

Harnessing the Substrate Promiscuity of Dioxygenase AsqJ and Developing Efficient Chemoenzymatic Synthesis for Quinolones

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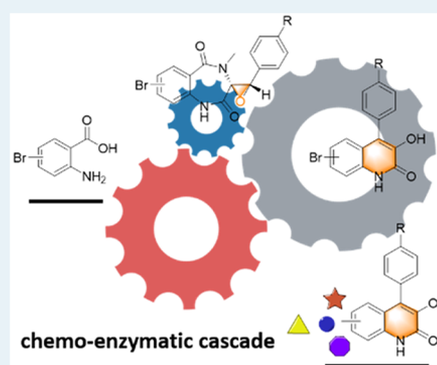
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Supporting Information

ABSTRACT: Nature has developed complexity-generating reactions within natural product biosynthetic pathways. However, direct utilization of these pathways to prepare compound libraries remains challenging because of limited substrate scopes, involvement of multiple-step reactions, and moderate robustness of these sophisticated enzymatic transformations. Synthetic chemistry offers an alternative approach to prepare natural product analogues. However, because of complex and diverse functional groups appended on the targeted molecules, dedicated design and development of synthetic strategies are typically required. Herein, by leveraging the power of chemoenzymatic synthesis, we report an approach to bridge the gap between biological and synthetic strategies in the preparation of quinolone alkaloid analogues. Leading by *in silico* analysis, the predicted substrate analogues were chemically synthesized. The AsqJ-catalyzed asymmetric epoxidation of these substrate analogues was followed by a Lewis acid-triggered ring contraction to complete the viridicatin formation. We evaluated the robustness of this method in gram-scale reactions. Lastly, through chemoenzymatic cascades, a library of quinolone alkaloids is effectively prepared.

KEYWORDS: Fe/2OG enzyme, AsqJ, chemoenzymatic synthesis, rearrangement, quinolone



Quinoline and quinolone alkaloids are secondary metabolites produced by several organisms with broad biological properties including antiallergic, antibacterial, antitumor, and antiviral activities.^{1–4} In both chemical and biological syntheses, quinoline and quinolone core serve as key intermediates in the preparation of these bioactive molecules.⁵ Among them, the skeleton of 3-hydroxyquinolin-2(1H)-one, viridicatin, has been recognized as a valuable scaffold in numerous natural products as well as synthetic compounds.^{3,6,7} Quite a few synthetic strategies have been developed for the construction of this skeleton including ring expansion of isatin and aryldiazomethane, Knoevenagel reaction followed by cyclization, condensation using 2-aminobenzaldehyde with chloroacetic anhydride or *N*-phenylacetoacetamide derivative, transition metal-catalyzed cross-coupling, and ring contraction of cyclohexenone.^{8–14} In nature, viridicatin is biosynthesized from *O*-methyl-L-tyrosine/phenylalanine and anthranilic acid through a series of enzymatic transformations involving condensation, methylation, desaturation, epoxidation, and rearrangement.¹⁵ Although the aforementioned strategies have their merits, the limitations of these methods to generate viridicatin analogues led us to consider developing an efficient method combining the tools employed by chemistry and nature.

AsqK, a methyl transferase domain-containing NRPS (nonribosomal peptide synthetase), catalyzes the cyclization

and methylation to form (4'-methoxy)cyclohexenone. An iron and 2-oxoglutarate (Fe/2OG) dependent oxygenase, AsqJ, catalyzes sequential desaturation and asymmetric epoxidation to furnish (4'-methoxy)cyclohexenone (2) via (4'-methoxy)-dehydrocyclohexenone (1).¹⁵ Finally, cyclohexenase, AsqI, triggers ring contraction to complete the biosynthesis of (4'-methoxy)-viridicatin (3) (Figure 1A).¹⁶ In comparison with the enzymatic synthesis, 1 and 5 can be chemically prepared through Perkin condensation (Figure 1B).⁹ However, chemical asymmetric epoxidation of 1 remains a synthetically challenging task. Herein, we demonstrated that utilization of AsqJ catalyzed epoxidation represents an attractive approach to prepare cyclohexenone analogues (6), particularly as the epoxidation requires only molecular oxygen and 2-oxoglutarate to enable the catalysis under mild conditions and generates succinate and CO₂ as coproducts. Using *in silico* analysis, we demonstrated that a Lewis acid can effectively convert cyclohexenone to viridicatin (2 → 3 and 6 → 7), which is analogous to the cyclohexenase-catalyzed reaction where a zinc

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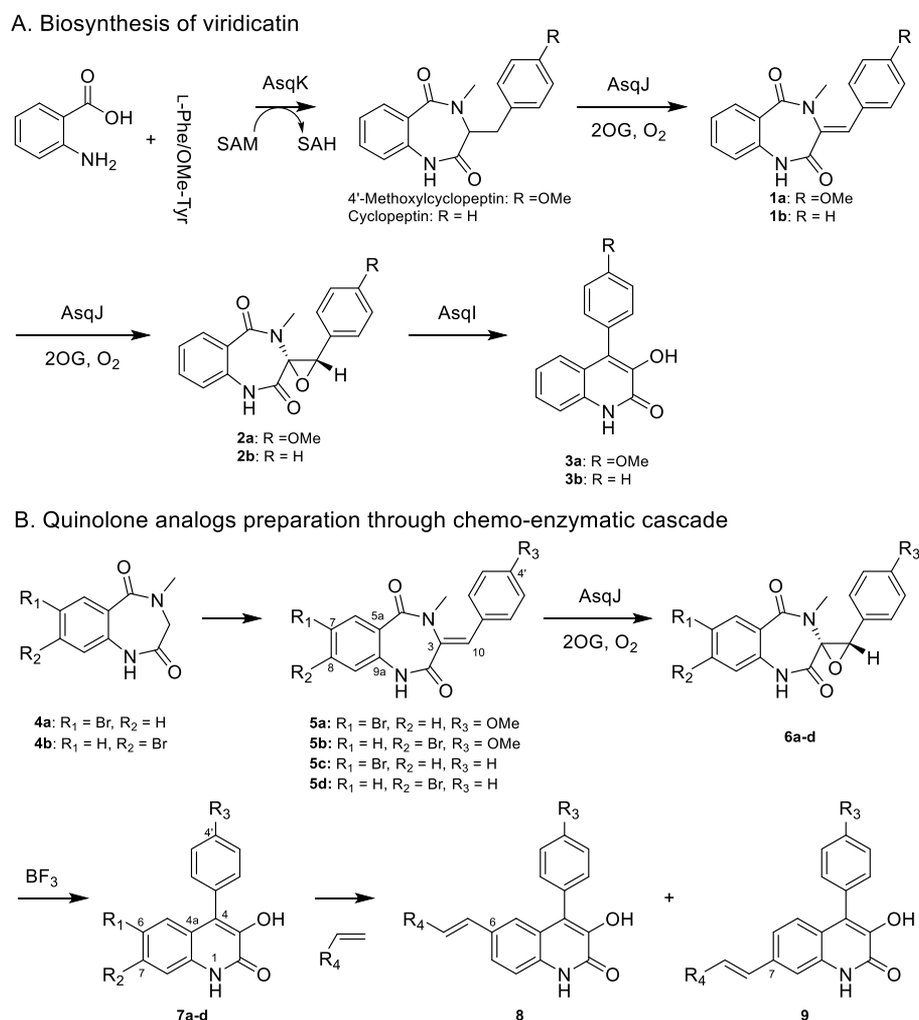


Figure 1. (A) Biosynthetic pathway of viridicatin and (B) preparation of quinolone analogs through chemoenzymatic transformations.

ion located at the active site triggers the rearrangement. Lastly, a cross-coupling reaction using halogenated viridicatin (7a–7d) yields a quinolone library (Figure 1B).

To harness the advantages provided by AsqJ, challenges including substrate scope, selectivity, and effectiveness toward substrate analogues have to be addressed. By inspecting the previously published crystallographic data on AsqJ,^{17–20} we found that the binding of 1 and analogues (5) are highly conserved. The two carbonyl oxygens from the diazepine-2,5-dione moiety of the substrate participate in hydrogen bonding interactions with the N atom of Asn70 and the backbone N atom of Met137, respectively. The aryl ring of diazepine-2,5-dione has hydrophobic interactions with Leu79, Met118, Met122, and Thr227. These four residues are part of the double stranded β -helix structure commonly present in Fe/2OG enzymes. In addition, the phenyl moiety has a π – π interaction with His134 and is also stabilized by hydrophobic interactions with Val72, Phe139, and Pro132 (Figure 2). To efficiently build a quinolone library through cross-coupling reactions, we consider brominated viridicatin (7) as our targeted molecules. We performed molecular dynamics (MD) simulations to assess the binding and potential steric effects. As revealed by 10 ns MD trajectory simulations carried out using 1a, 1b, and 5a–5d, all compounds exhibit a similar binding configuration in the MD equilibrated structures, which closely resembles those in the AsqJ crystal structures (Figures 2 and

S83–S86). In addition, calculations using alchemical ligand transformation^{21,22} and the Bennett Acceptance Ratio method^{23,24} suggest that the differences between the binding free energies of all compounds are relatively small. Compared with the native substrates (1a and 1b), the binding energies for 5b and 5d are 4.7 and 2.3 kcal/mol higher. On the other hand, the binding energies for 5a and 5c are lowered by 1.4 kcal/mol. More importantly, the distances between the iron center and the C10 of all compounds are kept in a narrow range of ~4.0–5.5 Å (Figure 2). According to the recent study of the AsqJ catalyzed epoxidation,²⁰ the iron-oxo of the ferryl intermediate reacts with the olefin moiety at the C10 position of the substrate. Thus, the Fe–C10 distances predicted by the MD simulations strongly suggest that 5a–5d would be suitable for AsqJ catalysis.

Given the potential substrate promiscuity assessed through *in silico* analysis, we began by synthesizing the halogenated dehydrocyclopeptin analogues (5a–5d, see Supporting Information (SI) for details). In parallel, AsqJ was heterologously expressed in *Escherichia coli* and purified using metal-affinity chromatography. Analytical-scale reactions with each substrate were carried out. Analogous to the native substrates, complete substrate consumption of 5c and 5d was detected in 4 h (Table 1, entries 5 and 6). Moderate conversions (~71–75%) were observed in the reactions using 5a and 5b (entries 3 and 4).

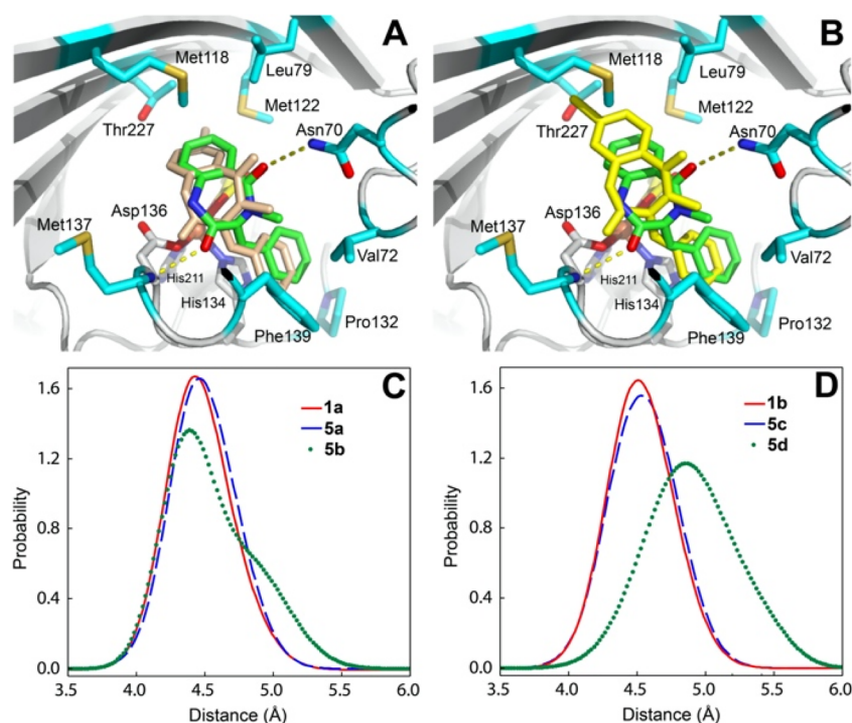


Figure 2. Top: Overlap of **1b** (green) in the AsqJ active site (PDB: 6KOE) with **5c** (wheat) (Panel A) or with **5d** (yellow) (Panel B) obtained from the MD equilibrated structures. The protein residues surrounding **1b** are highlighted. Bottom: The Fe–C10 distance distribution among **1a**, **5a**, and **5b** (Panel C), and among **1b**, **5c**, and **5d** (Panel D) based on MD simulations.

Table 1. Substrate Consumption Monitored by LC-MS

entry	substrate	4 h (%)	20 h (%)
1	1a	90	100
2	1b	100	100
3	5a	75	93
4	5b	71	92
5	5c	100	100
6	5d	96	97

For all substrates, longer reaction time led to almost complete substrate consumption.

Next, we assessed the reaction outcome and scalability of this biotransformation. Although AsqJ delivers **2a** and **2b** as the sole products, it is possible that non-native substrates may tilt the reaction outcomes. Fe/2OG enzymes have been reported to produce alternative products when substrate analogues are used. For example, in NapI and SyrB2, the reaction outcomes change from native desaturation and halogenation to hydroxylation.^{25,26} Therefore, we optimized the reaction conditions to achieve ~1000 turnovers for **5a**, **5c**, and **5d** and 250 turnovers for **5b**. Currently, the reason for the lower turnover using **5b** is unknown. In the scaled-up reactions, the loading of enzyme is at 0.1–0.4% molar ratio to the substrate. All reaction products are purified and characterized by detailed NMR analysis (Figure 3 and SI for details). As revealed by ¹H and ¹³C NMR, **6a**–**6d** display similar resonances as those of **2a** and **2b**. The up-shifting of the ¹³C signal (C7 or C8) is due to the introduction of the bromine. Based on HMBC and ¹H–¹H COSY correlations, the structures of **6a**–**6d** are determined. Furthermore, on the basis of the X-ray structure of **6c**, the newly installed chiral centers at C3 and C10 have *R* and *S* configurations (Figure 3). These results are consistent with the substrate binding configuration

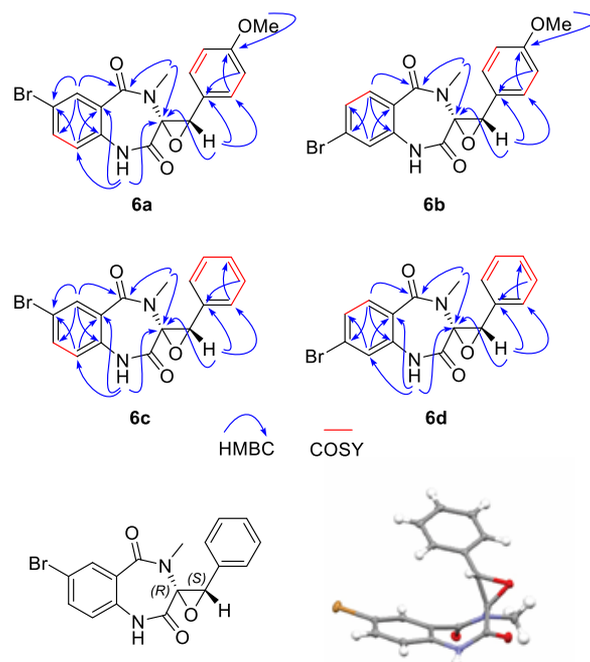


Figure 3. Structural characterization by selective correlations of HMBC and ¹H–¹H COSY and the X-ray structure of **6c**.

observed in the AsqJ structures and the aforementioned MD simulation predictions where the oxygen atom transfer by the ferryl intermediate is stereospecific.

Additionally, we determined the steady-state kinetics of the AsqJ catalyzed reactions using **1a**, **1b**, **5a**–**5d** (Figure S24 and Table S2). Comparing the steady-state kinetics of the two native substrates, **1b** has higher catalytic efficiency (k_{cat}/K_m of

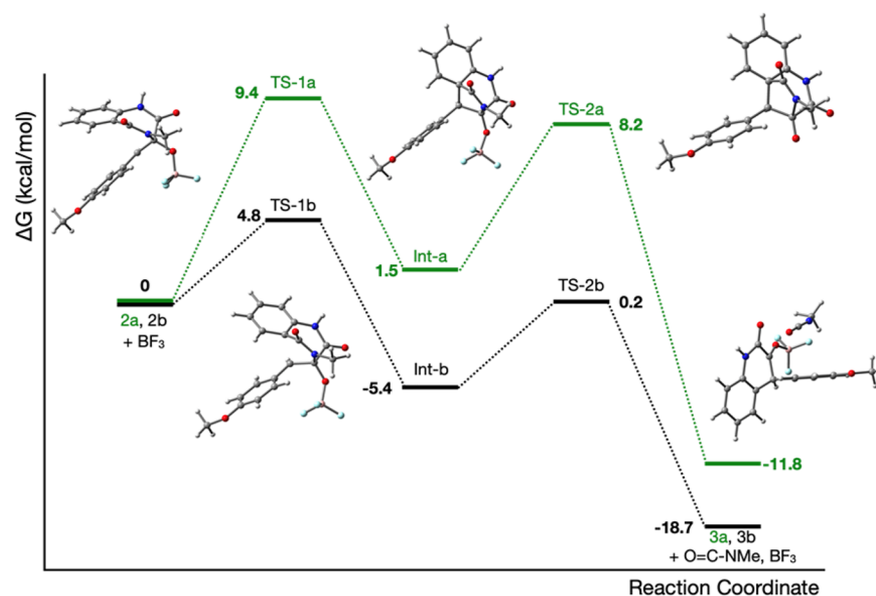


Figure 4. Reaction coordinate for the conversion of **2a** (green) and **2b** (black) to **3a** and **3b** in the presence of BF_3 . The numerical values indicate the calculated free energy differences of various species. The molecular structures of reactant, intermediate, product, and transition states of **2a** \rightarrow **3a** are shown.

1b/1a is ~ 5). When analogues (**5a–5d**) are used, K_m values are increased by 5–10 times. Among **5a–5d**, the K_m values of 7-brominated analogues (**5a** and **5c**) are ~ 2 times smaller than 8-brominated analogues (**5b** and **5d**). The k_{cat} values of **5a** and **5c** are similar or even slightly higher than those of the native substrates. Thus, the catalytic efficiency is not perturbed. However, for **5b** and **5d**, the overall enzyme efficiency is compromised. Thus, Br-substituent at 7 position (**5a** and **5c**) has minimal influence on AsqJ efficiency, whereas C8-bromination (**5b** and **5d**) reduces the efficiency by ~ 10 –100 fold. These results are consistent with the MD results where the binding affinity of **5b** and **5d** are ~ 4.7 and 2.3 kcal/mol higher than native substrates. Taken together, these observations demonstrate that AsqJ is flexible toward the 7- and 8-substitutions and is capable of catalyzing effective asymmetric epoxidation, which is compared favorably to the reported chemical epoxidation of dehydrocycloheptin.¹⁰

Toward preparation of the quinolone library, our next goal is to develop an efficient method to convert cyclophenin to viridicatin (**2** \rightarrow **3** and **6** \rightarrow **7**). In the reported AsqJ catalysis, the active site Zn ion serves as a Lewis acid to activate the epoxide and to facilitate an anti-Baldwin-type epoxide-opening 6-*endo-tet* cyclization.¹⁶ Followed by the elimination of methyl isocyanate, a keto–enol tautomerization leads to viridicatin formation (Figure S93). On the other hand, Groll et al. and Gulder et al. reported that the rearrangement is spontaneous once cyclophenin (**2**) is released from the active site of AsqJ.^{17,27} We first assess the plausible rearrangement pathways through *in silico* analysis. The DFT calculated reaction coordinate using **2a** and **2b** largely follows the above-mentioned mechanism (see the SI for the details). The C–C bond formation between C10 and C5a accompanied by epoxide ring opening generates a cyclized intermediate. Subsequently, elimination of a methylisocyanate and keto–enol tautomerization completes the reaction. Interestingly, in both aqueous and organic conditions (calculated by using the Polarizable Continuum Model),²⁸ **2a** and **2b** alone are unlikely to rearrange to **3a** and **3b** spontaneously. The calculated free

energy of the transition state connecting **2a/2b** to the cyclized intermediate is ~ 25 – 30 kcal/mol (ΔG^\ddagger) higher than the substrate state (Figure S87, S88). This result is consistent with the DFT calculations reported by Borowski et al.²⁹ and is less consistent with the reported spontaneous rearrangement phenomenon. To explore the possibility of using an Lewis acid as a potential catalyst, we calculated the effect of Lewis acids on the rearrangement reaction. The DFT calculations predict that a Lewis acid, such as BF_3 , would strongly interact with the epoxide oxygen of cyclophenin to enable epoxide ring opening in organic solvent, such as dichloromethane (Figure 4). Release of the ring strain of the epoxide leads to a better π – π stacking interaction between the two aryl rings of the substrate, which in turn significantly reduces the distance between C10 and C5a from ~ 3.3 to ~ 2.7 Å. As a consequence, the BF_3 bound reactant is structurally similar to the transition state structure (the C10 and C5a distance is ~ 2.0 Å) and dramatically reduces the barrier (ΔG^\ddagger) of the C10–C5a bond formation from ~ 25 – 30 to ~ 5 kcal/mol for **2b** (Figure 4). For the subsequent methylisocyanate elimination step, the reaction barrier is ~ 6 kcal/mol. Compared with **2b**, the methoxy group of **2a** seems to increase the transition state barriers for both steps to ~ 8 – 9 kcal/mol. Besides, other commonly used Lewis acids, such as AlCl_3 , show a similar effect (Figure S89) and brominated analogues (**6a–6d**) do not affect the overall reaction coordinate and energetics (Figures S90, S91). It is worthwhile to mention that **2b** obtained in the AsqJ crystal structure and **6c** have very different three-dimensional conformations (Figure S92). Because of the geometric constraints enforced by AsqJ, only the flat, or chairlike, form is retained in the active site. The folded, or boat-like, form becomes a dominant species once the epoxide is released from the active site of AsqJ. Our DFT calculations suggest that the folded conformation is energetically favorable by ~ 2 kcal/mol for all the compounds used in this study (Table S3).

We then set out to test the rearrangement reaction by monitoring the conversion of **2a** and **2b** using ^1H NMR. Within 120 min, no obvious substrate consumption can be

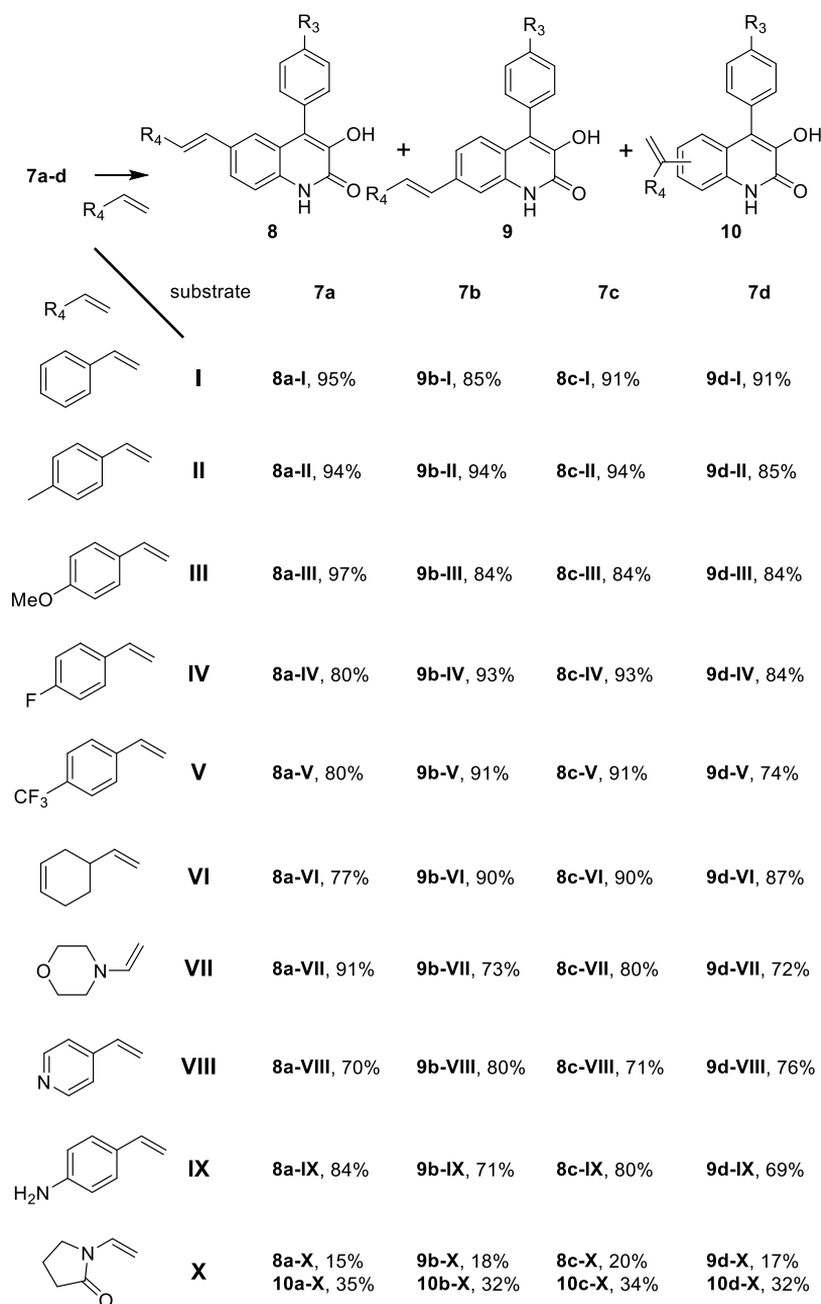


Figure 5. A library of quinolone analogues is prepared via chemoenzymatic reactions. The products synthesized using XI are shown in the SI (Figure S94).

observed under pH 5–8.5 (Figures S25, S26). These observations indicate the need of an acid for efficient rearrangement. To our delight, when using BF_3 in dichloromethane, through a thin-layer-chromatographic analysis, we observed a complete consumption of the substrates (2a and 2b) and the formation of (4'-methoxy)viridicatin (3a and 3b). Applying this method also affords brominated viridicatin analogues (7a–7d). Although the origin of anti-Baldwin rearrangement remains to be carefully investigated, these results highlight the advantage of our approach which allows for the effective preparation of viridicatin through tandem enzymatic and chemical reactions.

While enzymatic reactions are typically carried out on submilligram to milligram quantities to identify and characterize the reaction products, a potential challenge of preparative-

scale reactions can serve as a bottleneck to the enzymatic transformations being embraced for synthetic applications. We envisioned transitioning from sequential reactions/purification to a scalable platform could benefit synthetic usefulness. Toward this goal, reactions using AsqJ-containing lysate were investigated. Although complete consumption of the substrate was achieved, the components originated from the crude lysate severely affect the rearrangement efficiency. Taking advantage of robust expression and stability of AsqJ (~38 mg protein per liter of culture), the scale-up reactions were conducted using the purified enzyme. After AsqJ completed epoxidation, the crude product obtained via a simple ethyl acetate extraction was subjected to BF_3 catalyzed rearrangement to afford 7a–7d in gram scale with an overall yield of 57–91%.

To build quinolone library, we sought to leverage the reactivity of brominated viridicatin via cross-coupling reactions. We anticipated that a series of compounds can be accessed in one-step by reacting with cross-coupling partners. The 6-substituted vinyl quinolone skeleton is found in aflaquilonones, aspoquinolones, 4-phenyl-3,4-dihydroquinolones, and yaequinolones.^{30–33} However, the naturally occurring 7-vinyl substituted quinolones are rare. Herein, we explored the synthesis of a library of 6- and 7-vinyl substituted quinolones. To access the 6-substituted vinyl quinolones, **7a** and **7c** were reacted with substituted alkenes under Heck reaction conditions³⁴ to afford the desired product in reasonable yields (Figure 5 and Figure S94, **8a**-(I–XI), **8c**-(I–X)). Analogously, the 7-substituted vinyl quinolones were prepared using **7b** and **7d** as the substrates (**9b**-(I–XI), **9d**-(I–X)). It is worthwhile to mention that using 1-vinylpyrrolidin-2-one and D-limonene as the coupling partners (**X** and **XI**) produces the anticipated products as well as the terminal olefin products (**10a**-(X–XI), **10b**-(X–XI), **10c**-X, and **10d**-X). Production of **10a**-d-X likely originates from regioselectivity at the migratory insertion. Furthermore, formation of **10a**-XI and **10b**-XI is likely due to the regio-selectivity of the beta-hydride elimination.

In summary, the results described herein demonstrated the potential to combine the current tools employed by chemistry and nature for the preparation of important molecules. The substrate diversity available from chemical synthesis, the robustness of AsqJ catalyzed asymmetric epoxidation, a simple Lewis Acid-catalyzed rearrangement, and the cross-coupling reactions have provided a platform for developing a suit of quinolone analogues. As enabled by mechanistic and in silico analyses, the catalytic efficiency of AsqJ paired with the mild and environmentally friendly reaction conditions provide an opportunity to apply this biocatalyst in chemoenzymatic reaction cascades. Along with the development of a biomimetic Lewis acid method of ring contraction, viridicatin can be prepared in gram-scale quantity. Overall, this work sets an example to use tools employed by synthetic chemistry and nature for the effective preparation of quinolone alkaloids. It would be interesting to test the biological activities of newly synthesized compounds in future studies.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acscatal.1c01150>.

Experimental details, Figures S1–S94, and Tables S1–S3 (PDF)

DFT_xyz (ZIP)

X-ray data (CIF)

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Notes

The authors declare no competing financial interest.

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