Accepted Article

Title: Photochemical Methods for Peptide Macrocyclization

Authors: Laetitia Raynal, Nicholas C Rose, James R Donald, and Christopher Spicer

This manuscript has been accepted after peer review and appears as an Accepted Article online prior to editing, proofing, and formal publication of the final Version of Record (VoR). This work is currently citable by using the Digital Object Identifier (DOI) given below. The VoR will be published online in Early View as soon as possible and may be different to this Accepted Article as a result of editing. Readers should obtain the VoR from the journal website shown below when it is published to ensure accuracy of information. The authors are responsible for the content of this Accepted Article.

To be cited as: Chem. Eur. J. 10.1002/chem.202003779

Link to VoR: https://doi.org/10.1002/chem.202003779
Photochemical Methods for Peptide Macrocyclization

Laetitia Raynal*, Nicholas C. Rose*, James R. Donald*, Christopher D. Spicer*

*Authors contributed equally

Department of Chemistry, University of York, Heslington, York, YO10 5DD; York Biomedical Research Institute, University of York, Heslington, York, YO10 5DD.

*Correspondence addressed to: james.donald@york.ac.uk; chris.spicer@york.ac.uk

Abstract

Photochemical reactions have been the subject of renewed interest over the last two decades, leading to the development of many new diverse and powerful chemical transformations. More recently, these developments have been expanded to enable the photochemical macrocyclization of peptides and small proteins. These constructs benefit from increased stability, structural rigidity, and biological potency over their linear counterparts, providing opportunities for improved therapeutic agents. In this review, we provide an overview of both the established and emerging methods for photochemical peptide macrocyclization, highlighting both the limitations and opportunities for further innovation in the field.

1.1 Introduction

Peptides and small proteins (collectively, PSPs) represent an exciting and emerging frontier in drug discovery, combining the advantages of modularity and synthetic flexibility of small molecules (<500 Da) with the exquisite levels of selectivity and potency of biologics (>5000 Da).[1–3] PSPs are particularly prized for their capacity to engage in potent and selective interactions with protein surfaces, and hence their ability to modulate protein-protein interactions (PPIs) – therapeutic targets that were once considered “undruggable” using small molecules.[4,5] The peptidic nature of PSPs provides them with beneficial attributes such as high binding affinity, biocompatibility, and ease and modularity of synthesis, but is also their Achilles heel. Low cell permeability, poor metabolic stability, and unstable secondary structure formation as a result of conformational flexibility, all hinder clinical applications of PSPs.[6–9]

To overcome these limitations the design and synthesis of macrocyclic PSPs has become increasingly prominent.[1,4,8,10] Macrocyclization (defined as cyclization to form rings of ≥12 atoms) can occur through head-to-tail, head-to-sidechain, sidechain-to-tail or sidechain-to-sidechain coupling (stapling) (Figure 1), delivering cyclical products which possess a number of advantageous characteristics over their linear counterparts[11]: i) structures are rigidified, stabilising or enforcing peptide conformations that mimic elements of protein secondary structure.[12] α-Helices,[13–20] β-sheets,[21] and β-hairpin turns[22] which would otherwise be unstable can all be induced by cyclization; ii) cyclized peptides exhibit increased stability to proteolysis, thus prolonging their biological activity and improving pharmacokinetics. This stability
can result from a number of factors, including the poor fit of macrocycles into the active sites of endopeptidases,[23] resistance to the activity of exoproteases that preferentially cleave near peptide N- or C-termini,[11] or the formation of α-helices that are resistant to proteolysis due to the presence of a rigidifying, intramolecular, hydrogen-bonding network[17]; iii) binding efficiency for a target is often improved, an effect classically attributed to cyclic structures being held in conformations better disposed towards binding, with a resultant reduction in the entropic penalty to binding.[23,24] However, studies such as those by Martin and Spaller illustrate a much more complex picture, where pre-organization through macrocyclization may instead strengthen the enthalpic component of binding at the expense of entropy, highlighting the importance of treating macrocycle binding thermodynamics on a case by case basis,[25–28] and iv) cell membrane permeability may be improved, as side chains can be oriented around the macrocycle in a manner that shields polar atoms from the solvent medium, reducing the polar surface area of the peptide.[10]

Figure 1. Peptide macrocyclization modes and generalized layout of figures in this review.

Given the benefits of PSP macrocyclization, it is unsurprising that many such compounds occur naturally. A variety of macrocyclic linkages have been identified in natural products, including head-to-tail amide bonds, disulfide bridges, biaryls, and biaryl ethers.[29–33] Chemists have subsequently followed suit, utilizing analogous tactics to form cyclic PSPs, as well as developing a diverse toolkit of novel synthetic strategies for macrocyclization, including ring closing metathesis,[34] azide-alkyne cycloadditions,[35,36] other transition-metal catalyzed methods,[37] conjugate additions,[38] nucleophilic aromatic substitutions,[39] and multicomponent reactions[40,41]. These and other methodologies have been extensively reviewed.[3,18,19,23,37,42–47] Though the coupling chemistry of each of these strategies may differ greatly, common to all of these approaches are a set of challenges that must be overcome, chiefly coaxing the ground-state trans geometries of multiple amide bonds along the PSP backbone into a suitable conformation for cyclization,[48] and outcompeting deleterious oligomerization reactions.

In contrast to the methods outlined above, the use of photochemical methods for PSP cyclization had until recently been relatively underexplored. However, in line with a wider renaissance of the fields of photo- and radical chemistry, recent developments have brought this area to the fore. Photochemical strategies are highly
attractive for macrocyclization: reactive open-shell or photoexcited-state species are known to readily participate in macrocyclization via mechanisms that are fundamentally different to their two-electron ground-state counterparts, offering ample scope for novel cyclization manifolds that overcome some of the limitations of two-electron and transition metal catalysed methods.\(^{[49,50]}\) Given that most peptides are largely transparent to light of wavelengths >320 nm (and those without aromatic amino acids or disulfide bonds, >250 nm),\(^{[51]}\) photochemical methods allow targeted excitation of chromophores in a reaction mixture, affording the potential for both mild and highly selective processes to take place, that are compatible with complex biological systems.\(^{[52-54]}\) Furthermore, the precise control of photochemical processes afforded by manipulating the incident light, both spatially and temporally, offers much scope for applications of these emerging methods in biomedical settings. Herein, we therefore review the application of light-driven processes for PSP macrocyclization for the first time. It is our hope that through this summary of the field we will promote increased uptake of these powerful methodologies, and help stimulate further development in this exciting and rapidly evolving area.

2. Redox-neutral photochemical peptide macrocyclization

2.1 Introduction

Photochemical macrocyclization reactions can typically be grouped into two categories: i) redox-neutral processes where the photoexcited-state either directly takes part in the macrocyclization, or produces reactive species via atom transfer or bond rearrangement which then participate in cyclization; and ii) those where the photoexcited-state first engages in photoinduced electron transfer (PET), to provide radical, radical ion, or organometallic species that are necessary for cyclization. In this section, we will focus on the first class of reactions, where photochemical macrocyclization can be further categorized based on the specific behaviour of the photoexcited-state: hydrogen atom transfer (HAT); bond reorganization; or cycloaddition. Photoactivatable motifs are widespread in the bioconjugation field, however, in this review we will focus only on reactions with levels of chemoselectivity that allow their use for controlled cyclizations. Thus, widely used reactive handles in photoaffinity labelling, such as diazirines and aryl azides, which generate highly reactive species upon activation that can react in a promiscuous, largely non-selective manner, will not be discussed.

2.2 Macrocyclization triggered by photoinitiated hydrogen atom transfer (HAT)

2.2.1 Thiol-‘ene’ reactions

Thiols represent a versatile reactive handle for peptide and protein modification that can undergo a wide range of different chemistries. As a result, the modification of cysteine residues has been widely used for peptide macrocyclization via alkylation,\(^{[55,56]}\) arylation,\(^{[39]}\) and disulfide formation.\(^{[57]}\) Conjugate-addition reactions,\(^{[38,58]}\) also referred to as nucleophilic thiol-‘ene’ reactions, are particularly popular due to their chemoselectivity, mild reaction conditions, and rapid kinetics.\(^{[59]}\)
However, the need to use activated, electrophilic alkenes can lead to possible side reactions with other nucleophilic amino acids, particularly lysine.[60] The highly selective reaction of thiol radicals with unactivated alkenes, via a photo thiol-‘ene’ mechanism, is therefore an attractive alternative.[61]

Thiol radical formation (1, Figure 2) can be initiated through direct photolysis of the thiol with UV light. Although thyl radical generation is slow at wavelengths >280 nm (e.g. using a sunlamp with pyrex filter), this can be advantageous for chain processes where a low steady-state concentration of radicals helps suppress deleterious processes.[61] More often, a photoinitiator is employed and thyl radicals are generated via a rapid HAT between the thiol and either the photoexcited-state initiator directly, or daughter radicals stemming from its photo-decomposition (e.g. for HAT between •-CH₃ and EtSH, k₂₉₈K = ~5 x 10⁷ M⁻¹ s⁻¹).[62] Thiol radical 1 subsequently adds across an alkene C=C bond in an anti-Markovnikov manner, generating a carbon-centred radical 2. Abstraction of a hydrogen atom from another thiol generates the hydrothiolated product 3 and an additional thyl radical 1 facilitating chain propagation.

The steric and electronic nature of the alkene substituents play a significant role in governing the efficiency and outcome of this process. A fine balance exists between forward (and reverse) rates of thyl radical (1) addition to an alkene, S-H abstraction from another thiol by the resultant carbon-centred radical 2, and competitive addition of 2 to another alkene to afford off-cycle oligomerization/polymerization products.[63–65] In general, electron-rich alkenes typically react more rapidly, with terminal alkenes being similarly more reactive than internal analogues.[63] Norbornene and vinyl ether derivatives undergo exclusively hydrothiolation over oligo-/polymerization owing to particularly rapid thyl radical additions, as a consequence of strain release and polarity matching respectively, and fast S-H abstraction steps. Though the competitive formation of disulfide bonds is typically slow, recent reports by the Bowman group have highlighted the potential role of thiolate anions in accelerating this side reaction, through the formation of a metastable disulfide radical anion.[66] However, although this may be problematic in the context of thiol-‘ene’ polymerizations, the high effective concentration of thyl radical and alkene during intramolecular macrocyclization likely negates this effect.

Figure 2. Mechanism of the photo-activated thiol-‘ene’ reaction. A photoinitiator can be used to accelerate the formation of a thyl radical via S-H abstraction.
In the earliest example of photochemical thiol–ene' peptide macrocyclization, Aimetti et al. demonstrated the on-resin synthesis of an integrin-binding cyclic Arg-Gly-Asp (RGD) peptide.\textsuperscript{[67]} Irradiation of resin-bound peptides 4, containing unprotected cysteine residues, at 365 nm in dimethylformamide (DMF) led to cyclization with either allyloxycarbonyl (alloc) (4a) or amido-norbornene (4b) modified lysine residues in the presence of the photoinitiator 2,2-dimethoxy-2-phenylacetophenone (DMPA). As expected, given the reactivity profiles of different alkenes discussed above, higher yields and shorter reaction times were reported for 4b than 4a (37\% vs 24\% respectively). Notably, photoactivation was shown to be both quicker and more effective than the use of the thermally activated radical initiator azobisisobutyronitrile (AIBN). Since all other amino acids in the peptide sequence were protected, potential side reactions were minimized and the cyclized peptides 5a and 5b were efficiently obtained following cleavage. Macrocyclization of the solution-phase, fully deprotected peptide was also performed in methanol. Though conversions were comparable, the need for subsequent peptide purification led to overall lower yields.

![Diagram showing the synthesis and cyclization process]

Figure 3. First example of peptide macrocyclization via a photoactivated thiol–ene' reaction. On-resin cyclization onto both alloc and norbornene alkene partners was achieved in the presence of a DMPA photoinitiator.\textsuperscript{[67]}

Building upon this work, the same authors demonstrated that peptides could undergo further extension following on-resin thiol–ene' macrocyclization.\textsuperscript{[68]} This allowed the installation of a second reactive cysteine for subsequent conjugation of a cyclic-RGD motif to a multivalent polymer backbone. Importantly, a combination of cyclic and multivalent peptide presentation was found to act synergistically to enhance inhibition of fibrinogen binding to glycoprotein IIb/IIIa, a key integrin found on the surface of platelets, by up to two orders of magnitude.
The use of unactivated, aliphatic alkenes for on-resin photo thiol-‘ene’ macrocyclization was subsequently reported by Zhao et al.\[^{69}\]\ Peptides containing unnatural amino acids with but-4-ene, pent-5-ene, or hex-6-ene side chains (6a-c respectively) were cyclized with a cysteine residue at the i+4 position under 365 nm irradiation in DMF, with comparable yields (7a-c, 79-90%) (Figure 4). Of the radical initiators screened, 2-hydroxy-4’-(2-hydroxyethoxy)-2-methylpropiophenone (8, also known as IHT-PI 659) was found to give the highest conversion in the presence of 4-methoxyacetophenone (9), an additive that was observed to reduce by-product formation, facilitating product purification by HPLC.

---

**Figure 4.** On-resin macrocyclization of unactivated alkenes via a photoactivated thiol-‘ene’ reaction.\[^{69}\]

In 2015, Wang et al. reported the photo thiol-‘ene’ macrocyclization of peptides composed solely of natural amino acids via a novel solution-phase, two-component approach.\[^{70}\] A series of bifunctional dienes were used to link two cysteine residues under 365 nm irradiation, in the presence of a DMPA photoinitiator. Though the reaction was compatible with DMF, the highest yields were obtained in N-methyl-2-pyrrolidone (NMP). This two-component approach offers synthetic versatility, enabling the generation of a library of peptides cyclized with different diene linkers from a single peptide precursor. This is particularly important given the strong influence of cross-linker structure on peptide properties. For example, changes in lipophilicity have been shown to impact on the ability of cyclic peptides to cross phospholipid bilayers and therefore the potency of therapeutic peptides,\[^{71}\] while hydrogen bonding interactions have been shown to influence peptide conformation and target binding.\[^{72}\] As a result, the authors were able to develop bis-thioether cyclized peptide inhibitors of p53-HDM2 interactions that were able to induce the apoptosis of colorectal carcinoma cells with a comparable potency to previously reported hydrocarbon-linked peptides generated by ring-closing metathesis.\[^{73}\]
More recently, the same authors have developed this two-component approach further to enable the macrocyclization to be performed in water. To overcome challenges with aqueous solubility previously encountered by other groups, the water-soluble photo-initiator 2,2’-azobis[2-(2-imidazolin-2-yl)propane]-dihydrochloride (also known as VA-044) was employed to induce thiol-conjugation to water-soluble diallyl-urea (Figure 5). At pH 4 in aqueous solution, the reducing agent TCEP was found to greatly enhance the conversion of peptide to cyclized from 53% to 95% by minimising competing disulfide formation.

**Figure 5.** Aqueous photoactivated thiol-‘ene’ reaction, enabled by the use of the water-soluble initiator VA-044. 

Impressively, the macrocyclization of a 9 kDa, dicysteine, coiled-coil protein substrate was also demonstrated. Though the addition of TCEP was found to be detrimental in this scenario due to competitive desulfurization, in its absence the reaction was found to proceed effectively in mildly acidic (pH 4) acetate buffer containing denaturing guanidine hydrochloride (Gdn-HCl). This reduced disulfide formation and maximized cyclization efficiency respectively, leading to 90% conversion to stapled protein 14. Double cyclization of a tetra-cysteine mutant containing two separate i,i+7 cysteine pairs to generate, was also found to proceed with 80% conversion (Figure 6). Though not reported by the authors, in this scenario the presence of undesired linkages between the two separate cysteine pairs cannot be ruled out. However, it is likely that the spatial proximity of the i,i+7 residues would strongly favour cyclization between adjacent cysteines, even under denaturing conditions.
Figure 6. Aqueous di-cyclization of a tetra-cysteine protein mutant via photoinitiated thiol-'ene' chemistry.[74]

2.2.2 Thiol-'yne' reactions

In an analogous fashion to thiol-'ene' reactions, thiols can also react with alkynes via both nucleophilic and radical mechanisms. Photoinitiated thiol-'yne' reactions with unactivated alkynes therefore offer an alternative strategy for achieving PSP macrocyclization. Conjugation proceeds via a mechanism similar to that for thiol-'ene' reactions, comprising light-initiated formation of thiyl radical 17, addition to an alkyne to form 18 in an anti-Markovnikov manner, and subsequent hydrogen atom abstraction by vinyl radical 18 (Figure 7). However, the thiol-'yne' reaction differs in that it generates a vinyl sulfide product 19 that is itself susceptible to further modification in the presence of excess thiol, via a subsequent thiol-'ene' reaction. Though less widely used than the analogous 'ene' reaction, thiol-'yne' conjugations therefore offer intriguing possibilities for achieving double modification or dual functionality,[75] though these properties have yet to be exploited by the PSP macrocyclization community. Indeed, to date there has only been a single report of photoactivated thiol-'yne' mediated peptide macrocyclization.

Figure 7. Mechanism of the photo-activated thiol-'yne' reaction. A photoinitiator can be used to accelerate the formation of a thyl radical via S-H abstraction.

Tian et al. demonstrated the intramolecular photo thiol-'yne' cyclization of peptides containing an unnatural amino acid bearing a pent-5-ynyl side chain, and either cysteine (20a) or homocysteine (20b) at the i+4 position (Figure 8).[71] A screen of photoinitiators found IHT-PI 659 (8) and 2,2-dimethoxy-2-phenylacetophenone (DMPA, 21) to provide the highest conversions, with a strong preference for the formation of the Z-vinyl sulfide products 22a and 22b. The resultant peptides were designed to modulate intracellular interactions between the oestrogen receptor and its coactivators, a key target for the treatment of certain cancers and osteoporosis.
Importantly, the vinyl sulfide-containing macrocycle generated was proposed to provide enhanced rigidity, and to enhance $\alpha$-helical character over the alkyl sulfide linker that would be generated via an analogous thiol-‘ene’ cyclization. Furthermore, comparison to an all hydrocarbon linked analogue generated by ring-closing metathesis indicated that the vinyl sulfide contributed to greatly reduced membrane toxicity. Thus, this report highlights the importance of the cyclization linker structure in determining the properties, both physical and biological, of a cyclized peptide. Rather than being interchangeable with the thiol-‘ene’ reaction, photo thiol-‘yne’ reactions should be considered an important addition to the macrocyclization toolbox that can generate a distinct vinyl sulfide linkage that may possess unique properties.

**Figure 8.** Photo-activated thiol-‘yne’ macrocyclization to generate a vinyl sulfide-linked cyclic peptide.[71]

### 2.2.3 Benzophenone-methionine conjugation

The ability of certain ketones and aldehydes to form reactive species under UV irradiation has been widely exploited in organic synthesis and chemical biology. Benzophenones undergo photoexcitation on irradiation with longer wavelength UV light (350-360 nm),[76,77] and importantly, relative to other photoactivated groups such as diazirines and aryl azides, form intermediates that exhibit useful (albeit limited) levels of chemoselectivity. Mechanistically, the highly reactive diradical 23 generated following irradiation can abstract a hydrogen atom from an accessible C-H bond to generate an $\alpha$-hydroxy radical 24 and carbon-centred radical 25 (Figure 9). Pairing of these two species leads to the formation of a new carbon-carbon bond in tertiary alcohol 26.
In an attempt to understand and harness preferential reactivity, Deseke et al. studied the regio- and chemo-selective C-H abstraction of irradiated benzophenones with a panel of N-acetyl amino acid methyl esters.\cite{78} In acetonitrile, the highest reactivity was observed with glycine due to its readiness to undergo hydrogen atom abstraction from the $\alpha$-carbon (51\% conversion). Methionine was also found to be preferentially modified (40\% conversion), with alkylation occurring either at the $\gamma$- or $\epsilon$-carbon atoms adjacent to sulfur. Conversely, in a 4:1 mixture of pyridine:water methionine modification was found to be favoured (45\% conversion), with a corresponding drop in glycine modification (16\%). The authors proposed that this change in selectivity was due to competitive base-catalysed degradation of adducts formed at the $\alpha$-position of glycine. Though modification of these two amino acids was dominant, low levels of reactivity with several other amino acids was also observed. Indeed, only residues containing primary amides or carboxylic acids (aspartic acid, asparagine, glutamic acid and glutamine) were found to be inert.

Building on this preferential reactivity at glycine and methionine, Moretto et al. reported the first use of benzophenone photoactivation for intramolecular peptide cyclization in 2009.\cite{79,80} Hexapeptides 27 containing the unnatural benzophenone-based amino acid Bpa 28 and methionine were synthesized, and their positions in the peptide chain varied to study the effects of distance on macrocyclization efficiency (Figure 10). To prevent unwanted reactions at other sites in the peptide, the di-$\alpha$ substituted unnatural amino acid 2-aminoisobutyric acid (Aib, or $\alpha$-methylalanine) was installed at all other positions due to its known inertness to C-H abstraction, and ability to promote $\alpha$-helix formation. Cyclization was successful when methionine was placed at the i+1, i+2, i+3, or i+4 positions relative to Bpa, generating cyclized peptides 29. For an analogous nonapeptide 30, cyclization to the i-3 and i+3 positions was found to occur exclusively at the $\epsilon$-carbon of methionine, with the two diastereomers generated by the new chiral tertiary alcohol being formed in a $\sim$1:1 ratio.

**Figure 9.** Mechanism of the intermolecular reaction between a photoexcited benzophenone and an accessible C-H bond.
Figure 10. Macrocyclization via the conjugation of benzophenone to methionine under UV irradiation. By placing the unnatural amino acid Aib at all other positions, potential side reactions with α-hydrogens are prevented.[80]

Building upon this work, Wright et al. subsequently introduced a new Cα-tetrasubstituted, cyclic, benzophenone-based amino acid BpAib 31, with increased structural rigidity (Figure 11).[81] Though the photo-activated macrocyclization of this amino acid with methionine was demonstrated, efforts to exploit the benefits of decreased conformational freedom imparted by BpAib have yet to be reported.

Figure 11. Structures of the unnatural benzophenone-containing amino acids Bpa 28 and BpAib 31.

Lewandowska-Andralojc et al. have subsequently demonstrated that Bpa-methionine cyclization can take place between the two side chains of a cyclic dipeptide to form a rigid bridged macrocycle.[82] This study highlighted that the C-H abstraction step to form an α-hydroxy radical is in fact reversible. Furthermore, C-C bond formation was found to take place selectively at the methionine δ-carbon in order to minimize the significant ring strain imposed by cyclization. Thus, both steric and chemical factors are at play in dictating C-H selectivity.

The ability of benzophenones to cyclize with natural amino acids following irradiation is an advantage to their use. However, the lack of specificity resulting from off-target reactions with alternative amino acids, particularly glycine, is severely restricting. This is highlighted by the limited number of reports of benzophenone PSP macrocyclization, and the even more striking lack of diversity in the amino acids that have been integrated into the peptide substrates. Moreover, the formation of diastereomeric products, resulting from the creation of a new chiral center, may be
problematic, potentially requiring separation and complex purification to generate a homogeneous cyclized product. As such, the photo-activation of benzophenones as a means to control PSP macrocyclization is unlikely to find increasing prominence in the coming years, given the advantages of many of the other reactions presented in this review.

2.3 Photoactivation via bond reorganization

2.3.1 Photorearrangement of 7-nitroindoline amides

In 1999 Papageorgiou and co-workers proposed a mechanism for the solvolysis of 5-bromo-7-nitroindoline (Bni)-amides (32), and related 5-substituted derivatives, following photoactivation,[83,84] Irradiation of 32 with UV light generates a photoexcited-state 33, which undergoes N→O acyl transfer to generate the highly activated intermediate 34 (Figure 12). In water, 34 undergoes cleavage to generate 7-nitroso-indole 35 with the release of a free carboxylic acid. However, in the presence of a competing nucleophile or nucleophilic solvent, attack at the electrophilic carbonyl leads to cleavage of the Bni motif with the release of 7-nitroindoline 36.

The propensity to undergo hydrolysis under irradiation makes Bni-derivatives useful protecting groups, but also severely impacts their bioconjugation efficiency in aqueous media. However, in non-nucleophilic organic solvents, nucleophilic attack by amines can be used to induce amide formation following photoactivation. This chemistry has therefore been exploited for intramolecular photocyclization and in particular as a useful method for macrolactamization.[85] Indeed, compared to other macrolactamization strategies that require the use of added coupling agents, the irradiation of Bni-amides provides a facile means to induce cyclization without the need for additives.

The capacity of photoactivated Bni-amides to react with nucleophiles was exploited by Mifune et al. for PSP macrocyclization.[86] This approach is attractive as it
generates a native peptide bond and the resultant macrocycles are therefore able to mimic natural cyclic peptides. Indeed, the authors demonstrated that a synthetic pentapeptide 37, bearing a C-terminal Bni amide, could be cyclized in a 36% yield following irradiation at 365 nm under flow conditions, generating a previously reported cyclic RGD sequence 38 in situ that can act as a selective antagonist of the $\alpha_v\beta_3$ integrin receptor (Figure 13).\textsuperscript{[87–91]} The absence of excess activating agents or catalysts during cyclization greatly facilitated purification of the cyclized peptide, and the ability of the Bni-group to play a parallel role as a C-terminal protecting group during peptide synthesis was also highlighted by the authors as a notable advantage. However, the presence of a C-terminal Bni-amide also necessitated the use of solution-phase peptide synthesis. Though this was achieved using micro-flow peptide synthesis, such technologies are not as widespread as solid-phase approaches and the generality and translatability of this approach may therefore be limited at present.\textsuperscript{[92]} Post-synthesis derivatization is another possible route to achieve C-terminal Bni-amide installation for head-to-tail or sidechain-to-tail cyclizations but also presents synthetic challenges. In contrast, head-to-sidechain or sidechain-to-sidechain couplings would not experience these difficulties and the installation of a Bni-motif could plausibly be achieved in a straightforward manner on-resin. However, the need to also protect Asp, Lys, and Arg residues to prevent side reactions remains a significant limitation of this strategy, and as a result there remains only a single report of Bni-photoactivation in the peptide macrocyclization literature to date.

![Figure 13. Solution-phase head-to-tail macrocyclization of a peptide containing a C-terminal Bni-amide under UV irradiation.\textsuperscript{[86]}](image)

**2.3.2 Photorearrangement of tetrazoles**

Building on the pioneering work of Huisgen, in 2007 the Lin group reported the use of 2,5-diaryl tetrazoles as photoactivatable precursors for 1,3-dipolar cycloadditions.\textsuperscript{[93]} As these reactions are greatly accelerated in aqueous media, they have emerged as useful tools for photoactivated bioconjugation. Upon UV irradiation of tetrazole 39, expulsion of nitrogen gas generates a highly reactive nitrile imine 40 that can serve as a 1,3-dipole for cycloaddition with a suitable dipolarophile 41, most commonly an alkene, to afford a pyrazoline cycloadduct 42 (Figure 14). Cycloaddition proceeds with high regioselectivity, particularly with electron-deficient alkenes, generating 42 as a single regioisomer.\textsuperscript{[93]} When photolysis is rapid,
cycloaddition becomes rate-determining and reactions follow apparent second-order kinetics.

The chemical versatility of the 2,5-diaryl scaffold is attractive as it provides significant scope to tune the properties of the tetrazole and intermediate nitrile imine. For example, by variation of the aryl substituents X and Y, the wavelength sensitivity can be tuned within the UV region.\[^94\] Similarly, the rate of cycloaddition can be increased by selecting substituents that serve to raise the highest occupied molecular orbital (HOMO) energy of nitrile imine \[^40\].\[^95\] Monoaryl tetrazoles are also able to eliminate nitrogen under irradiation and can undergo efficient cycloaddition, albeit at a significantly reduced rate.\[^96\]

\[ \text{Figure 14. Mechanism of 2,5-diaryl tetrazole photoactivation to form a nitrile imine, and subsequent 1,3-dipolar cycloaddition with a suitable dipolarophile.} \]

The first use of this chemistry for peptide macrocyclization was reported by Madden et al. in 2009.\[^97,98\] Tetrazole- and (meth)acrylamide-based unnatural amino acids were introduced into a synthetic heptapeptide 43 and irradiated at 302 nm, to trigger nitrile imine formation and subsequent 1,3-dipolar cycloaddition to form 44 (Figure 15). Nitrile imines generated from both mono- and di-aryl tetrazoles were able to undergo macrocyclization with comparable efficiency, though side chain flexibility was found to be a major determinant of cyclization efficiency. Functionalized amino acids based on a shorter chain ornithine core (43a,b, n = 3, 15%) underwent cyclization with lower efficiency than those based on lysine (43c-h, n = 4, 40%). Similarly, macrocyclization efficiency was found to depend on the structure of both the tetrazole and alkene reactive partners. Electron donating substituents on the aryl tetrazole (R\(^2\) = OMe, 43e,f) greatly increased conversions by increasing the HOMO energy of the intermediate nitrile imine, therefore accelerating cycloaddition. Interestingly, methacrylamide derivatives (R\(^1\) = Me, 43f,h) were found to react more efficiently than the corresponding acrylamides (R\(^1\) = H, 43e,g) despite previous small molecule studies to the contrary.\[^94\] Acrylamides are known to possess increased reactivity towards dipolar cycloaddition, suggesting that conformational freedom is likely to play a significant role in dictating peptide cyclization efficiency. Interestingly, the conversion of monoaryl or diaryl tetrazoles was found to be comparable despite the lower reactivity of monoaryl derivatives, again suggesting a conformational influence on cyclization.\[^96\]
Figure 15. Macrocyclization of a tetrazole and alkene containing peptide under UV irradiation, via the formation of an intermediate reactive nitrile imine and subsequent 1,3-dipolar cycloaddition.\cite{97}

The high selectivity reported for nitrile imine-alkene cycloadditions would appear to make 2,5-di-aryltetrazoles interesting motifs for light-induced peptide macrocyclization. However, it is important to note that side reactions of the highly reactive nitrile imine with natural amino acid side chains have been reported.\cite{99} This may limit macrocyclization efficiencies in systems where the tetrazole and alkene reactive pair are unfavourably positioned, but does open up the intriguing possibility of using tetrazoles as light-activatable reagents for cyclization with proteinogenic amino acids.

2.4 Photoinitiated cycloaddition

In the previous section, photoactivation led to the generation of a reactive intermediate that could subsequently undergo cycloaddition. An alternative strategy is to exploit photoexcited-state molecules that can themselves directly undergo cycloaddition. For example, following photoexcitation with UV light, maleimide molecules readily undergo photochemical dimerization via a concerted [2+2] cycloaddition. In 2012, Tedaldi et al. reported that functionalized thiomaleimides, formed via a sequential addition-elimination reaction of thiols with bromomaleimides, were also able to undergo [2+2] cycloaddition under irradiation.\cite{100} As predicted by frontier molecular orbital theory, exo- head-to-head products are preferentially formed. The red-shifted absorbance of thiomaleimides relative to the parent maleimide is advantageous, enabling photoactivation with lower energy 365 nm light. This reaction was subsequently exploited for photoinduced peptide macrocyclization, as well as the re-bridging of native disulfide bonds in an antibody fragment.\cite{101} The utility of this chemistry was demonstrated by generating an analogue of the therapeutic cyclic peptide octreotide, a synthetic mimic of somatostatin. A key disulfide bridge is essential to the biological activity of octreotide, and the authors showed that linear di-thiomaleimide peptide 45 indeed showed very low biological activity. Upon irradiation at 365 nm to generate cyclized product 46, a partial recovery of activity was observed, albeit at levels < 10% of disulfide-bridged...
octreotide. Interestingly, it was noted that irradiation led to the formation of four major products of identical mass. This was proposed to be due to the formation of different diastereo- and regioisomers, in stark contrast to the high selectivity observed in the intermolecular reactions reported by Tedaldi et al. This serves as another indication that steric restrictions imparted by a peptide substrate can significantly influence regioselectivity.

![Diagram of peptide macrocyclization]

**Figure 16.** Thiomaleimide [2+2] cycloaddition under UV irradiation to generate a macrocyclic peptide.[101]

### 3. Photoinduced Electron Transfer in Peptide Macrocyclization

#### 3.1 Introduction

Photochemical macrocyclization can be initiated by quenching of a photoexcited-state chromophore through electron transfer (ET). This reaction manifold is distinct from the quenching mechanisms discussed in Section 2. Photoinduced electron transfer (PET) processes can generate radical ions from neutral starting materials, or neutral radicals from charged precursors. Reaction efficiency is governed by the propensity of these secondary reactive intermediates to undergo macrocyclization, versus deleterious side reactions or competing back electron transfer (BET) to reform the ground-state starting materials.[102–104] In this section we detail the emerging field of PET-mediated peptide macrocyclization, from its origins in UV-driven intramolecular PET, to applications of intermolecular, visible-light driven photoredox catalysis, a rapidly developing strategy that is impacting many areas of organic and biomolecular synthesis.

#### 3.2 Intramolecular PET initiated peptide macrocyclization
Intramolecular PET-initiated macrocyclizations on peptide substrates have exclusively exploited ET from donor functional groups to photoexcited phthalimides. Phthalimides are easily incorporated at the N-termini of peptide chains during solid-phase peptide synthesis, and upon photoexcitation become highly oxidizing ($E^{S1} = +2.1$ V, $E^{T1} = +1.6$ V).\cite{105} Griesbeck and co-workers performed important early work demonstrating the intramolecular cyclization of N-phthaloyl $\alpha$-amino acids under UV irradiation.\cite{106} Substrate 47 is prototypical of this approach – on photoexcitation to generate 48, the phthalimide chromophore underwent PET with the terminal carboxylate, generating a ketyl radical anion on the phthalimide, and a carboxyl radical at the end of the chain, 49 (Figure 17a). Rapid decarboxylation to form a primary alkyl radical 50 was followed by subsequent biradical intersystem crossing (a triplet state to singlet state transition via the spin flip of an electron) and ensuing cyclization via radical-radical coupling, to give macrocyclic amidol 51 in 81% yield. The authors crucially also illustrated the compatibility of this chemistry with substrates containing amide bonds (52) in the formation of 26-membered ring compound 53 (Figure 17b).

The first true PET-mediated PSP macrocyclizations were reported by Griesbeck et al. in 2002 (Figure 18).\cite{107,108} Under analogous conditions to those described above, tripeptides 54, composed of diglycine and a long-chain unnatural amino acid at either the C or N-termini, afforded 13- or 18-membered macrocycles 55 in synthetically useful yields. Interestingly, the triglycine derivative 54a successfully underwent decarboxylation but then failed to cyclize, instead affording solely the quenched linear N-methyl product in 28% yield. This behaviour was attributed to hydrogen bonding between the amide proximal to the N-terminus and the phthalimide unit, which was seemingly disrupted in substrates bearing long chain residues thus enabling cyclization. By exchanging the central glycine for its N-methylated analogue sarcosine (Sar), this H-bonding contribution could be removed and a Gly-Sar-Gly tripeptide underwent cyclization to afford a 9-membered ring in 35% yield.

**Figure 17.** First example of a PET-initiated macrocyclization by Griesbeck and co-workers. a) Mechanism of cyclization; b) Cyclization of an amide containing substrate.\cite{106}
The approach of employing substrates with N-alkylated proximal amides was subsequently extended to the first example of PET macrocyclization of a peptide composed solely of proteinogenic amino acids (Figure 19). Cyclization of the tetrapeptides (Pht)Gly-Sar-Gly-Gly \(56a\) and (Pht)Gly-Pro-Gly-Gly \(56b\) (Figure 19a/b) produced 12-membered ring compounds \(57a\) and \(57b\) respectively. Interestingly, \(57b\) was isolated as a single diastereoisomer, assigned through a combination of \(^1\)H NMR analysis and analogy to previously reported benzodiazepines.\(^{109}\) Finally, this methodology was applied to the macrocyclization of pentapeptide \(58\) (Figure 19c). Proline residues played a vital dual role in enabling cyclization of this substrate, by both removing the deactivating hydrogen bond at the second residue, and introducing a hairpin turn that facilitated cyclization by enhancing the proximity of the N- and C-termini. Unfortunately however, the presence of prolyl amide bond rotamers and the formation of diastereomers at the amidol position led to the isolation of cyclic tetramer \(59\) as a complex mixture of species.
Interestingly, in 2003 Yoon, Mariano and co-workers reported a similar example of PET-mediated decarboxylative macrocyclization on an all-glycine [(Pht)Gly-Gly-Gly-Gly] substrate.\textsuperscript{[110]} Though the proximal amide was not N-alkylated, and therefore substrate cyclization might not have been expected on the back of Griesbeck’s previous observations, a cyclic trimer was generated in 41% yield. However, the product was observed to be unstable, with complete decomposition being observed by $^1$H NMR spectroscopy over the period of 1 day. This behaviour was attributed to a deleterious amidol $\rightarrow$ amido ketone $\rightarrow$ intermolecular amidol pathway, ultimately forming insoluble oligomers. This process was successfully suppressed through the use of N-alkylated tertiary amide substrates (vide infra).

Vazdar and Basarić have subsequently applied PET-mediated decarboxylative cyclization to tetra- and penta-peptides 60 bearing N-terminal adamantyl phthalimides and C-terminal phenylalanines, or methoxylated analogues thereof (Figure 20).\textsuperscript{[111]} Products 61 were found to sit as open chain amides, rather than the amidol structures observed in previous reports, and hence are fully peptidic albeit containing an unusual, and potentially metabolically vulnerable phenyl ketone unit.\textsuperscript{[112]} Interestingly, the macrocycles were formed as single diastereomers (with the exception of 61d), with the newly formed chiral center found to be inverted between 17- or 20-membered ring sizes. The stereochemistry, assigned via a combination of NOESY spectra and molecular dynamics simulations, was determined to result from the conformation of the linear peptides which controlled the facial approach of the radical species during cyclization. Incorporation of additional Phe residues in 60b,d,f induced turns in the linear chains, and hence an opposite sense of approach. Possible epimerization post-cyclization was excluded through both computational studies and chemical evaluation through deuterium-labelling. Substrate-induced conformational control of a similar biradical macrocyclization to form an 11-membered ring product was previously observed by the authors in a separate study. In this instance however, a single stereogenic center in the precursor orchestrated the cyclization with complete stereofidelity through a chiral memory effect.\textsuperscript{[113]}

Figure 20. PET-initiated cyclization of N-adamantyl phthalimides to C-terminal phenylalanine derivatives. The stereochemistry of the resultant products was dictated by ring size.\textsuperscript{[111]}
Mechanistically, the macrocyclization of C-terminal Phe substrates \(60a\) and \(60b\) was proposed to proceed in a similar manner to that outlined in Figure 17a. Direct PET from the carboxylate to the photoexcited-state phthalimide \(62\) is followed by decarboxylation (\(63\) to \(64\)), triplet to singlet ISC, cyclization (\(64\) to \(65\)), and a final amidol to amido ketone ring expansion (\(65\) to \(66\)) (Figure 21). However, a mechanistic divergence was suggested for the C-terminal Phe(OMe) and Phe(OMe)\(_2\) substrates \(60c\) and \(60f\), with a more facile initial PET from the electron-rich arenes to the photoexcited-state phthalimide taking place, generating aryl radical cations of the type \(67\). Subsequent ET from the carboxylate to the aryl group would generate carboxyl radicals \(63\) which can then follow the established reactivity pattern. ET from the carboxylate to the more stable dimethoxy radical cations in \(67e\) and \(67f\) is slower than to the less stable monomethoxy variants \(67c\) and \(67d\), reflecting the differences in redox potentials between the two substrates and resulting in lower conversions to the cyclized products \(61e\) and \(61f\). Hence, through a balance of increased rates of initial PET from the arene to the photoexcited-state phthalimide \(62\), and a less stable radical cation intermediate \(67\) promoting rapid ET from the carboxylate, the highest yields were observed for the monomethoxy substrates \(60c\) and \(60d\). Generally, the isolated yields of the cyclic peptides \(61a\)–\(d\) generated via this methodology are rather low, seemingly reflecting difficulties in purification, which may call into question the synthetic utility (at least on scale) of a study that teaches us much about the nature of these cyclizations.

Figure 21. Mechanism of PET-initiated macrocyclization of N-adamantyl phthalimides to C-terminal phenylalanine derivatives. Phenylalanine-based substrates undergo an analogous PET process to that outlined in Figure 17. In contrast, mono- and di-methoxy substituted phenylalanines undergo an intermediate generation of an aryl radical cation.\(^{[111]}\)
Yoon, Mariano and co-workers introduced C-terminal N-trimethylsilylmethyl amides as alternative precursors for PET-mediated peptide macrocyclization in 2003.[110,114] All-glycine tri-, tetra-, and penta-cyclic peptides 68 were prepared in high yields upon UV irradiation of the N-trimethylsilylmethyl amides 69 in either acetonitrile/water mixtures or methanol (Figure 22). Protection of the peptide backbone as N-alkyl tertiary amides was required for the stability of the products.

Figure 22. Peptide macrocyclization via PET from C-terminal N-trimethylsilylmethyl amides to an excited state phthalimide.[110,114]

The authors suggest a mechanism of initial PET between the phthalimide and a proximal amide donor group, followed by amide cation radical migration (or hole transfer) along the peptide backbone to the N-trimethylsilylmethyl amide. Upon desilylation, triplet to singlet ISC and radical-radical cyclization can then occur.[110,114,115] This radical migration mechanism is supported by studies conducted by the authors on polyether- and polymethylene-linked systems, with macrocyclization yields being lower without electron donor atoms in the linker chain. Furthermore, in competition experiments on substrates containing branched chains, and hence two potential sites for reactivity, cyclization of polyether chains along which radical migration could take place was found to be favoured over hydrocarbon chain cyclization.[115] Desilylation of N-trimethylsilylmethyl amide radical cations is a rapid process, comparable in rate to the analogous decarboxylation and significantly faster than α-deprotonation of the intermediate amide radical cation.[116,117] Hence, cation radical migration is competitive with BET and α-deprotonation and therefore yields are largely independent of chain length. It is suggested that electrostatic preorganization of the phthalimide radical anion/N-trimethylsilylmethyl amide radical cation pair helps to overcome some of the entropic barrier to cyclization, lowering this energetic cost.

Collectively, these reports demonstrate that PET-initiated macrocyclization using excited-state phthalimide chemistry is potentially a simple and effective strategy to prepare peptide macrocycles. Synthetically useful yields and the possibility to control stereochemistry with substrate conformation are particular advantages of this approach. However, the use of UV light in combination with intramolecular single-electron transfer (SET) limits the composition of the precursor peptides to a subset of
amino acids which do not significantly absorb light at the same wavelengths as phthalimide, and which bear sidechains with oxidation potentials higher than that of the carboxylate donor at C-terminus. This greatly limits versatility and likely precludes the macrocyclization of peptides containing Asp, Cys, Glu, Met, Sec, Ser, Thr, Trp, or Tyr residues, or His and Lys in their free-base forms. This can be seen in the relatively limited diversity of peptide substrates that have been used. Yield-limiting hydrogen bonding, substrate instability, high dilution conditions, and a requirement for peptide N-terminal pre-functionalization (along with C-terminal for N-trimethylsilylmethyl amides) are also factors that move this strategy away from being an ideal photochemical strategy for peptide macrocyclization.

3.3 Peptide Macrocyclization Under Photoredox Catalysis

Photoredox catalysis has seen rapid growth in organic synthesis in the last decade, both in terms of application and capability.\(^{[118]}\) PET with a photoexcited photoredox catalyst \(\text{PC}^*\) under this manifold can take one of two possible courses (Figure 23): i) an oxidative quenching cycle whereby catalyst \(\text{PC}^*\) transfers an electron to an acceptor species \(A\), itself being oxidized to a form \(\text{PC}^{+1}\) which can then accept an electron from a donor \(D\) to return to the ground-state \(\text{PC}\); or alternatively, ii) a reductive quenching cycle proceeding via reduction of the photoexcited-state catalyst (\(\text{PC}^* \rightarrow \text{PC}^{-}\)) by donor \(D\), followed by oxidation back to the ground-state \(\text{PC}\) via ET to acceptor \(A\).\(^{[119]}\) Acceptor and donor molecules can be reagents, substrates, or intermediates generated during the reaction. Whether \(\text{PC}^*\) quenches oxidatively or reductively depends on the best match of the redox potentials of the excited-state catalyst relative to the species present in the reaction. Various catalytic structures can be exploited, including transition metal polypyridyl complexes,\(^{[119,120]}\) lanthanide ions,\(^{[121]}\) organic compounds,\(^{[122]}\) bulk semiconductors,\(^{[123,124]}\) or metal-organic-frameworks\(^{[125]}\).

![Figure 23. Possible catalytic quenching cycles of a photoexcited photoredox catalyst PC*](image)

When applied to PET-initiated PSP macrocyclization, photoredox catalysis offers several potential advantages over a direct intramolecular PET approach. Separating the chromophore from the substrate removes the necessity to pre-functionalize the PSP with a potentially disruptive unnatural motif. It also gives greater flexibility over chromophore structure, allowing the use of catalysts which absorb lower energy
visible light to which all natural amino acids are transparent. This makes photoredox catalysis more compatible with biological species that may suffer damage under UV irradiation.\textsuperscript{[53,126–128]} Furthermore, with indirect (and therefore asynchronous) ET from the donor to the acceptor through the intermediary of the photocatalyst, more complex and varied chemistries are accessible.\textsuperscript{[129–131]}

The first reports of peptide macrocyclization under photoredox catalysis came from the Noël group, as part of their development of photocatalyzed thiol oxidations to form disulfides.\textsuperscript{[132,133]} The proposed mechanism for this photoredox catalysed transformation commences with photoexcitation of the catalyst with white LEDs, followed by a reductive quenching event through proton-coupled electron transfer (PCET) from a thiol 70 to PC*, a proposal supported by the observation of increased yields in the presence of base (Figure 24).\textsuperscript{[132,133]} The resultant thyl radical 71, then undergoes disulfide bond formation to form 72, with the formal loss of a hydrogen atom. This is suggested to occur via sequential thiol deprotonation, thyl radical addition to the thiolate, and single-electron oxidation of the disulfide radical anion (potentially by superoxide radical anion).\textsuperscript{[61,132]} Closure of the catalytic cycle occurs through ET from the reduced form of the photocatalyst PC\textsuperscript{−1} to molecular oxygen, which acts as the terminal oxidant for this net-oxidative process. An alternative mechanism where thyl radical formation occurs via ET to singlet oxygen (\textsuperscript{1}O\textsubscript{2} - produced through catalyst-mediated photosensitization of ground-state \textsuperscript{3}O\textsubscript{2}) was discounted owing to the observation of low yields with Ru and Ir photocatalysts which are known to be capable of generating \textsuperscript{1}O\textsubscript{2}.\textsuperscript{[132]}

![Figure 24. Mechanism of photoredox formation of disulfide bonds via a reductive quenching cycle.](image)

This technology was applied to the synthesis of the cyclic peptide hormone oxytocin (73, Figure 25). Irradiation of the organic photoredox catalyst eosin Y under an oxygen gas flow in a continuous flow photoreactor (200 s residence time) offered a significant rate acceleration over batch conditions, owing to better light penetration and improved oxygen mixing which helped suppress side reactions (Figure 25, conditions A).\textsuperscript{[132]} Full conversion of linear precursor 74 was observed, with the formation of oxytocin 73 being accompanied with intermolecular disulfide peptide dimers in varying oxidized states, the formation of which was minimized by running the process at higher dilutions. The same group subsequently reported the use of the semi-conductor photocatalyst TiO\textsubscript{2} in batch, to mediate the cyclization of oxytocin 73 (Figure 25, conditions B). This heterogeneous photocatalyst is attractive as the
authors demonstrated that it could be removed by simple filtration or centrifugation, and reused up to ten times with no drop in yield. Though product formation was monitored by LC-MS, no further isolation or purification was performed and hence the question of synthetic viability remains unanswered in full. However, as a mild method (room temperature, neutral buffer, visible light) employing an easily separated and re-usable catalyst, to synthesize naturally occurring disulfide-bridged macrocycles directly from native peptides bearing two cysteines, this approach shows much promise and merits further investigation.

**Figure 25.** Photoredox-mediated peptide macrocyclization to form a disulfide linkage, catalyzed by a) eosin Y,*[^132] and b) TiO$_2$.[^133]

In 2014, MacMillan and co-workers published an influential paper outlining the Giese reaction (conjugate addition) of small molecule alkyl radicals, generated through the decarboxylation of carboxylic acids under visible-light photoredox catalysis, to electron-deficient alkenes.[^134] Notably, this chemistry was shown to be well-suited for radical generation from both N-carbamoyl α-amino acids and dipeptides. This strategy was subsequently extended to peptide macrocyclization by the same group in 2017, by incorporating a Michael acceptor radical trap at the peptide N-terminus (e.g. acrylamide 75, Figure 26).[^135] Mechanistically, PET from the carboxylate form of the C-terminal acid to the photoexcited catalyst PC* generates a carboxy radical that rapidly undergoes loss of CO$_2$ to produce α-amido radical 76. Following an intramolecular Michael addition, α-carbonyl radical 77 is reduced to an enolate species through ET from PC$^-1$, closing the photoredox catalytic cycle and affording macrocycle 78 after protonation. The macrocyclic products of the type 78 are fully peptidic, containing an unnatural γ-amino acid linkage. The non-canonical γ-aminobutyric acid linker is likely to be insensitive to the action of proteases, providing products with increased stability under physiological conditions.
When a series of N-acryloyl pentapeptides with a C-terminal glycine residue, 79a-g, were irradiated with a blue LED in the presence of the oxidising iridium photocatalyst \( \text{Ir} \left[ \text{dF(CF}_3\text{ppy)}_2(\text{dtbbpy})\text{PF}_6 \right] \), the corresponding macrocyclic peptides 80a-g were produced efficiently, although isolated yields varied which is a common observation in this field (Figure 27). The methodology was applied to peptides containing a broad range of amino acids, including non-canonical propargylglycine (Pra) and N-methyl alanine (N-Me-Ala). Notably, for examples with amino acids bearing polar sidechains protecting groups were utilized, potentially to suppress undesired nucleophilic and/or redox reactions. This lack of functional group tolerance may limit the widespread applicability of this chemistry at the present time. For substrates 79a and 79c, the addition of 10 mol% 2,4,6-triisopropylthiophenol to the reaction mixture was found to be beneficial to the yield. This effect was attributed to interception of the \( \alpha \)-carbonyl radical of the form 77 (Figure 26) via H-atom abstraction from the thiol at a rate competitive with retro-Michael addition and unwanted intermolecular oligomerization, negating the need for rate-limiting reduction by \( \text{PC}^{-1} \), to provide products 80a and 80c. In these instances, the catalytic cycle is instead closed via ET from \( \text{PC}^{-1} \) to the resultant thiol radical.
Figure 27. First example of a photoredox-mediated Giese reaction for peptide macrocyclization. \(^{a}\)10 mol\% 2,4,6-trisopropylthiophenol added. \(^{b}\)Pra = propargylglycine. \(^{c}\)N-Me-Ala = N-methyl alanine.\(^{[135]}\)

Expanding the scope of the reaction further, the authors explored alternative functionalities at the C- and N-termini. Peptides bearing C-terminal amino acids with \(\alpha\)-substituents (81a–f) were found to be well tolerated and underwent decarboxylative cyclization, however yields were reduced relative to terminal glycines (Figure 28). This may be attributable to increased rates of retro-Michael addition resulting from the increased stability of the substituted \(\alpha\)-amido radicals generated. The formation of diastereomeric product mixtures was found to occur with little control (81b and 81c). Notably, substrate 81c with a C-terminal glutamic acid underwent chemoselective \(\alpha\)-amino decarboxylation on account of the lower pKa and oxidation potential of the C-terminus relative to the \(\gamma\)-carboxylic acid.\(^{[136]}\) Though not demonstrated, analogous selectivity over aspartic acid side chains would also be expected for the same reasons. Unnatural di-\(\alpha\)-substituted cyclic amino acids at the peptide C-termini were able to form unusual spirocyclic peptide macrocycles (82d and 82e). At the N-terminus, only a single functionalized acrylamide motif was investigated, with an electronically activating phenyl group at the \(\alpha\)-position leading to the formation of macrocycle 82f in high yield and with good diastereoselectivity. The methodology performed well when applied to the synthesis of larger ring sizes, with remarkably little yield variation observed across the preparation of 8-, 10- and 15-membered cyclic peptides. Importantly, the authors also demonstrated the straightforward post-cyclization removal of acid-labile protecting groups, leading to the generation of the somatostatin-receptor agonist peptide COR-005 (83 in 47% isolated yield over two steps).
Figure 28. Alternative C- and N-terminal functionalities for the Giese macrocyclization of peptide substrates. \(^a\)N-Me-Leu = N-methyl leucine. \(^b\)Reaction performed in DMSO. \(^c\)ACC = 1-Aminocyclopropane-1-carboxylic acid. \(^d\)AC5 = Cycloleucine. \(^e\)Yield over two steps, following removal of protecting groups with TFA:PhOH:H\(_2\)O:iPr\(_3\)SiH (88:5:5:2) at 0 °C for 2 h.\(^{135}\)

3.4 Peptide macrocyclization under dual photoredox and transition metal catalysis

Advances in the field of photoredox catalysis have been mirrored in their application in PSP macrocyclization. At the cutting edge of this area is the merger of photoredox and transition metal catalysis. This extremely powerful combination opens up unprecedented chemical transformations that are well-suited to application in complex settings on account of the mild reaction conditions.\(^{131,137,138}\) Key to the success of this approach is the activation of transition metal complexes by a photocatalyst, either through redox modulation or energy transfer pathways, triggering mechanistic steps that would not be operative under transition metal catalysis alone.\(^{139–144}\)

The Sciammetta group disclosed a methodology for the etherification of peptidic alcohols with aryl bromides via C(sp\(^2\))–O cross-coupling under dual photoredox and
In addition to intermolecular ether formation, the reaction conditions were also shown to be amenable to macrocyclization of a series of N-terminal bromobenzamides, via aryl etherification of C-terminal serine derivatives. The authors proposed this methodology to target underutilized ether macrocyclic linkages, which they anticipated would overcome several of the shortcomings at times exhibited by other macrocycle chemistries, such as proteolytic instability (e.g. thioether linkages), and the poor cell-permeability of hydrogen bond-donor containing linkages (e.g. those with an N-H bond).

The proposed mechanism of this reaction was analogous to that previously reported by MacMillan and co-workers for the dual photoredox-nickel catalyzed arylation of small-molecule alcohols (Figure 29). Following the oxidative addition of a Ni$^{0}$ complex 84 into an aryl bromide 85, the resultant aryl-Ni$^{II}$ species 86 can undergo ligand exchange to form cyclic Ni$^{II}$ alkoxide 87. Single-electron oxidation of complex 87 via ET to the excited-state photocatalyst PC$^*$ generates a key Ni$^{III}$ aryl alkoxide 88 which, unlike Ni$^{II}$ aryl alkoxide complex 87, is unstable with respect to reductive elimination and the catalytic cycle therefore generates peptide macrocycle 89 and the Ni$^{I}$ species 90. Finally, ET to 90 from PC$^{-1}$ then regenerates Ni$^{0}$ species 84 and simultaneously closes both catalytic cycles.

![Figure 29. Mechanism of dual photoredox-nickel catalysis for the macrocyclic etherification of alcohols with aryl bromides.](image)

Using the isophthalonitrile-based organic photocatalyst 4DPAIPN 91 and a NiBr$_2$: glyme/dtbbpy transition metal catalyst system, in concert with quinuclidine as a yield boosting additive (an affect attributed to its capacity to act as an electron donor or shuttle), the authors were able to induce macrocyclization of a series of N-terminal m-bromobenzamide modified peptides 92a–f (Figure 30). All substrates contained a β-hairpin inducing D-Pro–L-Pro subunit to preorganize the linear peptide for cyclization via rigidifying hydrogen bonds, leading to the formation of cyclized peptides 93. In the absence of this dipeptide only trace yields of macrocycle formation were detected by UPLC-MS. The importance of these hydrogen bonds...
was supported by in silico conformational sampling via a distance geometry approach. For pentapeptide substrates with C-terminal serinamides (92a,b,e), competing O- and N-arylation was observed to provide ether and amide linked macrocycles which were separable by HPLC. This competitive cyclization favoured the ether over amide linkages by ratios of 6:4–7:3 (see Figure 30). For substrates with a C-terminal ester (92c) or glycaminide (92d) residues, presenting a single nucleophilic position, macrocyclization proceeded via exclusive etherification or amidation respectively. Variation of the peptide sequence to incorporate residues bearing protected polar functionality (92e) or increasing the ring size (92f) was also successful. Interestingly, macrocyclization in the presence of a free carboxylic acid was not demonstrated, potentially because of competing lactonization. This transformation has been demonstrated in an intermolecular sense via an energy-transfer mechanism under similar reaction conditions. Notably, the requirement for the protection of polar residues, and more importantly the presence of a turn-enforcing D-Pro–L-Pro subunit to achieve good cyclization efficiency are limitations that need to be addressed for this methodology to achieve broad synthetic applicability in peptide macrocyclization.

Figure 30. Dual photoredox-nickel catalysis for peptide macrocyclization.

4. Conclusion

In this review, we have detailed the photochemical strategies that have been exploited to induce PSP macrocyclization. These methods benefit from mild reaction conditions, the ability to selectively introduce energy, and the sheer variety of reaction modes available. Developments in this field have mirrored the recent renewed interest in photochemistry, and this looks set to continue as macrocyclic PSPs become increasingly prevalent in the drug discovery pipeline. The rapid expansion of new photochemical strategies in organic synthesis provides a rich source of potential new reactions to control cyclization. Indeed, there remains significant scope for innovation in the area, with a pressing need to both evolve existing strategies for cyclization, and develop new ones.
In particular, we see an urgent need for new reactions that can overcome limitations in amino acid tolerance. It is notable that in many of the examples presented within this review, the linear peptide substrates are composed of a very simple set of afunctional amino acids with hydrocarbon side chains. The resultant methodologies, while synthetically interesting, provide little scope to generate macrocyclic peptides with useful biological properties. A lack of functional group tolerance is also highlighted by the common reliance on fully protected peptides as substrates. Exceptions, such as thiol-‘ene’ cyclizations, must proceed with exquisite chemoselectivity to avoid side-reactions with the myriad of reactive functionalities found within unprotected, canonical amino acids. Recent developments in photoredox catalysis, enabling site-selective modification of even complex proteins, are therefore particularly exciting, though it should be noted that even then the presence of a sub-set of amino acids must be avoided.\cite{136,146} New reactions that can increase this selectivity still further would be invaluable to the bioconjugate community, and would provide an invaluable and generalisable tool for peptide macrocyclization.

Moreover, we anticipate increased use of reactions that exploit lower energy visible light sources to induce macrocyclization, with associated improvements in selectivity. The reliance of many of the reactions described here on high energy UV irradiation is inherently limiting to functional group tolerance.\cite{147} Photoredox catalysis driven by visible light is an important avenue to overcome this limitation, and we anticipate a greater application of this technology to the macrocyclization of PSPs. Other visible light-driven photochemical catalysis manifolds, such as energy transfer and photon upconversion, are also well-positioned for implementation in PSP macrocyclization. With these innovations the prospect of achieving general strategies for protein macrocyclization will become more realistic, adding to the fairly limited toolkit of photochemical reactions that have been exploited for this ambitious goal to date.\cite{74,101}

Finally, the development of photochemical reactions that can deliver unique macrocyclic linkages represents an important future challenge for the community. The diversity of reaction manifolds that are accessible through photochemical approaches is attractive given the importance of even subtle differences in peptide linker structure on biological properties, such as membrane permeability and target binding.\cite{71} New reactions that can be used to modulate the properties of the resultant cyclic peptides would therefore represent a valuable tool in the search for novel peptide therapeutics. Important challenges that need to be overcome in this regard include the stereocontrol of newly formed chiral centres, a difficult task in such a functionally-dense environment, and a reduced reliance on turn-inducing residues to aid cyclization.

We anticipate the benefits of light-mediated chemistry will be increasingly exploited in the future as the field matures. Exciting opportunities for spatial and temporal control over cyclization may enable applications in advanced biomedical
technologies, diversifying away from the traditional roles of cyclized peptides as therapeutic agents. As the journey towards more ‘ideal’ macrocyclization techniques that are readily applicable to PSP substrates and that tolerate ever increasing functionality continues, the future for photochemical macrocyclization methods looks bright.

5. Acknowledgements

L.R. and C.D.S. are grateful to the Rosetrees Trust for PhD Studentship funding (agreement A2413). N.C.R. and C.D.S. acknowledge PhD Studentship support through the EPSRC Doctoral Training Partnership. J.R.D. and C.D.S. thank the University of York for Pump Priming funding.

6. Author biographies

Laetitia Raynal studied Chemistry at the École Nationale Supérieure de Chimie de Montpellier, with a major in Materials chemistry. During her Masters, she did an internship at DSM in the Netherlands and a one-year research placement at the University of Geelong in Australia, graduating with a Diplôme d’ingénieur Chimiste in 2019. The same year, she was awarded a Rosetrees Trust PhD Scholarship with Dr Chris Spicer at the University of York, developing new approaches to enhance protein signalling.

Nicholas C. Rose received his MSci of Chemistry with a Year in Industry in 2019 from the University of Nottingham, following an industrial placement year in 2018 at Merck Chemicals Ltd. in Southampton. In 2019 he was awarded an EPSRC Doctoral PhD Scholarship with Dr. Christopher Spicer at the University of York. His current
research interests involve developing novel dynamic biomaterial conjugation systems for tissue engineering.

James studied Natural Sciences at the University of Cambridge, and then obtained his Ph.D. at the University of York with Prof. Richard Taylor. Following postdoctoral research at the University of Texas at Austin and the University of Oxford, he joined Prof. David MacMillan’s group at Princeton University. On returning to York, he has worked with Prof. Richard Taylor, Prof. Peter O’Brien and Dr Chris Spicer. His research interests centre on applications of photoredox catalysis in novel bond construction and the synthesis of molecules of medicinal relevance.

Chris is a Lecturer in Chemistry at the University of York, where his group is interested in the design and chemical synthesis of novel biomaterials and bioconjugates for tissue repair. Previously, he studied Natural Sciences at the University of Cambridge, before moving to the University of Oxford to undertake a PhD with Prof. Ben Davis. Chris went on to complete postdoctoral research with Prof. Molly Stevens, first at Imperial College London and then the Karolinska Institute in Stockholm.
Photochemical reactions have emerged to prominence over the past two decades, leading to the development of many new diverse and powerful chemical transformations. In this paper, the application of these established and emerging methodologies for peptide macrocyclization is reviewed, providing a new route to generate improved therapeutic constructs.

Recent developments have allowed photochemical reactions to move to the forefront of the search for new strategies for peptide macrocyclization. Here, these developments and emerging photochemistries are...
reviewed, highlighting the opportunities to further innovate in this rapidly expanding field.

Keywords
Peptides; Photochemistry; Macrocyclization;

7. References

[1] A. M. Ali, J. Atmaj, N. Van Oosterwijk, M. R. Groves, A. Dömling, Comput. Struct. Biotechnol. J. 2019, 17, 263.
[2] M. Moiola, M. G. Memeo, P. Quadrelli, Molecules 2019, 24.
[3] X. Jing, K. Jin, Med. Res. Rev. 2020, 40, 753.
[4] N. Tsomaia, Eur. J. Med. Chem. 2015, 94, 459.
[5] P. G. Dougherty, A. Sahni, D. Pei, Chem. Rev. 2019, 119, 10241.
[6] L. Otvos, J. D. Wade, Front. Chem. 2014, 2, 62.
[7] K. Fosgerau, T. Hoffmann, Drug Discov. Today 2015, 20, 122.
[8] T. A. F. Cardote, A. Ciulli, ChemMedChem 2016, 11, 787.
[9] A. C. L. Lee, J. L. Harris, K. K. Khanna, J. H. Hong, Int. J. Mol. Sci. 2019, 20, 2383.
[10] D. S. Nielsen, N. E. Shepherd, W. Xu, A. J. Lucke, M. J. Stoermer, D. P. Fairlie, Chem. Rev. 2017, 117, 8094.
[11] Á. Roxin, G. Zheng, Future Med. Chem. 2012, 4, 1601.
[12] T. A. Hill, N. E. Shepherd, F. Diness, D. P. Fairlie, Angew. Chemie - Int. Ed. 2014, 53, 13020.
[13] J. Garner, M. M. Harding, Org. Biomol. Chem. 2007, 5, 3577.
[14] L. K. Henchey, A. L. Jochim, P. S. Arora, Curr. Opin. Chem. Biol. 2008, 12, 692.
[15] A. D. De Araujo, H. N. Hoang, W. M. Kok, F. Diness, P. Gupta, T. A. Hill, R. W. Driver, D. A. Price, S. Liras, D. P. Fairlie, Angew. Chemie - Int. Ed. 2014, 53, 6965.
[16] T. K. Sawyer, V. Guerlavais, K. Darlak, E. Feyfant, RSC Drug Discov. Ser. 2015, 40, 339.
[17] Y. Tian, Y. Jiang, J. Li, D. Wang, H. Zhao, Z. Li, ChemBioChem 2017, 18, 2087.
[18] X. Li, Y. Zou, H. G. Hu, Chinese Chem. Lett. 2018, 29, 1088.
[19] K. J. Skowron, T. E. Speltz, T. W. Moore, Med. Res. Rev. 2019, 39, 749.
[20] D. A. Guaraccino, J. A. Riordan, G. M. Barreto, A. L. Oldfield, C. M. Kouba, D. Agrinsoni, Chem. Rev. 2019, 119, 9915.
[21] W. A. Loughlin, J. D. A. Tyndall, M. P. Glenn, T. A. Hill, D. P. Fairlie, Chem. Rev. 2004, 104, 6085.
[22] J. A. Robinson, Acc. Chem. Res. 2008, 41, 1278.
[23] A. K. Yudin, Chem. Sci. 2015, 6, 30.
[24] J. E. Bock, J. Gavenonis, J. A. Kritzer, ACS Chem. Biol. 2013, 8, 488.
[25] A. P. Benfield, M. G. Teresk, H. R. Plake, J. E. DeLorbe, L. E. Millsbaugh, S. F. Martin, Angew. Chemie - Int. Ed. 2006, 45, 6830.
[26] D. G. Udugamasooriya, M. R. Spaller, Biopolymers 2008, 89, 653.
[27] J. E. Delorbe, J. H. Clements, B. B. Whiddon, S. F. Martin, ACS Med. Chem. Lett. 2010, 1, 448.
[28] S. F. Martin, J. H. Clements, Annu. Rev. Biochem. 2013, 82, 267.
[29] E. M. Driggers, S. P. Hale, J. Lee, N. K. Terrett, Nat. Rev. Drug Discov. 2008, 7, 608.
[30] J. Xie, H. Jin, A. S. K. Hashmi, Chem. Soc. Rev. 2017, 46, 5193.
[31] F. Giordanetto, J. Kihlberg, J. Med. Chem. 2014, 57, 278.
[32] M. R. Naylor, A. T. Bockus, M. J. Blanco, R. S. Lokey, Curr. Opin. Chem. Biol. 2017, 38, 141.
[33] C. T. Walsh, ACS Infect. Dis. 2018, 4, 1283.
[34] S. J. Miller, R. H. Grubbs, J. Am. Chem. Soc. 1995, 117, 5855.
[35] M. Roice, I. Johannsen, M. Meldal, QSAR Comb. Sci. 2004, 23, 662.
[36] Y. H. Lau, Y. Wu, P. De Andrade, W. R. J. D. Galloway, D. R. Spring, Nat. Protoc. 2015, 10, 585.
[37] D. G. Rivera, G. M. Ojeda-Carralero, L. Reguera, E. V. Van der Eycken, Chem. Soc. Rev. 2020, 49, 2039.
[38] N. S. Robertson, S. J. Walsh, E. Fowler, M. Yoshida, S. M. Rowe, Y. Wu, H. F. Sore, J. S. Parker, D. R. Spring, Chem. Commun. 2019, 55, 9499.
[39] A. M. Spokoyny, Y. Zou, J. J. Ling, H. Yu, Y. S. Lin, B. L. Pentelute, J. Am. Chem. Soc. 2013, 135, 5946.
[40] A. Failli, H. Immer, M. Götz, Can. J. Chem. 1979, 57, 3257.
[41] A. V. Vasco, C. S. Pérez, F. E. Morales, H. E. Garay, D. Vasilev, J. A. Gavín, L. A. Wessjohann, D. G. Rivera, J. Org. Chem. 2015, 80, 6697.
[42] S. Sengupta, G. Mehta, Org. Biomol. Chem. 2020, 18, 1851.
[43] J. S. Davies, J. Pept. Sci. 2003, 9, 471.
[44] C. J. White, A. K. Yudin, Nat. Chem. 2011, 3, 509.
[45] Y. H. Lau, P. De Andrade, Y. Wu, D. R. Spring, Chem. Soc. Rev. 2015, 44, 91.
[46] J. Wu, J. Tang, H. Chen, Y. He, H. Wang, H. Yao, Tetrahedron Lett. 2018, 59,
325.

[47] L. Reguera, D. G. Rivera, Chem. Rev. 2019, 119, 9836.

[48] V. Martí-Centelles, M. D. Pandey, M. I. Burguete, S. V. Luis, Chem. Rev. 2015, 115, 8736.

[49] A. G. Griesbeck, Liebigs Ann. 1996, 1581.

[50] L. G. López-Valdeza, H. Zuleta-Prada, B. Reyes-Trejo, E. Cuevas-Yañez, Tetrahedron 2018, 74, 5416.

[51] S. Prasad, I. Mandal, S. Singh, A. Paul, B. Mandal, R. Venkatramani, R. Swaminathan, Chem. Sci. 2017, 8, 5416.

[52] G. Knör, Chem. - A Eur. J. 2009, 15, 568.

[53] S. H. Lee, D. S. Choi, S. K. Kuk, C. B. Park, Angew. Chemie - Int. Ed. 2018, 57, 7958.

[54] J. Li, H. Kong, C. Zhu, Y. Zhang, Chem. Sci. 2020, 11, 3390.

[55] Y. S. Or, R. F. Clark, J. R. Luly, J. Org. Chem. 1991, 56, 3146.

[56] H. Jo, N. Meinhardt, Y. Wu, S. Kulkarni, X. Hu, K. E. Low, P. L. Davies, W. F. DeGrado, D. C. Greenbaum, J. Am. Chem. Soc. 2012, 134, 17704.

[57] A. M. Leduc, J. O. Trent, J. L. Wittliff, K. S. Bramlett, S. L. Briggs, N. Y. Chirgadze, Y. Wang, T. P. Burris, A. F. Spatola, Proc. Natl. Acad. Sci. U. S. A. 2003, 100, 11273.

[58] B. P. Sutherland, B. M. El-Zaatari, N. I. Halaszynski, J. M. French, S. Bai, C. J. Kloxin, Bioconjug. Chem. 2018, 29, 3987.

[59] C. D. Spicer, B. G. Davis, Nat. Commun. 2014, 5, 4740.

[60] H. Chen, R. Huang, Z. Li, W. Zhu, J. Chen, Y. Zhan, B. Jiang, Org. Biomol. Chem. 2017, 15, 7339.

[61] F. Dénès, M. Pichowicz, G. Povie, P. Renaud, Chem. Rev. 2014, 114, 2587.

[62] E. Denisov, C. Crissostomos, A. Shestakov, T. Denisova, Int. J. Chem. Kinet. 2009, 41, 284.

[63] N. B. Cramer, S. K. Reddy, A. K. O'Brien, C. N. Bowman, Macromolecules 2003, 36, 7964.

[64] C. E. Hoyle, A. B. Lowe, C. N. Bowman, Chem. Soc. Rev. 2010, 39, 1355.

[65] B. H. Northrop, R. N. Coffey, J. Am. Chem. Soc. 2012, 134, 13804.

[66] D. M. Love, K. Kim, J. T. Goodrich, B. D. Fairbanks, B. T. Worrell, M. P. Stoykovich, C. B. Musgrave, C. N. Bowman, J. Org. Chem. 2018, 83, 2912.

[67] A. A. Aimetti, R. K. Shoemaker, C. C. Lin, K. S. Anseth, Chem. Commun. 2010, 46, 4061.

[68] A. A. Aimetti, K. R. Feaver, K. S. Anseth, Chem. Commun. 2010, 46, 5781.

[69] B. Zhao, Q. Zhang, Z. Li, J. Pept. Sci. 2016, 540.
[70] Y. Wang, D. H. C. Chou, Angew. Chemie - Int. Ed. 2015, 54, 10931.
[71] Y. Tian, J. Li, H. Zhao, X. Zeng, D. Wang, Q. Liu, X. Niu, X. Huang, N. Xu, Z. Li, Chem. Sci. 2016, 7, 3325.
[72] S. Chen, D. Bertoldo, A. Angelini, F. Pojer, C. Heinis, Angew. Chemie - Int. Ed. 2014, 53, 1602.
[73] F. Bernal, M. Wade, M. Godes, T. N. Davis, D. G. Whitehead, A. L. Kung, G. M. Wahl, L. D. Walensky, Cancer Cell 2010, 18, 411.
[74] Y. Wang, B. J. Bruno, S. Cornillie, J. M. Nogieira, D. Chen, T. E. Cheatham, C. S. Lim, D. H. C. Chou, Chem. - A Eur. J. 2017, 23, 7087.
[75] A. Massi, D. Nanni, Org. Biomol. Chem. 2012, 10, 3791.
[76] G. Dormán, H. Nakamura, A. Pulsipher, G. D. Prestwich, Chem. Rev. 2016, 116, 15284.
[77] S. S. Ge, B. Chen, Y. Y. Wu, Q. S. Long, Y. L. Zhao, P. Y. Wang, S. Yang, RSC Adv. 2018, 8, 29428.
[78] E. Deseseke, Y. Nakatani, G. Ourisson, European J. Org. Chem. 1998, 243.
[79] A. Moretto, M. Crisma, F. Formaggio, L. A. Huck, D. Mangion, W. J. Leigh, C. Toniolo, Chem. - A Eur. J. 2009, 15, 67.
[80] A. Moretto, M. Crisma, F. Formaggio, C. Toniolo, Biopolymers 2010, 94, 721.
[81] K. Wright, A. Moretto, M. Crisma, M. Wakselman, J. Mazaleyrat, F. Formaggio, C. Toniolo, Org. Biomol. Chem. 2010, 3281.
[82] A. Lewandowska-Andralojc, F. Kazmierczak, G. L. Hug, G. Hörner, B. Marciniak, Photochem. Photobiol. 2013, 89, 14.
[83] G. Papageorgiou, D. C. Ogden, A. Barth, J. E. T. Corrie, J. Am. Chem. Soc. 1999, 121, 6503.
[84] B. Amit, D. A. Ben-Efraim, A. Patchornik, J. Am. Chem. Soc. 1976, 98, 843.
[85] K. C. Nicolaou, B. S. Safina, N. Winssinger, Synlett 2001, 900.
[86] Y. Mifune, H. Nakamura, S. Fuse, Org. Biomol. Chem. 2016, 14, 11244.
[87] M. D. Pierschbacher, E. Ruoslahti, Proc. Natl. Acad. Sci. U. S. A. 1984, 81, 5985.
[88] M. D. Pierschbacher, E. Ruoslahti, Nature 1984, 3.
[89] M. Gurrath, J. Calvetd, H. Kessler, R. Tim, FEBS Lett. 1991, 29, 50.
[90] M. Gurrath, G. Muller, H. Kessler, M. Aumailley, R. Timpl, Eur. J. Biochem. 1992, 210, 911.
[91] K. M. Yamada, J. Biol. Chem. 1991, 266, 12809.
[92] A. Isidro-llobet, M. N. Kenworthy, S. Mukherjee, M. E. Kopach, K. Wegner, F. Gallou, A. G. Smith, F. Roschangar, J. Org. Chem. 2019, 84, 4615.
[93] Y. Wang, C. I. Rivera Vera, Q. Lin, Org. Lett. 2007, 9, 4155.
[94] Y. Wang, W. J. Hu, W. Song, R. K. V Lim, Q. Lin, Org. Chem. Front. 2008, 10, 3725.
[95] Y. Wang, W. Song, W. J. Hu, Q. Lin, Angew. Chem. Int. Ed. Engl. 2009, 48, 5330.
[96] J. Wang, W. Zhang, W. Song, Y. Wang, Z. Yu, J. Li, M. Wu, L. Wang, J. Zang, Q. Lin, J. Am. Chem. Soc. 2010, 132, 14812.
[97] M. M. Madden, C. I. Rivera Vera, W. Song, Q. Lin, Chem. Commun. 2009, 5588.
[98] M. M. Madden, A. Muppidi, Z. Li, X. Li, J. Chen, Q. Lin, Bioorganic Med. Chem. Lett. 2011, 21, 1472.
[99] Z. Li, L. Qian, L. Li, J. C. Bernhammer, H. V. Huynh, J. S. Lee, S. Q. Yao, Angew. Chemie - Int. Ed. 2016, 55, 2002.
[100] L. M. Tedaldi, A. E. Aliev, J. R. Baker, Chem. Commun. 2012, 48, 4725.
[101] D. A. Richards, S. A. Fletcher, M. Nobles, H. Kossen, L. Tedaldi, V. Chudasama, A. Tinker, J. R. Baker, Org. Biomol. Chem. 2016, 14, 455.
[102] A. G. Griesbeck, N. Hoffmann, K. D. Warzecha, Acc. Chem. Res. 2007, 40, 128.
[103] T. Kumpulainen, B. Lang, A. Rosspeintner, E. Vauthey, Chem. Rev. 2017, 117, 10826.
[104] Y. Wei, Q. Q. Zhou, F. Tan, L. Q. Lu, W. J. Xiao, Synth. 2019, 51, 3021.
[105] U. C. Yoon, P. S. Mariano, Acc. Chem. Res. 2001, 34, 523.
[106] A. G. Griesbeck, A. Henz, K. Peters, E. -M Peters, H. G. von Schnering, Angew. Chemie Int. Ed. English 1995, 34, 474.
[107] A. G. Griesbeck, T. Heinrich, M. Oelgemöller, J. Lex, A. Molis, J. Am. Chem. Soc. 2002, 124, 10972.
[108] A. G. Griesbeck, T. Heinrich, M. Oelgemöller, A. Molis, A. Heidtmann, Helv. Chim. Acta 2002, 85, 4561.
[109] A. G. Griesbeck, W. Kramer, J. Lex, Angew. Chem. Int. Ed. Engl. 2001, 40, 577.
[110] U. C. Yoon, Y. X. Jin, S. W. Oh, C. H. Park, J. H. Park, C. F. Campana, X. Cai, E. N. Duesler, P. S. Mariano, J. Am. Chem. Soc. 2003, 125, 10664.
[111] M. Sohora, M. Vazdar, I. Sović, K. Mlinarić-Majerski, N. Basarić, J. Org. Chem. 2018, 83, 14905.
[112] D. A. Griffith, D. W. Kung, W. P. Esler, P. A. Amor, S. W. Bagley, C. Beysen, S. Carvajal-Gonzalez, S. D. Doran, C. Limberakis, A. M. Mathiowetz, K. McPherson, D. A. Price, E. Ravussin, G. E. Sonnenberg, J. A. Southers, L. J. Sweet, S. M. Turner, F. F. Vajdos, J. Med. Chem. 2014, 57, 10512.
[113] T. Šumanovac Ramljak, M. Sohora, I. Antol, D. Kontrec, N. Basarić, K. Mlinarić-Majerski, Tetrahedron Lett. 2014, 55, 4078.
[114] D. W. Cho, U. C. Yoon, P. S. Mariano, Acc. Chem. Res. 2011, 44, 204.
[115] W. C. Dae, H. C. Jung, W. O. Sun, C. Quan, C. Y. Ung, R. Wang, S. Yang, P. S. Mariano, J. Am. Chem. Soc. 2008, 130, 2276.
[116] X. Zhang, S. R. Yeh, S. Hong, M. Freccero, A. Albini, D. E. Falvey, P. S. Mariano, J. Am. Chem. Soc. 1994, 116, 4211.
[117] Z. Su, P. S. Mariano, D. E. Falvey, U. C. Yoon, S. W. Oh, J. Am. Chem. Soc. 1998, 120, 10676.
[118] R. C. McAtee, E. J. McClain, C. R. J. Stephenson, Trends Chem. 2019, 1, 111.
[119] C. K. Prier, D. A. Rankic, D. W. C. MacMillan, Chem. Rev. 2013, 113, 5322.
[120] T. Koike, M. Akita, Inorg. Chem. Front. 2014, 1, 562.
[121] Y. Qiao, E. J. Schelter, Acc. Chem. Res. 2018, 51, 2926.
[122] N. A. Romero, D. A. Nicewicz, Chem. Rev. 2016, 116, 10075.
[123] A. Savateev, I. Ghosh, B. König, M. Antonietti, Angew. Chemie - Int. Ed. 2018, 57, 15936.
[124] J. Chen, J. Cen, X. Xu, X. Li, Catal. Sci. Technol. 2016, 6, 349.
[125] X. Yu, L. Wang, S. M. Cohen, CrystEngComm 2017, 19, 4126.
[126] J. Q. Liu, A. Shatskiy, B. S. Matsuura, M. D. Kärkäs, Synth. 2019, 51, 2759.
[127] C. Bottecchia, T. Noël, Chem. - A Eur. J. 2019, 25, 26.
[128] S. Angerani, N. Winssinger, Chem. - A Eur. J. 2019, 25, 6661.
[129] K. L. Skubi, T. R. Blum, T. P. Yoon, Chem. Rev. 2016, 116, 10035.
[130] J. C. Tellis, C. B. Kelly, D. N. Primer, M. Jouffroy, N. R. Patel, G. A. Molander, Acc. Chem. Res. 2016, 49, 1429.
[131] J. Twilton, C. C. Le, P. Zhang, M. H. Shaw, R. W. Evans, D. W. C. MacMillan, Nat. Rev. Chem. 2017, 1.
[132] A. Talla, B. Driessen, N. J. W. Straathof, L. G. Milroy, L. Brunsveld, V. Hessel, T. Noël, Adv. Synth. Catal. 2015, 357, 2180.
[133] C. Bottecchia, N. Erdmann, P. M. A. Tijsse, L. G. Milroy, L. Brunsveld, V. Hessel, T. Noël, ChemSusChem 2016, 9, 1781.
[134] Z. Zuo, D. W. C. Macmillan, J. Am. Chem. Soc. 2014, 136, 5257.
[135] S. J. McCarver, J. X. Qiao, J. Carpenter, R. M. Borzilleri, M. A. Poss, M. D. Eastgate, M. M. Miller, D. W. C. MacMillan, Angew. Chemie - Int. Ed. 2017, 56, 728.
[136] S. Bloom, C. Liu, D. K. Kölmel, J. X. Qiao, Y. Zhang, M. A. Poss, W. R. Ewing, D. W. C. C. Macmillan, Nat. Chem. 2017, 10, 205.
[137] Z. Zuo, D. T. Ahneman, L. Chu, J. A. Terrett, A. G. Doyle, D. W. C. MacMillan, Science 2014, 345, 437.
[138] J. C. Tellis, D. N. Primer, G. A. Molander, Science 2014, 345, 433.
[139] J. A. Terrett, J. D. Cuthbertson, V. W. Shurtleff, D. W. C. MacMillan, Nature 2015, 524, 330.
[140] E. R. Welin, C. Le, D. M. Arias-Rotondo, J. K. McCusker, D. W. C. MacMillan, Science 2017, 355, 380.
[141] B. J. Shields, B. Kudisch, G. D. Scholes, A. G. Doyle, J. Am. Chem. Soc. 2018, 140, 3035.
[142] L. Tian, N. A. Till, B. Kudisch, D. W. C. MacMillan, G. D. Scholes, J. Am. Chem. Soc. 2020, 142, 4555.
[143] S. I. Ting, S. Garakhyaraghi, C. M. Taliaferro, B. J. Shields, G. D. Scholes, F. N. Castellano, A. G. Doyle, ACS Appl. Mater. Interfaces 2020, 142, 5800.
[144] M. Yuan, Z. Song, S. O. Badir, G. A. Molander, O. Gutierrez, J. Am. Chem. Soc. 2020, 142, 7225.
[145] H. Lee, N. C. Boyer, Q. Deng, H. Y. Kim, T. K. Sawyer, N. Sciammetta, Chem. Sci. 2019, 10, 5073.
[146] X. Chen, F. Ye, X. Luo, X. Liu, J. Zhao, S. Wang, Q. Zhou, G. Chen, P. Wang, J. Am. Chem. Soc. 2019.
[147] Q.-Q. Q. Zhou, Y.-Q. Q. Zou, L.-Q. Q. Lu, W.-J. J. Xiao, Angew. Chemie - Int. Ed. 2019, 58, 1586.