Genetically Encoded 2-Aryl-5-carboxytetrazoles for Site-Selective Protein Photo-Cross-Linking

Yulin Tian,† Marco Paolo Jacinto,† Yu Zeng,‡ Zhipeng Yu,‡,# Jun Qu,†⊥ Wenshe R. Liu,‡⊥ and Qing Lin*‡⊺

†Department of Chemistry, State University of New York at Buffalo, Buffalo, New York 14260, United States
‡Department of Chemistry, Texas A&M University, College Station, Texas 77845, United States
⊺Department of Pharmaceutical Sciences, State University of New York at Buffalo, Buffalo, New York 14260, United States

ABSTRACT: The genetically encoded photo-cross-linkers promise to offer a temporally controlled tool to map transient and dynamic protein–protein interaction complexes in living cells. Here we report the synthesis of a panel of 2-aryl-5-carboxytetrazole-lysine analogs (ACTKs) and their site-specific incorporation into proteins via amber codon suppression in Escherichia coli and mammalian cells. Among five ACTKs investigated, N-methylpyrroletetrazole-lysine (mPyTK) was found to give robust and site-selective photo-cross-linking reactivity in E. coli when placed at an appropriate site at the protein interaction interface. A comparison study indicated that mPyTK exhibits higher photo-cross-linking efficiency than a diazirine-based photo-cross-linker, AbK, when placed at the same location of the interaction interface in vitro. When mPyTK was introduced into the adapter protein Grb2, it enabled the photocapture of EGFR in a stimulus-dependent manner. The design of mPyTK along with the identification of its cognate aminoacyl-tRNA synthetase makes it possible to map transient protein–protein interactions and their interfaces in living cells.

To map dynamic protein–protein interactions in living cells, a powerful chemical strategy involves the use of genetically encoded photo-cross-linkers that permanently link transient protein–protein interaction complexes with a burst of light. Based on the structures, the reported genetically encoded photo-cross-linkers contain one of the three moieties: phenyl azide such as pAzF,† benzophenone such as pBpa,‡ and diazirine such as AbK‡⊥ and their derivatives (Chart 1).†⊥ Although these genetically encoded photo-cross-linkers have proven to be valuable in the study of protein structure and function, they invariably cross-link with their interacting protein partners with no selectivity for any particular residue as the photogenerated reactive intermediate, i.e., the nitrene from phenyl azide, the diradical from benzophenone and the carbene from diazirine, inserts into a proximal C–H bond with appropriate distance and angle,‡⊥ making it difficult to predict a prior suitable positions for installation of the photo-cross-linker. In addition, the tandem mass spectrometry-based mapping of the interaction interface is complicated as any residue from the interacting protein partner can potentially participate in the photo-cross-linking.

Recently, we reported a new photoaffinity label based on 2-aryl-5-carboxytetrazole (ACT) with a size similar to benzophenone, which cross-links its target proteins via addition with a proximal nucleophile near the active site.‡⊥ Because ACT exhibits ligand-dependent selective photo-cross-linking, we envisioned that ACT may also serve as a new class of genetically encoded photo-cross-linkers for mapping transient protein–protein interaction interfaces. It is noted that a biocompatible proximity-driven nucleophilic substitution reaction between a genetically encoded N′-fluoroacetyllysine and cysteine was reported recently for mapping the protein–protein interaction interface.‡⊥ However, the occurrence of native cysteine at the protein–protein interaction interface is rather rare.‡⊥ Herein, we report the synthesis of a panel of ACT-lysine analogs (ACTK, Chart 1), and the identification of a new ACTK-specific pyrrolysyl-tRNA synthetase for site-specific incorporation of ACTK into proteins in Escherichia coli and in mammalian cells. One of the ACTK analogs, mPyTK, exhibited robust and site-selective photo-cross-linking of a GST dimer in bacteria. In a comparison study, mPyTK showed significantly higher cross-linking efficiency than AbK when both are incorporated at the same location of GST. Moreover, the mPyTK-encoded adapter protein, Grb2, showed a stimulus and position-dependent capture of its transient interaction partner, epidermal growth factor receptor (EGFR), in mammalian cells.

Because pyrrolysyl-tRNA synthetase (PyIrs) and its variants have shown tremendous versatility in charging various lysine derivatives into proteins site-selectively in bacteria, yeast and mammalian cells,†⊥ we decided to append ACT motif onto the ε-amino group via simple acylation reaction. For the synthesis of

Chart 1. Genetically Encoded Photo-Cross-Linkers

Previous work:

This work:

© 2017 American Chemical Society
ACTK analogs 1–4 in Chart 1, the key intermediate, ethyl 2-aryl-2H-tetraazole-5-carboxylate, was obtained through Cu(I)-catalyzed cross-coupling of ethyl 2H-tetraazole-5-carboxylate with the phenylaryliodonium salt14 (Schemes S1–S4 in Supporting Information). For PhTK (5), the Kakehi tetrazole synthesis was followed to give the ethyl 2-phenyl-2H-tetraazole-5-carboxylate intermediate15 (Scheme S5 in Supporting Information). Subsequent hydrolysis and coupling with Fmoc-lysine·HCl followed by removal of the protecting group afforded the mPyTKRS library, in which four residues surrounding the N-methylpyrrole-lysine side chain based on the crystal structure of Methanosarcina mazei PylRS in complex with Pyl-AMP16 (Y306L, L309, C348, and Y384; Figure 1a) were randomized, was subjected to successive rounds of the positive and negative selections.17 An MmPylRS mutant displaying the highest amber suppression efficiency in E. coli was identified that carries the Y306L/L309A/C348F/Y384F mutations and is hereafter referred to as mPyTKRS. A plasmid displaying the highest amber suppression efficiency in E. coli was identified that carries the Y306L/L309A/C348F/Y384F mutations and is hereafter referred to as mPyTKRS. A plasmid pEvol-mPyTKRS encoding mPyTKRS and tRNA\textsuperscript{Pyl}_{\text{CUA}} was then constructed and showed site-specific incorporation of mPyTK into superfolder green fluorescent protein (sfGFP) carrying an amber mutation at the Q204 position and a C-terminal His tag (Figure 1b). Interestingly, we found that mPyTKRS also exhibits polyspecificity and allows site-specific incorporation of other ACTK into sfGFP-Q204TAG with the level of expression following the order of mPyTK > PhTK > PyTK > TTK (Figure 1b). A 100 mM solution of the sfGFP-Q204mPyTK mutant in BL21DE3 cells in the presence of 1 mM mPyTK gave a protein yield of 0.8 mg L\(^{-1}\); subsequent mass spectrometry analysis verified the incorporation of mPyTK (Figure 1c and Figure S1 in the Supporting Information (SI)). The lower expression yield is likely due to reduced enzymatic activity of mPyTKRS as a higher yield, 17.8 mg L\(^{-1}\), was obtained when BocK was charged into the sfGFP-Q204 position using the wild-type PylRS under identical conditions (Figure S2 in SI). Next, we examined the efficiency of ACTK in photo-cross-linking. Schistosoma japonicum glutathione-S-transferase (S\textregistered GST) homodimer (Figure 2a), a system used widely for evaluating the genetically encoded photo-cross-linkers.\textsuperscript{1,2} Inspection of the GST dimer structure (PDB code: 1Y6E) revealed that the four interfacial residues, E52, F53, L66, and R74, are located at a distance of \(\sim 4–8\) Å from a potential nucleophilic residue from the opposite monomer (Figure S3 in SI). Thus, the GST mutants carrying mPyTK at these positions were expressed in BL21DE3 cells and purified by Ni-NTA affinity chromatography. To our surprise, only the GST-E52TAG mutant gave detectable expression (with a yield of 2.8 mg L\(^{-1}\)) based on SDS-PAGE, which was confirmed by LC-MS data (Figure S4 in SI). To probe whether mPyTK enables photo-cross-linking in vitro, the purified GST-E52mPyTK protein was irradiated with a handheld 302 nm UV lamp for 0, 1, 5, and 15 min on ice, and covalent dimer formation was monitored by SDS-PAGE. We observed time-dependent GST dimer formation for the E52mPyTK mutant with \(\sim 53\%\) yield at 5 min, but not for wild-type (Figure 2b), indicating that the ACT moiety is responsible for dimer cross-linking. For comparison, we expressed the GST-E52AbK mutant using the wild-type PylRS with a yield of 2.5 mg L\(^{-1}\) and
examined its photo-cross-linking reactivity. To our surprise, AbK exhibited very weak reactivity as the dimer band was detected only by Western blot (Figure S5 in SI) but not Coomassie blue (Figure 2b). Interestingly, a higher cross-linking yield (~79%) was obtained when E. coli cells expressing GST-E52mPyTK were directly photoirradiated, which could be attributed to higher intracellular concentration of the GST mutant (Figure 2c and Figure S6 in SI). Because mPyTKRS can charge other ACTKs into proteins site-selectively, we expressed SJGST mutants carrying PyTK, FTK, TTK and PhTK, respectively, at position S2 and compared their photo-cross-linking efficiency in E. coli cells. Based on Western blot analysis, only mPyTK, FTK and TTK showed the cross-linked dimer with the efficiency order of mPyTK > TTK > FTK (Figure 2d), presumably due to the highest electron density at N-methyl-pyrrrole ring, which helps to stabilize the photogenerated carboxy-nitrile imine and increase its lifetime in biological media.

Because ACT photoreacts with proximal nucleophilic residues on proteins, we sought to determine which nucleophilic residues on the opposite GST monomer might react with the photogenerated carboxy-nitrile imine intermediate. To this end, we built a model of the GST-E52mPyTK and surveyed the chemical environment surrounding mPyTK. Four nucleophilic residues (E92, M133, C139 and K141) were identified that are located 2.8−13.0 Å from the electrophilic nitrile imine carbon (Figure 3a). To determine which one of these four residues participates in the cross-linking reaction, we mutated these residues to alanine and examined the photo-cross-linking activity of the resulting mutants. We found the Glu92 → Ala mutation completely abolished the covalent dimer formation whereas other mutations had no effect (Figure 3b). This result is consistent with the proximity-driven reactivity as E92 is closest to mPyTK with a calculated distance between the carboxylate oxygen and the nitrile imine carbon of 2.8 Å (Figure 3a). Similar results were obtained when the alanine scan was conducted with the GST-E52-FTK mutant (Figure S7 in SI). We propose a photo-cross-linking mechanism in which the E92 carboxylate undergoes nucleophilic addition to the photogenerated carboxy-nitrile imine followed by 1,4-acyl shift (Figure 3c). The rearranged cross-linked structure was supported by tandem mass spectrometry data in which the two fragment ions derived from the two discrete cleavage pathways were positively identified (Figure S8 in SI). For comparison, we performed photo-cross-linking studies with the same set of alanine mutants of the GST-E52AbK (E92A, M133A, C139A and K141A). We did not observe prominent attenuation in GST dimer formation; unexpectedly, two alanine mutants (M133A and K141A) showed greater extent of dimer formation than the GST-E52AbK alone (Figure S7 in SI), presumably due to a remodeling of the interaction interface that alters the distance and/or angle of a suitable proximal C−H bond.

To examine whether the genetically encoded mPyTK can capture protein−protein interaction complexes in mammalian cells, we first confirmed that when mPyTKRS was expressed in HEK293T cells, it allowed site-selective incorporation of mPyTK into mCherry-TAG-EGFP based on confocal fluorescence microscopic analysis (Figure S9 in SI). We then introduced mPyTK into Grb2, an adaptor protein that links phosphorylated EGFR to the Ras signaling pathway through guanine nucleotide exchange factor Sos1 and was used previously for evaluating the genetically encoded photo-cross-linkers in mammalian cells. Inspect the crystal structure of the Grb2 SH2 domain in complex with a phosphotyrosine-containing heptapeptide ligand20 revealed that 9 residues surrounding the ligand, A91, D104, V105, Q106, F108, K109, L111, W121 and N143, could be mutated to mPyTK for potential photo-cross-linking with the SH2 ligands such as the cytoplasmic domain of EGFR (Figure 4a). Thus, HEK293T cells were cotransfected with pCMV-mPyTKRS-tRNA\textsuperscript{Pyl}\textsubscript{CUA} encoding mPyTK-specific PylRS and tRNA\textsuperscript{Pyl}\textsubscript{CUA} pCMV6-Grb2-myc-DDK encoding either wild-type or amber mutant with TAG codon substituted at any of the 9 positions with a C-terminal myc-DDK tag, and pcDNA3-EGFR-EGFP encoding full-length EGFR and a C-terminal EGFP tag, and protein expressions were carried out in DMEM medium supplemented with 10% FBS and 1 mM mPyTK. The cells were starved for 12 h before EGF stimulation and subsequent photoinactivation on ice. The cells were then lysed and the lysates were treated with protein tyrosine phosphatase 1B to hydrolyze the phosphotyrosine to obviate noncovalent Grb2-interacting protein complexes. The cross-linked Grb2-interacting proteins were immunoprecipitated with anti-Flag antibody and analyzed by sequential Western blots using antmyc and anti-EGFR antibodies. The cross-linked Grb2−EGFR complex was detected for 8 out 9 Grb2-mPyTK mutants; the D104mPyTK mutant gave the highest photo-cross-linking yield followed by
Figure 4. Photo-cross-linking of EGFR by the mPyTK-encoded Grb2 mutants in mammalian cells. (a) A close-up view of Grb2-SH2 domain in complex with phosphoryrosine-containing heptapeptide (PDB code: 1TZE), highlighting the proximal residues surrounding pY (rendered in cyan tube model) that were selected for the mPyTK mutagenesis. (b) Comparison of the efficiency of the mPyTK-containing Grb2-SH2 domain mutants for photo-cross-linking with EGFR in HEK293T cells. Cells were exposed to 302 nm UV light for 5 min before lysis. The Grb2-cross-linked proteins in cell lysates were immunoprecipitated with the anti-FLAG antibody-immobilized agarose beads. The samples were analyzed SDS-PAGE/Western blot and probed successively with anti-FLAG antibody (top panel) and anti-EGFR antibody (bottom panel).

In summary, we have synthesized a panel of 2-aryl-5-carboxy-$\text{N}^1$-tetrazole-based photo-cross-linkers and evolved a polyspecific pyrrolysil-tRNA synthetase variant that charges these ACT-lysine analogs site-selectively into proteins in *E. coli*. One of the most reactive, genetically encoded photo-cross-linkers, mPyTK, allowed site-selective photo-cross-linking of a GST dimer in vitro and in *E. coli* cells. In comparison studies, mPyTK exhibited a significantly higher efficiency than AbK when placed at the same location of the GST dimer interface. Moreover, mPyTK enabled covalent capture of the transient Grb2-interacting protein partner in mammalian cells in a stimulus-dependent manner. In view of their higher photo-cross-linking yields and unique cross-linking mechanism, these genetically encoded ACT-lysines should offer a powerful chemical tool to map transient protein–protein interactions underlining signal transduction pathways. Because tandem MS analysis revealed the characteristic fragments after photo-cross-linking, these ACT-lysines may also find applications in structural analysis of protein complexes.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.7b02615.

Supplemental figures and table, synthetic schemes, experimental procedures, characterization of new compounds (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**

Jun Qu: 0000-0002-1346-6809

Wenshe R. Liu: 0000-0002-7078-6534

Qing Lin: 0000-0002-9196-5718

**Present Address**

Sichuan University College of Chemistry, China.

**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

We gratefully acknowledge the NIH (GM088592 to Q.L., CA161158 to W.R.L.) for financial support. We thank Prof. John Koland at University of Iowa for providing the pcDNA3-EGFR-EGFP plasmid, and Carlo Ramil in Q.L. lab for cloning assistance.

**REFERENCES**

(1) Chin, J. W.; Santoro, S. W.; Martin, A. B.; King, D. S.; Wang, L.; Schultz, P. G. J. Am. Chem. Soc. 2002, 124, 9026–9027.

(2) Chin, J. W.; Martin, A. B.; King, D. S.; Wang, L.; Schultz, P. G. Proc. Natl. Acad. Sci. U. S. A. 2002, 99, 11020–11024.

(3) Ai, H. W.; Shen, W.; Sagi, A.; Chen, P. R.; Schultz, P. G. ChemBioChem 2011, 12, 1854–1857.

(4) Chou, C. J.; Uprety, R.; Davis, L. I.; Chin, J. W.; Deiters, A. Chem. Sci. 2011, 26, 480–483.

(5) Lin, S.; He, D.; Long, T.; Zhang, S.; Meng, R.; Chen, P. R. J. Am. Chem. Soc. 2014, 136, 11860–11863.

(6) Yang, Y.; Song, H.; He, D.; Zhang, S.; Dai, S.; Lin, S.; Meng, R.; Wang, C.; Chen, P. R. Nat. Commun. 2016, 7, 12299.

(7) Yanagisawa, T.; Hino, N.; Ihara, F.; Mukai, T.; Sakamoto, K.; Yokoyama, S. Mol. BioSyst. 2012, 8, 1131–1135.

(8) Hino, N.; Oyama, M.; Sato, A.; Mukai, T.; Ihara, F.; Hayashi, A.; Kozuka-Hata, H.; Yamamoto, T.; Yokoyama, S.; Sakamoto, K. J. Mol. Biol. 2011, 406, 343–353.

(9) Sato, S.; Mimatsu, S.; Sato, A.; Hino, N.; Sakamoto, K.; Umehara, T.; Yokoyama, S. Biochemistry 2011, 50, 250–257.

(10) Hernes, A.; Marjanovic, J.; Lewandowski, T. M.; Marin, V.; Patterson, M.; Miesbauer, L.; Ready, D.; Williams, J.; Vasudevan, A.; Lin, Q. J. Am. Chem. Soc. 2016, 138, 14609–14615.

(11) Kobayashi, T.; Hoppmann, C.; Yang, B.; Wang, L. J. Am. Chem. Soc. 2016, 138, 14832–14835.

(12) Bogan, A. A.; Thorn, K. J. J. Mol. Biol. 1998, 280, 1–9.

(13) Pan, W.; Tharp, J. M.; Liu, W. R. Biochim. Biophys. Acta, Proteins Proteomics 2014, 1844, 1059–1070.

(14) An, P.; Yu, Z.; Lin, Q. Chem. Commun. (Cambridge, U. K.) 2013, 49, 9920–9922.

(15) Wang, Y.; Lin, Q. Org. Lett. 2009, 11, 3570–3573.

(16) Kavran, J. M.; Gundlapalli, S.; O’Donoghue, P.; Englert, M.; Soll, D.; Steitz, T. A. Proc. Natl. Acad. Sci. U. S. A. 2007, 104, 11268–11273.

(17) Wang, Z. A.; Zeng, Y.; Kurra, Y.; Wang, X.; Tharp, J. M.; Vatansere, E. C.; Hsu, W. W.; Dai, S.; Fang, X.; Liu, W. R. Angew. Chem., Int. Ed. 2017, 56, 212–216.

(18) Li, N.; Batzer, A.; Daly, R.; Yajnik, V.; Skolnik, E.; Chardin, P.; Bar-Sagi, D.; Margolis, B.; Schlessinger, J. Nature 1993, 363, 85–88.

(19) Hino, N.; Okazaki, Y.; Kobayashi, T.; Hayashi, A.; Sakamoto, K.; Yokoyama, S. Nat. Methods 2005, 2, 201–206.

(20) Rahuel, J.; Gay, B.; Erdmann, D.; Strauss, A.; Garcia-Echeverria, C.; Furet, P.; Caravatti, G.; Fretz, H.; Schoepfer, J.; Grutter, M. G. Nat. Struct. Biol. 1996, 3, 586–589.

(21) Zheng, Y.; Zhang, C.; Croucher, D. R.; Soliman, M. A.; St-Denis, N.; Pascaulescu, A.; Taylor, L.; Tate, S. A.; Hardy, W. R.; Colwill, K.; Dai, A. Y.; Bagshaw, R.; Dennis, J. W.; Gingras, A. C.; Daly, R. J.; Pawson, T. Nature 2013, 499, 166–171.