Triple, Mutually Orthogonal Bioorthogonal Pairs through the Design of Electronically Activated Sulfamate-Containing Cycloalkynes

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ABSTRACT: Interest in mutually exclusive pairs of bioorthogonal labeling reagents continues to drive the design of new compounds that are capable of fast and predictable reactions. The ability to easily modify S-, N-, and O-containing cyclooctynes (SNO-OCTs) enables electronic tuning of various SNO-OCTs to influence their cycloaddition rates with Type I–III dipoles. As opposed to optimizations based on just one specific dipole class, the electrophilicity of the alkynes in SNO-OCTs can be manipulated to achieve divergent reactivities and furnish mutually orthogonal dual ligation systems. Significant reaction rate enhancements of a difluorinated SNO-OCT derivative, as compared to the parent scaffold, were noted, with the second-order rate constant in cycloadditions with diazacacetamides exceeding 5.13 M\(^{-1}\) s\(^{-1}\). Computational and experimental studies were employed to inform the design of triple ligation systems that encompass three orthogonal reactivities. Finally, polar SNO-OCTs are rapidly internalized by mammalian cells and remain functional in the cytosol for live-cell labeling, highlighting their potential for diverse in vitro and in vivo applications.

INTRODUCTION

Bioorthogonal chemistry represents a powerful strategy to monitor individual molecules in biological systems; indeed, within the past two decades, a multitude of reactions have been developed to fulfill this function.\(^1\) Nonetheless, a larger challenge involves the identification of mutually orthogonal bioorthogonal reagent pairs for simultaneous observation of multiple biomolecules in a cell.\(^2\) The accessibility of convenient, inexpensive, and electronically tunable "orthogonal bioorthogonal" reagent pairs remains a topic of intense interest, as new labeling tools continue to drive the design of new compounds that are capable of fast and predictable reactions. The ability to easily modify S-, N-, and O-containing cyclooctynes (SNO-OCTs) enables electronic tuning of various SNO-OCTs to influence their cycloaddition rates with Type I–III dipoles. As opposed to optimizations based on just one specific dipole class, the electrophilicity of the alkynes in SNO-OCTs can be manipulated to achieve divergent reactivities and furnish mutually orthogonal dual ligation systems. Significant reaction rate enhancements of a difluorinated SNO-OCT derivative, as compared to the parent scaffold, were noted, with the second-order rate constant in cycloadditions with diazacacetamides exceeding 5.13 M\(^{-1}\) s\(^{-1}\). Computational and experimental studies were employed to inform the design of triple ligation systems that encompass three orthogonal reactivities. Finally, polar SNO-OCTs are rapidly internalized by mammalian cells and remain functional in the cytosol for live-cell labeling, highlighting their potential for diverse in vitro and in vivo applications.

The 1,3-dipolar cycloadditions are popular reactions in bioorthogonal chemistry, particularly strain-promoted azide–alkyne cycloadditions (SPAAC; Figure 1a).\(^{1c,3b,4}\) The slow kinetics of unfunctionalized cyclooctynes has been addressed by electronically activating the alkyne via the installation of fluorine atoms at the propargyl carbon.\(^3\) For example, Bertozzi and co-workers showed that moving from monofluorinated to difluorinated alkyne (DIFO) resulted in a nearly 2 orders of magnitude increase in the rate of reaction with BnN\(_3\) from 0.0043 to 0.076 M\(^{-1}\) s\(^{-1}\).\(^{5a}\) The popular bioorthogonal reagent dibenzocyclooctyne (DBCO) utilizes benzene rings fused to the cycloalkyne to accelerate reaction with BnN\(_3\); however, further electronic tuning of this scaffold is challenging, due to limited derivatization sites.\(^6\) Other reported modifications to the parent DBCO yield reaction rates with BnN\(_3\) spanning less than an order of magnitude (0.31–0.90 M\(^{-1}\) s\(^{-1}\)).\(^7\) In addition to the lack of modularity, the strain-activated nature of this alkyne leads to issues with stability and chemoselectivity, limiting opportunities to develop mutually orthogonal reactions involving DBCO derivatives.\(^8\)

Recently, we developed a unique class of strained cycloalkynes that contain S-, N-, and O-heteroatoms embedded in the ring (SNO-OCTs).\(^9\) Our previous kinetic studies of the sulfamate-containing SNO-OCTs I–2 (Figure 1a) show second-order kinetic rate constants (k\(_s\)) in reactions with BnN\(_3\) that range from 0.0014 to 0.087 M\(^{-1}\) s\(^{-1}\), with polar protic solvents further enhancing the rate of 2 in the azide–alkyne cycloaddition to 0.13 M\(^{-1}\) s\(^{-1}\).\(^1k,9\) Despite the relaxed alkyne angles and reduced ring strain in 1–2, these SNO-OCTs exhibit a significant rate increase for cycloadditions, comparable to those seen with strain-activated 4-dibenzo-cyclooctyne (DIBONE) and DBCO. Density functional theory (DFT) calculations indicated our SNO-OCTs benefit from unique electronic activation of the alkyne via tailored anomer effects and hybridization, resulting in robust stability at high pH and in the presence of glutathione.\(^2b,9\) Furthermore, the presence of the sulfonyl group increases the polarity in the transition state.
which has also been shown to enhance the rates of cycloaddition. 8

The tunability of SNO-OCTs begs the question of whether these scaffolds can be tailored for divergent reactivity with diverse coupling partners. The Prescher group developed several mutually orthogonal dual ligation systems, most notably, a series of cyclopropenes that could be sterically tuned to achieve mutually orthogonal Type I and Type III cycloadditions (Figure 1b). 2c,d,3c Sterically encumbered 3,3-disubstituted-cyclopropenes prefer to react with nitrile imines, while shifting the bulky tetrazine. Azide cycloadditions were also studied computationally to determine if a triple SNO-OCT ligation system might achieve mutually exclusive orthogonal reactivities of SNO-OCTs for biological applications. As Type I, Type II, and Type III dipoles. Frontier molecular orbital (FMO) and natural bond orbital (NBO) 15 analyses were used to rationalize how derivatization influences alkyne electronics, while distortion/interaction analysis shed insight into how electronics and strain impact reactivity trends. 13 Computational insights guided the identification of potentially orthogonal SNO-OCT/dipole pairs; for example, the electron-deficient alkyne of difluorinated SNO-OCT 3 (df-SNO-OCT) was expected to display divergent reactivity relative to the parent 1 (Figure 1c). Successes in developing a new triple ligation system utilizing two SNO-OCT derivatives highlight the versatility of these scaffolds for chemoselective reactions with various dipoles by modifying the alkyne electronics. Finally, the use of a novel triple ligation system for protein labeling is demonstrated, and the application of SNO-OCTs as efficient cell-surface visualization agents is described.

■ RESULTS AND DISCUSSION

Computational Methods to Predict SNO-OCT Cycloaddition Reaction Rates. To facilitate rational design of promising SNO-OCT derivatives (see Figure S1−1), we sought to develop a computational approach capable of accurately predicting reaction rates and reactivity trends. As numerous SNO-OCT derivatizations can be envisaged, DFT calculations were needed to rapidly identify potential mutually orthogonal reactions and streamline experimental efforts. DFT calculations to study reactions of 1 employed a simplified model compound 4, where the n-pentyl chain was truncated to a methyl group for computational ease. Initial computations conducted at the M06-2X level of theory 13 with a 6-311+G(4d,p) basis set using Gaussian 16 14 showed predicted rates that deviated significantly from experimental values (Figure S1−4 and S1−5). Zuilhof also reported that M06-2X overestimates activation barriers in strain-promoted oxidation-controlled quinone (SPOCQ) cycloadditions and found that switching to the B97D functional gave calculated rates that aligned better with experimental data. 15 SNO-OCT rates were calculated at the B97D level of theory using a Conductor-like Polarizable Continuum Model for the solvent (see the SI for details). 16 B97D improved the agreement between computed and experimental k2 values (within ~1 order of magnitude); accordingly, further computational analyses were conducted utilizing the B97D level of theory to predict potential mutual orthogonalities of SNO-OCT derivatives.

SNO-OCT Derivatization to Increase Reaction Rates. Our ability to readily change the substitution pattern of the SNO-OCT scaffold stimulated efforts to electronically tune the alkyne to (1) improve rates of dipolar cycloadditions, (2) achieve mutually exclusive orthogonal reactivities of SNO-OCTs with Type I−III dipoles in [3 + 2] and [4 + 2] cycloadditions, and (3) highlight the ability to install multiple functional handles for appending reporter molecules and dyes to SNO-OCTs for biological applications. As Type I−III cycloadditions can be designed based on FMO considerations, the potential for mutually orthogonal reactivity may be realized if effective orbital matching is achieved. We hypothesized an electron-poor, electrophilic SNO-OCT derivative might engage in chemoselective reaction with a nucleophilic diazoacetamide dipole; furthermore, the reaction should be orthogonal to an electron-rich, electrophilic SNO-OCT derivative might engage in inverse electron-demand Diels−Alder (IEDDA) reaction between a more electron-rich SNO-OCT and a dipyrrolyl tetrazine. Azide cycloadditions were also studied computationally to determine if a triple SNO-OCT ligation system might transcend all three dipole classes.

NBO analyses 15 of geometry-optimized SNO-OCTs 2−4 in water were used to study the effects of derivatization on alkyne electronics. Multiple orbital interactions exist within the SNO-
OCT scaffold; these may differ greatly in their contributions to the alkyne electronics, depending on the nature of the substitution in the alkyl side chain or the propargylic substitution (Figures 2a and S1−2). The predicted SNO-OCT alkyne electrophilicities were determined based on the second-order perturbation energies of the relevant orbital interactions (Figure 2b and Table S1−1). As might be expected, SNO-OCT 4 and its hydroxylated counterpart 2, which lack fluorne at the propargylic site, contain electron-rich alkenes. Introduction of a single propargyl fluorne (3a) yields a less electron-rich alkyne, while a second fluorneation to furnish dF-SNO-OCT 3 contains the most electrophilic alkyne of the series. It is important to note the calculations suggest modification of the R group in 2 and 4 minimally impacts the π system of the alkyne, whereas modifications at the propargylic site adjacent to the reactive center lead to dramatic changes in electrophilicity.

Identifying Potential Orthogonal Reactivity via Computed Reaction Rates. While qualitative predictions of SNO-OCT/Type I−III dipoles that are likely to yield mutually orthogonal systems can be gleaned from Figure 2, reaction modeling could help select optimal cycloaddition partners. Transition state (TS) modeling of reactions between SNO-OCTs 2−4 and an example from each dipole class (Type I, methyl diazocacetamide D1; Type II, methyl azide D4; Type III, dipropyridyl tetratetrazine D3) were carried out to assess potential orthogonality (Figure 3). Molecular dynamics simulations were conducted on the initial transition state models to identify the lowest energy conformers (see the SI for details). Cycloadditions of nucleophile Type I dipoles should be accelerated with electron-deficient alkenes, due to a lowering of the LUMO and a reduced FMO energy gap. Indeed, difluorination in dF-SNO-OCT 3 was predicted to give a 267-fold increase in reaction with D1, as compared to 2 and 4 (Figure 3a,e). Distortion/interaction or activation strain analysis on calculated transition state models shed further insight into the key features of the SNO-OCT driving reactivity. Computed reactivities of 2−4 in reactions with methyl diazoacetamide show excellent correlation to the interaction energy required to reach the transition state geometry (Figure 3d, Eint Figure S1−15), and the calculated alkyne electrophilicity, with dF-SNO-OCT 3 showing the largest stabilizing interaction energy between the coupling partners (+3.1 kcal mol−1 relative to 4). This distinction can be rationalized by FMO considerations, as the most electron-deficient alkenes benefit from lower-lying LUMO levels; the presence of the propargylic fluorines stabilizes the LUMO (3, −2.22 eV compared to 2, −1.45 eV; 4, −1.39 eV; Figure S1−3).

Type III cycloadditions, as exemplified by IEDDA, should display faster reaction kinetics with electron-rich alkenes, due to destabilization of the HOMO. Previous computational studies on these systems investigated the transition states for either the cycloaddition alone or both the cycloaddition and N2 evolution. The cycloaddition TS is expected to be the kinetically relevant barrier, and the values reported herein correspond to the cycloaddition TS. Dipropyridyl tetratetrazine D3 (Figure 3b) was modeled, as the electron-withdrawing pyridyl groups enhance cycloaddition rates due to LUMO+1 stabilization of the dipole. As predicted, electron-rich SNO-OCTs 2 and 4 (HOMO levels −6.42 and −6.44 eV, respectively) exhibit the fastest kinetics, whereas dF-SNO-OCT 3 shows significantly lower reactivity due to HOMO stabilization (−7.03 eV). Hydroxylated SNO-OCT 2 was predicted to display a 2.7-fold rate enhancement as compared to 4. Reactivity trends showed moderate-to-excellent correlation to both distortion and interaction energies (Figures 3d and S1−16 and S1−17).

Based on reactions of SNO-OCTs with Type I and III dipoles, the potential for mutually orthogonal double ligation appeared promising. The next challenge was to identify a third orthogonal reaction, potentially between a SNO-OCT and a Type II dipole. Thus, reactions of 2−4 with azide D4 were modeled for comparison (Figure 3c). In Type II cycloadditions, reaction can occur via HOMO dipole−LUMO dipole or HOMO dipole−LUMO dipole interactions, depending on the corresponding FMO energy gaps. This dichotomy makes it challenging to correlate reactivity with alkyne electrophilicity; thus, reactivity trends in SNO-OCT reactions with azides were instead proposed to track with the relevant energy gaps. The dF-SNO-OCT 3 was expected to react 17-fold more quickly with D4 relative to 4, with a k2 value predicted to be over an order of magnitude higher. Indeed, this reactivity trend can be predicted by assessing FMO gaps for each SNO-OCT/azide pair. Higher electron density in 2 and 4 leads to HOMO destabilization, and as such, these SNO-OCTs react via a reduced HO−MO dipole gap. Conversely, the presence of the fluoro groups in dF-SNO-OCT 3 increases this gap through HOMO stabilization, thus the 3/Men reaction proceeds through a more favorable LUMO dipole−HOMO dipole interaction. Though dF-SNO-OCT 3 exhibits reduced dipole distortion energy (−0.6 kcal mol−1 relative to 4), consistent with bond relaxation (+0.8°), the corresponding alkyne distortion energy is elevated in comparison to parent SNO-OCT 4 (+0.52 kcal mol−1). Thus, the predicted acceleration for cycloadditions of azides with dF-SNO-OCT 3 is attributable to this pair having the highest stabilizing interaction energy (11.8 kcal mol−1). Indeed, the activation barriers show improved correlation with the interaction energies relative to the distortion energies (Figure S1−16).

Experiment Validates Computational Trends. The extent to which SNO-OCT substitution impacts the value of k2 in reactions with Type I−III dipoles was determined by experimental investigation of the reactivities of 2−4 and comparison to predicted values. SNO-OCTs 1 and 2 were prepared as previously reported, while 5 required a modified synthetic route, affording the product in 34% yield from propionaldehyde (see Scheme S2−1 for details). Computational results were corroborated in selected instances through experimental kinetic data using 1H NMR spectroscopy and
Figure 3. B97D/6-311+G(d,p) calculated transition state models for reactions of SNO-OCTs with (a) diazoacetamide, (b) tetrazine, and (c) azide. (d) Distortion/interaction analysis indicating the influence of distortion energy ($\Delta E_{\text{dis}}$) and interaction energy ($\Delta E_{\text{int}}$) on activation barriers for cycloadditions of Type I–III dipoles with various SNO-OCT derivatives. (e) Calculated and experimental second-order rate constants. Truncated SNO-OCTs 3 and 4 and dipoles D1 and D4 were used for computations; analogous SNO-OCTs 1 and 5 and dipoles D2 and D5 were used for experiments. Energies ($E_{\text{dis,SNO-OCT}}, E_{\text{dis,dipole}}, E_{\text{int}}, E_{\text{act}}$ and $\Delta G^\ddagger$) are reported in kcal/mol and second-order rate constants ($k_{\text{calc}}$ and $k_{\text{exp}}$) are in M$^{-1}$ s$^{-1}$. 

\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|}
\hline
SNO-OCT & $E_{\text{dis,SNO-OCT}}$ & $E_{\text{dis,dipole}}$ & $E_{\text{int}}$ & $E_{\text{act}}$ & $E_{\text{dis,SNO-OCT}}$ & $E_{\text{dis,dipole}}$ & $E_{\text{int}}$ & $E_{\text{act}}$ & $E_{\text{dis,SNO-OCT}}$ & $E_{\text{dis,dipole}}$ & $E_{\text{int}}$ & $E_{\text{act}}$ \\
\hline
4 & 3.46 & 16.2 & 12.7 & 6.96 & 6.73 & 17.8 & 24.2 & 0.41 & 2.53 & 15.2 & 9.48 & 8.21 \\
2 & 2.50 & 15.9 & 13.4 & 5.02 & 5.76 & 17.1 & 24.0 & -1.17 & 0.83 & 15.0 & 9.71 & 6.15 \\
3 & 3.55 & 14.4 & 15.8 & 2.21 & 7.68 & 20.2 & 23.2 & 4.71 & 3.05 & 14.6 & 11.8 & 5.86 \\
\hline
\end{tabular}
plate-reader techniques (Figure 3e). It is important to note that the plate reader was utilized for dipyridyl tetrazine reactions, due to solubility issues encountered in NMR kinetic experiments.

Calculated and experimental results of reactions between SNO-OCTs and alkyl diazoacetamides were particularly promising with respect to chemoselectivity and accelerated reaction rates (Figure 3e). A gen-difluorination led to a predicted rate enhancement of 267-fold moving from 4 to 3 in reaction with diazoacetamides (D1). Experimental determination of $k_2$ for the cycloaddition of 5 and D2 proved difficult; the reaction at 23 °C was too fast to accurately determine $k_2$ by NMR. VT-NMR showed formation of 5a with 80% conversion within 2 min at −32 °C (Figure S2−17). Despite the lack of accurate rate data, a series of competition reactions were performed using an excess amount of SNO-OCTs to provide an estimated $k_2 > 5.13 \text{ M}^{-1} \text{s}^{-1}$ (see Figure S2−16 and S2−17 for details), which falls within 1 order of magnitude from the computational prediction, thus reaffirming our approach to kinetic rate calculations.

Predicted reactions of 2−4 with dipyridyl tetrazine D3 indicate ~5350-fold rate enhancement in moving from electron-poor dF-SNO-OCT 3 to electron-rich parent 4; indeed, experimental observations corroborated this prediction, and no reaction was noted between 5 and D3. Though calculated rates for IEDDA with D3 showed a 2−3 orders of magnitude deviation from experimental rates, the computational method was successful in identifying two promising, mutually orthogonal reactions of SNO-OCTs. Further computational investigation into the origin of this deviation is underway to predict more accurate rates in IEDDA with D3 for future studies.

Calculated rates between azides and SNO-OCTs fall within 1 order of magnitude from experimental values, when conducting measurements in CD$_3$CN (Figure 3e). Furthermore, calculations indicated a 17× rate-enhancement moving from parent 4 to dF-SNO-OCT 3, compared to the experimentally observed 18× rate-enhancement in moving from 1 to 5, accurately predicting relevant experimental reactivity trends. Modified SNO-OCTs 4 and 1 also show predictable trends for azide/SNO-OCT cycloadditions, with a rate acceleration when more water was used (2:1 D$_2$O:CD$_3$CN, see Figure S2−8 for details), suggesting a potential hydrophobic effect contributing to rate acceleration.5,23

The inability of DFT methods to accurately model solvents is well-known. Zuilhof and co-workers also point out that their solvent model does not account for rate increases dependent on solvent identity.15 In our case, the B97D functional is sufficient to predict how electronic fine-tuning of the SNO-OCT influences cycloaddition rates, particularly in [3 + 2] cycloadditions.

**Development of Dual and Triple Ligation Systems with Custom SNO-OCTs.** With initial computational and experimental data in hand, we wanted to develop mutually exclusive bioorthogonal dual or triple ligations of SNO-OCTs with various dipoles, as dual ligation has been shown to be effective in probing complex native biological environments and details of subcellular mechanisms.24 Success would effectively expand the tool box available for multiple ligation of biologically relevant functionalities, and enable fine-tuning of the SNO-OCT properties for specific applications.

As predicted by DFT, SNO-OCT 5 gave excellent yields in reactions with diazoalkane D2 (94%) and azide D5 (96%); no reaction was noted with D3 (Scheme 1a). In validation of our initial hypothesis that a dual ligation system comprised of orthogonal SNO-OCTs and Type I/Type III dipoles could be achieved, a competition experiment using SNO-OCTs 1 and 5 along with dipoles D2 and D3 (Schemes 1b, S2−2a, and S2−2b and Figure S2−19) showed complete mutual orthogonality.25−28 Indeed, the excellent chemoselectivity supported the large rate enhancement predicted in moving from parent 4 to dF-SNO-OCT 3. In contrast, a dual ligation system with azide D5 instead of D3 gave significantly reduced chemoselectivity (Scheme S2−2c); this also matches computed reaction rates (Figure 3e).

The development of a triple ligation system using only custom SNO-OCTs proved challenging, as predicted by the computational rate data. Ideally, the three mutually exclusive pairs of reagents should display $k_2$ values differing by at least 2 orders of magnitude. Based on calculated relative rates of dF-SNO-OCT 3/D1 and parent 4/D1 (Figure 3e), this condition cannot be
explicitly met with our current SNO-OCT derivatives. Thus, a commercially available non-SPAAC boronic acid/hydrazine pair was introduced. Bane and co-workers have shown that boronic acid and hydrazine/hydrazide pairs are competent in fast bioorthogonal labeling of biologically relevant molecules; furthermore, the boronic acid system shows orthogonality to both IEDDA and SPAAC. This orthogonality was verified by NMR (Figure S2−13 and S2−21), as well as LC-MS analysis, which is more reliable than NMR methods when significant peak overlap occurs in complex mixtures. To our delight, the orthogonality in Scheme 1b translated to the triple ligation (Scheme 1c and Figure S2−22), with mutual orthogonality observed among all three reaction pairs by LC-MS.

**Protein Triple Ligation and Mammalian Cell-Labeling Assays.** Simultaneous multitarget in vitro and in vivo labeling is of considerable interest for advancing modern biotherapeutics, biological assay development, and basic science investigations. We sought to demonstrate the mutual orthogonality of our SNO-OCT bioorthogonal pairs in biological settings, as well as their capacity to enable triply orthogonal ligations as suggested by LC-MS assays (Scheme 1c). To that end, SNO−DF−Rho and SNO-C343 were synthesized from 5 and 1, employing either carboxylic acid or primary amine handles, as they are easily derivatizable with other biologically relevant functionalities (Scheme S2−3). Additionally, a previously reported cysteine-hydrazide Texas Red derivative was also adapted (Figure 4a, see the SI for more details). Based on the LC-MS experiment in Scheme 1c, these reagents should form the basis of a tripartite mutually orthogonal set, appropriate for demonstrating translatability to biologically relevant conditions. Indeed, one-pot labeling of three different proteins functionalized to present a diazoacetamide (Cyto c-AcDz), tetrazine (RNase A-Tez), and boronic acid-aldehyde (RNase A-BA) showed excellent chemoselectivity in agreement with small-molecule triple ligation results (Figure 4a). As anticipated, in situ gel fluorescence images revealed some bleed over between dyes, but minimal cross reactivity across orthogonal pairs,
demonstrating the chemoselectivity of the reagents (Figure S2−23).

Next, we sought to explore potential applications in cellulo. The cellular uptake properties of SNO−C343, SNO−DF−Rho, and three DBCO-derivatives were compared to assess internalization and localization of these molecules within Chinese hamster ovary (CHO)-K1 cells. We observed the rapid internalization of SNO−DF−Rho and SNO−C343 (Figures 4b, S2−24, and S2−25). Colocalization analysis of SNO−DF−Rho with MitoTracker Green revealed robust mitochondrial trafficking as shown by a Pearson’s correlation coefficient of R = 0.90 ± 0.01 (Figure S2−26). Comparison to an in-house synthesized DBCO−C343 derivative as well as two commercial conjugates, DBCO−PEG4−Rho110 and DBCO−Cy5, revealed enhanced internalization of SNO−OCT−dye conjugates relative to the commercial reagents upon incubation with CHO-K1 cells (Figure 4b). Internalization of DBCO−C343 occurred at comparable concentrations and gave staining patterns similar to those observed for SNO−C343 (Figures 4B and S2−28). Internalization experiments with DBCO−PEG4−Rho110 and DBCO−Cy5 conjugates illustrated indirect internalization of DBCO−PEG−Rho110, and ablation of internalization in the case of DBCO−Cy5, even at concentrations 5-fold greater than the maximum explored with SNO−OCT reagents (Figures 4B and S2−27). In addition to the orthogonality of SNO−DF−Rho and SNO−C343 reactivity with Type-I and Type-II dipoles, these unconjugated reagents are taken-up robustly by mammalian cells and thus have potential for in cellulo applications.

The mammalian glycocalyx forms a protective anionic coating anchored to the lipid bilayer.28 The structure and chemical properties of this bilayer play important roles in both normal and pathological biological processes, including cellular recognition, organismal development, cancer, and infectivity.29 In pioneering investigations, Reuter, Bertozzi, and their co-workers labeled the glycocalyx of live cells metabolically with synthetic carbohydrates.30 We sought to enlarge the tool-kit for glyocalyx analyses with the new SNO−OCT reagents. Metabolic labeling with Ac4ManDia23 was used to assess the reactivity of SNO−DF−Rho in a biological context. The concentration-dependent uptake and distribution of Ac4ManDia in CHO-K1 cells was assessed to identify the optimal concentrations for resolving the glyocalyx upon reaction with SNO−DF−Rho at 1 μM (Figure S2−29). Side-by-side comparison of staining patterns in CHO-K1 cells cultured in the absence or presence of Ac4ManDia revealed a difference in the distribution of the label. Specifically, SNO−DF−Rho when cells were metabolically labeled with Ac4ManDia (Figure 4c,d). In analogous flow-cytometry experiments, the robust labeling and internalization of cells metabolically labeled with Ac4ManDiaz was again observed, and without significant cell death (Figure S2−30). Together, these data demonstrate the reactivity of SNO−DF−Rho in cellulo and highlight the potential of SNO−OCT reagents for biological assays in living cells.

# CONCLUSIONS

The integration of computational and experimental studies led to custom heterocyclic cycloalkynes (SNO−OCTs) that display orthogonal reactivities with Type I and III dipoles. The cooperative effects of the propargylic and homopropargylic heteroatoms, which are separated by a sulfamate bridge, enable tuning of the alkyne electronics and alkyne distortion via remote hybridization and stereoelectronic effects. The addition of a gem-

difuoro group to the propargylic carbon of the SNO−OCT has several beneficial features, including a significant increase in the value of k2 for reactions with Type I−II dipoles and a greatly decreased preference for participation in IEDDA-type reactions. Inspired by computational investigations, experimental validation of a new mutually orthogonal dual ligation was realized through successful electronic fine-tuning of the SNO−OCT alkyne. The inclusion of a previously reported boronic acid/hydrazole bioorthogonal pair furnished a triple ligation system demonstrated to be mutually orthogonal. The development of highly chemoselective, bioorthogonal labeling reactions for double and triple ligations provides useful new tools for future in cellulo investigations. The utility of SNO-OCTs for biological applications was demonstrated through a series of protein and cellular labeling assays. The full synthetic potential of these strained alkynes is an area of ongoing investigation in our research groups, with the goal of creating easily accessible and robust bioorthogonal reagents to accelerate biological discovery.

# ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c06725.

- Characterization data, optimization tables, and additional substrates/catalysts, Part 1 (PDF)
- Characterization data, optimization tables, and additional substrates/catalysts, Part 2 (PDF)
- Molecular dynamics models of molecules as discussed in the text (ZIP)

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