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Selective release of a potent anticancer agent from a supramolecular hydrogel using green light†

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Light-triggered drug release from hydrogels is a promising method to improve efficiency of antitumor treatment, as an alternative to existing photodynamic therapies. Here we report a photochromic supramolecular low-MW hydrogel that can quickly and selectively release a physically encapsulated potent anticancer agent upon green light irradiation under physiological conditions.

Numerous advanced drug delivery systems are based on hydrogels, which can protect labile drugs from premature degradation, or maintain constant drug concentration in blood over prolonged periods of time. Macroscopic gels can be used in the form of implants, whereas microgels allow for oral and pulmonary delivery, and nanogels can be systemically administered into the bloodstream of patients.2,3 While covalent hydrogels are based on polymers (gelatin, alginate, agarose), supramolecular low-MW hydrogels4 can be formed by mutual non-covalent interactions of short linear oligopeptides,5-10 nucleobases,11 or urea derivatives.12,13 Hydrogels can be triggered by external stimuli, such as pH changes or enzymes, to release encapsulated drugs in a controllable fashion.14-17 Light is a particularly attractive stimulus in this context, 18 because it can be applied on living organisms with high spatiotemporal precision without their permanent contamination. Lightresponsive supramolecular hydrogels, 19-23 often tailored to particular applications like cargo release, 24,25 or shape memory,26 frequently contain molecular photoswitches structures that can reversibly interconvert between two photoisomers upon irradiation.27 The associated modulation of geometry, polarity, or other molecular properties may elicit macroscopic effects in various materials.28-33

Common photoswitches are usually triggered with UV light, but this frequency does not penetrate living cells or tissues over, hydrogels based on the bis-ortho-fluoroazobenzene pho-

tochrome (sensitive on green light) only formed above the

critical gelator concentration (CGC) of 40 g L⁻¹. Consequently,

complete gel dissipation required relatively long irradiation time (≥ 3 h).⁵⁵ The selectivity and timespan were sufficient for

a proof-of-principle demonstration of green-light control over bacterial growth by photocontrolled antibiotic release.

However, these parameters must be improved before thera-

efficiently. Therefore, photoswitches triggered with visible light are

developed for applications in complex biological systems.³⁴⁻⁴¹ In azobenzenes, such a bathochromic shift is usually associated with

loss of thermal stability. Notable exceptions are ortho-fluorinated

azobenzene derivatives, mutually switchable with green (trans \rightarrow

cis) and violet (cis \rightarrow trans) light between two photoisomers, which

are thermally (meta)stable at ambient conditions. 42 This motif was

peutic applications were conceivable for human subjects.

In this communication, we report a successful strategy towards decreasing the CGC, and increasing selectivity of the light-induced cargo release in photochromic supramolecular

hydrogels based on cyclic dipeptide gelators.

It is known that aromatic C-F bonds may generate new non-covalent interactions in proximity of the fluorine atoms.⁵⁶ We hypothesized that the tetra-*ortho*-fluorinated azobenzene

used in numerous light-triggered materials43-45 and biological applications.46-50 Our group has previously merged the excellent hydrogelation propensity of amphiphilic cyclic dipeptides⁵¹⁻⁵³ with the photoswitchable azobenzene motif. The resulting biocompatible photochromic hydrogels can reversibly dissipate to fluids upon irradiation (due to the reversible polarity change ensuing azobenzene photoisomerization) under physiological conditions, concomitantly release encapsulated bioactive substances.54,55 However, due to the non-covalent nature of the encapsulation, substantial spontaneous cargo diffusion ("leaking") from loaded gels in darkness was often observed, thus compromising selectivity of the light-triggered release. More-

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Paper

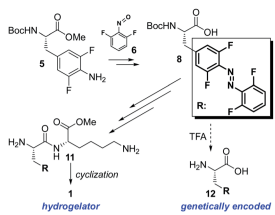
(TFAB) should improve supramolecular association of the fibre subunits and thus decrease the critical gelator concentration in comparison to the previously demonstrated hydrogelator, based on the bis-ortho-fluorinated azobenzene.55

The hydrogelator 1 (Fig. 1) has been prepared according to the synthetic route depicted in Schemes 1 and S1.† The crucial intermediate – bis-fluorinated (S)-phenylalanine analogue 5 was synthesized from 4-bromo-2,6-difluoroaniline (4) and the protected (R)-iodoalanine (3) using Negishi coupling. Mills reaction of 5 with in situ generated 2,6-difluoronitrosobenzene (6) and subsequent methyl ester hydrolysis provided the photochromic amino acid derivative 8 (>96% ee, Fig. S14†), which upon coupling to an (S)-lysine derivative and cyclization of the linear dipeptide 11 yielded the hydrogelator 1. Additionally, full deprotection of 8 may provide an enantioselective access to the amino acid 12, previously used as racemate for biosynthetic incorporation into luciferase protein using the amber codon suppression method.⁵⁰

Irradiation of 1 in aqueous solution (phosphate-buffered saline, pH 7.4 - a typical buffer for biological experiments, abbreviated below as PBS, composed of 8 g NaCl, 0.2 g KH₂PO₄, 1.15 g Na₂HPO₄, and 0.2 g KCl in 1 L of water) with green light (530 nm LED, 7 mW cm⁻²) until equilibration resulted in the mixture containing 89% of the cis-1. Further exposure of this solution on violet light (410 nm LED, 9 mW cm⁻²) produced a mixture containing 23% of the cis-1 (Fig. S1 and Table S1†). In darkness, the thermal $cis \rightarrow trans$ isomerization of 1 at r.t. is practically suppressed. At 60 $^{\circ}\text{C},$ the thermal isomerization of cis-1 ($t_{1/2}=70.9\pm0.6$ h at 60 \pm 2 °C, in MeCN, Fig. S2† left) is slightly faster than for the unsubstituted cis-TFAB ("F4", $t_{1/2}$ = 92 h at the same temperature in MeCN⁴²).

Another common limitation for in vivo applications of azobenzene derivatives in biological systems is their reduction with biogenic thiols to arythydrazines. The compound 1 is resistant (below 8% of degradation within 6 days of incubation, Fig. S2† right) to 5.0 mM reduced glutathione solution - an established model of the intracellular redox environment.

Fig. 1 Photochromic cyclic dipeptide trans-1 is a low-MW supramolecular hydrogelator, while its photoisomer cis-1 forms non-viscous fluids at the same concentrations; a photochromic hydrogel based on ${f 1}$ (here, 17 g L $^{-1}$ of ${f 1}$ in PBS, pH 7.4) reversibly dissipates to a fluid upon irradiation with green light.



Scheme 1 Schematic depiction (detailed on the Scheme S1†) of synthesis of the photochromic supramolecular hydrogelator 1 bearing the "TFAB" photoswitch (framed and abbreviated as R). Upon deprotection, the molecule 8 can also yield the TFAB-substituted (S)-alanine 12, which has previously been biosynthetically incorporated into proteins using the amber codon suppression technique. 50

The compound 1 forms stable and homogenous hydrogels in aqueous solutions under physiological conditions (PBS buffer, pH 7.4, Fig. 1) at the concentrations \geq 15 g L⁻¹ of 1 (Table S2†). Although this value is still over a magnitude higher above the most efficient UV-light-triggered super(organo)gelators (CGC < 0.1 wt%),21,22 our material enables drug release in water with visible light - not reported for supergelators yet.

After 30 minutes of irradiation with green light (530 nm, power input of 3 W, 7 mW cm⁻², 22 ± 2 °C), hydrogels containing 15 g L^{-1} or 17 g L^{-1} of 1 turned into homogenous fluids without any mechanical stimulation (Fig. 1 and S4†), while the hydrogel containing 20 g L⁻¹ of 1 needed slight mechanical stimulation (e.g. by shaking the vial). Gels containing 30 g L^{-1} of 1 or above withstood 60 min of irradiation under the same conditions without any visible phase transition. All the aforementioned fluids remained liquid for the period of at least one week when stored in darkness at room temperature. The fluid produced upon green-light irradiation of the hydrogel containing 20 g L⁻¹ of 1 has been subsequently irradiated with violet light (410 nm LED, 9 mW cm⁻²) for 60 min and incubated overnight (16 h) at room temperature in darkness. After that time, the sample turned into a transparent hydrogel again, with mechanical stability comparable to the non-irradiated sample (Fig. S5†). The internal fibrous structure of the hydrogels has been visualized using transmission electron microscopy (TEM) (Fig. S6†). We have also investigated gelation of 1 under a broader range of pH values in aqueous buffers at the constant concentration of 17 g $\mathrm{L^{-1}}$ of 1. At the pH 10, the material did not dissolve up to the boiling point and no gelation was observed upon cooling. Samples buffered to the pH 4, pH 6, and pH 8 formed homogenous solutions above c.a. 60 °C, and upon short boiling formed homogenous hydrogels within 1 h of cooling at room temperature. The hydrogel formed at pH 4 was mechanically unstable and dissipated upon slight shaking or vial inversion. Hydrogels formed at pH 6 and pH 8 exhibited similar mechanical and thermal stability as the hydrogel prepared at the pH 7.4 from 17 g L^{-1} of 1. (Fig. S3†).

Our next goal on the way to therapeutic relevance of our drug-releasing hydrogels was to minimize the passive cargo diffusion ("leaking") from non-irradiated gel samples, which previously varied between 5% and 30% of respective light-induced release rates. We noticed that the "leaking" is less pronounced for the cargo negatively charged under physiological conditions (*e.g.* DNA oligomers, ⁵⁴ or carboxylic acids ⁵⁵). This was attributed to electrostatic interactions with the basic, positively charged lysine side chains, likely exposed on the fiber surface of the gel matrix. This interaction would, in turn, slow down the diffusion of acidic cargo through cavities of the gel matrix.

Here we decided to assume a different approach. Fast self-healing of our gels (<1 min, based on rheological determination of the storage modulus G' recovery upon 100% deformation, Fig. S5 \dagger) indicates its highly dynamic nature, where supramolecular interactions between monomeric units are quickly reconstituted upon deformation. We hypothesized that a cargo bearing the cyclic dipeptide motif connected to an aromatic residue would be promptly incorporated into the fibrous structure of the hydrogel (Fig. 2), instead of being absorbed in the cavities between fibers. As photoisomerization breaks the fibrous structures, the non-photochromic cargo would then be released from the supramolecular network upon hydrogel erosion.

Many clinically approved drugs contain the cyclic dipeptide motif (2,5-diketopiperazine, DKP) bound to aromatic residues. They show a broad spectrum of activities – to name only tadalafil (the active ingredient of "Cialis®" – a drug against erectile dysfunctions), retosiban (an oxytocin receptor inhibitor used for the treatment of premature birth), or aplaviroc (a CCR5 entry inhibitor developed for treatment of HIV infections).⁵⁷

To test our hypothesis, we have selected a strong antiproliferative agent plinabulin (2) (Fig. 2), which is a promising anticancer drug candidate.⁵⁸ It acts as a low-nanomolar inhibitor of tubulin depolymerization, which hampers cytoskeleton dynamics. The resulting mitotic arrest stops cell division and

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Trans-1

Fig. 2 Schematic representation of supramolecular interactions between the gelator 1 and a potent anticancer agent plinabulin (2). Hydrogen bonding and hydrophobic interactions of aromatic rings, most likely stabilize drug molecules inside of the fibrous network of the hydrogel and prevent its leaking in darkness.

the cancer growth.⁵⁹ 2 shares structural similarities, in particular the cyclic dipeptide residue and an aromatic substituent, with the gelator 1. We have synthesized 2 following a literature procedure (Scheme S2†).⁶⁰

Afterwards we encapsulated 2 in the hydrogel composed of 17 g L⁻¹ of 1 in PBS pH 7.4, and characterized its release upon dissipation of the gel resulting from green light irradiation. To our delight, the cargo could be recovered almost quantitatively in solution upon light-driven gel dissipation within 30 min, while "leaking" from identical gels incubated in darkness was negligible (below 1%, Fig. 3). For comparison, an antibiotic ciprofloxacin (with similar MW and a structure unrelated to the gelator 1) encapsulated and released upon light-induced dissipation of the otherwise identical gel composition showed over 25% leaking in darkness within the same time period (Fig. S7†).

Due to limited solubility of plinabulin (2) in water, stock solutions of 2 were prepared in DMSO and diluted 100-fold in aqueous solutions for the final formulations. We determined the aqueous solubility of 2 (PBS pH 7.4, 1% DMSO) in absence of the gelator 1 to be $7.5 \pm 1.7 \, \mu \text{mol L}^{-1}$ (2.5 mg L⁻¹). Above this concentration, visible precipitation appeared within 60 minutes from diluting the DMSO stock solutions with PBS (for the original drug, the issue of low water solubility was addressed *e.g.* by covalent functionalization with hydrophilic PEG groups).

However, hydrogels (17 g L^{-1} of 1) comprising the compound 2 at final concentrations \leq 350 µmol L^{-1} (118 mg L^{-1}) (by adding respectively more concentrated DMSO stock solutions during the preparation process) were still transparent and without any precipitation upon prolonged time (>16 h). This additionally supports the hypothesis that 2 is being incorporated into the supramolecular structure of the hydrogel upon its formation. The dopant 2 still constitutes below 1% of the dry mass of the gelator 1 used in the respective composition,

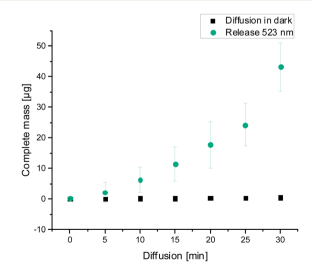


Fig. 3 Light-induced release of encapsulated plinabulin 2 (56 μ g) upon dissipation of the hydrogel (0.5 mL) containing 17 g L⁻¹ of 1 caused by irradiation with green light (530 nm LED, 7 mW cm⁻²) (green circles). Passive diffusion ("leaking") of 2 from the same hydrogel in darkness (black squares) is negligible. Suppression of the "leaking" is attributed to the supramolecular association of 2 and *trans-*1 in fibres.

therefore no significant alterations of the gel physical properties (melting temperature, or mechanical strength) have been observed.

An important factor in potential therapeutic applications of the demonstrated drug-releasing material is the toxicity of the drug delivery vehicle to human cells. This was determined for our gelator 1 using cell viability assays (MTT assays). We have treated one human cancerous (HeLa) and one non-cancerous (NHDF - fibroblasts) cell line with increasing concentrations of the gelator 1 as pure trans-isomer (the thermally stable form that forms hydrogels in water), or as a mixture obtained upon irradiation of trans-1 with green light until the respective photostationary state was achieved (this mixture corresponds to the dissipated hydrogel). In all but one combination (Fig. S9†) we observed no significant decrease in cell viability up to the concentrations of 1 mM of 1 (c.a. 0.45 g L^{-1} , slightly below the CGC values). Only in the case when cancerous HeLa cells were exposed to the pure trans-1, the cell viability strongly decreased above the concentration of 0.1 mM of the gelator 1.

To further assess applicability of our composition for therapeutic purposes, we considered that the irradiation power required for the gel dissipation in our experiments (<10 mW cm⁻²) is considerably lower than the safety limits determined for human tissues (200 mW cm⁻²).²⁴ In contrast to photodynamic therapies (PDT) used clinically to treat malignant cancers,⁶¹ tissues targeted with our gel composition would not need to be oxygenated. Furthermore, the clinically existing PDT infrastructure could be used for the release of 2 upon photodegradaton of our hydrogel.

In the future, the efficiency of our system will have to be confirmed *in vivo* (*e.g.* in model mice with tumor xenografts), before human therapeutic applications can be considered. Particularly suitable therapeutic targets for the currently presented composition seem to be retina cancers or squamous carcinoma (skin cancer), which appear in organs that can be efficiently penetrated with green light.

The composition demonstrated here could be further formulated as micro/nanogel (with eventual stabilization, e.g. by covering with biocompatible alginate), and injected into the bloodstream, or given upon inhalation. With the reported lownanomolar antitumor potency of plinabulin 2 ($IC_{50} = 15$ nM against human colon cancer HT-29 cells60), systemic introduction of such a microformulated hydrogel containing 0.35 mM of 2, followed by its rapid and close-to-quantitative light-driven discharge in the selected area, should produce sufficiently high local drug concentration to successfully eradicate neighboring tumor cells. At the same time, the remaining drug will circulate in darkness in its inactive form (entrapped in the gel fibers), eventually metabolized and cleared from the organism over a prolonged period of time, thus minimizing the systemic toxicity and resulting in better therapeutic effect in comparison to neat plinabulin.

The remaining challenge is modification of the photochromic segment of our system to cause the gel dissipation upon irradiation with red or near-IR frequencies (within the socalled "therapeutic window" of light, 650–900 nm), capable of deep penetration (>1 cm) of soft human tissues. This would enable *e.g.* a general strategy for targeting of solid tumors inside of human organs without surgical intervention with established and efficient cytotoxic agents, yet with minimizing systemic side effects. Therefore, in the future we will examine the applicability of *ortho*-alkoxy and *ortho*-chlorinated azobenzenes, which were efficiently switched upon red light irradiation, ^{39,40} in our hydrogel systems.

Conclusions

In conclusion, we demonstrated that the gelator 1, non-toxic for non-cancerous human cells below 1 mM concentrations, forms stable supramolecular hydrogels at and above concentrations of 15 g $\rm L^{-1}$ under physiological conditions and in the pH range 6-8. The gels can be reversibly dissipated to fluids with green light, and concomitantly release previously encapsulated cargo. Moreover, we identified a potent anticancer agent plinabulin 2 as an optimal therapeutic agent, which due to the structural similarity with 1 can be encapsulated and quickly released from the dissipating hydrogel upon irradiation, while its diffusion ("leaking") outside of the intact gel in darkness is negligible. The concentration of 2 in our supramolecular composition can be over 40-fold higher than its aqueous solutions, without disturbing the gel homogeneity. This additionally enhances delivery efficiency of 2 in therapeutic use.

Formulation of such compositions as injectable or inhalable micro/nanogels may enable systemic introduction into organisms. Subsequent cargo release with laser or LED green irradiation will produce a high local concentration of the cytotoxic agent. The local dose can be much higher than the concentration achieved upon simple drug injection, without causing severe side effects outside of the irradiation area, and thus enable faster and more efficient cancer eradication. Due to the penetration propensity of green light, eye and skin cancers remain the primary targets for this composition. In contrary to photodynamic therapies, the targeted tissues do not have to be oxygenated.

Extension of the described methodology to the red- or NIR-light-induced gel dissipation is also foreseen to enable targeting of solid human tumours on the organismal scale. The release of other drugs based on the cyclic dipeptide (DKP) motif will be also investigated.

Conflicts of interest

There are no conflicts to declare.

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