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CHEMISTRY

Excited state intramolecular proton transfer in hydroxyanthraquinones: Toward predicting fading of organic red colorants in art

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Compositionally similar organic red colorants in the anthraquinone family, whose photodegradation can cause irreversible color and stability changes, have long been used in works of art. Different organic reds, and their multiple chromophores, suffer degradation disparately. Understanding the details of these molecules' degradation therefore provides a window into their behavior in works of art and may assist the development of improved conservation methods. According to one proposed model of photodegradation dynamics, intramolecular proton transfer provides a kinetically favored decay pathway in some photoexcited chromophores, preventing degradation-promoting electron transfer (ET). To further test this model, we measured excited state lifetimes of substituted gas-phase anthraquinones using high-level theory to explain the experimental results. The data show a general structural trend: Anthraquinones with 1,4-OH substitution are long-lived and prone to damaging ET, while excited state intramolecular proton transfers promote efficient quenching for hydroxyanthraquinones that lack this motif.

INTRODUCTION

Exposure to light degrades many molecules when absorption of ultraviolet (UV) and visible wavelength photons places them in an energetically excited state prone to chemical and physical changes. Molecules susceptible to photodegradation are ubiquitous in both natural and synthetic systems, and undesired effects of this degradation can include a wide range of phenomena, such as the appearance of abnormal skin cells marking melanoma and the drop of efficiency in polymer solar cells via photobleaching. One notable form of organic photodegradation with relevance to the broad field of cultural heritage research is the fading of certain chromophores in works of art, which leads to visual change in a medium where meaning and value are often predicated on color.

Natural organic red colorants of either plant or scale insect origin are important traditional sources of red hues (1). These colorants have high tinting strength and are therefore present on artifacts in low concentration; as little as subnanogram concentrations of these colorants may be required to achieve the desired color saturation (2). Many traditional organic red colorants (including madder, alizarin, cochineal, lac dye, brazilwood, and dragon's blood) are compositionally similar: These anthraquinones (AQs) and hydroxyanthraquinones (HAQs) have long been used as lake pigments and contain primarily hydroxyl, carboxylic acid, and carbonyl moieties on a conjugated aromatic backbone. While commercial interest in AQs and HAQs has recently expanded to include pharmaceutical applications (3, 4), the use of natural AQ derivatives (see Fig. 1) as lake dyes (insoluble dye-mordant complexes, typically precipitated with Al or Ca cations) has long attracted the attention of fine artists and craftsmen, and these pigments continue to be used today. The irreversible fading of organic red colorants in art is well documented and greatly affects the perception of masterpieces from antiquity to

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the present day. For example, recent publications have highlighted alteration of organic red colorants in objects as disparate as medieval manuscript illuminations to paintings by Vincent van Gogh (5–7).

One of the simplest HAQ molecules to be used as a lake pigment is 1,2-dihydroxyanthraquinone (1,2-HAQ), also known as alizarin. Alizarin is produced naturally by multiple species related to *Rubia tinctorum* (common madder) where it is a primary chromophore along with 1,2,4-trihydroxyanthraquinone (1,2,4-HAQ), also known as purpurin (1). It is well documented in the cultural heritage and condensed phase scientific communities that alizarin is more photostable than purpurin (8–11). Condensed phase photodynamics studies of these AQs suggest that the difference in photostability may stem from an excited state pathway, specifically an excited state intramolecular proton transfer (ESIPT), in alizarin that dissipates photonic energy to regain a stable ground-state structure on a time scale much faster than the reaction rates of degradation (11, 12).

ESIPT constitutes one of the fastest reactions known (13, 14), and planar organic molecules containing acidic and basic functional groups connected by an intramolecular hydrogen bond often have photophysics driven by this proton transfer. In these molecules, excited state tautomerization occurs through the excitation of the ground-state enol form to an electronically excited intermediate where a rapid (subpicosecond time scale), often energetically barrierless, enol to keto proton transfer takes place. Radiationless decay follows this transfer, bringing the excited keto form back to the ground state, and the stable ground-state enol reforms through a barrierless back proton transfer (15). Molecules that undergo ESIPT are typically identified by dual band fluorescence with a large degree of Stokes shift from the ESIPT lower well equilibrium geometry.

Formative work examining this mechanism in AQs/HAQs was done in the 1980s (16); laser-induced fluorescent spectroscopy studies (17), including that of Flom and Barbara (15), determined that the presence of a dual 1,4-electron donating group (e.g., -OH) precludes dual fluorescence, likely indicating a high barrier toward ESIPT. This qualitative result was later supported by the nodal plane model by Nagaoka and Nagashima (18). More recent transient spectroscopic work has measured the rate of ESIPT (19–21) as a femtosecond process

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Fig. 1. The molecules of this study. Molecules having a 1,4-OH motif are above the images, and those without 1,4-OH motif are below them. Isomers within the LE ZPE corrected structure (labeled I) (~25 kJ/mol) are also shown with their relative energy values (kJ/mol) with respect to the LE form. Central to this figure is the common madder root and its primary chromophores alizarin and purpurin. Photo credit: Jacob Berenbeim, UCSB

and correlated this with pigment fading (11, 22). However, these are all condensed phase experiments where the effects of secondary molecules, such as solvent molecules or other pigment components, cannot be completely excluded, which is particularly important since intermolecular effects are widely known to affect relaxation (20, 23). To measure the unimolecular dynamics of the AQ and HAQ molecules require gas-phase experiments. Note that in the context of art materials, much of the available literature deals only with alizarin and purpurin as exemplars of the HAQ system (24). Therefore, there remains a need to examine a broader range of these important molecules in the gas phase to fully explicate the observed differences in the relative photostabilities of the basic chromophores themselves. By studying a larger set of isolated HAQs, it is possible to identify the key structural motifs that determine the photodynamics of this class of compounds and further explicate the observed photochemical behavior of alizarin and purpurin. Such a fundamental study can then inform a fuller understanding of the more complex systems found in works of art, which will include binding media, the support, and other pigments in addition to the organic red colorant in question.

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Here, we address this need and report an experimental and computational case study on the effects of proton transfer on the excited state lifetimes of seven related HAQs as isolated chromophores. We measured intrinsic lifetimes of a series of neutral HAQ molecules in a jet-cooled molecular beam by time-resolved, pump-probe, two-color resonant two-photon ionization (2C-R2PI) spectroscopy to elucidate the relaxation dynamics occurring at the lower limit of the excited state potential energy surface (PES). For each molecule studied, these data provide the first 2C-R2PI action spectra, building a library to allow their identification in works of art. We also obtained the excited state lifetimes from both the lowest energy (LE) vibronic transition and at higher internal energy (~500 to 1000 cm⁻¹) and partial mid-infrared (IR) characterization of the ground-state hydrogen bond vibrations with IR hole burning of the R2PI probe signal for selected compounds. These measurements allow us to study these chromophores in selected unique tautomeric forms. The resulting detailed vibrationally and tautomerically resolved excited state lifetime data reveal the intrinsic properties of the chromophores and serve as the basis for high-level quantum computational modeling of the excited state dynamics. We explore the implications of the resulting data for the photodegradation of these compounds. Our results show a trend of shorter excited state lifetimes for the structures for which there is evidence for ESIPT, as derived from dual fluorescence by Flom and Barbara (15) and others. We find that both phenomena correlate with structural motifs with a specific double hydrogen bonding pattern, and this observation provides experimental evidence to support predictions for the expected stability of HAQs beyond alizarin and purpurin in works of art.

RESULTS

Analytes studied and R2PI spectroscopy

Figure 1 shows the series of substituted HAQs analyzed. The analytes represent structures with and without the 1,4-OH substitution motif postulated to be determinative of ESIPT properties (11, 15). Several of these molecules are also found in artists' red lake pigments. The molecular structures shown in Fig. 1 are planar in all cases. In each molecule, the LE structure (indicated by the Roman numeral I) is that of the 9,10-AQ with carbonyl groups rather than hydroxyl moieties on the central ring of the backbone. The interpretation of R2PI and pump-probe spectra requires an understanding of the possible structures, which are most likely the LE isomers, present in the molecular beam. Figure 1 shows all isomers with energies calculated to be within 25 kJ/mol of the LE structure; isomeric forms with higher energies appear in the Supplementary Materials. The center of Fig. 1 shows the two primary common madder chromophores alizarin (1,2-HAQ) and purpurin (1,2,4-HAQ). Both have multiple low-energy isomers, as do 1,2,5,8-HAQ and 1,4-HAQ. Three of the molecules examined-1-HAQ, 1,5-HAQ, and 1,8-HAQhave only a single LE structure (the rotamers for each of these molecules are 52.3, 53.2, and 47.4 kJ/mol higher in energy than the forms shown, respectively). In the figure and throughout, we use the nomenclature of HAQ for all compounds, ignoring the more formal designations of DHAQ and THAQ for the di- and trihydroxy forms, respectively, as the numbered prefixes already indicate the number of hydroxyl substituents.

Figure 2 presents the R2PI spectra of the HAQs from Fig. 1. The origins, corresponding to S_{0-0} transitions, of molecules with a 1,4 motif are about 2000 cm⁻¹ lower in energy than those without the 1,4 motif.



Fig. 2. Picosecond 2C-R2PI spectra of each HAQ in this study (OPG + 213 nm). Intensities have been normalized.

The origins of 1,2,5,8-HAQ (19,661 cm⁻¹), 1,2,4-HAQ (19,845 cm⁻¹), and 1,4-HAQ (19,920 cm⁻¹) all lie within 300 cm⁻¹ of each other, and each is characterized by sharp, laser linewidth limited bands more than a ~1000 cm⁻¹ range. In contrast to this, the spectra for HAQs without the 1,4 motif are broader, suggesting shorter excited state lifetimes. Their origins—1-HAQ (21,645 cm⁻¹), 1,2-HAQ (21,748 cm⁻¹), 1,5-HAQ (21,321 cm⁻¹), and 1,8-HAQ (22,031 cm⁻¹)— are located over a larger energy spread, although still within ~700 cm⁻¹ of each other. The difference between these two groups of spectra suggests that the excited state properties strongly depend on the presence or absence of the 1,4-OH structural motif.

Pump-probe spectroscopy

We performed picosecond pump-probe measurements for each HAQ from Fig. 1 on both the origin transition and on one additional higher energy transition determined from the R2PI spectra. The measurement using the higher energy transition probed whether additional relaxation pathways become available at higher energies, which would likely be evidenced by a change in excited state lifetime. Figure 3 shows the effect of structure on excited state lifetime. The molecules with the 1,4-OH motif, shown at the top in Fig. 3A in a red box, have nanosecond lifetimes at their LE transitions. In contrast, those molecules without the 1,4-OH motif, shown at the bottom of Fig. 3A in a blue box, have subnanosecond lifetimes.

Coincidently, the primary madder chromophores (1,2,4-HAQ and 1,2-HAQ) have the longest and the shortest excited state lifetimes, respectively, of all molecules measured. Figure 3B shows the pump-probe traces of these two end-members (picosecond pumpprobe data for the other molecules are shown in fig. S1). Each of these traces can be adequately fit using a single exponential, providing the excited state lifetime of each molecule. When probed at the origin, 1,2,4-HAQ (purpurin) has an excited state lifetime of 5.4 ns, while 1,2-HAQ (alizarin) has an excited state lifetime of only 120 ps. Other HAQ series have lifetimes intermediate between these two; e.g., 1-HAQ has a measured lifetime of 290 ps, while 1,4-HAQ has a measured lifetime of 1.7 ns. In general, excitation at higher energies shortens the lifetime, as shown in Fig. 3A (black bars). 1,4-HAQ



Fig. 3. Results of picosecond pump-probe spectroscopy. (A) Pump-probe lifetime as measured at the origin transition (red bars) and at excess energy (black bars) arranged in order of decreasing measured lifetimes. The excess energy transition used is noted next to the black bars. (B) Pump-probe traces of the molecules with the longest and shortest measured excited state lifetime, 1,2,4-HAQ and 1,2-HAQ, respectively, plotted over 500 ps. The pump-probe data are fit to a curve (blue trace), which is the sum of a single exponential decay (green trace) convolved with a Gaussian component (red trace) representative of our instrument response function.

provides an exception to this trend, with a slight increase in excited state lifetime when measured at a transition $+655 \text{ cm}^{-1}$ from the origin (from 1.7 to 1.9 ns). This slight increase is likely an artifact due to the short observation period relative to the excitation lifetime. For completeness, we also carried out nanosecond pump-probe measurements on the two end-members (1,2,4-HAQ and 1,2-HAQ; data shown in fig. S2). These measurements show no component at longer time scale than those found in the picosecond measurements.

Intramolecular hydrogen bonding characterization and structural confirmation

To understand the hydrogen-bonding environment of the isomer of species in the molecular beam with and without the 1,4-OH motif, we performed IR hole burning. IR-UV results in mode II indicate that for 1,2,4-HAQ and 1,4-HAQ, the R2PI results are from a single conformation (see fig. S3).

We also performed hole burning in mode I on 1,2,4-HAQ, 1,4-HAQ, and 1,8-HAQ while probing the origin of each, as shown in Fig. 4. In all three spectra, the peak at \sim 3100 cm⁻¹ represents stretching modes associated with the -OH hydrogen bound to a carbonyl group. In the case of 1,8-HAQ, this peak is broadened by the competitive sharing of the carbonyl with intramolecular hydrogen bonds, 180° about the oxygen atom (9 position). In the case of 1,2,4-HAQ, the peak at 3570 cm⁻¹ results from the in-plane hydroxyl (2 position), which is hydrogen bound to the neighboring hydroxyl (1 position). The 470-cm⁻¹ separation is a direct measurement of different hydrogen bond environments: between carbonyl and hydroxy intramolecular bonded -OH stretches.

Figure S4 also shows calculated LE spectra convoluted with a Lorentzian linewidth of 3 cm⁻¹. These calculations reproduce the experimental patterns but not the exact frequencies, reflecting the fact that these are unscaled harmonic calculations. Future work with anharmonic calculations at a higher level may provide further details on structures.

Theoretical calculations: 1-HAQ and 1,4-HAQ

To computationally model the difference in excited state dynamics in the two structural motifs, with and without the 1,4 substitution,



Fig. 4. IR hole burning spectra. Spectra were taken in mode I of 1,2,4-HAQ, 1,4-HAQ, and 1,8-HAQ probed at their origin R2PI transition. a.u., arbitrary units.

we performed detailed calculations on the most basic representatives of the two structural families, namely, 1-HAQ and 1,4-HAQ. **1-HAQ: Ground and excited state**

Figure 5 shows the calculated energetic landscapes of 1-HAQ. The ground-state global minimum of the keto form [I in Fig 5, hereafter referred to as K(I)] with the proton attached to the O₁ oxygen atom is planar and stabilized by a single intramolecular hydrogen bond between two oxygen atoms as proton acceptor: O_1 —H... O_9 . The proton-transferred form [enol form, II in Fig. 5, hereafter referred to as E(II)] with the proton attached to the O₉ oxygen atom is not stable in the S₀ state, and its geometry optimization transfers the proton back to the O₁ oxygen atom to reform the global minimum form K(I). Possible rotation of the O₁H group could generate the second lowest S₀-state minimum, the keto rotamer (III in Fig. 5) structure. However, the O₁H rotation breaks the intramolecular hydrogen bond



Fig. 5. Potential-energy energy profiles of 1-HAQ. Profiles shown are of the S₀ state (black circles), the S₁($\pi\pi^*$) state (blue squares), and the S₁($n\pi^*$) state (red diamonds) of the 1-HAQ molecule as a function of the torsional reaction path (**A** and **C**) and the hydrogen transfer reaction path (**B**). Full lines (full symbols): Energy profiles of reaction paths determined in the same electronic state. Dashed lines (empty symbols) represent ground-state energy calculated for the geometry optimized in the given excited state $n\pi^*$ (red) or $\pi\pi^*$ (blue). Purple rectangles highlight the $n\pi^*$ and $\pi\pi^*$ intersections.

in the 1-HAQ structure, which destabilizes the rotated form compared to the global minimum K(I) by 0.52 eV (50 kJ/mol). Furthermore, the rotamer (III) minimum is separated from the global minimum by a S₀-state energy barrier of 0.16 eV (15 kJ/mol). The fourth form, IV with the proton attached to the O₉ atom, has much higher energy and can be excluded from this study. This energetic profile indicates that the 1-HAQ molecule should exist in the K(I) form in the ground state.

When 1-HAQ is irradiated, the ground-state K(I) form is photoexcited to the lowest $\pi\pi^*$ excited state, $S_2(\pi\pi^*)$ with vertical energy $\Delta E^{VE} = 3.38$ eV at the S₀-state minimum geometry. At this optimized ground-state geometry of K(I), S₁ is a dark $n\pi^*$ state below the $S_2(\pi\pi^*)$. The lowest $\pi\pi^*$ excited state does not have a stable minimum in the K(I) form. Therefore, as long as the system remains in the excited $\pi\pi^*$ state, the proton will transfer between the two oxygen atoms (from O₁ to O₉) along a barrierless path to form the proton-transferred $\pi\pi^*$ excited state form, E(II), as indicated by the minimum of the blue curve (full squares) in the central panel of Fig. 5.

As depicted in Fig. 5 (highlighted with purple rectangles), the $n\pi^*$ and $\pi\pi^*$ excited states intersect in the vicinity of the $S_1(\pi\pi^*)$ -state minimum, E(II). Consequently, the $n\pi^*$ state can be populated directly from the $\pi\pi^*$ excited state. The $n\pi^*$ state has two minima, at the K(I) and E(II) forms, and the adiabatic energies of both minima are lower than that of the $S_1(\pi\pi^*)$ E(II) minimum. Once in the nonfluorescent $n\pi^*$ state, the system seeks another decay channel. The S_1 - S_0 energy gap lowers for the $n\pi^*$ state to 1.38 eV for the E(II) excited state form, as depicted by the vertical purple line in Fig 5. As shown in Fig. 5, if the system is in $n\pi^*$ -E(II) minimum, rotation of the O₉H group may lead to further notable decrease of the S_1 - S_0 energy gap (see red traces) until S_1 and S_0 meet at a C-C-O₂-H dihedral angle of about 100°. This pathway provides an additional channel for excited state may make this process less efficient.

1,4-HAQ: Ground and excited state

The addition of the OH group to position 4 of the 1-HAQ molecule (i.e., the introduction of the 1,4-OH motif) results in substantial changes,

both structurally by forming a second intramolecular hydrogen bond and photophysically by changing the ordering of the excited states.

In contrast to 1-HAQ, 1,4-HAQ has two intramolecular hydrogen bonds: O1-H...O9 and O4-H...O10, which bridge the molecule on opposite sides and ensure planarity of the system. Figure 6 displays the two-dimensional PES of 1,4-HAQ, with panels (A), (B), and (C) showing the S_0 , S_1 , and S_2 states, respectively; panel (D) illustrates the possible tautomers of 1,4-HAQ, both single hydrogen transfers (KE and EK), and double hydrogen transfer (EE). The *x* axes of Fig. 6 (A to C) are the reaction coordinate for the hydrogen displacement from O_1 , while the y axes are the reaction coordinate for the hydrogen displacement from O_4 . Moving along the x axis moves the hydrogen from O_1 to O_9 , while the y axis moves the hydrogen from O_4 to O_{10} . Thus, the top right corner of each energy surface is the EE tautomer, the product of both hydrogen transfers. There exists a tautomer in each corner of each PES, labeled accordingly. In addition to the global minimum form-the KK form with protons attached to O1 and O4-the EE form, with the protons attached to O₉ and O₁₀, provides an additional local minimum, 0.36 eV higher in energy. As shown in Fig. 6A, the EE minimum is separated from the KK form by a relatively small S_0 -state energy barrier of ~+0.15 eV. The presence of this additional stable tautomer and the energy barrier (in contrast to the 1-HAQ system, which lacks an energy barrier) might be explained by the fact that any proton transfer usually requires shortening of the interatomic distance between two proton accepting centers (here, two oxygen atoms). While one interatomic distance, e.g., $O_1...O_9$, contracts, the other, i.e., $O_4...O_{10}$, must simultaneously lengthen. This effect is not energetically favorable, creating the barrier in S₀.

In these coordinates for the $\pi\pi^*$ (Fig. 6B), the KK tautomer is the minimum by ~0.15 eV. A single or double proton transfer to generate any other tautomer would be uphill and unfavorable. The vertical excitation energies, ΔE^{VE} , to the lowest excited $\pi\pi^*$ state of the KK and EE forms of the 1,4-HAQ molecule are 2.80 and 2.58 eV, respectively (for a complete tabulation of results, see tables S1 and S2). Both ΔE^{VE} values for 1,4-HAQ are lower than the respective value of 3.38 eV for 1-HAQ, which is consistent with the π -electron-donating character of the OH group (25), which should result in stabilization of the $\pi\pi^*$ states (Fig. 6C) in the absorption ladder of both tautomeric forms of 1,4-HAQ are at least 0.5 eV above the lowest $\pi\pi^*$ states and therefore inaccessible.

The properties of the optimized excited state forms of the molecules are gathered in table S1. KK and EE, respectively, each have the same ground state and $\pi\pi^*$ excited state geometries. Both excited state forms are almost isoenergetic. Moreover, a barrier of only 0.15 eV separates the S₁ states of these two forms. At the same time, the $n\pi^*$ state PES lies almost entirely above the fluorescent $\pi\pi^*$ state. Such a situation greatly reduces the probability for nonradiative decay for the 1,4-HAQ in comparison to the 1-HAQ system.

DISCUSSION

The experimental and computational results from this series of molecules demonstrate that the presence or absence of a hydroxyl substitution at the 4 position has profound effects on the spectroscopy and dynamics of substituted AQs. When interpreting spectral and pump-probe results, it is important to keep in mind that multiple



Fig. 6. PESs of 1,4-HAQ. (**A**) Minimum PES of the S_0 state, (**B**) the $S_1(\pi\pi^*)$ singlet excited state, and (**C**) the $S_1(n\pi^*)$ singlet excited state of the 1,4-HAQ molecule as a function of the hydrogen transfer reaction path as a function of two coordinates: $R(O_1...H)$ and $R(O_4...H)$. (**D**) Schematic indication of the structures at the four combinations of minimum and maximum *R* values. The location of each tautomer on the PES is also labeled.

tautomeric forms are possible (as shown in Fig. 1). In general, in jet cooling conditions, only the LE forms exist, and the mode II IR-UV data for 1,2,4-HAQ and 1,4-HAQ suggest those data to be exclusively from a single tautomer (although it is possible that signal arises from complimentary rotamers in Fig. 1). We assume that the other compounds also to be exclusively or at least predominantly in the LE tautomeric form, but the possible existence of other tautomers in the beam cannot be definitively excluded.

With this limitation in mind, the experimental and computational evidence can be summarized as follows: (i) The origin transition for HAQs with the 1,4 motif is about 2000 cm⁻¹ to the red of those without the 1,4 motif, as shown in Fig. 2. (ii) The excited state lifetime is approximately an order of magnitude shorter for those without the 1,4 motif, going from a few nanoseconds to a few hundred picoseconds at the origin transition, as shown in Fig. 3. Relative to the 1-OH motif, there is a 15× average longer lifetime for the molecular species with the 1,4-OH motif. (iii) In systems without the 1,4 motif (and therefore with short excited state lifetimes), proton transfer between the two oxygen atoms to form the proton-transferred $\pi\pi^*$ excited state form is likely, while in systems with the 1,4 motif, more energetically favorable forms are likely to be present, limiting the probability for nonradiative decay pathways.

The photobleaching effects of common madder chromophores, 1,2,4-HAQ and 1,2-HAQ, were recently investigated by Tan *et al.* (11) by counting emissive events in time (photoblinking) of these dyes under inert (N_2) irradiant conditions on glass. They measured

that alizarin is able to absorb and emit four times longer (and over many more events) than purpurin, which they argue to be due to a long-lived excited state of purpurin that degrades through electron injection to the glass slide. This decay pathway is consistent with the literature (19, 21, 26, 27). One cannot compare absolute lifetime values between gas-phase and condensed phase states, but the trend in the condensed phase correlates with the inherent lifetime trends presented here that 1,2-HAQ is ~45 times shorter lived at its vibrationless transition than 1,2,4-HAQ. The experimental data presented here seem to suggest that this pathway may be common to other substituted HAQs, with variations in the excited state lifetimes determined by the exact substitution arrangement.

Our theoretical investigation focused on the excited state potentials of 1-HAQ and 1,4-HAQ, since the experimental work showed that these motifs represent two distinct families of molecules. For 1-HAQ, the computations explain the experimentally observed short lifetimes by an energetically downhill process from the initially populated $\pi\pi^*$ state. Along the proton transfer coordinate, the $n\pi^*$ state is crossed twice, leading to the minimum of E(II). Following this pathway, the energy gap to the ground state is 1.38 eV, allowing for nonradiative decay, although there may be additional involvement of an out of plane torsion leading to a conical intersection. These dynamics are consistent with the relatively short lifetimes observed for the HAQs without a 1,4 motif. After reaching the ground state, it is a downhill path to transfer the proton back and complete the photocycle, recovering the K(I) tautomer.

The addition of the OH group to the 4 position of the HAQ skeleton results in several critical changes to ground and excited state profiles. First, in the 1,4-HAQ molecule, there is a second S₀-state minimum, the EE characterized by the two protons being transferred to the oxygens on carbons 9 and 10. In addition, the shape of the $\pi\pi^*$ excited state in 1,4-HAQ is much shallower than in 1-HAQ, and the photoexcitation of the KK form does not result in a barrierless proton transfer to the EE form as in 1-HAQ. The most prominent effect is that the $\pi\pi^*$ excited state PES lies below the $n\pi^*$ excited state PES, an effect of electron-donating property of the OH group, which tends to stabilize the $\pi\pi^*$ excited state versus the $n\pi^*$ state. This explains the red shift of 1,4 motif versus those molecules that lack it (25, 28). Because of the stabilization of the $\pi\pi^*$, the $n\pi^*$ is not accessible at the excitation energies used. This effect inhibits nonradiative decay from the excited $\pi\pi^*$ to S₀ via crossing to the $n\pi^*$, as is the case for 1-HAQ. A major effect of 4-OH substitution is related to the unique stabilization of the KK($\pi\pi^*$), lacking the downhill proton transferred minimum of the $\pi\pi^*$ excited state in 1-HAQ. The analogous initial $\pi\pi^*$ state in 1-HAQ undergoes proton transfer in a barrierless manner, leading to picosecond excited state lifetimes. However, this proton transfer shortens the interatomic distance between the two proton accepting oxygen atoms. With the 1,4 motif, while one interatomic distance, e.g., O1...O9, contracts, the other, i.e., O4...O10, simultaneously lengthens, creating a barrier in the excited state potential. As a result, we find a single nanosecond time scale lifetime for the molecules with a 1,4 motif, implying that instead of undergoing ESIPT, they relax from the initially populated $\pi\pi^*$ state. This model also explains why 1,5-HAQ and 1,8-HAQboth doubly hydrogen bound-behave photodynamically like 1-HAQ.

We investigated the possibility of longer-lived states such as triplets with a nanosecond ionization source for 1,2-HAQ and 1,2,4-HAQ but have not observed these states experimentally. However, Mohammed *et al.* (20) reported the formation of a long-lived triplet state with high quantum yield in a study of 1,8-HAQ in a series of nonpolar, polar aprotic, and polar protic solvents. Furthermore, work in this area is therefore justified to clarify whether long-lived triplets may form in some systems.

As seen by Flom and Barbara (15) and explained by Nagaoka and Nagashima (18), the 1-HAQ motif exhibits dual fluorescence due to ESIPT. As the main excited state pathway, this explains the photostability of molecules lacking the 1,4-HAQ motif observed in works of art (by, e.g., alizarin). In contrast, molecules with the 1,4-HAQ motif lack dual fluorescence and the corresponding ESIPT properties. At the same time, molecules with the 1,4-HAQ motif are noted for their photodegradation in works of art (e.g., purpurin). We have corroborated these phenomena both experimentally and theoretically in the gas phase, indicating that this is a fundamental characteristic of the AQ molecules used to create lake dyes and not due to other effects such as interactions with the substrate on which the dyes are precipitated, the binding media, or other pigments that may be present in the complex system of a work of art. Molecules with the 1-HAQ motif have ~15× shorter excited state lifetime than molecules with the 1,4-HAQ motif, explaining the former's photostability. The energetically downhill process of ESIPT accounts for the shorter lifetime of the 1-HAQ motif. Meanwhile, the geometries resulting from ESIPT of the 1,4 motif are all higher in energy than the initially excited, nonproton transfer state, making proton transfer highly unfavorable. Without this process available, the 1,4 motif has a much longer excited state lifetime and worse photostability.

Taken with the body of work done on UV radiation on anthracene by Mallakin et al.(29, 30) on the production of toxic species (i.e., reactive, typically ¹O₂) through the stepwise formation and further irradiation of AQs and HAQs and the work by Nagaoka and Nagashima (18) on the quenching of singlet oxygen $({}^{1}O_{2})$ by ESIPT HAQs, as well as the previous experimental and theoretical work done on alizarin and purpurin (8-12, 27), the data presented here form a complex picture of the degradation of the madder colorantsand AQ dyes more generally-in cultural heritage materials. These pigments need a way to dissipate photoenergy that they inherently will absorb from the environment. But without a viable energetic channel for decay, such as proton transfer or quenching of singlet oxygen, AQ chromophores with a 1,4-substitutional motif will tend to degrade at a faster pace than their photostable counterparts that lack this motif. Knowledge of the structural trend that drives the photochemical response therefore allows those who care for culturally significant materials to better predict their photosensitivity: If the specific chemical composition of the organic red dyes in an object can be determined and a 1,4-substitutional motif is prevalent (either in the only chromophore present or in high concentration in a mixture of several AQs), then a higher photosensitivity can be presumed than if the structure were lacking, regardless of how many chromophore(s) are present. The data shown here suggest that conservators and curators may therefore use information about the chemical structure to predict general trends in the photosensitivity of the objects in their care if colored with AQ dyes and protect those objects accordingly.

MATERIALS AND METHODS Experimental design

Standards of seven HAQs were purchased from Sigma-Aldrich and used without further purification. Standards were directly applied to individual graphite sample bars as a thin solid layer and examined in isolation to ensure spectral purity for 2C-R2PI and pump-probe measurements.

The instrument has been previously described in detail, and only a brief description of the experimental setup follows (*31, 32*). Samples were laser desorbed in vacuo directly in front of a pulsed molecular beam controlled by a piezo cantilever valve (*33*). The desorption laser was a tightly focused Nd:YAG laser (1064 nm, ~1 mJ/pulse), and the piezo cantilever valve operates at a 45-µs pulse duration with eight bars backing argon gas. The desorbed sample was adiabatically cooled by collisions with the argon jet expansion to between 10 and 20 K, and the molecular beam was skimmed before being intersected by laser beams and photoionized by 2C-R2PI. The subsequent ions were detected by a reflectron time of flight mass spectrometer (analyzer pressure, 2×10^{-6} torr; mass resolution, $m/\Delta m = 500$).

The 2C-R2PI spectroscopic and picosecond pump-probe delay measurements were performed with an Ekspla PL2251 Nd:YAG laser system producing ~30-ps laser pulses. The 355-nm output pumps an Ekspla PG401 tunable optical parametric generator (OPG) (output of 450 to 600 μ J/pulse and spectral linewidth of ~6 cm⁻¹). The sample was excited by the OPG and ionized by 213-nm fifth harmonic of the Ekspla PL2251 laser, which was mechanically delayed up to 600 ps before collineation with the OPG beam. A variable electronic (SRS DG645) delay between OPG UV laser and an excimer laser (193 nm, 1.5 to 2 mJ/pulse) was used for pump-probe measurements in the nanosecond time delay range.

For IR-UV double resonant spectroscopy (i.e., IR hole burning) a Laser Vision optical parametric oscillator/amplifier (mid-IR output over the range 3000 to 3600 cm⁻¹ of ~3 to 5 mJ/pulse and spectral linewidth of 3 cm⁻¹) precedes the R2PI pulse by 200 ns. This study used double resonant spectroscopy with two different pulse sequences: In mode I, the IR pump was scanned at a fixed UV probe wavelength, while in mode II, the UV was scanned with a fixed IR burn wavelength. In mode I, the UV laser wavelength was selected to correspond to a single vibronic transition, and the resulting 2C-R2PI signal depletes when the IR laser becomes resonant with the ground-state population. The resulting ion-dip spectrum therefore represents the ground-state IR spectrum of a single tautomer selected by the UV probe wavelength. This IR spectrum can be compared with calculated IR frequencies to determine the specific tautomer of the selected vibronic transition. In mode II, the IR laser wavelength was selected to correspond to a tautomer-specific vibrational resonance, and spectra were collected both with IR laser on and off. The difference spectrum identifies peaks in the UV spectrum that arise from the same tautomer.

Calculations for IR-UV double resonant spectroscopy

Calculations were performed with the Gaussian 09 program package (34). Starting structures for hydroxy derivatized anthracene-9,10-dione (HAQ) structures, tautomer and rotamer isomers, were optimized using the B3LYP hybrid functional with cc-pTVZ basis set. Relative zero-point-corrected energy (ZPE) values were used to predict the number of isomers in our molecular beam based on the rule of thumb that for a given species, isomers up to 20 kJ/mol of the LE structure were typically kinetically trapped laser desorption jet expansion. Past work on the nucleobase adenine showed that only the LE isomer was present, where the next LE isomer was calculated to be at ~33 kJ/mol higher energy than the one observed (35). Ground -state minima were confirmed by the absence of imaginary frequencies, and these geometries were later used to determine electronic transition state strengths for S_{1-4} by way of time-dependent density functional theory with the B3LYP hybrid functional with cc-pVTZ basis set. The simulated IR spectra arise from frequencies with harmonic intensities using a Lorentzian shape and a full width at half maximum of 3 cm⁻¹ and were presented without a spectral shift.

Calculations for construction of energy profiles

The ground-state minima forms of the 1-HAQ and 1,4-DHAQ were optimized by means of the MP2 method (36) using the ccpVDZ (37) correlation-consistent atomic basis set. The excited state geometries were optimized with the use of the same basis set while using the CC2 (38, 39) method as implemented in TURBOMOLE software package (40). In the calculation of the vertical excitation energies, ΔE^{VE} , mimicking the absorption spectra, performed on top of the MP2/cc-pVDZ-optimized S₀-state geometries, the CC2 (38, 39) method was used to evaluate the response properties.

To elucidate the photophysical mechanism on the molecular level, the important driving coordinates were appropriate for each system so that the ground- and excited-state minimum potential energy (MPE) profiles or surfaces could be constructed to estimate the ground- and excited-state energy barriers determining the photophysics of the both molecules. The 1-HAQ molecule was bound by a single hydrogen bond. In this case, one coordinate, the $R_1(O_1H)$ distance, is needed to be chosen as a driving coordinate to illustrate the photophysical mechanism of the photo-tautomerization process. The MPE profile for 1-HAQ was constructed in a way that for fixed given value of the $R_1(O_1H)$ distance, all the remaining nuclear degrees of freedom were optimized; once in the ground state and twice in the two excited states: $S_1(\pi\pi^*)$ and $S_1(n\pi^*)$.

The 1,4-DHAQ system has an additional intramolecular hydrogen bond binding the molecule, and more tautomeric forms were possible to be formed upon photoexcitation. A convenient method for illustrating the tautomerization process in this molecule was to construct the MPE surface spanning the two driving reaction coordinates describing the two intramolecular hydrogen bonds by the $R_2(O_2H)$ and $R_3(O_4H)$ distances. In that case, both the R_2 and R_3 coordinates were frozen for given values, while the rest of the parameters were optimized in the constructed MPE surface, separately for the ground (S_0) and the two excited states: $S_1(\pi\pi^*)$ and $S_1(n\pi^*)$ with the *Cs* symmetry constrain.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/ content/full/5/9/eaaw5227/DC1

- Fig. S1. Picosecond pump-probe spectra.
- Fig. S2. Nanosecond pump-probe spectra.
- Fig. S3. IR hole burning spectra in mode II with the burn laser set to 3090 cm^{-1} .
- Fig. S4. Hole burning spectra in mode I.

Table S1. Calculated properties of 1-HAQ and 1,4-HAQ.

Table S2. Emission properties of 1-HAQ.

Table S3. Emission properties of 1,4-HAQ.

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Excited state intramolecular proton transfer in hydroxyanthraquinones: Toward predicting fading of organic red colorants in art

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