Improved photodecarboxylation properties in zinc photocages constructed using *m*-nitrophenylacetic acid variants

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The methoxy- and fluoro-derivatives of *meta*-nitrophenylacetic acid (*m*NPA) chromophores undergo photodecarboxylation with comparable quantum yields (Φ) to unsubstituted *m*NPA, but uncage at red-shifted excitation wavelengths. This observation prompted us to investigate DPAdeCageOMe (2-[bis(pyridin-2-ylmethyl)amino]-2-(4-methoxy-3-nitrophenyl)acetic acid) and DPAdeCageF (2-[bis(pyridin-2-ylmethyl)amino]-2-(4-fluoro-3-nitrophenyl)acetic acid) as Zn²+ photocages. DPAdeCageOMe has a high Φ and exhibits other photophysical properties comparable to XDPAdeCage ({bis[(2-pyridyl)methyl]amino}{9-oxo-2-xanthenyl}) acetic acid), the best perforiming Zn²+ photocage reported to date. Since the synthesis of DPAdeCageOMe is more straightforward than XDPACage, the new photocage will be a highly competitive tool for biological applications.

We are developing decarboxylation reactions of metanitrophenylacetatic acid (mNPA) and xanthone (XAN) groups to design photocages that release Zn2+ upon irradiation with light.^{1, 2} Such photocaged complexes block the biological activity of metal ions until the Zn²⁺ release is initiated by exposure to light of a specific wavelength.3 We recently described DPAdeCage (1) and XDPAdeCage (2), two Zn²⁺ photocages that incorporate a dipyridyl amine (DPA) chelating group to selectively bind Zn²⁺ over other common loosely bound metal ions found in cells (Error! Reference source not found.).2 Irradiation at or near the λ_{max} initiates a photoreaction that results in the loss of the carboxylic acid functional group as CO₂, which shifts the Zn²⁺ binding equilibrium toward unbound metal ion (Scheme 1). The transformation from a 4-coordinate to a 3coordinate chelator leads to a 105-fold reduction in binding affinity (ΔK_d). For biological applications, a rapid synthesis to produce photocages in high yields would be ideal. The photophysics of XAN chromophores are superior to those of mNPA groups,4-6 but the low-yielding synthesis of the XAN scaffolding limits the pace of photocage development. To overcome these limitations, we functionalized several $m{\sf NPA}$ chromophores to develop DPAdeCage derivatives with improved photophysical properties. In addition to those improvements, the added functional groups facilitate selective

Fig 1 Structures and important photophysical properties of the *m*NPA-based DPAdeCage (1) and XAN-based XDPAdeCage (2), which are both effective Zn²⁺ photocages. The use of the XAN chromophore red-shifts the uncaging wavelength. The name DPAdeCage derives from dipicolylamine (DPA) decarboxylation (de) photocage(Cage) where X stand for xanthone in XDPAdeCage.

Scheme 1. Photodecarboxylation reaction of and Zn²⁺ release by DPAdeCage.

Designing effective photocages requires avoiding short wavelength excitation (λ_{max}), which can induce cellular photodamage, while still delivering sufficient energy to break chemical bonds. Compared to our mNPA-based DPAdeCage, the λ_{max} of XAN-derived XDPAdeCage is red-shifted by over 80 nm (Error! Reference source not found.). Furthermore, upon coordination to Zn2+ the photoreactivity of DPAdeCage is significantly reduced. The XAN chromophore offers significant photophysical advantages that include maintaining a high Φ upon Zn2+ binding. Accessing the XAN chromophore necessary to make XDPAdeCage requires a 2-step synthesis with a maximum 20% yield. By comparison, the nitration step to synthesize the mNPA chromophore has a much higher yield, increasing the total material obtained ($^{\sim}70\%$ yield of mNPA). The difference in reaction time, 24 h and 1 h respectively, also impacts the rate of photocage synthesis. We hypothesize that

nitration at the 3-position of the aromatic ring simplifying the purification procedure for the *m*NPA chromophore that will facilitate the rapid generation of photocages.

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the relative ease of making mNPA derivatives compared to the XAN chromophore, as well as the numerous readily available starting materials, will facilitate the rapid production of mNPA-based photocages for numerous applications. Furthermore, if functionalization of the mNPA group could shift the λ_{abs} sufficiently, we can prepare and screen Zn^{2+} photocages for biological applications more quickly as well as expand the toolbox of tunable chromophores.

Table 1. Summary of photophysical data of mNPA derivatives.

	R	σ_{p}^{a}	λ _{max} (nm)	ε (M ⁻¹ , cm ⁻¹)	$\Phi_{Photolysis}$
3a	OAc	0.31	317	1478	16.3
3b	Cl	0.23	315	1455	8.53
3c	F	0.06	311	1758	24.0
3d	HNAc	0	332	2123	10.2
3e	Н	0	272	7087 ^b	20.1
3f	OMe	-0.27	347	2589	22.6
3g	ОН	-0.37	427	2989	0.0
3h	NH_2	-0.66	425	3729	0.0
3i	NMe ₂	-0.83	460	5915	0.0

Measurements obtained in 40 mM HEPES buffer (pH 7.5, 100 mM KCI). Quantum yields were obtained in solutions containing 30% methanol to maintain photoproduct solubility. $^{\rm a}$ -Hammet constants indicating the relative electron withdrawing or donating properties of each R group. $^{\rm b}$ -3e lacks a red-shifted absorbance band; ϵ = 3264 M $^{\rm c}$ 1 cm $^{\rm c}$ 1 at λ_{300} , 356 M $^{\rm c}$ 1 cm $^{\rm c}$ 1 at λ_{350} .

In previous studies on ortho-nitrobenzyl photocages (oNB), introducing electron donating groups (EDGs) or electron withdrawing groups (EWG) produced photocages with different rates of photolysis, quantum yields and excitation wavelengths.⁷⁻¹¹ Tsien, Kaplan, and others reported functionalization of the oNB ring with methoxy groups shifted the λ_{max} to longer wavelengths, $^{7,\,8}$ suggesting we might benefit from a similar approach. Modification of the nitrobenzyl ring, however, can also impact the electronics of the aromatic ring and therefore the nature of the photoreaction. 10, 12 Unlike oNBs that undergo a Norrish type II photoreaction, 13, 14 the mechanism for photodecarboxylation of mNPA chromophores involves an electron transfer from the nitro group to the methylene bridge of the phenylacetate to initiate the photodecarboxylation. 15-17 This suggests increasing the electron-density around the aromatic ring would impede electron-transfer to the methylene bridge, which could reduce photolysis quantum yields (Φ). Thus, we predicted EDGs would exhibit reduced photoactivity compared to mNPAs functionalized with EWGs. By adjusting the electronic-structure of the mNPA chromophore we hoped to access a DPAdeCagederivative that will maintain a high Φ upon coordination to $\mathrm{Zn^{2+}}.$

A series of mNPA chromophores were prepared with various EDGs and EWGs at the 4-position of the aromatic ring (Error! Reference source not found.). We hypothesized that functional groups with heteroatoms would extend the conjugated π -system and red-shift the λ_{max} . Functionalization resulted in the

appearance of a new red-shifted absorbance feature that was not observed in the parent mNPA **3e** (Figure 2). The red-shifted feature appears as a shoulder of the strong UV band centred at circa 270 nm in most of the compounds, but the band shifts sufficiently in the methoxy- (**3f**) and hydroxy-derivatives (**3g**) to appear as a distinct new absorbance feature. While all the derivatives possess a weaker extinction coefficient (ϵ) of the large absorbance band below 300 nm compared to **3e**, the compounds also absorb more strongly at wavelengths greater than 320 nm (**Error! Reference source not found.**).

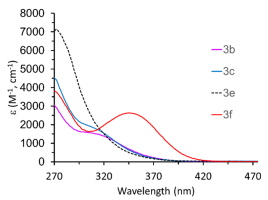


Fig. 2. Absorption spectra of 3b (-Cl), 3c (-F), 3e (-H) and 3f (-OMe) in aqueous solvent and simulated physiological conditions (40 mM HEPES buffer pH 7.5 100 mM KCl)

The photolysis quantum yields (Φ) were determined under simulated physiological conditions using LC/MS to monitor the loss of **3a-3i**. Despite the red-shifted band, the photoactivity of the acetyl (**3a**), chloro (**3b**), and amide-functionalized (**3d**) compounds is diminished relative to **3e**. The only EWG-containing derivative that exhibits a Φ similar to **3e** is the fluoroderivative **3c**. Although **3b** has a similar chloro group, the observed photoreactivity is diminished relative to **3c**. Notably the absorptivity of **3c** is stronger than **3b** across the entire spectrum and likely contributes to the observed difference in Φ (Error! Reference source not found.). While **3c** exhibits photoactivity comparable to XDPAdeCage, the λ_{max} is blueshifted by over 30 nm.

The mass peaks observed in the LC/MS spectrum of 3a-f after photolysis indicate tolyl and aldehyde derivatives analogous to those previously observed with mNPA-based photocages predominate the photoproducts (Scheme 1).1, 2 Compounds containing EDGs (3f-3i) exhibit a bathochromic shift in λ_{max} compared to the parent compound **3e**; however only **3f** retained photodecarboxylation activity. The hydroxy compound 3g exhibits a higher ϵ (2989 M⁻¹, cm⁻¹) and a more red-shifted absorbance (λ_{max} = 427 nm) but lacks photoactivity compared to the corresponding methoxy derivative (3f). Excited state proton transfer does not appear to be responsible for the inactivity of 3g as the compound remains photochemically inert under strongly basic conditions (pH 14). Although 3f contains an electron-donating methoxy group, the photodecarboxylation Φ and red-shifted λ_{max} is the most comparable to XDPAdeCage. While the methoxy functional group is electron-donating by resonance, inductive effects provide the most plausible explanation for the anomalous behaviour in the EDG series.

The model methoxy- and fluoro-compounds 3f and 3c exhibited the most comparable photophysical properties to XDPAdeCage, therefore we chose these chromophores to build new DPAdeCage derivatives (Error! Reference source not found.). Our initial attempts to prepare these photocages via our established route were unsuccessful due to oxidative decomposition during the final nitration reaction. Alternatively, starting with 3f and 3c provided 4 (DPAdeCageOMe) and 5 (DPAdeCageF) in 11.6% (5 steps) and 11.8% (4 steps) respectively. The only significant difference between the two synthetic pathways is the final ester hydrolysis. DPAdeCageF required acidic conditions at elevated temperature, while the basic conditions used to prepare DPAdeCage successfully provided DPAdeCageOMe. A combination of product decomposition in the harsher deprotection reaction as well as the more difficult isolation of DPAdeCageF from acidic solution, reduced the overall yield of the final product. A comparison of the synthetic pathways required to develop the new mNPA photocages and XDPAdeCage reveals both DPAdeCageOMe and DPAdeCageF can be more rapidly prepared and in greater amounts than XDPAdeCage.

$$R = \text{OMe, } \textbf{4} \text{ (DPAdeCageOMe)} \qquad R = \text{F, 5 (DPAdeCageF)} \\ \lambda_{\text{max}} \text{ (apo/Zn}^{2+}) = 342 \text{ nm/337 nm} \qquad \lambda_{\text{abs}} \text{ (apo/Zn}^{2+}) = 304 \text{ nm / 298 nm} \\ K_{\text{d}} \text{ Zn}^{2+} = 6.1 \text{ pM} \qquad \qquad K_{\text{d}} \text{ Zn}^{2+} = 7.8 \text{ pM} \\ \Delta K_{\text{d}} = 2.6 \times 10^4 \qquad \qquad \Delta K_{\text{d}} = 2.0 \times 10^4 \\ \Phi \text{ (apo/Zn}^{2+}) = 24.5/23.3 \qquad \Phi \text{ (apo/Zn}^{2+}) = 22.0/<1 \\ \end{pmatrix}$$

 $\begin{tabular}{ll} \textbf{Fig. 3.} Structures and important photophysical properties of DPAdeCageOMe (4) and DPAdeCageF (5). \end{tabular}$

Zinc-binding constants for DPAdeCageOMe and DPAdeCageF were determined by competitive titration with a high affinity zinc ligand compound 4-(p-pyridylazo)resorcinol (PAR). Under simulated physiological conditions (40 mM HEPES, 100 mM KCl, pH 7.5), DPAdeCageOMe (K_d 6.1 pM, Δ K_d 2.6 × 10⁴) and DPAdeCageF (K_d 7.8 pM, Δ K_d 2.0 × 10⁴) bind Zn²+ with similar affinity to DPAdeCage (K_d 0.8 pM). The slight difference in K_d has a minimal impact on the Δ K_d but both compounds should efficiently release Zn²+ following irradiation due to changes in the binding equilibrium. The minor difference observed in the K_d values is likely due to small inductive effects, caused by the addition of functional groups on the mNPA ring.

Table 1: Photophysical data for apo- and Zn²⁺-complexes of DPAdeCage, XDPAdeCage, DPAdeCageOMe and DPAdeCageF

	λ_{max} (nm)	ε (M ⁻¹ , cm ⁻¹)	$\Phi_{\text{Photolysis}}$
DPAdeCage	261	5120	40.0
[Zn(DPAdeCage)] ⁺	262	5230	5.0
XDPAdeCage	347	4730	20.0
[Zn(XDPAdeCage)] ⁺	347	5270	27.0
DPAdeCageOMe	262/342	9680/1686	24.5
[Zn(DPAdeCageOMe)] ⁺	262/337	6670/1737	23.3
DPAdeCageF	261/304	10871/1804	22.0
[Zn(DPAdeCageF)] ⁺	261/298	10983/1808	<1

Measurements obtained in 40 mM HEPES buffer (pH 7.5, 100 mM KCI). Quantum yields were obtained in solutions containing 30% methanol to maintain photoproduct solubility.

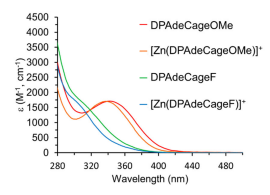


Fig. 4. Absorbance spectrum of DPAdeCageOMe/[Zn(DPAdeCageOMe)]* and DPAdeCageF/[Zn(DPAdeCageF)]* in aqueous solvent under simulated buffered conditions (40 mM HEPES buffer pH 7.5 and 100 mM KCI). Notably, DPAdeCageOMe has a well-defined, red-shifted absorbance band at 347 nm that is comparable to XDPAdeCage.

The photophysical properties of DPAdeCageOMe and DPAdeCageF are nearly identical to the parent compounds 3f and 3c under aqueous conditions (Error! Reference source not found.). Notably, the λ_{max} of both DPAdeCageOMe and DPAdeCageF are red-shifted relative to DPAdeCage. The methoxy-functionalized DPAdeCageOMe (342 nm) has a λ_{max} nearly identical to XDPAdeCage (347 nm). The addition of Zn²⁺ to the photocages results in a slight hypsochromic shift of λ_{max} , but relatively little change in absorptivity (Table 1). An analysis of the LC/MS spectrum of DPAdeCageOMe and DPAdeCageF following irradiation, indicate successful photodecarboxylation following irradiation. In the absence of Zn²⁺, both compounds appear to successfully release the carboxylate group based on the change in mass. In DPAdeCage, 2 a decrease in Φ is observed upon coordination to Zn2+ and DPAdeCageF exhibits an even more dramatic decrease upon Zn2+ coordination. Only extended irradiation times (>30 min) of [Zn(DPAdeCageF)]+ resulted in evidence of photolysis, which severely limits the potential applications of the photocage. The addition of Zn2+ does not appear to affect the resulting photoreaction DPAdeCageOMe; however, DPAdeCageF forms a wider slate of photoproducts in the presence of Zn²⁺ after long irradiation. Some higher mass, emissive photoproducts suggest that Zn2+ facilitates the formation of coupled DPAdeCageF products

TD-DFT calculations were performed to probe if the electronic structure contributed to the observed difference in photoreaction of [Zn(DPAdeCageOMe)]+ [Zn(DPAdeCageF)]*. The structure of DPAdeCageOMe, DPAdeCageF, [Zn(DPAdeCageOMe)]⁺ and [Zn(DPAdeCageF)]⁺ were optimized using DFT and the frontier molecular orbitals contributing to the electronic transitions found in the TD-DFT calculations were visualized. In all cases, the lowest energy excitation involves excitation from the HOMO to the LUMO. The LUMOs are localized around the nitro group and phenyl ring for all compounds, consistent with previous studies of mNPA chromophores (Error! Reference source not found.).17 We believe that promoting an electron into a LUMO localized on the nitro group is necessary for photolysis to occur, suggesting [Zn(DPAdeCageF)]⁺ should be capable of undergoing photodecarboxylation based on the calculated electronic structure. A closer examination of the observed oscillator strengths (f) representing the molar absorptivity of $Zn(DPAdeCageOMe)^+$ (f = 0.04) and $Zn(DPAdeCageF)^+$ (f = 0.01) reveal a nearly 4-fold decrease in the oscillator strength of the lowest energy transition for Zn(DPAdeCageF)+,which may explain a decrease in observed photolysis, although other factors probably contribute to the difference given the large drop in Φ observed experimentally.

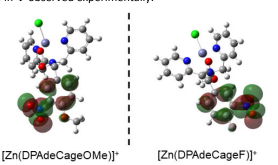


Fig. 5. Optimized geometry depicting the LUMO of [Zn(DPAdeCageOMe)]* and [Zn(DPAdeCageF)]* determined by DFT calculations (B3LYP/6-311+G(d)).

Through a study of model *m*NPA chromophores, we were able to identify derivatives with comparable photophysical properties compared to the decarboxylation reaction observed with XAN groups. Specifically, fluoro- and methoxy- groups introduce a distint red-shifted absorption band; however, only the methoxy derivative retains sufficient photodecarboxylation activity when integrated into a Zn²+ photocage. We were pleased to see DPAdeCageOMe maintains a high Φ when coordinated to Zn²+ and has nearly identical photocaging properties to XDPAdeCage. A methoxy group in the 4-position also provides a potential site for modification of future photocages through an ether linkage that should not impact Zn²+ binding significantly and will be the subject of future investigations.

Conflicts of interest

There are no conflicts to declare.

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