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Photosensitizer-free visible light-mediated goldcatalysed cis-difunctionalization of silylsubstituted alkynes†

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A new photosensitizer-free visible light-mediated gold-catalysed cis-difunctionalization reaction is developed. The reaction was chemoselective towards silyl-substituted alkynes with excellent regioselectivity and good functional group compatibility, giving a series of silyl-substituted quinolizinium derivatives as products. The newly synthesized fluorescent quinolizinium compounds, named JR-Fluor-1, possessed tunable emission properties and large Stokes shifts. With unique photophysical properties, the fluorophores have been applied in photooxidative amidations as efficient photocatalysts and cellular imaging with switchable subcellular localization properties. **EDGE ARTICLE**
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Substituted alkyne

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Introduction

Over the past years, homogeneous gold catalysis has become a powerful tool for the synthesis of complex organic molecules, particularly in the activation of C–C multiple bonds with gold catalysts acting as strong soft π -Lewis acids.¹ In recent years, instead of accessing hydrofunctionalized products, a promising cross-coupling strategy involving $Au(I)/Au(m)$ catalytic cycles has been developed to afford difunctionalized products.² With a significantly high redox potential of the $Au(I)/Au(m)$ couple, strong oxidants are required to access the oxidation of $Au(1)$ species, while the functional group compatibility could be hampered.^{3a} A complementary approach to overcome this barrier was developed by Glorius³ and Toste⁴ through merging gold catalysis with photoredox catalysis, which promoted oxidation of Au(1) species employing photosensitizers and aryl radicals generated *in situ* under irradiation. This approach inspired the development of diverse organic transformations including difunctionalization of alkenes, $3b,c,4b$ allenes^{5a} and alkynes,^{3e,5b,c} as well as C–C^{3d,4d,5d-f} and C–P^{4c} cross-coupling reactions. Recently, Hashmi and co-workers reported a visible light-mediated gold-catalysed oxyarylation of alkynes^{6a} and an

aryl-aryl coupling reaction^{6b} which could be conducted without addition of photosensitizers.

For alkyne difunctionalization reactions undergoing an antinucleophilic addition, trans-difunctionalized products are afforded in the majority of the examples.^{3c,e,5c,6a} However, visible light-mediated cis-difunctionalization of alkynes still remains largely unexplored.⁷ Inspired by a stereo- and regioselective gold-catalysed hydroamination of internal alkynes reported by Stradiotto et al.,⁸ we set out to combine visible light mediated $Au(i)/Au(m)$ catalysis with a plausible syn insertion pathway to achieve alkyne *cis*-difunctionalization products (Scheme 1).^{7b,c}

Development of organic fluorescent materials has become an emerging and important research area due to their wide applications in chemistry, biology and materials science.⁹ Compared to the intensive investigations on scaffolds such as fluorescein, BODIPY and, recently, Seoul-Fluor,¹⁰ only a few examples of cationic fluorophores have been reported, although the unique properties of fluorophores containing positive charge have been demonstrated in photocatalysis and cellular imaging.¹¹ Quinoliziniums, cationic aromatic heterocycles bearing a quaternary bridgehead nitrogen, were firstly investigated in alkaloid chemistry and recently employed as efficient DNA intercalators.¹² However, structure photophysical property relationship (SPPR) studies and applications in photocatalysis and cellular imaging of quinolizinium compounds still remain elusive.¹³

Along with our ongoing interest in gold-catalysed organic transformations,¹⁴ herein, we report a new photosensitizer-free visible light-mediated gold-catalysed alkyne cis-difunctionalization reaction affording a series of silyl-substituted fluorescent quinolizinium compounds. Spectroscopy experiments and DFT calculations were conducted to study the SPPR of the

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[‡] Dr Sharon Lai-Fung Chan designed and performed the spectroscopic studies and computational experiments of this research. Dr Ben Chi-Bun Ko designed the cellular imaging experiments.

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quinolizinium derivatives. Applications of the fluorescent quinolizinium compounds as efficient photocatalysts for photooxidative amidation and as fluorescent dyes for live cell imaging have also been conducted.

Results and discussion

We initiated the reaction screening by treatment of quinolinesubstituted aryl diazonium 1a (0.12 mmol, 1.2 equiv.), and (4 uorophenylethynyl)trimethylsilane 2a (0.10 mmol, 1 equiv.) with Ph_3PAuCl 3a (10 mol%) to afford silyl-substituted quinolizinium 4a in 71% yield (Table 1, entry 1). To our surprise, no cross-coupling product 4aa (reported by Toste et al. using phenyl diazoniums as substrates) was obtained in our experiments.^{4d} Besides, the reaction was exceptionally regioselective giving no regioisomer $4a'$. The gold $\scriptstyle\rm (i)$ catalyst $3b$ bearing comparatively electron-rich phosphine led to a lower yield of product formation (entry 2), while catalysts 3c–d bearing electron-poor phosphine gave higher yields of 4a (entries 3–4). No desired product was obtained when Ph_3PAuBF_4 3e was employed as a catalyst (entry 5). Using $gold(r)$ catalyst 3f possessing a diphosphine ligand led to product formation in 21% yield (entry 6). Only a trace amount of product was observed when simple AuCl 3g was used (entry 7). In addition, employing gold(m) catalysts 3h-j resulted in no product formation (entries 8–10). Without the addition of gold catalyst, no desired product was observed (entry 11), indicating that the gold (i) phosphine chloride catalyst played a crucial role in this transformation. Addition of the transition metal-based photocatalysts $Ru(bpy)_{3}Cl_{2}$, $Ru(bpy)_{3}(BF_{4})_{2}$ and $Ir(ppy)_{3}$ or the organic photocatalyst Rose bengal led to lower yields of the desired product (entries 12–15). A control experiment in the dark gave no product formation (entry 16). These results suggested that this reaction was conducted under photosensitizer-free reaction conditions. Additionally, only a trace amount of product was detected when the reaction was conducted in toluene, dichloromethane or methanol as the solvent (entries 17–19), and a lower yield (35%) of product was afforded when the reaction was carried out in open air (entry 20). Open Access Article. Published on 04 September 2017. Downloaded on 9/22/2024 6:11:29 PM. This article is licensed under a [Creative Commons Attribution-NonCommercial 3.0 Unported Licence.](http://creativecommons.org/licenses/by-nc/3.0/) **[View Article Online](https://doi.org/10.1039/c7sc02294h)**

With the optimized reaction conditions, we expanded the scope of this reaction by using various diazoniums (0.60 mmol, 1.2 equiv.) and alkynes (0.50 mmol, 1.0 equiv.) as substrates. Silyl-substituted alkynes 2a–q bearing ether, alkyl, halogen, aldehyde, carboxylic acid, cyanide, trifluoromethyl, nitro and hetero-aromatics as substituents were well tolerated with the

gold-catalysed alkyne cis-difunctionalization.

Table 1 Optimization of the reaction conditions and control experiments^a

Entry	Au cat.	Photo cat.	Light	N_2/air	Solvent	Yield ^b [%]
$\mathbf{1}$	3a		Blue ^c	N_2	CH ₃ CN	71
$\overline{2}$	3b		Blue ^c	N_2	CH ₃ CN	58
3	3c		Blue ^c	N_2	CH ₃ CN	75
4	3d		Blue ^c	N_2	CH ₃ CN	83
5	3e		Blue ^c	N_2	CH ₃ CN	n.d. ^d
6	3f		Blue ^c	N_2	CH ₃ CN	21
7	3g		Blue ^c	N_2	CH ₃ CN	Trace
8	3h		Blue ^c	N_2	CH ₃ CN	n.d. ^d
9	3i		Blue ^c	N_2	CH_3CN	n.d. ^d
10	3j		Blue ^c	N_2	CH ₃ CN	n.d. ^d
11			Blue ^c	N_2	CH ₃ CN	n.d. ^d
12	3a	$Ru(bpy)_3Cl_2$	Blue ^c	N_2	CH ₃ CN	9
13	3a	$Ru(bpy)_{3}(BF4)_{2}$	Blue ^c	N_2	CH ₃ CN	Trace
14	3a	$Ir(ppy)_3$	Blue ^c	N_2	CH ₃ CN	Trace
15	3a	Rose bengal	Blue ^c	N_2	CH ₃ CN	65
16	3a		Dark	N_2	CH ₃ CN	n.d. ^d
17	3a		Blue ^c	N_2	Toluene	Trace
18	3a		Blue ^c	N_2	CH_2Cl_2	Trace
19	3a		Blue ^c	N_2	CH ₃ OH	Trace
20	3a		Blue ^c	Air	CH ₃ CN	35

Reaction conditions: treatment of 1a (0.12 mmol), 2a (0.10 mmol) and gold catalyst 3a–j (10 mol%) with or without photocatalyst (5 mol%) in 5 mL of solvent under N₂ at room temperature for 16 h. $\frac{b}{b}$ Yield of 4a was determined by ¹⁹F-NMR using fluorobenzene as the internal standard.
^{*c*} Blue LEDs ($\lambda_{\text{max}} = 469$ nm) were employed as a light source. ^{*d*} n.d.: product formation could not be detected.

reactions giving quinolizinium products 4a–q in 31–69% yield. Increasing the steric bulkiness using ortho-disubstituted phenylethynylsilane 2r gave a trace amount of product. Using a triethylsilyl group gave quinolizinium 4u in slightly lower yield (45%). Further increasing the bulkiness employing tertbutyldiphenylsilyl-substituted alkyne 2t gave no product 4v. Interestingly, employing terminal alkyne phenylacetylene 2u, cyclohexylacetylene 2v or internal alkyne 1,2-diphenylethyne 2w, 1-phenyl-1-propyne 2x or 1-phenyl-1-butyne 2y gave no desired product 7a–e, suggesting that the reaction was chemoselective towards silyl-substituted alkynes. To support our hypothesis, alkyne 2z bearing diphenylethynyl and trimethylsilylethynyl groups was used, and only the trimethylsilylethynyl cis-difunctionalization product 4w was obtained in 46% yield, which convinced us of the chemoselectivity of this goldcatalysed transformation. Further expansion of the scope employing diazoniums 1b–d bearing different structure skeletons and heterocycles gave the desired products 5a–c and 6a in Scheme 1 Literature works and our strategy for visible light-mediated up to 60% yield. These results indicated that the reaction was

 a Reaction conditions: treatment of $1a-d$ (0.60 mmol), $2a-z$ (0.50 mmol) and 3a (10 mol%) in 5 mL of CH₃CN under irradiation (blue LEDs) and N_2 at room temperature for 16 h. $\frac{b}{c}$ Isolated yield.

very compatible with various diazoniums and highly selective towards silyl-substituted alkynes (Table 2).

To provide insight into this visible light-mediated goldcatalysed cis-difunctionalization reaction, stoichiometric reactions were set up by treatment of aryl diazonium 1a (0.10 mmol) with Ph_3PAuCl 3a (0.10 mmol) under irradiation for 0 to 240 min. ¹H-NMR monitoring revealed that aryl diazonium 1a was consumed and new signals appeared gradually from 0 to 60 min, suggesting the formation of plausible intermediates (Fig. 1a). Irradiation for a longer time gave a slight decomposition of the plausible intermediates. These results were consistent with those observed by $31P-NMR$ analysis, which indicated the gradual appearance of new signals at 31.0, 44.2 and 45.3 ppm (Fig. 1b). The reaction mixtures were further

Fig. 1 (a) ¹H-NMR studies on reaction mixtures $X_{0-240 \text{ min}}$ in CD₃CN; (b) $31P$ -NMR studies on reaction mixtures X_{0-240} min in CD₃CN.

treated with silyl-substituted alkyne 2a (1.5 equiv.) for 60 min in the dark, resulting in the disappearance of the newly observed signals after irradiation and concomitant formation of the signals of the quinolizinium product 4a (Fig. S3a and b in the ESI[†]). The yield of the product formed was monitored by 19 F-NMR analysis through the addition of fluorobenzene as the internal standard (Fig. S3c in the ESI†). In addition, regeneration of Ph_3PAuCl 3a could be recovered as a precipitate from the reaction mixtures.

Due to difficulties in the isolation of the reaction intermediates, we sought to investigate the plausible $Au(m)$ species through ESI-MS analysis by treatment of aryl diazonium 1a (0.12 mmol) with Ph_3PAuCl 3a (0.10 mmol) under irradiation for 1 h. The experimental results indicated that rather than the formation of $A(m/z = 204.0792)$ by the displacement of N₂ of 1a in ESI-MS analysis, plausible Au(III) intermediates **B** $(m/z = 698.1014)$ and $B'(m/z = 960.1910)$ were found (Fig. 2a). Further treatment of the afforded reaction mixture Y with silyl-substituted alkyne 2a (1.5 equiv.) in the dark for 1 h afforded reaction mixture Y' . ESI-MS analysis of reaction mixture Y' revealed that no signal for \bf{B} or \bf{B}' was observed, and the signal of the quinolizinium product $4a$ appeared, suggesting that both B and B' were consumed and reacted with 2a to form the quinolizinium product 4a (Fig. 2b). For a detailed understanding of the plausible Au(m) intermediates **B** and **B**^{\prime}, ESI-MS/MS analysis of species **B** (precursor ion $m/z = 698$) and **B**' (precursor ion $m/z =$

Fig. 2 (a) ESI-MS analysis of the reaction mixture Y; (b) ESI-MS analysis of the reaction mixture Y' .

Fig. 3 (a) UV/Vis absorption spectrum of aryl diazonium 1a; (b) fluorescence quenching of aryl diazonium 1a with Ph₃PAuCl 3a.

910) was conducted. Product phosphonium ions **B-I** $(m/z =$ 459.0483) and Au(I) species **B-II** ($m/z = 466.1634$) were found in the MS/MS analysis of species B (Fig. S4a in the ESI†). Formation of B-I was assumed to be ascribed to the reductive elimination of the $Au(m)$ species **B**, which was previously reported as a feasible deactivation pathway of phosphine-supported aryl $Au(m)$ complexes.¹⁵ In MS/MS analysis of species B' , product ions of **B** ($m/z = 698.0994$), **B-I** ($m/z = 466.1672$) and **B-II** ($m/z =$ 459.0518) were found (Fig. S3b in the ESI†). Results suggested that species B' was composed of Au(m) species **B** and triphenylphosphine and presumably formed by possible transmetallation.¹⁶ A control experiment under the same reaction conditions without irradiation led to no formation of the $Au(m)$ species B , B' or product 4a, suggesting that a light source was necessary for promotion of the $Au(I)/Au(m)$ transformation in this reaction.

To study the photosensitizer-free reaction conditions, we measured the UV/Vis absorption properties of aryl diazonium 1a and Ph_3PAuCl 3a. Spectroscopic analysis revealed that no absorption peak of Ph₃PAuCl 3a was observed at λ_{abs} > 395 nm (Fig. S6 in the ESI†), indicating that Ph_3PAuCl 3a might not be directly excited by the light source in this reaction which was consistent with literature work indicating that the photosensitizer was necessary to initiate the reaction. $3-5$ On the contrary, a tail of the lowest energy absorption peak of aryl diazonium 1a was found at λ_{abs} > 395 nm (Fig. 3a), suggesting that aryl diazoniums could be firstly excited to initiate the reaction. 17 To further support our hypothesis, we measured the difference of the fluorescence intensity before and after mixing Ph_3PAuCl 3a (1 equiv.) with 1a. Spectroscopic analysis indicated that the fluorescence of **1a** could be quenched $(I_q/I_0 = 0.68)$ after the addition of Ph₃PAuCl 3a, providing strong evidence for electron transfer from Ph₃PAuCl 3a to aryl diazonium 1a under irradiation (Fig. 3b). In addition, the estimated excited state reduction potential (E_{red}^*) of aryl diazonium 1a was 3.28 V, which was much higher than the redox potential of the $Au(1)/Au(m)$ couple $(E_0 = 1.41 \text{ V})$,¹⁸ indicating that Ph₃PAuCl 3a could be oxidized by the aryl diazonium 1a in the first step of the reaction.

Based on the aforementioned experimental results, a reaction mechanism is proposed as shown (Scheme 2). Under

Scheme 3 Rhodium-catalysed synthesis of quinolizinium compounds.

Fig. 4 X-ray crystal structure of 4b (front view, left view and vertical view).

irradiation, the aryl diazonium compound is firstly excited and subsequently reduced by single electron transfer from $Au(i)$ catalyst 3a to form an aryl radical with the generation of a $Au(II)$ species. The $Au(\Pi)$ species further recombines with the aryl radical to give $Au(m)$ intermediate **B**. The oxidation and addition on Au(I) catalyst 3a to form Au(II) intermediate **B** is also possibly a concerted reaction pathway rather than a two-step process.^{6a} The reaction quantum yield was found to be 0.91 (less than 1), suggesting that the radical chain process was not prominent in this transformation. Then, silyl-substituted alkyne is activated by Au(III) intermediate **B** through π -activation to give species C. After that, the Au–N bond is regioselectively inserted by the π activated alkyne to form cis vinyl gold species D. Then, reductive elimination provides a quinolizinium compound as the cisdifunctionalized product and regenerates the Au(I) catalyst 3a.

Pioneered by Cheng and co-workers,¹⁹ rhodium-catalysed C–H bond activation followed by annulation has been used as a powerful tool for the synthesis of disubstituted quinolizinium compounds. We sought to compare the newly developed approach of gold-catalysed cis-difunctionalization with rhodium catalysis. An optimized reaction employing 2-phenylquinoline (0.1 mmol, 1 equiv.) with internal alkyne 1,2-diphenylethyne $2w$ (0.1 mmol, 1 equiv.), $[Cp*RhCl₂]$ ₂ (5 mol%) and AgBF4 (0.1 mmol, 1 equiv.) in 1,2-dichloroethane under open air at room temperature for 16 h afforded diphenyl-substituted quinolizinium 7c in 68% yield. However, when 1-phenyl-2 trimethylsilylacetylene 2d was employed as a substrate, no desired product 4d was formed. To our knowledge, the present photosensitizer-free visible light-mediated gold-catalysed Edge Article. Common Access Article on Operation Common Comm

^a Absorption and emission properties were measured in CH₂Cl₂ at a concentration of 1×10^{-5} M. ^b Quantum yields were measured using fluorescein ($\Phi_F = 0.95$ in 0.1 N NaOH buffer) as a standard.

Fig. 5 HOMO and LUMO diagrams of 4d.

Table 4 Photooxidative amidation using quinolizinium compounds as $photocatalysts^a$

Entry	Photo cat.	$E_{\rm red}$ * (V)	Time (h)	Yield ^b [%]
1	4a	1.99	16	53
2	4b	1.97	16	50
3	4e	2.21	16	61
4	4f	2.22	16	59
5	4j	2.24	16	52
6	4p	2.07	16	53
7	Eosin Y	1.23 ^c	16	48
8	Fluorescein	1.25 ^c	16	47
9	Rose Bengal	1.18 ^c	16	32
10	$[Acr^+-Mes](BF_4)$	2.08 ^c	16	44
11	4e	2.21	48	86
12^d	4e	2.21	48	71 ^e

 a Reaction conditions: treatment of 8a (0.1 mmol, 1 equiv.), 9a (0.2 mmol, 2 equiv.), $Na₂CO₃$ (0.2 mmol, 2 equiv.) and photocatalyst (5 mol%) in 5 mL of CH₃CN under blue LEDs and air at room temperature. $\frac{b}{ }$ Yield was determined by ¹H-NMR using 1,3,5temperature. The was determined by R_{real}^* refers to ref. 9a trimethoxybenzene as the internal standard. c_{Head}^* refers to ref. 9a and literature cited there. d Reaction was performed by treatment of $(1 \text{ mmol}, 1 \text{ equiv.}), 9a (2 \text{ mmol}, 2 \text{ equiv.}), Na₂CO₃ (2 \text{ mmol}, 2 \text{$ equiv.) and photocatalyst (5 mol%) in 5 mL of $CH₃CN$ under blue LEDs and air at room temperature. ϵ Isolated yield.

alkyne cis-difunctionalization is the first example of a highly regioselective synthesis of silyl-substituted quinolizinium compounds (Scheme 3).

X-ray crystal structure analysis revealed a twisted conformation of quinolizinium 4b, with a torsion angle of 107.84 (C1–C2–C18–C19), suggesting that the molecule could be divided into two parts: a slightly distorted planar quinolizinium moiety and a phenyl moiety. This twisted structure suggested that the two moieties should be weakly coupled conjugatively (Fig. 4).

After synthesis of the new silyl-substituted quinolizinium compounds, we moved on to study their absorption and emission properties (Table 3). Interestingly, these compounds possessed the lowest energy absorption maxima at λ_{abs} > 395 nm, full color tunable emission properties (λ_{em} = 450 to 640 nm) in the visible light region and large Stokes shifts (up to

Table 5 Photooxidative amidation of aldehydes with secondary amines a,b

 a Reaction conditions: treatment of 8a-d (0.1 mmol, 1 equiv.), 9b d (0.2 mmol, 2 equiv.), Na₂CO₃ (0.2 mmol, 2 equiv.) and photocatalyst 4e (5 mol%) in 5 mL of CH3CN under blue LEDs and air at room temperature for 48 h. b Isolated yield.</sup>

 6797 cm^{-1}) with quantum yields up to 0.59. Incorporation of electron-donating groups at the para-position of the phenyl moiety or introduction of π -excessive heteroaromatics resulted in bathochromic shifts of the emission properties. Further bathochromic shifts were observed by the introduction of an electron-withdrawing ester group on the quinolizinium moiety. Substituents bearing heavy atoms (Cl, Br, I) led to lower quantum yields. A positive solvatochromism was found in compound 4b, which indicated an increase of dipole moment of the excited state compared to its ground state (Fig. S8 in the ESI†). An unexpectedly low quantum yield of 5a (0.02) bearing a dimethylamine substituent was observed, in which the fluorescence was proposed to be quenched by the amine group via intramolecular photo-induced electron transfer (PET). Cyclic voltammetry (CV) experiments indicated a quasireversible oxidation couple at $+1.08$ V (vs. SCE) of 5a which originated from the presence of the amine group and no similar peak was found in 5c (ESI†). Protonation of the amine group by measuring the emission in HCl/NaOH buffer (pH changing from 7 to 1) gave a \geq 100 fold enhancement of the emission intensity at a shorter wavelength (λ_{em} = 436 nm) which supported our hypothesis (Fig. S9 in the ESI†).

For a detailed investigation of the SPPR of the quinolizinium compounds, compound 4d was subjected to TDDFT

Fig. 6 Confocal fluorescence microscopic images of HeLa cells. (a) Sub-cellular localization of 5c; (b) subcellular localization of MitoTracker® red; (c) merged images of (a) and (b); (d) subcellular localization of 4l; (e) subcellular localization of LysoTracker® deep red; (f) merged images of (d) and (e); (g) subcellular localization of 4l; (h) subcellular localization of mRFP-Rab7; (i) merged images of (g) and (h).

calculations. Results reveal that the lowest energy absorption band at 435 nm is originated from HOMO \rightarrow LUMO transitions. The HOMO is composed of a π orbital of the quinolizinium and phenyl ring whereas the LUMO is composed of a π^* orbital of the quinolizinium ring. The low energy absorption band can be assigned as an admixture of $\pi \to \pi^*$ transitions within the quinolizinium ring and $\pi \to \pi^*$ transitions from the phenyl to quinolizinium ring. Hence, bathochromic shifts were observed by the introduction of an ester group at C9 (LUMO was dominant) and the introduction of electron-donating groups at C21 (HOMO was partially dominant) (Fig. 5).

In line with our interest in amide synthesis,²⁰ we envisioned that the newly developed quinolizinium fluorophores could be utilized as photocatalysts for photooxidative amidation of aldehydes and secondary amines. We first investigated the catalytic activities by treatment of 4-nitrobenzaldehyde 8a (0.1 mmol, 1 equiv.), piperidine 9a (0.2 mmol, 2 equiv.), $Na₂CO₃$ (0.2 mmol, 2 equiv.) and selected photocatalysts (5 mol%) in $CH₃CN$ under irradiation (blue LEDs) and air for 16 h. Interestingly, the catalytic activities of quinoliziniums were comparable or even better than the currently used photocatalysts (Table 4). The reaction was optimized using 4e as a photocatalyst to afford amide 10a in 86% yield after 48 h. Expansion of the scope employing different aldehydes and secondary amines gave amides 10b–g in up to 84% yield (Table 5). Further investigations revealed that the quinolizinium compounds possessed tunable and high excited state reduction potentials $(E_{\text{red}}^* = 1.97 \text{ to } 2.23 \text{ V})$, which were comparable to the wellknown photocatalyst 9-mesityl-10-methylacridinium [Acr⁺-Mes $[BF₄]$ developed by Fukuzumi.²¹ To our knowledge, this is

the first example of employing quinoliziniums as photocatalysts for visible light-mediated photoredox catalysis.

The feasibility in cellular imaging was demonstrated by incubation of HeLa cells with $2 \mu M$ of the fluorescent quinolizinium compounds $4a-b$, d , f , h , $l-o$, $s-t$, w and $5c$, respectively (Experimental details in the $ESI⁺$). Confocal fluorescence microscopic images revealed that the quinoliziniums were selectively localized in cytoplasm with no transportation into the nucleus, which was different from the known examples of quinoliziniums.²² Compound 5c and 4l were chosen for colocalization studies to track the subcellular localization. Compound 5c was specifically localized in the mitochondria (Fig. 6a–c), which could be attributed to the presence of the positively charged quinolizinium skeleton, directing it towards the mitochondria membrane with -180 mV potential. Interestingly, apart from localization in the mitochondria, compound 4l bearing a nitro substituent localized mainly in the lysosome and partially in late the endosome (Fig. 6d–i). The subcellular localization of the quinoliziniums could be switched by simply modifying the substituents, and these compounds would be amenable for the design of specific molecular probes for individual organelle imaging.

Conclusions

In summary, we have developed the first photosensitizer-free visible light-mediated gold-catalysed cis-difunctionalization reaction with high chemoselectivity and excellent regioselectivity for modular synthesis of a series of silyl-substituted quinolizinium compounds. Control experiments as well as a combination of NMR, ESI-MS and spectroscopic analysis indicate a plausible visible light-mediated $Au(1)/Au(m)$ catalytic transformation involving a regioselective syn insertion of silylsubstituted alkynes. Additionally, we have studied applications of the newly synthesized silyl-substituted quinolizinium compounds in photooxidative amidation and cellular imaging. The efficient modular synthesis and unique photophysical properties of the quinolizinium compounds will open up a new direction in gold catalysis, photoredox catalysis and molecular imaging.

Conflicts of interest

M.-K. Wong, J.-R. Deng and N. C.-H. Lai applied patents on quinolizinium compounds 4a–q, s–u, w and 5a–c.

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