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Analyzing Site Selectivity in Rh₂.esp₂-Catalyzed Intermolecular C–H Amination Reactions

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ABSTRACT: Predicting site selectivity in C–H bond oxidation reactions involving heteroatom transfer is challenged by the small energetic differences between disparate bond types and the subtle interplay of steric and electronic effects that influence reactivity. Herein, the factors governing selective Rh₂.esp₂-catalyzed C–H amination of isoamylbenzene derivatives are investigated, where modification to both the nitrogen source, a sulfamate ester, and substrate are shown to impact isomeric product ratios. Linear regression mathematical modeling is used to define a relationship that equates both IR stretching parameters and Hammett σ⁺ values to the differential free energy of benzylic versus tertiary C–H amination. This model has informed the development of a novel sulfamate ester, which affords the highest benzylic-to-tertiary site selectivity (9.5:1) observed for this system.

INTRODUCTION

Discriminate control over product selectivity in carbon–hydrogen (C–H) bond functionalization reactions represents one of the great challenges in modern synthetic chemistry.¹ The high energy barriers to C–H bond cleavage (on the order of 98 kcal mol⁻¹) contrast the small energetic differences that bias enantio- and chemoselective C–H bond functionalization (∆∆G° of ~2 kcal mol⁻¹ for >20:1 selectivity). Given the small differences in transition state free energies that modulate isomeric product ratios, it is often difficult to distinguish the steric and electronic factors that influence reaction selectivity. Identification of such factors, however, can prove invaluable for tailoring catalyst and reagent structures to afford greater control over reaction outcomes.

The Du Bois group recently reported an intermolecular Rh-catalyzed C–H amination protocol and demonstrated that oxidation of isoamylbenzene (a) results in benzylic-to-tertiary (B:T) product ratios that are dependent upon the choice of sulfamate ester b (Figure 1).¹ The relationships between steric and electronic factors that contribute to these disparate outcomes are not obvious from the trends in selectivity. Specifically, sulfamate ester b₁, R = CH₂C₂Cl₂, yielded the highest degree of B:T selectivity (8:1), while substitution to R = CH₂t-Bu (b₂), a steric homologue, resulted in reduced benzylic selectivity (4:1). An equally intriguing result was obtained from the evaluation of sulfamate ester b₃, R = CH₂CF₃, which yields equimolar amounts of the two products. Similar losses in selectivity were observed for both electron-poor (b₄, R = 2,6-F₂C₆H₃, 1.5:1) and electron-rich (b₅, 4-t-BuC₆H₄, 1:1) aryl sulfamate esters.

An archetypical physical organic technique for identifying features that influence product selectivity as a function of substituent changes is linear free-energy relationship (LFER) analysis.² Pioneered by Hammett for electronic analysis of meta- or para-substituted benzene rings³ and adopted by Taft⁴ and, later, Charton⁵ for steric effect analyses, these techniques have been broadly applied to interrogate reaction outcomes.⁶ While these classic LFER parameters have been instrumental in a variety of contexts, often illuminating mechanistic details by relating log(K) to empirically derived electronic or steric constants (where K may represent relative rate and equilibrium constants, ratios of enantiomers and constitutional isomers, etc.), LFERs also bear significant limitations;⁷ namely, there are a modest number of reactions that can be successfully modeled using Hammett or Taft/Charton parameters alone.⁸,⁹

Over the last several years, the Sigman laboratory has investigated the use of discretely measured molecular parameters (vide infra) as opposed to those derived from relative-rate experiments (e.g., Hammett and Taft values) for nonclassic free-energy relationship analysis, relating these parameters to ∆∆G° for differential transition state interrogation.⁹b,¹¹ As the data from the Rh-catalyzed C–H amination lacks obvious explanation, commonly employed free-energy relationships are not likely capable of delineating the entangled effects of the sulfamate ester on site selectivity. Therefore, we have turned to a recent discovery that specific infrared (IR) molecular vibrations represent a broadly applicable, yet uniquely descriptive, parameter set.¹⁰a IR vibrations can be computationally calculated for any molecule, the result of which is a tailored parameter set that is capable of describing the distinct nature of each reactive species.

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Herein, we exploit the intrinsic ability of IR vibrations to describe the inherent molecular properties of sulfamate ester nitrene precursors in selective Rh₂(esp)₂-catalyzed amination of benzylic versus tertiary C–H bonds. Using IR-derived descriptors to quantitate steric and electronic selectivity determinants, we apply linear regression modeling to identify the sulfamate ester features responsible for differential nitrene induction of the Rh₂(esp)₂-catalyzed amination of isoamylbenzene (Figure 1a).

Each of the sulfamate esters depicted in Figure 2 was evaluated in the Rh₂(esp)₂-catalyzed amination of 1. Of particular interest, chlorine substitution (2e–2g) has a pronounced effect on product selectivity with trichloromethyl sulfamate esters (2k, 2l) yielding B:T ratios of ∼9:1, regardless of the proximity of this group to the −SO₂NH₂ moiety. Relative to R = nBu (2d), this same trend is maintained for di- and monochloromethyl sulfamate esters, where B:T ratios average 5.3:1 (2i, 2j) and 4.5:1 (2e–2g), respectively.

The insensitivity of B:T selectivity to chain length is a general trend observed throughout the data set. The influence of steric effects on selectivity becomes apparent when the sulfamate ester bears a branched α-carbon (i.e., sulfamates prepared from secondary alcohols). Specifically, selectivity for the benzylic insertion product increases for di- and monochloromethyl sulfamate esters (2h, 2l) relative to EtOSO₂NH₂ (2b, 5.9:1). A marked change in the product ratio is noted when halogen substituents are introduced in these secondary alcohol-derived sulfamate esters (2h, 2l). For example, a reaction performed with (CF₃)₂CHOSO₂-NH₂ produces nearly equal amounts of the benzylic and tertiary product (2m–2o).

**Parameter Selection.** Collectively, the data portrayed in Figure 2 reflect an ill-defined role for steric and electronic modulation of the sulfamate ester on product selectivity. Steric influences manifest principally in the narrow dimension of branched versus nonbranched sulfamate groups. Additionally, while inclusion of electronegative halogen atoms clearly alters product selectivity, the effect cannot be ascribed entirely to electronic differences in nitrenoid reactivity. These general features of the amination reaction significantly complicate quantitative free-energy modeling of selectivity. Classic steric parameters, such as Taft² and Charton values and Winsstein–Holness (A) values³, derived from relative-rate and conformation equilibration experiments, respectively, treat substituent steric bulk as a spherical unit.⁴b Therefore, this treatment has the disadvantage of averaging the nuances of substituent asymmetry and length-to-length ratios into a single-value representation of steric effects.
In the development of free-energy relationships describing selectivity, it is precisely these subtleties that are responsible for the differential transition state energies leading to isomeric product ratios, as predicated by the Curtin–Hammett principle. Verloop innovatively approached this deficiency in the description of steric effects through the development of Sterimol parameters (Figure 3). This parameter set gives dimensional specificity to the description of steric bulk through three subparameters: B1 (minimum radius), B5 (maximum radius), and L (length). Comparisons of nPr and t-Bu demonstrate a deficiency in the Sterimol parameters, where sterically distinct groups are similarly described.

While the effectiveness of Sterimol parameters in various contexts has been successfully demonstrated, this steric descriptor still lacks information about the position along L at which steric bulk resides. For example, as depicted in Figure 3, Sterimol measures of the CH2 substituent are 1.52 (B1), 4.18 (B5), and 4.89 (L). Comparatively, the Sterimol system describes nPr, a group with its own distinct apparent steric bulk, as nearly isosteric with CH2 and t-Bu, measuring 1.52 (B1), 3.49 (B5), and 4.92 (L). A similar parameter deficiency occurs for electronic description. The presence of R-group chlorine atoms, particularly trichloromethyl, generally enhances selectivity (in the absence of branching), independent of the chlorine atom distance from the –NH2 group of the sulfamate moiety. This observation cannot be explained through the use of the ubiquitous electronic descriptor, pKa, or any descriptor of inductive effect.

While the reactive oxidant believed to be involved in the Rh-catalyzed amination is a nitrene (f, Figure 1b), our computed vibrational data are from the sulfamate ester and not the nitrenoid. As noted above, the differential energy between nitrenoid transition states (ΔΔG°‡) is responsible for benzylic versus tertiary amination ratios. Our working hypothesis, for which we provide supporting evidence, is that modifications to the nitrene precursor (i.e., sulfamate ester) commensurately impact molecular properties of the selectivity-defining transition states (vide infra). This is an important qualification, which allows ground state IR frequencies and intensities to be computed for the simplest of these species, the sulfamate ester. This approach significantly reduces the computational effort, making the methodology tractable. In order to proceed with free-energy relationship model development, we identified a group of IR vibration parameters as potential selectivity descriptors. From such a set, stepwise linear regression analysis is performed, whereby the descriptors...
are statistically whittled down to a subset of parameters that best mathematically relates features of the sulfamate ester to $\Delta \Delta G^i$ (equating to $-RT \ln(\text{tertiary/benzylic})$, where $R$ is the ideal gas constant and $T$ is temperature). As each sulfamate ester is characterized with many disparate vibrational modes, we chose those vibrations that were consistently identified in our computations (i.e., major vibrational modes) and assumed to significantly impact the Rh-nitrene selectivity profile. Given these criteria, four vibrations were chosen as potential descriptors of selectivity: O–S–N asymmetric stretch ($\nu_{\text{OSN}}$), C–O stretch ($\nu_{\text{CO}}$), $\text{SO}_2$ symmetric stretch ($\nu_{\text{SO}_2\text{sym}}$), and $\text{SO}_2$ asymmetric stretch ($\nu_{\text{SO}_2\text{asym}}$). Figure 4 depicts a simulated IR spectrum for sulfamate ester 2a ($R = \text{Me}$) and highlights both the calculated frequencies and intensities of these four vibrations, giving a total of eight vibration-derived descriptors that were used for regression analysis.

**Model Development.** Prior to developing a mathematical relationship between selectivity and the identified vibrational frequencies and intensities, we first applied design of experiments (DoE) principles to our initial 20-membered sulfamate ester library (Figure 2).\(^{16}\) DoE tenants dictate that the most robust mathematical models are developed from data sets that are systematically varied. Thus, eight sulfamate esters were selected (terming the DoE set and noted with asterisks and bolded in Table 1) that quantitatively sample the observed range of B:T ratios and qualitatively represent a distribution of steric and electronic perturbations.

In addition to examining sulfamate substituent effects on B:T selectivity, we have also varied the electronic structure of the isomylbenzene substrate. After preparing a traditional Hammett series ($R = \text{OMe}$ (1a), $t$-Bu (1c), Br (1d), CF$_3$ (1e)), this library was subjected to oxidation reactions with each of the eight DoE-set sulfamate esters. (See Figure 5 for a description of this library was subjected to oxidation reactions with each of the eight DoE-set sulfamate esters. (See Figure 5 for a description of the two sets of experiments that did not yield measurable data.))

Table 1. Training Set (Entries 1–23), External Validations (Entries 24–61), and Predictions (Entries 62–64, Bold)

| entry | R | $R'$ | $\Delta \Delta G^i$ (kcal/mol) | $\Delta \Delta G'^i$ (kcal/mol) | meas. B/T |
|-------|---|-----|-----------------|-----------------|---|
| 30    | CH$_2$Cl$_2$ & 4-H | 1.33 | 0.92 | 4.8 ± 0.1 |
| 31    | (CH$_2$)$_2$Cl & 4-H | 0.99 | 0.90 | 4.6 ± 0.1 |
| 32    | CH$_2$-Bu & 4-H | 1.15 | 0.86 | 4.3 ± 0.3 |
| 33    | CH$_2$Cy & 4-H | 0.89 | 0.84 | 4.2 ± 0.1 |
| 34    | nBu & 4-H | 1.05 | 0.77 | 3.7 ± 0.1 |
| 35    | CH(CF$_3$)$_2$ & 4-H | 1.19 | 0.06 | 1.1 ± 0.1 |
| 36    | CH$_3$Cl$_2$ & 4-Bu | 1.96 | 2.27 | 476 ± 0.3 |
| 37    | CH$_2$CF$_3$ & 4-Bu | 2.61 | 2.33 | 52.8 ± 0.1 |
| 38    | nPr & 4-Bu | 1.38 | 1.29 | 9.0 ± 0.3 |
| 39    | (CH$_2$)$_2$Cl & 4-Bu | 1.44 | 1.47 | 12.2 ± 0.1 |
| 40    | (CH$_2$)$_2$-Bu & 4-Bu | 1.21 | 1.20 | 7.7 ± 0.2 |
| 41    | CH(CH$_2$Cl)$_2$ & 4-Bu | 1.18 | 1.51 | 13.1 ± 0.1 |
| 42    | CH$_2$Pr & 4-Bu | 1.21 | 1.28 | 8.8 ± 0.5 |
| 43    | (CH$_2$)$_2$Cl & 4-Bu | 1.25 | 1.51 | 13.0 ± 0.5 |
| 44    | CH$_2$Cl$_2$ & 4-Br | 1.37 | 1.16 | 7.2 ± 0.1 |
| 45    | CH$_2$CF$_3$ & 4-Br | 1.04 | 1.15 | 7.1 ± 0.1 |
| 46    | nPr & 4-Br | 0.88 | 0.90 | 4.6 ± 0.3 |
| 47    | (CH$_2$)$_2$Cl & 4-Br | 0.88 | 1.03 | 5.8 ± 0.2 |
| 48    | (CH$_2$)$_2$-Bu & 4-Br | 0.77 | 0.97 | 5.2 ± 0.2 |
| 49    | CH$_2$Pr & 4-Br | 0.82 | 0.98 | 5.3 ± 0.4 |
| 50    | (CH$_2$)$_2$Cl & 4-Br | 0.77 | 1.15 | 7.1 ± 0.3 |
| 51    | CH$_2$CHCl$_2$ & 4-OMe | 2.23 | 2.40 | 59.3 ± 0.4 |
| 52    | (CH$_2$)$_2$Cl & 4-OMe | 1.88 | 2.21 | 47.2 ± 0.5 |
| 53    | nBu & 4-OMe | 2.01 | 1.74 | 19.3 ± 0.5 |
| 54    | nBu & 3-Cl | 0.56 | 0.63 | 2.9 ± 0.2 |
| 55    | (CH$_2$)$_2$Cl & 3-Cl | 0.55 | 0.52 | 2.4 ± 0.1 |
| 56    | CH$_2$Pr & 3-Cl | 0.59 | 0.49 | 2.3 ± 0.1 |
| 57    | (CH$_2$)$_2$Cl & 4-Ph | 1.33 | 1.68 | 17.3 ± 0.2 |
| 58    | nBu & 4-Ph | 1.27 | 1.39 | 10.7 ± 0.2 |
| 59    | CH$_2$CF$_3$ & 3-Br | 1.33 | 1.41 | 10.9 ± 0.4 |
| 60    | (CH$_2$)$_2$Cl & 3-Br | 1.17 | 0.92 | 4.8 ± 0.1 |
| 61    | nBu & 3-Br | 1.12 | 0.79 | 3.8 ± 0.2 |
| 62    | CH$_2$CF$_2$CF$_3$ & 4-H | 1.38 | 1.32 | 9.5 ± 0.2 |
| 63    | (CH$_2$)$_2$CF$_3$ & 4-H | 1.43 | 1.26 | 8.5 ± 0.1 |
| 64    | CH$_2$(CH$_2$Me)$_2$CH$_2$Cl & 4-H | 1.17 | 1.06 | 6.1 ± 0.1 |
H(1), CF₃(1b)), we subjected the 23-membered training set (Table 1, see Figure 5 for an explanation of the data point omitted) to a standard stepwise linear regression algorithm (see Supporting Information for details).²¹ Using this algorithm, which facilitates statistical exploration of the relationship between vibrational parameters, $\sigma^+$, and $\Delta \Delta G^\ddagger$, the equation depicted in Figure 6a can be formulated. To evaluate the accuracy of this model, we compare predicted and measured $\Delta \Delta G^\ddagger$ in Figure 6b, which demonstrates a high level of correlation between experimental values and model predictions. Leave-one-out (LOO) analysis was also performed to evaluate the robustness of the model (Figure 6c).²² The slope and $R^2$ values, which are close to unity, are positive indicators of the model’s accuracy.

**External Validation of the Model.** A third measure of model strength was determined by externally validating the model with data points not part of the training set. Of the original 20-membered library, 12 sulfamate esters, which were not members of the DoE set, were evaluated with isoamylbenzene (1). The robustness of the model for describing substrate variation was evaluated with five isoamylbenzene derivatives: 1-t-Bu-4-isopentylbenzene (1c), 1-bromo-4-isopentylnbenzene (1d), 1-Cl-3-isoamylbenzene (1e), 1-Ph-4-isoamylbenzene (1f), and 1-t-Bu-3-isoamylbenzene (1g). The complete external validation set is tabulated in Table 1. Graphical representation of this data (red squares, Figure 6b) demonstrates the overall good agreement between predicted $\Delta \Delta G^\ddagger$ values and experimental measurements. An obvious outlier between predicted and measured $\Delta \Delta G^\ddagger$ values occurs with sulfamate ester 2n, (CF₃)₂CHOSO₂NH₂. We hypothesize that this highly electron-deficient, sterically large sulfamate ester may be forced to adopt conformations not accessible to other nitrene sources in the defining C=N bond forming event. It is also possible that 2n facilitates C=H amination through a mechanistic pathway that differs from that of other sulfamate esters. Future investigations of reactions with 2n are warranted; use of this reagent was discontinued for the remainder of this study.

Figure 6. (a) Normalized mathematical relationship, derived from tabulated training set in Table 1, describing differential free energy of benzyl (B)-to-tertiary (T) amination. R: ideal gas constant, T: 23 °C. (b) Predicted versus measured $\Delta \Delta G^\ddagger$ plot of training set and external validations. Grayed data point, designated as an outlier, represents R = CH(CF₃)₂, R’ = H. (c) Leave-one-out (LOO) analysis.

Figure 7. (a) Representation of increasing C=O stretch frequency ($\nu_{CO}$) versus sulfamate ester R group. (b) Representation of increasing intensity of O=S=O asymmetric stretch ($I_{OSO}$) versus sulfamate ester R group. Grayed columns highlight model-informed predictions 2u (R = CH₂CF₂CF₃) and 2v (R = CH₂(CF₂)₂CF₃).
Analysis of the Model. We have capitalized on the robustness of our model, which relies on both vibrational data and substrate σ⁺ parameters, to predict new sulfamate structures that display a higher propensity toward benzylic C–H insertion. As the relationship in Figure 6a is a normalized equation, the magnitude of the coefficients yield information about the relative influence of each parameter on selectivity. Notably, the overriding selectivity determinant, σ⁺, is associated with the strength of the benzylic C–H bond (vide supra). Perhaps unsurprisingly, the C–O frequency (ν(CO)) of the sulfamate ester also plays a prominent role in this model. Included as a single term and, again, within a cross-term, ν(CO) is the shortest conduit from the O-alkyl substituent to the sulfamate moiety. The vibrational frequency of the C–O bond will reflect changes in the substituent groups on the alkyl chain, which alter the force constant and/or reduced mass (components of vibrational modes).

It is particularly intriguing to find that the C–O stretching vibration is coupled with the Hammett descriptor, σ⁺, in the optimized selectivity model. This result suggests a synergistic relationship between the nitreneoid and the isoamylbenzene substrate, indicative of a defined intermolecular interaction between these two species. While the precise nature of this interaction is unclear, we considered illuminating the origin of the σ⁺ trends by assessing sulfamate esters according to increasing vibrational frequencies (Figure 7a). Qualitatively, we observed that the more halogenated sulfamate esters showed greater ν(CO) values. In accordance with this trend, more polarized bonds vibrate with energetically higher frequencies. Greater differential electronegativity across a bond increases the bond force constant and, thus, its vibrational frequency.23

Patterning in a similar manner our analysis of the other vibration-related parameter, I_OSN, revealed in our model, we constructed Figure 7b, which displays sulfamate ester R groups according to increasing O–S–N asymmetric stretch intensities. Organizing the data in this manner, we observe that variation in I_OSN is primarily characterized by increases in distal steric bulk and by halogenation. These qualitative trends served to inform our use of the developed model as a tool for predicting new sulfamate esters that yield improved B:T ratios.

Application of the Model. We have computationally evaluated several sulfamate derivatives that included electro-negative atoms and variation in chain-length; most of these, however, were not predicted to afford improved site selection. In contrast, sulfamate esters 2u (R = CH2CF2CF3) and 2v (R = CH2(CF2)2CF3) were identified using our model, as these two reagents were expected to give enhanced levels of the benzylic oxidation product. In practice, the predicted selectivities closely matched those measured, with sulfamate ester 2u effecting the highest degree of site-selection observed for amination of isoamylbenzene (1) (Figure 8a). The enhancement of selectivity achieved by changing the sulfamate from Cl3CCH2OSO2NH3 to CF3CF2CH2OSO2NH2 (2u), albeit modest, is striking given the apparent electronic similarities and steric differences between these two reagents.

The identification of 2u and 2v by consideration of both ν(CO) and I_OSN (gray columns, Figure 7) highlights the predictive utility of our model. Of note, the calculated IR frequencies and intensities of these nitrene sources do not represent the highest observed values in the sulfamate ester library. This is rationalized by considering the interdependency of the terms derived from vibrational modes, since these are intrinsically linked. Thus, maximizing the value of ν(CO) alone does not guarantee proportionate increases in ΔΔG° values. This underscores the balance that is achieved in the developed model (see equation in Figure 6a) between the selectivity-enhancing effects of the ν(CO) and I_OSN parameters (positive coefficients) and the potentially deleterious effect on site selection of the (ν(CO))(σ⁺) cross term (negative coefficient).

As a final step, we have evaluated the performance of sulfamate ester 2u on a preparative scale (0.5 mmol) with isoamylbenzene (1). The benzylic product from this reaction was obtained in 58% yield with the same level of B:T site-selectivity (9.4:1) that was noted in the original evaluation process (0.3 mmol scale). The reaction of 2u with substrate 5 shows even higher levels of site selectivity in favor of the benzylic amine product (60% yield). Finally, oxidation of a more
sophisticated polycyclic substrate, 6, is demonstrated to give exclusively the product of secondary, benzylic oxidation in 55% yield.

In summary, the subtle interplay of steric and electronic effects in the Rh-catalyzed C–H amination of isoamylbenzenes has been evaluated with a wide-range of sulfamate esters. Product selectivity in these reactions can be effectively modeled using a combination of a classical Hammett parameter and computed IR vibrational data. Of particular interest is the ability to deconstruct the model and use this information to extrapolate to new sulfamate esters, one of which offers the highest performance, to date, for this intermolecular Rh-catalyzed C–H amination reaction. Current efforts are underway to apply this modeling approach to examine other selectivity challenges in C–H functionalization reactions and to use the results of these investigations to deduce relevant transition state models.

ASSOCIATED CONTENT

## Supporting Information

Experimental procedures, characterization data for new substances, mathematical modeling details. This material is available free of charge via the Internet at http://pubs.acs.org.

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### Notes

The authors declare no competing financial interest.

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