Supplementary Information

"Tracking excited state decay mechanisms of pyrimidine nucleosides in real time"

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Supplementary Note 1:
Bibliographic survey on the proposed photoinduced decay mechanisms in pyrimidines

Over the past years many experimental studies have been performed to shed light in the fate of photoexcitations in DNA/RNA [Mid09, Koh10, Cre04, Sai07, Che14], which has relevant implications in the preservation of genetic code. Pioneering transient absorption (TA) spectroscopy studies by Kohler’s group [Pec00, Pec01] revealed sub-picosecond decay times for nucleosides in water. These results were further confirmed by femtosecond fluorescence up-conversion (FU) measurements [Peo01, Gus11]. The decay channels of pyrimidines and the involvement of $n\pi^*$ excited states are still the matter of a heated debate and several hypotheses exist [Mid09, Cre04, Imp16].

While most theoretical studies in gas phase assign the lowest excited state (ES) to a $n\pi^*$ transition in uracil (Ura), 5-methyl-uracil (5mUra), also known as thymine (Thy) and their respective nucleosides uridine (Urd) and 5-methyl-uridine (5mUrd), there is a general agreement that the solvent modulates the interplay between the $\pi\pi^*$ and $n\pi^*$ states, destabilizing the $n\pi^*$ state with respect to the bright $\pi\pi^*$. Due to the dark nature of the $n\pi^*$ state, it is not easy to obtain a direct experimental proof of its energy shift in order to assess its involvement in the ES deactivation [Imp16, He04].

Among the five solvated nucleosides, pyrimidines present longer lifetimes than purines [Oni02, Har07, Imp16, Gus02] Usually, longer time constants have been reported for 5mUra and 5mUrd, compared to Ura and Urd [Pec01, Peo01, Gus02, Har07, Gus06, Pol14, Kwo08, Pil18, Xue16, Kob12, Pro16]: this fact is in itself intriguing, given the close molecular structure and similar potential energy surfaces of both systems[Pep17]. Experimental TA and FU spectroscopy measurements agree on describing their excited state decays with one or two sub-picosecond components and a further component of several picoseconds [Pec01, Peo01, Gus11, Gus02, Oni02, Har07, Gus06, Pol14, Xue16, Kob12, Pro16, Coh04, Buc15]. Additionally, significantly longer lifetimes have been reported for pyrimidine nucleosides with respect to free bases, suggesting that in pyrimidine systems sugar substitution could alter significantly the lifetime of an excited nucleobase [Con18, Har07].

In this complex context, three hypotheses have been proposed for the assignment of the experimental time constants.
1.1 Direct internal conversion (IC) to the ground state ($\pi\pi^* \rightarrow S_0$)

Early works assigned a unique time constant of $\sim$600-700 fs to internal conversion [Peo01, Pec01]. Improvements in the temporal resolution have meanwhile revealed an additional time constant as short as $\sim$100 fs in all pyrimidines [Buc15, Can05, Bri19, Che13, Pep17, Gus06, Har07, Oni02] suggesting that ICs may occur much faster than originally anticipated. In support of the $\pi\pi^* \rightarrow S_0$ direct IC hypothesis, theoretical studies have disclosed a conical intersection (CI) seam characterized by ring-puckered structures with hydrogen (Ura, Cyd), methyl (5mUra), amino (Cyd) and/or carbonyl out-of-plane bent groups [Buc15, Nak13, Har08, Pep18].

A further $\pi\pi^* \rightarrow S_0$ decay channel has been hypothesized in the case of Ura and Thy [Pep17, Ric14, Nac11, Bar10, Gha18, Bar13]: a ring opening mechanism characterized by the $N_1-C_2$ bond breaking, which could potentially lead to photoproducts formation.

Recent molecular dynamics simulation of 5mUrd/5mUra in water support the hypothesis that the entire population follows the $\pi\pi^* \rightarrow S_0$ decay path (including both ring puckering and carbonyl bending mechanisms) in the sub-picosecond time range [Buc15, Nak13, Har08], assigning the few ps lifetime to the dissipation of the excess of energy of the electronic ground state to the solvent, by vibrational cooling. The $n\pi^*$ state is found to be strongly destabilized and conversely does not play any role in the decay path, in contrast to what happens in gas phase. The hypothesis is further supported by recent liquid jet time-resolved photoelectron spectroscopy in combination with XMS-CASPT2/CASSCF/AMBER calculations, providing a time constant for the $\pi\pi^*$ decay of ca. 400 fs [Eri19].

1.2 IC to an intermediate dark state ($\pi\pi^* \rightarrow n\pi^*$)

Although the involvement of a dark state of $n\pi^*$ nature is very probable for pyrimidines in gas phase, as confirmed both in experimental [He03, He04, Yu16, Wol19] and theoretical [Imp16, Sto16, Per06, Mai17, Wol19, Par19] studies, the role of dark states in condensed phase is still controversial. According to the hypothesis of the involvement of the $n\pi^*$ state in solution, the shortest time constant is to be assigned to the CI mediated decay to the intermediate dark state, with lifetimes ranging from few hundred femtoseconds to few hundred picoseconds [Kob12, Pro16, Xue16, Coh04, Bri19, Cer18, Liu18].

Regarding the deactivation of the $^1n\pi^*$ state, currently three mechanisms are hypothesized:
1.2.1 Direct IC to the ground state \( (n\pi^* \rightarrow S_0) \)

Occurs on sub-ps to few ps time scale through an energetically higher lying \( ^1n\pi^*/S_0 \) CI seam [Bri19, Lis18].

1.2.2 Decay to a triplet state \( (n\pi^* \rightarrow T_1 \rightarrow S_0) \)

Occurs on a sub-ps to ps time scale, facilitated by the energetic proximity of states of different multiplicity in the vicinity of the \( ^1n\pi^* \) minimum and the large spin orbit couplings (mainly with the \( ^3\pi\pi^* \)), in agreement with El-Sayed’s rule [Kwo08, Wol19, McN97]. The triplet itself is proposed to decay to the ground state on a ps to ns time scale [Pil18]. Support for the formation of the triplet state has been provided by dynamics calculations for Ura. [Ser07, Ric14, Gon10]

1.2.3 Re-population of the \( \pi\pi^* \) state \( (n\pi^* \rightarrow \pi\pi^* \rightarrow S_0) \)

Occurs on a tens to hundreds of ps time scale. Support for this mechanism is provided by a recent computational work [Con18] on stacked thymidines which shows that the \( \pi\pi^* \leftrightarrow n\pi^* \) exchange could involve a molecular distortion very sensitive to the sugar substitution, thus potentially rationalizing the longer lifetimes observed in nucleosides, compared to the nucleobases.

1.3 ISC from the bright \( \pi\pi^* \) to an intermediate triplet \( ^3n\pi^* \) state \( (\pi\pi^* \rightarrow ^3n\pi^*) \)

This decay process has been documented recently for the solvated uridine monophosphate [Bri19] on the basis of femtosecond broadband TA spectroscopy experiments. A trifurcation process from the bright \( \pi\pi^* \) state is proposed, where the system decays within 200 fs simultaneously to the ground state, to the singlet \( n\pi^* \) and to the triplet \( ^3n\pi^* \) state which subsequently funnels the population to the energetically lower lying \( ^3\pi\pi^* \) triplet state, which survives on the ns time scale.
Supplementary Note 2:
Theoretical background and protocols for acquisition of simulation parameters

2.1 Classical molecular dynamics

Classical molecular dynamics simulations for Urd and 5mUrd dissolved in water were performed with the Amber 12 suite of programs [Fer13,Cas12] utilizing the Parm99 force field and the all_nucleic02.lib libraries for RNAs. Periodic boundary conditions were applied to a periodic box of water molecules described by the TIP3P force field [Jor83]. The hydrogen-containing bonds were restrained by the SHAKE algorithm [Ryc77] while the water geometry was rigidized by the SETTLE scheme [Miy92]. Non-bonding and electrostatic interactions were evaluated with a cutoff of 9.0 Å making use of the particle mesh Ewald method for quantification of the long-range electrostatics. The whole system, including the nucleoside moiety, was equilibrated from 0 to 300°K for 1 ns while maintaining a constant volume and pressure (1 atm). A production run was then performed for 100 ns recording snapshots every 200 fs. Analysis of the geometrical changes along the trajectories is presented in [Pep17].

It is well known that in mono-nucleosides two predominant conformers, syn and anti, can be populated. A cluster analysis performed along the trajectories gives as most probable geometry a syn conformation that is stabilized by an intramolecular hydrogen bond between the CH$_2$OH group on the sugar and the C=O group in the nucleobase. To select the initial geometry for subsequent QM/MM refinement and generation of initial conditions we performed a cluster analysis based on a Root Mean Square (RMS) coordinate deviation analysis on the nucleoside moieties over all snapshots recorded along the dynamics run within a 1.5 Å difference using the MMTSB toolbox of Amber 12 [Fei01]. We obtained two clusters, denoted A and B, with syn/anti populations relating to 73/27 and 62/38 for Urd and 5mUrd, respectively. The snapshot with the geometrical parameters closest to the centroid (i.e., the mean position) of the structures obtained in cluster A, was selected for further QM/MM refinement.

2.2 QM/MM geometry refinement

QM/MM calculations were executed with the COBRAMM suite, developed in our group interfacing various QM codes with the MM software AMBER.
Supplementary Figure 1: High/Medium/Low layer partitioning of Urd (left) and 5mUrd (right) in the QM/MM refinement protocol. The High layer comprises the 5mUra, the Medium layer comprises the sugar and all waters in 5 Å distance from the center of mass of the nucleobase (gray color). The rest of waters are kept frozen in the Low layer.

[Alt07,Wei18]. A High Layer (HL) / Medium Layer (ML) / Low Layer (LL) partitioning was applied to a spherical droplet centered at the nucleoside with a radius of 12 Å (containing 260 waters), obtained from the cubic box (Figure 1). The HL (QM region) comprises the nucleobase. The water molecules in 5 Å distance from the center of mass of HL were included in the movable ML. The remaining water molecules were kept fixed in the LL. The ground state minimum was obtained at the Møller-Plesset second order perturbation theory (MP2) level as implemented in the Molcas 8 package [Aqu16] through its interface with COBRAMM. The ANO-L basis set was used [Wid90], adopting a valence double-ζ contraction (i.e. 3s2p1d for non-hydrogen and 2s1p for hydrogen atoms).

2.3 Wigner sampling

For each system 500 initial conditions were generated using the Wigner sampling technique based on a frequency calculation at the MP2 optimized structure through an interface with a stand-alone script, part of the quantum molecular dynamics program JADE [Du15], which considers temperature effects and mode-specific excitation. Consequently, sampling was performed at room temperature (300 K). High frequency modes, belonging to C-H and N-H stretchings, were excluded from the sampling. In the case of 5mUrd two
normal modes with low frequencies (< 100 cm\(^{-1}\)) involving methyl group
twisting along the CH\(_3\)-C\(_5\) \(\sigma\)-bond were excluded from the production sam-
pling as it was ascertained through preliminary sampling that displacement
along such modes breaks the harmonic approximation and distorts exces-
sively the geometry. Each of the sampled geometries was subjected to a 10
ps long MM equilibration run in which the QM region was kept fixed through
harmonic constraints. This strategy allows the mobile MM region to adapt
to each sampled geometry, thereby removing the bias of the water arrange-
ment of the initial snapshot. The water coordinates and velocities of the last
snapshot of the MM equilibration run were then combined with the coordi-
nates and velocities of the sampled nucleoside geometry to form a snapshot
used to initiate the mixed quantum classical dynamics simulations.

2.4 Linear absorption spectroscopy

The electronic spectrum of two systems was computed at the CASSCF/CASPT2
level with the Molcas 8 software package through its interface with CO-
BRAMM. For the active space size dependence tests the full valence-\(\pi\) active
space comprising 10 electrons in 8 orbitals (i.e. CAS(10,8)) was extended
by four and eight virtual \(\pi\)-orbitals, respectively. The restricted active space
modification of the CASSCF protocol (i.e. RASSCF) allowing for up to four
simultaneous excitations was used. In the following we will refer to these ac-
tive spaces as RAS(4,5|0,4,7) and RAS(4,5|0,0|4,11)\(^1\). The excited state
computations were averaged over two and four states, respectively. Addition-
ally, the CAS(10,8) and RAS(4,5|0,0|4,7) were augmented by the two lone
pairs of the oxygens to disentangle the effect of the \(n\pi^*\) transitions on the lin-
ear absorption (LA) spectra, thus giving CAS(14,10) and RAS(4,7|0,0|4,7).
In these cases the excited state computations were averaged over six states.
The CASPT2 computations were performed both in the single-state
and multi-state flavor. All calculations were performed with the generally con-
tracted basis set ANO-L adopting valence double-\(\zeta\) contractions.

For each system the spectra were computed utilizing the transition ener-
gies and dipole moments computed at 500 snapshots generated via Wigner
sampling according to the formula

\[ \sum_{\text{states}} \frac{\text{transition energy}}{\text{dipole moment}} \]

\(^1\)In the RASSCF method the molecular orbitals and electrons are divided into three
active spaces: RAS1—RAS2—RAS3. We adopt the following notation: \((i,j|k,l|n,m)\) with
\(i\) and \(j\) the maximal number of holes and occupied orbitals in RAS1, \(k\) and \(l\) the number
of electrons and orbitals in RAS2, and \(n\) and \(m\) the number of simultaneous excitations
and virtual orbitals in RAS3.
\[ I(\omega) \propto \frac{1}{N_{\text{traj}}} \sum_{k} \sum_{e} \mu_{eg}^{2}(R_{k})g(\Delta E_{eg}(R_{k}) - \hbar \omega) \]  \hspace{1cm} (1)

where \( k \) iterates over the number of realizations, \( R_{k} \) denotes the geometrical coordinates and \( f \) denotes a state from the manifold of excited states. The intensity is convoluted with a normalized line shape function \( g(\Delta E_{eg}(R_{k}) - \hbar \omega) \) peaked at the transition energies and broadened by a phenomenological constant \( \sigma \):

\[ g(\Delta E_{eg}(R_{k}) - \hbar \omega) = \exp \left( -\frac{(E^{(e)}(R_{k}) - E^{(g)}(R_{k}) - \hbar \omega)^{2}}{2\sigma^{2}} \right) \]  \hspace{1cm} (2)

For \( \sigma \) we chose a value of 0.1 eV to match the experimental LA spectra. State-specific spectra were generated by grouping transitions according to the leading configuration of the ES wave function (i.e. \( n \rightarrow L \) for \( n\pi^{*} \) and \( H \rightarrow L \) for \( \pi\pi^{*} \)). For the purpose of the discussion we use the terms configuration and state interchangeably.

### 2.5 Non-adiabatic mixed quantum-classical dynamics simulations

For each system, out of the 500 snapshots generated via Wigner sampling, 60 geometry realizations were selected for running mixed quantum-classical molecular dynamics simulations with the condition that the transition energy of the lowest \( \pi\pi^{*} \) state computed at the SS-CASPT2/SA-4-RASSCF(4,5|0,0|4,11) level falls under the envelope of the pump pulse, as this level is shown to give excellent agreement with experiment (see Figure 11e). Molecular dynamics simulations following Newton’s equations of motion for the nuclei and utilizing hybrid QM/MM gradients were performed at the full-\( \pi \) SS-2-CASPT2/SA-2-CASSCF(10,8) level of theory for 500 fs in the case of Urd and for 1000 fs in the case of 5mUrd with a time step of 1.0 fs applying the Tully’s fewest switches surface hopping algorithm [Tul90] with the Tully-Hamnes-Schiffer (THS) scheme [Ham94] using the COBRAMM’s parallel environment for computing numerical gradients. The state averaging covered the ground and the lowest \( \pi\pi^{*} \) state (\( S_{1} \)). The movable (MM) water layer was extended (now comprising all water molecules within 5 Å from the HL) to accommodate distortions of the aromatic ring such as puckering, as predicted by static calculations [Pep17].

Gradients were computed numerically through a two-point finite differentiation formula using the COBRAMM’s parallel environment. To speed up the numerical computations of the SS-RASPT2 gradients we made use of the
fact that the perturbative correction is applied independently on each root. Thus, while at the reference point all states are corrected at SS-RASPT2 level, during the computation of the gradient only the state that drives the nuclear dynamics (i.e. the photoactive state) was taken into account. Care was taken for possible state swapping at each displaced geometry in case of near-degeneracy at the RASSCF level by following the nature of the wave function. This was facilitated by computing the wave function overlap between reference and displaced geometries within the RASSI utility [Mal86] of Molcas. The stability of the simulations was monitored by COBRAMM. If

a) an element the numerical gradient exceeds a threshold (default 0.5)

b) the ratio of the lengths of the gradient vectors at two consecutive steps (i.e. $||g(t)||/||g(t-dt)||$) and, at the same time, the deviation of the total energy from its value at the previous step (i.e. $E_{tot}(t-E_{tot}(t-dt))$) exceed certain thresholds (defaults 1.3 and 0.5 kcal/mol, respectively)

c) a single point computation fails to converge

the gradient is discarded and the propagation of the nuclei is repeated at the last stable geometry, first with a half step (0.5 fs) and, in case the gradient is discarded again, with a double step (2.0 fs).

Following the THS scheme the expression for the time-derivative coupling (TDC) between states $i$ and $j$

\[
\sigma_{ij} = \frac{dR}{dt} d_{ij} = \frac{dR}{dt} \langle \Psi_i(r,t) | \frac{d}{dt} \Psi_j(r,t) \rangle = \frac{\langle \Psi_i(r,t) | \frac{d}{dt} \Psi_j(r,t) \rangle}{TDC} \tag{3}
\]

is approximated to finite differences and the change in the electronic wave functions is resolved by computing overlap integrals between the adiabatic wave functions at different time steps. We generalized the TDC formulation of Barbatti (Eq. 9 in ref. [Pit09]) to arbitrary time steps $\Delta t$:

\[
\sigma_{ij} = \frac{1}{2\Delta \tau} \left( 1 - \frac{0.5 \Delta \tau + \Delta t}{0.5 \Delta \tau + 0.5 \Delta t} \right) \left( \langle \Psi_i(t-\Delta t) | \Psi_j(t-\Delta t-\Delta \tau) \rangle - \langle \Psi_j(t-\Delta t) | \Psi_i(t-\Delta t-\Delta \tau) \rangle + \langle \Psi_j(t-\Delta t) | \Psi_i(t-\Delta t) \rangle - \langle \Psi_i(t) | \Psi_j(t-\Delta t) \rangle \right) \tag{4}
\]

where $\Delta t$ and $\Delta \tau$ are the time steps used to propagate the time-dependent Schrödinger equation between step $k-1$ and $k$, and between step $k-2$ and $k$.
1, respectively. Note that eq. 4 simplifies to eq. 9 in ref. [Pit09] when $\Delta \tau = \Delta t$. The individual terms are obtained from wave function overlaps, which are computed at the RASSCF level (i.e. using the RASSCF wavefunctions) through the RASSI utility [Mal86] of Molcas and further scaled by the ratio of the SS-RASPT2 and RASSCF energy gaps $(E_{j}^{\text{CASSCF}} - E_{i}^{\text{CASSCF}})/(E_{j}^{\text{SSPT2}} - E_{i}^{\text{SSPT2}})$, which follows from the relation:

$$
\sigma_{ij} = \langle \Psi_{i}(t)^{\text{SSPT2}} \mid \frac{d}{dt} \Psi_{j}(t)^{\text{SSPT2}} \rangle = \frac{dR}{dt} \langle \Psi_{i}(t)^{\text{SSPT2}} \mid \frac{d}{dR} \Psi_{j}(t)^{\text{SSPT2}} \rangle \approx \frac{v \cdot d_{ij}^{\text{SSPT2}}}{E_{j}^{\text{SSPT2}} - E_{i}^{\text{SSPT2}}} = \frac{v \cdot \langle \Psi_{i}(t)^{\text{CASSCF}} \mid \nabla_{R} \hat{H} \mid \Psi_{j}(t)^{\text{CASSCF}} \rangle}{E_{j}^{\text{CASSCF}} - E_{i}^{\text{CASSCF}}} \cdot \frac{E_{j}^{\text{CASSCF}} - E_{i}^{\text{CASSCF}}}{E_{j}^{\text{SSPT2}} - E_{i}^{\text{SSPT2}}} =
$$

(5)

In the above derivation following equality is assumed:

$$
\langle \Psi_{i}(t)^{\text{SSPT2}} \mid \nabla_{R} \hat{H} \mid \Psi_{j}(t)^{\text{SSPT2}} \rangle \approx \langle \Psi_{i}(t)^{\text{CASSCF}} \mid \nabla_{R} \hat{H} \mid \Psi_{j}(t)^{\text{CASSCF}} \rangle
$$

(6)

Effectively, when the SS-RASPT2 correction to the SA-CASSCF energies decreases the energy gap between a pair of states, the value of the time-derivative coupling is increased uniformly and linearly. In addition, a wave function following algorithm allows to identify state swapping at the SS-CASPT2 level. The THS scheme allows to work with relatively large time steps (such as 1.0 fs) and performs better in the region of $\pi\pi^{*}/n\pi^{*}$ near-degeneracy compared to the original Tully scheme based on non-adiabatic coupling vectors.

Tully’s fewest switches surface hopping algorithm is known to suffer from artificial coherence effects. A decoherence correction, originally proposed by Truhlar and co-workers [Zhu05] and realized by Persico et al. [Gra07] is used in the present case as a countermeasure. Kinetic energy scaling (for total energy conservation) after a hopping event is performed along the velocity vector.

2.6 Critical appraisal of the method of choice

Our tests showed that the pronounced $n\pi^{*}/\pi\pi^{*}$ wavefunction mixing at the CASSCF level makes dynamics simulations considering oxygen lone pairs in the active space unstable both at single state (SS) and at the multi state
(MS) levels, the cause lying in the numerical calculation of the energy gradients, exhibiting non-physically large energy changes (and, hence, forces) for infinitesimal geometry displacements in regions of strong wavefunction mixing at the CASSCF level. Despite being able to substantially improve the stability of the simulations with the aforementioned sanity checks, e.g. through doubling the time-step in order to over-step the regions of strong mixing (see Supplementary Note 2.5) it was impossible to maintain energy conservation over the period of several hundred femtoseconds as often strong wavefunction mixing was encountered in extended regions of the potential energy surface, that cannot be overcome even by tripling the size of the time step. Therefore, we were left with no other choice but to exclude the oxygen lone pairs from the active space. The participation of the omitted $n\pi^*$ state in the deactivation mechanism was explored ad-hoc (see Supplementary Note 2.7).

A crucial step in the execution of the simulations is thus to make sure that the oxygen lone pairs do not enter the active space. This was realized in the following way. At the first step of each trajectory it was assured that the active space contains only $\pi$-orbitals. At every following step $t$ a reference CASSCF calculation was run, re-optimizing the molecular orbitals of the previous step $t - \Delta t$ without any restrictions. Subsequently, the nature of each state $i$ was tested by computing the norm of its overlap with the states at the previous time step following the formula $||i|| = \sum_j \langle i(t) | j(t - \Delta t) \rangle$ with elements obtained from the overlap matrix $S(t - \Delta t \rightarrow t)$. If the value for the overlap is above a certain threshold ($75\%$), meaning that the character of the state is essentially conserved, we proceed with the calculation of the CASPT2 energies and gradients. Instead, if the value for the overlap is below the threshold, we assume that the active space was contaminated with a lone pair. This generally happens when the $n\pi^*$ state is considerably stabilized at the CASSCF level (as in the case of elongation of the carbonyl bonds) so that rotating a lone pair at the place of the completely bonding $\pi$-orbital would lower the total state-averaged energy of the system. In such case the CASSCF calculation is repeated once again (using the molecular orbitals of the previous step), this time preventing orbital rotations between the lone pairs and the remaining orbitals. CASPT2 energies and gradients at this time step are then all computed in the same manner. Benchmarking showed that "freezing" the lone pairs for a few steps has practically no effect on the outcome of the dynamics. Moreover, it was found that simulations with continuously "frozen" lone pairs (i.e. forced in a sup-symmetry block) begin to deviate from simulations with "free" lone pairs after about 50 fs.
A set of 50 trajectories was run with both CASPT2 flavors (SS-2-CASPT2/SA-2-CASSCF(10,8) and MS-4-CASPT2/SA-4-CASSCF(10,8)) for Urd (see Figure 18). The comparison shows that, while in the case of SS-CASPT2 the majority of trajectories have decayed to the ground state after 200 fs, the dynamics at the MS-CASPT2 level is considerably slower with half of the trajectories not having decayed within the simulation period. In fact, we found only two cases where the MS-CASPT2 dynamics decays faster to the ground state compared to the SS-CASPT2 dynamics (e.g. Figure 18e). While in some cases single state and multi state simulations agree fairly well (e.g. Figure 18a,b), in most cases we observe that the system roams in the CI region at the MS-CASPT2 level opposite to trajectories at the SS-CASPT2 level which decay in a ballistic fashion (e.g. Figure 18c,d). The reason for this discrepancy is an artifact of the MS-CASPT2 approach with small active spaces that gives rise to relatively large off-diagonal $S_0-S_1$ elements in the MS-CASPT2 Hamiltonian (an order of magnitude larger than the value of 0.002 suggested in the seminal paper on MS-CASPT2 [118]). While not posing a problem in the FC region where the $S_0-S_1$ energy gap is large, these couplings become important in the cases when the two states come energetically close. In a direction in which the CASSCF wave functions mix, the MS-CASPT2 routine over-estimates the level of un-mixing thus pushing the two states apart (e.g. Figure 18c,d), introducing an artificial barrier in the excited state and preventing an efficient decay. To demonstrate the tendency of MS-CASPT2 to overestimate the splitting we document the distribution of energy gaps at the SS-CASPT2 and MS-CASPT2 level at the CIs encountered along the MD simulations in Figure 2. Thereby, we selected the geometry with the lowest $S_1-S_0$ energy gap along each trajectory (which in most cases coincides with the geometry at which the hopping occurs) and computed SS-2-CASPT2 and MS-2-CASPT2 energies with two active spaces, the full-$\pi$ CAS(10,8) used in the simulations and a large RAS(4,5,0,0|4,11) augmented with eight virtual orbitals. It can be seen that, while the energy gap at the SS-CASPT2 level does not change as a function of the active space (Figure 2,a), it is very much affected at the MS-CASPT2 level where the splitting is halved (Figure 2,b). The splitting is directly related to the magnitude of the MS-CASPT2 couplings at both levels (Figure 2,c). The only way for the excited state to decay to the ground state is in the form of a diabatic crossing where the two states do not mix at the CASSCF level but the system has to roam longer on the $\pi\pi^*$ potential energy surface to find such a crossing. Comparison with the experimental transient data clearly shows that the majority of the population has decayed to the ground state within 200 fs and that the dynamics is completed within 500 fs, in better agreement with the SS-CASPT2 results. Finally, we note that, even if the system spends
considerably longer time in the excited state at the MS-CASPT2 level, the $S_2$ state was found to interfere in the dynamics only in a few cases.

Supplementary Figure 2: Comparison between the distribution of energy gaps at the conical intersections encountered along the MD simulations: a) SS-2-CASPT2/CAS(10,8) (red) and SS-2-RASPT2/RAS(4,5|0,0|4,11) (black) levels; b) at MS-2-CASPT2/CAS(10,8) (red) and MS-2-RASPT2/RAS(4,5|0,0|4,11) (black) levels. c) comparison at the MS-CASPT2 couplings between $S_0$ and $S_1$ at the MS-2-CASPT2/CAS(10,8) (red) and MS-2-RASPT2/RAS(4,5|0,0|4,11) (black) levels.
2.7 How to estimate the $n\pi^*$ involvement

For the reasons outlined in the previous section we could not obtain energy conservation during the molecular dynamics simulation when the oxygen lone pairs were included in the active space. Therefore, these simulations cannot directly shed light on the involvement of the $n\pi^*$ state in the ultrafast (sub-ps) non-adiabatic dynamics. However, it is well known that the lowest $\pi\pi^*$ and $n\pi^*$ states are energetically close and, as elucidated in the main text, there is still a controversy in the literature regarding the importance of the $n\pi^*$ state for trapping excited state population. In order to estimate the percentage of trajectories likely to undergo non-adiabatic population transfer to another excited state before decaying to the ground state we:

- analyzed the extent of $\pi\pi^*/n\pi^*$ mixing in the FC region relying on the MS-6-RASPT2/SA-6-RASSCF(4,5|0,0|4,7) calculations, the level which agrees best with the experimental LA spectrum (Figure 11h); to this aim, we computed the overlap of the electronic states at the equilibrium and at each of the realizations used to initiate the molecular dynamics;

- computed the energies of further four excited states above the lowest $\pi\pi^*$ state at the MS-6-RASPT2/SA-6-RASSCF(4,5|0,0|4,7) level of theory (two $n\pi^*$ and two $\pi\pi^*$ excited states) along the trajectories run at the SS-2-CASPT2/SA-2-CASSCF(10,8) level and analyzed the evolution of the energy gap between the lowest $\pi\pi^*$ state and the higher lying excited states in a diabatic representation;

- performed ad-hoc non-adiabatic Tully fewest switches surface hopping dynamics simulation (10 repetitions for 50 trajectories for Urd and 5mUrd each) in a diabatic basis to resolve the percentage of trajectories prone to a non-adiabatic population transfer to another excited state;

Below we outline the details of the diabatization procedure and of the non-adiabatic Tully fewest switches surface hopping dynamics in the diabatic picture. In a closed system diabatic $\phi^d$ and adiabatic $\phi^a$ states are connected via a transformation

$$|\phi^d\rangle = |\phi^a\rangle \mathbb{U}$$

(7)

diabatic states are defined as the combination of the adiabatic states that resemble as much as possible the reference states. At the beginning of the simulation ($t = 0$), it is assumed that the diabatic and adiabatic states coin-
At every time step of the simulation the (local) transformation matrix \( U(t - ∆t → t) \) is constructed via Löwdin orthogonalization of the overlap matrix \( S(t - ∆t → t) = \langle φ^a(t - ∆t) | φ^a(t) \rangle \). In particular, we calculate the inverse of the overlap matrix \( S^{-1}(t - ∆t → t) \). The relation
\[
1 = S(t - ∆t → t)S^{-1}(t - ∆t → t) = \langle φ^a(t - ∆t) | φ^a(t) \rangle S^{-1}(t - ∆t → t) = \langle φ^a(t - ∆t) | φ^d(t) \rangle
\]
demonstrates that the inverse of the overlap matrix rotates the adiabatic basis at time step \( t \) to ensure maximal overlap with the wavefunctions at the previous time step \( t - ∆t \). Thus, following the definition of the adiabatic-to-diabatic transformation, \( S^{-1}(t - ∆t → t) \) equals the local transformation matrix \( U(t - ∆t → t) \). The transformation matrix \( U(0 → t) \) connecting the adiabatic states at time step \( t \) to the references ones at the beginning of the simulation is computed by propagating the local transformation matrices.

\[
U(0 → t) = U(0 → ∆t)U(∆t → 2∆t)U(2∆t → 3∆t) \ldots
\]

Thereby, in order to accurately perform the propagation one must keep track of relative signs of orbitals. In practice this is done by monitoring the sign of the diagonal elements of the local overlap matrices \( S(t - ∆t → t) \). Finally, the adiabatic-to-diabatic transformation matrix \( U(0 → t) \) is applied to the (diagonal) adiabatic Hamiltonian to transform it into a diabatic basis.

\[
H^d(t) = U^T(0 → t)H^a(0 → t)U(0 → t) = U^T(0 → t)\langle φ^a(t) | \hat{H} | φ^a(t) \rangle U(0 → t) = \langle φ^d(t) | \hat{H} | φ^d(t) \rangle
\]

The diabatic-to-adiabatic transformation holds for a closed electronic system, i.e. the states in the vector \( φ \) are separated from any higher lying states along the trajectory. In practice this condition is normally not fulfilled for any sub-space of states due to the high density and near degeneracy of excited states which leads to discontinuities due the occasional appearance of intruder states (i.e. not defined at the point of reference) and incomplete mapping to the states of reference. The incompleteness affects the local transformation matrix \( U(t - ∆t → t) \), an error which is then propagated in the computation of \( U(0 → t) \). To circumvent this obstacle we make use of the notion that

\[\text{This is not always the case. In fact, for uracil we observed in about 10% of the realizations a pronounced } ππ^* - nπ^* \text{ wavefunction mixing already in the FC point. These trajectories are excluded from the diabatization procedure.}\]
states which are energetically well separated are expected to mix negligibly\(^3\), hence we introduce a threshold (10 kcal/mol) for the energy difference between adiabatic states \(i\) and \(j\) above which the two states are assumed to not mix. Thus, the matrix elements \(U_{ij}(t - \Delta t \rightarrow t)\) and \(U_{ji}(t - \Delta t \rightarrow t)\) of the local transformation matrix are set to 0.0. The same rationale is applied to the global transformation matrix \(U(0 \rightarrow t)\). Unlike the local transformation matrix, the global transformation matrix retains memory of the state order with respect to the FC point (point of reference). Thus, if at a time step \(t\) all energy gaps would be above the threshold, \(U(t - \Delta t \rightarrow t)\) would equal the identity matrix, whereas \(U(0 \rightarrow t)\) could contain 0 on the diagonal and ±1 on the off-diagonal thus indicating state swapping\(^4\). Furthermore, we apply the diabatization to a sub-space of the lowest three roots (being ground state, \(\pi\pi^*\) and \(n\pi^*\) in the FC point), thereby assuring that the photoactive \(\pi\pi^*\) state is described continuously and nearly completely in this sub-space\(^5\). In practice, along a given trajectory the third root could change nature (e.g. the \(n\pi^*\) state occasionally gets energetically destabilized in favor of the second \(\pi\pi^*\) state). We observe that this happens when the energy gap to the lowest \(\pi\pi^*\) state is far above the selected threshold. Thus this state swapping has no effect on the population in the photoactive \(\pi\pi^*\) state. Instead, we obtain a new set of reference wavefunctions (ground state, lowest \(\pi\pi^*\) and second \(\pi\pi^*\) states). Thus, the diabatization is not performed on a closed system of states along an entire trajectory. Instead, we obtain information about how often and how strongly the photoactive \(\pi\pi^*\) state mixes with the closest lying excited state whose nature may vary. In the case of Urd the third root is almost exclusively a \(n\pi^*\) state, however in 5mUrd we observe frequent swapping between \(n\pi^*\) and the second \(\pi\pi^*\) state.

Having the diabatic Hamiltonian at each time step we are in a position to perform ad-hoc non-adiabatic dynamics simulations in the diabatic basis. Thereby, we realize that the simulations performed at the SS-2-CASPT2/SA-2-CASSCF(10,8) level of theory (i.e. in the absence of lone pairs in the active space and thus of \(n\pi^*\) states in the averaging procedure) can be regarded as performed in the diabatic picture, i.e. following the \(\pi\pi^*\) state, thereby disregarding its mixing with any \(n\pi^*\) or higher lying \(\pi\pi^*\) states. We can thus

---

\(^3\)Similar approximation is frequently adopted in non-adiabatic dynamics in the adiabatic representation in order to reduce the effort in computing non-adiabatic coupling elements.

\(^4\)\(U(0 \rightarrow t)\) is related to the identity matrix by \(2 \times 2\) rotations (with an angle \(\phi = 90°\)) on each pair of swapped states.

\(^5\)We have assured that the photoactive \(\pi\pi^*\) state never becomes higher than the third root in the calculations. Furthermore, we observe that the photoactive \(\pi\pi^*\) state always mixes with only one other state at a time.
re-run the trajectories, re-using at each time step the velocities for the nuclei computed at the SS-2-CASPT2/SA-2-CASSCF(10,8) level, thereby propagating the electronic coefficients $c_i$ using the diabatic Hamiltonian obtained with aforementioned adiabatic-to-diabatic transformation

$$i\hbar \frac{\partial c_i}{\partial t} = \sum_j H^d_{ij}(t)c_j. \quad (11)$$

Thereby, we adopt the approximation that the occasional mixing of the $\pi\pi^*$ state with close-by electronic states does not affect the dynamics. This approximation is supported by the empirical observation on both systems under investigation that state mixing is a rare (in time) a highly localized (in space) event. At every time step the hopping probability between diabatic states is calculated according to the formula

$$P_{i \rightarrow j} = \max \left[ 0, \frac{2\Delta t}{c_i^*c_i} \Im(c_j^*c_i)H^d_{ji} \right] \quad (12)$$

and a transition from surface $i$ to surface $j$ (i.e. a hopping event) is realized through the usual stochastic procedure. An estimate of the percentage of trajectories that depart from the $\pi\pi^*$ state before its decay to the ground state is finally obtained by re-running the set of trajectories multiple times and averaging over the number of hopping events. Thereby, it is assumed that the system does not return on the $\pi\pi^*$ surface once it hopped to another surface. Thus, the obtained estimate could be regarded as an upper limit for the population transfer to other excited states.

### 2.8 Simulation of transient absorption spectra

On the basis of the simulations transient absorption spectra are generated after computing the excited state electronic structure (energies $E$ and transition dipole moments $\mu_{ij}$) of Urd and 5mUrd at the SS-15-CASPT2/SA-15-CASSCF(10,8) level of theory at every time step. In practice, at each delay time $t_d$ between pump and probe, the transient spectrum is calculated at the coordinates as a superposition of stimulated emission to the ground state and photoinduced absorption to higher electronic states assuming simple vertical transitions.

$$I(\omega, t_d) \propto \frac{1}{N_{\text{traj}}} \sum_{k} \left[ \sum_f \mu_{ef\leftarrow e}^2(R_k(t_d)) g(\Delta E_{fe}(R_k(t_d)) - \hbar \omega) ight. \\
- \left. \mu_{eg\leftarrow g}^2(R_k(t_d)) g(\Delta E_{eg}(R_k(t_d)) - \hbar \omega) \right]$$

(13)
where $k$ iterates over the number of trajectories, $R_k(t_d)$ denotes the geometrical coordinates at time $t_d$, $e$ denotes the state which drives the semi-classical trajectory (i.e. either $S_1$ or the ground state) and $f$ denotes a state from the manifold of excited states accessible with the probe pulse. The intensity is convoluted with a normalized line shape function $g(\Delta E_{ij}(R_k(t_d)) - \hbar \omega)$ peaked at the transition energies and broadened by a phenomenological constant $\sigma$:

$$g(\Delta E_{ij}(R_k(t_d)) - \hbar \omega) = \exp\left(\frac{-E(i)(R_k(t_d)) - E(j)(R_k(t_d)) - \hbar \omega)^2}{2\sigma^2}\right)$$

(14)

For $\sigma$ we chose a value of 0.15 eV to remove statistical noise. The final spectrum has been uniformly blue-shifted by 0.3 eV resulting in a better match with the experiment. This correction is justified by the analogous red-shift observed in the LA spectra computed with CAS(10,8) with respect to the experiment (see Supplementary Note 3.2 for details).

Polarization control was incorporated by introducing a weighting factor $c(\mu_\alpha, \mu_\beta, E_a, E_b)$ to each contribution in the inner sum in eq. 13, functions of the angles between the incident electric fields $E_a$ (pump) and $E_b$ (probe) and between the coupled transition dipole moments $\mu_\alpha$ and $\mu_\beta$ of the system.

$$c(\mu_\alpha, \mu_\beta, E_a, E_b) = \langle (\mu_\alpha \cdot E_a)^2 (\mu_\beta \cdot E_b)^2 \rangle = \frac{1}{15} \left[ 2 - \cos^2 \Theta_{E_a,E_b} + \cos^2 \Theta_{\mu_\alpha,\mu_\beta} (3 \cos^2 \Theta_{E_a,E_b} - 1) \right]$$

(15)

eq . 15 results from the general equation derived by Hochstrasser for four pulses and four interacting dipoles[Hoch01]. Thereby, $\mu_\alpha = \mu_{eg}(R_k(0))$ is the transition dipole moment of the $S_1$ (bright state) at time $t_d = 0$ fs which interacts with the pump pulse, whereas $\mu_\beta = \mu_{fe}(R_k(t_d))$ or $\mu_\beta = \mu_{eg}(R_k(t_d))$ denotes the transition dipole moments of the states coupled to the photoactive states via the probe pulse at time $t_d$. For parallel arrangement of the pump and probe pulse ($\Theta_{E_a,E_b} = 0^\circ$) eq. 15 simplifies to:

$$c(\mu_\alpha, \mu_\beta) = \frac{1}{15} \left[ 1 + 2 \cos^2 \Theta_{\mu_\alpha,\mu_\beta} \right]$$

(16)

whereas for orthogonal arrangement ($\Theta_{E_a,E_b} = 90^\circ$) eq. 15 reads

$$c(\mu_\alpha, \mu_\beta) = \frac{1}{15} \left[ 2 - \cos^2 \Theta_{\mu_\alpha,\mu_\beta} \right]$$

(17)

At the so called "magic angle" set up (pump and probe have an angle of 54.7°) the signal intensity becomes independent of the angle between the
dipole moments (the third term in eq. 15 vanishes as $3 \cos^2(54.7°) - 1 = 0$). Thus, the intensity depends only on the magnitude of the dipole moments.

Finite pulse duration is accounted for ad-hoc by convoluting the transient absorption signal $I(\omega, t_d)$ with a Gaussian function in the time domain

$$I(\omega, t_d) = \int_{-\infty}^{+\infty} d\tau \, I(\omega, \tau) \exp\left(-\frac{(\tau - t_d)^2}{2\sigma^2}\right)$$

(18)

A standard deviation $\sigma = 11$ fs was used (matching the experimental full-width-half-maximum of 25 fs).

**Supplementary Note 3:**

**Simulation results**

3.1 **Wigner sampling analysis**

Below we present an analysis of the Wigner sampling regarding potential and kinetic energy redistribution, number of movable water molecules, temperature effects and several relevant coordinates.

3.1.1 **Uridine (Urd)**

Figure 3,a shows a representative snapshot of the Urd sampling with corresponding High/Medium/Low layer partitioning used for the mixed quantum-classical molecular dynamics simulations with waters within 5 Å distance to atoms of the uracil nucleobase (High Layer) being added to the movable Medium layer. On average 45 water molecules are included in the Medium layer.

The kinetic energy in the vibrational ground state of Urd is 3.12 eV at 0 K when all modes are considered in the sampling, whereas the kinetic energy reduces to 1.91 eV when only the lowest 69 modes are considered. The sampling at 0 K correctly reproduces this value (see Figure 4a) with the kinetic energy computed from the sampled velocities according to the formula:

$$E_{\text{kin}} = \frac{mv^2}{2}.$$  

(19)

Thereby, 0.75 eV of the kinetic energy is accumulated by the uracil nucleobase (see Figure 4b). Performing the same sampling at room temperature (300 K) increases the average kinetic and potential energies by ca. 0.18 eV for Urd and 0.07 for uracil (see Figure 4a,b).
Supplementary Figure 3: Left: High/Medium/Low layer partitioning used for MD simulations; waters in 5 Å distance to Uracil (High Layer) added to the movable layer; Right: distribution of water molecules in all 500 snapshots generated in the Wigner sampling.

Supplementary Figure 4: Distribution of kinetic energies in Urd (left) and Uracil (right) for 0K and 300K.

At 300 K the average N1-C2 bond length is 1.38 Å (see Figure 5,a) with few snapshots exhibiting elongation of up to 1.50 Å. The room temperature sampling does not lead to a more pronounced bond elongation compared to the sampling at 0 K. The distribution of dihedrals is Gaussian around the average values of 180 °(H_{10}-C_6-C_5-C_4 and H_9-C_5-C_6-N_1) / 0 °(H_{10}-C_6-C_5-H_9) (see Figure 5,b solid lines). The H_{10}-C_6-C_5-C_4 and H_9-C_5-C_6-N_1 dihedrals show fluctuations of ±20 °around the equilibrium value. The H_{10}-C_6-C_5-H_9 dihedral is more flexible showing fluctuations of ±30 °around the equilibrium value. The snapshots selected for the MD simulations which fall within the envelope of the pump pulse (277-266 nm) do not show any preferential geometrical distortion compared to the complete set of 500 snapshots (see Figure 5,b dotted lines).
Supplementary Figure 5: Distribution of several characteristic coordinates in uracil in all 500 snapshots (solid lines) and in the window of pumping [277-266 nm] (dotted lines).

3.1.2 5-methyl-uridine (5mUrd)

Figure 6,a shows a representative snapshot of the 5mUrd sampling with corresponding High/Medium/Low layer partitioning used for the mixed quantum-classical molecular dynamics simulations with waters within 5 Å distance to atoms of the 5mUra nucleobase (High Layer) being added to the movable Medium layer. On average 46 water molecules are included in the Medium layer.

Supplementary Figure 6: Left: High/Medium/Low layer partitioning used for MD simulations; waters in 5 Å distance to Uracil (High Layer) added to the movable layer; Right: distribution of water molecules in all 500 snapshots generated in the Wigner sampling

The kinetic energy in the vibrational ground state of 5mUrd is 3.50 eV at 0 K when all modes are considered in the sampling, whereas the kinetic
energy reduces to 2.10 eV when only the lowest 74 modes are considered. The sampling at 0 K correctly reproduces this value (see Figure 7a). Thereby, 0.95 eV of the kinetic energy is accumulated by the 5mUra nucleobase (see Figure 7b). Performing the same sampling at room temperature (300 K) increases the average kinetic and potential energies by ca. 0.17 eV for 5mUrd and 0.07 for 5mUra (see Figure 7a,b).

Supplementary Figure 7: Distribution of kinetic energies in 5mUrd (left) and 5mUra (right) for 0K and 300K

At 300 K the average N1-C2 bond length is 1.38 Å (see Figure 8,a) with few snapshots exhibiting elongation of up to 1.50 Å. The room temperature sampling does not lead to a more pronounced bond elongation compared to the sampling at 0 K. The distribution of dihedrals is Gaussian around the average values of 180 °(H10-C6-C5-C4 and CH3-C5-C6-N1) / 0 °(H10-C6-C5-CH3) (see Figure 8,b solid lines). The H10-C6-C5-C4 and CH3-C5-C6-N1 dihedrals show fluctuations of ±20 °and ±10 °around the equilibrium value, respectively. The H10-C6-C5-CH3 dihedral shows fluctuations of ±30 °around the equilibrium value. The snapshots selected for the MD simulations which fall within the envelope of the pump pulse (276-266 nm) do not show any preferential geometrical distortion compared to the complete set of 500 snapshots (see Figure 8b, dotted lines).
3.1.3 Comparison

At room temperature 5mUrd has on average 0.18 eV more kinetic energy at disposal compared to Urd (2.27 vs. 2.09 eV) as seen by the shift of the distribution of kinetic energies (Figure 9,a). The 5mUra nucleobase has on average 0.20 eV more kinetic energetic at disposal compared to uracil (1.02 vs. 0.82 eV) (Figure 9,b).
Supplementary Figure 10: Comparison of the distribution of several characteristic coordinates in Urd (solid lines) and 5mUrd (dotted lines).
3.2 Linear absorption spectra

In the following we present the LA spectra of Urd and 5mUrd computed at various levels of theory. In particular, we benchmarked active space, number of the states in the state-averaging procedure and RASPT2 flavor (single state vs. multi-state).

3.2.1 Uridine (Urd)

Figure 11a,b compares the LA spectrum of Urd computed at the SS-4-CASPT2/SA-4-CASSCF(10,8) level with active space comprising all $\pi$-orbitals of the system (black solid line) to the experimental spectrum (purple solid line). It is seen that both methods tend to red-shift the absorption energy of the $\pi\pi^*$ bands by ca. 0.30 eV ($\sim$ 20 nm at this wavelength). State-assignment (relying on the coefficients of the leading configuration state functions, CSFs) demonstrates that the lowest absorption band (peaking at 260 nm (280 nm) in the experiment (simulation)) is dominated by the H→L transition (red solid line). Including the lone pairs of the oxygens in the active space, thus using CASSCF(14,10), allows to compute the contribution of the lowest $n\pi^*$ states (blue solid line in Figure 11c,d). As expected the GS→ $n\pi^*$ transition is essentially dark and it does not affect significantly the LA spectrum. Increasing the active space with additional virtual orbitals leads to a much improved agreement with the experiment (Figure 11e-h). In the case of the $\pi$-only active space we could increase the active space in a RASSCF fashion by eight additional orbitals thus using a RASSCF(4,5|0,0|4,11) (Figure 11e-h), instead in the case of the active space augmented with lone pairs we could afford to add four virtual orbitals thus giving rise to a RASSCF(4,7|0,0|4,7). In both cases the $\pi\pi^*$ bands exhibit a blue-shift with respect to the smaller active spaces. Thereby, the multi-state treatment gives a slightly better agreement with experiment. As elucidated in the methodology sections we used these findings to define the criteria for selecting the snapshots for the mixed quantum-classical dynamics simulations, namely the snapshots which fall within the spectral envelope of the pump pulse (green solid line in Figure 11) at the SS-4-CASPT2/SA-4-CASSCF(4,5|0,0|4,11).

3.2.2 5-methyl-uridine (5mUrd)

Figure 12a,b compares the LA spectrum of 5mUrd computed at the SS-4-CASPT2/SA-4-CASSCF(10,8) level with active space comprising all $\pi$-orbitals of the system (black solid line) to the experimental spectrum (purple solid line). It is seen that both methods tend to red-shift the absorption energy of the $\pi\pi^*$ bands by ca. 0.35 eV ($\sim$ 25 nm at this wavelength).
Supplementary Figure 11: Linear absorption spectra of Urd obtained with various levels of theory.
State-assignment (relying on the coefficients of the leading CSFs) demonstrates that the lowest absorption band (peaking at 266 nm (290 nm) in the experiment (simulation)) is dominated by the H→L transition (red solid line). Including the lone pairs of the oxygens in the active space, thus using CASSCF(14,10), allows to compute the contribution of the lowest nπ* states (blue solid line in Figure 12c,d). As in the case of Urd the GS→ nπ* transition is essentially dark and it does not affect significantly the LA spectrum. Increasing the π-only active space with eight additional virtual orbitals, thus using RASSCF(4, 5|0, 0|4, 11) (Figure 12e-f), and with four additional virtual orbitals in the case of the active space augmented with lone pairs, thus using RASSCF(4, 7|0, 0|4, 7) (Figure 12g-h), leads to a much improved agreement with the experiment. As elucidated in the methodology sections we used these findings to define the criteria for selecting the snapshots for the mixed quantum-classical dynamics simulations, namely the snapshots which fall within the spectral envelope of the pump pulse (green solid line in Figure 12) at the SS-4-CASPT2/SA-4-CASSCF(4, 5|0, 0|4, 11).
Supplementary Figure 12: Linear absorption spectra of 5mUrd obtained with various levels of theory.
Supplementary Figure 13: $\pi\pi^* - n\pi^*$ energy gap distribution in Urd (left) and 5mUrd (right).

3.2.3 Comparison

Figure 13 compares the distributions of $n\pi^* - \pi\pi^*$ energy gaps in Urd (left) and 5mUrd (right) computed with the small CASSCF(14,10) (red and blue solid lines) and large CASSCF(4,7|0,0|4,7) (green and black solid lines) at both single-state and multi-state RASPT2 level. A minor fraction of the samples (18% in the case of SS-CASPT2/CASSCF(14,10), 11% in the case of SS-RASPT2/RASSCF(4,7|0,0|4,7)) shows an inverted state order with the $n\pi^*$ being the lowest electronic state. At MS-CASPT2 level the number of snapshots with inverted state order is reduced with respect to the single state variant to 6% and 9%. Overall, it can be clearly seen that the majority of snapshots exhibit an $n\pi^*$ state well above the $\pi\pi^*$ state in the FC point slightly varying between 0.35-0.45 eV with the level of theory. Importantly, this demonstrates that the small active space seems to reproduce properly the $n\pi^*$ energy gap despite the overall red-shift of the transition energies seen in the corresponding spectra (Figure 11a,b)). With 0.50-0.60 eV the $n\pi^* - \pi\pi^*$ energy gap is even more pronounced in 5mUrd (Figure 13, right). A fraction of the samples (19% at SS-CASPT2/CASSCF(14,10)) shows an inverted state order. At MS-CASPT2 level the number of snapshots with inverted state order is reduced to 4%.
3.3 SS-CASPT2/SA-2-CASSCF(10,8) trajectory plots

In the following we plot the evolution of the individual representative trajectories computed for Urd and 5mUrd.

3.3.1 Uridine (Urd)

Figure 14 shows ten representative trajectories which were run for up to 500 fs. We observe a hop to the ground state in 53 of all 59 trajectories, with hops occurring as early as 50 fs (e.g. 14a,e), i.e. within two oscillations of the C-C double bond vibration. In total, we document 15 trajectories that hop on a sub-100 fs time scale, 21 trajectories which hop between 100-200 fs, 12 trajectories which hop between 200-300 fs and five trajectories which hop on a longer time scale (300-500 fs). Three trajectories do not hop on the time scale of the simulation (Figure 14i,j). The decay to the ground state occurs on all cases through H_{9} out-of-plane puckering (Figure 21), whereas we observe a clear preference for down-puckering (41 trajectories, Figure 14a-d) with respect to up-puckering (12 trajectories, Figure 14e-h).

The CASPT2 dynamics with the full valence-π active space are overall stable, showing a tolerable increase of the total energy over the course of the simulation. There are several factors that play a role in this issue. Apart from the 1.0 fs time step, which is insufficient to describe fast stretching dynamics or the large amplitude dynamics of the ”hot” ground state, the active space size also plays a significant role. Increasing the active space by two extra-valence orbitals reduces substantially the deviation of the total energy, in particular in the region of the crossing (Figure 15). Hence, the CAS(10,10) trajectories were utilized in the stochastic analysis and spectra simulations instead of their CAS(10,8) counterpart. The improved accuracy comes at the price of an increased cost in the computation, therefore only trajectories which show total energy deviation above 10 kcal/mol on the excited state (8 in Urd) were repeated with the larger active space. We note that the two sets of trajectories, SS-CASPT2/SA-2-CASSCF(10,8) (red) and SS-CASPT2/SA-2-CASSCF(10,10) (black) in Figure 15, behave qualitatively similarly on a short time scale (up to 300-400 fs). As a consequence the photophysics of Urd is not affected.
Supplementary Figure 14: Representative trajectories for Urd
Supplementary Figure 15: Comparison between representative trajectories computed at the SS-2-CASPT2/SA-2-CAS(10,8) (red) and at the SS-2-CASPT2/SA-2-CAS(10,10) (black) levels of the theory in Urd
3.3.2 5-methyl-uridine (5mUrd)

Figures 16 show ten representative trajectories which were run for up to 1000 fs. Overall, we observe a hop to the ground state in 40 of all 59 trajectories. Thereby, we document only one trajectory which hops to the ground state earlier than 200 fs (Figure 16a) where the decay is facilitated through N1-C2 bond breaking on a 100 fs time scale (conical intersection shown in Figure 22). In the majority of the remaining cases the decay to the ground state is facilitated through CH3 out-of-plane puckering (conical intersections shown in Figure 22), whereas we observe a clear preference for down-puckering (31 trajectories, Figure 16,b-d) with respect to up-puckering (15 trajectories, Figure 16e-g). The hopping events are evenly distributed between 200 and 1000 fs with 12 hops between 200-400 fs, 11 hops between 400-600 fs, 7 hops between 600-800 fs and 10 hops between 800-1000 fs. In 19 trajectories no hop was encountered on the time scale of the simulation (Figure 16i,j). Thereof, in 7 the CI region is reached but no hop is observed (Figure 16i). In 3 of these cases the CI region is reached towards the end of the simulation (trajectories 49, 51 and 52). In the other 4 cases we observe a O8-puckering causing the simulation to break due to a pronounced lone pair mixing in the active space (trajectories 53, 55, 57 and 58, Figure 16h). These 7 trajectories are regarded as decaying, thus resulting in a total a 47 hopping trajectories within the time scale of the simulation.

The CASPT2 dynamics with the full valence-π active space are overall stable, although with more pronounced deviations compared to Urd. As in the case of Urd, in the ground state the total energy increase is more pronounced due to the large gradient. Increasing the active space by two extra-valence orbitals reduces substantially the deviation of the total energy (Figure 17). Hence, the CAS(10,10) trajectories were utilized in the stochastic analysis and spectra simulations instead of their CAS(10,8) counterpart. Only trajectories which show total energy deviation above 10 kcal/mol on the excited state (16 in 5mUrd) were repeated with the larger active space. We note that the two sets of trajectories, SS-CASPT2/SA-2-CASSCF(10,8) (red) and SS-CASPT2/SA-2-CASSCF(10,10) (black) in Figure 17, behave qualitatively similarly on a short time scale (up to 200-300 fs). However, due to the significantly longer life time of 5mUrd the dynamics diverge beyond 400 fs. The CAS(10,10) dynamics show an accelerated decay, albeit still in the few-hundred femtosecond range.
Supplementary Figure 16: Representative trajectories for 5mUrd
Supplementary Figure 17: Comparison between representative trajectories computed at the SS-2-CASPT2/SA-2-CAS(10,8) (red) and at the SS-2-CASPT2/SA-2-CAS(10,10) (black) levels of the theory in 5mUrd.
3.4 SS-2-CASPT2/SA-2-CASSCF(10,8) vs. MS-4-CASPT2/SA-4-CASSCF(10,8) in Urd

In Urd we run 49 trajectories at the MS-2-CASPT2/SA-2-CASSCF(10,8) level of theory. Figure 18 shows a comparison between the single- and multi-state description for several representative trajectories. Whereas in 10 trajectories we observe good agreement between both CASPT2 flavors (e.g. Figure 18a,b), in the majority of cases (34) the system decays faster at the SS-CASPT2 level (e.g. Figure 18c,d). Thereby, the MS-CASPT2 profiles exhibit the aforementioned $S_1$-$S_0$ energy splitting in the region of the conical intersection, the reason for the slowed down decay. Finally, Figure 18e documents one of the two cases in which the MS-CASPT2 dynamics decays faster.
Supplementary Figure 18: Comparison between representative trajectories computed at the SS-2-CASPT2/SA-2-CASSCF(10,8) level (left) and MS-4-CASPT2/SA-4-CASSCF(10,8) level (right) in Urd.
3.5 MS-CASPT2/SA-6-CASSCF(14,10) plots along trajectories: adiabatic vs. diabatic picture

For reasons elucidated in Supplementary Note 2.6 the lone pairs of the oxygen atoms were not included in the active space used to perform the mixed quantum classical molecular dynamics simulations (thus denoted $\pi$-only active space). The involvement of higher lying excited states was estimated ad-hoc by computing the energetics along the trajectories at the MS-6-RASPT2/SA-6-RASSCF(4, 5|0, 0|4, 7) level, thus considering four electronic states above the lowest $\pi\pi^*$ state and analyzing the probability for population transfer therein.

Figures 19a-f (Urd) and 20a-f (5mUrd) show a detailed analysis for several representative trajectories. The individual panels contain the following information:

(A) shows the energetics of the ground (red) and $S_1$ (blue) states at the $\pi$-only SS-2-CASPT2/SA-2-CASSCF(10,8) level (solid lines), as well as the energetics of the diabatic states $D_0$, $D_1(\pi\pi^*)$ and $D_2$ resulting from the diabatization (sec. 2.7) of the lowest three states computed at the MS-6-RASPT2/SA-6-RASSCF(4, 5|0, 0|4, 7) level (dotted lines) along the $\pi$-only trajectories.

(B) documents the evolution of the $D_1(\pi\pi^*)$-$D_0$ energy gap (diagonal elements of the diabatic Hamiltonian)

(C) documents the evolution of the $D_2$-$D_1(\pi\pi^*)$ energy gap (diagonal elements of the diabatic Hamiltonian)

(D) shows the evolution of the magnitude of the $D_0 \rightarrow D_1(\pi\pi^*)$ (black) and $D_0 \rightarrow D_2$ (green) transition dipole moments.

(E) documents the population dynamics in the diabatic states $D_0$, $D_1$ and $D_2$ (blue, black and red solid lines, respectively).

We remind that the nature of $D_2$ can change along the trajectories as the $n\pi^*$ and second $\pi\pi^*$ states often swap order due to their energetic proximity. This state swapping can be recognized by inspecting the magnitude of the $D_0 \rightarrow D_2$ transition dipole moment in panel (D). While the photoactive state $D_1(\pi\pi^*)$ shows a constant value around 2 Debye which decreases to zero towards the crossing region, the oscillator strength of $D_2$ often exhibits sudden jumps between zero and 2 Debye, an indicator of its variable nature. This does not affect the dynamics on the photoactive state when it happens outside of the energy window considered to give rise to a notable state mixing.
(D$_2$-D$_1$ gap below 10 kcal/mol). Thus, panel (D) shows the effective probability for the population to depart from the photoactive $\pi\pi^*$ state to the energetically closest electronic state.

3.5.1 Uridine (Urd)

**Analysis of wavefunction mixing in the FC region** The wavefunction analysis in the Franck-Condon region shows that six out of 56 trajectories ($\sim10\%$) exhibit pronounced $\pi\pi^* - n\pi^*$ mixing which makes it impossible to perform initial assignment of the adiabatic states. In 18 of the remaining 50 trajectories (1/3) we observe a weak mixing (with $n\pi^*$ CSF coefficients in the $\pi\pi^*$ wavefunction of up to 0.3). In the remaining 32 trajectories the $\pi\pi^*$ and $n\pi^*$ states could be clearly identified, and showed no significant mixing. In one case the $n\pi^*$ was found to be the lowest excited state in the Franck-Condon point. Based on this analysis, we can conclude that roughly 5% of population (half of the trajectories exhibiting strong wavefunction mixing) would immediately adopt $n\pi^*$ due to the wavefunction mixing in the Franck-Condon region.

**Analysis of the $n\pi^*$ energetics along the $\pi$-only trajectories** Including the oxygen lone pairs in the active space seemingly affects strongly the energy profiles of the electronic states (compare solid vs. dotted profiles in panels (A) of Figure 19a-f). The trajectories with and without lone pairs in the active space often exhibit considerably differing profiles (e.g. 19c). However, the $\pi\pi^*$/ground state energy gaps remain quasi-identical (see panel (B) in Figure 19a-f) except for the crossing region where the MS-CASPT2 calculations result in large state splitting as discussed earlier (sec. 2.6). This indicates that the energy profiles of $S_0$ and $S_1$ are modulated in the same manner by the presence of the oxygen lone pairs in the active space.

Regarding the relative positions of the states D$_1(\pi\pi^*)$ and D$_2$, it is evident that they are close in the FC region (often within 10 kcal/mol, see also Figure 13) but diverge towards the conical intersection. We observe oscillatory dynamics for the D$_2$-D$_1$ energy gap with a period of ca. 20 fs (see panel (C) in Figure 19a-f which correlates to the period of the oscillation of the double bonds, thereby often reducing the gap to a few kcal/mol. Nevertheless, an inversion of the energies of D$_1$ and D$_2$ is observed only on three occasions (e.g. Figure 19e,f). Furthermore, we generally observe a weak and temporally confined mixing of the diabatic states (indicated by the small off-diagonal elements in the diabatic Hamiltonian). Thus, overall, Urd rarely explores regions of state-mixing and when this is the case it spends therein only a few femtoseconds.
Non-adiabatic dynamics (including the \( n\pi^* \) state) in the diabatic representation  Finally, we used the diabatic profiles and propagated the amplitudes of the electronic wavefunction starting with amplitude 1.0 in the \( \text{D}_1(\pi\pi^*) \) state. Panels (E) of Figure 19a-f show the evolution of the population along each trajectory\(^6\). Due to the weak wavefunction mixing, the population of \( \text{D}_1 \) rarely deviates from unity and, consequently, we find low probability for population transfer to another excited state. In ten repetitions of the dynamics of the entire set of trajectories we found that on average 15\% of the trajectories depart from the \( \text{D}_1(\pi\pi^*) \) state before a funnel to the ground state is reached.

Conclusion  Thus, based on the above analysis we estimate that in Urd, pumped at 275 nm, an upper limit of 20\% of the population could potentially evolve on a different electronic state (an \( n\pi^* \) or a higher lying \( \pi\pi^* \) state). The \( \text{D}_2-\text{D}_1 \) energy gap dynamics exhibit a high-frequency oscillatory behaviour which can be traced back to double bonds vibrations in the aromatic ring.

\(^6\)In order to explore the hopping probability along the entire trajectory hopping was suspended.
Supplementary Figure 19: Analysis of the diabatic dynamics along six representative trajectories of Urd.
3.5.2 5-methyl-uridine (5mUrd)

Analysis of wavefunction mixing in the Franck-Condon region  The wavefunction analysis in the Franck-Condon region shows that, in contrast to Urd, 5mUrd does not exhibit pronounced $\pi\pi^* - n\pi^*$ wavefunction mixing. In fact, in no case do we find strong mixing, while we observe a weak mixing (with $n\pi^*$ CSF coefficients in the $\pi\pi^*$ wavefunction of up to 0.3) in seven out of the 54 analyzed trajectories ($\sim 13\%$). This trend is in line with the larger $\pi\pi^* - n\pi^*$ energy gap found for 5mUrd (0.50-0.55 eV) compared to Urd (0.35-0.40) as shown in Figure 13. Furthermore, we observe slightly more often (six cases) state swapping between the lowest $n\pi^*$ state and the second $\pi\pi^*$ state, which was encountered three times for Urd. This finding is in line with the predicted stronger involvement of the second $\pi\pi^*$ state in the photophysics of the 5mUrd in comparison to Urd [Pepino17]. However, we emphasize that we do not observe mixing of the lowest two $\pi\pi^*$ states. We expect this to be the case when pumping in the blue part (e.g. 260-240 nm) of the lowest absorption band in the LA spectrum (see Figure 12). Based on this analysis, we can conclude that immediate population of the $n\pi^*$ state is improbable in the Franck-Condon region.

Analysis of the $n\pi^*$ energetics along the $\pi$-only trajectories  Most of the conclusions drawn for Urd apply also in the case of 5mUrd. The $\pi\pi^*/$ground state energy gaps remain quasi identical in all cases except for the crossing region where the $\pi\pi^*/$ground state degeneracy is often lifted and the splitting can reach up to 50 kcal/mol (panels (B) in Figure 20,a-f). Regarding the relative positions of the $D_1(\pi\pi^*)$ and $D_2$ we observe state mixing even more rarely compared to Urd. Inversion of the energies of $D_1$ and $D_2$ is again observed only at three occasions and is temporarily confined to a few femtoseconds. In contrast to Urd the magnitude of the $D_0$-$D_2$ transition dipole moment frequently exhibits sudden jumps between zero and ca. 2 Debye, an indicator of the constant swapping of $n\pi^*$ and the second $\pi\pi^*$ states (panels (D) in Figure 20,a-f).

Non-adiabatic dynamics (including the $n\pi^*$ state) in the diabatic representation  Finally, we used the diabatic profiles and propagated the amplitudes of the electronic wavefunction starting with an amplitude 1.0 in the $D_1(\pi\pi^*)$ state. Panels (E) of Figure 20,a-f show the evolution of the population along each trajectory$^7$. Due to the larger (in comparison to Urd)

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$^7$In order to explore the hopping probability along the entire trajectory hopping was suspended.
energy gap to the higher lying state leading to a weak wavefunction mixing, the population of D$_1$ rarely deviates from unity and, consequently, we find lower probability for population transfer to other excited states compared to Urd. In ten repetitions of the dynamics of the entire set of trajectories we found that on average merely 1% of the trajectories depart from the $\pi\pi^*$ state before the funnel to the ground state is reached.

**Non-adiabatic dynamics (including the $n\pi^*$ state) in the adiabatic representation** Considering the fact that the character of the S$_1$ state is dominated by the $\pi\pi^*$ configuration throughout the entire dynamics$^8$, we performed also ad-hoc non-adiabatic fewest switches surface hopping dynamics in the adiabatic representation in order to remove the bias of the diabatization procedure. To that aim we run the trajectories, re-using at each time step the velocities for the nuclei computed at the SS-2-CASPT2/SA-2-CASSCF(10,8) level$^9$ and propagated the amplitudes of the electronic wavefunction starting with an amplitude 1.0 in the S$_1$ state. At each step time-derivative couplings $\sigma_{ij}$ were computed based on wave function overlaps (sec. 2.5) and hopping was realized through the usual stochastic procedure. The simulations confirm the findings of the dynamics performed in the diabatic representation, namely, that the S$_2$ state is not involved in the non-adiabatic dynamics. In ten repetitions of the dynamics of the entire set of trajectories we found that the population of S$_1$ rarely deviates from unity and on average merely 1% of the trajectories depart from the S$_1$ state before the funnel to the ground state is reached.

**Conclusion** Thus, based on the above analysis we estimate that in 5mUrd, pumped at 275 nm, the involvement of further excited state ($n\pi^*$ or higher lying $\pi\pi^*$) is marginal (around $\sim$ 1%).

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$^8$S$_1$ mixes weakly with higher lying electronic states along the dynamics and inversion of the D$_1(\pi\pi^*)$/D$_2$ energies is virtually not observed.

$^9$Essentially, we assume that the S$_1$ does not mix with higher lying electronic states and their energetic vicinity does not affect the potential energy surface topology of S$_1$.  

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Supplementary Figure 20: Analysis of the diabatic dynamics along six representative trajectories of 5mUrd.
3.6 Conical intersections

3.6.1 Uridine (Urd)

Figure 21 shows superimposed CIs encountered along the 59 trajectories of Urd. The decay to the ground state occurs in all cases through $H_9$ out-of-plane puckering, whereas we observe a clear preference for down-puckering with respect to up-puckering.

Supplementary Figure 21: Conical intersections in Urd: a) $C_3$ out-of-plane down-puckering (46); b) $C_3$ out-of-plane up-puckering (13). Points of degeneracy along the same trajectory separated by more than 20 fs were considered as different conical intersection geometries.

3.6.2 5-methyl-uridine (5mUrd)

Figure 22 shows superimposed CIs encountered along the 59 trajectories of Urd. The decay to the ground state occurs through three different mechanisms, the dominant one being the CH$_3$ out-of-plane puckering, whereas we observe a clear preference for down-puckering with respect to up-puckering. O$_8$ out-of-plane puckering and N$_1$-C$_2$ bond breaking are secondary channels.
Supplementary Figure 22: Conical intersections in Urd: a) H$_2$ out-of-plane down-puckering (31); b) H$_2$ out-of-plane up-puckering (15); c) O$_4$ out-of-plane puckering (4); d) N$_1$-C$_2$ bond breaking (1). Points of degeneracy along the same trajectory separated by more than 20 fs were considered as different conical intersection geometries.
3.7 Analysis of vibrational coherences

To obtain a better idea of the ES vibrational dynamics of Urd and 5mUrd we performed a normal mode analysis. To this aim, we optimized the nuclear coordinates in the $S_1 (\pi\pi^*)$ state utilizing the same QM/MM set up outlined in Supplementary Note 2.2 and using the SS-2-CASPT2/SA-2-CASSCF(10,8)/ANO-L[321,21] level of theory of the High Layer. The optimized $S_1$ geometries superimposed over the ground state equilibrium geometries are shown in Figure 23. We note a small distortion of the planarity of the aromatic ring, more pronounced in the 5mUrd. Subsequently, a frequency calculation was performed (numerically) at the same level of theory. All frequencies resulted positive, thus confirming the stationary nature of the obtained geometries.

Starting from the definition of the normal mode matrix

$$ P^\dagger = M^{1/2} \nabla \tilde{q} Q $$

(20)

where $P$ is a matrix whose columns are the normal modes of the system $\tilde{q}_k$ expressed in normalized mass-weighted Cartesian coordinates $Q$ and $M$ is a diagonal matrix with the nuclear masses, one can reformulate it in terms of finite differences

$$ P^\dagger \Delta \tilde{q} = M^{1/2} \Delta Q $$

(21)

Thus, $\Delta \tilde{q}$ represents the array of displacements $\tilde{d}_i$ along the normal modes of the system when the difference is taken with respect to the reference point on the $i$—the electronic state. Rearranging gives the final working equation

$$ \tilde{d}_i = P^{-1} M^{1/2} \Delta Q $$

(22)

which allows to estimate how much every normal mode $\tilde{q}_k$ has to be displaced to connect two points in Cartesian coordinate space of the potential energy surface of the $i$—th electronic state. $\Delta Q$ is the difference in Cartesian coordinates between two geometries. The displacements $\tilde{d}_i$ can be related to spectroscopic parameters like the Huang-Rhys factors $S_{ik}$ or the reorganization energy $\lambda_{ik}$ of the system:

$$ S_{ik} = \frac{d_{ik}^2 \omega_k}{2\hbar} \quad \text{and} \quad \lambda_{ik} = \frac{d_{ik}^2 \omega_k^2}{2\hbar}, $$

(23)

where $\omega_k$ is the frequency of the $k$–th normal mode. With the normal mode analysis attention should be paid to global translational and rotational degrees of freedom which have to be removed prior to computing the difference in Cartesian coordinates. To this end we followed an iterative procedure.
Table 1: Reorganization energy in selected normal modes of Urd

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relying on the vectors of inertia in order to minimize the distance in space between two geometries as outlined in ref. [Kur01b].

In Tables 1 and 2 we list the modes accumulating the majority of the reorganization energy (> 90 cm$^{-1}$). Figure 24 shows the three modes accumulating the most reorganization energy.

The NMA shows that the 775 cm$^{-1}$ breathing mode characterized by a large amplitude of the $N_1C_6C_5$ angle (numeration according to Figs. 5 and 8) is the dominant mode in the mid-frequency range of both systems (< 1000 cm$^{-1}$). This observation is in agreement with the Fourier transform of the residuals of the transient absorption spectra (Figure 3 of the main text). The ring breathing mode, together with the $C_5C_6$ (1044 cm$^{-1}$) and $C_5C_6$ (1852 cm$^{-1}$) stretch modes dominates the excited state vibrational dynamics of Urd. We note the significant red-shift of the frequency of the $C_5C_6$ stretch mode with respect to its ground state value of ca. 1700 cm$^{-1}$. Notably, in 5mUrd the $C_5C_6$ stretching is not among the active modes in the excited state. Instead, a notable amount is inserted in the symmetric CO stretching (1672 cm$^{-1}$), not directly involved in the hydrogen out-of-plane bending deformation facilitating the internal conversion to the ground state.

To correlate the findings of the NMA with the results of the Fourier transform analysis of the residuals of the transient absorption spectra (Figure 3 in the main text) we traced the evolution of the main geometrical deformation associated with the 770 cm$^{-1}$ breathing mode, namely the $N_1C_6C_5$ angle, along the dynamics. Figure 25 shows heat maps of the fluctuation of the $N_1C_6C_5$ angle for Urd (top) and 5mUrd (bottom) as a function of time.
Supplementary Figure 23: Overlaid structures of the ground state and $\pi\pi^*$ excited state equilibrium geometries in Urd (left) and 5mUrd (right).

Table 2: Reorganization energy in selected normal modes of 5mUrd

<table>
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Supplementary Figure 24: Modes accumulating the most reorganization energy in Urd (top) and 5mUrd (bottom)

Supplementary Figure 25: Heat maps of the fluctuation of the $N_1C_6C_5$ angle (left) and their Fourier transform for Urd (top) and 5mUrd (bottom).
Fourier transform of the dynamics (until 250 fs for Urd and until 600 fs for 5mUrd) confirms that that $N_1C_6C_5$ angle exhibits a coherent oscillation with ca. 750 cm$^{-1}$ frequency, as predicted by the NMA.

Similarly, we also traced the dynamics of the $C_5C_6$ bond and the $H_9/CH_3C_5C_6H_{10}$ hydrogen/methyl out-of-plane bending involved in the internal conversion to the ground state. Figure 26 and 27 show the evolution of the $C_5C_6$ bond length and $H_9/CH_3C_5C_6H_{10}$ dihedral angle during the MD in Urd (top) and 5mUrd (bottom). Regarding the $C_5C_6$ bond length we observe immediate elongation to 1.65 Å after populating the $\pi\pi^*$ state, followed by coherent oscillatory dynamics in the first few hundred femtoseconds. As Urd rapidly decays to the ground state the $C_5C_6$ bond length is restored to its ground state equilibrium value of 1.38 Å within 0.5 ps. As 5mUrd remains in the excited state on a picosecond time scale we observe oscillations around 1.45 Å in the first 0.4 ps, i.e. the value of the $C_5C_6$ bond length in the $\pi\pi^*$ equilibrium. After 0.4 ps a shift of the average $C_5C_6$ length to smaller values is visible. Fourier transform of the dynamics (until 250 fs for Urd and until 600 fs for 5mUrd) shows that in Urd the stretching is governed in the excited state by a single mode around 1000 cm$^{-1}$. This matches perfectly with the 1040 cm$^{-1}$ active mode revealed by the NMA (Figure 24, top). In 5mUrd the stretching contributes to three modes, ca. 750 cm$^{-1}$, 1000 cm$^{-1}$ and 1250 cm$^{-1}$ which correspond to modes 7,9 and 10 in Table 2 whose Cartesian deformations show contribution by the $C_5C_6$ stretching.

The plot of the $H_9C_5C_6C_{10}$ dihedral angle evolution in Urd (Figure 27, top left) shows that it takes about 50-75 fs to distribute energy in the hydrogen out-of-plane bending mode. Between 50-250 fs we observe a scattered plot due to the decay to the ground state through the puckered conical intersection with values around 90° and -90°. 5mUrd takes considerably longer to distribute energy in the methyl out-of-plane bending mode (Figure 27, bottom left), i.e. around 250-300 fs. Puckering events are scattered in the plot.

We recognize that the aromatic ring exhibits a symmetry plane with respect to the hydrogen/methyl out-of-plane bending which is resembled in the two sets of conical intersections with the ground state which can be reached. Thus, in Figure 27, right, we plot the evolution of the absolute value of the dihedral $H_9/CH_3C_5C_6H_{10}$. This helps to recognize signs of coherent ultrafast deformation of the planarity of the aromatic ring in Urd where the hydrogen out-of-plane bending reaches values of up to 50-60° already after 25 fs before

\[^9\text{We note that the symmetry is broken in the presence of the surrounding (sugar and solvent) which might explain the preferred population of one of the two conical intersections in each nucleoside)}\]
going back to $0^\circ$ at 50 fs. The period associated with this ring deformation matches well the period (55 fs) associated with the 600 cm$^{-1}$ peak appearing in the Fourier transform maps of Urd shown in Figure 3 of the main draft suggesting that this may be a spectral signature of the coherent ring deformation observed at early times in simulations. Thereby, a candidate for a driving mode is a 600 cm$^{-1}$ hydrogen out-of-plane bending, immediately activated in the FC region. After the first period (ca. 50 fs) the coherence is partially destroyed and the individual trajectories reach the conical intersection region within a time window of three periods of the mode (most of the trajectories decay between 75 fs and 225 fs). Correspondingly Fourier transform of the heat maps (not shown) does not reveal clear vibrational signatures.
Supplementary Figure 26: Heat maps of the fluctuation of the C₅C₆ bond length (left) and their Fourier transform for Urd (top) and 5mUrd (bottom).

Supplementary Figure 27: Heat maps of the fluctuation of the H₉/CH₃C₅C₆H₁₀ dihedral (left: using real values; right: using absolute values) for Urd (top) and 5mUrd (bottom).
Supplementary Note 4:  
Materials and Steady-State Spectroscopy

5mUrd (97% purity) and Urd (99% purity) were purchased from Sigma-Aldrich and used as received. A phosphate-buffered saline (PBS) solution was prepared by dissolving 3.6 g of sodium dihydrogen phosphate and 4.26 g of sodium hydrogen phosphate in ultrapure water to obtain a pH 7.4 and a concentration of 15 mM. The 5mUrd and Urd in PBS solutions were prepared to obtain a concentration of 24.2 and 27.6 mM. Different concentrations (from 0.4 mM to 30 mM) were tested without finding significant differences in their absorption spectra.

The steady-state absorption spectra were measured at room temperature using a V-570 Jasco spectrophotometer. They were background corrected by subtracting the solvent spectrum measured at the same experimental conditions. The absorption spectra of the samples used for the experiments reported in the paper are shown in Figure 28.

Supplementary Figure 28: Linear absorption spectra for Urd (blue) and 5mUrd (red) together with the pump pulse spectrum (purple).
Supplementary Note 5:
Transient absorption spectroscopy

5.1 Transient absorption spectra

Figure 29 shows the transient absorption spectra at different time delays between pump and probe (parallel polarizations) for 5mUrd (left) and Urd (right).

In 5mUrd the SE signal progressively decays within 1 ps presenting a red-shift. This decay is concomitant with that of the photo-induced absorption band peaked at 2 eV (PA1). A third spectral feature around 4.2 eV (PA2) is formed within 100 fs and decays blue-shifting on the picosecond time scale.

In Urd the SE band completely vanishes within 200 fs, followed by the formation of PA band (PA2) which reaches its maximum in about 300 fs. A photo-induced absorption signal in the visible (PA1) progressively decays in 500 fs.

Supplementary Figure 29: Transient absorption spectra at selected time delays for a) 5mUrd and b) Urd. The vertical line at 3.65 eV stands for the two probe spectral ranges used in the experiment.
5.2 Measurements on the pure solvent

Figure 30 shows the dynamics at selected energies of the phosphate buffer solution. Transient absorption spectra at different time delays between pump and probe (parallel polarizations) are reported in figure 31. No signal from solvated electrons was noticed under our experimental conditions.

Supplementary Figure 30: Dynamics of the PBS solvent at selected energies.
5.3 Measurements at magic angle and orthogonal polarizations

Figures 32 and 33 present the experimental and theoretical transient absorption maps for Urd at magic angle and crossed polarizations, respectively. In addition to the signals identified in the main text, a photo-induced absorption band (PA3) becomes visible when using orthogonal polarizations, due to the partial suppression of the SE signal.

Figures 34 and 35 present the experimental and theoretical transient absorption maps for 5mUrd at magic angle and crossed polarizations, respectively. In addition to the signals identified in the main text, a photo-induced absorption band (PA3), which follows the same trend as PA1, becomes visible for perpendicular polarizations.

Supplementary Figure 31: Transient absorption spectra at selected time delays for the PBS solvent.
Supplementary Figure 32: Experimental (left) and simulated (right) transient absorption spectra maps and selected dynamics of Urd with the polarization of pump and probe at magic angle.
Supplementary Figure 33: Experimental (left) and simulated (right) transient absorption spectra maps and selected dynamics of Urd with orthogonal polarizations of pump and probe.
Supplementary Figure 34: Experimental (left) and simulated (right) transient absorption spectra maps and selected dynamics of 5mUrd with the polarization of pump and probe at magic angle.
Supplementary Figure 35: Experimental (left) and simulated (right) transient absorption spectra maps and selected dynamics of 5mUrd with orthogonal polarizations of pump and probe.

5.4 Components of the simulated transient spectra

Figures 36 and 37 present individual components in the simulated transient absorption maps of Urd and 5mUrd at magic angle. Specifically, signals involving the ground state (SE and PA2, labels according to Figs. 1 and 2 of the main text) are shown separately from the remaining PA signals. We have deliberately used the same scale as in the total maps (Figures 32 and 34). This allows to appreciate the destructive overlap between signals of opposite sign, as well as the presence of characteristic $S_1$ fingerprint PA which remain covered in the total spectra. Note that the PA2 signal in the spectra of 5mUrd in Figure 37 is dominated by excited state PA at early times and by ground state PA at later times.
Supplementary Figure 36: Components of the simulated transient absorption spectra (with the polarization of pump and probe at magic angle) of Urd; left: Stimulated emission (SE) and photoinduced absorption of the ground state (PA2); right: excited state (S_1) photoinduced absorption signals PA1, PA3 and PA4. Due to destructive overlap PA3 and PA4 are not visible in Figure 1 of the main text. Same scale used as in the total map (Figure 32).

Supplementary Figure 37: Components of the simulated transient absorption spectra (with the polarization of pump and probe at magic angle) of 5mUrd; left: Stimulated emission (SE) and photoinduced absorption of the ground state (PA2); right: excited state (S_1) photoinduced absorption signals PA1, PA2 and PA3. Note that signal PA2 is dominated by excited state PA at early times and by ground state PA at later times. Due to destructive overlap PA3 is not visible in Figure 2 of the main text. Same scale used as in the total map (Figure 34).
5.5 Measurements in the picosecond time scale

A complementary set of measurements was taken at longer time scales. Figures 39 and 40 present the maps of Urd (up to 150 ps) and 5mUrd (up to 5 ps). In both measurements, the transient spectra were recorded at magic angle. While in the first case we observe a long living signal (which survives beyond our time window), in the second case the signals in this spectral range have vanished within 5 ps.

Supplementary Figure 38: Transient absorption spectra map and selected dynamics of Urd up to 150 ps.
Supplementary Figure 39: Transient absorption spectra of Urd at selected time delays.

Supplementary Figure 40: Transient absorption spectra map and selected dynamics of 5mUrd up to 5 ps.
Supplementary Note 6:
Data analysis

6.1 Fitting procedure

The following model was used to fit the experimental transient absorption dynamics at selected probe photon energies:

\[
h(t) = g(t) + A_{\text{rise}} \left[ 1 - \exp \left( -\frac{t}{\tau_{\text{rise}}} \right) \right] \left[ A_1 \exp \left( -\frac{t}{\tau_1} \right) + A_2 \exp \left( -\frac{t}{\tau_2} \right) \right] H(t)
\]

where \( H(t) = u(t) \otimes f(t) \) is the convolution of the Heaviside function, \( u(t) \), and the instrumental response function, \( f(t) \). The latter was approximated by a Gaussian, whose temporal width was a free parameter in the model. The function \( g(t) \) models the coherent artifact [Lor02]:

\[
g(t) = A_{\text{art}} \left[ t \exp \left( -2 \left( \frac{t}{\tau} \right)^2 \right) - (t - \tau_{\text{GVD}}) \exp \left[ -2 \left( \frac{t - \tau_{\text{GVD}}}{\tau^2} \right)^2 \right] \right]
\]

being \( \tau \) the pump pulse duration and \( \tau_{\text{GVD}} \) the difference of passage times through the medium between the pump and probe pulses.

The following table reports the time constants found for the dynamics of Urd reported in the main text. Note that we in addition we included an additional offset term to account for the hundreds of picoseconds signal.

<table>
<thead>
<tr>
<th>Energy (eV)</th>
<th>( \tau_{\text{rise}} ) (fs)</th>
<th>( \tau_1 ) (fs)</th>
<th>( \tau_2 ) (fs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.0</td>
<td>–</td>
<td>125</td>
<td>–</td>
</tr>
<tr>
<td>3.6</td>
<td>–</td>
<td>80</td>
<td>1000</td>
</tr>
<tr>
<td>4.1</td>
<td>150</td>
<td>1000</td>
<td>–</td>
</tr>
</tbody>
</table>

Table 3: Time constants for dynamics at selected probe photon energies in Urd

The same procedure was applied for 5mUrd finding the following time constants:
Table 4: Time constants for dynamics at selected probe photon energies in 5mUrd

<table>
<thead>
<tr>
<th>Energy (eV)</th>
<th>$\tau_{\text{rise}}$ (fs)</th>
<th>$\tau_1$ (fs)</th>
<th>$\tau_2$ (fs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.0</td>
<td>–</td>
<td>135</td>
<td>1160</td>
</tr>
<tr>
<td>3.6</td>
<td>–</td>
<td>70</td>
<td>770</td>
</tr>
<tr>
<td>4.1</td>
<td>70</td>
<td>–</td>
<td>1600</td>
</tr>
</tbody>
</table>

Supplementary Figure 41: Experimental data (blue) and fit (red) for selected dynamics.
6.2 Excited-state vibrational modes

In order to extract the oscillatory pattern a Fourier low-pass filter was applied to the transient absorption spectra maps. The first 100 fs were removed in order to avoid artifacts in the 2D Fourier map due to the presence of the coherent artifact. To keep a good signal-to-noise ratio, we analyze a temporal window up to 400 fs in Urd and 1 ps in 5mUrd. The residual maps are shown in Figure 42. In both cases we observe a phase jump coinciding with the maximum of the SE signal, thus confirming that the oscillations belong to vibrational motions of the excited state wave-packet.

Supplementary Figure 42: Residuals showing an oscillatory pattern in 5mUrd (left) and Urd (right).
Supplementary Note 7:

References

