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Site-specific in vivo protein SUMOylation via translational incorporation of a proximity-reactive pyrrolysine analogue†

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Here, we present a novel strategy that integrates genetic-code expansion and proximity-induced crosslinking to achieve site-specific in vivo SUMOylation. This approach involves incorporating the unnatural amino acid 2-chloroacetyl-NE-lysine (CIAcK) into the target protein using MmFAcKRS1, a previously reported pyrrolysyl-tRNA synthetase mutant that we have repurposed for CIAcK incorporation. Once incorporated, CIAcK can be specifically targeted to react with a cysteine engineered at the C-terminus of SUMO variants leading to a chemically SUMOylated protein. This reaction is proximity-induced, and preferentially promoted when the two reactive groups are in close spatial proximity. We therefore leverage the natural affinity of SUMO for SUMO-interacting motifs (SIMs) on target proteins to generate the targeted SUMO conjugation. Using this approach, site-specific SUMO-conjugates have been produced for two distinct proteins in cells, thus demonstrating its potential as a strategy for helping to dissect the role of SUMOylation in its native cellular context.

SUMOylation is a reversible post-translational modification that involves the covalent attachment of small ubiquitin-like modifiers (SUMOs) to the ε-nitrogen of lysine residues on target proteins. Mirroring the ubiquitination system, it employs a dedicated cascade of enzymes. The heterodimeric SUMO-E1 activating enzyme (SAE1/SAE2) initiates the process by activating SUMO and transferring it to the SUMO-E2 conjugating enzyme (Ubc9), which then attaches SUMO to the target protein with or without a specific SUMO-E3 ligase.¹

SUMOylation is important in diverse cellular processes, regulating protein stability, localization, activity, and interactions with other macromolecules.2 It also plays essential roles in mediating cell proliferation and stress responses,³ and has been implicated in various human diseases, including neurodegenerative disorders,⁴ cancer, 5,6 and cardiovascular diseases. 7,8 In cancer cells, the

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SUMOylation cascade is frequently upregulated,⁵ and contributes to their proliferation by facilitating decatenation and adaptation to stress.^{5,6} Conversely, SUMOylation exhibits protective effects in synucleinopathies, with SUMOylated α-synuclein (αSyn) displaying an inhibitory effect on αSyn aggregation, 9,10 suggesting a promising therapeutic strategy for these disorders.11

Despite its importance, studies of SUMOylation in vivo have been hampered by a lack of control over the specific lysine to be modified. Many substrates possess multiple lysines susceptible to SUMOylation, and thus blocking the dominant site can lead to compensatory modifications at secondary sites. This significantly complicates the production of site-specific SUMOconjugates via enzymatic methods.

To address this challenge, multiple chemical approaches including disulfide exchange, 12 Cu(1)-catalyzed azide-alkyne cycloaddition, 13 thiol-ene reaction, 14 and oxime ligation 15 have been developed. While these methods can generate site-specific SUMOylated proteins, most require in vitro settings or are sensitive to reducing conditions within cells, rendering them unsuitable for studying SUMOylation in vivo. An approach to perform site-specific in vivo SUMOylation would be beneficial, as it would facilitate studies aimed at dissecting the in vivo consequences of SUMOylation at different positions on the same protein. A recent attempt to achieve this used genetic-code expansion and sortase.16 Through the genetic incorporation of an azido-protected glycylglycyl-lysine and sortase-mediated transpeptidation, site-specific SUMOylation was achieved in living cells.

Here we present an alternative method for in vivo sitespecific SUMOylation. Our approach utilizes a proximityinduced crosslinking strategy that avoids the need for sortase co-expression. The key innovation lies in the genetic incorporation of 2-chloroacetyl-Ne-lysine (ClAcK; single letter code 'X'), a non-canonical amino acid equipped with a 2-chloroacetyl group that can readily form thioether bonds with a proximal thiol moiety (Fig. 1). Previous research has successfully employed the S_N2 reaction between cysteine and the 2-chloroacetyl group for

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Fig. 1 Genetic code expansion. L-Pyrrolysine (Pyl; O) and 2-chloroacetyl-Nε-lysine (ClAcK; X).

affinity-guided peptide conjugation.^{17,18} The balanced electrophilicity of the 2-chloroacetyl group allows it to remain inert towards distant cysteines within the protein while efficiently reacting with the proximal cysteine. Furthermore, once incorporated in a peptide, its moderate electrophilicity minimizes unwanted reactions with free thiol-containing compounds, making the reaction suitable for conjugation within living cells. 19 One potential concern, however, is the potential depletion of ClAcK due to the high concentrations (~10 mM) of glutathione in E. coli²⁰ and mammalian cells.²¹ Indeed, in vitro studies revealed that 2 mM ClAcK reacts with 10 mM reduced glutathione with a half-life of \sim 3 hours at pH 7.4 (Fig. S1, ESI†). Despite this, as detailed below, we were able to successfully implement ClAcK for SUMO-conjugation in E. coli. Presumably the reservoir of extracellular ClAcK present in the growth medium is sufficient to replenish the cellular ClAcK levels at a concentration suitable for translational incorporation, making it feasible for use in cell-based studies.

For translational incorporation of ClAcK into proteins using the UAG codon, we employed the pyrrolysine translational machinery with a modified Methanosarcina mazei PylRS mutant, MmFAcKRS1, that was previously identified by Kobayashi et al. for the genetic incorporation of Ne-fluoroacetyllysine (FAcK).²² While FAcK can also react with proximity-confined thiols, its reactivity is limited, with the yield of intramolecular crosslinking in calmodulin estimated at only 14%.22 We hypothesized, however, that ClAcK with a more reactive chloroacetamide group,23 should result in enhanced crosslinking efficiencies while remaining mild enough to mediate the proximity-induced reaction needed for site-specificity.

Since ClAcK and FAcK differ only by one atom, our initial investigation focused on determining whether MmFAcKRS1 could accept ClAcK as a substrate. Using the mCherry reporter assay, where mCherry fluorescence reflects tRNA-synthetase charging efficiency, we confirmed that MmFAcKRS1 can indeed incorporate ClAcK into mCherry at the UAG codon in a specific and dose-dependent manner (Fig. 2a), while having low readthrough for endogenous amino acids (Fig. 2b). To assess the efficiency of ClAcK incorporation by MmFAcKRS1 relative to the benchmark (PylRS for BocK), we compared their readthrough efficiencies for these respective ncAAs. It was found that MmFAcKRS1 exhibited only slightly lower readthrough efficiency for ClAcK than PylRS for BocK (Fig. S2, ESI†).

An initial assessment of the reactivity of the 2-chloroacetylhandle with a thiol group via "proximity-induced reaction"

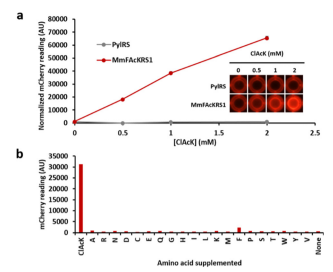


Fig. 2 mCherry reporter assay. (a) Comparison of PylRS and MmFAcKRS1 readthrough of ClAcK. BL21(DE3) cells were transformed with a reporter plasmid containing either PylRS or MmFAcKRS1, along with PylT and an mCherry gene with Lys55 mutated to an amber stop codon. Transformants were grown in a 24-well plate in the presence of varying concentrations of [CIAcK] in M9 minimal medium. Normalized mCherry fluorescence was calculated by dividing the mCherry signal by the OD600. (b) Evaluation of background readthrough by MmFAcKRS1 using the 20 canonical amino acids. BL21(DE3) cells were transformed with the plasmid pDuet-PylT-MmFAcKRS1-KanR-K46X-mCherry-K55UAG and grown in M9 minimal medium supplemented with 2 mM of each of the 20 canonical amino acids

within live cells was performed using E. coli arginine decarboxylase (EcADC) as a model system. EcADC naturally exists as a non-covalent tetramer, comprising two associated dimers, thus the generation of a covalent intermolecular crosslink between subunits should be easily confirmed by SDS-PAGE analysis. Based on the previously determined crystal structure,24 we engineered EcADC-K240X/D535C by introducing ClAcK at position 240, which is proximal to the mutated Cys residue on an adjacent monomer (Fig. S3a, ESI†). We then co-expressed the EcADC-K240X/D535C with PylT/MmFAcKRS1 in the presence of 2 mM ClAcK in E. coli (Fig. S3b, ESI†). SDS-PAGE confirmed the formation of the covalent dimer with a crosslinking efficiency of $\sim 83\%$, thereby demonstrating the robust reaction between ClAcK and Cys in vivo (Fig. S3c and d, ESI†).

We next investigated its application for site-specific in vivo SUMOylation. Studies show that nearly 90% of SUMO binders are also SUMOylation targets, with the SUMO-interacting motif often mediating the outcome of SUMOylation.^{25,26} Given the functional relationship between SUMO-SIM interaction and SUMOylation,^{27,28} we hypothesized that the SUMO-SIM interaction could be used to naturally position the C-terminus of SUMO close to native SUMOylation sites on the target proteins, enabling "proximity-induced" SUMO conjugation.

We first tested this idea with thymine-DNA glycosylase (TDG). To generate the SUMO-TDG conjugate, we engineered the C-terminus of human SUMO1 from its native sequence QTGG to QTC (SUMO(QTC)). This introduced a reactive Cys for a thioether linkage with the site-specifically incorporated ClAcK

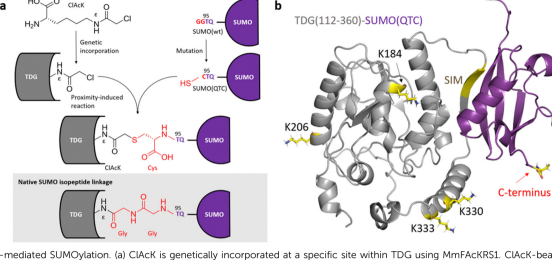


Fig. 3 CIACK-mediated SUMOylation. (a) CIACK is genetically incorporated at a specific site within TDG using MmFAcKRS1. CIACK-bearing TDG then reacts with a co-expressed SUMO variant, SUMO(QTC), harbouring a C-terminal cysteine through a proximity-induced SN2 reaction in cytosol upon expression. Notably, the linkage of the SUMO-conjugate generated by the Cys-ClAcK crosslink possesses identical chain length as the native SUMO isopeptide linkage, though it contains an additional carboxylate group from cysteine. (b) Illustration of SUMO(QTC) non-covalently bound to TDG(112-360). The model was generated using AlphaFold 2.2.0.²⁹

on TDG, while maintaining chain length as the native SUMOvlation linkage via diglycine motif deletion (Fig. 3a). Subsequently, SUMO(QTC) was co-expressed with TDG-K330X-His6, which has ClAcK incorporated at the native SUMOylation site, in E. coli. The formation of the SUMO-TDG conjugate was confirmed by both mass spectrometry and western blotting analysis. LC-MS detected the intact protein with a mass corresponding to the combined mass of SUMO(QTC), TDG(112-360)-His6, and the linker (Fig. S4, ESI†), while LC-MS/MS further identified the SUMO C-terminal tryptic peptide with a modification corresponding to the mass of the 7-residue TDG tryptic peptide containing TDG-K330X (Fig. S5, ESI†). Western blotting analysis also detected the conjugate (Fig. 4a), and suggested a 15.7% SUMO-conjugation yield. This compares favourably to

the 5-10% SUMO-conjugation yield reported for sortylation in mammalian cells.¹⁶

Previous studies have shown that truncating the intrinsically-disordered N-terminus of SUMO (ΔN-SUMO) can improve its SIM-mediated binding affinity to its SUMOylation targets. 11,30 To explore whether this tighter SUMO-SIM interaction could be used to enhance the conjugation efficiency, we designed a truncated SUMO1 variant, ΔN-SUMO(QTC), lacking the N-terminal 14 residues of SUMO1 and containing the C-terminal QTGG to QTC mutation and co-expressed it with TDG-K330X-His₆ in E. coli. As expected, the conjugation yield increased from 15.7% to 34% (Fig. 4a), supporting the role of the SUMO-SIM interaction in directing the SUMO C-terminal thiol for "proximity-induced" crosslinking with

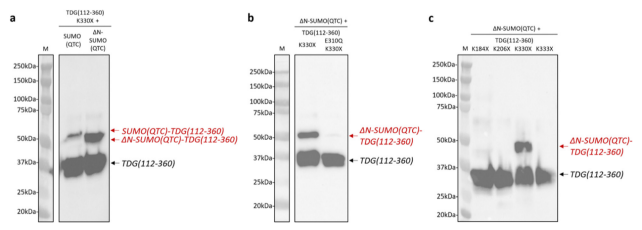


Fig. 4 Western blotting analysis of ClAcK-mediated TDG SUMOylation with anti-His antibody. (a) Comparison of the conjugation yield between $SUMO(QTC) \ and \ \Delta N-SUMO(QTC) \ to \ TDG(112-360)-His_6 \ with \ Clack \ installed \ at \ K330. \ (b) \ Comparison \ of \ the \ conjugation \ yield \ between \ TDG(112-360)-His_6 \ with \ Clack \ installed \ at \ K330. \ (b) \ Comparison \ of \ the \ conjugation \ yield \ between \ TDG(112-360)-His_6 \ with \ Clack \ installed \ at \ K330. \ (b) \ Comparison \ of \ the \ conjugation \ yield \ between \ TDG(112-360)-His_6 \ with \ Clack \ installed \ at \ K330. \ (b) \ Comparison \ of \ the \ conjugation \ yield \ between \ TDG(112-360)-His_6 \ with \ Clack \ installed \ at \ K330. \ (b) \ Comparison \ of \ the \ conjugation \ yield \ between \ TDG(112-360)-His_6 \ with \ Clack \ installed \ at \ K330. \ (b) \ Comparison \ of \ the \ conjugation \ yield \ between \ TDG(112-360)-His_6 \ with \ Clack \ installed \ yield \ yie$ K330X-His₆ and the SIM-defective TDG(112-360)-E310Q-K330X-His₆ with Δ N-SUMO(QTC). (c) Conjugation of Δ N-SUMO(QTC) to TDG(112-360)-His₆ with CIACK installed at either the native SUMOylation site (K330) or the different alternate sites. The full gel images can be found in the "full gels and blots" section of the ESI.†

ClAcK. To further validate this role, we introduced a known SIM-disrupting mutation (E310Q) into the TDG-K330X-His₆ mutant.31 Co-expression of the SIM-defective mutant with ΔN-SUMO(QTC) resulted in markedly reduced amount of conjugated product (~1.4%) on western blot (Fig. 4b and Fig. S6, ESI†). This confirms that the SUMO-SIM interaction is directly responsible for guiding the C-terminus of SUMO to K330.

Since the SUMO-conjugation efficiency in our method is heavily influenced by the SUMO-SIM interaction, we wondered if our approach could be extended for use in mapping the SUMOylation landscape. To explore this, we designed TDG mutants with ClAcK incorporated at alternate sites (Fig. 3b), including proximal (K333), intermediate (K184), and distal (K206) positions relative to the native SUMOylation site (K330). Notably, despite successful ClAcK incorporation, none of these mutants formed crosslinks with ΔN-SUMO(QTC) (Fig. 4c and Fig. S7-S9, ESI†). This suggests that our method is highly selective for the native SUMOvlation site.

To better understand the differences between the SUMO-TDG conjugate generated by our method and the enzymatically SUMOylated TDG, we treated these conjugates with Ulp1 SUMO protease (Fig. S10, ESI†). While the enzymatically SUMOylated TDG can be completely deSUMOylated by Ulp1 treatment, the ClAcK-mediated conjugate remained intact, thus confirming the resistance of the thioether linkage to SUMO protease cleavage. Furthermore, we investigated the impact of ClAcKmediated SUMOylation on the substrate binding activity of TDG. Similar to enzymatic SUMOylation, 32,33 ClAcK-mediated SUMO conjugation at K330 of TDG resulted in a weakening of substrate binding (Fig. S11, ESI†).

To demonstrate the versatility of our in vivo ClAcK-mediated SUMOylation strategy, we extended its application to other SUMO substrates. Unlike TDG, which is ordered and has a single major SUMOylation site, α-synuclein (αSyn) is intrinsically disordered and possesses multiple native SUMOylation sites.9 K96 and K102 are the dominant sites, accounting for over 50% of SUMO-αSyn conjugates, while at least 9 of its remaining 13 lysines are reported as minor SUMOylation sites.9 It was therefore intriguing to test if our method could successfully conjugate SUMO to the various acceptor sites on αSyn.

Since αSyn is predominantly SUMOylated at K96 and K102, we first introduced ClAcK at these two positions. We co-expressed SUMO(QTC) with either aSyn-K96X or aSyn-K102X in E. coli and confirmed successful conjugation at both sites (Fig. 5 and Fig. S12, ESI†). To assess whether SUMO-SIM interaction likewise drives the conjugation in αSyn, we tested the conjugation efficiency of ΔN-SUMO(QTC), which has higher binding affinity to SIM, by co-expressing it with αSyn-K96X. Consistent with the results for TDG, ΔN-SUMO(QTC) produced a higher αSyn-K96X conjugation yield (~48.5%) compared to SUMO(QTC) (\sim 22.5%) (Fig. S13, ESI†).

Encouraged by these results, we expanded our investigation to other sites on αSyn. We selected four alternate sites for ClAcK incorporation (Fig. S14, ESI†): K10, a minor SUMOylation site on the N-terminus of αSyn; V52, a residue on SIM2 of αSyn; 11 and A91, and G93, proximal sites with varying distances from

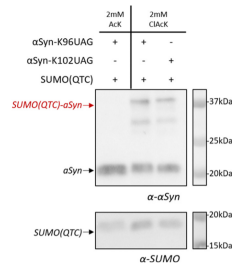


Fig. 5 ClAcK-mediated SUMOylation of αSyn at its two dominant native SUMOylation sites (K96 or K102). Boiled cell lysates of BL21(DE3) co-expressing SUMO(QTC) and different aSyn variants were analyzed by western blotting using either anti- αSyn or anti-SUMO antibodies. AcK (Nε-acetyl-lysine). Conjugation yield determined by ImageJ: αSyn-K96X, 27.6%; αSyn-K102X, 25%. The full gel images can be found in the "full gels and blots" section of the ESI.†

the native SUMOylation site (K96). Each of these mutants was co-expressed with Δ N-SUMO(QTC) in *E. coli* and analysed by western blot using anti-αSyn.

As expected, ΔN-SUMO(QTC) was successfully conjugated to αSyn-K10X. However, contrary to our expectation, all other mutants modified at non-SUMOylation sites also exhibited robust crosslinking (Fig. S15-S17, ESI†). This unexpected promiscuity contradicts our observation with TDG, where SUMO(QTC) reacted solely with ClAcK at native SUMOylation sites. To reconcile these findings, we propose two possibilities: (1) the specificity of SUMO(QTC) for native SUMOylation sites is context dependent. For structurally disordered targets like αSyn, increased accessibility to alternate sites due to its conformational flexibility could lead to promiscuous conjugation; or (2) the observed non-native conjugation sites could represent cryptic SUMOylation targets. Mutation of these sites to lysine might reveal their latent potential for enzymatic SUMOylation.

To probe deeper into the concept of cryptic SUMOylation targets, we focused on site V52, owing to its further distance from the dominant native SUMOylation sites (K96 and K102). We engineered αSyn-V52K-K96R-K102R by replacing V52 with lysine and mutating the two dominant SUMOylation sites to arginine. Purified αSyn-V52K-K96R-K102R was then incubated with a complete enzymatic SUMOylation system (Fig. S18, ESI†). LC-MS/MS confirmed V52K as a genuine SUMOylation target (Fig. S19, ESI†). This suggests that the crosslink between ClAcK and SUMO(QTC) can accurately predict the positions of cryptic enzymatic SUMOylation sites.

To the best of our knowledge, the mutation of V52 to lysine in aSyn is the first reported instance of an engineered, enzymatic SUMOylation site. Interestingly, upon analysis with JASSA,³⁴ the mutated sequence was found to exhibit low inherent potential for SUMOylation (Table S1, ESI†). Considering that both our crosslinking experiment (Fig. S15, ESI†) and molecular dynamic simulation (see ESI†) revealed that SUMO-SIM interaction favours positioning of the SUMO C-terminus near residue 52 of aSyn, we propose that its unexpected SUMOylation is primarily driven by SUMO-SIM interaction, similar to SIM-dependent SUMOylation.²⁸

Conclusions

In summary, we describe a straightforward approach for generating site-specific SUMO-conjugates in cells by leveraging native SUMO-SIM interactions to precisely position SUMO for efficient crosslinking at the native SUMOylation sites. Using this approach, we demonstrate the successful conjugation of SUMO to two distinct proteins, thymine-DNA glycosylase and α-synuclein at their native SUMOvlation sites. Furthermore, we have illustrated the ability of this method for mapping the SUMOylation landscape and for unveiling previously unknown cryptic SUMOylation sites.

While these current studies are demonstrated in E. coli, our method for site-specific in vivo SUMOylation is likely adaptable to mammalian cells. Efficient reaction between the chloroacetyl handle and targeted cysteine has been demonstrated in the cytosol of human breast cancer cells. 19 Additionally, the PylT/ MmPylRS system for ClAcK incorporation has also been shown to be orthogonal in mammalian cells.35 We envision that translating our system to mammalian cells could enable in vivo SUMOylation studies with unprecedented spatial resolution.

Author contributions

Conceptualization, M. K. C.; methodology, Y. H. C.; investigation, Y. H. C.; writing - original draft, Y. H. C.; writing - review & editing, M. M. L. and M. K. C.; funding acquisition, M. K. C.; supervision, M. M. L. and M. K. C.

Conflicts of interest

There are no conflicts to declare.

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