Radical-Based Synthesis and Modification of Amino Acids

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Amino acids (AAs) are key structural motifs with widespread applications in organic synthesis, biochemistry, and material sciences. Recently, with the development of milder and more versatile radical-based procedures, the use of strategies relying on radical chemistry for the synthesis and modification of AAs has gained increased attention, as they allow rapid access to libraries of novel unnatural AAs containing a wide range of structural motifs. In this Minireview, we provide a broad overview of the advancements made in this field during the last decade, focusing on methods for the de novo synthesis of α-, β-, and γ-AAs, as well as for the selective derivatisation of canonical and non-canonical α-AAs.

1. Introduction

Amino acids (AAs) are widely used in applications across several scientific fields; for example, they can be used as ligands in transition-metal catalysis,[1] as key components in polymers and materials,[2] or as synths for the preparation of biologically active molecules and peptidomimetic drugs.[3] To fully exploit the versatility offered by AAs, it is important to have efficient and straightforward methods to access structural motifs beyond those exhibited by proteomic AAs. To this end, chemists have developed a myriad of efficient and straightforward strategies for the synthesis of unnatural amino acids (UAAs) using enzymatic or transition-metal-catalysed processes.[4] A less explored approach, however, is the use of methods exploiting one-electron pathways.[5]

Radical chemistry offers exciting and highly attractive approaches to access new chemical space in a rapid fashion. This is due, in part, to the plethora of synthetic precursors available to generate open-shell species.[6] During the last decade, mild and versatile radical-based methods have been developed to access a wide range of structural motifs in organic synthesis. An application that is becoming ever more prominent is the use of radical strategies to gain rapid access to libraries of novel UAAs or for the site-selective modification of peptides and proteins.[7]

The aim of this Minireview is to provide a broad overview of the advances made in this field during the last decade. For clarity, it is organised in two sections: de novo synthesis, which deals with strategies to access the main three classes of AAs, namely α-, β-, and γ-AAs, and modification, which describes strategies for the derivatisation of canonical and non-canonical AAs. Furthermore, to avoid significant overlap with recent reviews,[8] special attention is paid to non-photoredox-mediated strategies.

2. De Novo Synthesis

Arguably, α-, β-, and γ-AAs constitute the most common types of AAs. In this section, strategies to access these key structural motifs, either in their racemic or enantiomeric forms, are discussed.

2.1. α-Amino Acids

The synthesis of α-AAs using radical reactions can be broadly classified in three categories: addition of an open-shell species to an imine, which can afford racemic or enantiomeric α-AAs depending on the N-substituents of the imine, C(sp³)−H aminations, or use of CO₂ as a C₃ building block.

2.1.1. Addition to Imines

In 2015, Inoue and co-workers reported a BEt₃-mediated procedure for the synthesis of α,β-diamino acid derivatives starting from α-aminoacyl tellurides 1, which can be readily prepared from the corresponding α-AA and diphenyl telluride, as well as ethyl glyoxylate oxime 2 (Scheme 1A).[9] The key step in this reaction is the homolytic cleavage of the C–Te bond in 1, which leads to the formation of an acyl radical intermediate that undergoes facile decarbonylation to deliver a highly stabilised α-amino radical species.[10] Subsequent addition to 2, followed by protonation furnishes the desired α,β-diamino acid. This strategy can also be applied to the synthesis of γ-AAs if acrylates are used as radical acceptors.

A more direct method for the synthesis of α,β-diamino esters using α-AAs and 2 was reported by Ye and co-workers (Scheme 1B).[11] This light-mediated procedure employs an acridinium-based photocatalyst (PC) to access the key α-amino radical intermediate directly from α-AAs.

Mariano, Wang, and co-workers reported in 2019 a light-mediated, decarboxylative procedure for the synthesis of C-glyco-α-AAs (Scheme 2).[12] This catalyst-free method proceeds by the addition of C-centred glycosyl radicals, generated from the corresponding N-hydroxyphthalimide (NHP) derived redox-active esters (RAEs) 3, to α-imino ester derivatives 4. The transformation shows a broad scope in terms of both the saccharide and imine motifs. Although the reaction is completely stereoselective regarding the α-oxo stereocentre, only modest levels of diastereoselectivity were observed for the α-amino stereocentre. The key step of the reaction is a photoinduced electron transfer (PET) between the excited Hantzsch ester (HE) and the NHP-derived RAE to afford glycosyl radical intermediate 5, which subsequently

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Minireviews

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adds to 4 to deliver the targeted C-glyco-α-AA after a hydrogen atom transfer (HAT) step.

Recently, Nishikata, Yazaki, Ohshima, and co-workers developed a Cu-catalysed method for the synthesis of quaternary α-AA derivatives from tertiary alkyl halides and AA-derived Schiff bases (Scheme 3A).13 The transformation proceeds through the radical–radical coupling of a tertiary and anazaallyl radical, both generated by the Cu catalyst. This mild method shows a broad scope and functional group (FG) tolerance.

Scheme 1. Syntheses of α,β-diamino esters.

Scheme 2. Synthesis of C-glyco-α-AAs.

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Adrián Gómez-Suárez graduated from the University of Santiago de Compostela in 2009. In 2014 he was awarded his PhD from the University of St. Andrews for his studies on the chemistry of dinuclear gold complexes under the supervision of Prof. Steven P. Nolan. For postdoctoral research he moved to the WWU Münster to work with Prof. Frank Glorius. In 2018 he started his habilitation at the Bergische Universität Wuppertal, under the mentorship of Prof. Stefan F. Kirsch. His research interests include radical chemistry, catalysis, and photochemistry.
An alternative metal-free method for the synthesis of quaternary α-amino esters was reported by Nam and Jang (Scheme 3B).[14] This procedure employs BEt₃ as a radical initiator to generate open-shell species from alkyl iodides. The generated radical intermediates undergo radical addition to hydrazine derivatives of α-keto esters to produce the desired products. Notably, this transformation works better when using H₂O as the solvent and under air. The authors rationalised this by the hydrophobic effect and hydrogen bonding between the interface of the water and the substrate, which would accelerate the reaction.

Since it is possible to form imines in situ from a combination of amines and carbonyl motifs, multicomponent reactions for the synthesis of α-AAs have also been designed.

In 2018, Lu, Gong, and co-workers presented a three-component, one-pot reaction for the formation of masked α-amino aldehydes (Scheme 4A).[15] The reaction proceeds through an AIBN-initiated radical chain mechanism, and affords the desired α-amino aldehydes after the addition of an α-oxo radical, generated from 1,3-dioxolane, to an in situ generated imine.

Later, the same group developed a light-mediated, redox-neutral strategy to access γ-azide-α-AAs (Scheme 4B).[16] This three-component, one-pot reaction proceeds through the addition of an electrophilic azide radical (generated from Me₃SiN₃) to an enol ether to afford nucleophilic radical intermediate 6, which can subsequently react with the imine to deliver the targeted product.

The use of chiral auxiliaries on the N-substituent of the imine allows for the formation of enantiopure α-AAs through diastereoselective radical additions. The most common auxiliary employed in these processes is the mesitylsulfinyl group, as its large steric demand allows for high levels of diastereocntrol during the key radical-addition step.

In 2018, Baran and co-workers reported a Ni-catalysed method for the diastereoselective synthesis of α-AAs by using NHP-derived RAEs as radical precursors (Scheme 5A).[17] These RAEs undergo facile single-electron transfer (SET) with Ni²⁺ species to give alkyl radicals, which add diastereoselectively to chiral imine 7 to afford the targeted α-AAs. The utility of this mild and scalable method was highlighted by its impressive scope, with broad FG tolerance, including the derivatisation of natural products and pharmaceuticals.

Recently, Kärkäs and co-workers expanded this strategy by using photoredox catalysis (PRC; Scheme 5B).[18] This procedure complements Baran’s method by using simpler radical precursors (carboxylic acids vs. NHP-derived RAEs) and employing an organic-based photocatalyst.

In 2018, Shenvi and co-workers reported the use of unactivated olefins as alkyl-radical precursors for the synthesis of α-AAs (Scheme 6). α-AAs through diastereoselective radical additions. The most common auxiliary employed in these processes is the mesitylsulfinyl group, as its large steric demand allows for high levels of diastereocntrol during the key radical-addition step.
2.1.3. CO$_2$ as a C$_1$ Building Block

Recently, the use of CO$_2$ as a C$_1$ source to access α-AAs has attracted increasing attention.[21] Studies by Prikhod’ko, Walter, Py et al.,[22] as well as Radosevich and co-workers[23] and Mita and Sato,[24] focused on the use of (super)stoichiometric metal reductants (SmI$_2$, Mg, or Mn) to generate α-amino radicals that can react with CO$_2$ to access α-AAs. A similar strategy was reported by the groups of Yu[25] and Walsh,[26] through the use of PRC. In this case, the key α-amino radicals, generated after single-electron reduction of imines, are further reduced to the corresponding carbanion, which is subsequently trapped by CO$_2$ to yield α-AAs (Scheme 8A).

Scheme 8. CO$_2$ as a C$_1$ building block to access α-AAs.

2.1.2. C(sp$^3$)–H Amination

Zhang and co-workers reported a Co-catalysed approach for the synthesis of α-AAs by intramolecular radical C(sp$^3$)–H aminations (Scheme 7).[20] In this work, a Co$^{III}$-porphyrin complex was used to generate an N-centred radical, which, through a 1,6-HAT process, generates a C-centred radical. Subsequent intramolecular α-amination affords the targeted cyclic α-AA. The selectivity towards the less hydridic C–H bond is achieved via the formation of a nucleophilic Co$^{III}$-nitrene radical intermediate. The use of enantiopure starting materials allows for high levels of diastereorecontrol over the reaction.

Scheme 7. α-AA synthesis by C(sp$^3$)–H amination.

2.1.1. Ni-catalysed synthesis of α-AAs – Baran 2018

NiOAc (25.0 mol%) Zn (3.0 equiv.) NMP (0.2 M) RT, 6 h

Scheme 5. Enantiopure α-AAs by radical decarboxylations.

Scheme 6. Synthesis of α-AAs by MHAT.

Scheme 8A. α-AA synthesis via reductive coupling of imines with CO$_2$.

Scheme 8B. α-AA synthesis via photoredox-activation of CO$_2$ in continuous flow – Jenckens 2016.

Scheme 8C. α-AA synthesis via carboxylation of unactivated alkenes with CO$_2$ – Yu 2020.
Finally, the groups of Jamison[27] and Yu[28] have reported the synthesis of \(\alpha\)-AAs by \(\alpha\)-C–H functionalisation. Whereas Jamison’s strategy proceeds by a radical–radical coupling of \(\alpha\)-amino radicals with \(\text{CO}_2\), Yu’s proceeds by an intramolecular 1,5-HAT, which generates the key \(\alpha\)-amino radical that can be subsequently reduced to the corresponding carbanion and trapped by \(\text{CO}_2\) (Scheme 8B,C).

### 2.2. \(\beta\)-Amino Acids

The synthesis of \(\beta\)-AAs by radical methods remains highly rare.

In 2018, Jahn and co-workers reported a diastereoselective strategy for the asymmetric synthesis of \textit{anti-}\(\beta\)-amino-\(\alpha\)-(aminoxy) esters and amides (Scheme 9).[29] The reaction proceeds through a polar, asymmetric azo-Michael addition of lithium amides onto \(\alpha,\beta\)-unsaturated carboxylic acid derivatives, followed by a diastereoselective radical recombination with the persistent free radical TEMPO, which acts as an oxygen source. The resulting oxygen protecting group (TEMP) is stable towards acidic, basic, hydric, and hydrogenolytic conditions, but can be readily deprotected using zinc and acetic acid. Remarkably, these conditions are compatible with \(\text{BuMe}_2\text{Si}\) (TBDMS) or Boc protecting groups without causing epimerization.

Recently, Zard reported the synthesis of \(\beta\)-AAs using xanthates as radical precursors (Scheme 10).[30,31] This lauroyl peroxide promoted transformation proceeds by the radical addition of \(\beta\)-phthalimido-\(\alpha\)-xanthylpropionic acid derivatives to vinylic olefins or heteroaromatic molecules. When using the free acid, spontaneous decarboxylation of the intermediate occurs in some cases, thereby leading to the formation of \(\beta\)-heteroarylethylamines.

### 2.3. \(\gamma\)-Amino Acids

The main strategy for the synthesis of \(\gamma\)-AAs is the addition of \(\alpha\)-amino radicals to acrylate derivatives. The key nucleophilic \(\alpha\)-amino radical species can be accessed by several approaches, such as decarboxylation of \(\alpha\)-AAs,[32] proton-coupled electron-transfer (PCET) reduction of imines,[33] or HAT to \(\alpha\)-AAs or benzylic amines.[34]

![Scheme 9. Asymmetric synthesis of \(\beta\)-amino-\(\alpha\)-(aminoxy) esters and amides.](image)

![Scheme 10. \(\beta\)-AA synthesis using xanthates.](image)

![Scheme 11. Syntheses of \(\gamma\)-AAs by radical decarboxylations.](image)
In 2014, MacMillan and co-workers reported their seminal work on the photoredox-mediated Giese reaction (Scheme 11A).[32a] By employing an Ir-based PC, it is possible to induce a single-electron oxidation of α-amino carboxylates to generate acyloxy radicals, which undergo rapid decarboxylation to afford α-amino radical intermediates. These intermediates subsequently add to acrylates to deliver the targeted γ-AAs. Variations of this method were later reported by the groups of Rüping[32b] and König[32c] with the latter using NHP-derived RAEs as radical precursors (Scheme 11B,C). In addition, MacMillan and co-workers have exploited this approach to accomplish peptide macrocyclisations[32e] and site-selective bioconjugation of proteins.[32f]

Recently, Yoshimi and co-workers expanded MacMillan’s procedure for the synthesis of ring-constrained γ-AAs by a radical addition/intramolecular radical cyclisation sequence (Scheme 11D).[32d]

Dixon and co-workers reported an elegant strategy for the synthesis of primary α-tertiary amines by the α-C@H functionalisation of α-secondary amines (Scheme 12A).[33] This bio-inspired method proceeds through the reaction of quinone with primary amines to generate ketimine intermediates in situ, which can readily react with C-centred nucleophiles. The versatility of this transformation was demonstrated by its broad scope. In addition, it is possible to expand this concept to the synthesis of γ-AAs. It was shown that the in situ generated ketimine intermediate can be reduced by HE and an excited PC through a PCET process to produce a nucleophilic α-amino radical, which can then undergo addition to acrylates to yield γ-AAs.

Recently, Cresswell and co-workers reported a new method based on PRC for the synthesis of α-tertiary amines by the α-C–H alkylation of unmasked primary amines with different Michael acceptors (Scheme 12B).[34] The reaction is promoted by an organophotocatalyst and “Bu₂NN₃ as the HAT catalyst. The authors demonstrated the potential of this transformation with an impressive scope and showing its applicability in continuous-flow syntheses.

In 2017, Xie and Zhu reported a diastereoselective strategy for the synthesis of five-membered heterocyclic γ-AAs (Scheme 13).[35] The key step in this light-mediated method is a HAT reaction between a thyl radical and the benzyl C–H bond of a benzyl-protected amine, which results in the formation of a benzyl α-amino radical intermediate. A subsequent 5-exo-trig radical cyclisation onto an α,β-unsaturated carbonyl moiety delivers the targeted γ-AA. A variety of heterocyclic γ-AAs were synthesised in moderate to good yields and diastereoselectivities.

Following the seminal work by Phipps and co-workers on asymmetric Minisci-type reactions,[36] Zheng and Studer reported an elegant approach for the synthesis of enantioenriched γ-AAs (Scheme 14).[37] This light-mediated, three-component radical cascade reaction proceeds in a highly chemo-, regio-, and enantioselective fashion. The key for the reaction to succeed is the philicity of the open-shell intermediates: SET reduction of bromoacetate derivative 9 generates electrophilic radical intermediate 10, which subsequently adds to the electron-rich N-vinylacetamide. This affords a nucleophilic α-amino radical species 11, which adds to the activated electron-poor heteroarene. The is the enantiodetermining step of the reaction and it is controlled by the use of a chiral Brunsted acid catalyst.
3. Modifications

This section highlights radical-based modifications of canonical and non-canonical AAs. Cysteine derivatisations, which have recently been reviewed,[8b] as well as deaminative lysine modifications, which are highly rare, are not included.[38]

3.1. Aspartic (Asp) and Glutamic Acid (Glu)

The primary carboxylate motif present at the β- or γ-position of Asp and Glu, respectively, can be exploited through radical decarboxylative processes to provide a wide range of UAAs. These transformations can be mediated by light or transition metals.

3.1.1. Light-Mediated Procedures

Building upon the radical spirocyclisation procedure developed by Guindeuil and Zard,[39] as well as the use of NHP-derived RAES as radical precursors by Schermann and Overman,[40] Reiser and co-workers developed in 2013 a procedure for the synthesis of (spiro)annelated furans using PRC.[41] To demonstrate the utility of this strategy, the authors carried out the synthesis of (S)-(+)−lycoperic acid in seven steps, using commercially available 5-bromofurfural and (S)-Asp dimethyl ester as starting materials (Scheme 15A).

Interestingly, in the proposed reaction mechanism, Ir(ppy)₂(dtbbpy)PF₆ interacts with the phthalimide moiety of the substrate through an energy-transfer event, rather than the more common SET process.

In 2014, Jamison and co-workers developed a light-mediated method for the synthesis of polycyclic quinoxaline derivatives using PhI(O₂CR)₂ reagents, which are readily accessed from the corresponding carboxylic acids and PhI-(OAc)₂ as RAES.[42] To further highlight the versatility of this procedure, the authors conceived the synthesis of UAAs by reacting Asp and Glu derivatives with the quinoxaline core under photoredox conditions, which provided the target products in moderate yields (Scheme 15B).

Finally, Fu and co-workers reported a photoredox-mediated synthesis of chiral UAAs using NHP-derived Glu and Asp RAES as radical precursors (Scheme 15C).[43] The key step of the method is the quenching of the excited Ru-based PC by either HE or (Pr₂NEt)₂ to generate a highly reducing Ru⁺ species, which can engage in SET with the RAE to deliver the required alkylic radical. This can then react with Michael acceptors,[43a] alkynyl sulfones,[43a] or diselenides[43b] to give the targeted UAA.
3.1.2. Transition-Metal-Mediated Procedures

The groups of Weix[44] and Baran[45] independently reported in 2016 that NHP-derived RAESs easily react with Ni\textsuperscript{I} species by SET, thereby inducing the formation of alkyl radicals, which can be subsequently engaged in cross-coupling reactions.\[46\] Since then, Baran and co-workers have demonstrated that Fe- and Cu-based catalysts are also competent in these processes.\[47,48\]

This highly versatile and powerful strategy has been applied to the modification of Asp and Glu derivatives through arylation,\[44,47,49\] alkylation,\[45\] alkenylation,\[50\] alkylation,\[51\] borations,\[52,53\] Giese reactions or Barton decarboxylation,\[53\] and even C-isotope exchanges.\[54\] These reactions afford moderate to good yields of the desired UAA, and, in many cases, proceed without racemization of the \(\alpha\)-amino stereocentre (Scheme 16). This strategy can also be exploited for the selective modification of Asp and Glu residues on peptides.

Baran and co-workers reported the use of CITU as a coupling reagent to generate RAESs.\[55\] This commercially available reagent opened the door for the synthesis and modification of peptidic RAESs, both in the solid phase and in solution, to afford a straightforward procedure for the introduction of non-natural side chains in peptides (Scheme 17A).

Recently, Weix and co-workers reported a sophisticated Ni-catalysed decarboxylative cross-coupling method for the synthesis of ketones, starting from two different types of ester derivatives, such as NHP-derived RAESs and \(S\)-2-pyridyl thioesters (Scheme 17B).\[56\] Both esters can be readily accessed from carboxylic acids, which broadens the scope of the method. The key to the reaction is that one of the esters acts as a radical precursor—RAE—while the other acts as an acyl surrogate—thioester. The high versatility and FG tolerance of the method was showcased by solid-phase modification of several Asp and Glu residues on peptides.

3.2. Dehydroalanine (Dha)

Dha derivatives are highly versatile synthons that usually behave as Michael acceptors.\[57\] Dha modifications by radical pathways can be broadly classified depending on whether the reaction affords a racemic or enantioenriched \(\alpha\)-AA.

3.2.1. Racemic Procedures

Several methods have been developed for Giese-type reactions of Dha using RAESs,\[58\] carboxylic acids,\[59\] imines,\[60\] ketals,\[61\] trifluoroborates,\[62\] alkyl halides,\[63\] or thioesters\[64\] as radical precursors.

In 2018, Baran and co-workers reported a modification of his Ni-catalysed Giese reaction,\[59\] which allowed the use of NHP-derived RAESs as radical precursors for the synthesis of DNA-encoded libraries.\[58\] The procedure must be carried out under highly diluted conditions (1 mM) and on a small scale (0.01 mmol) to account for the poor solubility of DNA.
in water. The reaction showed broad scope, with examples using a Dha derivative as a radical acceptor affording a wide range of UAAs in high yields (Scheme 18A).

Recently, Shah et al. reported a light-mediated, metal-free, decarboxylative method for the synthesis of UAAs (Scheme 18B). This one-pot, two-step procedure uses readily available 1, 2, or 3 carboxylic acids as radical precursors, and an N-Boc-Dha derivative (12) as radical acceptor, to deliver a wide range of unprotected UAAs.

Organoborate species are versatile radical precursors in PRC, which have been employed for the modification of Dha derivatives. de Bruijn and Roelfes reported a light-mediated alkylation of Dha residues in antimicrobial peptides using an Ir-based PC and trifluoroborate derivatives (Scheme 19A). Molander and co-workers reported a photocatalysis-mediated three-component reaction for the synthesis of α-fluorinated UAAs using alkyl trifluoroborates, 12, and Selectfluor (Scheme 19B). The key for this procedure to proceed is the electron-withdrawing character of the N-protecting group in 12, as it is essential to favour the attack of the nucleophilic alkyl radical on 12, rather than its engagement in a fluorne-transfer reaction with Selectfluor.

Dixon and co-workers have recently reported a couple of light-mediated methods for the modification of Dha derivatives using imines and ketals as radical precursors. The first is a three-component procedure where in situ generated imines are reduced by an Ir-based PC to generate α-amino radicals that add to 12, thereby affording UAAs bearing 1,3-diamine motifs (Scheme 20A). Although the products are generated in good to excellent yields, the method displays low to moderate diastereoselectivities.

Lewis acids can be used to generate oxocarbenium species from ketals. Dixon and co-workers have cleverly exploited this reactivity to develop a procedure for the synthesis of α-tertiary ethers (Scheme 20B). The key step in this process is the reaction of Me3SiOTf with a ketal to in situ generate an oxocarbenium species, which is subsequently reduced by an Ir-based PC to afford α-alkoxy radicals. These add to 12 to afford UAAs. As in the previous transformation, low to moderate diastereoselectivities are observed.

Recently, several methods have been reported for the modification of Dha using alkyl halides. In 2016, Davis and co-workers designed a bio-orthogonal strategy for the side-chain modification of proteins (Scheme 21A). The procedure proceeds through the selective addition of alkyl radicals, generated from a combination of alkyl bromides/iodides and NaBH4, in aqueous solution, to Dha residues.

In 2019, Brandhofer and García Mancheño reported a straightforward light-mediated method for the derivatisation of Dha derivatives (Scheme 21B). Although the majority of the reactions were carried out using fluorinated alkyl halides, arylsulfonyl chlorides or NHP-derived RAEs are also suitable precursors in this Ru-catalysed process. To highlight the versatility of the method, peptide modifications containing Dha residues were carried out.
Finally, Scanlan and co-workers reported various chemo-selective strategies for the synthesis of cysteinyl peptide thioesters through the addition of thioesters to Dha residues. This transformation can be efficiently accomplished by either ionic or radical-mediated pathways, and can be carried out under aqueous buffered conditions at neutral pH. The radical approach uses UVA light and 4′-methoxyacetophenone as a photosensitizer to promote the generation of the key thyl radical intermediate.

### 3.2.2. Diastereoselective Procedures

The Beckwith-Karady alkene 13 is a chiral Dha derivative developed in the early 1990s. Over the years, 13 has been employed to establish highly diastereoselective routes for the synthesis of UAAAs. With the advent of PRC, the use of 13 in radical processes has experienced a resurgence. As a result, methods using alcohols, heteroaryl halides, amines, C(sp³)-H bonds, or α-keto and carboxylic acids as radical precursors have been developed.

In 2017, Jui and co-workers reported a practical and scalable light-mediated synthesis of heteroaryl α-AAs (Scheme 22A). The key step in this photoredox-mediated procedure is the single-electron reduction of heteroaryl halides by a highly reducing IrIII species, generated from the excited-state quenching of an IrIII-based PC with HE. The method displays a broad FG tolerance and scope, and it is readily amenable to large-scale synthesis. Examples using both 12 and 13 as radical acceptors were shown.

A year later, the same group reported the synthesis of UAAAs and peptides by a light-mediated aminoalkylation method (Scheme 22B). The key step of the transformation is the generation of an amine radical cation, by the oxidation of tertiary amines by an excited Ir-based photocatalyst, which, after deprotonation, affords α-amino radicals that can engage...
in radical additions to 13. This method can be applied for the late-stage modification of complex molecules, as well as, for the modification of Dha residues in peptides.

In 2018, Gaunt and co-workers disclosed an elegant light-mediated strategy for the synthesis tertiary alkylamines. This three-component method combines dialkyl amines with carbonyl compounds (either aldehydes or ketones) to give iminium ions in situ. These ions are subsequently reduced by a PC to afford α- amino radical intermediates that can engage in Giese-type reactions with a wide range of Michael acceptors. The procedure presents a broad scope and FG tolerance, with several examples of using 13 as a radical acceptor, and affords UAA derivatives in moderate to good yields and excellent diastereoselectivities (Scheme 23). The versatility of this method was later highlighted by the rapid synthesis of several natural products.

Recently, the groups of Gómez-Suárez, Wang, and Shubert independently reported a photoredox-mediated decarboxylative synthesis of UAAAs using 13 as a radical acceptor (Scheme 24). Whereas the procedure developed by Gómez-Suárez and co-workers presents a broad scope with a variety of aromatic and aliphatic α-keto acids, as well as aliphatic carboxylic acids as radical precursors, the approach developed by Wang and co-workers allows the synthesis of α-deuterated UAAAs and the method developed by the Shubert group facilitates the modification of peptides under metal-free conditions. All three methods allow the straightforward synthesis of a wide range of UAAAs in good to excellent yields and high diastereoselectivities.

### 3.3. Glycine (Gly)

The direct C–H functionalisation of Gly has been a greatly sought-after route for the facile synthesis of UAAAs. In 2015, You and co-workers reported a Ni-catalysed strategy for the C–H benzylolation of Gly (Scheme 25). This transformation proceeds by coordination-activation of a 2-picolinamido -amino ester with a high-valent NiIII species. The system avoids formation of an imine intermediate through direct trapping of a benzyl radical species by the amine. A wide range of β-aromatic α-AAs can be accessed by using a variety of (hetero)aryl methane, arylenes, or aryl ethers.

![Scheme 23. Three-component synthesis of UAAAs.](image)

![Scheme 24. Decarboxylative functionalisations of 13.](image)

![Scheme 25. Ni-catalysed C–H benzylolation of Gly.](image)
Very recently, Xu and co-workers reported a remarkable light-mediated, catalyst-free C–H alkylation of Gly using alkylpyridinium salts as alkyl radical precursors (Scheme 26).\(^{[73]}\) The key step in the reaction is the formation of an electron-donor-acceptor (EDA) complex between Gly and the pyridinium salt, which upon irradiation undergoes an intermolecular charge transfer to generate an alkyl radical—after homolytic cleavage of the N–C bond—and a Gly-derived α-amino radical. These two species can undergo a radical–radical coupling, thereby allowing the straightforward derivatisation of Gly under very mild conditions. This was highlighted by the selective derivatisation of Gly residues on several peptides.

A different strategy for the modification of Gly involves its derivatisation in situ to generate imines that can be further functionalised.\(^{[74]}\)

In 2015, Wu and co-workers described a dual photoredox/Co-catalysed Gly modification that takes advantage of this approach (Scheme 27 A).\(^{[74a]}\) In this procedure, the in situ generated imine is further derivatised by a nucleophilic attack from a β-keto ester/indole derivative to access highly functionalised UAAs.

Recently, Yang et al. have also exploited this strategy to develop an enantioselective Gly derivatisation procedure (Scheme 27 B).\(^{[74b]}\) This light-mediated method uses a dual photoredox/Cu-catalysed strategy to generate the key imine intermediate, which is subsequently involved in a proline-catalysed asymmetric Mannich-type reaction to produce a wide range of enantioenriched UAAs.

\[ \text{Scheme 26. Deaminative Gly modification.} \]

\[ \text{Scheme 27. Gly derivatisations by in situ generated imines.} \]

### 3.4. Leucine (Leu)

Leu modifications by C–H functionalisation of the isopropyl motif through HAT reactions have been targeted.

In 2015, Di Rocco, Britton, and co-workers reported a light-mediated procedure for the preparation of a fluorinated Leu derivative employed in the synthesis of a cathepsin K inhibitor used in the treatment of osteoporosis (Scheme 28 A).\(^{[75]}\) The reaction involves photodissociation of a decatungstate catalyst, which produces an excited-state intermediate that abstracts a hydrogen atom from the γ-position of Leu. The newly formed C-centred radical undergoes subsequent fluorine atom transfer with N-fluorobenzensulfonylimide (NFSI).

Later, the groups of Britton and Schaffer partnered to expand this strategy to the synthesis of 18F-labeled peptides, by the C–H fluorination of Leu residues in unprotected, unmodified peptides (Scheme 28 B).\(^{[76]}\) The selectivity of this reaction for the γ-position of Leu, combined with the tolerance of charged, polar, and hydrophobic AAs is particularly notable.

In 2016, Chen and co-workers developed a system for the functionalisation of tertiary alliphatic C–H bonds of several substrates, including Leu (Scheme 28 C).\(^{[77]}\) By using Zhdankin’s reagent, Ru(bpy)_3Cl_2, and visible-light irradiation, the reaction allows azidation of Leu under mild conditions. By adding LiCl or Bu_4NBr to the reaction, halogenated Leu derivatives can also be prepared.

Later, the Chen group expanded this method to the hydroxylation and amidation of tertiary and benzylic C–H...
bonds, including Leu, thereby suggesting a possible efficient method for the late-stage modification of biomolecules.\textsuperscript{[78]}

### 3.5. Phenylalanine (Phe)

Phe presents two main vectors for derivatisation by radical-based methods: the benzylic C–H bond and the aromatic ring.

Several reports have dealt with the fluorination of the benzylic C–H bond in Phe using Selectfluor\textsuperscript{®} as the fluorinating source.\textsuperscript{[79]} Among them, the most relevant is the light-mediated fluorination of Phe residues on peptides reported by Lectka and co-workers in 2016 (Scheme 29).\textsuperscript{[80]} Although the method afforded poor diastereoselectivities, the products could be successfully separated using flash chromatography. In peptides presenting a wide range of C–H bonds susceptible to HAT reactions, the reported transformations showcased a remarkable selectivity towards the fluorination of benzylic C–H bonds in Phe residues.

Another strategy to modify Phe is the functionalisation of aromatic C–H bonds through amination reactions. Ritter and co-workers reported a method for \textit{para}-selective C–H amination by charge-transfer-directed radical substitution of aromatic systems in a one-pot, two-step reaction (Scheme 30A).\textsuperscript{[81]} The method showed a broad substrate scope and was used for the selective \textit{para}-amination of phthalimide-protected Phe in reasonable yield and with complete stereoretention.

Recently, Leonori and co-workers expanded the scope of \textit{para}-selective C–H aminations using PRC (Scheme 30B).\textsuperscript{[82]} The key step of the process is the generation of a highly electrophilic aminyl radical, which selectively adds at the \textit{para} position of the arene. The method can be applied for the selected derivatisation of Phe residues in tetrapeptides.
methyl acrylate and a PC (Scheme 32). The reaction can be carried out with other Michael acceptors, and represents an innovative process for activation of the β-C(sp³)–H bond of Trp for residue-specific peptide modifications.

3.7. Vinyl and (Homo)allylglycine

Despite its use as a starting material for several reactions, the application of vinyl-Gly as a substrate in radical reactions has scarcely been investigated, possibly because of the presence of a labile, tertiary, and allylic hydrogen atom. To overcome this limitation, Zard and co-workers investigated the intermolecular radical addition of xanthates to protected vinyl-Gly. This afforded UAAs in respectable yields and with complete stereoretention at the α-amino position (Scheme 33).

In 2019, Jackson and co-workers reported a radical hydrofluorination of unsaturated AAs, including allyl-Gly derivatives (Scheme 34). This study expanded the scope of the radical method reported by Barker and Boger for the addition of HF to unactivated alkenes. The combination of Fe³⁺/NaBH₄ and Selectfluor or NaN₃ resulted in enantioselectively pure protected fluorinated or azidated AAs in reasonable yields, without the need to remove protecting groups or use additional toxic reagents.

In addition, the groups of Shenvi as well as Ruffoni and Leonori have reported several methods where allylglycine is part of the reaction scope. Its modification usually proceeds smoothly and in moderate to good yields. Finally, Deming and co-workers have shown that it is possible to functionalise alkene side chains on poly(homoallylglycine) through radical thio-ene reactions. The incorporation of homoallyl Gly residues into polypeptides allowed alteration/control of the peptide-chain conformations.

4. Conclusions and Outlook

During the last decade, significant advances have been made in the synthesis and modification of AAs by using radical-based strategies. This progress has been driven by the development of milder and more general catalytic strategies for the generation of open-shell species. As a result, straightforward procedures for the synthesis of α-, β-, and γ-AAs, both in racemic and enantiopure forms, as well as methods for the chemoselective modification of AA residues in peptides, have been reported. Despite this progress, there is still room for improvement. For example, the main use of α-AAs in syntheses involving radical chemistry is as readily available...
precursors of ω-amino radicals by decarboxylative transformations,[63] which obliterates their stereocchemical information. Moreover, although substantial developments have been made for site-selective peptide modification using radical chemistry, bioorthogonal methods remain scarce. At the current pace of innovation, it is expected that methods capable of addressing these challenges will emerge in the near future. The radical-based synthesis and derivatisation of AAs is an exciting field of research with a healthy present and a brilliant future, full of opportunities for innovation.

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Conflict of interest

The authors declare no conflict of interest.

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