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# BODIPY phototether enables oligonucleotide cyclization and subsequent deprotection by tissue-transparent red light<sup>†</sup>

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meso-methyl-BODIPY photocages release a leaving group upon visible light irradiation but often lack thermal stability. In turn, our thermally-stable, red-shifted BODIPY phototether allows oligonucleotide cyclization, preventing complementary strand hybridization. Hybridization resumes upon red-NIR irradiation, disconnecting the phototether by oxidative cleavage, which is easily monitored by a blue shift in fluorescence.

Oligonucleotides are short nucleic acids that hybridize to a complementary sequence, with various biological functions.<sup>1</sup> Access to oligonucleotides by solid phase synthesis has prompted the development of many biological and medical diagnostic technologies.<sup>2</sup> Moreover, oligonucleotides can be designed to target specific genes and are, as such, promising candidates for targeted therapy.<sup>3,4</sup> Native oligonucleotides, however, show low stability and are degraded by nucleases limiting their applications. This problem can be overcome by protecting oligonucleotides against decomposition, including through cyclization.

Cyclization of oligonucleotides blocks their ability to hybridize with a complementary strand and, therefore, masks their function. In fact, transient cyclization with subsequent release is an efficient strategy of gene regulation.<sup>5</sup> This process can be controlled using a photoactivatable tether (phototether)<sup>6</sup> to temporarily connect two segments of oligonucleotides and subsequently release them by light in a defined time, space, and concentration.<sup>5</sup>

Despite their potential, photocage applications as phototethers are still incipient because phototethers must absorb in the tissue-transparent window and have low toxicity and defined photoreactivity while remaining thermally stable under cyclization conditions. Only a few photocages, such as cyanine<sup>7,8</sup> and anthracene,<sup>9</sup> have been introduced so far, though with limited stability and absorption properties.<sup>10</sup> To meet these requirements, BODIPY photocages stand out as promising alternatives.

BODIPY photocages can be readily synthesized and derivatized,<sup>11</sup> displaying multimodal photoreactivity.<sup>12</sup> Furthermore, their spectral properties change upon irradiation, which enables photorelease monitoring.<sup>13</sup> However, the biological use of the most common *meso*-methyl substituted BODIPY photocages is limited by the low thermal stability of their C-heteroatom bond, suffering from hydrolysis and reductive homolytic cleavage.<sup>13,14</sup> Therefore, we hypothesized that a more stable, *meso*-phenyl  $\pi$ -extended BODIPY phototether could be designed to be cleaved by the recently described photooxidative styryl cleavage induced by singlet oxygen.<sup>13,15,16</sup>

In this study, we aimed at developing a BODIPY phototether used for oligonucleotide cyclization and subsequent cleavage by red light. As a result of photoxidizing one of the styryl moieties, irradiation shortened the conjugated  $\pi$ -system, resulting in a blue shift in absorption and emission monitored by multichannel fluorescence.<sup>13</sup> Photocleavage of cyclized oligonucleotides should not affect the biological environment because the singlet oxygen produced by the excited state fluorophore is consumed in this process. A 15-mer DNA oligonucleotide was cyclized and subsequently photoreleased, showing marked changes in hybridization with a complementary strand in melting point measurements, thereby demonstrating the potential applicability of this concept.

We synthesized phototether **1** (Scheme 1) by a Knoevenagel condensation of an azide-containing aromatic aldehyde and *meso*-phenyl-substituted BODIPY. The synthesis started from 4-(2-hydroxyethoxy)benzaldehyde **2**, which was treated with  $PPh_3I_2$  and imidazole to obtain the iodinated analogue **3**. Then, iodine was substituted with azide yielding **4**. BODIPY scaffold **5** 

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was prepared by condensation of 2,4-dimethylpyrrole with benzoyl chloride and further complexation with  $BF_3 \cdot OEt_2$ . Knoevenagel condensation of 5 with an excess of 4 provided phototether 1.

Subsequently, we synthesized a model 15-mer DNA oligonucleotide: 5' m1-GCA TAA ATA AAG GTG-m2 3' (A), modified with propargyl groups at the 3' and 5' ends (m = alkynemodification; for details, see Supporting Information (SI)). Oligonucleotide A was further cyclized by BODIPY phototether 1 to 1A via a copper-catalyzed azide-alkyne cycloaddition click reaction in a DMSO-water mixture (Fig. 1A). To avoid head-tail dimerization or polymerization, the oligonucleotide sequence was decorated with alkyne groups attached to both ends, whereas 1 was symmetrically decorated with two azido groups allowing cyclization via triazole formation. The click reaction was conducted under high dilution conditions ( $c \sim 0.7$  mM), favouring the intermolecular cyclization between compound 1 and oligonucleotide A over intermolecular polymerization. Both 1 and 1A were thermally stable allowing for chromatographic purification.

Due to its limited aqueous solubility, photophysical and photochemical data for **1** were measured in a MeOH:DCM mixture (9:1,  $\nu/\nu$ ). The BODIPY phototether proved to be a strong chromophore, with excellent absorption in the red region ( $\lambda_{abs,max} = 636$  nm) and fluorescent properties ( $\Phi_f =$ 0.16, Table 1). Although **1** aggregated in aqueous media (see SI for more details), it did not require any structural modification by water-solubilizing groups for oligonucleotide cyclization. Attaching **1** to **A** produced water-soluble, cyclized oligonucleotide **1A**, whose spectroscopic properties in water were comparable to those of **1**. The emission maximum of **1A** was 662 nm and hence red-shifted by 10 nm in relation to **1**.

We designed our BODIPY phototether based on photooxygenation cleavage of styryl-substituted BODIPY.<sup>13,15,16</sup> As in cyanine photocages, the electron-rich double bonds of 1 reacted with singlet oxygen produced by the excited dye through sensitization. This oxidative process yielded a dioxetane intermediate 1-diox, which subsequently decomposed to two aromatic aldehydes, 1-irr and a benzaldehyde derivative (Fig. 1B). We found 1-diox and its oxidation products to accumulate within the irradiation time frame (Fig. S14, ESI<sup>†</sup>), which limits achieving full conversion from 1 to 1-irr. However, in 1A photolysis, the preference for the formation of 1A-irr is driven by entropy, which promotes the linearization of the tethered cycle, facilitating thermal decomposition of 1A-diox (dioxetane analogue of cyclic oligonucleotide 1A) to 1A-irr, thereby cleaving the cyclized oligonucleotide (Fig. 1C). Because the molar masses of 1A-diox and 1A-irr are identical, the photocleavage of 1A cannot be studied by mass spectrometry. Therefore, the oligonucleotide recovery was validated through spectroscopy and melting point measurements (vide infra).

Irradiation of **1** was followed by UV/vis and fluorescence spectroscopy (Fig. 2A). The absorption and emission maxima of the photoproduct were significantly hypsochromically shifted (572 nm and 588 nm respectively). Accordingly, oligonucleotide



Fig. 1 (A) Oligonucleotide cyclization by a BODIPY phototether *via* a Cu<sup>1</sup>-catalysed coupling reaction. (B) Uncaging mechanism of BODIPY phototether **1**. (C) Uncaging of cyclized oligonucleotide **1A**. (D) Left: change in the color of **1** upon irradiation; right: change in the emission of **1** upon irradiation (excited by UV light).

 Table 1
 Spectroscopic and photochemical properties of the studied compounds

Compd.	$\lambda_{abs}{}^a$	$\varepsilon_{\max}^{\ \ b}$	$\lambda_{\rm em}{}^c$	$\Delta  ilde{ u}^d$	${\Phi_{ m r}}^e  imes 10^{-6}$	$arPsi_{ m r}\left({ m O}_2 ight)^{e} imes10^{-6}$	$\Phi_{r} \bullet \epsilon_{max}$	$\Phi_{ m f}$
$1^f$	636	138 000	652	386	$0.32\pm0.1$	n.d.	0.044	$0.16\pm0.01$
$1A^g$	648	125 000	662	326	$1.2 \pm 0.1$	$1.5\pm0.1$	0.15	n.d. <sup><i>h</i></sup>
1A irr <sup>g</sup>	572	n.d. <sup>h</sup>	588	475	n.r. <sup>i</sup>	n.r. <sup>i</sup>	n.r <sup>i</sup>	n.d. <sup>h</sup>

<sup>a</sup> Absorption maximum in nm. <sup>b</sup> Molar absorption coefficient at the absorption maximum in M<sup>-1</sup> cm<sup>-1</sup>. <sup>c</sup> Fluorescence maximum in nm. <sup>d</sup> Stokes shift in cm<sup>-1</sup>. <sup>*e*</sup> Photorelease quantum yield,  $\Phi_r$  aerated and  $\Phi_r(O_2)$  saturated with  $O_2$ . <sup>*f*</sup> Measured in the mixture of dichloromethane/methanol (1:9, v/v). <sup>*g*</sup> Measured in PBS. <sup>*h*</sup> n.d. = not determined. <sup>*i*</sup> n.r. = not relevant.



Fig. 2 (A) Normalized absorption and emission spectra of 1A and 1A-irr in PBS, and (B) photorelease quantum yields upon irradiation with 617 nm liaht

1A photocleavage could be easily monitored by a change in the absorption and emission (Fig. 1D).

The cyclized oligonucleotide 1A underwent photorelease in PBS with the quantum yield of  $1.2 \times 10^{-6}$ . Irradiating oxygensaturated solutions of 1A increased the photoreaction quantum yield by 1.2-fold, showing the oxygen effect. The quantum yield of 1 was ~4-fold lower than that for 1A (Fig. 2B), most likely due to the solvent effect (different polarity, lower oxygen solubility in methanol<sup>17</sup> ( $c = 4.15 \ 10^{-4} \ M$ ) than in water<sup>18</sup> (c = $1.22 \ 10^{-3} \text{ M}$ ) and to entropy effects of cyclization. The excellent absorption properties of both 1 and 1A compensate for the low photorelease quantum yields, resulting in acceptable uncaging cross-sections of 0.044 and 0.15, respectively (Table 1).

The photocleavage of the cyclized oligonucleotide 1A was followed by HPLC (Fig. 3A) until quantitative deprotection.

The starting material was consumed and resulted in the formation of 1A-irr as a main product. To demonstrate the ability to control the hybridization of A by light, melting points were measured after the addition of an equimolar amount of a complementary strand. While the melting points of A and 1Airr were almost identical (51.6 and 52.8 °C, respectively, Fig. 3B and C), the melting point of cyclized 1A was not detected due to the absence of duplex formation (Fig. 3D). This finding proves that the cyclization of oligonucleotide blocked its ability to hybridize to a complementary strand and the subsequent photorelease restored this property.

We demonstrated that the BODIPY phototether enables temporary oligonucleotide cyclization and subsequent photorelease by red-NIR light. Thermally stable and easily accessible, the BODIPY phototether is activated by light in the tissuetransparent window, and its distribution and release are easily



Fig. 3 (A) Zoomed-in LC-MS chromatograms extracted at 254 nm: red starting material 1A. Yellow: product 1A-irr. Melting points (5-7 consecutive measurements) of (B) linear 15-mer oligonucleotide A, (C) uncaged oligonucleotide 1A-irr and (D) cyclized oligonucleotide 1.

monitored by fluorescence spectroscopy. A 15-mer DNA oligonucleotide can be temporarily cyclized, blocking its hybridization ability. This function is resumed upon irradiation with red-NIR light that cleaves the phototether with acceptable uncaging cross-sections. Photocleavage is oxygen-dependent suggesting that oxygen plays a key role in this process.

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### Conflicts of interest

There are no conflicts to declare.

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