A Photoactivatable Formaldehyde Donor with Fluorescence Monitoring Reveals Threshold To Arrest Cell Migration

Lukas P. Smaga,† Nicholas W. Pino,† Gabriela E. Ibarra,† Vishnu Krishnamurthy,‡ and Jefferson Chan*†‡

†Department of Chemistry and Beckman Institute for Advanced Science and Technology, University of Illinois at Urbana—Champaign, Urbana, Illinois 61801, United States
‡Department of Biochemistry, University of Illinois at Urbana—Champaign, Urbana, Illinois 61801, United States

Supporting Information

ABSTRACT: Controlled light-mediated delivery of biological analytes can enable the investigation of highly reactive molecules within living systems. As many biological effects are concentration dependent, it is critical to determine the location, time, and quantity of analyte donation. In this work, we have developed the first photoactivatable donor for formaldehyde (FA). Our optimized photoactivatable donor, photoFAD-3, is equipped with a fluorescence readout that enables monitoring of FA release with a concomitant 139-fold fluorescence enhancement. Tuning of photostability and cellular retention enabled quantification of intracellular FA release through cell lysate calibration. Application of photoFAD-3 uncovered the concentration range necessary for arresting wound healing in live cells. This marks the first report where a photoactivatable donor for any analyte has been used to quantify intracellular release.

Chemical tools that enable controlled perturbation of complex biological networks represent a powerful approach for the mechanistic evaluation of biological signaling. For instance, enzyme inhibitors can disrupt metabolic pathways and alter the cellular levels of an associated analyte. Although direct supplementation of the cell culture media can yield the desired effect, this strategy precludes analytes that exhibit high reactivity and short biological lifetimes. To address these limitations, a diverse palette of donors based on spontaneous, enzymatic, and light-mediated activation have been designed to mask reactive analytes until release. The latter class, photoactivatable donors, provides the greatest control because the intensity of light, duration of irradiation, and focal point can be adjusted. Photoactivatable donors for reactive oxygen, nitro, sulfur, and carbonyl species, as well as metal ions, have been reported. Recent developments have led to photoactivatable donors equipped with the ability to monitor analyte release through fluorescence or photoacoustic readouts. Despite these advances we have recognized that there are two major voids in this area. First, there are no photoactivatable donors (with or without a readout) for biological aldehydes, such as formaldehyde (FA). Second, no studies have leveraged the unique monitoring capabilities of donors with internal readout to quantify analyte release within living systems.

Recently, mounting evidence has surfaced that suggests that FA is not merely a byproduct of cellular metabolism. On the contrary, FA was hypothesized to be a signaling molecule. and has been shown to be involved in the synthesis of purines via the one-carbon cycle. At elevated levels (in the mM range) FA is associated with aging and pathologies ranging from cancer to Alzheimer’s disease. The cellular concentration that marks the transition from physiological to pathological roles of FA is currently unknown. In this work, we have developed a series of photoactivatable FA Donors (photoFADs) that are equipped with the ability to report on FA release via fluorescence enhancement. We selected the most suitable donor based on photocaging kinetics, turn-on response, biocompatibility, subcellular localization, and cellular retention to evaluate the FA concentration range required to arrest wound healing in HEK293 cells.

Each of the photoFADs features a photolabile nitrobenzyl group connected to a silicon-xanthene dye through an acetal linkage (Figure 1a). Alkylated silicon-based fluorophores are nonfluorescent due to the perturbed electronic push–pull system, as well as donor-photoinduced electron transfer (d-PeT) quenching from the proximal nitro group. Irradiation with light induces cleavage of the photocage to afford an unstable hemiacetal intermediate that we hypothesize will

Figure 1. (a) Chemical structures of the photoFAD series. (b) Mechanism of light-mediated FA release.

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spontaneously collapse to generate FA and the corresponding dye (Figure 1b). Our first-generation donor, photoFAD-1, displayed a 40-fold fluorescence signal enhancement following activation; however, photobleaching of the dye was observed, which would preclude reliable in cellulo quantification (Figures S1–S2). We reasoned that replacing the 2-methyl substituent on the pendent aryl ring with a trifluoromethyl moiety would lead to higher quantum yields and less photobleaching.28 The resultant donor, photoFAD-2, exhibited improved photostability, as well as greater fluorescence turn-on (205-fold) (Figure S3). Unfortunately, when photoFAD-2 was evaluated in HEK293 cells, we observed that the dye was not retained relative to the unactivated donor (Figure S4a). This was addressed in photoFAD-3 (Scheme 1) by installing electron-withdrawing chloro substituents on the silicon xanthene core to decrease the pK_a of the phenol from 6.9 to 5.6 (Figure S5).30 This modification resulted in greater cellular retention because the dye is predominantly in an anionic state under physiological conditions, which reduces passive diffusion through the cell membrane.

The synthesis of photoFAD-3 began with the chlorination of silicon xanthone 1 using N-chlorosuccinimide to yield 2. The free hydroxyl groups were protected as silyl ethers using tert-butyldimethylsilyl chloride to afford 3 in 33% over two steps. 1-Bromo-2-(trifluoromethyl)benzene was treated sequentially with tert-butyllithium and 3 to yield the dye 4 after acidic workup. Compound 4 was then alkylated with 5-[1-(chloro-fluoromethyl)-6-nitro-1,3-benzodioxole (CNDB) to afford photoFAD-3 in 45% yield.

Scheme 1. Synthesis of photoFAD-3

In vitro characterization of photoFAD-3 revealed that it is nonfluorescent in its unactivated form (Figure 2a). The quantum yield of 4 on the other hand was determined to be 60% (Table S1). This corresponds to a 139-fold fluorescence turn-on response in vitro (k_on = 137 nM/min) (Figure 2b and 2c). Irradiation with light over 30 min did not cause significant photobleaching (Figure S2). Of note, the total exposure to light does not exceed 3 min in our experiments. When activated a 10.4-fold fluorescence enhancement was observed in HEK293 cells (Figure S6). Moreover, we demonstrate spatial control by activating photoFAD-3 in a small area within a larger field of cells (Figure S7).

Owing to the novel combination of the photolabile nitrobenzyl group, acetox linkage, and fluorescent dye it was essential to verify that FA was being released upon photoysis. To address this, we performed mass spectrometry analysis on an activated solution of photoFAD-3. The experimental results confirmed release of 4 (m/z = 467.3) (Figure S8). Furthermore, we confirmed the formation of FA using an isotopologue of photoFAD-3 (13C-photoFAD-3). Activation of 13C-photoFAD-3 in deuterated methanol resulted in a 13C NMR signal at 91.10 ppm, which corresponds to the resonance of the hemiacetal species in methanol (Figure S9).

Prior to application in living cells, we evaluated its chemo- and photostability. Neither prolonged incubation with a serum-containing medium (Figure S10a) nor exposure to ambient light (Figure S10b) resulted in significant activation. Additionally, photoFAD-3 was stable against a panel of thiols and reactive oxygen species (Figure S11a), as well as across a range of pH values (Figure S11b). We were delighted to observe that photoFAD-3 and 4 did not induce any notable cytotoxicity in HEK293 cells (Figure 2d) and that both compounds are localized in the cytoplasm (Figure S12).

We next turned our attention to establishing a protocol to quantify FA release from photoFAD-3 in living systems. Specifically, photoFAD-3-stained HEK293 cells were subjected to different irradiation times and imaged using fluorescence microscopy (Figure 3a). The cells were then lysed, and the signal was quantified using fluorescence spectroscopy (Figure S14a). Compound 4 was used to establish a calibration curve to correlate the intensity of the cell lysates to a known concentration of 4 (Figure S14b). By determining the number of cells within each sample, we were able to calculate the FA concentration released per cell at different irradiation times (eq S1). For instance, 6.20 attomol (10^-18) of FA were released in a single cell upon 3 min of photoactivation. We determined the average diameter of HEK293 cells in suspension to be ~14.5 μm, which corresponds to a cell volume of ~1.60 pL (10^-12). Using this information, we determined that 4.0 μM FA was released inside each cell after 180 s of irradiation, assuming that the resultant FA will persist throughout activation. It is important to note that our quantification procedure is not limited to fluorescence microscopy. For example, we were able to perform experiments with an IVIS fluorescence imager with a larger imaging window (e.g., the entire well of a 6-well plate) (Figure 3b). Results from these experiments were consistent with those obtained using a fluorescence microscope (3.9 μM increase of FA concentration upon 3 min of photoactivation). Together, these experiments highlight the utility of photoFAD-3 for consistent results across multiple imaging platforms.
Finally, we applied photoFAD-3 to determine the threshold required to exceed the cell’s ability to detoxify FA. For these studies we synthesized a control reagent (Ctrl-photoFAD-3) consisting of the dye connected directly to the nitrobenzyl photocage. Because Ctrl-photoFAD-3 does not feature the acetoxymethyl (AM) ether (Figure S15b). Incubation of HEK293 cells with Ctrl-photoFAD-3 or photoFAD-3 and irradiated (0–180 s). These cells were subsequently used for a wound healing assay which reports on cell migration and proliferation. The rate of wound closure was monitored over the course of 24 h (Figure 4a). We did not observe any effect on wound healing when Ctrl-photoFAD-3 was activated (Figure S17). In contrast, activation of photoFAD-3 resulted in an irradiation time-dependent decrease of wound healing (Figure 4b). Specifically, FA concentrations below 2.2 μM per cell did not significantly affect recovery, whereas higher concentrations resulted in a decreased propensity to close the wound. At the longest irradiation time, cells lost their ability to adhere to the plate (Figure 4c). Moreover, dose-dependent activation of Ctrl-photoFAD-3 strongly suggests that the nitrosoacetophenone byproduct and the released dye do not have an apparent impact on wound healing (Figure S18).

Inspired by these results, we explored whether common prodrug and probe delivery strategies that release FA through esterase-mediated cleavage of masking groups (e.g., acetoxymethyl (AM) ether) would yield similar results. This led to the design of AM-FAD-3 that consists of the dye capped with an AM ether group (Figure S15b). Incubation of HEK293 cells with AM-FAD-3 under standard conditions did in fact impair wound healing to the same extent (Figure S19). This observation indicates that FA-releasing masking groups could aid drug efficacy through a secondary effect beyond simple increase of bioavailability, if FA levels overwhelm the intracellular detoxification mechanisms.

In conclusion, we have developed the first photoactivatable FA donor that enables perturbation of intracellular FA levels with high spatiotemporal control. This is the first report where a photoactivatable donor equipped with a reporter has been used to quantify in cellulo release. Prior to this work, calibration was only performed in vitro which does not indicate how much of an analyte is being delivered into cells. Our quantification approach is readily compatible with various instruments. We were able to reveal that an increase of 2.2 μM FA is sufficient to impair wound healing for HEK293 cells. Additionally, we were able to show that intracellular FA buildup through cleavage of prodrug masking groups can reach this concentration range, which could enhance observable drug efficacy. Our future work will focus on the use of photoFAD-3 to study the role of FA in stem cell aging. In addition to providing a tool to study FA biology, we envision this work will set a new standard of using the readout of photoactivatable donors to determine the concentration of cellular release.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.9b11899.

Details regarding the materials, instrumentation, data processing, synthetic procedures as well as NMR spectra, supporting figures, tables, and equations PDF

**AUTHOR INFORMATION**

**Corresponding Author**

*jeffchan@illinois.edu

**ORCID**

¥ Lukas P. Smaga: 0000-0001-8659-3326
Nicholas W. Pino: 0000-0002-7887-9399
Gabriela E. Ibarra: 0000-0002-6591-8651
Jefferson Chan: 0000-0003-4139-4379

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Notes
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REFERENCES


