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A two-photon activatable amino acid linker for the induction of fluorescence

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A new one- and two-photon activatable fluorophore based on ATTO565 was developed using a photolabile linker that simultaneously acts as quencher. It is especially interesting for protein and peptide applications because it can be incorporated by standard peptide chemistry. The application of the new fluorogenic construct in super-resolution microscopy of antibody conjugates is shown.

New fluorescence imaging probes are of high interest for bioimaging at the subcellular level. For the control of fluorescence within biological or artificial systems in time and space many different photoactivatable fluorophores were developed in the last years.¹ These caged fluorophores are weakly or non-fluorescent molecules. Irradiation with the appropriate wavelength releases the fluorescent dye. The use of light as external trigger has the great advantage of being a non-invasive manner to control the time and area of the fluorescence if the wavelength is carefully chosen.

Photoactivatable fluorophores are also the key for single-molecule super-resolution techniques.² In these techniques, the resolution limit of light microscopy is bypassed by a temporally confined detection of single fluorophores, the determination of their position and the reconstruction of an image from single-molecule coordinates.^{3,4} Organic fluorophores exhibit a high photon yield and typically show a better performance than fluorescent proteins, however, photoswitching often requires adding redox chemicals to the image buffer which are cytotoxic. In order to achieve the best results in single-molecule super-resolution imaging, it is thus desirable to develop caged organic fluorophores which exhibit a high contrast ratio between the fluorescent and the quenched

state.

There are different strategies for the development of caged organic fluorophores. A simple but very effective strategy is to attach a photolabile group to the fluorophore at a position that interrup the conjugated π -system and suppresses fluorescence. Since the first reports in this field (using fluorescein derivatives),⁵ the strategr was also applied to other chromophore classes like rhodamines,⁵ acridinone,⁷ rosamine,⁸ coumarins,⁹ carborhodamines¹⁰ and borodipyrromethane (BODIPY).¹¹ Also the efficiency and the properties of the photolabile groups were enhanced in different studies.¹² . 1 recent publications the use of 2-diazoketone as very small caging group for fluorophores was presented. Irradiation of the emolecules releases nitrogen and causes a rearrangement reaction resulting in the recovery of fluorescence. This strategy was applied to a variety of different fluorophores and their application in super-resolution microscopy was shown.¹³

Other approaches that generate light activatable fluorophores without the use of caging groups trigger for instance photoinitiated oxidation of a ZnSalen from a quenched thioether a fluorescent sulfoxide.¹⁴ Another example of a quenched cyanine dye used the photocleavage of a C-Br bond to restore the fluorescence. The fluorescence is quenched by the bromides by the induction of intersystem crossing.¹⁵ A completely different strateg *t* uses photoactivated click chemistry of a tetrazol with different dipolarophiles to form fluorescent cycloadducts. Irradiation *t* tetrazole induces the elimination of nitrogen. The resulting high *t* reactive nitrile imine can form fluorescent products wit...

The release of nitrogen was also used in the design c photoactivatable push-pull fluorophores.¹⁷ The electron donating amino group is masked as azide moiety. Upon illumination, the ryl azide releases nitrogen and is transformed to an aryl amine, thus restoring fluorescence. The compatibility of these systems wit super-resolution imaging was also demonstrated.¹⁸

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⁺ Electronic Supplementary Information (ESI) available: Synthetic procedures and characterization of the new compounds, one- and two-photon illumination procedures, widefield and super-resolution microscopy details. See DOI: 10.1039/x0xx00000x

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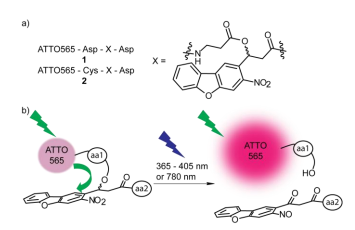


Fig. 1 a) Small peptides 1 and 2 with the new linker in the middle and the fluorophore on the N-terminus were synthesized. b) Photo cleavable activation of ATTO565 can be performed with various wavelengths. The quenching effect is indicated with the green arrow.

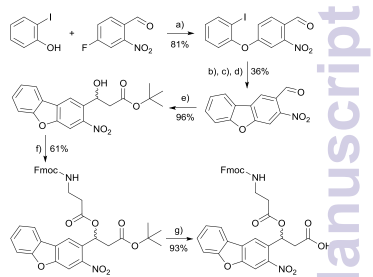
An alternative way to trigger fluorescence with light is to attach a fluorophore and a quencher in close proximity. The photolabile groups act either as linker between fluorophore and quencher¹⁹ or the cages force a structure in a special conformation. Upon irradiation separation of fluorophore and quencher takes place and fluorescence is restored. For instance, both of these techniques were used to follow mRNA distribution in cells with molecular beacons.²⁰

In this work we present a new strategy where the photolabile group is not directly attached to the fluorophore. Our newly synthesized linker molecule combines the properties of a quencher and a photolabile group (Fig. 1). This linker is especially interesting for protein and peptide applications because it can be incorporated by standard peptide coupling chemistry. We prepared small peptides containing the new linker and investigated the photochemical properties of the new constructs with one-photon excitation (shown here for 365 – 568 nm) and two-photon excitation (shown here for 780 nm). To the best of our knowledge the new linker is the first example for a photolabile quencher.

Furthermore the new construct offers the possibility to be used as strand break that can be activated with IR light. Strand breaks are prominent tools to control biomolecules such as oligonucleotides²¹ and peptides²². Future application in which the linker could trigger self-assembly processes²³ or control the activity of enzymes and proteins with high spatial resolution²⁴ are possible.

We observed in a previous study, that the nitrodibenzofuran (NDBF) caging group possesses the ability to quench the fluorescence of the rhodamine fluorophore ATTO565 presumably by contact quenching.²⁵ The NDBF group is known to be cleaved with UV light and also above 400 nm.²⁶ NDBF possesses a good two-photon absorption cross section, which allows to trigger the photoreaction with IR light.²⁷

The first step of the synthesis was reproduced from an already published procedure.²⁸ 4-Fluoro-2-nitro-benzaldehyde was coupled with 2-iodophenol in an Ullmann reaction. The resulting aldehyde was protected as an acetal and the dibenzofuran core was generated by palladium-catalysed cross coupling. After acidic deprotection, the aldehyde was transferred to an alcohol by



Scheme 1 Synthesis scheme of the new linker molecule. a) CuBr, K_2 b) Ethyleneglycol, p-toluenesulfonic acid. c) Cs₂CO₃, Pd(OAc)₂. d) HCl. e) Rieke-Zn *tert*-butyl chloroacetate. f) Fmoc- β -alanine, EDC, DMAP. g) TFA. For deta procedures and analytical data see ESI[†].

Reformatzki reaction and further coupled to $Fmoc-\beta$ -alanine. Find deprotection of the carboxylic acid resulted in the new linker which can then be used in standard Fmoc peptide coupling chemistry (Scheme 1).

First, we checked if base-induced elimination is a problem in furtiapplications of the new linker and confirmed that it is sufficiently stable in 20% (v/v) piperidin in DMF – a typical cleavage condition for Fmoc groups in solid-phase peptide synthesis (ESI⁺).

To study our new linker it was embedded into two short peptides and **2** (Fig. 1). Both contained aspartic acid residues for solubility purposes and an optional cysteine (peptide **2**) for maleimic coupling. The N-terminus was labelled with an ATTO565-Nhydroxysuccinimide (NHS) active ester. Peptide **1** was soluble small volumes of aqueous Tris and HEPES buffered saline (HBS) buffers (pH 7.5) to at least 2 mM. We observed less than degradation of the linker over 170 h in the dark and at room temperature in HBS buffer.

The uncaging quantum yield of peptide **1** was determined to be 0.13 at 365 nm with ferrioxalate actinometry.²⁹ A significary fluorescence increase of this compound was observed upon uncaging (81 ± 6 fold increase in the absolute fluorescence, Fig. 2. 22 fold increase in fluorescence quantum yield, ESI⁺). This makes the new construct very interesting for bioimaging applications. The dye could also be photoactivated with a widefield microscope at 405 nm illumination and even with 568 nm (ESI⁺).

To prove that the NDBF group incorporated into the peptides is stintwo-photon active, molecule **2** was immobilized in a maleim¹ 2containing hydrogel. The irradiation was performed with a lister scanning microscope at 780 nm in a defined square region with increasing laser power (Fig. 3). Fig. 3b (middle) shows that indee 1 an excellent three-dimensional activation could be obtained lift two-photon excitation. At high laser intensities bleaching of ATTO565 in the middle of the focal plane was observed (Fig. 3¹, right).

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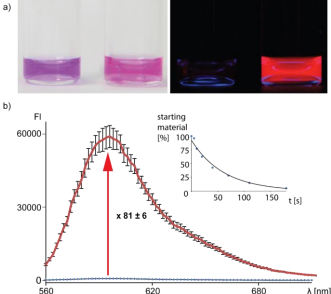


Fig. 2 Fluorescence increase after irradiation of peptide **1** at 365 nm with a 250 mW LED. a) Photographs of 44 nmol peptide in 200 μ L HBS buffer before and after irradiation for 3 min under white light (left picture) and with a TLC lamp at 366 nm (right picture). b) Fluorescence spectra of the new construct before (blue line) and after UV illumination (red line). The inlet shows the time course of the uncaging reaction determined by HPLC.

With the correct settings two-photon irradiation offers the possibility to activate and bleach the fluorophore consecutively. Due to the activation with IR light this procedure allows high tissue penetration depths with little phototoxicity.

To make the new construct a valuable marker for biomolecules, peptide **2** was reacted with 6-maleimidohexanoic acid N-hydroxysuccinimide ester, a bifunctional linker. The resulting NHS ester was coupled to an antibody and used for staining of microtubules of U2OS cells via immunofluorescence. These

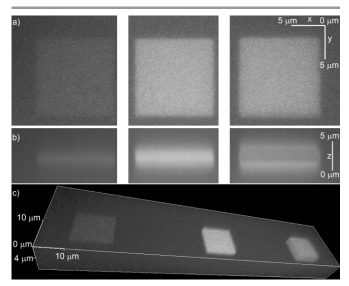


Fig. 3 Two-photon activation of the fluorescence of 2 in a hydrogel. Panels a) – c) show different projections of the same sample. Irradiation was performed at 780 nm in a laser scanning microscope. The fluorescence was recorded continuously at 561 nm. The left square was irradiated with 6 mW, the middle with 14 mW, the right one with 20 mW, each time 20 scans. The fluorescence increases (left squares to the middle ones) until bleaching starts (right squares).

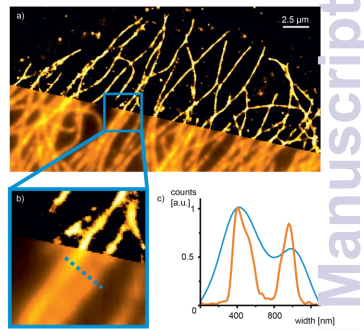


Fig. 4 Conventional and super-resolved image of the microtubulin network of an U₁, cell stained via immunofluorescence with the photoactivatable peptide **2** antiboc conjugate. a) Reconstructed super-resolved image (upper right) and diffractionlimited widefield image (lower left). b) Magnified view of the boxed region highlighte in a). c) Cross-sectional intensity profile through neighboring microtubulin filaments marked in b) (blue dotted line). The intensity profile demonstrates the resolu¹ improvement from the widefield (blue line) to the super-resolved image (orange line).

filamentary structures are an ideal target to evaluate the suitabilit of a fluorophore for super-resolution microscopy following the concept of single-molecule localization.³

We demonstrated one-photon activation of the antibody conjugate with a short 405 nm laser pulse and continuous illumination with 568 nm (ESI⁺, Fig. S5). Furthermore, we illuminated the sample with 568 nm only and found a consecutive activation and bleaching or blinking of single peptide **2** molecules at appropriate laser intensities (ESI⁺, Fig. S6). We used standard localization algorithes to fit a 2D Gaussian function to the diffraction-limited intensity profile of each single emitter in order to determine their center of mass. Reconstruction of these localizations in a 2D histogram yielded an image with a significantly improved spatial resolutio (Fig. 4) and an experimental localization uncertainty of 20.4 nm determined by nearest neighbor analysis.³⁰ Notably, we thereby demonstrate single-molecule super-resolution imaging with synthetic fluorophores without the need of chemical additives in the imaging buffer.⁴

In summary, we developed the first photolabile quencher for ATTO565 with a light/dark contrast of 81:1. It can be incorporated into peptides and acts both as quencher and strand break. The ϵ sy way of functionalization of the linker with other molecules like hydrophilic amino acids results in a good solubility in aqueous buffers (pH 7.5). The linker shows very little degradation under these conditions. Irradiation can be performed with UV light, als pabove 400 nm which is more favourable for living cells. Additionally, the construct is two-photon accessible that allows very high spatications. The linker provides a new tool in the field of bioimaging. It reduces the number of modifications needed

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generate a photoactivatable fluorophore quencher system from three (fluorophore, quencher and photolabile group) to two (fluorophore and new linker molecule). In addition, the high contrast between the fluorescent and dark state makes this probe ideally suited for single-molecule super-resolution microscopy. In the future we will investigate the mechanism of the quenching effect and develop an application of the new linker to other fluorophores.

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