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HIGHLIGHT

Recent Applications of Biphotonic Processes to Organic Synthesis

Jorge Castellanos-Soriano, a Jorge C. Herrera-Luna, a David Díaz Díaz, b,c,d M. Consuelo Jiméneza and Raúl Pérez-Ruiza*

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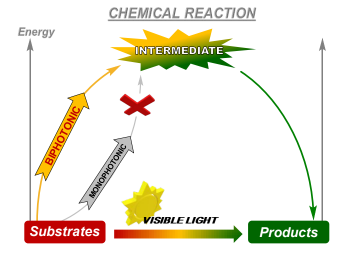
Currently, evolution of chemical transformations by visible light irradiation is highly desirable from cost, safety, availability, and environmental friendliness points of view. Besides, activation of less reactive substrates under very mild conditions becomes one of the most challenging tasks in organic synthesis. However, the insufficient energy provided by one photon of visible light for their activation definitely makes necessary the development of new protocols together with the design of new photocatalytic systems to overcome this limitation. In this context, the implementation of biphotonic processes has been found to be a solution of these drawbacks. This new mechanistic paradigm holds great potential for high-energy demanding bond activations, expanding the accesible reactivity window. Here, we wish to highlight the recent applications of biphotonic processes to organic synthesis.

Sunlight strikes our planet every day with more energy than we consume in a whole year. The question arises why the chemistry community has not used the sunlight as synthetic tool for developing photochemical processes during almost one century. ‡ The main reason lies in the inability of most organic compounds to absorb light in the visible range. Moreover, electronically excited states are often only available upon irradiation with shorter wavelengths of ultraviolet (UV) light. These high-energy photons can therefore cause uncontrolled photodecomposition processes, a factor that has limited the broad utility of photochemical synthesis in the fabrication of complex organic molecules. For that purpose, many researchers have explored ways to efficiently employ visible light energy for the activation of organic molecules;1 regarding, for example, the design, synthesis and development of photocatalysts that can absorb visible light and mediate the desired chemical transformations by rapid electron transfer processes. Activation of molecules with visible light offers the possibility of reaction pathways which are otherwise impossible to occur with classical nonphotochemical strategies.2

Another fascinating aspect of photochemistry boosted by visible light is the use of photons as "traceless and green reagents", rendering photochemical processes green and sustainable. In this context, the last decade has witnessed the fantastic growing in the field of Organic Photocatalysis using visible light as energy source, emerging as a new and powerful tool for activating molecules.³

However, the scope of photocatalytic bond activations is limited not only by the energy of one visible photon but also by the

energetic losses suffered by the photocatalysts such as intramolecular charge transfers or internal conversions.3h A plausible approach to expand the accessible reactivity window to less activated substrates is the use of photon-upconversion (UC),4 which converts low-energy into higher-energy radiation either by the two-photon absorption (TPA) or the triplet-triplet annihilation (TTA) mechanisms. 5 This technology has provided successful applications in diverse scientific fields ranging from energy to biology.⁶ Although the utilization of biphotonic processes as a potential synthetic tool is still in its infancy, it has gained great momentum over the past five years and has triggered a renewed interest toward photochemistry in general. Development of biphotonic technology for addressing critical bond activations or electron transfers in organic synthesis has been found to be advantageous while retaining the benefit of mild reaction conditions using lower-energy visible light (Scheme 1).



Scheme 1. Conceptual scheme of this Highlight article: application of biphotonic processes in organic synthesis.

^a Departamento de Química, Universitat Politècnica de València (UPV), Camino de Vera s/n, 46022, Valencia, Spain. Email: <u>raupreru@qim.upv.es</u>

b. Institut für Organische Chemie, Universität Regensburg, Universitätsstr. 31, 93053 Regensburg, Germany.

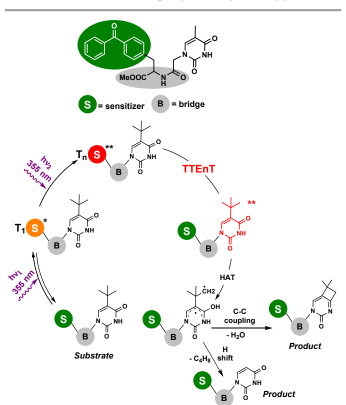
C Departamento de Química Orgánica, Universidad de La Laguna, Avda. Astrofísico Francisco Sánchez 3, 38206 La Laguna, Tenerife, Spain.

d Instituto de Bio-Orgánica Antonio González, Universidad de La Laguna, Avda. Astrofísico Francisco Sánchez 2, 38206 La Laguna, Tenerife, Spain.

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In this Highlight article, we therefore focus the attention on the existing examples of this emerging concept where several biphotonic processes such as **TPA**, **TTA** and consecutive photoinduced electron transfer (**ConPET**) have been successfully applied to chemical transformations.

Photosensitized Two-Photon Absorption (TPA) Reactions: The TPA mechanism implies an intramolecular simultaneous absorption of two photons process via a virtual state,7 and relaxation would lead to an emission with greater frequency than those of the absorbed photons. Despite **TPA** requires the use of high excitation intensity by lasers beams with concentrated peaks, numerous potential applications in biophotonics, optics and nano-micro fabrication have been reported.⁵ However, the implication of **TPA** processes in organic transformations is quite rare, presumably because of difficulties in experimental designs. Nevertheless, some examples can be found in literature focused mainly on the photosensitized biphotonic irradiation for DNA damage. Miranda and coworkers8 demonstrated the photochemistry of an upper thymine-like triplet states ($n\pi^*$ triplet). The proposed mechanism engaged the biphotonic excitation benzophenone by high energy laser pulses and a successful intramolecular sensitization. To avoid energetic losses, a dyad containing both the photosensitizer and the acceptor linked by an amide bridge was synthesized (Scheme 2). Thus, the benzophenone T_n formed by biphotonic excitation transferred its energy to thymine-like T_2 ($n\pi^*$), allowing the subsequent Norrish-Yang reaction to obtain the corresponding products. In this vein, the same research group has very recently published

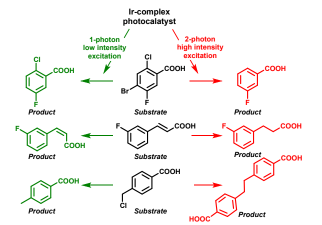


Scheme 2. Photoreaction of the benzophenone-modified nucleobase dyad. Involvement of a TPA process upon laser flash photolysis (λ_{exc} = 355 nm).

the intermolecular version of this reaction. Again benzophenone chromophore as photosensitizer and tert-butyluracil as modified pyrimidine base were utilized; however, the strategy was now different. After the first UVA (355 nm) photon absorbed by the sensitizer, triplet-triplet energy transfer (TTEnT) occurred, giving rise to the nucleobase triplet which was able to absorb the second photon (355 nm) reaching a T_n state. The viability of this strategy was supported by two model reactions: (i) the Norrish–Yang photocyclization of the tert-butyluracil and (ii) the photohydration of its uracil analogue, lacking the tert-butyl substituent.

Carbazole-based compounds were also found to be effective biphotonic sensitizers for the DNA photodamage. Description in the DNA photocleavage was successfully achieved in the presence of the photosensitizers upon irradiation with not only visible light but also with 800 nm NIR light via a **TPA** process. The proposed reaction mechanism involved a H atom abstraction (HAT) by N-centred radicals (type I mechanism) under anaerobic conditions. Accordingly, subsequent computational studies on these findings evidenced the great efficiency of these compounds as sensitizers in the NIR region; however, the rationalization of the mechanism of action revealed the involvement of solvated electrons ($e^-_{(aq)}$) produced by a spontaneous photoinonization that in turn were the responsible of DNA strand cleavage. DNA

Within this respect, Wegner and co-workers also observed that efficient two-photon excitation of a metal complex generated a $e^-_{(aq)}$ upon ionization. This fact allowed them to control the reactivity over several photochemical reactions relying on the light intensity. Hence, one-photon mechanism (low-powered intensity) permitted energy transfer from the photocatalyst to the substrates achieving debromination, cis-trans isomerization or HAT; on the contrary, employment of high-powered light intensity (TPA) electron transfer from the $e^-_{(aq)}$ took place, affording in this case dechlorinated, hydrogenated and dimer products, respectively (Scheme 3).



 $\textbf{Scheme 3.} \ \ \textbf{Control over the reactivity relying on one- or two-photon mechanism}.$

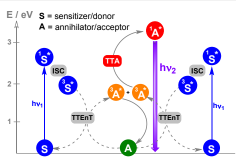
Regarding challenged dechlorination reactions, Yamaji and coworkers carried out two-colour two-laser photolysis on several chlorinated diketones.¹³ Results revealed the formation of halogen-free diketones as sole products by Cl elimination

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mechanism which the upper triplet $T_n\ (n\ge 2)$ state was involved. Finally, chemical reactions implying quinoline-based photoremovable protecting groups using **TPA** processes have been recently proven as a powerful tool for physiological studies. Thus, photolysis at longer wavelengths (740 nm light from a Ti:Sapphire laser) allowed to efficient release of homopiperonylic acid in high yield under simulated physiological conditions.

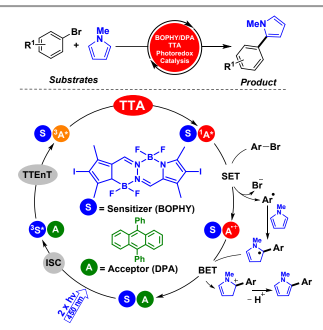
Photocatalytic Reactions by Triplet-Triplet Annihilation (TTA): Photon upconversion based on TTA between bimolecular organic systems has become one of the most attractive wavelength conversion technologies. Various combinations of compounds showing TTA properties can be found in literature. Scheme 4 depicts the photochemical events associated to this synchronized biphotonic process which includes intersystem crossing (ISC), TTENT, TTA and upconverted fluorescence (hv₂).



Scheme 4. Photon upconversion technology based on TTA. After the absorption of low-energy photons (hv₁), sensitizer triplet is produced by intersystem crossing (ISC) from its singlet. Subsequently, TTEnT from sensitizer triplets to the acceptor (Dexter mechanism) occurs. When two acceptor molecules in their triplet states are capable to collide during their lifetimes, a higher singlet energy level is formed by TTA and, consequently, generates delayed upconverted fluorescence (hv₂).

This transformation from two low-energetic photons ($2\times hv_1$) to one higher-energy photon (hv₂) has been already productively applied in diverse scientific areas of research;16 however, organic synthetic procedures using TTA processes has only recently been developed. For the first time, TTA was embedded into chemical reactions involving electron transfer activation of challenging organic molecules.¹⁷ Thus, we pioneered this elegant concept investigating the photophysical and photochemical aspects of combining TTA events of two simple organic dyes (diketone and oxazole) with reductive activations of aryl bromides. 17a This framework involved detailed spectroscopic, theoretical, and synthetic studies and ultimately led to the development of a new set of photocatalytic redox hydrodebrominations. Concomitantly with this work, we also contributed with the first intragel aerobic photoreduction of aryl halides catalysed by platinum(II) octaethyl-porphyrin (PtOEP) and 9,10-diphenylanthracene (DPA) as TTA system in supramolecular gel networks. 17b Thus, the gel network provided a stable microenvironment for the challenging multi-step process under aerobic conditions, room temperature and without additional additives. These results demonstrated that low weight molecular (LWM) gelators could be used as confined reaction media or micro/nanoreactors, providing the background for more demanding photophysical processes.

We further developed this methodology to more complex processes such as C–C coupling reactions. In particular, photocatalytic aromatic functionalization of N-methyl pyrrole was achieved by a metal-free TTA system based on diiodoBOPHY-like derivative (BOPHY) and DPA. ¹⁸ The reaction reported from-good-to-high product yields and the mechanistic aspects of this procedure were elucidated by means of combining product analysis, spectroscopic data and computational studies (Scheme 5). It is worth mentioning that an economic-friendly blue laser pointer (λ_{exc} = 445 nm ± 10, 2 W) was employed that facilitated the management of the optimal conditions instead of using the laser flash photolysis technique.



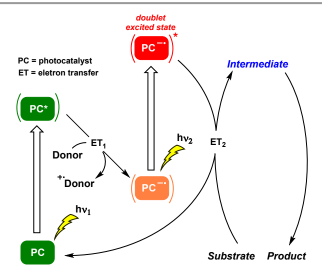
Scheme 5. Aromatic functionalization of N-methyl pyrrole photocatalyzed by **TTA** technology.

Among biological applications, near-infrared (NIR) radiation was also used for several organic transformations by utilizing the photophysical process of TTA. 19,20 Orange or blue light were accessed from low-energy NIR light (λ_{exc} = 730 nm) by pairwise combination of the TTA couple. 19 For instance, the delayed emission released from NIR-to-orange upconversion system (Pd-complex/diketopyrrolopyrrole derivative sensitizer/annihilator) was absorbed by a photocatalyst (eosin Y or Rose Bengal) that subsequent photocatalyzed lowdemanded energy reactions such as dehalogenation of α bromoacetophenone, amine oxidation or C-N bond activation. Besides, combination of a NIR absorbing sensitizer Pt-complex with tetratertbutylperylene as blue-emitter was adapted to the prototypical [Ru(bpy)₃]₂+-catalysed reaction: the intramolecular [2+2] cyclization of enones, obtaining moderate yields (48%). This TTA system was also employed directly for catalysing the cyclization of dienyl azides to pyrroles (80% yield) or the polymerization of methyl methacrylate. On the other hand, a photochromic reaction involving a C-N bond breaking was successfully achieved by TTA, where NIR light irradiation (λ_{exc} = 635 nm) was also implemented.²⁰

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Despite proper **TTA** systems in aqueous media are barely known,²¹ monodechlorination of trichloroacetate in airsaturated aqueous solution was successfully achieved by means of **TTA** technology.^{21d} This procedure constituted a promising green strategy for further photochemical transformations in homogeneous aqueous solution.

Consecutive Photoinduced Electron Transfer (ConPET) Reactions: Plants fabricate autotrophically their own energy from water, carbon dioxide and visible light. This combination, the renowned photosynthesis, follows a two-step photon excitation Z-scheme mechanism where accumulative photon absorption is involved. Surprisingly, the application of Z-scheme photosynthesis concept to activate substrates with high-energy demanding bonds for organic synthesis is very recent (Scheme 6).



Scheme 6. ConPET mechanism, that mimics the Z-scheme, adapted to organic transformations. Selective excitation of PC led to its excited state which is quenched by an appropriate donor generating the PC radical anion. Then, excitation of PC radical anion results in the formation of excited PC radical anion (doublet excited state), a highly reducing agent.

The kick-off work engaging these two terms is attributed to König and co-workers.²² Photogeneration of radical anions of aryl halides (Br, Cl) was achieved by a ConPET process involving a doublet excited state. An organic perylene diimide (PDI)-based photocatalyst was capable to accumulate the energy of two visible light photons in order to reduce stable aryl halides, generating the corresponding aryl radicals which were trapped by hydrogen atom donors or by specific nucleophiles to form C-C bonds. Thus, this outstanding chemical model mimicked the Z scheme in biological photosynthesis and permitted the photocatalytic conversion of less reactive chemical bonds in organic synthesis. Subsequently, the same PDI-based photocatalyst was incorporated into metal-organic polymer. This heterogeneous approach facilitated the ConPET process for the visible light-driven dehalogenation of aryl halides reducing markedly the irradiation times.²³ Regarding chromophorediimide derivatives, a heterogeneous strategy based on synergistic effects of ConPET and HAT processes was applied to aryl halides photoreduction. A new polyoxometalateincorporated naphthalenediimide (NDI)-based metal-organic framework (MOF) was designed and, after visible light irradiation, the host–guest MOF could stably generate NDI*-that was the critical intermediate for obtaining the photoreduced products. Strategically, diimide-like photocatalysts required *a priori* tedious synthesis, in addition to low solubility in organic solvents and sometimes employment of elevated temperatures. Moreover, alternative investigations related to the PDI system might suggest other mechanistic interpretation. ²⁵

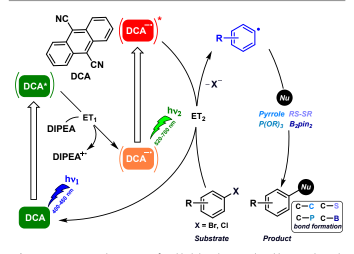
Various organic, inexpensive, and soluble dyes such as 1,8-dihydroxyanthraquinone (Aq-OH), rhodamine 6G (Rh-6G) or 9,10-dicyanoanthracene (DCA) were found to be suitable photocatalysts for **ConPET** processes. For instance, Aq-OH turned into its coloured radical anion or semiquinone anion in the presence of triethylamine upon one visible light photon irradiation. Then, excitation of these species by visible light led to an electron transfer process, activating aryl halides that reacted in dehalogenation or C–C bond-forming reactions.²⁶ To date, organic photocatalyst Rh-6G has been most widely used for organic synthesis.²⁷ The first work in this row described the selective aromatic functionalisation of arenes and

the selective aromatic functionalisation of arenes and heteroarenes by controlling the bond activation through lightcontrol regulation of redox potentials.^{27a} Subsequently, a series of synthetic applications photocatalyzed by Rh-6G through the ConPET process were reported. Thus, synthesis of pyrrolo-[1,2a]quinolines and ullazines was provided in one pot in moderate to good yields (35-75%).^{27b} Heteroaromatic biaryls were synthesized in moderate to excellent yields (41-91%) and, remarkably, this reaction worked for a broad range of brominated electron-rich heteroarenes and chlorinated heteroarenes bearing electron withdrawing groups.^{27c} A photo-Abzurov reaction for the construction of aryl phosphonates with consistent yields was also achieved by this methodology through the C-P bond formation.^{27d} Moreover, this Rh-6G-ConPET protocol was found to be an alternative strategy of the C-H arylation procedures using halogenated nucleobases under UV-light irradiation.^{27e} Combination of ConPET (Rh-6G) with metal catalysis (lanthanide ions) permitted the activation of even chlorobenzenes in order to form C-C and C-P bonds.27f Finally, it was demonstrated that gel networks could act as reaction media for aerobic photocatalyzed C-C coupling reactions by Rh-6G through ConPET mechanism, providing similar results than those obtained under inert atmosphere conditions.27g

Spectroscopic investigations gained some insights into the mechanistic aspects of this **ConPET** process using Rh-6G. Hence, fluorescence spectroscopy showed that Rh-6G radical anion could be generated from the singlet or triplet excited state of Rh-6G, relying on the sacrificial donor and varying their concentrations.^{27h} Very recently, the fate of the excited Rh-6G radical anion with femtosecond spectroscopy was studied.²⁷ⁱ The lifetime of the excited Rh-6G radical anion (350 fs) was found to be too short for diffusion-controlled electron transfer to the substrate. Therefore, authors postulated that the excited Rh-6G radical anion generated e⁻_(aq) as an additional step in the photocatalytic cycle, which was now the responsible of the aryl halide reductions.

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In this context, the excited DCA radical anion lifetime was previously ranged in the ns scale and it displayed an exceptionally high reducing potential of *ca.* –3 V.²⁸ Therefore, DCA possessed all prime qualifications for its use as a strongly reducing photocatalyst. Indeed, photocatalytic aromatic substitutions of poor reactive aryl halides by DCA were found to be operated via **ConPET** mechanism.²⁹ Thus the resultant excited DCA radical anion readily effected C–C, C–P, C–S, and C–B bond formations. Detailed information of the reaction mechanism of this biphotonic catalytic process was supported by product analysis, spectroscopic measurements, and computational studies (Scheme 7).



Scheme 7. Aromatic substitutions of aryl halides photocatalyzed by DCA through **ConPET** mechanism using cold white LEDs (410-700 nm). Fluorescence studies revealed quantitative formation of DCA radical anion that absorbs green light, giving rise to its doublet excited state; this species was capable to inject one electron to the aryl halides.

In summary, biphotonic technology has emerged as a powerful tool for addressing important bond activations or electron transfer processes in organic synthesis under mild conditions and using lower-energy visible light. These biphotonic processes include **TPA**, **TTA** and **ConPET** reactions, which have been used to trigger, for instance, reductive activations of aryl bromides, C–C or C–heteroatom coupling reactions, dehalogenation of aryl halides and functionalisation of arenes and heteroarenes. The success of this approach lies on a judicious selection of the photocatalytic system to achieve the required photoregulation of the involved redox potentials. In the coming years, we foresee a significant increase in the use of this technology to assist modern organic synthesis.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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Notes and references

‡ More than one century ago, the Italian chemist G. Ciamician published a *Science* ("The Photochemistry of the Future") where he recognized that sunlight could be utilized as a promoter of organic reactions (see G. Ciamician, *Science*, 1912, **36**, 385). Inspired by the ability of plants to make use of solar energy, he was the first scientist to investigate the photochemical reactions in a systematic way. His predictions on the advantages of utilizing solar energy to convert it into fuels has earned him to be considered a pioneer of modern photochemistry (In 1998, this paper was cited for the first time in a JRC journal: see for instance D. Gust, T. A. Moore, A. L. Moore. Mimicking bacterial photosynthesis. *Pure & Appl. Chem.* 1998, **70**, 2189. Source: SciFinder®).

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