

Electronic excitation and structural relaxation of the adenine dinucleotide in gas phase and solution†

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Introduction

In the last few years, several experiments studying the photo-physics of DNA fragments showed a strong difference between the excited state behaviour of nucleobase stacks as opposed to isolated nucleobases or mononucleotides. Whereas all isolated nucleobases decay on a picosecond timescale,^{1–3} additional transients with a lifetime of 10–100 ps and even nanosecond components were found in single and double stranded DNA oligonucleotides.^{4–9} Several possible reasons have been put forward to explain this intriguing situation. As one option, processes were considered where the excitation was strictly localized on one DNA base and it was conjectured from quantum chemical computations that the DNA environment could have a significant influence in this case.¹⁰ However, in a dynamical study no significant increase in lifetime due to the stacking interaction was found.¹¹ Secondly, hydrogen bonding was considered as a major factor. In this sense a proton-coupled electron transfer between Watson–Crick (WC) paired bases was considered as a possible decay channel.^{12–14} Furthermore, dynamics simulations emphasized the role of structural effects related to hydrogen bonding.^{11,15,16} Thirdly, stacking interaction was considered as the main influence on the collective excited state behaviour in DNA. The importance of stacking

was highlighted by the observation that single and double stranded DNA oligonucleotides showed very similar transients after photoabsorption, highlighting the relative unimportance of the second strand.⁴ Experimental signatures for two types of processes due to stacking interactions were found pertaining to (i) electron dynamics leading to excitation energy transfer,^{8,17,18} and (ii) structural relaxation leading to a trapping of the defects.^{4,19} The nature of the trapping sites in the second case, which remains a major open question, will be the main focus of this study. These sites were initially assumed to be charge transfer exciplexes.^{19,20} But also computational evidence for neutral or mixed exciplexes was given.^{21,22} Aside from their structure, also the fate of such exciplexes is unknown. There was experimental evidence for a decay through charge recombination¹⁹ and computational evidence for deactivation mediated by a shortening of the intermolecular distance,²³ but also a restoration of localized states has been considered.²⁴ In light of these unknowns, further studies are needed to fully understand this important phenomenon.

Having recently analysed the effects of base–base stacking on the absorbing states in DNA,²⁵ it is the aim of this paper to consider structural relaxation processes. It has been observed by experiment that long-lived transients are already present in the ribodinucleotides (diribonucleoside monophosphates) and, in particular, that the adenine dinucleotide (ApA, see Fig. 1) shows identical bleach recovery signals as compared to longer adenine strands.¹⁹ Moreover, it was pointed out that after UV excitation in adenine strands the excitation is localized on not more than two bases after 1 ps and that there is not even conclusive experimental evidence for invoking delocalized absorbing states.²⁶ Furthermore, there is recent computational evidence that the absorbing states are rather localized.^{25,27} Specifically, the RNA (rather than DNA)

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†Electronic supplementary information (ESI) available: Cartesian coordinates, selected intermolecular distances, total energies and depictions of the molecular structures used in this work (S1–S7), extended analysis of the gas phase potential curve (S8), and NTOs at the S1 minima (S9–S11). See DOI: 10.1039/c3pp50032b



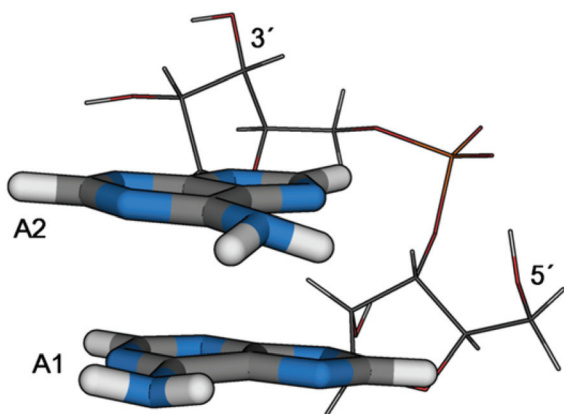


Fig. 1 Depiction of the adenine dinucleotide (as optimized at the MP2/SV(P)-SV level in solution). The labelling of the two units (A1, A2) and the position of the 5'- and 3'-ends of the backbone are indicated.

dinucleotide ApA was chosen in order to be able to match a detailed experimental investigation of this system.¹⁹ However, we believe that due to similar stacking geometries the presented results should also be relevant to DNA analogues. In light of all these considerations we conclude that ApA is a highly interesting model system possessing many parallels to larger RNA and DNA oligonucleotides.

When studying structural relaxation and excited state dynamics in DNA fragments, special challenges are present related to (i) the system size, (ii) a treatment of the environment, in particular for describing charge transfer states, (iii) providing a reliable description even for strongly distorted structures, (iv) giving an accurate interaction potential, especially for exciplexes with short intermolecular separations, and (v) an instructive analysis of the nature of the excited states. A number of computational strategies with different merits and shortcomings in light of these demands are, in principle, applicable and a wide range of types of calculations have been performed. These amount to wavefunction based *ab initio* calculations,²¹ time-dependent density functional theory (TDDFT) using a polarizable continuum model,^{22,28,29} semi-empirical methods,^{23,30} and model Hamiltonians.³¹ For more information about calculations performed on DNA fragments see also ref. 25 and 32. This study is based on *ab initio* theory in an attempt to eliminate uncertainties related to unusual molecular structures, which may not be accounted for properly in approaches relying on parameterization such as, aside from purely semi-empirical methods, also the empirical dispersion correction for DFT³³ or the M06 method.³⁴ The algebraic diagrammatic construction to second order (ADC(2))³⁵ using the resolution of the identity approximation (RI)³⁶ is used as an efficient and reliable method for the description of excited state energies, properties, and gradients (see also ref. 25 and 37 for experiences on DNA fragments with this method and the related second order coupled cluster (CC2) method). These calculations are supplemented by multi-reference configuration interaction (MR-CI) computations, which provide reference values for strongly distorted geometries and structures with a

small S_0/S_1 gap, cases in which the reliability of a single reference method is limited. Solvation effects are treated using explicit solvation in an electrostatic quantum mechanics–molecular mechanics (QM/MM) coupling scheme. To allow for averaging over several different solvent configurations the concept of an averaged solvent electrostatic potential (ASEP)³⁸ was used. Mutual polarization between the core region and the solvent is treated in an iterative manner. A newly developed scheme for analysing excitonic and charge transfer (CT) interactions was used to allow for a meaningful description of the excited states even in challenging cases of partially delocalized orbitals and several interacting configurations.³⁹

Computational details

Following ref. 19 the adenine ribodinucleotide (diriboadenosine monophosphate, ApA) was considered. This system was solvated in 3000 water molecules and the negative charge of the phosphate group was counterbalanced with one Na^+ ion. RI-ADC(2)³⁵ calculations were performed in TURBO-MOLE^{36,40,41} using the SV(P),⁴² TZVP, TZVPP, and QZVPP basis sets.⁴³ MR-CISD calculations were carried out with COLUMBUS^{44–46} using the 6-31G* basis set⁴⁷ where CI expansions of about 200 million configurations were considered by applying the parallel version^{48,49} running on 64 CPUs. All MR-CISD results are reported with the size consistency correction suggested by Pople (+P).⁵⁰

Electrostatic embedding, QM/MM calculations⁵¹ were performed following ref. 25 and 52. For the MM interactions the Amber-99 forcefield⁵³ as implemented in TINKER⁵⁴ was used. The different potentials were combined using the Newton-X^{55–57} hybrid gradient facility.⁵² Initial structures were generated using PACKMOL.⁵⁸ The QM region consisted either of the whole ApA system or just the two adenines. The remaining system (*i.e.* the water molecules, the Na^+ ion and in the second case also the backbone) was treated as point charges using Amber-99 values. The connection of the adenine molecules to the sugar/phosphate backbone in the second case was achieved through the link hydrogen atom technique,^{51,59} where the link hydrogen was placed on the broken C–N bond but moved closer to the N atom with the distance reduced by a factor of 0.68922 obtained by considering individual optimizations of the isolated base and the nucleotide. The charge of the carbon atom forming the link atom on the MM side was set to zero, and to maintain the original neutral charge of the overall system, the excess charge (*i.e.*, the sum of the deleted charge of the MM-link atom and the partial charge corresponding to the atoms in the respective QM region) were equally distributed onto the three atoms bonded to the MM-link atom (*cf.* ref. 60).

Specifically the following levels were considered with or without additional solvation in water:

- “SV(P)-SV” denoting that the whole dinucleotide was in the QM region with the two adenine molecules described with the SV(P) basis set and the backbone with SV,
- “SV(P)-MM”, “TZVPP-MM”, “QZVPP-MM”, *etc.* denoting that the adenine molecules were considered at the



respective basis set while the backbone was considered only as MM point charges,

- “SV(P)-x” denoting that the backbone was not considered at all, used for test purposes.

Gas phase geometry optimizations were carried out at the SV(P)-MM and TZVP-MM levels allowing for a more efficient treatment. Optimizations in aqueous solution were performed in general with the whole dinucleotide in the QM region (SV(P)-SV) to simplify the computation of interaction terms.

The solvated QM/MM single point and geometry optimization calculations were performed using an averaged solvent electrostatic potential (ASEP, *cf.* ref. 38, 61 and 62). The idea of this approach is to combine the (properly scaled) point charges of several MD frames to obtain an averaged potential describing the solvent influence. Mutual polarization of the solvent field and the central molecule can be obtained by using a coupled method where charges obtained in the QM calculation are fed back to the MD.⁶¹ Geometry optimizations in such a potential actually amount to approximate free energy optimizations with respect to the solvent coordinates.⁶² An important consideration for a practical application of this approach is that the number of point charges has to be reduced for the QM/MM calculation, in particular when gradients are considered. For this purpose we use a coarse graining of the point charges (in a somewhat different approach as presented in ref. 38). Our method considers a dynamical threshold, which for point charges i and j takes on the value

$$d_{ij,\max} = \left(C \frac{r_i + r_j}{2} - r_0 \right)$$

which depends on the distances r_i and r_j of these point charges to the origin. The parameters r_0 and C represent the radius of the cavity and the fineness of the reduction procedure, respectively. Following an iterative procedure, any pair of charges of the same sign with a separation below $d_{ij,\max}$ was combined to one point charge with the summed charge located at the charge weighted center of the two charges. Charges of different signs were not combined in order to preserve their resulting dipole moments. This procedure preserves a very detailed description of the charge distribution close to the molecule of interest while efficiently retaining only large scale moments deriving from charges farther away. A more detailed description is given in ref. 63. For single point calculations this potential could be directly used. For geometry optimizations also a van der Waals (vdW) potential was included at the MM level. In the computations reported here, always 100 MD frames were used, obtained from an MD run with sampling every 0.5 ps. The parameter values $r_0 = 7.0$ Å and $C = 0.02$ were chosen. The average vdW potential was obtained by considering 25 of those MD frames, retaining only molecules with a maximum distance to the origin of 11 Å, and computing the vdW interaction to the central region. To obtain a consistent potential, the vdW terms had to be scaled due to the presence of several snapshot solvent configurations. This was achieved by scaling the well depth parameter ϵ of the water oxygen by a factor of 1/625 (while the parameter for

hydrogen was left at zero). This had the effect that all vdW interactions, which were computed as geometric means, were scaled by 1/25. Using these fixed point charges and vdW spheres as an external potential, the geometry of ApA was optimized. After the geometry optimization new atomic charges were fitted to the molecular electrostatic potential of the respective state of interest using Turbomole and the fitting procedure of ref. 64. With these new charges a new MD run was performed to yield a new ASEP, which was in turn used for a geometry optimization of the core region. The procedure was iterated until no more significant changes in geometries or charges were observed, requiring usually two or three cycles (depending on the quality of the starting structure and charges).

To represent the intermolecular separation between the two adenine molecules, the mutual distance between the C6 atoms, *i.e.* the C atoms bonded to the respective amino groups (Fig. 2), labelled d_{C6C6} , was in most cases considered. Additional geometric parameters are presented in the ESI.†

In some test calculations the counterpoise (CP) correction⁶⁵ for basis set superposition error (BSSE) was used, considering gas phase calculations at the SV(P)-MM, TZVP-MM, and QZVPP-MM levels. However, due to the problems of this approach for correlated methods (as described below) the main results reported are all without CP correction. The CP correction term was computed as

$$\Delta E_{CP} = -E(A1[A2]) + E(A1) - E(A2[A1]) + E(A2) \quad (2)$$

where $E(A1)$ is the energy of a QM calculation of adenine A1 alone and $E(A1[A2])$ the energy of a QM calculation of A1 where additionally the basis functions of A2 were present, and *vice versa*. The ΔE_{CP} term was added to the hybrid QM/MM energy for BSSE correction. CP corrected gradients for geometry optimizations were simply obtained by combining the gradients of the individual terms with the normal QM/MM gradient expression using the Newton-X hybrid gradient facility.

For the analysis of the excited states a scheme developed in ref. 39 (partially based on work by Luzanov and Zhikol⁶⁶ as well as Tretiak and Mukamel⁶⁷) was applied. The scheme proceeds by analyzing the transition density matrix in the atomic orbital AO basis. Using different summations over the blocks of this matrix, several different quantities can be computed:

- POS, the position of the excitation;
- PR, the excitonic participation ratio, representing delocalization;

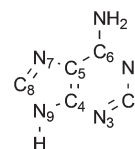


Fig. 2 The numbering scheme used in the adenine molecule.



- CT, the total amount of charge separation (including charge resonance and directed transfer contributions); and
- CT_{net} , the net charge shifted.

In addition it is also of interest to consider the natural transition orbitals (NTO),^{68–70} which can be obtained from a singular value decomposition of the transition density matrix. Aside from pictorial representations, also the NTO participation ratio PR_{NTO} was considered, counting the number of non-vanishing singular values.

Results and discussion

Franck–Condon point and vertical excitations

In this section the ground state potential around the Franck–Condon (FC) point, the vertical excitations, and effects of solvation will be discussed. In addition, several methodological points related to the reliability of our method will be considered.

In a first step the gas phase ground state potential of the system and the adequacy of the present methods to describe it were considered. Particular focus was laid on BSSE as it has been pointed out that this may have important effects for the binding energies of excimers.^{21,71} For this purpose structures were first optimized at the MP2 level considering the adenine molecules at the TZVP level and the backbone as an MM potential (*i.e.* the MP2/TZVP-MM level in the notation described above). Optimizing at this level without CP correction, a tightly stacked minimum “Stack A” with $d_{\text{C6C6}} = 3.00$ Å (as a representative stacking distance) was found. By contrast, when the CP correction was applied the stacking distance was increased significantly, yielding “Stack B” with $d_{\text{C6C6}} = 3.42$ Å. Furthermore, with CP correction, an additional “V-shaped” minimum stabilized by about 0.05 eV with respect to “Stack B” was found exhibiting $d_{\text{C6C6}} = 4.31$ Å. These structures are depicted in the ESI (S1–S3†). To understand these discrepancies and to get more insight into the ground state potential, an interpolation in internal coordinates between these three structures was performed and potential curves at the MP2 level using the SV(P), TZVP, and QZVPP basis sets with and without CP correction were computed and plotted against the respective energy of the V-shaped system (Fig. 3). By considering the same set of structures for these different cases, a direct comparison of the different methods is possible. However, it should be noted that these curves have been obtained by linear interpolation between the three structures (see above) and are not geometrically relaxed curves. Using the QZVPP basis set a very flat potential curve was obtained with and without CP correction showing differences between the minima and maxima on the curve of only ~0.05 eV. The situation is significantly different with the TZVP basis set. Without CP correction “Stack A” exists as a minimum at 0.13 eV below the energy of the V-shaped structure. By contrast, when applying the CP correction at this level “Stack A” is destabilized lying 0.14 eV above the V-shaped structure. Both values differ

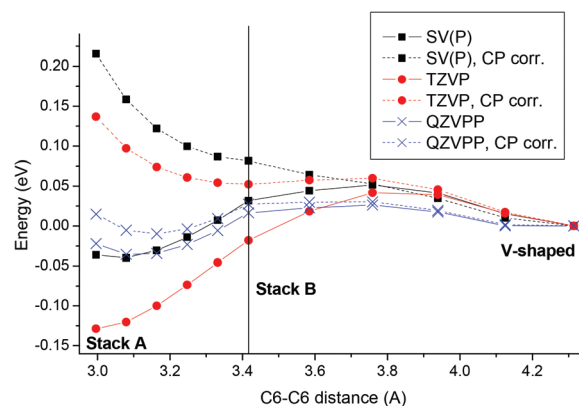


Fig. 3 Interpolated ground state potential energy curves for ApA plotted against the intermolecular C6–C6 distance for three basis sets considered with (dotted lines) and without (solid lines) CP correction.

strongly from the QZVPP reference. The results are even somewhat further off when applying the CP correction at the SV(P) basis set level, yielding a destabilization of “Stack A” by 0.22 eV. Interestingly the uncorrected SV(P) curve shows a very similar behaviour to the QZVPP curves exhibiting a similar flat potential, which is only somewhat biased toward the tightly stacked “Stack A” structure.

The strong basis set dependence of CP corrected MP2 interaction energies is in agreement with previous experience.⁷² However, an interesting observation in the present context is that the CP correction may even worsen the result. This phenomenon in the context of the different curves will now be discussed briefly. Firstly, the results suggest that as expected BSSE at the TZVP level is rather strong, explaining the drop in energy at lower stacking distances. Secondly, the good performance of the SV(P) basis set may be related to the fact that this basis set is simply too small to have significant superposition at the stacking distances considered here or possibly to other kinds of error compensation. Thirdly, a possible reason of the overcompensation of the CP correction has been given in ref. 73. Briefly, the virtual space in the “ghost” calculation (containing 265 orbitals in the SV(P) case) is larger than in the dimer calculation (230 orbitals) due to the smaller number of electrons present. This phenomenon, in addition to different energetics of these orbitals following from missing electron repulsion, leads to additional terms in the MP2 energy expression providing an overestimation of the CP correction term at the correlated level.

In summary we have shown that at the MP2/QZVPP-MM level the system possesses a rather flat potential energy surface with respect to a variation in the degree of stacking, suggesting that a wide distribution of such structures should exist at room temperature in agreement with experimental interpretations.^{19,26} At the MP2/SV(P)-MM level without CP correction the flatness of the QZVPP-MM surface is reproduced remarkably well. Therefore, this basis set and no CP correction will be used for geometry optimizations in the further course of this study. In addition, results will be verified at the QZVPP-MM



level. As described for example in ref. 74 the MP2 method tends to overestimate somewhat the interaction strength in nucleobase stacks and therefore yields somewhat too tightly stacked structures. For example, binding energies are too large by ~ 0.1 eV and the stacking distances are too small by ~ 0.1 Å as compared to CCSD(T) results (which are in turn about 0.1 Å shorter than experimental values).^{74,75}

We have recently discussed the UV absorption of oligonucleotides in detail.²⁵ Therefore, in this contribution the absorbing states will be considered only briefly. For this purpose the ground state structure of ApA was optimized in solution at the MP2/SV(P)-SV level using an ASEP solvent field included through electrostatic embedding as explained in more detail in the Computational details section and the vertical excitations were computed (Table 1). Aside from excitation energies and oscillator strengths, also descriptors characterizing the electronic excitation as defined in the Computational details section were determined. The first four states were of $\pi\pi^*$ character, all showing at least partial delocalization ($PR > 1.5$). The brightest state at this level was S_4 followed closely in oscillator strength by S_3 . Then two $n\pi^*$ states localized on either molecule follow. The S_7 and S_8 states are combinations of opposing CT states, yielding charge resonance states, *i.e.* states with a high amount of charge separation ($CT \approx 0.75$) but a low amount of net transfer ($|CT_{net}| < 0.30$), also characterized by a concurrent high delocalization degree ($PR \approx 1.9$). It may be noted here that the current computational protocol neglects to include dynamic solvent polarizability of the surrounding water, which may somewhat stabilize the CT states and increase net charge transfer as compared to the results presented in Table 1. But due to the small dynamic polarizability of water this effect should be rather small. After the CT states, two more localized $n\pi^*$ states follow. An observation to be made at this point is that many of the states are not pure Frenkel excitons but contain small CT contributions, showing that already at the Franck–Condon point, orbital interactions mixing these types of states (see also ref. 76) come into play. This is in agreement with our recent study on alternating DNA strands.²⁵ Another interesting fact is that the delocalized states exhibit a PR_{NTO} significantly larger than one, which is consistent with the statement that several configurations are needed to describe excitonic delocalization (see also ref. 39, 77 and 78).

To get a more detailed insight into the effect of the environment and of the computational model chosen, the adenine dinucleotide was considered at three different levels: (i) as two isolated molecules (SV(P)-x), (ii) two molecules with the backbone included at the MM level (SV(P)-MM), and (iii) the full system at the QM level (SV(P)-SV). Furthermore, models (ii) and (iii) were considered with and without H_2O solvation. In Fig. 4 the individual effects of these models on the first six states are summarized. In all of these cases the geometry optimized in solution (as used in Table 1) was considered. The largest effect visible is the shift of the $n\pi^*$ states due to H-bonding. Such a shift is a well known phenomenon in these types of systems.^{25,79–81} Further trends are that an inclusion of the backbone through point charges (moving from SV(P)-x to SV(P)-MM) somewhat destabilized the $n\pi^*$ states as well. A full QM treatment of the backbone (moving to SV(P)-SV) had only a minor effect, somewhat lowering all the excitation energies. But in summary the glycosidic bond has a rather low influence (*cf.* ref. 81). In solution the difference between SV(P)-SV and SV(P)-MM is somewhat enhanced: the $n\pi^*$ state on A2 is shifted down by a significant amount when the backbone is only considered at the MM level. However, a more detailed

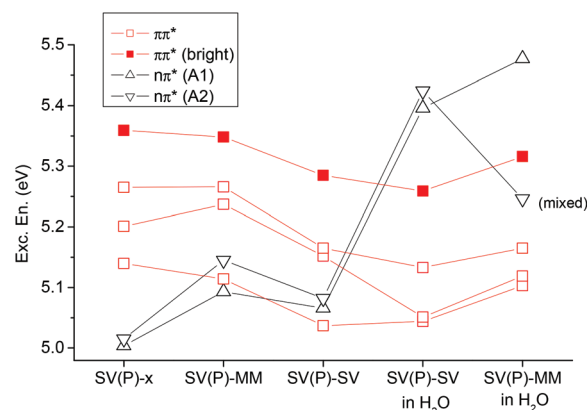


Fig. 4 Vertical excitation energies of the first six excited states of ApA using different models: two isolated adenine molecules (SV(P)-x), the system with the backbone treated at the MM level (SV(P)-MM), the full system at the QM level (SV(P)-SV), and the latter two levels with an additional consideration of water solvation.

Table 1 Excitation energies (ΔE , eV), oscillator strengths (f), descriptors and type assignments for the first 10 excited states of ApA in aqueous solution computed at the ADC(2)/SV(P)-SV level in the ground state minimum geometry

State	ΔE	f	POS	PR	CT	CT_{net}	PR_{NTO}	Type
S_1	5.044	0.033	1.26	1.61	0.13	0.02	2.12	$\pi\pi^*$
S_2	5.051	0.085	1.77	1.56	0.06	0.00	1.58	$\pi\pi^*$
S_3	5.133	0.160	1.51	2.00	0.09	-0.03	2.32	$\pi\pi^*$
S_4	5.259	0.189	1.46	1.99	0.10	-0.04	2.41	$\pi\pi^*$
S_5	5.396	0.004	1.03	1.05	0.04	0.01	1.03	$n\pi^*$
S_6	5.424	0.014	1.97	1.06	0.04	-0.03	1.03	$n\pi^*$
S_7	5.742	0.072	1.48	1.85	0.75	-0.28	1.73	CT ($\pi\pi^*$)
S_8	5.914	0.031	1.53	1.92	0.76	0.21	1.88	CT ($\pi\pi^*$)
S_9	5.931	0.002	1.05	1.10	0.07	0.04	1.06	$n\pi^*$
S_{10}	6.018	0.002	1.94	1.12	0.07	-0.03	1.12	$n\pi^*$



analysis of the character of this state revealed that the large apparent shift is probably enhanced by mixing with the $\pi\pi^*$ states. In summary we can state that the standard link-atom model used here is reliable but in critical cases the results may have to be checked carefully. An alternative link atom model for these types of systems including also two atoms of the adjacent sugar into the QM region has been suggested.¹⁰ Such an approach would, however, significantly increase the size of the QM region and add complications due to the need of two link atoms per nucleobase. Fortunately, in the further course of the present study we were able to perform full QM calculations on the whole dinucleotide; the link-atom models are only used for increased basis sets and MR-CISD calculations.

Finally, basis set effects on the vertical excitations will be discussed (Fig. 5). The general trend, a rather uniform decrease of excitation energies when the basis set is increased, agrees with previous experience.^{25,82} This decrease amounts to ~ 0.1 – 0.3 eV when going from SV(P) to QZVPP and is present for all states considered. However, the lowest $\pi\pi^*$ state is affected to a somewhat larger extent than the others. Furthermore, for the larger basis sets the brightest state is stabilized and becomes S_2 .

Excited state geometry relaxation

The second part of this paper will be concerned with processes happening in the excited state subsequent to photoabsorption. In this context it is of particular interest to investigate how interactions between the two nucleobases may give rise to new excited state minima and possible decay channels. In general, a large manifold of structures and a number of modes need to be considered. Here, we will focus on well-defined stacked structures considering that they are connected to the long-lived experimental transients²⁶ whereas unstacked structures will trivially decay according to monomer-like pathways. As far as coordinates are concerned, we will consider the intermolecular distance between the two C6 atoms on the bases (*cf.* Fig. 2) d_{C6C6} since this coordinate showed large alterations in the unconstrained optimizations (shown below). The study will focus on excited state minima, which are well accessible with the

present computational methods, whereas intersections between electronic states will only be considered indirectly.

First, the ApA system was analysed in gas phase. For this purpose a potential curve was computed where the d_{C6C6} coordinate was fixed at different values between 2.0 and 2.9 Å and the geometry of the remaining system was optimized for the S_1 state. Considering these geometries, the adiabatic energies of the S_0 – S_2 states relative to the ground state minimum geometry were computed (Fig. 6). An important feature is the flatness of the S_1 curve over the whole range considered, down to a distance of $d_{C6C6} = 2.0$ Å where the ground state surface is of course already strongly repulsive. To understand this phenomenon a more detailed inspection of the involved excited states is needed. At larger intermolecular separations the S_1 state is described by a local $n\pi^*$ transition on A1 ($PR \approx 1$, $CT \approx CT_{net} \approx 0$, $PR_{NTO} \approx 1$). This state possesses a very low oscillator strength ($f \approx 0.005$). There is no significant change down to $d_{C6C6} = 2.4$ Å, only that the PR and CT values increase slightly as the π^* orbital acquires partial delocalized character. The vertical excitation energy always stays above 3.0 eV. Structurally the system exhibits an out-of-plane motion of the amino group bonded to the C6 atom on A1 whereas almost no changes occur in A2 (see Fig. 7(a)).

When the d_{C6C6} distance is decreased to 2.30 Å and below, the geometry optimization converges to a different type of structure. The amino group on A1 is still out-of-plane, similar to the $d_{C6C6} \geq 2.40$ Å case, but now also the planarity of A2 is lifted (see Fig. 7(b)) yielding a structure similar to the $\pi(AA)^*$ minimum in solution as described below. The structures

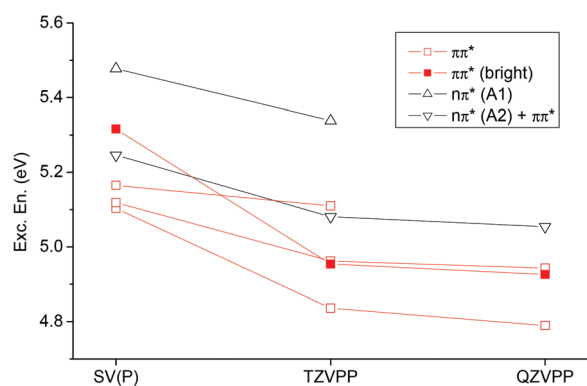


Fig. 5 Vertical excitation energies of the first six excited states of ApA in aqueous solution considering different basis sets (always considering the backbone at the MM level).

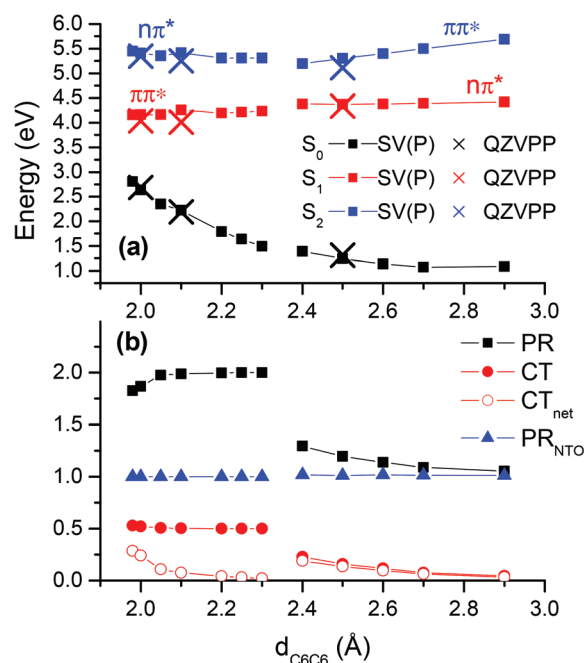


Fig. 6 Relaxed potential energy curve computed for the S_1 state of ApA at the ADC(2)/SV(P)-MM level in gas phase: (a) energies for the first three electronic states at this level (squares) and at the ADC(2)/QZVPP-MM level (crosses) for selected points; (b) statistical descriptors for the S_1 state.



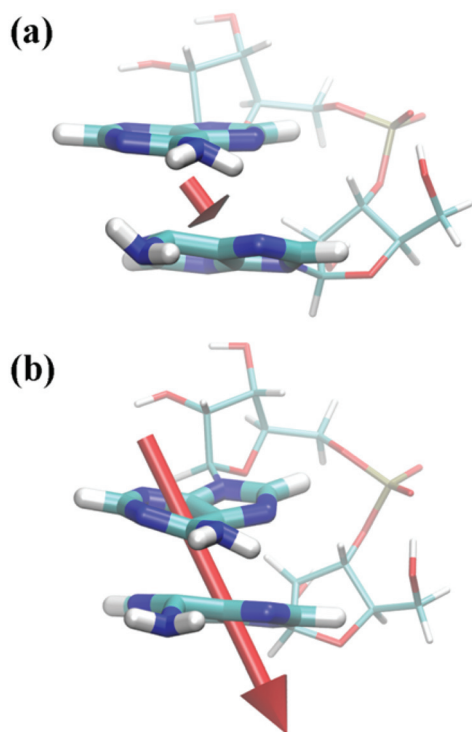


Fig. 7 Depiction of ApA structures optimized at the ADC(2)/SV(P)-MM level in gas phase on the S_1 surface for fixed d_{C6C6} distances of (a) 2.6 Å and (b) 2.2 Å. Transition dipole moments corresponding to (a) $f = 0.005$ and (b) $f = 0.035$ are plotted as well.

exhibit a strong interbase “tilt” and “roll”. This can for example be understood from the fact that when d_{C6C6} was fixed at 2.20 Å, the intermolecular distances between the neighbouring atoms were much larger with $d_{C5C5} = 2.50$ Å and $d_{N1N1} = 2.89$ Å. The states in this area can be characterized as delocalized $\pi\pi^*$ states ($PR \approx 2$). A more detailed consideration of their excited state structure reveals that their nature is different from any of the states at the FC point as presented in Table 1. This may be immediately seen by considering the number of independent transitions needed to describe the excited state (*i.e.* the PR_{NTO} value). At the FC point there were always several transitions involved for delocalized states ($PR_{NTO} \approx 2$), in agreement with the model of two independent transitions needed to form a delocalized Frenkel exciton.^{39,77,78} By contrast the S_1 state computed for structures on this potential curve exhibited always $PR_{NTO} \approx 1.00$ despite the delocalization present for $d_{C6C6} < 2.30$ Å. In fact, this difference could already be seen in the canonical orbital representation, which showed that the excited state was always largely dominated by a simple transition between the highest occupied molecular orbital (HOMO) and the lowest unoccupied MO (LUMO).[†] A different way to analyse these states can be achieved by considering the charge transfer character. The

values $CT \approx 0.5$ and $CT_{net} \approx 0$, which are present for $2.05 \text{ Å} \leq d_{C6C6} \leq 2.25 \text{ Å}$, show that this is a state with 50% charge separation but no net transfer of charge. Another way to interpret this is by the presence of a homogeneous excited state over the whole system with an equal probability of the *electron* and *hole* being on the same or different fragments, a condition which is in turn consistent with the simple picture of only one transition. This situation is very similar to the naphthalene dimer.³⁹ In that case we could show that the excimer state could formally be constructed as a mixture between the Frenkel excitonic and charge resonance states present at larger intermolecular separations. However, at the excimer minimum the state was of neither type and was determined by direct interactions between the orbitals on the two molecules, rather than just electrostatic interactions. Due to the $\pi\pi^*$ nature and the lack of charge transfer of this excimer state in ApA it possesses appreciable oscillator strength (*e.g.* $f = 0.035$ for $d_{C6C6} = 2.2 \text{ Å}$). An interesting phenomenon arising from the excited state structure is an out-of-plane polarization of the transition moment (Fig. 7(b)). It should be noted here that $\pi\pi^*$ states in planar systems show strict in-plane polarization for symmetry reasons. Furthermore, if these states are coupled by purely electrostatic interactions to form Frenkel excitons, the polarization vectors still have to remain parallel to the molecular planes of the aromatic molecules. In this sense the out-of-plane transition moments are an observable manifestation of the unusual excited state structure described here. An extended analysis of this potential curve is presented in the ESI (Fig. S8†). In particular, it should be noted that for the $\pi\pi^*$ state orbital interactions come into play already at $d_{C6C6} = 2.90 \text{ Å}$.

In a one-dimensional picture one would expect to notice a lowering of the S_1/S_2 gap to an avoided crossing as these two states exchange their character around $d_{C6C6} = 2.35 \text{ Å}$. Such a feature is only rudimentarily visible in Fig. 6 where the S_1/S_2 gap always stays above 0.8 eV. This fact is probably related to the multidimensional nature of the potential surface and the fact that the optimization procedure moves the structures away from the avoided crossing when relaxing the S_1 state of the respective character. In fact, when searching for a signature of the S_1/S_2 avoided crossing a different type of constrained local S_1 minimum was found in the area between $2.25 \text{ Å} \leq d_{C6C6} \leq 2.35 \text{ Å}$. Those structures were determined by strong ring puckering of C6 on A1 and the amino group was shifted out-of-plane pointing in a direction almost perpendicular to the ring, a structure resembling the C6-puckered intersection in the monomer,⁸³ only that the virtual orbital was partially delocalized here ($PR = 1.33$). The S_0/S_1 gap was lowered to $\sim 0.5 \text{ eV}$ (while the S_1/S_2 gap moved up to $\sim 1.5 \text{ eV}$). The MR-CISD reference value was somewhat higher at 1.1 eV, which is, however, still in agreement with the picture of a close neighbourhood of an S_0/S_1 intersection structure. This finding suggests that monomer-like decay pathways to the ground state may play a role even in tightly stacked structures.

Fig. 6 shows that the S_1 potential is very flat and in fact no minimum related to the exciplex could be located by unconstrained optimizations. By contrast, these optimizations lead

[†]The presence of only one transition between two canonical orbitals is a sufficient but not a necessary criterion for $PR_{NTO} = 1$. If more than one transition is present they may still disappear due to the NTO transformation.



to a decrease of d_{C6C6} below 2 Å and a subsequent strong destabilization of the ground state with very small S_0/S_1 gaps and a concomitant failure of the ADC(2) method. These results suggest that a new deactivation channel arises in the adenine dimer, which is accessed through decreasing the intermolecular distance between the respective C6 atoms to below 2.0 Å, similar to results obtained by semi-empirical calculations,²³ only that in those cases the mutual C2 distance was shortened to below 2.0 Å. In ref. 23 a non-reactive deactivation between the two adenine molecules was observed. By contrast, an analogous intersection between two thymine molecules may be

responsible for photodimerization in this case.⁸⁴ The importance of this type of intersection and the relation to potential photoproduct formation in different base stacks requires further investigations.

To verify our results against basis set effects, selected calculations at the ADC(2)/QZVPP-MM level were performed (Fig. 6, crosses). These showed very good agreement with the SV(P) results, highlighting in particular the relative unimportance of BSSE in this case. MR-CISD calculations performed at the structures with $d_{\text{C6C6}} = 2.0$ Å and $d_{\text{C6C6}} = 2.1$ Å gave excitation energies, which were very similar to the ADC(2)/SV(P)-MM results (about 0.2 eV higher) highlighting that also multi-reference effects do not play an important role.

The computation of potential curves in solution is not feasible with the present computational protocol due to the iterative nature of the procedure (see the Computational details section for more information). Therefore, the focus was on locating different S_1 minima in solution. Three minima possessing entirely different character were found. They are depicted in Fig. 8 and the major properties are collected in Table 2. The first minimum, which will be denoted AA*, is described by a local $n\pi^*$ excitation on A2 (POS = 1.97, PR = 1.06, orbitals shown in Fig. S9†). This nucleobase (Fig. 8, top) shows puckering around the C2 atom exhibiting a very similar structure to a corresponding S_1 minimum in the isolated adenine molecule (see *e.g.* ref. 83). The value of d_{C6C6} used as a measure of the intermolecular separation is 3.05 Å, which is similar to the ground state distance at this level, also reflecting that there is only negligible interaction.

The second S_1 minimum $n(\text{AA})^*$ is related to the structures present in the gas phase potential curve for $d_{\text{C6C6}} \geq 2.4$ Å (Fig. 6, right part). However, the molecules are now moved closer together in the geometry optimizations, in particular around the C6 atoms, leading to $d_{\text{C6C6}} = 2.21$ Å and the structure could indeed be called an exciplex. The vertical excitation energy of 2.438 eV shows the strong energetic stabilization in the excited state. As opposed to the gas phase, the excited state is strongly polarized here, showing $\text{CT}_{\text{net}} = 0.41$. This value follows from an excitation from the n orbital on A1 to a π^* orbital which is delocalized between both adenine molecules, resulting also in partial delocalization (PR = 1.53, see also Fig. S10†).

The third minimum denoted $\pi(\text{AA})^*$ is related to the gas phase potential curve at distances $d_{\text{C6C6}} \leq 2.30$ Å (Fig. 6, left part). It is structurally similar to $n(\text{AA})^*$, only that the distortions of the rings are enhanced. A strong exciplex interaction leads to a decrease of the separation to $d_{\text{C6C6}} = 2.00$ Å and to a vertical excitation energy of 1.843 eV. The excitation is described by a single transition between a π and π^* orbital

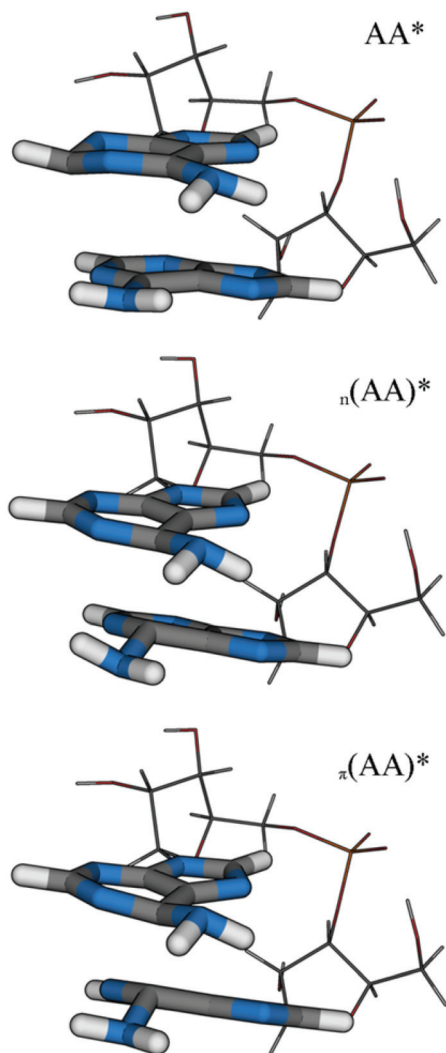


Fig. 8 Molecular structures of the different minima of ApA located by optimizations at the ADC(2)/SV(P)-SV level in aqueous solution.

Table 2 Intermolecular d_{C6C6} distances (Å), adiabatic S_0 energies (eV), excitation energies (ΔE , eV), oscillator strengths (f), descriptors and type assignments for the first excited states at different S_1 minimum geometries of ApA in aqueous solution computed at the ADC(2)/SV(P)-SV level

Struct.	d_{C6C6}	S_0 En.	ΔE	f	POS	PR	CT	CT_{net}	PR_{NTO}	Type
AA*	3.05	1.2	3.221	0.016	1.97	1.06	0.06	−0.06	1.01	$n\pi^*$
$n(\text{AA})^*$	2.21	2.2	2.438	0.010	1.25	1.53	0.45	0.41	1.00	$n\pi^*$
$\pi(\text{AA})^*$	2.00	2.3	1.843	0.061	1.40	1.77	0.53	0.32	1.00	$\pi\pi^*$



both delocalized over both adenine molecules (Fig. S11†). Being of $\pi\pi^*$ character, this structure shows an appreciable oscillator strength of 0.061 and it can be expected that its fluorescence should be visible. The transition moment has again a strong out-of-plane component similar to the gas phase calculation (*cf.* Fig. 7(b)). As described above, the presence of only one transition ($PR_{\text{NTO}} = 1.00$) in a strongly delocalized state ($PR = 1.77$) is in contrast to the picture of a Frenkel exciton and only orbital interactions can lead to this situation, highlighting the marked difference between the character of the state at the exciplex structure and any state at the FC point. This situation is also represented by $CT = 0.53$. Due to solvation the excited state is partially polarized ($CT_{\text{net}} = 0.32$). The comparable gas phase structure at $d_{\text{C6C6}} = 2.00$ Å exhibits $\Delta E = 1.534$, $CT_{\text{net}} = 0.24$ and an S_0 energy of 2.633 eV (as presented in Fig. 6). It thus appears that the solvent has an important effect on the structural relaxation leading to exciplex structures, which are more strongly polarized and not so strongly distorted as opposed to the gas phase.

The relative adiabatic ground state energies are presented in Table 2 as well (these were converged up to a few tenths of an eV in the iterative procedure described in the Computational details section). When adding these to the vertical energies given also in the table, it follows that the AA^* and ${}_n(AA)^*$ minima are located at about 4.5 eV with respect to the ground state minimum or in other words they are reached by a relaxation of about 0.7 eV from the bright excited states at the FC point. The ${}_n(AA)^*$ structure is situated lower at around 4.1 eV.

In a next step we aim to verify the results in light of reference calculations (Table 3). The backbone was treated at the MM level in these calculations and it was first established that this only had a minor effect, verifying that the ADC(2)/SV(P)-MM results always reproduced the full QM (SV(P)-SV) ones well, deviating by at most 0.1 eV in excitation energies. Increasing the basis set to the QZVPP-MM level somewhat lowered the excitation energies (by about 0.1–0.2 eV) but produced no systematic changes. MR-CISD excitation energies were always somewhat higher, which is however a known phenomenon for this method related in part to size consistency errors present in a system of this size and to orbital relaxation (see *e.g.* ref. 85 and 86). It is however worth noting that at least the energetic

trend is reproduced here. For comparison, also calculations at the PBE0⁸⁷ level were performed, showing quite good agreement with the ADC(2) results.

In agreement with ref. 21 and 22 we find a number of excited state minima of different degrees of delocalization and charge separation located down to adiabatic energies of about 4 eV relative to the ground state minimum. This presence of a number of different minima and plateaus is consistent with our observation that the potential surface in the ground (Fig. 3) as well as excited (Fig. 6) states is very flat. In ref. 21 and 22 these minima exhibit vertical S_0/S_1 energy gaps above 2.5 eV and stacking distances above 2.8 Å, whereas the ${}_n(AA)^*$ minimum in this study exhibits a gap of only 1.8 eV, which derives from a stronger decrease of the intermolecular separation and an out-of-plane movement of the amino groups. Precisely answering the question of how closely the two adenine molecules approach each other is quite challenging as accurate intermolecular potentials are difficult to compute. In ref. 21 the CASSCF method was used for geometry optimizations (while CASPT2 was only used for single point calculations). Due to a lack of dynamical electron correlation the CASSCF method is not expected to describe dispersion well and should therefore underestimate interaction energies and overestimate binding distances. Furthermore, it is also quite difficult to describe those interactions correctly with density functional theory. Whereas the M06-2X functional used in ref. 22 has been designed to give good interaction energies in the case of standard complexes, it is not clear whether good results can also be obtained for interatomic distances far below 3 Å. By contrast, we use the correlated *ab initio* methodology, which is, at least in general, able to provide good interaction energies. The MP2 method has been described as producing reasonably accurate binding energies, only that dispersion energies may be somewhat overestimated.⁷² Additionally, the state loses its single reference character giving a value of 0.063 for the D1 diagnostic,⁸⁸ which is exceeding the recommended value (0.04). In summary, it can be said that we could show strong evidence for exciplex structures determined by short C6–C6 distances and strong structural distortions even if the interaction strength may be somewhat overestimated. However, it should also be recognized that due to the flatness of the potential energy curves shown in Fig. 6, especially the location of the minima could change by a few tenths of an Å.

The existence of a strong exciplex interaction in ApA is certainly in agreement with the reported slow excited state decay of this system.¹⁹ However, in contrast to the original experimental interpretation we find that in this system interactions due to orbital overlap play a dominant role while charge transfer is small. A similar mechanism could also explain the increased stability reported for ApG as opposed to purine–pyrimidine stacks, a phenomenon which could not be accounted for when simply considering ionization potentials and electron affinities of the individual bases.¹⁹

An emission energy below 2 eV, which is found for the ${}_n(AA)^*$ exciplex in this study, is smaller than the values of fluorescence maxima reported from experimental studies of

Table 3 Vertical excitation energies (eV) of the S_1 and S_2 states at three different S_1 minimum geometries of ApA in aqueous solution considering different levels of theory

			AA^*	${}_n(AA)^*$	${}_n(AA)^*$
Vert. S_0-S_1	ADC(2)	SV(P)-SV	3.221	2.438	1.843
	ADC(2)	SV(P)-MM	3.259	2.476	1.898
	ADC(2)	QZVPP-MM	3.157	2.351	1.715
	MR-CISD	6-31G*-MM	3.651	3.008	2.295
	PBE0	TZVP-MM	3.307	2.533	1.908
Vert. S_0-S_2	ADC(2)	SV(P)-SV	4.267	3.258	2.821
	ADC(2)	SV(P)-MM	4.370	3.316	2.859
	ADC(2)	QZVPP-MM	4.188	3.081	2.735
	MR-CISD	6-31G*-MM	—	3.523	—
	PBE0	TZVP-MM	4.080	3.105	3.018



poly(dA) and d(pApA), which are around 3 eV.^{6,89–91} It appears therefore that the binding distances may be somewhat too short here. Due to the steep increase in the ground state potential with decreasing intermolecular distance (*cf.* Fig. 6) even small geometric changes may result in a large difference from an energetic viewpoint. In spite of this difference in quantitative terms our study sheds new light on excimer formation and highlights the importance of orbital interactions, which have rarely been considered in previous investigations. In particular, an out-of-plane polarization of the fluorescence (*cf.* Fig. 7), which has been reported from experiment,⁹⁰ is difficult to explain when only pure Frenkel exciton and CT states are considered.

Conclusions

A detailed *ab initio* QM/MM study of the excited state potential energy surfaces of ApA in solution was performed using the ADC(2) and MR-CISD methods. First, the Franck–Condon point was analyzed. The flat ground state surface represents the structural flexibility well known for dinucleotides. For the vertical excitations Frenkel excitonic states as well as charge transfer states were found with only a small mixing between these two types (less than 10%). As a next step, structural relaxation in gas phase and in solution was considered. In gas phase a potential curve against the intermolecular d_{C6C6} distance was computed showing a very flat potential down to $d_{C6C6} = 2.0$ Å. Three different minima on the S_1 surface of different excitation characters ($n\pi^*$ and $\pi\pi^*$) as well as varying degrees of delocalization (up to about 80%) and charge transfer (up to about 0.4 e) were located. This emergence of several S_1 minima of different character and of flat potential surfaces in between them agrees well with previous studies.^{21,22} A new outcome of this study is the presence of exciplex minima with significantly decreased intermolecular separations, in particular between the C6 atoms, reaching in our calculations distances of $d_{C6C6} \leq 2.20$ Å. The electronic structure at these minima was analyzed in detail using methods recently developed for this purpose.³⁹ The main result of this analysis is that the minima are neither of Frenkel excitonic nor of pure charge transfer type but that orbital interactions lead to coherent homogeneous excited states, which differ significantly from any states present in the Franck–Condon region. These states were in turn dynamically polarized by interactions with the solvent, which led to a net charge transfer of about 0.3–0.4 e for these states. The most stable minimum was of $\pi\pi^*$ character, which in connection with the low degree of charge separation led to the presence of appreciable transition strength for this structure. Interestingly the transition showed a high degree of out-of-plane polarization.

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