cm}^{-1}$ (up to this value we can increase the energy in the dissociative degree of freedom) and

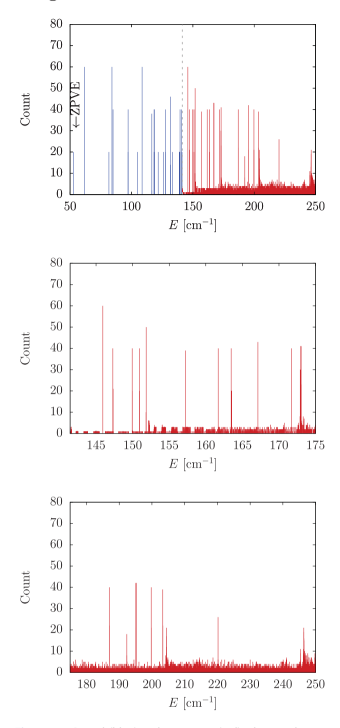


Figure 10. Bound (blue) and resonance (red) vibrational states of CH₄·Ar indicated by the peaks in the stabilisation histogram. measured from the global minimum of the PES. (For some triply degenerate states an artificial split can be observed in the histogram due to imperfect convergence. An unambiguous symmetry assignment is established through a CRD analysis (see Table 9).)

the CR energies of the $[n, n]_{00}$ functions with n = 1, 2, and 3 with ca. 10 cm^{-1} , 30 cm^{-1} , and 60 cm^{-1} , respectively. Assuming simple additivity of these energies, we may understand the occurrence of long-lived states dominated by $[1, 1]_{00}$ to ca. 100 cm^{-1} (however, we do not

Table 9. Energy of the long-lived (J = 0) vibrational resonances of CH₄·Ar computed with GENIUSH, the PES of Ref. [1], and the stabilisation technique. The vibrational states are given with respect to the zero-point vibration energy. Each state is characterised with its $T_d(M)$ irrep label and the dominant (and minor) coupled-rotor functions.

			Coupled-rotor states $(J = 0)$		
n	<i>E</i> [cm ⁻¹]	Г	dominant	minor	
J0.41–43	93.06	F_2	[1,1] ₀₀		
J0.44-46	94.44	F_2	[3,3] ₀₀	[2,2] ₀₀	
J0.47–49	97.08	F_2	[1,1] ₀₀	[2,2]00	
J0.50-51	98.06	F_2	[1,1] ₀₀	[2,2]00	
J0.52–54	98.98	F_2	[1,1]00		
J0.55–56	104.36	Е	[2,2] ₀₀		
J0.57–58	108.82	E	[3,3] ₀₀		
J0.59–61	110.60	F_2	[3,3] ₀₀	[2,2] ₀₀	
J0.62–63	114.25	E	[2,2] ₀₀		
J0.64–65	118.82	E	[2,2] ₀₀		
J0.66-67	120.03	E	[2,2] ₀₀		
J0.68–69	134.08	E	[2,2] ₀₀	[3,3] ₀₀	
J0.70–71	139.46	Е	[3,3]00	00	
J0.72–74	142.34	F ₂	[1,1]00	[2,2] ₀₀	
J0.75–77	150.45	F_2	[3,3] ₀₀	[2,2]00	

have any simple explanation for the J0.73–75, F_2 state), as well as occurrence of long-lived states dominated by $[2, 2]_{00}$ or $[3, 3]_{00}$ up to ca. 130–140 cm⁻¹.

The symmetry assignment by CRD and the major and minor coupled rotor contributions are listed in Table 9.

5. Summary

Rovibrational computations have been carried out for the methane-argon dimer using a new, *ab initio* atommolecule potential energy surface [1] and the general rovibrational programme, GENIUSH [11,12]. This PES, with a well depth of $-D_e = -141.47 \text{ cm}^{-1}$, supports twenty bound states (counting degenerate states only once). The energy of the lowest accessible vibrational state to the system is $-D_0(GM) = -88.60 \text{ cm}^{-1}$, measured from the dissociation asymptote. Although there are two different minimum structures, with a 26.41 cm⁻¹ energy separation, most of the computed (ro)vibraitonal states cannot be unambiguously assigned either to the global or to the secondary minimum. The computed rovibrational states were analyzed using two limiting models and detailed symmetry considerations.

The traditional picture of molecular rotations and vibrations almost completely breaks down for this weakly bound, floppy system. We could identify the complete tunnelling splitting only for the zero-point vibrational state of the global minimum. Vibrational parent states could be determined only for the lowest few rotational states with J=1 rotational quantum number. Interestingly, the lowest-energy rotational state does not correspond to the rotational excitation of lowest-energy

vibrational state but it corresponds to the second lowest vibrational state. This rotational state and its vibrational parent represent a 'formally' negative-energy rotational excitation, i.e. the rotationally excited state has a lower energy than its parent vibration.

These unusual features, which appear strange in terms of the traditional picture of rotating-vibrating molecules, can be well understood using a model which considers the rotating-vibrating complex in terms of coupled quantum rotors of rigidly rotating methane and the effective diatom (atom-molecule separation). All vibrational bound and low-energy, long-lived resonance states as well as most of the J=1 rovibrational states can be assigned within this coupled-rotor picture.

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Disclosure statement

No potential conflict of interest was reported by the authors.

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