



# Versatile Open-Source Photoreactor Architecture for Photocatalysis Across the Visible Spectrum

Philip P. Lampkin, Blaise J. Thompson, and Samuel H. Gellman\*

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**ABSTRACT:** Adoption of commercial photoreactors as standards for photocatalysis research could be limited by high cost. We report the development of the Wisconsin Photoreactor Platform (WPP), an opensource photoreactor architecture potentially suitable for general adoption. The WPP integrates inexpensive commercial components and common high-intensity LEDs in a 3D-printed enclosure. Dimensions and features of WPP reactors can be readily varied and configurations easily reproduced. WPP performance is evaluated using literature transformations driven by light of disparate wavelengths.



Modern photocatalytic methods allow for formation of otherwise inaccessible products under mild conditions.<sup>1</sup> This capability has generated significant interest in the field of visible-light photocatalysis<sup>2</sup> and led many laboratories to explore incorporation of photocatalysis into their work.

Photochemical transformations can be significantly affected by small changes in experimental configuration.<sup>3</sup> The number of photons absorbed by the reaction mixture depends, in part, on the intensity and emission profile of the light source as well as the physical surroundings of the reaction vessel.<sup>4</sup> These factors make careful apparatus design and documentation essential for reproducible photoreaction outcomes and reliable reaction discovery.

Many approaches have been used to deliver photons to reaction vessels, but no single, standardized approach has seen widespread adoption to date.<sup>4</sup> Operational variation can hinder the reproduction of reported transformations, the description of which may include only minimal characterization of light source or documentation of experimental setup.<sup>4,5</sup> Entry of new researchers into this field and introduction of photoreactions into the chemistry curriculum should be facilitated by photoreactor platforms that enable accurate reproduction of the apparatus employed in a published study, facilitate apparatus customization for new applications in photocatalysis, and streamline the documentation of apparatus modifications. The work reported here is intended to achieve these goals.

Over the past few years, several commercial photoreactors designed to address problems outlined above have been reported.<sup>3,5,6</sup> More recently, an open-access 3D-printed enclosure for temperature-controlled photoreactions utilizing expensive commercial light-emitting diode (LED) lamps was detailed by Schiel and co-workers.<sup>7</sup> These reactors integrate a

rigid enclosure and exchangeable, high-power photon sources to improve reproducibility and reliability relative to ad hoc experimental setups. However, the adoption of these reactors as standards can be limited by their high cost. This problem is compounded by the significant cost of acquiring multiple proprietary photon sources when different emission profiles are required.<sup>8</sup>

Here, we describe the Wisconsin Photoreactor Platform (WPP), an economical source of high-performance photoreactors that can be easily fabricated, readily modified, and precisely documented (Figure 1). This platform provides reactors constructed from commercial components in a 3Dprinted enclosure. The WPP is designed around inexpensive surface-mount LEDs as the light source. All design files are opensource to maximize transferability. A specific reactor with bespoke physical parameters can be fabricated by an experimentalist with no prior experience in less than a day (instructions in Supporting Information). The WPP is modular and allows photoreactor capabilities to be expanded by use of a well-established array of compatible electronic peripherals. These factors combine to provide a versatile architecture that could contribute to the standardization of photochemical protocols and enhance the discovery and optimization of new photochemical reactivity.

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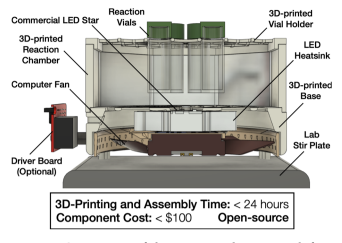
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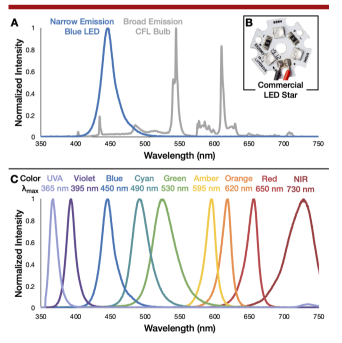
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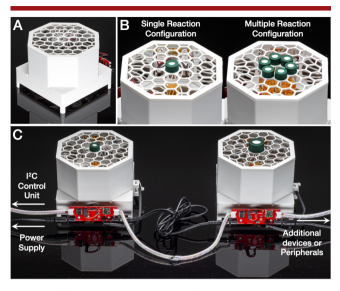
**Figure 1.** Cutaway view of the Wisconsin Photoreactor Platform architecture with labeled components. Estimated component cost is in U.S. dollars as of June 2021.

We viewed the photon source as the most important component of the WPP. An ideal source would be limited to the wavelength range necessary to drive an intended transformation. Sources that emit light in a narrow wavelength range minimize undesired heating and side reactions.<sup>9</sup> Therefore, we were drawn to single-color LEDs rather than broadly emitting light sources, such as compact fluorescent lights (CFL) (Figure 2A). High-intensity LEDs with narrow emission profiles across the visible range are available in inexpensive commercial packages (Figure 2B,C). Access to defined but diverse photon sources is important for new reaction discovery, as demonstrated by recent reports of red and near-infrared (NIR) light photocatalysts.<sup>10,11</sup> Commercial high-intensity LEDs have seen use in several recent studies of photocatalysis.<sup>12</sup>



**Figure 2.** (A) Comparison of emission spectra of 450 nm Cree, Inc. XT-E royal blue LED and ALZO Digital full spectrum CFL bulb. (B) Industry-standard LED star mounted with 450 nm Cree, Inc. XT-E LEDs. (C) Emission spectra of commercial LEDs purchased from manufacturers Cree, Inc., Luxeon, and Inolux.

A 3D-printed enclosure was designed to house a LED star and multiple reaction vials (Figure 3A). As shown in Figure 1, this



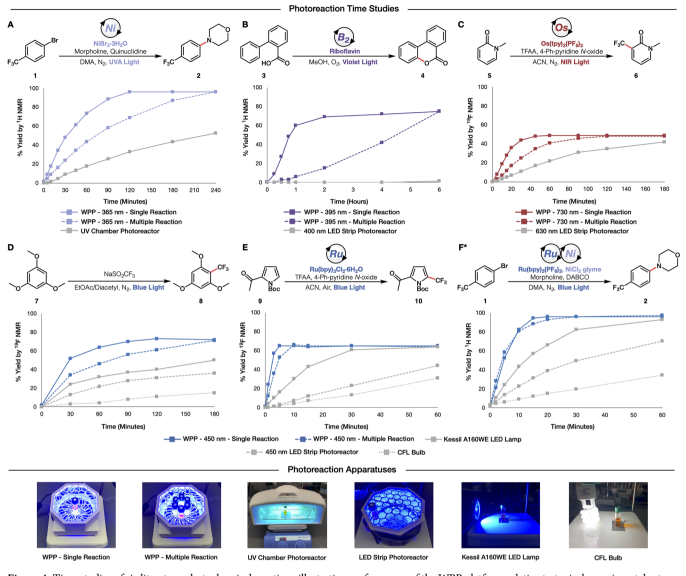
**Figure 3.** (A) Assembled WPP device. (B) Single and multiple reaction configurations of a WPP device fitted with a 4 mL vial holder and reaction chamber module. (C) Illustration of WPP expandability using optional driver boards.

enclosure is composed of a compact base, a reaction chamber, and a vial holder. These three components fit together rigidly, which ensures reproducibility of reaction vessel placement relative to the photon source. Chambers and vial holders with different physical parameters can be designed, printed, and exchanged with one another, which allows the experimentalist to use a variety of vessels. These designs can be documented in research publications to facilitate transferability to other laboratories. We provide printable modules for various common reaction vessels and a light-blocking cover for WPP devices in the Supporting Information.

The vial holders we have designed allow illumination of several vials equally, for rapid screening of experimental variables, or a single vial, for greatest light exposure (Figure 3B). The photoreactor incorporates a commercial aluminum heatsink and low-profile computer fan to cool the LED and reaction vessels. Standard lab stir plates can be used with the small-footprint WPP architecture.

To support kinetics studies or screening of reaction conditions, we designed a custom driver board to control operation of multiple WPP devices simultaneously. Reactors fitted with these boards can be connected in-series to a single power source and control unit (Figure 3C). The control unit can then "supervise" the light intensity and fan speed of each WPP device via I<sup>2</sup>C, a commonly used protocol for communication among digital integrated circuits.<sup>13</sup> If extended functionality is needed, any I<sup>2</sup>C-compatible peripherals could be connected to the driver board for use with the WPP architecture. Many I<sup>2</sup>Ccompatible devices and sensors are commercially available, including thermocouple adapters for temperature monitoring.<sup>1</sup> For work not requiring such control and expandability, a standard 1000 mA LED driver can be used. A simplified driver board offering control over light intensity using only a potentiometer is described in the Supporting Information.

Our intention is to make the WPP architecture suitable as an open standard, with specific reactors easily producible by any pubs.acs.org/OrgLett



**Figure 4.** Time studies of six literature photochemical reactions illustrating performance of the WPP platform relative to typical experimental setups. All reactions were conducted in 4 mL vials with  $\sim$ 1 to 2.5 mL of reaction volume. For exact conditions and procedures, see the Supporting Information. Yields were determined by <sup>1</sup>H or <sup>19</sup>F NMR analysis of crude reaction mixtures using mesitylene or trifluorotoluene as an internal standard. (\*) The "WPP – 450 nm – Single Reaction" benchmark was conducted in a 4 °C refrigerator at 90% light intensity.

chemist. Toward that end, we have included our enclosure and electronics design files as well as detailed component sourcing and fabrication instructions in the Supporting Information. A "living" online repository to which users can contribute custom WPP modules is provided.<sup>15</sup> 3D printers and 3D-printing services are readily available. We have found that the enclosure can be fabricated using a variety of affordable 3D printers.

Commercial small-scale printed circuit board fabrication is inexpensive and widely available. Our driver boards can be assembled quickly without specialized tools. These features should encourage adoption of the WPP.

To establish the versatility and reliability of the WPP architecture relative to typical experimental setups, we applied this approach to six reactions from the recent photocatalysis literature that are driven by light of disparate wavelengths. Photoreaction apparatuses used for comparison were derived from setups reported in the literature (see Supporting Information). WPP device performance across both the single reaction and multiple reaction configurations shown in Figure 3B was evaluated.

In presenting the results in Figures 4 and 5, we do not mean to imply that the WPP is the best way to conduct any particular photochemical reaction. Instead, we offer these data to suggest that use of the WPP platform is likely to provide good results for a wide range of photochemical transformations. In addition, because the specifics of a WPP device can be easily varied and documented, results achieved with this approach can be readily reproduced in other laboratories. These features may be particularly useful for researchers seeking to initiate studies in this important field.

The first "benchmark" process was the C–N cross-coupling of 4-bromobenzotrifluoride and morpholine via the photoexcitation of nickel-amine complexes with ultraviolet A (UVA) light, as described by Miyake et al. (Figure 4A).<sup>12a</sup> In both single- and multiple-reaction configurations, a WPP device fitted with a 365 nm LED star decreased reaction time relative to a more conventional apparatus fashioned from a commercial UV light

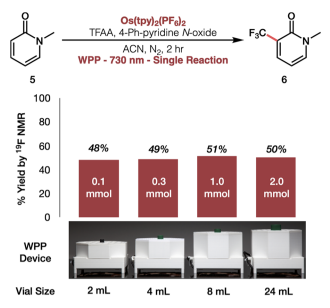


Figure 5. Results of scale-up trials. Reactions conducted at indicated scales using listed reaction vessel sizes over 2 hours. Yields were determined by <sup>19</sup>F NMR analysis of crude reaction mixtures using trifluorotoluene as an internal standard.

curing chamber.<sup>16</sup> With the WPP approach, reaction completion was reached in 2 or 4 h with 96% yield for the single- or multiple-reaction configuration, respectively, reproducing the yield reported by Miyake. The UV chamber achieved only 53% yield after 4 h of illumination.

Synthesis of 3,4-benzocoumarin using violet light and the vitamin photocatalysis strategy detailed by Gilmour et al. was examined next (Figure 4B).<sup>126</sup> Using a standard 400 nm LED strip reactor, based on the design of Stephenson et al.,<sup>17</sup> we obtained <5% yield after 8 h of continuous illumination. In contrast, a WPP device with a 395 nm light source provided 75% yield across single- and multiple-reaction configurations after 8 h, matching the literature yield and indicating the robust performance of the WPP architecture in this transformation. The single-reaction configuration achieved 72% yield after only 2 h.

Inspired by the recent report of Rovis et al. on the development of NIR photocatalysts,<sup>11</sup> we examined their Osphotocatalyzed trifluoromethylation of *N*-methyl-2-pyridone using a WPP device with a 730 nm LED star (Figure 4C). Both WPP configurations provided faster product formation than did a 630 nm LED strip photoreactor, despite stronger absorption of the Os(tpy)<sub>2</sub>(PF<sub>6</sub>)<sub>2</sub> photocatalyst at 630 nm.<sup>11</sup> The trifluoromethylated product was generated in 48% yield with both WPP configurations after 2 h of illumination, while the LED strip reactor provided 35% yield. The yield obtained using a WPP device closely agreed with the yield reported by Rovis and coworkers.<sup>11</sup>

We note that the comparison setups we used for the reactions in Figure 4A–C are not the specialized setups described in the original reports. Our goal in this work is to show that the WPP is effective across a range of photoreactions rather than to determine whether this architecture provides the best possible way to conduct any particular photoreaction.

We explored reactor efficacy with three blue light photoreactions previously used by MacMillan et al. to benchmark a commercial photoreactor.<sup>3</sup> We first tested our platform's performance in catalyzing the trifluoromethylation of 1,3,5trimethoxybenzene using the method reported by Li and coworkers (Figure 4D).<sup>18</sup> A WPP device with 450 nm LEDs yielded product faster and in higher yield than did conventional experimental setups involving a Kessil A160WE LED lamp, a 450 nm LED strip reactor, or a CFL bulb. Six reactions of this type simultaneously carried out using a WPP device in the multiple reaction configuration exhibited a consistent reaction profile, indicating the reliability of the WPP architecture (Figure S22). The same trend was observed when the protocol of Stephenson et al. for trifluoromethylation of 2-acetyl-N-Bocpyrrole was evaluated (Figure 4E).<sup>19</sup> Finally, the metallophotoredox-catalyzed C-N cross-coupling of 4-bromobenzotrifluoride and morpholine of MacMillan et al. revealed a similar trend (Figure 4F).<sup>20</sup> Across all three blue-light reactions, both WPP configurations offered superior performance relative to the conventional experimental setups tested and similar performance to that reported for the photoreactor described by MacMillan and co-workers (Figures S24-S26). In all cases, yields obtained using a WPP device matched those achieved with a commercial photoreactor.

As an additional test of architecture generality, we evaluated WPP performance across a series of typical laboratory reaction scales (Figure 5). Using the photocatalytic trifluoromethylation of *N*-methyl-2-pyridone as a testbed, we conducted reactions at 0.1, 0.3, 1.0, and 2.0 mmol scale in a WPP device fitted with a 730 nm LED star. Enclosure modules for each scale were designed to standardize reaction vessel placement across all trials. In all cases, product was generated in ~50% yield after 2 h. Increases in reaction scale up to 20-fold relative to the initial 0.1 mmol trial reaction were accommodated without loss of performance.

We have developed an open-source photoreactor platform that enables rapid progress in cutting-edge photocatalysis research and has a low barrier for adoption. The Wisconsin Photoreactor Platform is inexpensive, adaptable, and highly featured; therefore, the WPP should foster standardized experimental protocols and reproducibility. We have demonstrated the favorable performance of WPP devices across a series of benchmark photoreactions using UVA, violet, NIR, and blue light as well as in reaction scale-up. The WPP architecture meets the need for a standardized approach to experimental apparatus that is versatile, reliable, can be precisely documented and easily reproduced, and is economically accessible to a broad community of researchers.

# ASSOCIATED CONTENT

#### **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.orglett.1c01910.

- Photon source characterization data, experimental details, and NMR spectra (PDF)
- Wisconsin Photoreactor Platform apparatus fabrication and operation instructions (PDF)
- Wisconsin Photoreactor Platform project repository at time of publication (ZIP)

# AUTHOR INFORMATION

#### **Corresponding Author**

Samuel H. Gellman – Department of Chemistry, University of Wisconsin, Madison, Wisconsin 53706, United States; orcid.org/0000-0001-5617-0058; Email: gellman@ chem.wisc.edu

#### Authors

- Philip P. Lampkin Department of Chemistry, University of Wisconsin, Madison, Wisconsin 53706, United States;
   orcid.org/0000-0002-6908-9619
- Blaise J. Thompson Department of Chemistry, University of Wisconsin, Madison, Wisconsin 53706, United States; orcid.org/0000-0002-3845-824X

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.orglett.1c01910

## Notes

The authors declare no competing financial interest.

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## REFERENCES

(1) (a) Skubi, K. L.; Blum, T. R.; Yoon, T. P. Dual Catalysis Strategies in Photochemical Synthesis. *Chem. Rev.* 2016, *116* (17), 10035–10074.
(b) Yoon, T. P. Photochemical Stereocontrol Using Tandem Photoredox-Chiral Lewis Acid Catalysis. *Acc. Chem. Res.* 2016, *49* (10), 2307–2315. (c) Romero, N. A.; Nicewicz, D. A. Organic Photoredox Catalysis. *Chem. Rev.* 2016, *116*, 10075–10166. (d) Prier, C. K.; Rankic, D. A.; MacMillan, D. W. C. Visible Light Photoredox Catalysis with Transition Metal Complexes: Applications in Organic Synthesis. *Chem. Rev.* 2013, *113* (7), 5322–5363. (e) Strieth-Kalthoff, F.; James, M. J.; Teders, M.; Pitzer, L.; Glorius, F. Energy Transfer Catalysis Mediated by Visible Light: Principles, Applications, Directions. *Chem. Soc. Rev.* 2018, *47* (19), 7190–7202.

(2) Shaw, M. H.; Twilton, J.; MacMillan, D. W. C. Photoredox Catalysis in Organic Chemistry. J. Org. Chem. 2016, 81 (16), 6898–6926.

(3) Le, C. C.; Wismer, M. K.; Shi, Z. C.; Zhang, R.; Conway, D. V.; Li, G.; Vachal, P.; Davies, I. W.; MacMillan, D. W. C. A General Small-Scale Reactor to Enable Standardization and Acceleration of Photocatalytic Reactions. *ACS Cent. Sci.* **2017**, *3* (6), 647–653.

(4) Bonfield, H. E.; Knauber, T.; Lévesque, F.; Moschetta, E. G.; Susanne, F.; Edwards, L. J. Photons as a 21st Century Reagent. *Nat. Commun.* **2020**, *11*, 804.

(5) Bonfield, H. E.; Mercer, K.; Diaz-Rodriguez, A.; Cook, G. C.; McKay, B. S. J.; Slade, P.; Taylor, G. M.; Ooi, W. X.; Williams, J. D.; Roberts, J. P. M.; Murphy, J. A.; Schmermund, L.; Kroutil, W.; Mielke, T.; Cartwright, J.; Grogan, G.; Edwards, L. J. The Right Light: De Novo Design of a Robust Modular Photochemical Reactor for Optimum Batch and Flow Chemistry. *ChemPhotoChem.* **2020**, *4* (1), 45–51.

(6) Bissonnette, N. B.; Ryu, K. A.; Reyes-Robles, T.; Wilhelm, S.; Tomlinson, J. H.; Crotty, K. A.; Hett, E. C.; Roberts, L. R.; Hazuda, D. J.; Jared Willis, M.; Oslund, R. C.; Fadeyi, O. O. Design of a Multiuse Photoreactor To Enable Visible-Light Photocatalytic Chemical Transformations and Labeling in Live Cells. *ChemBioChem* **2020**, *21* (24), 3555–3562.

(7) Schiel, F.; Peinsipp, C.; Kornigg, S.; Böse, D. A 3D-Printed Open Access Photoreactor Designed for Versatile Applications in Photoredox- and Photoelectrochemical Synthesis. *ChemPhotoChem.* 2021, 5 (5), 431–437.

(8) The Penn PhD Photoreactor MK 2, the precursor of which was reported in ref 3, is available with a 450 nm light source from Sigma Aldrich Millipore for \$4910 USD as of 2021. Photon sources emitting 420 or 365 nm light are sold separately for \$752 USD each. See: https://www.sigmaaldrich.com/catalog/product/sial/z744035.

(9) Protti, S.; Ravelli, D.; Fagnoni, M. Wavelength Dependence and Wavelength Selectivity in Photochemical Reactions. *Photochem. Photobiol. Sci.* **2019**, *18* (9), 2094–2101.

(10) (a) Mei, L.; Veleta, J. M.; Gianetti, T. L. Helical Carbenium Ion: A Versatile Organic Photoredox Catalyst for Red-Light-Mediated Reactions. J. Am. Chem. Soc. 2020, 142 (28), 12056–12061. (b) Bilger, J. B.; Kerzig, C.; Larsen, C. B.; Wenger, O. S. A Photorobust Mo(0) Complex Mimicking [Os(2,2'-Bipyridine)3]2+and Its Application in Red-to-Blue Upconversion. J. Am. Chem. Soc. 2021, 143 (3), 1651– 1663.

(11) Ravetz, B. D.; Tay, N. E. S.; Joe, C. L.; Sezen-Edmonds, M.; Schmidt, M. A.; Tan, Y.; Janey, J. M.; Eastgate, M. D.; Rovis, T. Development of a Platform for Near-Infrared Photoredox Catalysis. *ACS Cent. Sci.* **2020**, *6* (11), 2053–2059.

(12) The following references describe using high-intensity commercial LEDs in custom experimental setups for photocatalysis research: (a) Lim, C. H.; Kudisch, M.; Liu, B.; Miyake, G. M. C-N Cross-Coupling via Photoexcitation of Nickel-Amine Complexes. J. Am. Chem. Soc. 2018, 140 (24), 7667–7673. (b) Morack, T.; Metternich, J. B.; Gilmour, R. Vitamin Catalysis: Direct, Photocatalytic Synthesis of Benzocoumarins via (-)-Riboflavin-Mediated Electron Transfer. Org. Lett. 2018, 20 (5), 1316–1319. (c) Bosque, I.; Bach, T. 3-Acetoxyquinuclidine as Catalyst in Electron Donor-Acceptor Complex-Mediated Reactions Triggered by Visible Light. ACS Catal. 2019, 9 (10), 9103–9109. (d) Elliott, L. D.; Kayal, S.; George, M. W.; Booker-Milburn, K. Rational Design of Triplet Sensitizers for the Transfer of Excited State Photochemistry from UV to Visible. J. Am. Chem. Soc. 2020, 142 (35), 14947–14956.

(13) Prabhu, G. R. D.; Urban, P. L. Elevating Chemistry Research with a Modern Electronics Toolkit. *Chem. Rev.* **2020**, *120* (17), 9482–9553.

(14) I<sup>2</sup>C-compatible peripherals are available from vendors:
(a) https://www.sparkfun.com/.
(b) https://www.adafruit.com.
(c) https://www.seeedstudio.com/.

(15) Users of the WPP can contribute custom modules, electronics and other designs to the WPP repository for others to use. See https://github.com/uw-madison-chem-shops/wisconsin-photoreactor.

(16) Aung, T.; Liberko, C. A. Bringing Photochemistry to the Masses: A Simple, Effective, and Inexpensive Photoreactor, Right out of the Box. J. Chem. Educ. 2014, 91 (6), 939–942.

(17) Douglas, J. J.; Albright, H.; Sevrin, M. J.; Cole, K. P.; Stephenson, C. R. J. A Visible-Light-Mediated Radical Smiles Rearrangement and Its Application to the Synthesis of a Difluoro-Substituted Spirocyclic ORL-1 Antagonist. *Angew. Chem., Int. Ed.* **2015**, *54* (49), 14898–14902.

(18) Li, L.; Mu, X.; Liu, W.; Wang, Y.; Mi, Z.; Li, C. J. Simple and Clean Photoinduced Aromatic Trifluoromethylation Reaction. J. Am. Chem. Soc. 2016, 138 (18), 5809–5812.

(19) Beatty, J. W.; Douglas, J. J.; Miller, R.; McAtee, R. C.; Cole, K. P.; Stephenson, C. R. J. Photochemical Perfluoroalkylation with Pyridine N-Oxides: Mechanistic Insights and Performance on a Kilogram Scale. *Chem.* **2016**, *1* (3), 456–472.

(20) Corcoran, E. B.; Pirnot, M. T.; Lin, S.; Dreher, S. D.; Dirocco, D. A.; Davies, I. W.; Buchwald, S. L.; Macmillan, D. W. C. Aryl Amination Using Ligand-Free Ni(II) Salts and Photoredox Catalysis. *Science* **2016**, 353 (6296), 279–283.