Site-Selective Acylation of Natural Products with BINOL-Derived Phosphoric Acids

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Supporting Information

ABSTRACT: The site-selective acylation of a steroidal natural product 19-hydroxydehydroepiandrosterone catalyzed by 1,1′-Bi(2-napthol)-derived (BINOL) chiral phosphoric acids (CPAs) is described. Systematic variation and multivariate linear regression analysis reveal that the same steric parameters typically needed for high enantioselectivity with this class of CPAs are also required for site-selectivity in this case. Density functional theory calculations identify additional weak CH–π interactions as contributors to site discrimination. We further report a rare example of site-selective acylation of phenols through the evaluation of naringenin, a flavonoid natural product, using CPA catalysis. These results suggest that BINOL-derived CPAs may have broader applications in site-selective catalysis.

KEYWORDS: site-selective catalysis, acylation, chiral phosphoric acids, modeling, noncovalent interactions

Chiral phosphoric acids (CPAs) and phosphates based on the BINOL scaffold have been used extensively for a myriad of asymmetric transformations, suggesting their ability to effectively induce significant energetic differences between diastereomeric transition states across different reaction manifolds. Similarly, site-selective functionalization of natural products requires significant energetic differentiation between structurally distinct transition states, with the additional challenge of overcoming innate selectivity. Despite their successful employment as asymmetric catalysts, there have been limited cases of using BINOL-derived phosphoric acids in selective natural product modifications. Notably, Nagorny and co-workers demonstrated that CPAs are capable of distinguishing between different alcohols on polyketide antibiotics for site-selective glycosylation, and others have reported that CPAs can be effective for site-selective acetalizations of compounds with more than one hydroxyl group. Other classes of organocatalysts have been used in both enantio- and site-selective acyl transfers, such as peptides, borinic and boronic acids, covalent scaffolding catalysts, and benzotriazoles. Chiral metal complexes have also been employed. These precedents suggest that the same catalyst features enabling asymmetric induction may be translated to site-recognition in complex molecule settings. Furthermore, site-selective functionalization with CPAs may be achievable through strategic manipulation of the noncovalent substrate–catalyst interactions known to be crucial for asymmetric induction with CPAs. Herein, we report the site-selective acylation of two natural products using BINOL-derived CPAs as catalysts and demonstrate that the same catalyst features characteristically required for high enantioselectivity also enable high site-selectivity.

We first examined the site-selective acylation of 19-hydroxydehydroepiandrosterone (I, 19-OH-DHEA), a steroidal natural product with a hydrocarbon backbone. Evaluation of the acylation of I under common acylating conditions revealed that the secondary and primary alcohols in the neopentyl framework have comparable innate reactivity (Table 1, entries 1–2).

Inspired by Takasu’s report on the kinetic resolution of alcohols catalyzed by BINOL-derived phosphoric acids, we subjected I to acidic acylating conditions under the influence of catalyst 5a (Ar = 2,4,6-tricyclohexylphenyl, TCYP), which led to an increase in selectivity, from 1.2:1 to 8.5:1 in favor of acylation of the secondary alcohol. In this case, the chirality of the catalyst influences reactivity and selectivity—both the (S)-enantiomer of the catalyst 5b and an achiral counterpart, 5c, gave lower selectivity (Table 1, entries 3–5). Notably, 5b is also a less active catalyst, giving a lower yield of the acylated products compared with 73% yield with the “matched” catalyst 5a. Further optimization of the reaction conditions resulted in a 15:1 selectivity when catalyst 5a was used (Table 1, entry 6).

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Because site-selectivity is determined by the relative energy barriers of the two competing acylation pathways originating from the two alcohols,
16 we reasoned that multivariate linear regression (MLR) analysis previously adopted for studying BINOL-derived phosphoric acid-catalyzed enantioselective reactions
12d,17 could be applied here to identify catalyst features contributing to site-selectivity. Thus, we evaluated the acylation reaction with a panel of catalysts containing different substitution patterns at the R1−R3 positions from which we identified catalysts giving site-selectivities of up to 50:1 (Figure 1A).

To define the catalyst parameters important for selectivity, density functional theory (DFT) optimizations were performed for the catalyst panel at the M06-2X/def2-TZVP level of theory.
2d,17 From these optimized geometries, we collected steric and electronic parameters including Sterimol values, natural bond orbital (NBO) charges, and infrared (IR) vibrations (see Supporting Information for all parameters collected).
18,19 Comparing the collected parameters to the measured site-selectivity using a stepwise linear regression algorithm revealed a statistical correlation (R2 = 0.92, intercept = 0.07) with the terms R1B, R1L, and the NBOp charge (Figure 1B). The NBOp parameter is likely describing the hydrogen bonding capacity between the catalyst (phosphoric acid) and substrate (the primary or secondary alcohol), because the NBOp and NBOO measures are collinear (see Supporting Information for additional models). The minimum width of the R1 substituent, R1B, points to the importance of having bulky groups proximal (R2 and R1L) to the phosphoric acid moiety. The length of the R3 substituent, R1L, describes the importance of functionalization at the remote (R2, R3) positions for fine-tuning of site-selectivity.
50 An absence of these features resulted in lower selectivity (Figure 1). These same parameters were also present in previous MLR analyses of CPA-catalyzed enantioselective transformations, indicating that similar catalyst features required for high enantioselectivity
12e,17c are also beneficial for high site-selectivity in this case.

Previous computational models of CPA-catalyzed transformations have highlighted the contribution of steric bulk proximal to the phosphoric acid moiety to high enantioselectivity.
1a,b,d,e,17c,v About 19−21 To develop a model for understanding the importance of this catalyst feature in site-selective acylation, we first investigated several CPA catalyzed acylation mechanisms using DFT transition-state calculations (see Supporting Information for details). From these calculations, the most energetically favored pathway proceeded by a bifunctional activation of acetic anhydride and substrate simultaneously by the CPA as depicted within Figure 1C. Using these computational results and inspiration from previous structural descriptions of enantioselective CPA transformations, we propose that catalysts without adequate proximal bulk resulted in a less-defined catalyst pocket (Figure 1, 5d−5m). This in turn provides minimal energetic distinction between the transition states leading to 2 and 3 (Figure 1C, top).
1a,20 In contrast, the presence
1a,19 of bulky groups shapes the catalyst pocket such that differences arise upon association of each alcohol to the phosphoric acid. In considering the acylation of the primary alcohol, steric interactions with the bulky groups on the catalyst reduce the number of low-lying productive conformations. This effect is less pronounced when the secondary alcohol associates with the phosphoric acid, such that there are more productive conformations that avoid steric interactions with 2,6-disubstituents (Figure 1C, bottom).

An interesting trend emerged from the data obtained within this particular case as a function of variation of R3 group on observed site-selectivity. Although incorporating R3 substituents is generally beneficial for site-selectivity as indicated from the R1L term in the MLR analysis, the effect varied depending on the type of substituent. Aryl groups at R3 resulted in a dramatic improvement in site-selectivity (5n vs 5q−5s), while alkyl groups provided a modest enhancement (5n vs 5o). We hypothesized that the differences in selectivity were the result of attractive NCIs between the remote R3-phenyl substituent in 5q and the substrate, which are not captured by the Sterimol term in the model. Additionally, we reasoned that these NCIs should be present to a greater extent in the transition state leading to 2 than in the transition state leading to 3. To probe this hypothesis, we performed transition state analysis on the acylation reaction using 5q and a truncated 1 (see Supporting Information for computational methods). Multiple low-lying TS leading to products 2 and 3 are present. Boltzmann averaging of all TS leading to the formation of 2 and 3 resulted in a computed selectivity of 1.4 kcal/mol, consistent with the experimental results observed (Figure 2, 1.84 kcal/mol, see Supporting Information for details).
22 The relative abundance and strength of attractive NCIs between the relevant sites of interaction within TS2 and TS1 were then assessed using second-order perturbation theory (Supporting Information).
21,22 We found that CH−π interactions between the π system of the R3-phenyl of 5q and the C−H of the substrate are more abundant within TS2 than in TS1 (Supporting Information).

We next investigated the influence that the hydrocarbon framework of 1 has on site-selectivity. To examine this, 6, a derivative of 1 in which the alkene was removed by hydrogenation, was synthesized and subjected to acylation with the optimized conditions using catalyst 5q. A substantial decrease in selectivity was observed (8.8:1 for 6 vs 28.1:1 for 1, Scheme 1). This result prompted us to use TS analysis to
assess if this diminished selectivity resulted from weaker or less-abundant NCI’s between the R3-phenyl of 5q and 6. The calculated TS reproduced the experimental selectivity well (experimental ΔΔG‡ 1.1 kcal/mol, computed ΔΔG‡ 1.0 kcal/mol). The TS leading to 7 and 8 either completely lack or contain limited weak interactions with the R3-phenyl of 5q (Supporting Information). These results indicate that subtle changes in the conformation of the substrate can significantly affect the attractive noncovalent interactions needed for site-selectivity, highlighting the challenges associated with achieving site-selectivity in complex molecules.

As a final step, we sought to expand BINOL-derived CPA-catalyzed site-selective acylation to a different class of natural products with a drastically different framework. Derivatives of flavonoids are under study to optimize their wide range of biological activities, but few examples exist for catalyst- or reagent-controlled site-selective O-functionalization of phenols. We thus targeted the acylation of naringenin, a triphenol-containing flavonoid. Previous literature reports on the acylation and alkylation of naringenin with various acyl halides leverages the higher acidity of the phenol para to the ketone to achieve site-selectivity (pKₐ of C7-OH = 7.5, pKₐ of C4’-OH = 8.4). Consistent with this, we found that acylation of 9 under basic conditions is highly selective for the formation of 11 (Table 2, entry 1). Because the mechanism of CPA-catalyzed acylation does not likely involve deprotonation of the phenol prior to acylation (see Supporting Information), we hypothesized that acylation with phosphoric acids could provide a different selectivity profile. Indeed, using diphenylphosphoric acid (DPP) as the catalyst reverses the selectivity, albeit with low reactivity (Table 2, entry 2, 18% yield of acylated products). A higher temperature is required for acylation of the phenols compared with alcohols (45 °C vs 4 °C), but both reactivity and site-selectivity can be enhanced with catalysts 5k and 5u. Significant catalyst control of site-selectivity was achieved using the catalyst containing 3,3′-substituents with the bulkiest groups in the 2,3,4,6-positions, giving a selectivity of 7.0:1 for 10 (catalyst 5u, entry 5).

In summary, BINOL-based chiral phosphoric acids have enabled the site-selective acylation of a diol-containing steroidal natural product. MLR analysis reveals that the same catalyst features that create a defined catalyst pocket required for high enantioselectivity are also required for site-selectivity.
in this case. DFT calculations point to CH−π interactions between the natural product and catalyst as additional factors contributing to site-selectivity. Both analyses indicate that strategic structural modification of the substituents on the CPA could improve site-selectivity. Furthermore, using this mode of acylation catalyzed by CPAs, we achieved a rare example of site-selective acylation of phenols in a flavonoid natural product. Collectively, these findings suggest that this class of catalysts may have broader applications in site-selective catalysis.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscatal.9b03535.

Experimental details, procedures, compound characterization data, computational details, and copies of NMR spectra of new compounds (PDF)

X-ray crystallographic data for 6 (CIF)

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**Notes**

The authors declare no competing financial interest.

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(24) Another challenge associated with site-selective functionalizations is achieving selectivity for all reactive sites. In this case, we were not able to obtain site-selectivity for 3 with this catalyst system.


(27) For examples of the complementary catalytic enantioselective desymmetrizations of bis(phenols), see: (a) Lewis, C. A.; Gustafson, J. L.; Chiu, A.; Balsells, J.; Pollard, D.; Murray, J.; Reamer, R. A.; Hansen, K. B.; Miller, S. J. A Case of Remote Asymmetric Induction in the


