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Real-time observation of sub-100-fs charge and energy transfer processes in DNA dinucleotides†

Using as showcase the DNA dinucleotide 5'-dTpdG-3', in which the thymine (T) is located at the 5' end with respect to the guanine (G), we study the photoinduced electronic relaxation of coupled chromophores in solution with an unprecedented refinement. On the one hand, transient absorption spectra are recorded from 20 fs to 45 ps over the 330-650 nm range with a temporal resolution of 30 fs; on the other hand, quantum chemistry calculations determine the ground state geometry of the 4 possible conformers with stacked nucleobases, the associated Franck–Condon states, and map the relaxation pathways leading to excited state minima. Important spectral changes occurring before 100 fs are correlated with concomitant $G^+ \to T^-$ charge transfer and $T \to G$ energy transfer processes. The lifetime of the excited charge transfer state is only 5 ps and the absorption spectrum of a long-lived $n\pi^*T$ state is detected. Our experimental results match the transient spectral properties computed for the anti-syn conformer of 5'-dTpdG-3', which is characterized by the lowest ground state energy and differs from that encountered in B-form duplexes.

Introduction

Excited state relaxation in DNA is studied in relation with its damage induced by absorption of UV radiation¹ and in view of future developments of DNA-based label-free optoelectronic devices. ^{2,3} In both cases, charge and energy transfer processes play a key role. For example, the formation of the carcinogenic (6–4) photo-adducts, ⁴ as well as charge carrier generation by low-energy photons, ⁵ involve the population of an excited charge transfer (CT) state, ⁶ in which an atomic charge has been transferred between two stacked nucleobases. Similarly, excitation energy transfer (ET) among nucleobases has been shown to provoke remote photodamage. ⁷

Numerous transient absorption (*TA*) studies have characterized the decay of *CT* states in various types of DNA multimers, ranging from dinucleotides and single strands to

duplexes and guanine quadruplexes (G-Quadruplexes).8-22

However, the formation of these states from the excited states initially populated upon photon absorption (Franck-Condon

states) has not been observed in real time. Likewise, ET was inferred from the fluorescence anisotropy values detected once

the process has occurred.²³ The main reason for the lack of direct experimental observation is that these processes are

(often referred as exciplexes/excimers). However, this approach becomes problematic when wave packets evolve along potential energy surfaces (*PES*). The latter do not fulfil the requirements for exponential decays, which are valid when the interactions of the chromophore with its environment are uniform in the three-dimensional space. They fall instead in the category of restricted geometries, in which dynamics has been discussed for a large variety of processes and systems.²⁴⁻²⁹

Here, we report the first comprehensive study monitoring the progression of electronic excitations from the Franck-Condon states to the *PES* minima in DNA multimers. We use as a showcase the DNA dinucleotide 5'-dTpdG-3', abbreviated as **TG**, in which the thymine (T) is located at the 5' end and the guanine (G) at the 3' end. The investigation of dinucleotides by

faster than the time resolution of the ultrafast spectroscopy setups used to date in these studies, which is typically in the 150–300 fs range.

A second, albeit equally important, difficulty is conceptual. So far, the data derived from *TA* experiments have been analysed using multiexponential functions, and time constants were correlated with monomer excited states and *CT* states (often referred as exciplexes/excimers). However, this approach becomes problematic when wave packets evolve along potential energy surfaces (*PES*). The letter do not fulfil the requirements

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*TA*⁸⁻¹⁴ spectroscopy has proved helpful in disentangling processes in larger systems, in which a multitude of interactions are operative. In addition to being a convenient model system, the 5'-dTpdG-3' sequence is frequently encountered in G-Quadruplexes. Thymines located at the 5' position with respect to the G-core have important implications on the geometry adopted by these four-stranded structures.³⁰

Our TA experiments are performed with an unprecedented combination of temporal resolution and spectral coverage;31 TA spectra (TAS) following resonant photoexcitation by ≈25 fs pulses at 266 nm are recorded from 20 fs to 45 ps over the 330-650 nm range with a temporal resolution of ≈30 fs, as discussed in Section 1 of the ESI.† In parallel, quantum chemistry calculations determine the associated Franck-Condon excited states of the four possible conformers with stacked nucleobases, and map the relaxation pathways leading to excited state minima for all of them. The TAS corresponding to the most stable conformer are also computed. We show that the evolution from the Franck-Condon states toward these minima is manifested by an ultrafast spectral shift occurring with halftime $(\tau_{1/2})$ of 50 fs. Moreover, along with the formation of the $G^+ \to T^- CT$ state, we detect an ultrafast $T \to G ET$. In a more general way, this is the first comprehensive study of photoinduced processes on bichromophoric systems performed with such refinement, i.e. detecting several transient species over a broad spectral range at early times and identifying them via the shapes of their TAS.

Results and discussion

Ground state properties

Fig. 1a shows the steady-state absorption spectrum of TG together with that of an equimolar mixture of the mononucleosides thymidine (dT) and deoxyguanosine (dG), their intensity being normalized at the peak wavelength of the laser pulse (266 nm). At this wavelength the molar absorption coefficients of dT and dG are the same.32 The maximum of the TG spectrum is slightly blue-shifted with respect to that of monomers (254 vs. 257 nm) and its intensity is somewhat higher, suggesting chromophore coupling. Base stacking in DNA is commonly detected by changes in the molar absorption coefficient upon heating (hypochromism/hyperchromism). This temperature dependence is explained by the fact that orbital overlap among stacked nucleobases leads to coupling between $\pi\pi^*$ and CT transitions. 33,34 Such changes are indeed observed in the difference of the absorption spectra recorded for TG at 95 °C and 23 °C (Fig. 1b), confirming base stacking. As reported in the literature, 9,14 these changes in dinucleotides do not exceed a few percent.

In order to determine the stacking patterns, we optimized the four possible stacking geometries, *anti-anti*, *anti-syn*, *syn-anti* and *syn-syn* (Fig. 2), where *anti* and *syn* refer to the position of each nucleobase (in the order T-G) with respect to the angle of the glycosidic bond (Fig. S2†), associated with the deoxyribose moiety.

The energy of the *anti-syn* conformer is lower, respectively, by 0.06, 0.19 and 0.17 eV than those of the three other

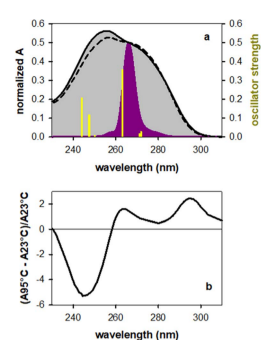


Fig. 1 (a) Steady-state absorption spectra recorded for TG (solid lines) and an equimolar mixture of dT and dG (dashed lines) normalized at the peak wavelength (266 nm) of the exciting laser pulse (violet); yellow sticks correspond to the electronic transitions computed for the anti-syn TG conformer. (b) Difference between the steady-state absorption spectra recorded for TG at 95° (A_{95 °C}) and at 23 °C (A_{23 °C}), divided by (A_{23 °C}).

conformers (Fig. 2 and Table S1†). These values are higher than the thermal energy at room temperature (0.025 eV), making *anti–syn* **TG** the most stable conformer. Therefore, we show detailed results for this conformer. However, in order to get a global picture, we also compute the Franck–Condon excited states and the associated relaxation pathways for the *syn–anti* and *syn–syn* conformers, and refine those of the *anti–anti* **TG**, whose properties were reported in ref. 35.

The properties of the 6 Franck-Condon states of anti-syn TG are shown in Table 1 and the difference in the electronic density between their ground and excited state in Fig. S3.† Among them, S₃, S₅ and S₆, having an oscillator strength higher than 0.1, are most likely to be populated by the exciting laser pulse. S₃ is a $\pi\pi^*T$ state, encompassing a very weak (0.08 a.u.) $G^+ \to T^-$ CT. A larger CT (0.57 a.u.) characterizes the S₅ state, while S₆ results from a mixing of the $\pi\pi^*G$ (La) state with weak CT (0.20 a.u.). The omnipresence of the CT is understandable, because G and dT are the nucleobases with the highest oxidation and the lowest reduction potential, respectively.36 All 6 states are represented as sticks in Fig. 1a. We note that the energy of the S5 state coincides with that of the $\pi\pi^*(Lb)$ state in the free guanosine chromophore, which is readily excited at 267 nm.^{32,37–39} Interestingly, no $\pi\pi^*G(Lb)$ was identified for anti-syn TG in the examined spectral range.

The properties of the Franck–Condon states of the three other conformers are presented in Tables S2, S3 and S4.† By comparing these states determined for all four conformers, we remark that the *CT* character is favoured when at least one base

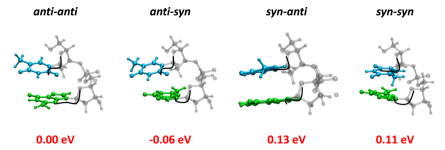


Fig. 2 Ground state structures of the four possible conformers with stacked nucleobases computed for TG and their relative energies (ΔG , in red). Thymine in blue; guanine in green. The dihedral angles that determine the *syn* and *anti* conformations are indicated in black.

Table 1 Properties of the Franck–Condon states determined for *anti*–syn **TG** at the PCM/M052X/6-31G(d) level of theory. VAE: vertical absorption energy; f: oscillator strength; δ : charge transfer character

0.0295 0.0192 0.3573 0.0093 0.1188	0.01 0.02 0.08 0.01 0.57
	0.1188

presents syn orientation. Indeed, no state with significant (>0.3 a.u.) CT is encountered below the S_{10} for the anti-anti conformer, whereas for the anti-syn and syn-syn conformers the S_5/S_6 states are characterized by a CT contribution. This trend is more pronounced in the case of the syn-anti conformer, for which the lowest-lying bright $\pi\pi^*$ states (S_2 and S_4) exhibit modest (0.3–0.4 a.u.) CT character.

Transient absorption spectra and dynamics

The experimental TAS recorded for TG, presented in Fig. 3, consist of a broad photoinduced absorption (PA) band whose shape and intensity evolve with time. Their comparison with the TAS of an equimolar mixture of dT and dG at selected times reveals important differences (Fig. 3a), further confirming the existence of significant electronic coupling between the nucleobases, in line with the steady-state spectra in Fig. 1. In particular, the negative signal, stemming from stimulated emission (SE) of the thymidine chromophore (Fig. S4†), already reported in the literature, 11,40 is completely absent from the TAS of the dinucleotide. Thus, it appears that the contribution of the $\pi\pi^*T$ state, either correlated with unstacked nucleobases or present as a localized state within stacked ones, is significantly lower than what is expected for a solution containing 50% dT chromophores. However, we cannot rule out the presence of a weak SE, peaking below 400 nm, which may overlap with the UV PA band and decrease its intensity. Consequently, at early times, we focus mainly on the spectral changes in the visible spectral range.

Focusing on the evolution of the **TG** *TAS*, we observe that, at early times (Fig. 3b), their intensity increases with the time, while the maximum of the low-energy band moves from 602 to

565 nm. The dynamics of the shift can be approximated by a power law $\alpha t^{-0.04}$ (Fig. 4a), the $\tau_{1/2}$ being 50 fs. After 100 fs, the intensity in the visible starts decreasing slowly, while the shift toward shorter wavelengths persists, albeit at a significantly slower rate (Fig. 3c). In this case, the spectral shift can be described by a mono-exponential function with a time constant of 0.61 ± 0.08 ps (Fig. 4b). The maximum of the low-energy band is located at 530 nm at 3 ps. After this time, when the monomer signals have practically disappeared, the decay of the low-energy band is also described by a mono-exponential function with a time constant of 5.44 ± 0.03 ps (Fig. 4c). Following this decay, a long-lived background *PA* remains (Fig. 3a).

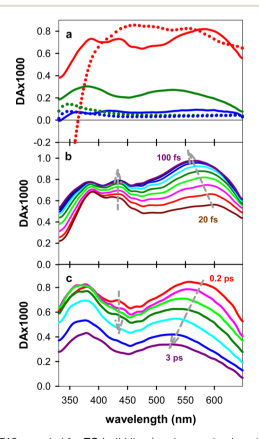


Fig. 3 TAS recorded for TG (solid lines) and an equimolar mixture of dT and dG (dots). (a) 50 fs (red), 5 ps (green) and 40 ps (blue). (b) From 20 to 100 fs with 10 fs steps. (c) 0.2 ps (red), 0.3 ps (pink), 0.4 ps (light green), 0.6 ps (dark green), 1 ps (cyan), 2 ps (blue) and 3 ps (violet).

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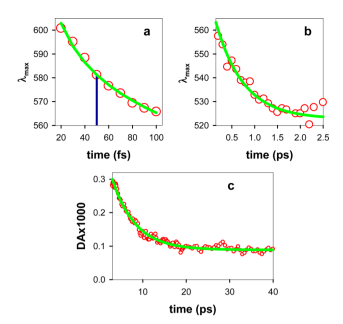


Fig. 4 Photo-induced dynamics (red circles) observed for TG over 3 timescales. (a) and (b) peak position (λ_{max}) of the low-energy band. (c) Decay of the *TA* signal at 570 nm. Green lines correspond to fits with a power law (a) and mono-exponential (b and c) functions. The blue line in (a) indicates $\tau_{1/2}$.

Computed relaxation paths

In order to get an insight into the origin of the above-described spectral evolution, we computationally mapped the deactivation pathways for the first 5 low-energy Franck–Condon states, plus the one exhibiting the largest charge transfer. For anti–syn TG, we identified three minima in the PES of the first excited states: min- $\pi\pi^*G$ (La), min-CT and min- $n\pi^*T$ (Table S5†). The pathway leading from S₃, which has predominantly $\pi\pi^*T$ character, to min- $\pi\pi^*G$ corresponds to a T \rightarrow G ET. S₅ leads to min-CT, where an electric charge of 0.81 a.u. has been transferred from G to T. The min- $n\pi^*T$ is reached following internal conversion from the S₂ and S₄ states, and, since the former state is not completely dark, we also expect weak population of this minimum.

The various relaxation pathways are schematically illustrated in Fig. 5; details are given in Table S5.† We stress that during the optimization procedure more than one reaction coordinate was varied. However, for two of the minima, one coordinate undergoes larger modifications than the others: the min- $\pi\pi^*G$ strongly distorts the C_2 -NH $_2$ coordinate whereas the min- $\pi\pi^*T$ moves the O_4 out of the plane (atom labelling in Fig. S5†). In the case of the min-CT there is a global rearrangement of the interbase distance and displacements, plus some modifications in the thymine aromatic ring.

Similar calculations were performed for the three other conformers. The results are summarized in Tables S6, S7 and S8.† For *anti–anti* **TG**, three minima, min- $\pi\pi^*G$ (La), min- $\pi\pi^*T$ and min- $n\pi^*T$, were also determined. In agreement with the higher stability of the CT states at the Franck–Condon region, mentioned above for the "syn" conformers, a min-CT was

optimized for the *syn-anti* conformer, while for *syn-syn* **TG** a conical intersection between the first excited and the ground state exhibiting *CT* character was found.

The *TAS* computed for the Franck–Condon states of *anti–syn* **TG** with the highest oscillator strength (Tables 1, S_3 , S_5 and S_6 †) are presented in Fig. S6† together with the corresponding minima. We remark a clear trend: the spectra of the minima are all blue-shifted with respect to those of the initial states.

Assignment of the observed spectral evolution

Next, we search the fingerprints of the above-described relaxation pathways in the experimental *TAS*, starting from the long times. We also discuss these data in comparison with a *TA* study on 5'-dGpdT-3'dinucleotide,¹¹ whose polarity, *i.e.* the order according to which two nucleobases are connected *via* the phosphodiester backbone, is opposite than that of **TG**.

The longest living transient species detected in 5'-dGpdT-3' was attributed to the thymine triplet state ($^3\pi\pi^*T$). However, the spectral shape of ${}^3\pi\pi^*T$, studied by ns flash photolysis and peaking at 350 nm,41 is very different than our TAS at 40 ps (Fig. 6a), exhibiting three well-defined absorption bands in the probed wavelength range (see Fig. S7a† for a comparison). These bands are absent from the 40 ps TAS determined for an equimolar mixture of dG and dT, shown on an appropriate scale in Fig. S8,† and from the ${}^3\pi\pi^*G(La)$ spectrum, peaking at 370 nm.42 The 40 ps TAS (Fig. 6a) resembles instead the spectrum computed for the min- $n\pi^*T$ state (Fig. 6b), which also exhibits three bands. The lifetime of the $n\pi^*$ state in thymidine monophosphate is 130 ps,43 explaining why it appears as a constant background in our measurements. We have not found any reference in the literature on the broad band TA spectra of $n\pi^*T$ or $n\pi^*G(La)$ states.

We assign the peak at 530 nm decaying with a time constant of 5.4 ps to a CT state. To obtain a rough evaluation of the CT state spectrum, we can consider an equimolar mixture of the guanosine radical cation and the thymidine radical anion, whose TAS have been obtained, respectively, by ns flash photolysis44 and pulse radiolysis.45 The resulting TAS, shown in Fig. S7b,† presents an absorption band around 505 nm. The overlap with that of the CT state of TG is not expected to be perfect for three reasons. (i) Not an entire atomic charge is transferred in the CT state (Table S5†), and as the amount of the transferred charge increases, the spectrum of the CT state shifts toward shorter wavelengths (Fig. S6b†). (ii) In a previous experimental and theoretical study, we reported that the spectral shape of the CT state in dinucleotides is affected by their polarity. 13 (iii) In pulse radiolysis experiments electrons give rise to additional transient species absorbing in the UV. The spectrum computed for the min-CT of anti-syn TG is shown in Fig. 6b, while the experimental TAS at 5 ps, when the contribution from localized $\pi\pi^*$ states has practically vanished (Fig. 3a) in Fig. 6a; the comparison between the two spectra is improved after subtraction of the long-lived background PA at 40 ps. The 5 ps TAS spectrum exhibits an intense band at 380 nm and a second one in the visible, in good agreement with the computed min-CT spectrum (Fig. 6b).

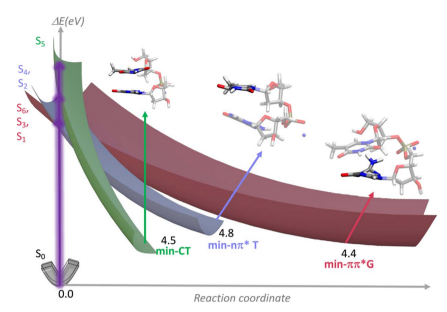


Fig. 5 Schematic representation of the *PES* associated with the 6 Franck-Condon states of *anti-syn* **TG** leading to three minima, whose structure and relative energy ΔE (eV) are also shown.

The measured lifetime of 5.44 ± 0.03 ps is about half of that reported for 5'-dGpdT-3' (10–14 ps).^{11,12} This is not surprising because the DNA polarity affects the excited state relaxation.¹³ Thus, the lifetime of the *CT* state in 5'-dApdG-3' (112 ps) was found to be shorter than that in 5'-dGpdA-3' (170 ps).¹³

The build-up of a small PA band at \sim 445 nm observed in Fig. 3b and its disappearance within 2–3 ps (Fig. 3c) are assigned, respectively, to T \rightarrow G ET and to the decay of the $\pi\pi^*G(La)$ state. Indeed, the dT TAS does not exhibit such a band (Fig. S4†), while it is present in the 50 fs TAS of dG. A previous theoretical study assigned this spectral feature to a minimum in the PES of the first excited state of dG. Although a build-up at this wavelength is also observed for the free dG, it is significantly smaller compared to that found for TG (Fig. 7a). Moreover, the maximum intensity is reached within 60 fs for dG, as compared to 90 fs for TG. These build-up times correspond to relaxation processes and are not limited by the time resolution, as attested by Fig. S1.†

According to our calculations, population of the min-CT, whose PA bands are located at shorter wavelengths with

respect to those of the initial Franck–Condon state S_5 (Fig. S6b†), is expected to induce a blue shift, in line with what is observed in Fig. 3b. The associated non-exponential dynamics (Fig. 4a), is explained by the fact that the process takes place within the restricted geometry defined by the *PES*. We remark that, in these spectral and temporal ranges, the *TAS* recorded for an equimolar mixture of mononucleotides do not exhibit any significant change either in the position or in the intensity of their maximum (Fig. S9†).

The slower blue shift of the low-energy band (Fig. 3c and 4b) is explained by the decay of the $\pi\pi^*G(La)$ state. This is better illustrated by overlaying the position of λ_{max} between 0.2 and 2.5 ps with the dG decay averaged within the considered spectral range, 530–560 nm (Fig. 7b). It is further supported by the similar time constants derived by fits with mono-exponential functions of the spectral shift (0.61 \pm 0.08 ps; Fig. 4b) and the dG decay (0.64 \pm 0.03 ps). We stress that the latter value is not representative of the entire dG dynamics, which is very complex; $^{37-39,46,47}$ it simply reflects its behaviour within the

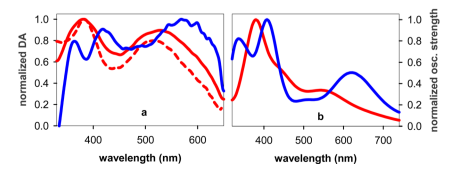


Fig. 6 Comparison between experimental *TAS* recorded for TG (a) with those computed for anti-syn TG (b); (a) *TAS* at 40 ps (blue) and 5 ps (red); the dashed red line corresponds to the difference between the *TAS* at 5 and 40 ps. (b) min-n π *T (blue) and min-CT (red).

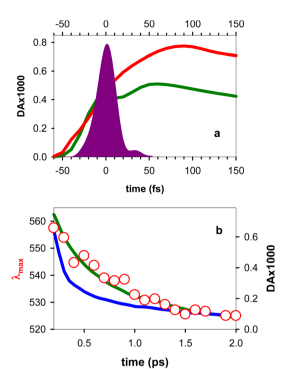


Fig. 7 (a) Rise of the *TA* signals at 445 nm recorded for **TG** (red) and a dG solution with equivalent concentration (green) at 445 nm; the temporal profile of the laser pulse is shown in violet. (b) Comparison of the dynamics observed for the shift of the low-energy band in the **TG** *TAS* (red circles) with the decays of dG (green) and dT (blue), averaged over the spectral range of the shift (530–560 nm).

spectral and temporal limits corresponding to Fig. 7b, where it is also clear that the dT decay follows a different pattern.

So far, we discussed the experimental TAS in the light of the relaxation pathways computed for anti–syn **TG**. Now we examine whether the other less stable conformers could be responsible for the three excited state PES minima that have been associated with the observed spectral evolution. The min-n π *T is reached in all of them. The min-CT is encountered only for syn-anti **TG**.

The pathway corresponding to $T \rightarrow GET$ is detected for the *syn-syn* conformer.

Although anti–syn is the most stable **TG** conformer in the ground state and the fingerprints corresponding to the minima of its PES are detected in the experimental TAS, we cannot rule out that the other conformers are also present in solution, albeit with a much lower concentration. Similarly, it is quite possible that we have a contribution from unstacked nucleobases, mainly characterized by localized excited states; yet the lack of a negative signal due to the SE of $\pi\pi^*T$, which does not correspond to a PES minimum in anti–syn **TG**, indicates that this contribution is rather low.

Global analysis

The multitude of the above discussed processes, associated with 3 minima in the 3 PES of each conformer and 2 for the unstacked nucleobases, giving rise to a complex network of excited states, is one reason why we have not based our analysis on global fits. The second and most important reason for avoiding analysis with exponential functions over the entire time domain explored in our experiments is related with the physics of the processes. As mentioned in the introduction, an excited state decays exponentially when its interactions with the environment are uniform within the three-dimensional space. In low-dimensional spaces the dynamics tend to be slower, described, for example, by stretched exponentials or exhibiting power-law patterns.^{25,48} To correctly describe the dynamics in a given system, specific theoretical analysis and/or numerical simulations are required. We stress that the power-law function used to fit the spectral shift in Fig. 4a is a phenomenological approach, simply showing that the dynamics of the process is not exponential.

In any case, we did perform a global analysis within the framework of sequentially evolving species. Fig. S10† reports the evolution associated spectra (*EAS*) for an equimolar mixture of dT and dG. Five time constants (120 fs, 250 fs, 800 fs, 2.1 ps and 2 ns) are needed to fit the data, numerically close to those

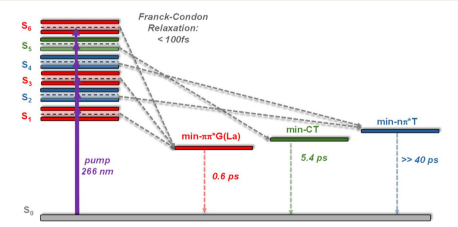


Fig. 8 Schematic drawing of the Franck–Condon excited states and the corresponding minima computed in the *PES* of *anti–syn* **TG**. In italics: dynamics determined from the *TAS* evolution. The relaxation time leading to min- $n\pi$ *T has not been observed, because the associated signal is too weak and blurred by those corresponding to other excited states; the 0.6 ps indicated for the decay of min- $\pi\pi$ *G(La) corresponds only to the minimum absorbing in the red.^{37–39,47}

reported by Kwok and co-workers for dT (150 fs and 760 fs)⁴⁰ and dG (200 fs, 790 fs and 2.0 ps)³⁸ using a lower temporal resolution. However, the similar time evolution of the individual nucleosides does not allow for their spectral discrimination within the studied equimolar mixture. As a result, instead of identifying distinct spectral signatures for each nucleoside, an average spectrum is obtained.

In the case of **TG**, the use of 4 exponential functions (50 fs, 380 fs, 4.5 ps and 2 ns) provides acceptable fits (see *EAS* in Fig. S11†). The spectral features corresponding to the time constants of 4.5 ps and 2 ns resemble those attributed to min-CT and min-n π *T (Fig. S12†) but the evolution on shorter times, in particular, that corresponding to the complex decay of $\pi\pi$ *G(La), is not correctly resolved.

Conclusion

This article reports an important breakthrough in the study of photoinduced excited state relaxation in coupled nucleobases in solution. The novelties concern both the approach and the obtained results. Two main methodological innovations need to be underlined. On the one hand, this is the first observation of transient absorption spectra and dynamics on the sub-100-fs timescale. It was rendered possible by the exceptional combination of sensitivity and temporal resolution of our femtosecond TA setup. On the other hand, the interpretation of the results has been based on a battery of quantum chemistry calculations on the most stable geometry in the ground state, the Franck-Condon states, their evolution toward the excited state minima and the corresponding TAS. In this way, we circumvented the global fits with exponential functions, which are inadequate to describe evolution along potential energy surfaces.

Several compelling results emerged from our investigation.

- (i) Our computations showed that the most stable conformation of TG involving stacked nucleobases (anti–syn) differs from that adopted by the T/G step in B-form duplexes (anti–anti), which has been implicitly considered for the interpretation of previous femtosecond measurements on dinucleotides. ^{11,13} The anti–syn conformation is encountered in G-Quadruplexes. ³⁰
- (ii) The minima corresponding to relaxation pathways computed for the *anti–syn* **TG**, summarized in Fig. 8, are detected in the experimental *TAS*.
- (iii) According to *TA* measurements, the *CT* state is reached within 100 fs, following non-exponential dynamics and decays with a lifetime of 5.4 ps. This the shortest lifetime ever determined for DNA dinucleotides, ⁸⁻¹⁴ and, probably, for any donor-acceptor pair of aromatic chromophores.
- (iv) The *PES* leading to min- $\pi\pi^*G(La)$ corresponds to T \to G *ET*. The associated spectral evolution occurs within 90 fs. Such an ultrafast *ET* could be facilitated by a very small *CT* between the nucleobases characterizing the initial Franck–Condon state (Table 1). So far, ultrafast *ET* in DNA was considered to proceed exclusively *via* exciton states.²³
- (v) Several detailed studies, dealing with monomeric pyrimidines, reported population of $n\pi^*$ states. ^{43,49,50} But so far, no

experimental studies identified such states in dinucleotides or larger DNA multimeric systems composed of the major nucleobases.

The use of quantum chemistry calculations allowed a qualitative assignment of the dominant processes. In this way, we circumvented the global fits with exponential functions which, although they bring precious information on longer timescales, are inadequate to describe wave packet evolution along *PES* and/or systems involving a large number of transient species. Yet, the question of dynamical models that are appropriate to describe these processes remains open and requires specific work. We hope that our experimental results will inspire the future development of less time-consuming methods than those currently available that can be applied on large molecular systems.

Finally, an interesting perspective is to examine whether the ultrafast $T \to G$ energy transfer, detected in TG, also occurs in G-Quadruplexes. If this were the case, positioning peripheral thymines at the 5' end with respect to the guanines of the G-core should further enhance their fluorescence. Such an enhancement is important because the intrinsic fluorescence of G-Quadruplexes is currently scrutinized in view of biosensors development. 3,51,52

Methods

Transient absorption setup

Ultrafast TA experiments were conducted using a custom-built setup, employing an amplified Ti:Sapphire laser (Coherent Libra) that generates 100 fs pulses at a central wavelength of 800 nm, operating at a repetition rate of 1 kHz.31,53 A fraction of the fundamental laser beam is first frequency-doubled and then directed to drive a broadband visible non-collinear optical parametric amplifier (NOPA). The output of the NOPA consists of broadband visible pulses, which are subsequently using chirped dielectric mirrors. compressed compressed pulses are further frequency-doubled in a 20-µmthick β-barium borate crystal, resulting in broadband UV pump pulses that can be tuned within the range of 250-300 nm. The generated UV pump pulses, fully characterized through two-dimensional spectral interferometry, are compressed to 24 fs (FWHM) duration with the help of a prism pair⁵⁴ (Fig. S13†). For the purpose of the experiment, the UV pump pulses are tuned to 266 nm. Broadband probe pulses, with spectrum spanning from 320 nm to 650 nm, were obtained by white-light continuum generation, focusing a portion of the laser fundamental beam onto a slowly moving 2-mm-thick CaF2 plate. Pump and probe pulses are non-collinearly focused on the sample under study, with spot sizes of 180 μm and 95 μm, respectively. The relative pump and probe polarizations are adjusted to the magic angle (54.7°). The pump fluence is kept at 100 μJ cm⁻², resulting in differential absorption (DA) signals lower than 10⁻³; in this way, the coherent artefact and solvated electrons stemming from two-photon ionization of the solvent are minimized. The transmitted probe spectrum is recorded by an optical multichannel analyser, operating at the full laser repetition rate. The temporal resolution is estimated to be sub**Edge Article**

30-fs. At 20 fs, although spectral intensities are not fully developed, spectral shapes in the visible are meaningful. During the experiment 6 mL of the solution kept flowing through a 1-mmthick quartz cell controlled by a pump. For a detailed discussion of the temporal resolution in the UV TA experiments, see Section 1 in the ESI.†

Key experimental points

Dinucleotides are difficult to purify and residual mononucleotides may "contaminate" their proprieties. Therefore, purchased from Eurogentec, TG was purified by reverse phase HPLC and tested by MALDI-TOF (Fig. S14†). It was dissolved in phosphate buffer ($\sim 0.12 \text{ mol L}^{-1}$, pH 7.0). The concentration of absorbed photons (8 \times 10⁻⁶ mol L⁻¹) was much lower than the dinucleotide concentration ($\sim 4 \times 10^{-3}$ mol L⁻¹), precluding two-photon absorption. Experiments on the monomeric chromophores were performed using the mononucleosides dT and dG in water, instead of the corresponding mononucleotides, which have a great propensity to aggregate in solutions containing salts.55

Computational details

We used quantum mechanical (QM) calculations, based on the Density Functional Theory (DFT) with the M052X functional,56-58 the 6-31G(d) basis set. The solvent effect was taken into account via an implicit polarizable continuum model (PCM),59 which was shown to reproduce experimental trends in a satisfactory way. 60,61 In our computations we included one Na+ ion per dinucleotide so as to keep the system electrically neutral.

The vertical absorption energies of the different excited states and the corresponding PES were characterized using the above-described methodologies, but resorting to the timedependent version (TD-DFT). The charge transfer character was computed by a simple Mulliken population analysis in terms of δq , i.e., the difference between the charges in the excited state and in the ground state.

For the TA spectra, the transition dipole moments between the excited states were obtained by a multifunctional analyser (multiwfn program).62 For comparison with the experimental spectra, the computed ones were shifted by -0.65 eV. Such a shift, typical for this level of calculation, matches the difference between the energy computed for the lowest bright state (La) of dG in water and the corresponding value derived from deconvolution of the experimental spectrum (Fig. 2 in ref. 63). Subsequently, we applied a phenomenological broadening via a Gaussian function with half width at half maximum of 0.2 eV.

Data availability

The data that support the findings of this study are available in the ESI† and the on: https://doi.org/10.5281/zenodo.11453876. They are also available upon request from the corresponding authors.

Author contributions

Conceptualization: LMF, GC, DM; data curation: VP, LMF; formal analysis: VP, LMF, EB, DM; funding acquisition: GC, MM, DM; investigation: VP, LMF, LU; methodology: LMF, GC, MM; supervision: GC, MM; validation: all authors; visualization: VP, LMF, EB, DM; writing original draft: VP, LMF, DM; writing review & editing: VP, LMF, GC, EB, DM.

Conflicts of interest

There are no conflicts to declare.

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