1	Fluorescent amino acids as versatile building blocks for chemical biology
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3	Zhiming Cheng <sup>1</sup> , Erkin Kuru <sup>2</sup> , Amit Sachdeva <sup>3</sup> , Marc Vendrell <sup>1†</sup>
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5	<sup>1</sup> Centre for Inflammation Research, University of Edinburgh, Edinburgh, UK
6	<sup>2</sup> Department of Genetics, Harvard Medical School, Boston, USA
7	<sup>3</sup> School of Chemistry, University of East Anglia, Norwich, UK
8	
9	<sup>†</sup> email: <u>marc.vendrell@ed.ac.uk</u>
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11	Abstract   Fluorophores have transformed the way we study biological systems, enabling non-
12	invasive studies in cells and intact organisms, which increase our understanding of complex
13	processes at the molecular level. Fluorescent amino acids (FlAAs) have become an essential
14	chemical tool because they can be used to construct fluorescent macromolecules, such as
15	peptides and proteins, without disrupting their native biomolecular properties. Fluorescent and
16	fluorogenic amino acids with unique photophysical properties have been designed for tracking
17	protein-protein interactions in situ or imaging nanoscopic events in real-time with high spatial
18	resolution. In this Review, we discuss advances in the design and synthesis of FlAAs and how
19	they have contributed to the field of chemical biology in the past 10 years. Important areas of
20	research that we review include novel methodologies to synthesize building blocks with
21	tunable spectral properties, their integration into peptide and protein scaffolds using site-
22	specific genetic encoding and bio-orthogonal approaches, and their application to design novel
23	artificial proteins as well as to investigate biological processes in cells by means of optical
24	imaging.

# 26 [H1] Introduction

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Fluorescence-based techniques have revolutionized our understanding of biological processes, as they allow researchers to examine the localization, trafficking and activity of biomolecules in cells, tissues and even whole intact organisms in a non-invasive manner.<sup>1-4</sup> Many natural peptides and proteins have key roles in various biological processes; however, directly visualizing these molecules is hampered by the fact they generally do not exhibit strong fluorescence emission. Whereas this low 'background' fluorescence boosts the sensitivity of spectroscopy and imaging experiments, its weak levels highlight the need for labelling

strategies that can facilitate the detection of these macromolecules. Fluorescent labelling of 35 peptides and proteins has been successfully used in numerous ways, from the widely used 36 expression of fluorescent protein fusions (such as green fluorescent protein<sup>5</sup> (GFP) and 37 subsequent derivatives<sup>6,7</sup>) to the fusion of proteins with self-labelling tags (such as SNAP, 38 CLIP and HALO tags<sup>8-10</sup>) or modification with specific peptide sequences that can bind to 39 small molecules with high affinity (such as FIASH and ReASH<sup>11,12</sup>). All of these approaches 40 41 have led to remarkable biological discoveries, but they typically involve a large structural modification of the peptides or proteins of interest, which can alter their biomolecular 42 43 properties.

Non-natural fluorescent amino acids (FIAAs) have been developed as chemical 44 alternatives for the derivatization of peptides and proteins in a less disruptive manner. Owing 45 to their small size and similarity to the natural residues found in polypeptides, they can be used 46 to fluorescently label macromolecules while retaining their overall function with minimal 47 perturbation of the native protein structure. The toolbox of FIAAs expanded considerably in 48 the early 2000s, partly due to the development of numerous FIAAs<sup>13</sup> with unique optical 49 properties, including environmental sensitivity<sup>14-16</sup>, responsive to metal chelation<sup>17</sup>, tunable 50 fluorescence emission and prolonged lifetime<sup>18</sup>). Importantly, these modular building blocks 51 can be introduced at specific sites in small peptides using solid-phase peptide synthesis<sup>19</sup> 52 (SPPS) or into larger proteins by genetic encoding<sup>20</sup> to generate fluorescent, native-like 53 biomolecules, enabling biological experiments that would be intractable by other means, 54 including analysis of protein conformational changes, monitoring of protein-protein 55 56 interactions and activity studies in live cells.

Over the past 10 years, a substantial increase in the number of versatile synthetic 57 strategies (such as multicomponent reactions<sup>21</sup>, metal-catalysed reactions<sup>22</sup>, light-induced 58 transformations<sup>23</sup> and bioorthogonal chemistry<sup>24</sup>) to fine-tune the chemical structure of 59 fluorophores<sup>25</sup> and natural building blocks has accelerated the design and preparation of new 60 FIAAs. In this Review, we provide an overview of different chemical transformations that have 61 been used to generate novel building blocks with bespoke fluorescent properties and to 62 facilitate their site-specific integration into peptides and proteins. We also discuss how the 63 range of applications of FlAAs has extended to include many biological assays in vitro, in cells 64 and in vivo, which has been facilitated by a rapid expansion in the use of spectroscopic and 65 imaging-based techniques. Finally, we review the versatility of FIAAs from the point of view 66 of an end-user and present several examples of the application of FIAAs in chemical biology, 67

- 68 including molecular recognition studies, live-cell imaging and applications in synthetic biology
- 69 and microbiology (**Figure 1**).

#### 73 [H2] Fluorescent analogues of natural amino acids

Three canonical amino acids, (tryptophan, tyrosine and phenylalanine, are fluorescent and thus might have potential as naturally-occurring fluorophores<sup>26</sup>; however, their optical properties (such as excitation and emission wavelengths, brightness and photostability) are suboptimal for most biological assays. Tryptophan, the most commonly used fluorophore among natural amino acids, absorbs and emits in the ultraviolet range (absorption wavelength  $(\lambda_{abs}) = 280$  nm; emission wavelength  $(\lambda_{em}) = 346$  nm) and its fluorescence quantum yield is  $\sim 20\%^{27}$ .

Initial attempts to improve the fluorescent properties of tryptophan yielded 81 azatryptophans as potential isosteric substitutes in proteins<sup>28</sup>. Although 4-azatryptophans have 82 a longer emission wavelength ( $\lambda_{em}$ = 425 nm) than tryptophan and a large Stokes shift 83 (>130 nm), they are typically less bright than tryptophan. Alternative approaches have involved 84 the preparation of cyanotryptophans, which have improved fluorescence quantum yields 85 (approaching  $\sim 50\%^{29}$ ) owing to the incorporation of the strong electron withdrawing cyano 86 87 group. Cyanotryptophans can be synthesized from commercially available cyanoindoles, and the red-shifted emission wavelength of 6-cyanotryptophan (1a) ( $\lambda_{em}$ = 370 nm) enabled its use 88 for Förster resonance energy transfer (FRET) experiments to study protein–DNA interactions. 89 90 Subsequent studies identified 4-cyanotryptophan (1b) as an amino acid with improved optical properties<sup>30</sup>. A key step in the synthesis of 4-cyanotryptophan is the palladium (Pd)-catalysed 91 incorporation of the cyano group at position 4 of L-tryptophan. Remarkably, the emission 92 maximum of 4-cyanotryptophan is in the blue visible region ( $\lambda_{em} = 405$  nm) and it has a high 93 quantum yield (~80%), good photostability and a long fluorescence lifetime (~13.7 ns), which 94 95 makes it an interesting building block for spectroscopic and microscopic measurements of proteins, such as the study of peptide-membrane interactions<sup>31</sup>. The structural similarity of 96 97 cyanotryptophans to tryptophan also makes them an attractive platform to explore enzymebased reactions, facilitating the preparation of enantiopure tryptophan analogues by 98 99 derivatization at multiple aromatic positions, using, for example, tryptophan synthase S  $(TrpS)^{32}$  or variants of its  $\beta$ -subunit  $(TrpB)^{33}$ . 100

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Extended tryptophan analogues have been also designed by conjugation of homocyclic and heterocyclic systems to the indole moiety, and include tricyclic tryptophan analogues (2)

derived from pyrrolo(iso)quinolines<sup>34</sup>, tryptophan derivatives obtained by hydrogen-mediated 104 coupling of 1,2,3-triazoles to indoles<sup>35</sup> (**3a**) and a series of 2-arylated tryptophans (**3b**) obtained 105 by Pd-mediated C-H functionalization<sup>36</sup>. Cross-coupling reactions (such as Suzuki-Miyaura 106 couplings) have also been effective for preparing C4-substituted tryptophans<sup>37</sup>, although the 107 effect of these substitutions on the optical properties of tryptophan has not been reported. Of 108 note, extended tryptophans have a red-shifted emission wavelength ( $\lambda_{em} = 420-480 \text{ nm}$ )<sup>34-36</sup> 109 and some have been incorporated in *Escherichia coli* dihydrofolate reductase (DHFR) using in 110 *vitro* translation, resulting in minimal disruption of DHFR function<sup>38</sup> and thus demonstrating 111 the potential of these molecules for monitoring conformational changes in macromolecular 112 113 structures.

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#### 115 [H2] Non-natural FIAAs

Despite the remarkable progress in the synthesis of fluorescent analogues of natural amino acids, numerous non-natural FlAAs with different optical properties have been developed (**Figure 2**). The synthetic approaches to generate non-natural FlAAs can be broadly categorized into those that involve appending fluorescent moieties to amino acids and those that involve *de novo* construction of amino acids with integrated chromophores (**Figure 2**).

[H3] Palladium catalysis. Pd-catalysed reactions are one of the most commonly used 121 approaches for the synthesis of non-natural FIAAs. Biphenyl derivatives of phenylalanine (4a) 122 have been prepared by Suzuki coupling of biphenyl boronic acids with iodophenylalanine<sup>39,40</sup>. 123 124 Their emission maximum is red-shifted relative to that of phenylalanine (~340 nm versus 282 nm, respectively) and they have high quantum yields (~70%). Suzuki couplings have also been 125 used in the synthesis of fluorescent D-amino acids (FDAAs), including 4-acetamidobiphenyl 126 groups (4b), which have a slightly longer emission wavelength ( $\sim$ 380 nm)<sup>41</sup>. Heck reactions 127 have been employed to extend the  $\pi$ -conjugation of tyrosine<sup>42</sup>, yielding styryl-containing 128 tyrosine analogues (5) with tunable fluorescence quantum yields (from 10% to 90%) and 129 emission maximathat cover the entire visible spectrum ( $\lambda_{em}$  = 400–800 nm). Sonogashira 130 coupling reactions have been used to append aryl-containing acetylene derivatives to 131 benzoxazol-5-yl-alanine to produce FlAAs (6) with a high quantum yield and emission maxima 132 in ultraviolet and blue wavelengths ( $\lambda_{em} = 340-450$  nm)<sup>43</sup>. Similarly, xanthone structures ( $\lambda$ -133  $_{em}$  = 380 nm) have been incorporated into the alanine core by Negishi couplings<sup>44</sup> (7a). The 134 135 high efficiency of Pd-catalysed couplings (such as the Buchwald-Hartwig reaction) has also 136 facilitated the synthetic optimization of widely used non-natural FIAAs, such as acridon-2-

- 137 ylalanine (ACD) (7b), which is one of the brightest blue-emitting FlAAs reported to date<sup>45</sup>.
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[H3] C-H activation. C-H activation is another synthetic methodology that has contributed 139 substantially to the development of non-natural FIAAs, particularly for the construction of new 140 tryptophan-based amino acids. Trp-BODIPY (8a) was the first BODIPY-containing tryptophan 141 analogue, which has excellent photophysical properties ( $\lambda_{abs} = 500 \text{ nm}$ ;  $\lambda_{em} = 530 \text{ nm}$ ) and is 142 compatible with SPPS<sup>46</sup>. A similar approach was used to prepare Trp(redBODIPY) (**8b**), which 143 has a red-shifted maximum excitation and emission wavelength ( $\lambda_{abs}$  = 560 nm,  $\lambda_{em}$  = 144 590 nm)<sup>47</sup>, and also was extended for the synthesis of styryl-derivatized tryptophan analogues 145 by C–H olefination at the C2 position of the indole moiety<sup>48</sup>. Given that the extension of  $\pi$ -146 conjugation systems often leads to improved photophysical properties, as described above with 147 Heck reactions<sup>42</sup>, this strategy has the potential to yield FIAAs with novel optical properties. 148 Pd-catalysed sp<sup>3</sup> C-H activation has also been used to append green and red-fluorescent 149 BODIPY dyes to the side chains of alanine and phenylalanine (9)  $(\lambda_{em.} = 510-625 \text{ nm})^{49}$  and to 150 the C2 position of tryptophan by a novel ruthenium(II)-catalysed C–H alkylation<sup>50</sup>. 151

152 [H3] Copper and rhodium catalysis. Further examples of metal-catalysed reactions include the Ullmann-type reaction for coupling the fluorophore pyrene to the phenol group of tyrosine 153 (10a) using copper catalysis ( $\lambda_{em}$  = 390 nm; quantum yield ~40%)<sup>51</sup>, or the widely used copper-154 catalysed azide-alkyne cycloaddition (CuAAC) to conjugate standard fluorophores, such as 155 coumarins -( $\lambda_{em}$  = 380 nm), benzothiadiazoles ( $\lambda_{em}$  = 470 nm), fluorescein ( $\lambda_{em}$  = 510 nm), 156 dansyl ( $\lambda_{em} = 520 \text{ nm}$ ), nitrobenzodioxazoles (NBDs;  $\lambda_{em} = 530 \text{ nm}$ ), and naphthalimides ( $\lambda_{em} = 530 \text{ nm}$ ). 157 550 nm), to the side chains of different amino  $acids^{52-54}$  (11). Fluorescent arylalanines (12) 158  $(\lambda_{em} = 400 \text{ nm}; \text{ quantum yield } \sim 60\%)$  have been synthesized via the formation of C–C bonds 159 between several polyaromatic hydrocarbons and alanine using rhodium catalysis<sup>55</sup>. 160

[H3] Subheading. Conventional reactions involving the formation of amides<sup>56-61</sup> (13), 161 carbamates<sup>62</sup> (14) or thioureas<sup>57,58,63</sup> (15) have also been explored for modifying the side chains 162 of different amino acids with various fluorophores. Other reactions include Knoevenagel-like 163 condensation to derivatize aminodicarboxylate  $\beta$ -ketoesters with pyrene<sup>64</sup> (16) or synthesis of 164 a green-fluorescent FlAA (17) ( $\lambda_{em}$  = 540 nm) by coupling 4-dimethylamino-1-naphthalenes to 165 methyl (2S)-5-(dimethoxyphosphoryl)-4-oxo-2-(tritylamino)-pentanoate via a Horner-166 Wadsworth-Emmons reaction<sup>65</sup>. The simplicity of some reactions and the availability of 167 168 properly derivatized fluorescent building blocks has also facilitated the preparation of 169 collections of FIAAs. For example, using Michael addition, a small library of D-cysteine FIAAs (18) with thiol-containing fluorophores was generated<sup>66</sup>, and maleimide-4-aminophthalimide<sup>67</sup> 170 (19) was coupled to L-cysteine. Furthermore, the Fukuyama–Mitsunobu reaction was used to 171 couple 6-acyl-2-naphthylamine to L-serine to produce 3-(6-acetylnaphthalen-2-ylamino)-2-172 aminopropanoic acid (ANAP) (20), a building block with remarkable optical properties ( $\lambda_{em}$ = 173 490 nm, quantum yield ~50%) that is widely used for fluorescent labelling of proteins<sup>68</sup>. NBD-174 like fluorophores have been conjugated by a nucleophilic aromatic substitution (S<sub>N</sub>Ar) reaction 175 to nucleophilic groups (21) in amino acid side chains (such as 3-amino-alanine)<sup>58,69</sup>, whereas 176 phospholyl(borane) amino acids (22) have been produced by nucleophilic substitution of 177 fluorescent phospholide anions with iodo-derivatized amino acids<sup>70</sup>, which enabled the 178 synthesis of a range of phospholyl derivatives with fluorescence properties covering a broad 179 spectrum of emission maxima wavelengths from ultraviolet to green ( $\lambda_{em.}$ = 340–530 nm). 180

[H3] Flavone-based FlAAs. Alternative synthetic approaches have involved the *de novo* 181 design of FlAAs by building fluorophores into an amino acid structure (Figure 3). For 182 example, flavone-based amino acids (23) (Figure 3a) are produced by constructing the 183 fluorophore 3-hydroxychromone from tyrosine by aldol condensation, followed by oxidative 184 cyclization (for example, the Algar–Flynn–Oyamada reaction)<sup>71-73</sup>. Of note, flavone FlAAs 185 show excited-state intramolecular proton transfer (ESIPT) fluorescence, and they only 186 minimally perturb the structure of peptides owing to their small size. Another FIAA with a 187 188 built-in fluorophore, benzoacridone-modified alanine, was synthesized by an Ullmann-type coupling between 3-chloro-2-naphthoic acid and aminophenylalanine, followed by cyclization 189 of 2-aminonapthoic acid to yield green-fluorescent benzoacridone amino acid ( $\lambda_{em} = 500-550$ 190 nm, quantum yield  $\sim 50\%$ )<sup>74</sup>. Furthermore, a non-natural FIAAs have been synthesised by 191 building aryl-substituted pyridyl, pyrazole, benzotriazole and pyrazoloquinazoline (24) 192 heterocyclic chromophores on aspartic acid and asparagine structures ( $\lambda_{em} = 348-460 \text{ nm}$ )<sup>75-78</sup> 193 (Figure 3b). 194

195 **[H3] Coumarin based FIAAs.** The simplicity of the coumarin scaffold (that is, 1-benzopyran-196 2-one) has facilitated the preparation of FIAAs integrating this fluorescent structure. Coumarin-197 based FIAAs can be prepared by Pechmann condensation, either between an amino acid 198 containing a  $\beta$ -ketoester unit and a phenol derivative or between tyrosine and an ethyl 199 acetoacetate derivative<sup>79,80</sup>. Alternatively, 7-(hydroxycoumarin-4-yl)ethylglycine is prepared 198 by Pd-catalysed cross coupling between 2,4-dimethoxyphenylboronic acid and amino acids 199 containing  $\beta$ -ketoester groups<sup>81</sup>. The brightness of coumarin-based FIAAs, with quantum yields reaching 70%, together with their fairly small size, have made them excellent buildingblocks for multiple applications in chemical biology.

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### 205 [H1] Bioactive fluorescent peptides

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### 207 [H2] Studying biomolecular interactions

Peptides are excellent scaffolds for biological studies, as they can be used to monitor 208 highly specific molecular interactions with a broad range of biomolecules. Fluorescent peptides 209 have traditionally been synthesized by coupling fluorophores to reactive groups (initially to 210 amines, carboxylic acids and thiols<sup>82</sup>, but now also to imidazoles<sup>83</sup> or phenols<sup>84</sup>) on the side 211 chains of amino acids or in conveniently placed spacers. However, in some cases, this synthetic 212 approach can alter the native conformation of the peptide or its biological properties (such as 213 functional activity, binding affinity and/or subcellular localization)<sup>85</sup>. One synthetic alternative 214 to minimize the effect of fluorescent labelling is to embed FlAAs within the sequence of the 215 peptides (to avoid the modification of polar groups, such as amines, carboxylic acids, and 216 thiols, which may be crucial for its bioactivity) or to optimize suitable spacers<sup>86</sup>. These 217 methodologies rely on the chemical robustness and flexibility of SPPS, which enables the 218 219 efficient preparation of highly diverse peptides, including those that incorporate non-natural FlAAs<sup>87</sup>. 220

221 [H3] Monitoring protein-protein and protein-DNA interactions. Most biological processes rely on the interaction of proteins with their binding partners. In this context, 222 223 fluorescent peptides have been developed to better understand protein-based interactions at the molecular level and to help in the discovery of new drugs to modulate them. For example, 224 flavone-based solvatochromic FlAAs were incorporated at specific sites in a peptide sequence 225 from the human immunodeficiency virus-1 (HIV-1) nucleocapsid protein during SPPS<sup>71,73</sup>. 226 These FIAAs have hydration-sensitive dual emission maxima ( $\lambda_{em.}$ = 430 nm and 530 nm), 227 which allowed the interaction between the nucleocapsid protein and nucleic acids to be 228 monitored to gain insights into the HIV-1 viral life cycle (Figure 4a). Other fluorescent 229 structures have been explored as solvatochromic amino acids with high sensitivity to their 230 surrounding microenvironments. For example, phthalimide-based amino acids have been 231 integrated into peptide structures to study dynamic protein-protein interactions, as in the case 232 of the PDZ domain<sup>88</sup> (Figure 4b), one of the most abundant protein interaction domains in 233 eukaryotes, or calmodulin<sup>89</sup>, a major calcium-signal transduction protein. Subsequently, novel 234

fluorogenic peptides have been used as conformational reporters of calmodulin activity<sup>90,91</sup>. In 235 this case, calmodulin-binding peptides were first identified by in vitro selection using tRNA 236 carrying an NBD-based amino acid<sup>90</sup>, which was later replaced with a 4-*N*,*N*-dimethylamino-237 1,8-naphthalimide amino acid that has a similar emission wavelength ( $\lambda_{em} = 530$  nm) but a 238 remarkably enhanced fluorogenic behaviour (up to 100-fold fluorescence increase after binding 239 to calmodulin)<sup>91</sup>. Furthermore, a fluorogenic peptide containing the environmentally-sensitive 240 1,5-naphthyridin-2(1H)-one amino acid has been used to obtain ratiometric fluorescence 241 242 measurements at two emission wavelengths (370 nm and 480 nm) in response to conformational changes in calmodulin<sup>61</sup>. 243

[H3] Monitoring peptide-membrane interactions. Fluorogenic peptides can also be designed 244 to study peptide-membrane interactions, which are crucial for peptide-based therapeutics and 245 transfection reagents. Cationic peptides, including antimicrobial peptides, are suitable for this 246 purpose, given their high membrane-binding activity. Fluorogenic analogues of melittin, an 247 active component of honey bee venom, have been used to study the orientation of melittin in 248 lipid-rich membranes, revealing details about the varying insertion depths of different residues 249 (that is, flavone FIAAs) and that melittin is oriented parallel to the surface of cell membranes<sup>72</sup>. 250 251 Similar approaches have been used to study the membrane binding of neuropeptides, which function as messengers between neurons. Analysis of the fluorescence of two analogues of the 252 insect neuropeptide helicokinin I (containing the FIAAs  $\beta$ -(4'-hydroxy-'-benzoyl) (ALB;  $\lambda_{em}$ = 253 360 nm) or 6-dimethyl-amino-2-acyl-naphthalene (ALD;  $\lambda_{em}$  = 530 nm)) interacting with 254 255 dodecylphosphatidylcholine micelles was used to develop a model of helicokinin I binding to membranes<sup>92</sup>. Furthermore, Fmoc-L-4-cyanotryptophan was incorporated in the membrane-256 interacting pH-(low) insertion peptide (pHLIP) during SPPS to measure peptide-membrane 257 binding constants using FRET experiments with the universal membrane stain DiO (ref.<sup>31</sup>) 258 (Figure 4c). 259

[H3] Analytical applications. Fluorogenic peptides have also been used as analytical tools for 260 the fluorometric detection of specific biomolecules. For example, peptide aptamers containing 261 an environmentally-sensitive NBD FIAA were evolved in vitro using ribosome display to 262 identify those that detect verotoxin, a protein that is produced by E. coli and is associated with 263 haemolytic uraemic syndrome<sup>93</sup>. Furthermore, a 7-azatryptophan-containing peptide has been 264 used for detection of haeme levels in cells<sup>94</sup> using FRET-based measurements based on the 265 overlap between the emission spectrum of 7-azatryptophan ( $\lambda_{em.}$ = 400 nm) and the absorbance 266 267 of the haeme group. Peptides containing other fluorophores have also been used in FRET

assays to measure protease activity<sup>69</sup>, including benzoacridone amino acids to detect active caspase 3 (ref.<sup>74</sup>), a key enzyme in the apoptotic pathway.

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# 271 [H2] Live-cell optical imaging

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Optical microscopy has allowed researchers to examine the movement of biomolecules 273 inside the cell with high spatial and temporal resolution. Fluorescent peptides are perfectly 274 suited for optical imaging, as they can target specific proteins in cells and also contain optical 275 reporters (that is, FIAAs) that are easily detected using fluorescence microscopes<sup>95</sup>. To speed 276 up the detection of pulmonary infections, fluorogenic Trp-BODIPY-containing antimicrobial 277 peptides were designed for visualization of the deadly fungal pathogen Aspergillus fumigatus 278 in *ex vivo* human lung tissue<sup>96</sup> (Figure 4d). This residue is an optimal reporter as it retains the 279 molecular recognition of the native tryptophan and also provides a fluorogenic readout on 280 specific binding to fungal cells. Other fluorogenic antimicrobial peptides have enabled *in situ* 281 detection of bacteria in explanted whole human lungs using real-time optical 282 endomicroscopy<sup>97</sup>. The versatility of Trp-BODIPY has led to the preparation of peptide 283 imaging agents for additional targets. For example, a fluorogenic cyclic peptide that mimics 284 285 milk fat globule EGF factor 8 (MFGE8; also known as lactadherin), a protein with high binding affinity for phosphatidylserine, has been used to image the release of apoptotic bodies into the 286 extracellular space during programmed cell death<sup>98</sup>. A red analogue, Trp-redBODIPY, has 287 been used to prepare fluorogenic cyclic peptides that bind to keratin 1 (KRT1). These peptides 288 289 were synthesized using Pd-catalysed C–H activation during SPPS, a labelling approach that is 290 less disruptive than conventional lysine derivatization and enabled imaging studies to 291 understand the interaction between immune cells and cancer cells in aggressive breast cancer tumours<sup>47</sup>. 292

293 Imaging studies can be also performed to acquire functional readouts from cells. For example, a fluorogenic coumarin-based amino acid ( $\lambda_{em}$  = 460 nm), the first phosphotyrosine-294 mimetic FIAA, was used to report the endogenous phosphatase activity of protein tyrosine 295 phosphatases in live cells<sup>99</sup> (Figure 4e). The wash-free imaging capabilities of fluorogenic 296 peptides makes them valuable tools for applications where samples must be rapidly analysed 297 with few processing steps, such as in clinical diagnostics or metabolic engineering. For 298 example, NBD-aminophenylalanine (NBD-amPhe) has been used as a building block for the 299 preparation of epithelial cell adhesion molecule (EpCAM)-binding peptides that can detect 300

301 circulating tumour cells in the blood<sup>100</sup>. Furthermore, fluorogenic peptide aptamers have been

302 used to image in real time the production of paramylon (a carbohydrate granule similar to

- starch) by the microalga *Euglena gracilis*, thereby opening new opportunities in metabolic
- 304 engineering<sup>101</sup>.

### 305 [H1] Fluorescent D-amino acids in bacteria

### 306 [H3] Visualizing bacterial cell wall growth

Bacterial cells contain two main types of macromolecules that are assembled from 307 amino acids — namely, proteins, which consist of L-amino acids, and peptidoglycans, which 308 contain both L-amino acids and D-amino acids. Peptidoglycans are complex polymers that form 309 310 the cell wall of bacteria and coordinate multiple important processes, including cell growth and division<sup>102</sup>. Their biological importance has made peptidoglycans the target of many 311 antibiotics, and peptidoglycan biosynthesis has become an area of extensive research for the 312 discovery of new antimicrobial drugs<sup>103</sup>. The discovery of fluorescent D-amino acids (FDAAs) 313 has provided researchers in this field with non-invasive probes to visualize key steps during 314 the peptidoglycan biosynthesis in bacterial cells<sup>104,105</sup>. 315

Taking advantage of the inherent promiscuity of taxonomically-diverse bacterial to 316 incorporate D-amino acids as peptidoglycan metabolites<sup>106-108</sup>, modified D-amino acids have 317 been used to specifically label sites of new cell wall growth in real time<sup>109,110</sup>. Structurally-318 diverse FDAAs, including different reporters of varying size and optical properties, have been 319 synthesised<sup>111</sup> (Figure 5a). For example, ethynyl-D-alanine (EDA), azido-D-alanine (ADA) 320 and dipeptides, such as ethynyl-D-alanyl-D-alanine (EDA-DA), have been used to label 321 peptidoglycans in different species of bacteria by cycloadditions with fluorophores. These 322 bioorthogonal approaches require two reaction steps that can compromise cell viability; 323 however, these 'clickable' D-amino acidsare small and thus are minimally disruptive and 324 generally compatible with multiple reporters. For example, the small size and biological 325 stability of AlexaFluor 488-conjugated EDA-DA enabled the demonstration that 326 peptidoglycans are present in the cell wall of the human pathogen *Chlamydia trachomatis*<sup>111</sup>. 327 Alternatively, FDAAs have been employed for single-step labelling of peptidoglycans during 328 329 their biosynthesis in bacterial cells, for example, by the metabolic incorporation of FDAAs into 330 new peptidoglycans to label bacteria with minimal perturbation of the cells and simpler 331 protocols (that is, fewer washing steps). The good selectivity for bacteria and straightforward application of FDAAs has opened multiple research avenues in different biological 332

applications, ranging from *in vivo* analysis of the gut microbiota in mice<sup>112</sup> and in human fecal
 samples<sup>113</sup> to light-induced strategies for killing pathogenic bacteria<sup>114</sup>.

335 [H3] Multi-colour and multiplexed imaging

The simple, modular design of FDAAs and their convenient synthesis by condensation 336 337 of standard fluorophores to D-amino acids (such as D-lysine and D-diaminopropionic acid), has not only facilitated their diversification with a broad range of fluorophores but has also enabled 338 imaging experiments to answer questions about bacterial growth and division with high spatial 339 resolution. For example, live-cell imaging experiments in which three differently-coloured 340 FDAAs, NBD-3-amino-D-alanine (NADA;  $\lambda_{em} = 538$  nm), tetramethylrhodamine-D-lysine 341 (TDL;  $\lambda_{em} = 565 \text{ nm}$ ) and 7-hydroxycoumarin-D-alanine (HADA;  $\lambda_{em} = 450 \text{ nm}$ ), were time-342 343 pulsed to record the chronological steps of cell wall growth in Streptomyces venezuelae (Figure **5b**). This approach has also been used to monitor the production of peptidoglycans in single 344 bacterial cells in different environments, (such as during antibiotic treatment<sup>115</sup> or 345 transplantation<sup>109</sup>, and to track dynamic interactions between different bacterial species. In a 346 notable example of the latter, the predation cycle of Bdellovibrio bacteriovorus (a small 347 bacterial species that preys on larger bacteria) and its prey E. coli was studied by super-348 resolution imaging of the multiplexed FDAAs HADA, BODIPY-3-amino-D-alanine (BADA; 349  $\lambda_{\rm em} = 512$  nm) and tetramethylrhodamine-3-amino-D-alanine (TADA;  $\lambda_{\rm em} = 565$  nm)<sup>116</sup>. 350

Numerous studies have investigated the mechanism by which small and large FDAAs 351 can be incorporated into peptidoglycans of multiple bacterial species. Most FDAAs behave as 352 substrate analogues of solvent-accessible, periplasmic peptidoglycan transpeptidases, which 353 are responsible for the assembly of the cell wall material during growth and division<sup>117,118</sup>. For 354 example, HADA was used as an activity-based probe of peptidoglycan transpeptidases to 355 monitor the spatial distribution of new cell wall versus old cell wall in live *Bacillus subtilis*<sup>117</sup>. 356 Owing to their multiplexing capabilities, FDAAs can also be combined with fluorescently-357 tagged peptidoglycan biosynthetic proteins for multi-colour imaging studies. For example, 358 metabolic incorporation of HADA was combined with the localization of key proteins in cell 359 division (for example, FtsZ) in evolutionarily distinct rod-shaped bacterial species (such as B. 360 subtilis<sup>119</sup> and *E. coli*<sup>120</sup>), and MreB was colocalized with an EDA-DA-labelled peptidoglycan 361 ring during cell division in *Chlamydia trachomatis*<sup>121</sup>. Of note, these studies support a new 362 363 paradigm in which bacterial division progresses directionally around the cell, in contrast to the previously described model of uniform cell division. 364

365 [H3] Tools for high-throughput screenings

However, macromolecular labelling of peptidoglycan structures with conventional 366 FDAAs has constraints on temporal resolution due to the washing steps that are needed to 367 reduce the background fluorescence from unincorporated free amino acids. These technical 368 limitations prompted the design of a series of FDAAs that are non-fluorescent in low viscosity 369 environments (that is, when the FDAA is moving freely in aqueous media) and fluoresce 370 strongly when their intramolecular conformation is constrained (for example, after 371 incorporation into peptidoglycans)<sup>122</sup> (Figure 5c). These so-called rotor-FDAAs (RfDAAs) 372 consist of an electron-donating tetrahydroquinoline core structure coupled to electron-373 374 withdrawing carboxy-2-cyanovinyl groups to create efficient push-pull systems (Figure 3c). The resulting water-soluble amino acids Rf420DL ( $\lambda_{em} = 420 \text{ nm}$ ), Rf470DL ( $\lambda_{em} = 470 \text{ nm}$ ) 375 and Rf490DL ( $\lambda_{em}$  = 490 nm) have enabled wash-free and real-time imaging of peptidoglycan 376 biosynthesis, making possible the first high-throughput in vitro assay to probe the activity of 377 peptidoglycan transpeptidases. The imaging capabilities of these FDAAs will facilitate their 378 application in biological studies in which monitoring biological events in real time is of 379 extreme importance, including morphogenesis and drug screening studies. 380

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#### 382 [H1] Genetically-encoded FIAAs

### 383 [H3] Encoding FIAAs into proteins

Proteins can be endowed with fluorescence properties by numerous methods, including by enzymatic and chemical site-specific labelling with optical reporters and by *in vitro* translation of proteins with fluorescent tags<sup>123-127</sup>. These methods have undoubtedly aided the study of complex biological processes but they do suffer from some limitations, including potential off-target fluorescence and restriction to *in vitro* assays. Bearing this in mind, the genetic encoding of FIAAs has rapidly evolved to enable the generation of artificial proteins that might overcome some of these challenges<sup>128</sup>.

391 Site-specific incorporation of non-natural amino acids by genetic encoding in cells is generally achieved by assigning an amber stop codon to the non-natural amino acid and supplying the 392 393 cells with an exogenous aminoacyl-tRNA synthetase (aaRS) and its corresponding transfer 394 ribonucleic acid (tRNA). Importantly, the exogenous aaRS and tRNA must be orthogonal and 395 not cross react with the endogenous aaRSs and tRNAs of the host cell. Then, the orthogonal aaRS specifically couples the orthogonal tRNA to the non-natural amino acid so that the 396 'charged' tRNA is used by the ribosome for site-specific incorporation of the non-natural 397 amino acid (Figure 6a). Rapid advances in this technology have led to the efficient synthesis 398

of artificial proteins containing non-natural amino acids in live cells and animals<sup>129-131</sup>. To date, 399 more than 150 non-natural amino acids, including bioorthogonal, photoreactive and 400 photocaged amino acids, as well as amino acids with electron paramagnetic resonance (EPR), 401 infrared (IR) and nuclear magnetic resonance (NMR) labels, have been genetically encoded in 402 different organisms<sup>132</sup>. The development of organisms with synthetic genomes<sup>133</sup> and the 403 evolution of quadruplet-codon-decoding ribosomes<sup>134,135</sup> will expand the applicability of new 404 405 genetically-encoded FIAAs with different chemical structures and/or additional photophysical 406 properties to label proteins.

#### 407 [H3] Genetically-encoded FIAAs in prokaryotes

The FlAAs that have been genetically incorporated into proteins (Figure 6b) can be 408 broadly classified by whether they are expressed in prokaryotic or in eukaryotic cells. 409 Following the discovery and application of L-(7-hydroxycoumarin-4-yl)ethylglycine (CouA; 410  $\lambda_{em.}$  = 380 nm in neutral form;  $\lambda_{em.}$  = 450 nm in anionic form), acridon-2-ylalanine (Acd;  $\lambda_{em.}$  = 411 446 nm) and 4-biphenyl-l-phenylalanine (terphenylA;  $\lambda_{em} = 342$  nm) were developed as 412 additional FIAAs for site-specific incorporation into proteins in E. coli<sup>20,45,136</sup>. In these cases, 413 evolved mutant tyrosyl-aaRSs from the archaeon Methanococcus jannaschii (MjTyrRS) were 414 415 used to generate 'charged' tRNAs inside the cell, which resulted in successful ribosomal incorporation. Of note, the *Mj*TyrRS–*Mj*tRNA pair is orthogonal in prokaryotes and has been 416 evolved for incorporation of several other non-natural amino acids. 417

CouA is synthesized by incorporating a  $\beta$ -keto ester in the amino acid side chain and then 418 reacting it with resorcinol by Pechmann condensation $^{20}$ . The ease of synthesis and efficient 419 420 incorporation of CouA into proteins has facilitated the widespread application of CouA in biological studies. As the emission spectrum of CouA overlaps with the excitation spectrum of 421 422 cyan fluorescent protein (CFP), CouA was used to engineer a new CFP with a long Stokes shift of ~110 nm (ref.<sup>137</sup>). In this case, the precise incorporation of CouA at 20 Å from the CFP 423 fluorophore results in efficient FRET so that CFP fluorescence (at 426 nm) is detected when 424 CouA is excited (at 365 nm) (Figure 6c). Furthermore, the sensitivity of CouA to pH and 425 polarity changes has also been exploited to produce artificial proteins with readouts that are 426 sensitive to environmental conditions<sup>138</sup>. For example, this approach was used to investigate 427 428 the difference in substrate specificity between the haloalkane dehalogenases DhaA and DbjA<sup>139</sup>. Steady-state and time-resolved fluorescence measurements of dehalogenase mutants 429 430 containing CouA at specific sites revealed greater hydration in DbjA than in DhaA, which correlates with the substrate specificity of the two proteins. CouA has also been genetically 431

432 encoded into the transient receptor potential cation channel subfamily V member 1 (TRPV1)

to study the dynamics of receptor activation, with changes in the fluorescence being correlated

434 with the opening and closing of the channel on binding of capsaicin<sup>140</sup>.

# 435 [H3] Genetically-encoded FIAAs in eukaryotes

436 Lysine derivatives of 7-hydroxycoumarin (Lys-Cou) have been genetically encoded into proteins using evolved mutants of Methanosarcina barkeri pyrrolysyl RS (PylRS)-437 tRNA<sub>CUA</sub><sup>141</sup>. The PylRS–tRNA pair<sup>142,143</sup> has become an extremely useful tool for genetic 438 encoding of non-natural amino acids because it is orthogonal to prokaryotes and eukaryotes 439 and therefore can be used in cells of evolutionarily diverse origins, including E. coli, 440 Saccharomyces cerevisiae, mammalian cells and even whole intact organisms, such as the 441 nematode worm *Caenorhabditis elegans* and the fruit fly *Drosophila melanogaster*<sup>144</sup>. ACD 442 has also been encoded in fluorescent proteins for expression in prokaryotes. The high quantum 443 yield (~95%, compared with 63% for CouA), efficient synthesis and high photostability of 444 ACD are some of its key advantages. 445

Among the FIAAs that have been genetically encoded in eukaryotes, the most notable 446 are dansyl and prodan derivatives, such as dansylalanine ( $\lambda_{em} = 540 \text{ nm}$ ) and ANAP ( $\lambda_{em} = 490$ 447 nm)<sup>145-147</sup> (Figure 6b). These FlAAs have been incorporated into proteins expressed in S. 448 cerevisiae using evolved mutants of the E. coli-derived leucyl-RS-leucyl-tRNA<sub>CUA</sub> pair, which 449 450 is orthogonal to those found in eukaryotic cells. A remarkable feature of these amino acids is that their emission maximum and quantum yield can vary drastically depending on the polarity 451 of the medium; for example, the quantum yield of dansyl-based amino acids vary from 50% 452  $(\lambda_{em} = 510 \text{ nm})$  in ethyl acetate to <10% ( $\lambda_{em} = 578 \text{ nm}$ ) in water<sup>148</sup>; the emission maxima for 453 454 ANAP are 490 nm in water and 420 nm in ethyl acetate. This sensitivity to the dielectric constant of the surrounding environment has been exploited to investigate the dynamics of 455 different protein structures. For example, ANAP was used to map the regions of the voltage-456 gated potassium ion channel ion channel that are inaccessible to chemical labelling<sup>149</sup> (**Figure** 457 6d). First, the FIAA was genetically encoded into different positions in the ion channel 458 sequence, either the extracellular and intracellular regions, and then electrophysiology and 459 460 fluorescence measurements were performed concurrently to identify the regions of the protein that cooperatively lead to opening of the channel. Similarly, ANAP has been also used to 461 examine the catalytic activity of a voltage-sensing phosphatase<sup>150</sup> and to gain insights into 462 protein-protein interactions using FRET measurements, as in the case of the pro-apoptotic 463 protein BAX and the anti-apoptotic protein HSP70 (ref.<sup>151</sup>). Different mutants of BAX with 464

ANAP encoded at three different positions in the protein were tested for binding to HSP70 fused to yellow fluorescent protein (YFP), so that FRET signals were detected only when the two proteins were in close proximity. This FRET signal was used as a reporter of the interaction between BAX and HSP70 and helped to discern the mechanism of small molecule-induced inhibition of this interaction. These examples highlight how genetically-encoded FlAAs can be used to gain insights into protein dynamics and folding, as well as the interactions between proteins and other biomolecules.

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### 473 [H1] Conclusions and outlook

Advances in the chemical derivatization of heterocyclic building blocks and 474 fluorophores have accelerated the design and synthesis of FIAAs with improved properties for 475 chemical biology studies. To date, the unique reactivity of the indole moiety<sup>152,153</sup> has resulted 476 in the synthesis of multiple tryptophan analogues by different transformations (such as Pd-477 catalysed couplings and C-H activation). Site-specific modification of other amino acids, 478 including, amongst others, histidine<sup>154,155</sup>, phenylalanine<sup>49</sup> and tyrosine<sup>156,157</sup>, will help to 479 enable other residues with complementary functionalities to be fluorescently labelled and, 480 ultimately, lead to broader chemical diversity. Similarly, the emergence of new chemistry in 481 482 the field of synthetic fluorophores will enable the construction of FlAAs with additional properties. These properties include not only physicochemical properties, such as small 483 size<sup>158,159</sup> to reduce the effect of labelling in peptides and proteins, but also optical features for 484 spectroscopy and imaging experiments, including near-infrared emission<sup>160,161</sup>, super-485 resolution capabilities<sup>162,163</sup>, photoactivatable behaviour<sup>164,165</sup> or suitability for multimodal 486 imaging (for example, optoacoustics and positron emission tomography)<sup>166,167</sup>. A greater 487 number and variety of FIAAs will create avenues to identify new roles for peptides and proteins 488 in biological systems. FIAAs with longer emission wavelengths will improve the penetration 489 490 depth that is required for in vivo imaging studies, and photoswitching FIAAs may allow researchers to image the localization and trafficking of proteins using super-resolution 491 microscopy. New FIAAs might also find important applications in other areas of chemistry in 492 which they may have been underexplored to date, such as the material sciences, where 493 494 fluorescent building blocks embedded in supramolecular structures could provide dynamic readouts to characterize the formation and properties of new materials in real time, including 495 nanofibres and peptide hydrogels<sup>168,169</sup>. 496

497 For biological applications, the design of new methodologies in synthetic biology will498 facilitate the integration of new modular blocks into different macromolecular structures. An

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499	area	that will benefit substantially from these developments is genetically-encoded FIAAs.	
500	Altho	ough currently available genetically-encoded FIAAs hold huge potential for studying the	
501	dynamics, folding and biomolecular interactions of proteins, their number and types are		
502	limited. Most fluorophores absorb at very short wavelengths that require high energy		
503	excitation and lead to limited tissue penetration and poor signal-to-background ratios, [which		
504	precludes their use in whole organisms. Non-natural amino acids showing good spectral		
505	overlap with existing fluorophores or fluorescent proteins would be also extremely useful to		
506	build FRET pairs within proteins and thereby study dynamic conformational changes with		
507	increased resolution. The UV excitation wavelengths of current FIAAs also reduce their		
508	compatibility with photocaging groups, which are typically cleaved when illuminated at those		
509	wavelengths (for example, 365 nm) <sup>170,171</sup> . Therefore, synthetic routes that are able to red-		
510	shift the excitation and emission wavelengths <sup>172</sup> of non-natural FIAAs would have a		
511	remarkable impact on the design of new artificial proteins. However, this synthetic effort will		
512	need to be matched with advances in the biological counterpart, where the identification of		
513	new aaRS-tRNA pairs that can recognize and incorporate such molecules will be essential		
514	for th	ne successful encoding of novel FlAAs into protein structures.	
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516	Refe	rences	
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# 963 Author contributions

- All authors contributed equally to the preparation of this manuscript.
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# 966 Competing interests

- 967 The authors declare no competing interests.
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### 979 **Figure legends**

**Figure 1**. Applications of FIAAs in chemical biology. The design and synthesis of new

981 fluorescent amino acids (FIAAs) has created multiple opportunities for *in vitro* biomolecular

recognition assays (part **a**), solid-phase peptide synthesis (SPPS) of fluorescent peptides for

983 optical imaging (part **b**), genetic encoding of fluorescent building blocks into proteins (part **c**)

and microbiology experiments to study bacterial growth (part **d**). Images in part **b** reproduced

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**Figure 2. Representative non-natural FIAAs developed in the past decade**. Chemical structures and basic spectral properties of fluorescent analogues of tryptophan (part **a**) and fluorescent amino acids (FIAAs) obtained by appending fluorophores to amino acids using a broad range of chemical reactions (part **b**).  $\phi_F$ , fluorescence quantum yield; Pd, palladium; Rh, rhodium; CuAAC, copper-catalysed azide–alkyne cycloaddition.

992 Figure 3. Synthetic schemes for the *de novo* preparation of non-natural FIAAs. a | Design of flavone-based fluorescent amino acids (FIAAs) building on tyrosine derivatization<sup>71-73</sup>. **b** 993 Pyrazoloquinazolines derived from L-aspartic acid as one-photon and two-photon 994 fluorophores<sup>78</sup>. c | Synthesis of rotor fluorogenic D-amino acids (RfDAAs) for labelling 995 in bacterial cells<sup>122</sup>. Boc, *tert*-butoxycarbonyl; 996 peptidoglycan structures DCM, 2,3-dichloro-5,6-dicyano-1,4-benzoquinone; 997 dichloromethane; DDQ, DIPEA, N,Ndiisopropylethylamine; DMF, *N*,*N*-dimethyl formamide; Fmoc-OSu, *N*-(9-998 999 fluorenylmethyloxycarbonyl)succinimide; TFA, trifluoroacetic acid ; THF, tetrahydrofuran; Trt-Cl, triphenylmethyl chloride; TSTU, N,N,N',N'-tetramethyl-O-(N-succinimidyl)uronium 1000 tetrafluoroborate. 1001

Figure 4. Bioactive fluorescent peptides for structural studies and optical imaging. a 1002 Schematic representation of the interaction between the HIV-1 nucleocapsid peptide W37 1003 (labelled with the flavone-based fluorescent amino acid (FlAA) M3HFaa) and 1004 1005 oligonucleotides. The graph shows the fluorescence emission spectra of the W37–M3HFaa 1006 peptide on binding to different nucleic acids. **b** | Schematic illustration of the binding between 1007 a fluorescent peptide containing the FIAA 4-DMAP and a PDZ domain. The graph shows the 1008 fluorescence emission spectra of the 4-DMAP-containing peptide in the unbound state and 1009 bound to the PDZ domain. c | Illustration of the pH-(low) insertion peptide (pHLIP) containing

1010 4-cyanotryptophan (which acts as a FRET donor, and DiO-stained large unilamellar vesicles (LUVs), which act as FRET acceptors, for the study of peptide-membrane interactions. The 1011 1012 graph depicts the FRET measurements taken at 512 nm with increasing concentrations of labelled pHLIP. **d** | Chemical structure of the Trp-BODIPY-labelled cyclic peptide PAF26 and 1013 1014 live-cell confocal imaging of Aspergillus fumigatus over time. Scale bar: 2.5 µm. Multi-photon 1015 fluorescence microscopy images of ex vivo human lung tissue after incubation with RFPexpressing A. fumigatus (red) and the Trp-BODIPY-labelled PAF26 peptide (green). Second 1016 harmonic generation images are shown in cyan. Scale bar: 10 µm. Fluorescence lifetime images 1017 1018 depicting autofluorescent tissue structures (white arrows) and a PAF26-stained A. fumigatus hypha (yellow arrows). Scale bar: 20 µm. e | pER peptide containing a self-immobilizing 1019 coumarin FIAA that mimics phosphotyrosine (pTyr) to report endogenous activity of protein 1020 1021 tyrosine phosphatases (PTPs). The graph depicts time-dependent fluorescence measurements 1022 of the pER peptide before and after ultraviolet (UV) irradiation (365 nm) in HeLa cell lysates. 1023 Fluorescence microscopy images of HeLa cells after incubation with pER and UV irradiation to visualize intracellular PTP activity (green: pER peptide, red: commercial tracker). Part a 1024 1025 reproduced with permission from ref. 73, American Chemical Society. Part b was reproduced with permission from ref. 88, American Chemical Society. Part c reproduced with permission 1026 1027 from ref. 31, The Royal Society of Chemistry. Part d reproduced with permission from ref. 96, Springer Nature Limited. Part e reproduced with permission from ref. 99, The Royal Society 1028 1029 of Chemistry.

1030 Figure 5. Fluorescent D-amino acids for studying bacterial growth. a | Peptidoglycans are polymers of L- and D-amino acids and sugars, and a major component of bacterial cell walls. 1031 The peptidoglycan layer is considerably thicker in Gram-positive than in Gram-negative 1032 bacteria. **b** | Chemical structures and optical properties of 'clickable' and fluorescent D-amino 1033 1034 acids (FDAAs) used for labelling peptidoglycan structures in Gram-positive and Gram-1035 negative bacteria. c | Time-lapse multicolour microscopy images of the bacterium *Streptomyces* 1036 by pulsed-incubation with NBD-3-amino-D-alanine (NADA; venezuelae green), 1037 tetramethylrhodamine-D-lysine (TDL; red), and 7-hydroxycoumarin-D-alanine (HADA; blue). 1038 d | Schematic representation of the fluorescence activation of the rotor FDAA Rf470DL on peptidoglycan labelling and fluorescence microscopy images of *Bacillus subtilis* depicting 1039 1040 peptidoglycan labelling using Rf470DL (red) or HADA (cyan) before and after washing. Part c reproduced with permission from ref. 110, Wiley VCH. Part d reproduced with permission 1041 1042 from ref. 122, Springer Nature Limited.

1043 Figure 6. Genetically encoded site-specific incorporation of FIAAs in live cells. a | Nonnatural amino acids are site-specifically incorporated into proteins by assigning an amber stop 1044 1045 codon (UAG) to non-natural amino acids and supplying the cells with an exogenous aminoacyltRNA synthetase (aaRS) and its corresponding tRNA, which charges the tRNA with the non-1046 natural amino acid. **b** | Chemical structures of genetically-encoded fluorescent amino acids 1047 (FIAAs) and their optical properties. c | Adjusting the photophysical properties of a fluorescent 1048 1049 protein. Site-specific incorporation of L-(7-hydroxycoumarin-4-yl)ethylglycine (CouA) at position 39 in cyan fluorescent protein (CFP), resulting in efficient FRET between CouA and 1050 1051 CFP. When CouA (in blue) is excited at 360 nm, emission from the CFP fluorophore (in cyan) at 476 nm is observed. **d** | Mapping ion channel dynamics. 3-(6-acetylnaphthalen-2-ylamino)-1052 1053 2-aminopropanoic acid (ANAP) is site-specifically incorporated into both extracellular and intracellular positions (black circles) in a voltage-gated potassium ion channel. Voltage-1054 dependent changes in the fluorescence emission of ANAP revealed information about the 1055 regions of the ion channel that move cooperatively on opening of the channel. Red spheres in 1056 1057 the structure are potassium ions. Part c adapted with permission from ref. 137, American Chemical Society. Part d adapted with permission from ref. 149, National Academy of 1058 1059 Sciences USA.

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#### 1061 ToC blurb

Fluorescent amino acids are widely used as building blocks for non-perturbative labelling of peptides and proteins. This Review covers recent advances in the design and synthesis of FlAAs with bespoke optical properties for different applications in biological studies.