

Tertiary Amine Coupling by Oxidation for Selective Labeling of Dimethyl Lysine Post-Translational Modifications

Benjamin Emenike, Patrick Czabala, Jonathan Farhi, Jagannath Swaminathan, Eric V. Anslyn, Jennifer Spangle, and Monika Raj*



Cite This: <https://doi.org/10.1021/jacs.4c00253>



Read Online

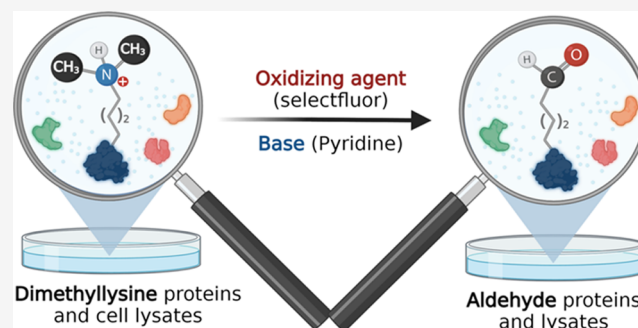
ACCESS |

Metrics & More

Article Recommendations

Supporting Information

ABSTRACT: Lysine dimethylation (Kme_2) is a crucial post-translational modification (PTM) that regulates biological processes and is implicated in diseases. There is significant interest in globally identifying these methylation marks. Unfortunately, this remains challenging due to the lack of robust technologies for selectively labeling Kme_2 . To address this, we present a chemical method named tertiary amine coupling by oxidation (TACO). This method selectively modifies Kme_2 to aldehydes using Selectfluor and a base. The resulting aldehydes from Kme_2 were then functionalized using reductive amination, thiolamine, and oxime chemistry. We successfully demonstrated the versatility of TACO in selectively labeling Kme_2 peptides and proteins in complex cell lysate mixtures with varying payloads, including affinity tags and fluorophores. We further showed the application of TACO chemistry for the identification of Kme_2 sites at a single-molecule level by fluorosequencing. We discovered novel 30 Kme_2 sites, in addition to previously known 5 Kme_2 sites, by proteomics analysis of TACO-modified nuclear extracts. Our work establishes a unique strategy for covalently modifying Kme_2 , facilitating the global identification of low-abundance Kme_2 -PTMs and their sites within complex cell lysate mixtures.



INTRODUCTION

Lysine dimethylation (Kme_2) dynamics constitute a crucial epigenetic event heavily involved in gene expression regulation and various biological processes.^{1–3} Despite increasing interest in this post-translational modification (PTM) and its pivotal role in diverse biological events and diseases beyond chromatin organization, its global identification remains elusive (Figure 1a). This is largely due to the lack of pan-specific chemical tools for labeling dimethyllysine (Kme_2). Attempts to identify and characterize lysine dimethylation Kme_2 have relied heavily on the use of affinity reagents such as antibodies and methyl-binding domains, MBD.^{4–7} However, these reagents fail to globally detect Kme_2 sites due to their sequence-specific nature and limited specificity for different lysine methylation states. Antibodies are further constrained by the unequal abundance of lysine methylation states, reducing the fidelity in measuring methylation levels (Figure 1b). These challenges can be partially attributed to no significant change in the charge and negligible change in the bulk, hydrophobicity, and physicochemical properties of lysine upon the addition of methyl groups (Figure 1b).^{4,8–10} Furthermore, affinity-based methods are unable to detect trypsin-digested Kme_2 fragments because they require flanking amino acids on either side of Kme_2 for efficient recognition.^{11,12} Alternatively, mass spectrometry (MS)-based proteomics for detecting these low-abundant Kme_2 PTMs

require additional modification to reduce sample complexity and minimize false positive due to misassignment resulting from amino acid combinations with similar masses to two methyl groups (e.g., Ala vs Val and Cys vs Met) (Figure 1b).^{10,13,14} Collectively, none of the existing methods can selectively label Kme_2 in a specific manner, underscoring the necessity for robust chemical methods for this purpose. Herein, we introduce a pioneering chemical strategy, termed tertiary amine coupling by oxidation (TACO), designed for the selective labeling of tertiary amines (Kme_2) to aldehydes regardless of the surrounding amino acid sequence (Figure 1c).^{15–18} We demonstrated the broad applicability of TACO chemistry by selectively labeling Kme_2 on peptides, proteins, nucleosomes, and whole cell lysates using various payloads like affinity tags and fluorophores, independent of the sequence, adjacent PTMs, or multiple Kme_2 modifications.

We demonstrated the potential of TACO chemistry in identifying Kme_2 sites at the single molecule level through

Received: January 7, 2024

Revised: March 22, 2024

Accepted: March 25, 2024

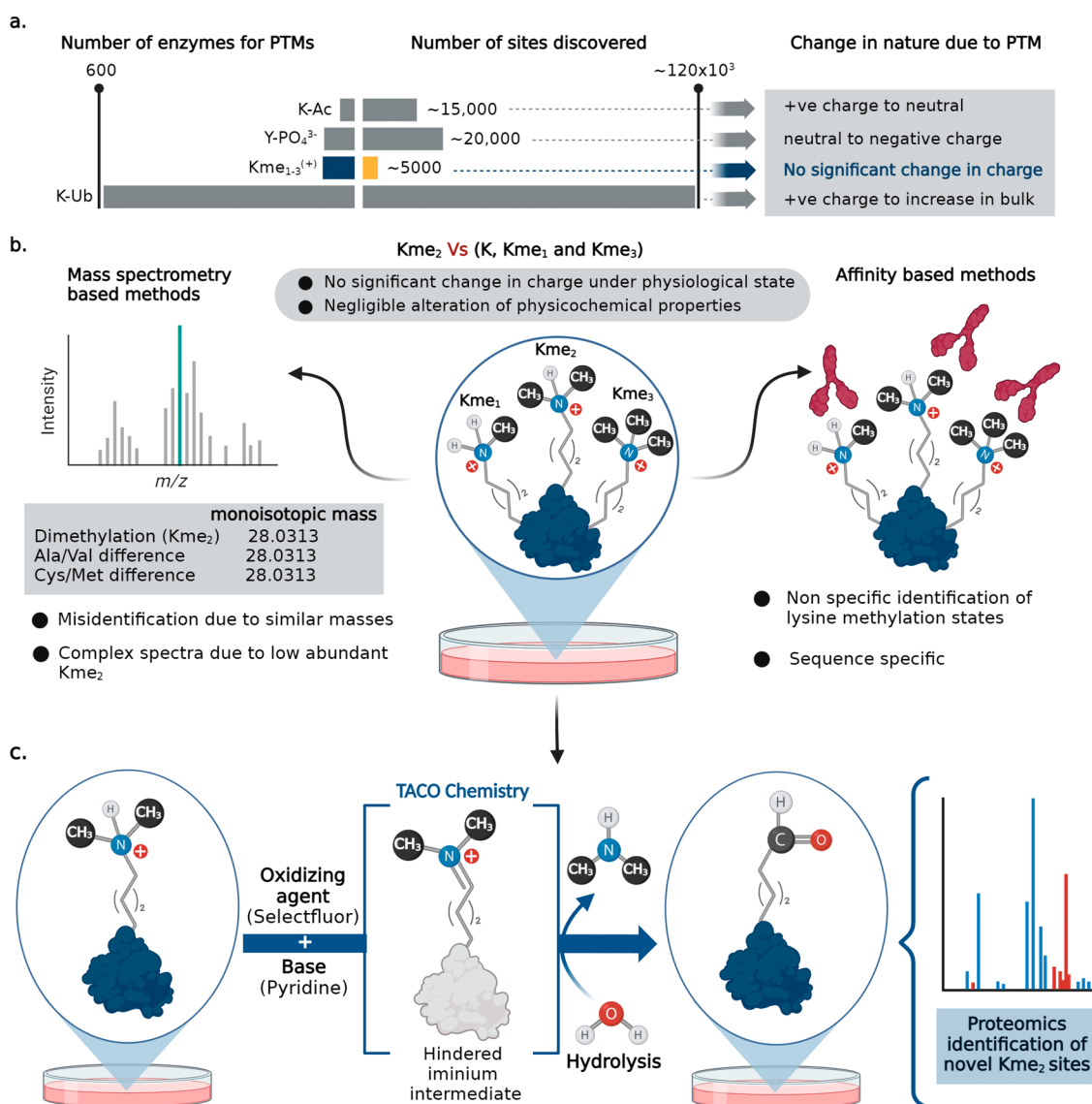


Figure 1. Different lysine methylation states and methods to profile dimethyl lysine. (a) Number of lysine methylation states discovered so far are much lower than enzymes responsible for such PTMs. The major challenge in identifying lysine methylation is due to the negligible change in the charge and physicochemical properties. (b) Limitations of the affinity-based method and MS in identifying lysine methylation. (c) This work: Development of the oxidative coupling chemical reaction for the selective modification of Kme₂ by hydrolysis of iminium ions generating clickable aldehyde sites for identification by proteomics and fluorosequencing.

fluorosequencing. Additionally, we applied TACO chemistry to label and enrich Kme₂ proteins, subsequently conducting a proteomic analysis on the labeled samples. This led to the discovery of 30 novel Kme₂ sites. Notably, there are no chemical methods available in the literature for the site-selective labeling of Kme₂. The capability to selectively label Kme₂ sites using various tags in a complex mixture renders this technology exceptionally valuable for identifying and quantifying Kme₂ biomarkers across diverse biological samples.

RESULTS AND DISCUSSION

Development of TACO. The tertiary amine coupling by oxidation (TACO) reaction capitalizes on the tendency of tertiary amines to generate highly electrophilic iminium ions under oxidative conditions. These ions undergo rapid hydrolysis, leading to the robust conversion of tertiary amines to aldehydes under physiological conditions (Figure 1c).^{15,19} We explored a variety of oxidizing reagents such as Selectfluor,

tropylium tetrafluoroborate, N-bromo succinimide (NBS), diethyl azodicarboxylate (DEAD), and *t*BuOOH (TBHP) with FeCl₃ under hydrolytic conditions on a model peptide FKme₂V 1a (Figures 2a and Supporting Figure 1). Of all the reactions, Selectfluor (2 equiv) resulted in the formation of an aldehyde product, FKme₂(CHO)V 2a, with 9% conversion in 1 h in sodium phosphate buffer (10 mM, pH 7) at room temperature. However, this reaction also generated small amounts of monomethyllysine FKme₁V 2a', which coeluted with the starting peptide FKme₂V 1a as determined by LCMS (Supporting Figure 1). We postulated that the formation of aldehyde 2a and monomethyllysine 2a' resulted from the nucleophilic attack of water on the more hindered and less hindered iminium ion, respectively (Supporting Figure 2). We did not observe the formation of any aldehyde product in the presence of other oxidants such as tropylium, NBS, DEAD, and (TBHP + FeCl₃), due to their poor water compatibility and instability in aqueous conditions. To enhance the conversion of

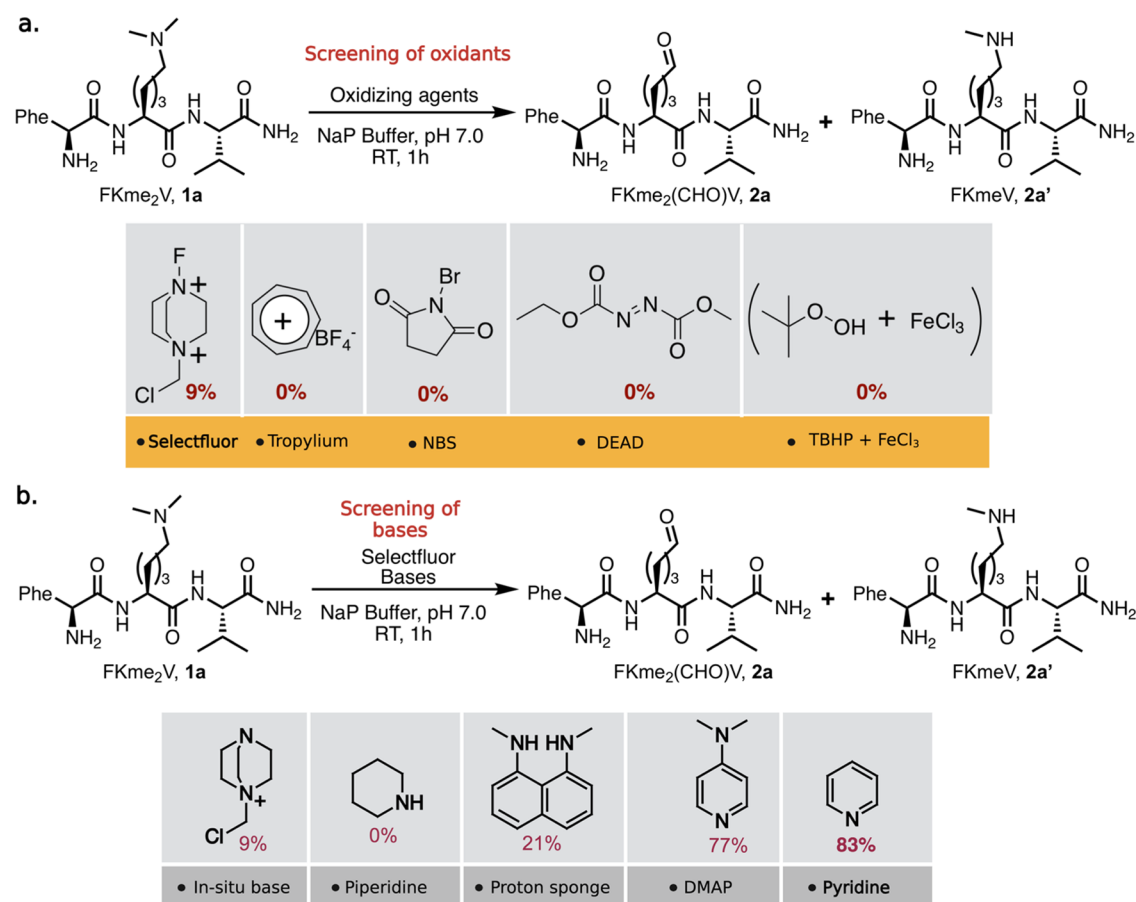


Figure 2. Optimization of the TACO chemistry. (a) Development of the TACO reaction for modification of dimethyllysine Kme₂ in peptide FKme₂V **1a** by exploring varying oxidizing reagents. (b) Evaluation of bases and optimization to the peptide aldehyde FKme₂(CHO)V **2a**. Reaction conditions: To 6–10 mM of dimethyllysine peptide in 300 μ L of 10 mM sodium phosphate buffer (NaP, pH 7.0) were added a base (5 equiv) and oxidizing reagent (2 equiv), and the reaction mixture was stirred for 1 h at room temperature.

the dimethyl lysine peptide to the peptide aldehyde, we introduced bases stronger than those generated in situ by Selectfluor. Screening various bases such as piperidine, proton sponge, DMAP, and pyridine, we observed high conversion to the peptide aldehyde product FKme₂(CHO)V **2a** with DMAP (77%) and pyridine (83%) (Figures 2b and Supporting Figure 3). We attribute the high conversion to the planarity of pyridine and DMAP that enables the abstraction of the hindered methylene proton of dimethyl lysine, resulting in the formation of iminium ions followed by hydrolysis to generate an aldehyde. Other bases such as piperidine, DABCO-like base from Selectfluor, and proton sponge are too bulky to remove the proton from the hindered carbon of dimethyl lysine. Next, we generated high amounts of peptide aldehyde (OAc)-FKme₂(CHO)V **2b** from peptide (OAc)FKme₂V **1b** under optimized reaction conditions and characterized it by NMR (Supporting Figure 4).

Chemoselectivity Studies for Peptide Aldehyde.

Chemoselectivity studies for peptide aldehyde revealed the specificity of the TACO reaction for generating aldehyde products with Kme₂ only, as demonstrated by varying peptides FXV with different lysine methylation states (where X = K, Kme, and Kme₂) and reactive amino acids (where X = H, D, S, M, Y, C, R, N, W) (Figures 3a and Supporting Figure 5). Our observations indicated the oxidation of Met and Cys, fluorination of Trp and His, and no formation of quinone and the ipso-fluorinated product with tyrosine under the reaction

conditions.^{21–24} To further evaluate the effect of these side adducts on the enrichment of Kme₂-generated aldehyde sites, we synthesized four dimethyllysine tripeptides with methionine (FKme₂M), histidine (FKme₂H), tryptophan (FKme₂W), and cysteine (FKme₂C) and subjected them to the optimized TACO reaction conditions. The reactions generated the corresponding aldehyde adducts, FK(CHO)M(sulfoxide), FK(CHO)H-(fluorine), FK(CHO)W(fluorine), and FK(CHO)C(disulfide) in high conversions (Figures 3b and Supporting Figure 6). Further, treatment of aldehyde adducts with hydroxylamine and amino-thiol (cysteine methyl ester) generated near-quantitative conversions to the oxime and thiazolidine products. These data strongly validate the negligible effects of TACO-generated adducts on downstream chemoproteomics application on Kme₂ in complex mixtures (Figures 3b and Supporting Figure 6).

Application of TACO for Diversification of Dimethyllysine on Histone Peptides. To demonstrate the application of TACO for diversification of histone peptides, we conducted reactions on various peptides of different sizes and amino acid compositions, including histone H3.3 peptide fragments frequently methylated at K4, K9, K27, and K36, known to regulate biological processes and implicated in disease states.^{25–27} Using solid-phase peptide synthesis, we synthesized the H3.3 peptide fragments Kme₂4K9 (ARTKme₂QTARKS) **1c**, Kme₂4Kme₂9 (ARTKme₂QTARKme₂S) **1d**, Kme₂9K14 (ARTKme₂STGGKA) **1e**, propargyl amine containing peptide AKme₂(CHO)GSKAF(PrG)A (where PrG = propargyl gly-

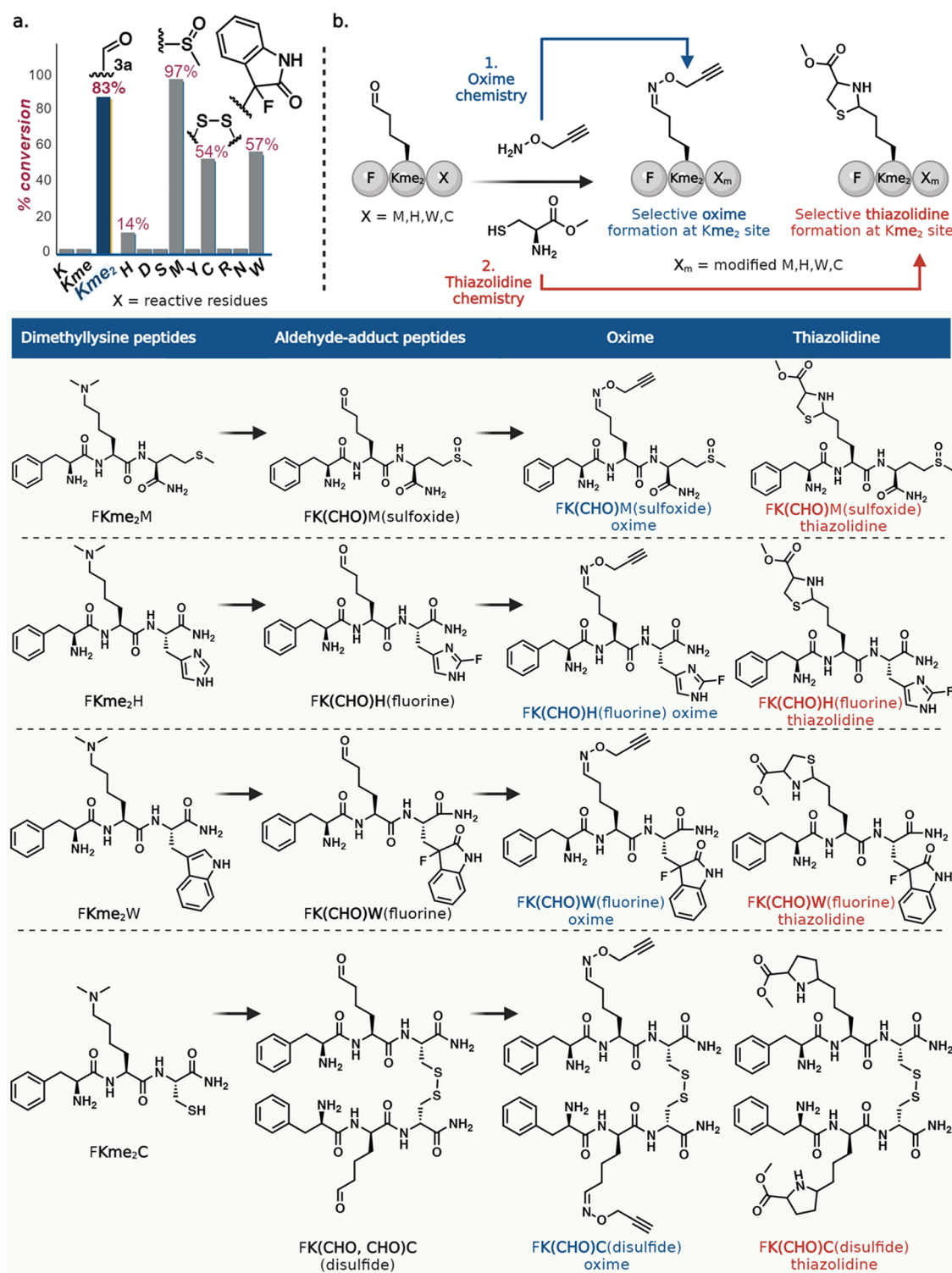


Figure 3. Evaluation of chemoselectivity of the TACO reaction. (a) Chemoselectivity studies of the TACO reaction with varying tripeptides FXV, where X = K, Kme, Kme₂, H, D, S, M, Y, C, R, N, and W. Only the Kme₂ residue modified to an aldehyde under the reaction conditions. Side adducts were observed with histidine (fluorination), methionine (sulfoxide), tryptophan (fluorination and oxygenation), and cysteine (disulfide). (b) Selective derivatization of Kme₂-generated aldehydes on tripeptides FKme₂X, where X = M, H, W, and C, in the presence of side adducts on histidine, methionine, cysteine, and tryptophan using oxime and thiazolidine chemistry in NaP buffer. The presence of TACO-generated adducts did not interfere with subsequent modification of dimethyl lysine sites, thus indicating the negligible effect of the adducts on downstream application of TACO for profiling of lysine dimethylome.

cine) **1f**, and a negative control peptide K4K9 (ARTKSTGG-KA) **1g**. Under optimized conditions, all Kme₂-containing peptides underwent modification to aldehyde peptide products (Figures 4a and Supporting Figure 7). Intriguingly, the H3.3

peptide fragment Kme₂4Kme₂9 (ARTKme₂QTARKme₂S) **1d**, containing two Kme₂, produced the double aldehyde product ARTKme₂(CHO)QTARKme₂(CHO)S **2d** with >98% conversion (Supporting Figure 7). Conversely, the negative control

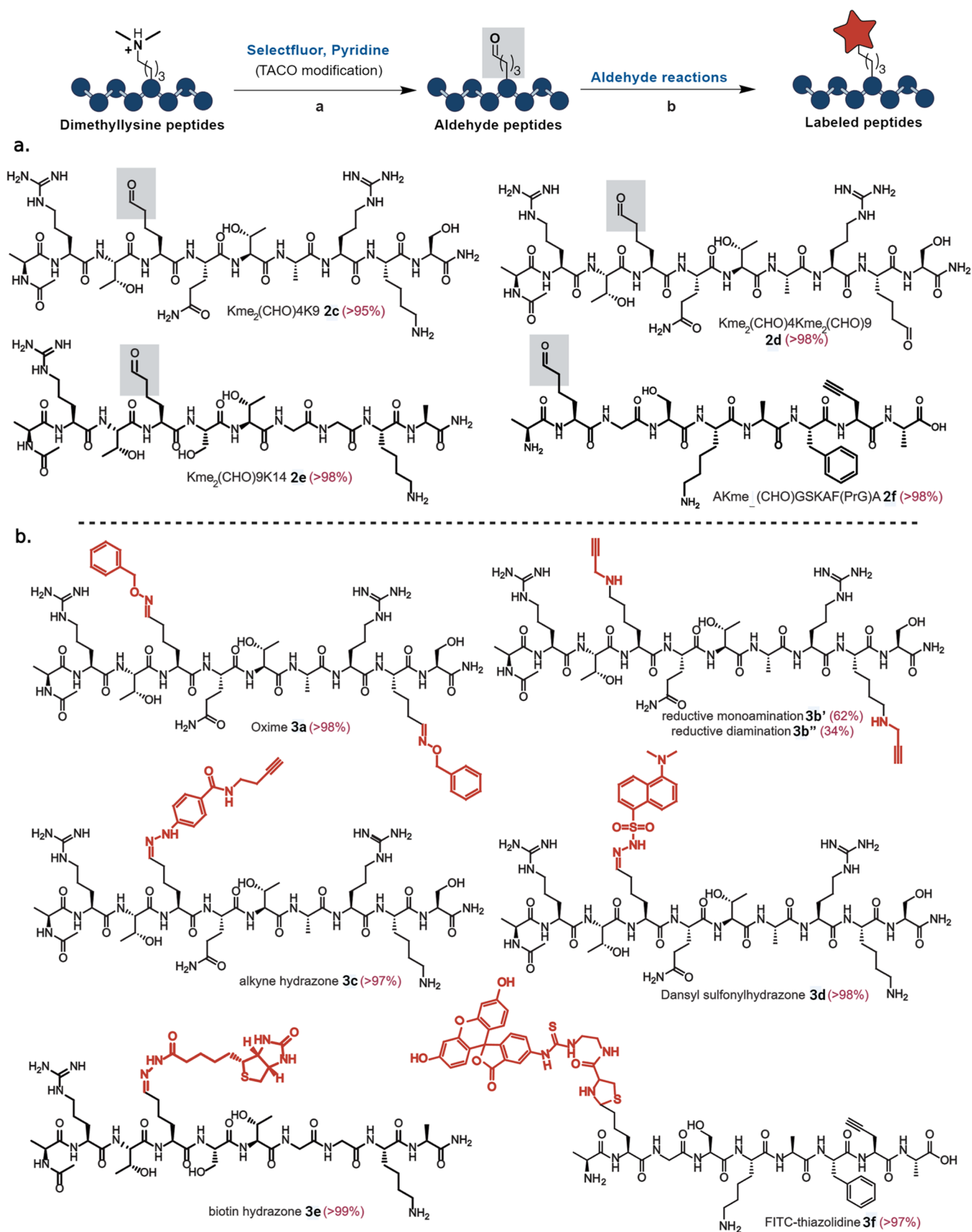


Figure 4. Application of the TACO reaction for Kme_2 and further derivatization with varying payloads. (a) TACO-mediated modification of histone H3.3 peptide (1c–1f) fragments with Kme_2 to aldehydes (2c–2f). Reaction conditions: To 6–10 mM of dimethyllysine peptide in 300 μL of 10 mM sodium phosphate buffer (NaP, pH 7.0) were added pyridine (5 equiv) and Selectfluor (2 equiv) and the reaction mixture was stirred for 1 h at room temperature. (b) Late-stage diversification of peptide aldehydes with varying chemistries and payloads.

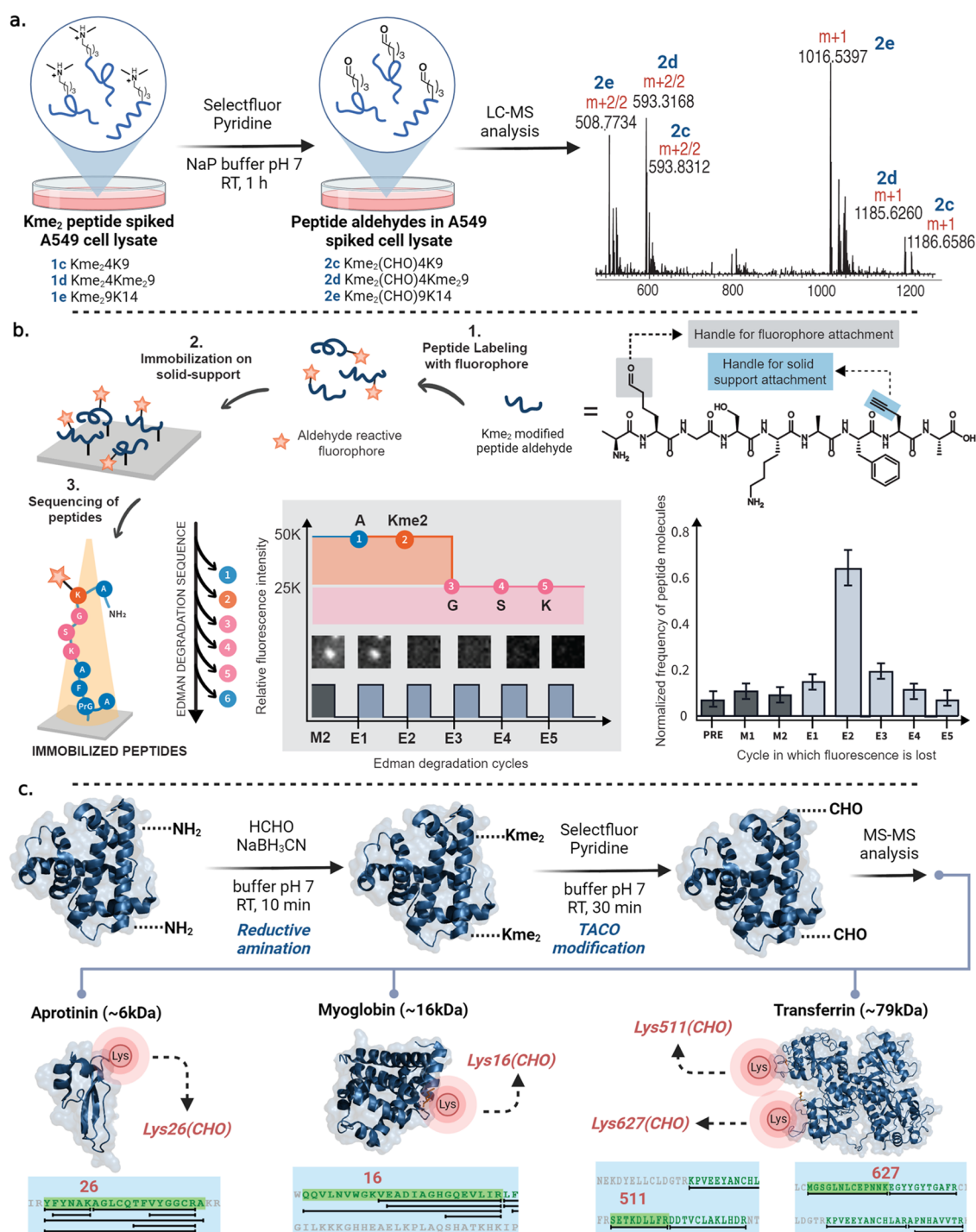


Figure 5. Labeling and identification of Kme₂ peptides and proteins in a complex mixture. (a) Selective modification of Kme₂ peptides in a complex mixture of A549 lung cancer cell lysate to peptide aldehydes in a quantitative manner. Reaction conditions: To 100 μg of A549 whole cell lysate in 200 μL of NaP buffer pH 7.0 was added 0.1 mg of each histone peptide followed by addition of pyridine (30 equiv with respect to peptides) and selectfluor (10 equiv with respect to peptides). The reaction mixture was stirred for 1 h. (b) Application of TACO in the identification of Kme₂ sites on a peptide by single molecule fluorosequencing. The left panel provides an exemplary view of the single fluorescent peptide molecule through cycles of Edman degradation. The right panel indicates the normalized counts of the labeled Kme₂ peptide where its fluorescence intensity was lost. (c) Selective modification of chemically generated Kme₂ proteins by TACO chemistry followed by the identification of Kme₂ sites on proteins by LC-MS/MS analysis.

peptide K4K9, ARTKSTGGKA **1g** without Kme₂, remained unmodified (Figures 4a and Supporting Figure 7). These results unequivocally confirmed the application of TACO toward dimethyllysine Kme₂, marking the first chemical method developed for Kme₂ modification.

We further diversified peptide aldehydes using aldehyde-specific reactions, such as oxime, hydrazone, thiazolidine chemistry, and reductive amination (Figure 4b). For instance, the histone peptide ARTKme₂(CHO)QTARKme₂(CHO)S **2d** with double aldehyde groups underwent modifications with

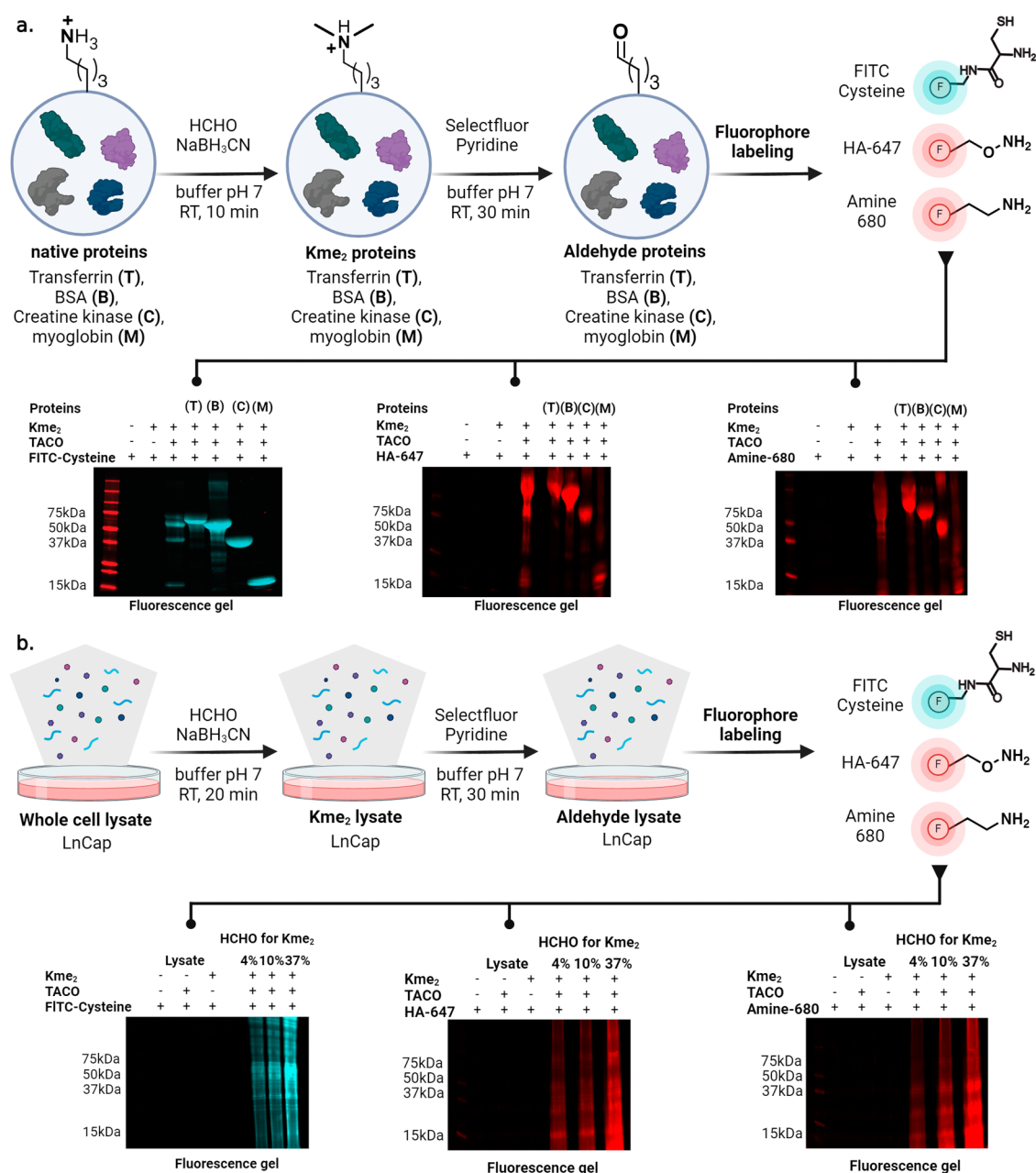


Figure 6. Selective labeling of Kme₂ proteins by TACO chemistry using varying fluorophores. (a) Selective modification of chemically generated Kme₂ proteins to aldehyde proteins by TACO chemistry followed by labeling with varying fluorophores using different aldehyde chemistries as analyzed by running SDS PAGE. No fluorophore labeling was observed without the TACO reaction. (b) Selective modification of chemically generated Kme₂ proteins in cell lysate to aldehyde proteins by TACO followed by fluorophore labeling and in-gel analysis. High fluorescence was observed in cell lysate in a dose-dependent manner, indicating high amounts of Kme₂-proteins. No fluorophore labeling was observed on cell lysate without the TACO reaction or without Kme₂-modified cell lysate.

benzylhydroxylamine, generating the oxime product **3a** in >98% yield (Figures 4b and Supporting Figure 8). Additionally, this double-aldehyde peptide was treated with propargylamine and sodium cyanoborohydride, resulting in reductive amination products, single modification **3b'** (62%) and double modification **3b''** (34%) in high conversions (96%) (Figures 4b and Supporting Figure 9).

Next, we functionalized Kme₂4K9-aldehyde (ARTKme₂(CHO)QTARKS) **2c** with alkyne- and dansyl-tagged hydrazines to generate hydrazone products **3c** and **3d** in >97% and >98% conversions, respectively (Figure 4b and Supporting Figures 10 and 11). Biotin hydrazine treatment of

the peptide aldehyde (ARTKme₂(CHO)STGGKA) **2e** resulted in 99% conversion to biotin peptide hydrazone **3e** (Figures 4b and Supporting Figure 12). Concurrently, introducing both an affinity handle and a fluorescent dye into a peptide (AKme₂(CHO)GSKAF(PrG)A, where PrG = propargyl glycine) **2f** via derivatization with FITC-cysteine yielded the thiazolidine conjugation product **3f** in >97% conversion (Figures 4b and Supporting Figure 13). Moreover, treatment of peptide aldehyde FKme₂(CHO)V **2a** with cysteine resulted in thiazolidine conjugation with high conversion (>99%, Supporting Figure 13). These outcomes underscore the broad scope and

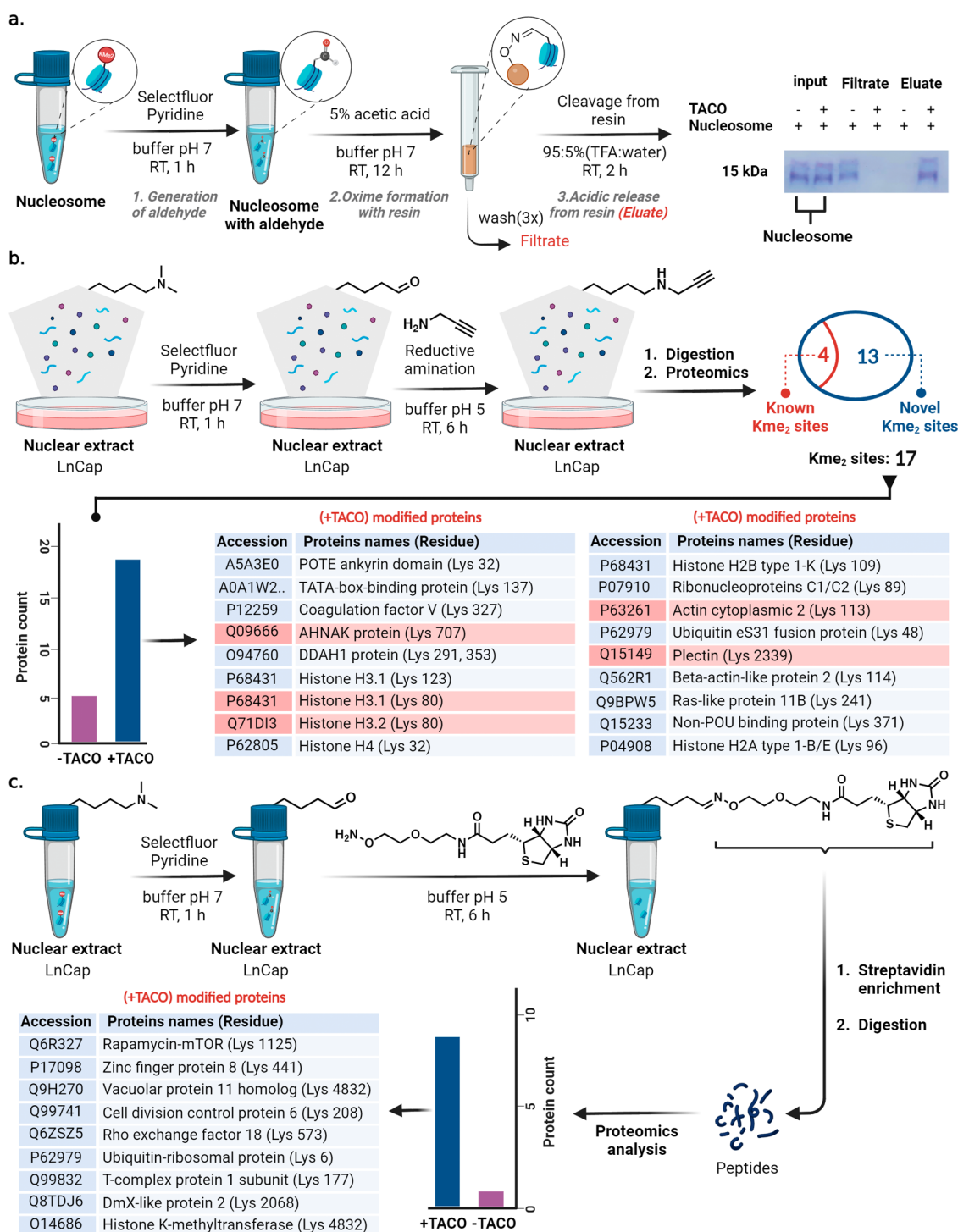


Figure 7. Discovery of naturally occurring Kme_2 proteins and Kme_2 -sites from a complex cell lysate was carried out by TACO chemistry. (a) Selective modification of naturally occurring dimethyl lysine in nucleosomes using TACO chemistry followed by the enrichment of Kme_2 -generated aldehyde proteins using hydroxylamine resin and release of trapped proteins under acidic conditions. The filtrate contains proteins remaining after the enrichment process with and without TACO chemistry on nucleosomes, while the eluate comprises proteins enriched from the nucleosomes with and without TACO chemistry using a solid support. The filtrate and eluate were analyzed by SDS PAGE analysis. The gels showed the enrichment of Kme_2 proteins from nucleosomes under TACO reaction conditions. (b) Selective modification of Kme_2 proteins by TACO chemistry followed by trapping of Kme_2 -generated aldehyde proteins by reductive amination and proteomic analysis. Twelve novel Kme_2 proteins and thirteen new Kme_2 sites were identified (novel = blue, known = red). In the negative control, five proteins were identified, which vary from the TACO-treated samples due to the presence of natural occurring allysine on proteins. (c) Enrichment of TACO-modified Kme_2 proteins followed by proteomic analysis. Nine low-abundance Kme_2 proteins were discovered with nine new Kme_2 sites.

efficiency of TACO chemistry in selectively modifying and diversifying Kme_2 -containing peptides with various payloads.

Selective Labeling of Kme_2 Peptides by TACO in a Complex Cell Lysate Mixture. To evaluate the robustness of our TACO technique in labeling Kme_2 peptides within a

complex mixture, we spiked cell lysates from epithelial carcinoma A549 and H1792 lung cancer cell lines (100 μg) with three different histone H3.3 peptides (0.1 mg each): Kme₂4K9 (ARTKme₂QTARKS) **1c**, Kme₂4Kme₂9 (ARTKme₂QTARKme₂S) **1d**, and Kme₂9K14 (ARTKme₂STGGKA) **1e** and **1c**, **1d**, and **1g**, respectively. Both complex mixtures were incubated with Selectfluor and pyridine for 1 h, followed by LC-MS analysis (Figure 5a, A549 Supporting Figures 14 and 15). All peptides containing Kme₂ (**1c–1e**) were successfully converted to the corresponding peptide aldehydes (**2c–2e**) without any detectable traces of the unmodified starting peptides in the complex cell lysate mixtures (Figure 5a, A549 Supporting Figures 14 and 15). Notably, peptide **1g**, K4K9 (ARTKSTGGKA), lacking Kme₂, showed no modification in the H1792 cell lysate (Supporting Figure 15).

Identification of Kme₂ Sites by TACO at a Single-Molecule Level Using Fluorosequencing. To determine the site of Kme₂ in a peptide at a single-molecule level, we modified a model peptide, AKme₂GSKAF(PrG)A (where PrG = propargyl glycine), by TACO chemistry to peptide aldehyde, AKme₂(CHO)GSKAF(PrG)A. Next, we functionalized the Kme₂-aldehyde group with an Atto647N fluorophore by using dithiolane chemistry under acidic conditions and subjected it to a fluorosequencing workflow including immobilization on the azide-functionalized microscopic slide using PrG on a peptide, AKme₂^{Atto647N}GSKAF(PrG)A, by click chemistry (Figures 5b and Supporting Figure 16). In the case of biologically derived peptides, the C-terminus will be used for immobilization using a photoredox catalyst that can differentiate oxidation potential of the C-terminus from the side chains of Asp and Glu, leading to C-terminal immobilization exclusively.²⁸ Next, the fluorophore-labeled immobilized peptide was subjected to several rounds of Edman's degradation including one prerun and two rounds of mock Edman's degradation (M1-M2) with all the reagents except phenylisothiocyanate followed by analysis using a total internal reflection fluorescence (TIRF) microscope (Figure 5b).²⁹ No decrease in the fluorescence was observed under the mock conditions, suggesting the high stability of the Kme₂-modified fluorophore under Edman's degradation conditions requiring TFA and pyridine. Next, Edman's degradation was performed on a peptide with phenylisothiocyanate followed by the analysis of fluorescence by TIRF after each round. A significant decrease in the fluorescence was observed after the second round of Edman's degradation, confirming the site of Kme₂ at a single-molecule level. Subsequent rounds of Edman's degradation did not lead to a change in the fluorescence. There are no other chemical methods for the identification of sites of Kme₂ PTM by any other single-molecule protein sequencing SMPS techniques.³⁰ Future studies will be focused on discovering proteins containing Kme₂ and the sites of Kme₂ on a proteome-wide scale by fluorosequencing. The work under this direction is currently underway in our laboratory.

Modification of Proteins and Whole Cell Lysate by TACO. To demonstrate the compatibility of TACO chemistry with proteins, we chemically modified lysines (K) to dimethyl lysine (Kme₂) on proteins of varying molecular weights, aprotinin (A, 5 kDa), myoglobin (M, 16 kDa), and transferrin (T, 79 kDa) using reductive amination. Proteins were treated with a solution of formaldehyde and sodium cyanoborohydride in citrate buffer for 10 min. Subsequently, Kme₂ proteins were subjected to TACO chemistry to generate aldehyde proteins. The MS/MS analysis of digested aldehyde proteins identified K26 (aprotinin), K16 (myoglobin), K511 (transferrin), and

K627 (transferrin) as Kme₂ modification sites (Figures 5c and Supporting Figure 17). Moreover, four proteins of varying molecular weights and 3 D structures, transferrin (T), BSA (B), creatine kinase (C), and myoglobin (M), were subjected to reductive amination followed by TACO chemistry to produce aldehyde proteins. These aldehyde proteins were incubated with various fluorophores containing diverse reactive aldehyde functional groups (FITC-cysteine, hydroxylamine (HA)-647 and amine-680) and analyzed via in-gel fluorescence (Figure 6a). The results showed the selective labeling with aldehyde proteins only (lanes 3–7, lane 3 = mixture of all four proteins, Figures 6a and Supporting Figure 18). No fluorophore labeling was observed with native proteins and Kme₂ proteins (lanes 1, 2, Figures 6a and Supporting Figure 18). This result demonstrated the ability of TACO chemistry to selectively modify Kme₂ sites on proteins to aldehydes, which can then be further modified with varying payloads.

To further demonstrate the robustness of TACO chemistry to modify proteins within cell lysates, we treated prostate cancer cell lysate (LnCap) with reductive amination reagents (formaldehyde and sodium cyanoborohydride) in a dose-dependent manner (4, 10, and 37%), followed by TACO chemistry and subsequent labeling with FITC-cysteine, hydroxylamine (HA)-647, and amine-680 fluorophores (Figures 6b and Supporting Figure 19). Analysis and quantification of in-gel fluorescence results clearly showed selective fluorophore labeling of aldehyde-modified cell lysate and a dose-dependent increase in the fluorescence intensity from 4 to 37% of reductive amination reagents (with Cys-FITC, 37% lane = 1.0, 10% lane = 0.582, 4% lane = 0.420; with HA-647, 37% lane = 1.0, 10% lane = 0.513, 4% lane = 0.459; with amine-680, 37% lane = 1.0, 10% lane = 0.672, 4% lane = 0.373) (lanes 4–6, Figures 6b and Supporting Figure 19). No fluorescence was observed with untreated LnCap cell lysate, Kme₂-modified cell lysate, and TACO-treated cell lysate without Kme₂ modification (lanes 1–3, Figure 6b and Supporting Figure 19), confirming the high selectivity of TACO chemistry for Kme₂.

Selective Enrichment of Kme₂ Proteins from Nucleosomes by TACO. To explore the capability of TACO chemistry in the selective modification and enrichment of naturally occurring Kme₂ on histones, we exposed nucleosomes from prostate cancer cell lysate (LnCap) to Selectfluor and pyridine for 1 h, followed by the enrichment of aldehyde proteins via oxime chemistry using hydroxylamine-functionalized resin in pull-down experiments. The resin was thoroughly washed with solvents to remove noncovalently bound proteins (filtrate). Subsequently, enriched proteins were released from the resin under acidic conditions (95% TFA in water).

SDS-PAGE analysis of the cell lysate (filtrate) postenrichment and protein release from the solid support (eluate) distinctly demonstrated the efficient capture and release of Kme₂-modified protein aldehydes (Figure 7a, lanes 4 and 6, Supporting Figures 20 and 21). Repetition of this reaction in duplicate showed consistent gel profiles across all replicates, confirming the reproducibility, selectivity, and robustness of the TACO approach in tagging and enriching naturally occurring Kme₂ proteins from a complex nucleosome mixture. Enrichment of nucleosomes by hydroxylamine resin without the TACO reaction failed to capture proteins (lane 3 (filtrate) and lane 5 (eluate), Figure 7a). Lanes 1 and 2 represent total nucleosomes with and without TACO reagents, respectively (Figure 7a).

Identification and Discovery of Kme₂ Proteins and Kme₂ Sites by TACO from a Complex Cell Lysate. To

demonstrate an exciting application of TACO chemistry for identifying and discovering naturally occurring Kme₂ proteins, we treated nuclear extracts obtained from LnCap cells with TACO chemistry followed by labeling of Kme₂-generated aldehyde proteins with propargylamine using reductive amination. We performed this reaction to increase the stability of the adducts under proteomics conditions and to add a tag for identification. By digesting and analyzing the cellular mixture via proteomics, we discovered 13 Kme₂ sites that had not been previously identified, along with four Kme₂-sites previously reported by other methods³¹ (Figures 7b and Supporting Figure 22). The treatment of nuclear extract of MCF-10A, a preadenocarcinoma cell, with TACO chemistry led to the identification of nine Kme₂ sites with one known Kme₂ site, K240 of actin, α cardiac muscle 1 (Supporting Figure 22). Notably, none of the identified Kme₂ proteins in MCF-10A cells match with Kme₂ proteins observed in LnCap cancer cells, thus suggesting the potential of this approach in discovering dimethyllysine biomarkers. Moreover, proteomics analysis of nuclear extracts without TACO modification (negative control) identified five proteins and sites, differing from those observed in the TACO-treated sample. We attribute this modification to the presence of allysine, an endogenous aerobic oxidation aldehyde product of lysine.³²

Furthermore, we developed an enrichment protocol for trapping Kme₂-generated aldehyde proteins obtained through TACO chemistry to identify low-abundant Kme₂ proteins from a complex nuclear extract mixture of LnCap cells. Employing biotin-derivatized alkoxyamine probes to label aldehyde proteins, followed by streptavidin enrichment and digestion of enriched proteins, led to the discovery of nine novel low-abundant Kme₂ proteins and identification of nine new Kme₂ sites (Figures 7c and Supporting Figure 23). Taken together, these results demonstrate the potential of TACO chemistry to serve as a chemical platform for the identification and discovery of the lysine dimethylome.

CONCLUSIONS

In summary, we have developed TACO, a versatile chemical method for modifying dimethyllysine Kme₂ to an aldehyde group within peptides, proteins, and cell lysates under physiological conditions. The TACO chemistry involves the oxidation of dimethyl lysine/Kme₂ by Selectfluor, followed by hydrolysis of the resulting iminium ions to generate aldehydes. Our study demonstrates the robustness of TACO chemistry for selectively labeling of Kme₂-containing peptides, proteins, and cell lysates with diverse payloads, including affinity tags such as alkyne, biotin, and various fluorophores, by diverse aldehyde chemistries in high conversions (>90%). Moreover, we showcased the compatibility of TACO chemistry with fluorosequencing, thus demonstrating the potential of identifying Kme₂ sites at a single-molecule level. Applying TACO chemistry to the nuclear extract of cancer and preadenocarcinoma cells, we discovered 30 novel Kme₂ sites, in addition to 5 previously reported sites using MS/MS analysis. This demonstrates the capability of TACO chemistry to enable the identification of Kme₂ proteins and their sites on a proteome-wide scale, through either MS-based proteomics or single molecule fluorosequencing. We compared proteome profiling of diseased cells to healthy controls, and none of the identified Kme₂ proteins in MCF-10A cells match with Kme₂ proteins observed in LnCap cancer cells, thus suggesting the potential of this approach in discovering Kme₂ proteins and Kme₂ sites as

disease biomarkers. The work in these directions is currently underway in our laboratory. We believe that our chemical approach will complement existing detection methods, significantly expanding the chemical toolbox available for epigenetics research.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/jacs.4c00253>.

List of proteins (TACO) (XLSX)

Optimization of the TACO reaction with varying oxidizing reagents and nucleophiles; reaction procedures; procedure of TACO reaction with peptides; proteins; cell lysates; nucleosomes; fluorosequencing; enrichment protocol; proteomic analysis and product characterization by NMR; HPLC; HRMS and computational data; and raw images for fluorosequencing uploaded to Zenodo (PDF)

AUTHOR INFORMATION

Corresponding Author

Monika Raj – Department of Chemistry, Emory University, Atlanta, Georgia 30322, United States; orcid.org/0000-0001-9636-2222; Email: monika.raj@emory.edu

Authors

Benjamin Emenike – Department of Chemistry, Emory University, Atlanta, Georgia 30322, United States

Patrick Czabala – Department of Chemistry, Emory University, Atlanta, Georgia 30322, United States; orcid.org/0000-0002-2365-3181

Jonathan Farhi – Department of Radiation Oncology, Emory University School of Medicine, Atlanta, Georgia 30322, United States

Jagannath Swaminathan – Department of Chemistry, University of Texas at Austin, Austin, Texas 78712, United States

Eric V. Anslyn – Department of Chemistry, University of Texas at Austin, Austin, Texas 78712, United States; orcid.org/0000-0002-5137-8797

Jennifer Spangle – Department of Radiation Oncology, Emory University School of Medicine, Atlanta, Georgia 30322, United States; orcid.org/0000-0002-0181-3244

Complete contact information is available at: <https://pubs.acs.org/doi/10.1021/jacs.4c00253>

Author Contributions

The article was written through contributions of all authors. All authors have given approval to the final version of the article.

Notes

The authors declare no competing financial interest.

B.E. and M.R. are the inventors of a United States patent application (PCT/US2022/076782) of Emory University covering the methodology of the TACO reaction for modification of Kme₂ and its application in global profiling and identification of Kme₂ sites from a complex biological system. J.S. and E.V.A. are cofounders and shareholders of Erisyon, Inc. E.V.A. serves on the scientific advisory board of Erisyon.

ACKNOWLEDGMENTS

This research was supported by the NIH (No. 1R35GM133719-01) and NSF (Grant No. CHE-2108774) to M.R. and Research Scholar Grant, RSG-22-025-01-CDP, from the American Cancer Society to M.R. The research was supported by the NIH (No. R35 GM149308-01) to E.V.A. J.S. work was supported through a sponsored research agreement from Erisyon to University of Texas (E.V.A.). The authors thank Heather D Stout for running duplicates of the fluorosequencing experiment. All the images are created with biorender.com.

ABBREVIATIONS

Kme₂, lysine dimethylation; PTM, post-translational modification; TACO, tertiary amine coupling by oxidation; Kme₁, monomethyl lysine; MS, mass spectrometry; TIRF, total internal reflection fluorescence; MBD, methyl-binding domains; PrG, propargyl glycine; DEAD, diethyl azodicarboxylate

REFERENCES

- (1) Moss, T. J.; Wallrath, L. L. Connections between epigenetic gene silencing and human disease. *Mutat. Res., Fundam. Mol. Mech. Mutagen.* **2007**, *618*, 163–174.
- (2) Egorova, K. S.; Olenkina, O. M.; Olenina, L. V. Lysine methylation of nonhistone proteins is a way to regulate their stability and function. *Biochemistry* **2010**, *75*, 535–548.
- (3) Yao, X.; Shen, W. Crucial function of histone lysine methylation in plant reproduction. *Chin. Sci. Bull.* **2011**, *56*, 3493–3499.
- (4) Moore, K. E.; Carlson, S. M.; Camp, N. D.; et al. A general molecular affinity strategy for global detection and proteomic analysis of lysine methylation. *Mol. Cell* **2013**, *50*, 444–456.
- (5) Harrison, E. E.; Carpenter, B. A.; Louis, L. E., St.; Mullins, A. G.; Waters, M. L. Development of “Imprint-and-Report” dynamic combinatorial libraries for differential sensing applications. *J. Am. Chem. Soc.* **2021**, *143*, 14845–14854.
- (6) Guo, A.; Gu, H.; Zhou, J.; Mulhern, D.; Wang, Y.; Lee, K. A.; et al. Immunoaffinity enrichment and mass spectrometry analysis of protein methylation. *Mol. Cell. Proteomics* **2014**, *13*, 372–387.
- (7) Emenike, B.; Nwajiobi, O.; Raj, M. Covalent chemical tools for profiling post-translational modifications. *Front. Chem.* **2022**, *10*, No. 868773.
- (8) Polevoda, B.; Sherman, F. Methylation of proteins involved in translation. *Mol. Microbiol.* **2007**, *65*, 590–606.
- (9) Bremang, M.; Cuomo, A.; Agresta, A. M.; Stugiewicz, M.; Spadotto, V.; Bonaldi, T. Mass spectrometry-based identification and characterisation of lysine and arginine methylation in the human proteome. *Mol. BioSyst.* **2013**, *9*, 2231–2247.
- (10) Wang, Q.; Wang, K.; Ye, M. Strategies for large-scale analysis of non-histone protein methylation by LC-MS/MS. *Analyst* **2017**, *142*, 3536–3548.
- (11) Zoghbi, H. Y.; Beaudet, A. L. Epigenetics and human disease. *Cold Spring Harbor Perspect. Biol.* **2016**, *8*, No. a019497.
- (12) Busby, M.; Xue, C.; Li, C.; Farjoun, Y.; Gienger, E.; Yofe, I.; et al. Systematic comparison of monoclonal versus polyclonal antibodies for mapping histone modifications by ChIP-seq. *Epigenet. Chromatin* **2016**, *9*, No. 49.
- (13) Young, N. L.; DiMaggio, P. A.; Plazas-Mayorca, M. D.; Baliban, R. C.; Floudas, C. A.; Garcia, B. A. High throughput characterization of combinatorial histone codes. *Mol. Cell Proteomics* **2009**, *8*, 2266–2284.
- (14) Nwajiobi, O.; Mahesh, S.; Streety, X.; Raj, M. Selective triazeneation reaction (STaR) of secondary amines for tagging monomethyl lysine post-translational modifications. *Angew. Chem., Int. Ed.* **2021**, *60*, 7344–7352.
- (15) Daniels, M. H.; Hubbs, J. The development of a Selectfluor-mediated oxidative Mannich reaction. *Tetrahedron Lett.* **2011**, *52*, 3543–3546.
- (16) Kumaraswamy, G.; Murthy, A. N.; Pitchaiah, A. FeCl₃-catalyzed oxidative allylation of sp² and sp³ C-H bond adjacent to a nitrogen atom: easy access to homoallyl tertiary amines. *J. Org. Chem.* **2010**, *75*, 3916–3919.
- (17) Niu, M.; Yin, Z.; Fu, H.; Jiang, Y.; Zhao, Y. Copper-catalyzed coupling of tertiary aliphatic amines with terminal alkynes to propargylamines via C–H Activation. *J. Org. Chem.* **2008**, *73*, 3961–3963.
- (18) Xu, X.; Li, X. Copper/diethyl azodicarboxylate mediated regioselective alkylation of unactivated aliphatic tertiary methylamine with terminal alkyne. *Org. Lett.* **2009**, *11*, 1027–1029.
- (19) Hauser, A.; Bohlmann, R. Preparation of aldehydes by oxidation of benzylic amines with selectfluor (F-TEDA-BF₄). *Synlett* **2016**, *27*, 1870–1872.
- (20) Seki, T.; Fujiwara, T.; Takeuchi, Y. A facile procedure for synthesis of 3-[2-(N,N-dialkylamino)ethyl]-3-fluorooxindoles by direct fluorination of N,N-dialkyltryptamines. *J. Fluorine Chem.* **2011**, *132*, 181–185.
- (21) Emenike, B.; Donovan, J.; Raj, M. Multicomponent oxidative nitrile thiazolidination reaction for selective modification of N-terminal dimethylation posttranslational modification. *J. Am. Chem. Soc.* **2023**, *145*, 16417–16428.
- (22) Karam, O.; Martin-Mingot, A.; Jouannetaud, M.-P.; Jacquesy, J.-C.; Cousson, A. Efficient oxidative ipso-fluorination of para-substituted phenols using pyridinium polyhydrogen fluoride in combination with hypervalent iodine(III) reagents. *Tetrahedron* **2004**, *60*, 6629–6638.
- (23) Pravst, I.; Stavber, S. Fluorination of 4-alkyl-substituted phenols and aromatic ethers with fluoroxy and N-F reagents: Cesium fluoroxysulfate and N-fluoro-1,4-diazonia-bicyclo[2.2.2]octane dication salts case. *J. Fluorine Chem.* **2013**, *156*, 276–282.
- (24) Jiang, Z.; Ni, T.; Wei, C.; Tian, S.; Li, Y.; Dai, L.; Liu, H.; Zhang, D. Selectfluor-promoted direct fluorination at the 4- or 5-position of imidazole derivatives. *Synlett* **2013**, *24*, 215–218.
- (25) Lowe, B. R.; Maxham, L. A.; Hamey, J. J.; Wilkins, M. R.; Partridge, J. F. Histone H3 mutations: An updated view of their role in chromatin deregulation and cancer. *Cancers* **2019**, *11*, No. 660.
- (26) Bjerke, L.; Mackay, A.; Nandhabalan, M.; Burford, A.; Jury, A.; Popov, S.; et al. Histone H3.3 mutations drive pediatric glioblastoma through upregulation of MYCN. *Cancer Discovery* **2013**, *3*, 512–519.
- (27) Papillon-Cavanagh, S.; Lu, C.; Gayden, T.; Mikael, L. G.; Bechet, D.; Karamboulas, C.; et al. Impaired H3K36 methylation defines a subset of head and neck squamous cell carcinomas. *Nat. Genet.* **2017**, *49*, 180–185.
- (28) Zhang, L.; Floyd, B. M.; Chilamari, M.; Mapes, J.; Swaminathan, J.; Bloom, S.; Marcotte, E. M.; Anslyn, E. V. Photoredox-catalyzed decarboxylative C-terminal differentiation for bulk-and single-molecule proteomics. *ACS Chem. Biol.* **2021**, *16*, 2595–2603.
- (29) Swaminathan, J.; Boulgakov, A. A.; Hernandez, E. T.; Bardo, A. M.; Bachman, J. L.; Marotta, J.; et al. Highly parallel single-molecule identification of proteins in zeptomole-scale mixtures. *Nat. Biotechnol.* **2018**, *36*, 1076–1082.
- (30) Restrepo-Pérez, L.; Joo, C.; Dekker, C. Paving the way to single-molecule protein sequencing. *Nat. Nanotechnol.* **2018**, *13*, 786–796.
- (31) Kapell, S.; Jakobsson, M. E. Large-scale identification of protein histidine methylation in human cells. *NAR: Genomics Bioinf.* **2021**, *3*, No. lqab045.
- (32) Serra-Bardenys, G.; Peiro, S. Enzymatic lysine oxidation as a posttranslational modification. *FEBS J.* **2022**, *289*, 8020–8031.