Nitrene transfer reactions represent one of the key reactions to rapidly construct new carbon-nitrogen bonds and typically require transition metal catalysts to control the reactivity of the pivotal nitrene intermediate. Herein, we report on the application of imиноiodinanes in amination reactions under visible light photochemical conditions. While a triplet nitrene can be accessed under catalyst-free conditions, the use of a suitable photosensitizer allows the access of a nitrene radical anion. Computational and mechanistic studies rationalize the access and reactivity of triplet nitrene and nitrene radical anion and allow the direct comparison of both amination reagents. We conclude with applications of both reagents in organic synthesis and showcase their reactivity in the reaction with olefins, which underline their markedly distinct reactivity. Both reagents can be accessed under mild reaction conditions at room temperature without the necessity to exclude moisture or air, which renders these metal-free, photochemical amination reactions highly practical.
Photochemistry is a classic discipline in chemistry and early reports date back even to the beginning of the 20th century forecasting the potential of photochemical synthesis on the challenges of modern societies. It features light as the main source of energy to conduct chemical transformations and can thus be regarded as a key pillar in the development of sustainable approaches and reduced ecological footprint. Today, photochemistry is one of the most rapidly developing fields in chemistry and the past decades have witnessed the development of major milestones and concepts, such as dye-sensitized solar cells, polymerization chemistry, protein labeling, photoredox catalysts, and classic photochemical applications that leverage photochemistry as one of the key technologies in the advancement of all chemical disciplines.

In organic synthesis, photochemistry constitutes an important strategy to access reactive intermediates via excited state chemistry of catalysts or reagents. Significant advances have been made in the context of organic synthesis methodology, for example in the utilization of radicals in the presence of photoredox catalysts, and carbenes via the photolysis reaction of appropriate reagents. The access and application of nitrenes or analogs therof under visible light photochemical conditions however remains largely underestimated although it would allow for direct amination reactions with vast potential in modern drug synthesis. Such photochemical nitrene transfer reactions would significantly expand currently available concepts in classic metal-catalyzed nitrene transfer reactions and provide pathways under mild conditions to the pivotal nitrene intermediate. Today, essentially two strategies have been developed to access nitrenes under photochemical conditions. One concept harnesses the use of metal-catalyzed nitrene transfer reactions under UV light photochemical conditions to facilitate the formation of metal-nitrene intermediates. Another concept employs highly specialized, tailor-made nitrene transfer reagents, relying on strong UV-light for their photolysis. UV-light can however significantly reduce reaction efficiency and increase by-product formation. Yet, UV light still remains a prerequisite to access pivotal (metal-)nitrene intermediate under photochemical conditions. The use of visible light is thus in high demand to overcome these limitations, to enable sustainable nitrene transfer reactions and to develop strategies for amination reactions (Fig. 2).

In this work, we show that iminoiodinanes are suitable, bench-stable reagents that can be used to access either a triplet nitrene or a nitrene radical anion intermediate under photochemical or photocatalytic conditions, respectively. We showcase the reactivity of these monovalent nitrogen-based reactive intermediates in the reaction with olefins that can lead either to aziridination or C-H amination reactions. Experimental and computational studies are discussed to rationalize for the observed experimental data.

Results

Reaction optimization. The utilization of simple, readily available nitrene transfer reagents can be regarded as a prerequisite for the realization of photochemical amination reactions. We therefore decided to initially study the photochemical properties of iodinane and phenyl azide, which are commonly applied reagents in metal-catalyzed nitrene transfer reactions. While phenyl azide did not show absorbance in the visible light region, a range of different, readily available iminoiodinanes revealed a weak absorbance in the visible light region as determined by both experiment and theory (Fig. 2a and Supplementary Fig. 1). An important observation was made for the electron distribution of the HOMO and LUMO. While the HOMO of 1a is predominantly associated to electron density at the nitrogen atom (Fig. 2a), the electron distribution in the LUMO is shifted to the iodine atom (Fig. 2a and Supplementary Fig. 6), which could facilitate photolysis reactions and formation of nitrene intermediates.

Based on this observation, we investigated the reaction of α-methyl styrene with iminoiodinane under photochemical conditions (Table 1). Indeed, a very efficient amination reaction was observed in the presence of blue light (470 nm) that lead to the selective formation of the C-H functionalization product without accompanying by-products from aziridination. Other light sources proved by far less efficient, which might be related to weaker absorption or side-reactions due to the high-energy UV light. A surprising observation was made when switching to photocatalytic reaction conditions. Using simple [Ru(bpy)_3]Cl_2 as photocatalyst, a complete reversal of reactivity was observed and selective aziridination to yield 11a occurred. Other photocatalysts, such as [Ru(bpy)_2](PF_6)_2, iridium-based photocatalysts, or organic dyes did not alter the reactivity and the C-H functionalization product was formed selectively. Only in the case of Eosin Y as photocatalyst, a mixture of C-H amination (10a) and aziridination (11a) was observed. Further optimization steps included investigations on the solvent, concentration, reaction stoichiometry, yet no further improvements were observed (Supplementary Table 1).

Comparative study on the reaction mechanism. To further explore this divergent reactivity, we examined control experiments to provide an understanding of the underlying reaction
mechanism. No notable association of α-methyl styrene 12a or phenyl iodide with iminoiodinane 1a were observed by 1H-NMR studies, suggesting no association of reagents. Similarly, no notable decomposition of phenyl iodide under 470 nm blue light irradiation was observed within 4 h, which suggests that iodine species do not participate as hidden catalysts in the reaction.

Further studies on the reaction mechanism involved theoretical calculations to better rationalize this photochemical nitrene transfer reaction (Fig. 3). We therefore conducted calculations using (TD)-DFT methods to rationalize the photochemical reaction of iminoiodinane 1a and DFT methods for the catalytic reaction involving photosensitizers. These calculations reveal that photochemical excitation of 1a leads to formation of an excited state 1a* that remains in singlet state. This excited state can now undergo two different pathways: A) relaxation on the singlet spin surface leads to the direct formation of a singlet nitrene that features a very short N-O distance, which can be interpreted as a stabilization of the low-valent nitrene with the lone pair of an oxygen atom of the pendant sulfonyl group. B) relaxation involving inter system crossing (ISC) results in the formation of a triplet intermediate 1a-T, which features a very long N-I bond, close to a non-bonding situation. Scanning of different N-I bond lengths indicated that further elongation of the N-I bond proceeds in a barrier-free fashion to directly lead to a triplet nitrene intermediate. This triplet nitrene intermediate is the energetically favored intermediate and can alternatively be accessed from the high-lying singlet nitrene via inter system crossing. This theoretical study, and photochemistry of iminoiodinane 1a now rationalizes for the formation of a triplet nitrene intermediate under photochemical conditions and shows a marked difference to the ground state reactivity. Under photocatalytic conditions, calculations are in line with our previous report and a very facile reduction of the iodinane, leading to iodinane radical anion INT1. This anion features a very long N-I bond that is close to non-existent and therefore a rapid cleavage of the N-I bond can occur in a barrierless process to give a nitrene radical anion.

For further analysis and understanding of the reactivity of triplet nitrene and nitrene radical anion, we examined their structural and electronic properties. The calculations show that the nitrene radical anion possesses a higher electron density at the nitrogen atom compared to the triplet nitrene, which in turn leads to higher nucleophilicity of the nitrene radical anion (Supplementary Figs. 4, 7, and 8). For stabilization of the negative charge, the sulfonyl group plays an important role as an electron acceptor leading to mesomeric stabilization of the nitrene radical anion. The aromatic ring plays only a minor role in stabilization of the nitrene radical anion. The analysis of bond lengths further supports this mesomeric effect within the nitrene radical anion.

Despite of the additional negative charge, the S-N bond shortens upon formal reduction from triplet nitrene to nitrene radical anion from 1.71 Å to 1.56 Å and thus resembles more an S-N double bond in the case of the nitrene radical anion, which is also reflected by bond order analysis (Supplementary Fig. 5).

Further exploration concerned the evaluation of the reaction mechanism and an understanding of the distinct reactivity of triplet nitrene and nitrene radical anion by theory and experiment (Fig. 4). In the case of the triplet nitrene, we examined two different reaction pathways using DFT calculations (Fig. 4a, right). A first pathway involves hydrogen atom transfer from α-methyl styrene to the triplet nitrene intermediate, yet an activation free energy of 14.7 kcal mol⁻¹ and the formation of two separate radicals that need intermolecular radical recombination renders this pathway not feasible (Supplementary Fig. 11, TS6). A second pathway involves the addition of the triplet nitrene to α-methyl styrene via a TS3 with an activation free energy of only 6.1 kcal mol⁻¹, which is significantly favored over the hydrogen transfer pathway. This addition product can undergo intersystem crossing to give an open shell singlet species that can either cyclize to give the aziridine 11a, or undergo hydrogen atom transfer to give the C-H functionalization product 10a. Analysis of the respective transition states reveals significant steric hindrance of rotation around the central C-C bond, which renders cyclization energetically unfavorable over hydrogen atom
Table 1: Investigations on the photochemical and photocatalytic reaction iminoiodinane 1a with \( \alpha \)-methyl styrene 12a.

| Entry | Light Source | % Yield (1a) | % Yield (1b) |
|-------|--------------|--------------|--------------|
| 1     | 385 nm       | 22           | 40           |
| 2     | 470 nm       | 70           | 90           |
| 3     | 530 nm       | 40           | 58           |
| 4     | White light  | 29           | 45           |
| 5     | In the dark  | –            | n.r.         |
| 6     | 470 nm –     | 35           | 51           |
| 7     | Ru(bpz)\(_3\)(PF\(_6\))\(_2\) | 35           | 51           |
| 8     | Ir(ppy)\(_3\) | 27           | n.r.         |
| 9     | (Ir\((dF(CF\(_3\))ppy\)](dtbpy)PF\(_6\)) | 30           | –            |
| 10    | 4-CzlPN      | 51           | –            |
| 11    | Eosin Y      | 34           | 45           |
| 12    | In the dark  | –            | n.r.         |

Reaction conditions: 12a (1 mmol, 5 equiv.), 1a (0.2 mmol) and the respective photocatalyst (1 mol%) were dissolved in 2.0 mL DCM under air atmosphere. The mixture was irradiated with the light source (3 W) indicated for 4 h at room temperature.

Application in synthesis. We next turned our attention towards applications of the above protocols in amination reactions (Fig. 5). The photochemical C-H functionalization reaction of \( \alpha \)-methyl styrenes 12 proceeded smoothly and the allylamines products 10a-q were obtained in good isolated yield, without formation of by-products from aziridination (Fig. 5a). Halogens, electron-withdrawing or -donating groups in all positions of the aromatic ring were compatible and the corresponding products were obtained in high isolated yields (10a-o). Importantly, ortho-substitution (10m, n) had a slightly detrimental effect on the product yield. Similarly, different sulfonyl groups were tolerated under the present photochemical reaction conditions and the allyl amines products were obtained in high isolated yield (10p, q). When applying this photochemical protocol to cyclic, trisubstituted, cyclic olefins, 1-alkynyl-1-methyl or aliphatic 1,2-disubstituted olefins, the C-H amination products 10r-v were selectively obtained in good isolated yield. Then, the photocatalytic protocol
was employed in the aziridination of α-methyl styrenes (Fig. 5b).

We examined a similar range of substituents, including halogen, alkyl, cyano, ether or alkoxy substituents, at both the sulfonyl group and the benzene ring of the α-methyl styrene, which proved compatible under photocatalytic conditions. Only in the case of ortho-substitution slightly reduced reaction yields were obtained (11i–k). Most notably, in this case also different α-alkyl-substituted styrene derivatives, such as a very bulky tert-butyl group (11s), and nucleophilic N-heterocycles (11p, q) were well tolerated to yield the corresponding aziridines in high yield, which are commonly very challenging substrates in aziridination reactions. Furthermore, a range of trisubstituted, cyclic olefins, and further examples of 1,1-disubstituted olefins were studied under the present photocatalytic conditions to yield the aziridine products 11l–11ad.

Finally, we embarked on the reaction of simple styrene derivatives under both photochemical and photocatalytic conditions (Fig. 6). In both cases, aziridination reaction was observed in high yields, with the photocatalytic reaction being slightly superior over the photochemical reaction. Different sulfonyl groups as well as a range of electronically and sterically distinct substituents at the aromatic ring of the styrene component proved compatible (22a–t). Most notably, even in the presence of two sterically demanding ortho-chloro substituents (22t) the desired aziridine product was obtained in good isolated yield. Further studies involved the reaction of cyclic olefins, such as indene, which smoothly reacted to the aziridine product (22u) in high isolated yield. An important difference in the reactivity of nitrene and nitrene radical anion was however observed in the reaction with aliphatic olefins (Fig. 6b). While only a poor yield was obtained under photochemical conditions, the photocatalysis approach via the nitrene radical anion proved superior and high yield in the aziridination reaction of aliphatic olefins was observed (23a–i).

Further examples under investigation concerned the utilization of 1,2-disubstituted olefins. In this case, different aryl-alkyl and alkyl-alkyl disubstituted olefins smoothly underwent selective aziridination reaction to afford the trans-aziridines 20a–i in high yield.

The aziridination was then examined by DFT calculations, which suggest a similar pathways as in the aziridination reaction of α-methyl styrene for both photochemical and catalytic conditions (cf. Fig. 3 and Supplementary Figs. 12, 13). However, a distinct difference in the conformational flexibility of the diradical intermediate INT3 (cf. Fig. 4a, right) was observed. The absence of the α-methyl group renders the cyclization step to the aziridine much more favorable and it can proceed with a very low activation free energy of only 1.4 kcal mol$^{-1}$. The direct comparison of the respective transition states show that the α-methyl group results in a conformational twist and that prevents efficient aziridination as it can be seen in the torsional angle around the N-C-C-CAr bond (Supplementary Fig. 11).

In summary, we herein report on amination reactions of olefins with iminoiodinane reagents under purely photochemical and photocatalytic reaction conditions. Depending on the reaction conditions, these amination reactions proceed via chemically distinct monovalent, nitrogen-based reactive intermediates that in turn lead to different reaction products in the reaction with α-methyl styrenes. While triplet nitrenes can be accessed via direct photolysis of iminoiodinanes and lead to C-H functionalization, a nitrene radical anion is formed under photocatalytic conditions and leads to aziridination reaction. We studied the formation, properties, and reactivity of triplet nitrene and nitrene radical anion using computational calculations to rationalize the observed reactivity in the reaction with olefins, which was further validated in control experiments. We conclude with studies on the application of these amination reactions in the reaction with a diverse set of olefins, ranging from α-substituted styrenes, styrenes, towards aliphatic olefins.

**Methods**

**General method for the photochemical reactions.** In an oven-dried tube (10 mL), equipped with a magnetic stirring bar, iminoiodinane (0.2 mmol, 1.0 equiv) and alkene (5.0 equiv) are dissolved in 2 mL DCM under air atmosphere. The reaction is stirred and irradiated with a 3 W LED lamp (3 cm distance) for 4 h. A cooling fan is used to maintain room temperature (25–28 °C). The product was obtained after column chromatography using n-hexane/EtOAc as eluent.
Fig. 4 Control experiments and theoretical calculations on the formation of the nitrene intermediate; calculations were performed at the SMD(DCM)-
(U)M06-2X-D3/def2-TZVP/(U)M06-2X-D3/def2-SVP level of theory. 

a Energy surface of the reaction of triplet nitrene and nitrene radical anion. Gibbs energies are given in kcal mol\(^{-1}\).

b Reaction with deuterated \(\alpha\)-methyl styrene.

c Reaction with \(\alpha\)-ethyl styrene.

d Photocatalytic reaction with \(\beta\)-methyl styrene.

e Photochemical reaction with \(\beta\)-methyl styrene.
**General method for the photocatalytic reactions.** In an oven-dried tube (10 mL), equipped with a magnetic stirring bar, iminoiodinane (0.2 mmol, 1.0 equiv), alkene (5.0 equiv), and catalyst (1 mol%) are dissolved in 2 mL DCM under air atmosphere. The reaction is stirred and irradiated with a 3 W LED lamp (5 cm distance) for 4 h. A cooling fan is used to maintain room temperature (25–28 °C). The product was obtained after column chromatography using n-hexane/EtOAc as eluent.

**Computational details.** All calculations were performed using the Gaussian 16 series of programs. All structures were optimized at the (U)M06-2X level of C-H functionalization (10) and aziridination (11).

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**Fig. 5 Scope of the nitrene transfer reactions in the photochemical C-H functionalization and photocatalytic aziridination reaction via nitrene radical anions.**

**a** Photochemical reaction with α-methyl styrenes. **b** Photocatalytic reaction with α-methyl styrenes.
theory in combination with D3 dispersion corrections\textsuperscript{34}, in which all atoms were described with the def2-SVP basis set\textsuperscript{35}. Analytical frequency calculations were carried out at the same level of theory in order to confirm each stationary point as either an intermediate (no imaginary frequencies) or a transition state (only one imaginary frequency). Key transition-state structures were confirmed to connect corresponding reactants and products by intrinsic reaction coordinate (IRC) calculations\textsuperscript{36,37}. The electronic energy was then refined using def2-TZVP basis set\textsuperscript{35} at the (U)M06-2X level on the optimized geometries in combination with D3

Fig. 6 Scope of the photochemical and photocatalytic nitrene transfer reactions with styrenes and aliphatic olefins. a Evaluation of styrene derivatives. b Evaluation of aliphatic olefins c evaluation of 1,2-disubstituted olefins.
dispersion corrections. Solvation energies in dichloromethane ($\varepsilon = 8.93$; for acetonicitrile: $\varepsilon = 35.688$) were evaluated by IEFPCM calculations with radii and non-electrostatic terms for SMD solvation model based on the optimized structures. Time-dependent (TD)-DFT calculations were carried out on the optimized structures of the PhINTs to obtain the absorption wavelength.

**Data availability**

The authors declare that the data supporting the findings of this study, including computational details, experimental details and compound characterization, are available within its Supplementary Information. All data available on request from the corresponding author.

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