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Highly Diastereoselective Functionalization of Piperidines by Photoredox Catalyzed α-Amino C–H Arylation and Epimerization

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ABSTRACT: We report a photoredox catalyzed α -amino C–H arylation reaction of highly substituted piperidine derivatives with electron deficient cyano(hetero)arenes. The scope and limitations of the reaction were explored, with piperidines bearing multiple substitution patterns providing the arylated products in good yields and with high diastereoselectivity. In order to probe the mechanism of the overall transformation, optical and fluorescent spectroscopic methods were used to investigate the reaction. By employing flash-quench transient absorption spectroscopy, we were able to observe electron transfer processes associated with radical formation beyond the initial excited state $Ir(ppy)_3$ oxidation. Following the rapid and unselective C–H arylation reaction, a slower epimerization occurs to provide the high diastereomer ratio observed for a majority of the products. Several stereoisomerically pure products were re-subjected to the reaction conditions, each of which converged to the experimentally observed diastereomer ratios. The observed distribution of diastereomers corresponds to a thermodynamic ratio of isomers based upon their calculated relative energies using density functional theory (DFT).

INTRODUCTION

Piperidines are by far the most prevalent of all heterocycles found in drugs.¹ For example, the piperidine substructure is present in the blockbuster antidepressant paroxetine, morphine along with its congeners, many of the antihistamine class of drugs, and tofacitinib used in the treatment of arthritis and ulcerative colitis.¹ Substitution about the piperidine scaffold is extremely common,^{1a} and for this reason, efficient new methods for the diastereoselective elaboration of the piperidine framework have the potential to greatly facilitate the discovery of new pharmaceutical agents.

Photoredox catalysis utilizing transition metalpolypyridyl complexes has provided a powerful strategy to access novel reactivity manifolds.² Although the utility of α amino radicals has long been appreciated,³ the MacMillan

Scheme 1. Photoredox catalyzed α-amino C-H arylation

A. Previously reported α-amino C-H arylation (MacMillan, 2011)

$$R^{1} \xrightarrow[R^{2}]{} H^{+} + ArCN \xrightarrow{Ir(ppy)_{3}, NaOAc} R^{1} \xrightarrow[R^{2}]{} R^{2} \xrightarrow{R^{3}} Ar$$

B. This work: Diastereoselective C-H arylation of highly substituted piperidines



group provided a seminal study using a transition metal photoredox catalyst for the synthesis of α -branched amines via α -amino radical coupling with electron deficient cy-ano(hetero)arene derivatives (Scheme 1A).^{4,5} Following this report, many groups have developed transformations using α -amino radicals as the reactive intermediate for additions to alkenes and other unsaturated π -bonds,⁶ cross-coupling with (hetero)arenes,⁷ and other coupling partners.⁸

Achieving stereoselective transformations is one of the major challenges for photoredox catalysis because radicals serve as key reactive intermediates.⁹ In this regard, significant advances have been realized for the asymmetric synthesis of α -branched amines via α -amino radical intermediates using photoredox catalysts often in combination with other modes of catalysis.¹⁰ Approaches for the diastereoselective synthesis of α -branched amines to produce compounds that incorporate two or more stereogenic centers via α-amino radical intermediates have also been developed.^{10c,11} However, the elaboration of complex molecules incorporating stereogenic center(s) requires that diastereoselectivity be achieved relative to the pre-existing stereogenic center(s). This type of stereoselective transformation has rarely been explored for photoredox catalyzed α -amino radical reactions.11b,12,13

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Although the reactivity of α -amino radicals can introduce challenges for achieving stereoselective transformations, the low inversion barrier for these radicals also provides new opportunities for stereoselective synthesis by epimerization. Pioneering studies by Bertrand and coworkers demonstrated that a thiyl radical could mediate the racemization of benzylic amines through reversible hydrogen abstraction.¹⁴ Very recently, Knowles and Miller have achieved an impressive photo-driven deracemization of cyclic ureas based on the use of excited-state redox events.^{10a} Moreover, in a photoredox catalyzed reverse polarity Povarov annulation to give disubstituted tetrahydroquinolines, an increase in diastereoselectivity to 18:1 was observed when the initial 10:1 ratio of product diastereomers was resubmitted to the reaction conditions.^{11e,15,16}

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Herein, we report the development of a highly diastereoselective Ir(III) photoredox catalyzed reaction cascade that proceeds by α -amino C–H arylation of densely functionalized piperidines **1** followed by epimerization at the α -position to give the most stable stereoisomer **2** (Scheme 1B). Notably, differentially substituted piperidine derivatives with up to four stereogenic centers were effective substrates, providing the products with generally high diastereoselectivity relative to the pre-existing stereogenic centers. We propose mechanisms for both the C-H arylation reaction and epimerization steps based on an array of spectroscopic studies and time courses for product formation. Our results are consistent with an unselective photoredox C–H arylation followed by product epimerization leading to the observed diastereoselectivity. The diastereomers underwent oxidation with comparable rate constants, and therefore, product oxidation kinetics were not responsible for the epimerization diastereoselectivity. We subjected the separate diastereomers to the photoredox-catalyzed reaction conditions, and each yielded the experimentally observed distribution of diastereomers suggesting that the reaction is under thermodynamic control. The calculated relative energies of the diastereomers using density functional theory (DFT) correlate with the observed diastereomer ratios.

RESULTS AND DISCUSSION

Efficient Preparation of Piperidine Starting Materials 1. We first developed a facile two-step route to diastereomerically pure, densely substituted piperidines 1 from readily available precursors (Scheme 2). Previously, we reported a Rh(I)-catalyzed C-H activation/electrocyclization cascade to furnish highly substituted 1,2-dihydropyridines 4 from imines 3 and internal alkynes.¹⁷ Catalytic hydrogenation of dihydropyridines 4 with Pd/C then provides piperidines 1 (see Supporting Information for optimization Table S1 and experimental details).¹⁸ In all cases, only the all-cis stereoisomer was detected and was isolated in 63% average overall yield for the two step process.

Optimization and Scope of C–H Arylation. To optimize the yield and diastereoselectivity of the C–H arylation reaction, we explored a variety of conditions for arylating trisubstituted piperidine **1a** ($R^1 = H$, $R^2 = Me$, $R^3 = R^4 = Et$) with 1,4-dicyanobenzene (DCB) to give **2a** (Tables S2 and S3 in the Supporting Information). Ir(ppy)₃ (ppy = 2-phenylpyridine) was determined to be the optimal photocatalyst for Scheme 2. Highly diastereoselective two-step synthesis of densely substituted piperidines



the α -arylation, in agreement with MacMillan's earlier report.^{4,19} Of particular note, we were able to employ the piperidine as the limiting reagent to give **2a** in high yield and with high diastereoselectivity (Table 1). For this transformation, arylation unambiguously provided the syn isomer as determined by X-ray crystallography.

The optimized conditions were next applied to a number of piperidine derivatives using DCB as the coupling partner (Table 1). Piperidines with the strained cyclopropyl ring (**2b**) and the linear propyl chain (**2c**) at R² were effective substrates in the reaction. A phenyl substituent could also be incorporated at the R² position to provide product **2d** in good yield and with high diastereoselectivity, although a longer reaction time of 16 h was required.

Trisubstituted piperidines with a different substitution pattern also provided the arylated products in high yield and diastereoselectivity as exemplified for 2e-2k. However, for this substitution pattern, the anti-stereoisomer was obtained, as unambiguously determined by X-ray crystallography for 2e. Piperidines with cyclopentyl (2f), Boc protected piperidine (2g), and isopropyl (2h) at R⁴ were all effective substrates in the reaction. Notably, arylation of the *N*-Boc piperidine in **2g** does not occur under our reaction conditions, due to the thermodynamically unfavorable oxidation of electron-deficient carbamates.²⁰ Different N-aryl substituents were also evaluated, with products 2h-2k each obtained in good to high yields and with excellent diastereoselectivities. However, for alkyl, acyl and toluenesulfonyl substituents on the piperidine nitrogen, no coupling was observed (for a full list of unsuccessful reactants, see Table S4 in the Supporting Information).

Arylation of a tetrasubstituted piperidine gave the fully substituted piperidine product **2l** in high yield and with 85:15 diastereoselectivity for the anti isomer.²¹ A monosubstituted piperidine also arylated in high yield to give product **2m** with modest diastereoselectivity and with a preference for the syn isomer.²¹

Using piperidine **1a**, we explored the scope with respect to the cyano(hetero)arene coupling partner (**2n-2q**). Heterocycles, such as for 4-cyanopyridine, provided product **2n** in moderate yield and with high selectivity. The desired products were also obtained in moderate to good yields for other electron deficient cyanoarenes, including a para substituted morpholine amide (**2o**), phthalide (**2p**) and ethyl ester (**2q**). Arylation with the electron-rich 4-methoxybenzonitrile and the electron-neutral cyanobenzene were not successful, consistent with the redox potentials of these derivatives²² and with MacMillan's initial report on tertiary aniline arylation.^{23,4} MacMillan has successfully coupled heteroaryl chlorides with unhindered tertiary anilines;²⁴

however, with our more hindered, substituted piperidines less than 10% coupling occurred with 2-chlorobenzothiazole (for a full list of unsuccessful reactants, see Table S4 in the Supporting Information). All of the cyano(hetereo)arene coupling partners were additionally explored with a trisubstituted piperidine with a different substitution pattern and provided products **2r-2u** in moderate to good yields and with excellent diastereoselectivities in all cases.

The large majority of piperidines that we investigated for arylation were efficiently prepared by face selective hydrogenation of DHPs 4 to give the all-syn substituted diastereomer 1 (see Scheme 2). To demonstrate the potential generality of the approach to other stereoisomers, we also evaluated the arylation of diastereomer 5 (eq 1), which dis-

played the 3-ethyl group anti to the two other alkyl substituents (for the synthesis of **5**, see the Supporting Information). Under the standard reaction conditions, product **2v** was obtained in good yield and with 84:16 diastereose-lectivity.

We further explored functional group compatibility of the reaction by evaluating readily accessible 4-substituted piperidines (Table 2). These unhindered piperidine substrates are capable of over-addition, which introduces an added challenge for obtaining high product yields. For photoredox mediated arylations of unhindered tertiary anilines, over-addition is typically minimized by using excess

Table 1. Substrate scope^a



^{*a*}Isolated yields on 0.2 mmol scale; *dr* was determined by ¹H NMR analysis of the purified product and was within a single percentage point relative to the crude *dr*. ^{*b*}X-ray structure shown with anisotropic displacement ellipsoids at the 50% probability level. The picryl sulfonate counterion and hydrogen atoms are omitted for clarity. ^{*c*}87:13 *dr* prior to isolation determined by crude ¹H NMR analysis. ^{*d*}16 h reaction time. ^{*e*}2 mol % Ir(ppy)₃, 72 h reaction time. ^{*f*}[Ir(dtbbpy)(ppy)₂]PF₆ instead of Ir(ppy)₃.

of the tertiary amine substrate.⁴ We instead chose to employ the DCB coupling partner in a 1:1 stoichiometry because the substituted piperidines are the more expensive of the two inputs. Even with this stoichiometry, the acetamide substituted piperidine provided the arylated product 2w in excellent yield though with modest diastereoselectivity. Products were also obtained in reasonable yields and modest diastereoselectivities for hydroxy (2x), hydroxymethyl (2y), and methoxymethyl (2z) substituents at the 4-position.

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Mechanistic Investigations. To better understand the mechanism of product formation, we investigated the reactivity of the Ir(ppy)₃ photosensitizer and DCB intermediates using optical and fluorescent spectroscopies. Upon irradiation at 450 nm, the Ir complex undergoes a metal-to-ligand charge transfer followed by rapid intersystem crossing to give the long-lived triplet state that engages in single electron transfer.25 Based on the reduction potential of *Ir(ppy)₃ (* $E_{1/2}$ (Ir ^{IV/*III}) = -1.73 V vs. SCE in MeCN)²⁶ and DCB (DCB/DCB⁻ E_p = -1.61 V vs. SCE in MeCN),²⁷ we would expect an oxidative quenching mechanism to be operative, generating Ir^{IV} and the DCB⁻⁻ in the first step as depicted in Figure 1A. Indeed, luminescence quenching studies revealed fast electron transfer (ET) between DCB and *Ir(ppv)₃, occurring with a rate constant $k_{quench} = 2.9 \times 10^9$ M⁻¹s⁻¹ in *N*,*N*-dimethylacetamide (DMA). In contrast, high concentrations of piperidine 1m caused minimal change in the *Ir(ppy)₃ lifetime (see Figure S2 in the Supporting Information). These data, and the experimental excess of DCB, support an initial excited state ET producing Ir(ppy)3+ and DCB⁻⁻ and are also in agreement with MacMillan's previous observations.4

In order to examine the reactivity of these photoproducts, we pursued the visible absorption spectra for each. We obtained authentic absorption spectra for Ir(ppy)₃⁺ from spectroelectrochemical oxidation of Ir(ppy)₃ in DMA (see Figure S3 in the Supporting Information), and found authentic spectra for the DCB⁻⁻ in the pulse radiolysis literature.²⁸ We fortuitously observed an isosbestic point for the Ir(ppy)₃ to Ir(ppy)₃⁺ conversion at 346 nm, very close the λ_{max} of the DCB⁻⁻ species. This allowed us to observe absorption changes at 346 nm that correspond only to the evolution of DCB⁻⁻.

The reactivity of the Ir(ppy)₃⁺ and DCB⁻ species is not visible in luminescence quenching studies, and so we turned to flash-quench transient absorption spectroscopy (TA) to observe the ground state reactivity of these photogenerated species. In the absence of piperidine substrate, Ir(ppy)₃⁺ and DCB⁻ recombine at or near the diffusion limit (Figure 1C), which we can follow at 346 nm for the DCB⁻ decay and at 390 nm for the Ir(ppy)₃⁺ decay (Figure 1B, S4 and S5 in the Supporting Information). This recombination follows an equal concentration bimolecular kinetic model (see Supporting Information). With added piperidine **1m**, Ir(ppy)₃⁺

Table 2. Examining scope with 4-substituted piperidines^a



^{*a*}Isolated yields on 0.2 mmol scale; *dr* was determined by ¹H NMR analysis of the purified product and was within a single percentage point relative to the crude *dr*. ^{*b*}1:1 *dr* prior to isolation determined by crude ¹H NMR analysis. ^{*c*}59:41 *dr* prior to isolation determined by crude ¹H NMR analysis.

is reduced back to Ir(ppy)3 more rapidly, consistent with piperidine oxidation by $Ir(ppy)_{3^+}$ to form the aminium radical cation 7m and Ir(ppy)₃ (Figure 1A, S5 in the Supporting Information). In agreement with this observation, increasing concentration of piperidine 1m leads to larger residual absorbance for the DCB⁻ at 346 nm because the Ir(ppy)₃⁺ that is consumed by piperidine oxidation is unable to recombine with DCB⁻⁻ (Figure 1C, S5 in the Supporting Information). Piperidine oxidation by Ir(ppy)₃⁺ is consistent with the reduction potentials of the two species, $(E_{1/2}(\text{Ir }^{\text{IV/III}}) = 0.77 \text{ V vs.})$ SCE in MeCN) for the Ir(ppy)₃ catalyst,²⁶ versus the aminium cation (E_p = +0.71 V vs. SCE in MeCN for N,N-dimethylaniline).²⁹ From the piperidine concentration dependent reformation of Ir(ppy)₃ observed at 390 nm, we estimate a rate constant for piperidine oxidation $k_{ox.1m}$ = 2.4 x 10⁷ M⁻¹s⁻¹ in DMA (see Figure S6 in the Supporting Information).

Under the conditions of our luminescence quenching and TA measurements, the photoinduced formation of DCB⁻ and the piperidine radical cation seems plausible. We did not observe further radical coupling reactions involving the persistent DCB⁻ by TA or by transient absorption IR experiments, specifically searching for new CN containing photoproducts. We therefore turned to quantum yield measurements to determine the plausibility of a closed-loop photoredox mechanism under the C–H arylation conditions. We used Scaiano's method³⁰ to determine the quantum yield of the reaction between **1a** and DCB. The quantum yield was calculated to be $\Phi = 0.5$ at early conversions, indicating that the photoredox catalyzed cycle is reasonable, although radical chain pathways cannot be ruled out.³¹

Taken together, these findings support the proposed mechanism shown in Figure 1D, which is consistent with MacMillan's original hypothesis.⁴ Ir^{III} is excited by blue light to produce the excited state *Ir^{III}, which transfers an electron to DCB to generate the Ir^{IV} species and the DCB radical anion. ET from the piperidine **1m** to the Ir^{IV} regenerates Ir^{III}. The formed amine radical cation **7m** undergoes deprotonation at the α position by either NaOAc or the piperidine starting material or product, to deliver the α -amino radical **8m**, which can undergo a radical-radical coupling with the persistent DCB radical anion.³² Following coupling to form



Figure 1. Mechanistic studies (A) Reactions studied by TA and luminescence spectroscopies. (B) TA data collected at 390 nm with 10 mM DCB, 40 μM Ir(ppy)₃, and increasing concentrations of **1m**. The laser flash leads to rapid oxidation of the Ir(ppy)³⁺, yielding a large negative absorbance change because the Ir(ppy)₃⁺ species has a smaller ε₃₉₀ than Ir(ppy)₃ (Figure S3). **1m** oxidation by Ir(ppy)₃⁺ is observed as a return in the absorbance at 390 nm as Ir(ppy)₃⁺ is reduced to Ir(ppy)₃. Residual DCB⁻ absorbance was observed at higher concentrations of **1m**. (C) TA spectroscopy at 346 nm to monitor DCB⁻ reactivity. (D) Proposed photoredox cycle.

9m, extrusion of cyanide furnishes the arylated piperidine **2m** and closes the catalytic cycle.

Epimerization studies. We sought to better understand the origins of the high diastereoselectivity observed under the reaction conditions, especially given that our postulated mechanism suggests that the C–C bond forming step is a radical-radical coupling.^{9b} Surprisingly, when monitoring the progress of the reaction of piperidine **1a** and DCB over time, we observed that the initial reaction is not stereoselective (Figure 2). In fact, at 16 min, the ratio of the syn to anti isomer is 50:50 with 75% of the product already formed. Over the course of the remaining 2 h, the anti product isomer **2a-anti** epimerizes to the observed syn diastereomer **2a-syn**, which is ultimately obtained in >95:5 dr.^{11e} Additionally, if the reaction is irradiated for 16 min followed by stirring in the dark for the remaining 2 h, the dr is 56:46 slightly favoring **2a-anti**, indicating that epimerization is a light driven process (see Table S5 in the Supporting Information).

In an effort to elucidate the mechanism of epimerization, we isolated the anti diastereomer **2a-anti** (>95:5 dr) and subjected it to the reaction conditions (Table 3). Notably, we



Figure 2. Time course studies of the C–H arylation of **1a** and follow-up epimerization of **2a**.

obtained only the syn diastereomer **2a-syn** (>95:5 *dr*) in 85% yield after 2 h (Table 3, entry 1). Epimerization was next evaluated upon removing different reaction components. Complete epimerization to the syn diastereomer **2asyn** was still obtained when NaOAc was removed (Table 3, entry 2), indicating that NaOAc does not play a significant role in the epimerization mechanism. When DCB was removed, only 21% of the product epimerized, with the unreacted anti isomer **2a-anti** recovered in 67% yield (Table 3, entry 3). Similar results were obtained when both DCB and NaOAc were removed (Table 3, entry 4). These observations suggest that significant epimerization only occurs under conditions that were shown to produce Ir(ppy)₃⁺ upon irradiation.

To further explore this hypothesis, we postulated that a more oxidizing photocatalyst could directly form the piperidine radical cation by excited state electron transfer, without the need for initial quenching by DCB. To this end, we employed [Ir(dtbbpy)(ppy)₂]PF₆ (dtbbpy = 4,4'-di-*tert*-butyl-2,2'-bipyridine, *E_{1/2}(Ir^{III/II}) = 0.66 V vs. SCE in MeCN), whose excited state reduction potential is significantly more positive than that of Ir(ppy)₃.^{25,33} Indeed, subjecting the anti diastereomer **2a-anti** to [Ir(dtbbpy)(ppy)₂]PF₆ under blue light, and in the absence of DCB, delivers the major diastereomer **2a-syn** as the only detectable product in 72% yield (Table 3, Entry 5). These results are consistent with an epimerization mechanism that proceeds through initial product piperidine oxidation (Table 3).

Based on these collective results, we propose the following mechanism for epimerization under the optimized reaction conditions with $Ir(ppy)_3$ (see Table 3): Excited state quenching by DCB gives potent oxidant $Ir(ppy)_3^+$. ET from **2a-anti** and perhaps also **2a-syn** followed by deprotonation yields α -amino radical **8a**. This intermediate then equilibrates to the **2a-syn** isomer through subsequent reprotonation and ET.

We next investigated whether or not the diastereoselectivity was a result of kinetic control as has been observed for other photoredox-catalyzed isomerization reactions.^{34,10a} Specifically, one possibility for kinetic control would require that the piperidine diastereomers undergo

Table 3. Evaluating epimerization conditions



entry ^a	conditions	% recovered 2a-anti	% yield 2a-syn
1	Standard conditions	<5	85
2	No NaOAc	<5	74
3	No DCB	67	21
4	No NaOAc, no DCB	62	15
5 ^b	No NaOAc, no DCB, [Ir(dtbbpy)(ppy)2]PF6	<5	72

*^a*Yields determined by ¹H NMR spectroscopy with 2,6-dimethoxytoluene as the external standard. *^b*16 h reaction time.

initial ET oxidation at different rates leading to steady-state concentrations of the diastereomers (Figure 3). To evaluate this hypothesis, we measured the rate constants for oxidation of stereoisomerically pure 2a-syn and 2a-anti by Ir(ppy)₃⁺ using TA spectroscopy. To account for the observed >19:1 2a-syn:2a-anti ratio, kinetically controlled selectivity would require that the rate constant for **2a-anti** oxidation $(k_{\text{ox,anti}})$ is at least 19 times larger than the rate constant for **2a-syn** ($k_{ox,syn}$), which would lead to depletion of 2a-anti and enrichment of 2a-syn under our reaction conditions. However, we determined that $k_{\text{ox,anti}}$ and $k_{\text{ox,syn}}$ were the same within error, indicating that the relative rate constants for piperidine oxidation were not responsible for the observed high selectivity. We can conclude that the epimerization is not kinetically controlled by initial oxidation to the aminium radical cations 7a.

We therefore investigated whether or not the epimerization reaction could be under thermodynamic control. The proton transfer between 7a-syn and 7a-anti could generate an equilibrium mixture of diastereomers which controls the observed distribution of 2a diastereomers. Such an equilibrium should be achievable starting with either diastereomer. To probe this possibility, we examined piperidine **2m**, which gave a quantifiable mixture of diastereomers under the reaction conditions (Scheme 3). We subjected both 2m-syn and 2m-anti separately to epimerization with the photocatalyst [Ir(dtbbpy)(ppy)₂]PF₆. After 90 min of irradiation of the syn diastereomer **2m-syn**, a 70:30 (syn:anti) mixture of stereoisomers was obtained in 91% yield. Furthermore, subjecting the anti diastereomer 2manti to the same conditions, gave a 78:22 (syn:anti) mixture of isomers and 94% recovery. Taken together, these results

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Figure 3. A) Oxidation reactions investigated for stereoisomerically pure **2a** diastereomers. B) Plot of observed oxidation rate constant versus **2a** concentration, where the slope is k_{ox} for the respective diastereomers. $k_{ox,anti}$ and $k_{ox,syn}$ are within error of each other.

suggest that the observed diastereoselectivity can be approached from either diastereomer, which is consistent with a thermodynamically controlled process. However, this result does not rule out that the observed selectivity is due to a kinetically controlled steady state.^{10a}

Scheme 3. Equilibration of 2m-syn and 2m-anti diastereomers^a



*^a*Yields determined by ¹H NMR spectroscopy with 2,6-dimethoxytoluene as the external standard.

Further evidence for a thermodynamically controlled epimerization was obtained using DFT calculations to determine the relative stability of various arylated piperidine diastereomers (Table 4). We reasoned that the relative stability of the **2-syn** and **2-anti** diastereomers and the **7-syn** and

7-anti aminium radical cation diastereomers would be similarly affected by the piperidine substituents. The calculated relative free energies of 2-syn and 2-anti piperidine isomers correlate with the observed diastereomer ratios. For the less substituted piperidine **2m**, the **2m-syn** isomer was calculated to be 0.6 kcal/mol lower in energy than the 2manti isomer, consistent with the modest 63:37 syn:anti diastereoselectivity (Table 4, Entry 1). For piperidine 2a, the 2a-anti isomer is calculated to be 1.3 kcal/mol higher in energy than the experimentally obtained 2a-syn isomer (Table 4, Entry 2). This preference is likely due to the 2a-anti isomer having two axial alkyl substituents in the lowest-energy conformation (ethyl at R⁴ and methyl at R²), while the 2a-syn isomer only has one. For piperidine 2e, the difference in energy is more pronounced, with the 2e-syn isomer disfavored by 4.3 kcal/mol (Table 4, Entry 3). In this case, the aryl substituent and the ethyl group at R⁴ are calculated to be axial in the lowest energy conformation of **2e-syn**. Typically, aryl substituents have much higher A values than secondary alkyl carbons,³⁵ likely causing the **2e-anti** isomer to be significantly more favored.

Table 4. Calculated relative energies

entry ^a	substrate	exptl. <i>dr</i> syn/anti	exptl. ΔG_{anti} - ΔG_{syn}	calculated ΔG_{anti} - ΔG_{syn}
1	2m	63:37	0.3 - 0.7	0.6
2	2a	>95:5	>1.8	1.3
3	2e	>5:95	<-1.8	-4.3

^{*a*}Level of theory: ω B97X-D/6-311++G(d,p), CPCM (DMA) // ω B97X-D/6-31G(d), CPCM (DMA). Exptl. = experimental.

CONCLUSIONS

We have described the highly diastereoselective α -amino C–H arylation of densely substituted piperidine derivatives. This study represents a rare example of utilizing photoredox catalysis to promote diastereoselective transformations of complex molecules containing pre-existing stereogenic centers. Key to the generally high selectivity was an epimerization reaction that followed a rapid and non-selective C–H arylation. The observed selectivities for the overall transformation correlate to the calculated relative stabilities of the diastereomers. We anticipate that epimerization processes should be applicable to efficient, diastereoselective syntheses of many different classes of nitrogen heterocycles. Towards this end, efforts are underway in our labs.

ASSOCIATED CONTENT

The Supporting Information is available free of charge on the ACS Publications website at DOI:

Experimental procedures, characterization data, crystallographic data, optical and fluorescence quenching studies, and Cartesian coordinates of all computed structures.

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Notes

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The authors declare no competing financial interest.

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chiral dehydroalanine gave 5-oxazolidinones with high
diastereoselectivity; however, for aminoalkyl radical additions
providing α-branched amines, little to no diastereoselectivity
was observed at the branched site, see: Ref 11d.

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