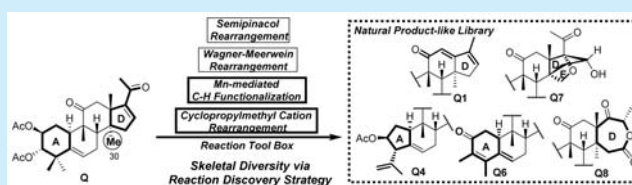


Synthesis of a Small-Molecule Library with Skeletal Diversity from Hemslecin A via the Reaction-Discovery Strategy

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S Supporting Information

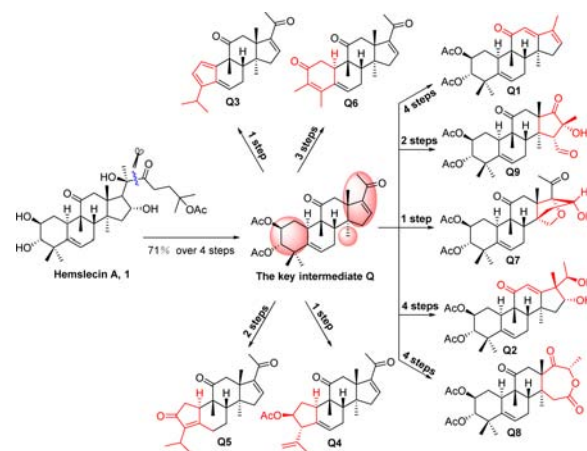
ABSTRACT: An efficient reaction tool box was developed for the synthesis of skeletally diverse and stereochemically complex templates for a small-molecule library based on the common synthon **Q**, which was prepared from hemslecin A in four steps. The reaction tool box comprises three acid-promoted rearrangements: semipinacol, Wagner–Meerwein, and cyclopropylmethyl cation rearrangements. More importantly, a Mn-mediated C–H oxidation was developed to achieve a high level of complexity, which provides a new entry for C–H functionalization of inert angular methyl groups in the chemistry of triterpenes. Our reaction-discovery strategy based on hemslecin A provides a basis for the inherent chemistry of triterpenes and could be applied for the further transformation of triterpenes.



Diversity-oriented synthesis (DOS) has become different to break through the limitation of traditional library synthesis by sampling new chemical space for small-molecule collections that exhibit a range of bioactivities.¹ The objective of DOS is to develop a synthetic scheme whereby each step opens up multiple opportunities for diversification that could be used to create natural product-like and/or drug-like small molecules with diverse molecular structures, each substantially different from the parent compound.^{1–3} An approach involving the Beckmann rearrangement and Beckmann fragmentation to complex natural products (steviol and isosteviol) has been demonstrated by the Georg group.⁴ Hergenrother et al. reported that tetracyclic diterpene gibberellic acid and steroid adrenosterone could be converted into a small collection of complex and diverse scaffolds using a ring-distortion strategy.⁵ Moreover, fumagillol has been selectively remodeled into a series of perhydroisoindoles and perhydroisoquinolines through sequential ring-opening reactions with amines using a reaction-discovery-based strategy by the Porco and Snyder groups.^{2a,c} These methods are inspired by nature's approach to creating certain complex natural products using a common intermediate to generate scores of compounds that differ substantially from one another.

Hemslecin A (**1**), a highly oxygenated tetracyclic triterpene,^{6,9} is characterized by its densely functionalized and stereochemistry-rich framework, which includes five contiguous stereogenic centers and carbocyclic moieties with each bearing reactive functional groups (Scheme 1). Additionally, hemslecin A exhibits diverse pharmacological activities, such as cytotoxicity,^{7,9} anti-HBV activity,⁸ anticancer activity,⁷ and anti-inflammatory activity.^{7,8} Thus, far, most studies of hemslecin A have been focused on the transformations of functional groups;^{6,8,9} skeletal transformations applied to hemslecin A have rarely been

Scheme 1. Concise Synthesis of a Small-Molecule Library from Hemslecin A



reported, likely because of the synthetic challenges arising from its structural complexity and its multiple reactive functionalities. Inspired by Porco's and Snyder's work² and by our continuous investigation of the bioactive diversity of terpenes,¹⁰ we herein disclose a concise synthesis of structurally complex and diverse derivatives of hemslecin A by an efficient reaction tool box that includes a number of rearrangement reactions and a Mn-mediated C(sp³)–H oxidation.

As illustrated in Scheme 1, the general synthetic strategy to achieve these molecules started from the preparation of the

Received: June 7, 2016

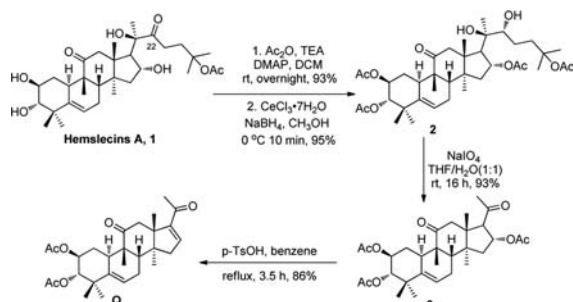
Published: August 3, 2016



requisite intermediate **Q** using hemslecin A as the starting material. The functional groups of the key intermediate **Q** provide a wide and controlled synthetic platform that can be strategically manipulated to synthesize novel, diverse, and complex chemical scaffolds within four steps (**Q1–Q9**).

The synthesis commenced with **1**, which is naturally abundant and commercially available (Scheme 2). Treatment of **1** with

Scheme 2. Synthesis of Key Intermediate **Q**

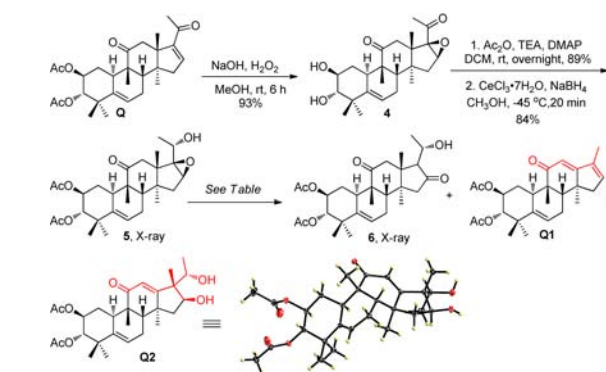


Ac₂O in the presence of 4-dimethyl-aminepyridine (DMAP) at room temperature (rt) afforded a triacetylated intermediate. Subsequent regioselective and stereoselective reduction of the C22-carbonyl group with NaBH₄ in MeOH at 0 °C gave the vicinal diol **2** in 88% yield over two steps. Oxidative cleavage of the diol on the side chain was then induced using NaIO₄ in THF/H₂O (1:1, v/v) to give ketone **3** in 93% yield,⁹ which subsequently underwent a smooth β -elimination in the presence of catalytic amounts of *p*-TsOH in refluxing benzene, readily resulting in the corresponding intermediate **Q** in 86% yield.¹¹ Thus, the common synthon **Q** with the necessary functional groups was prepared in 71% yield over four steps.

With the critical intermediate **Q** in hand, we initiated a reaction-discovery-based strategy to rapidly create complex and diverse small molecules focusing on ring D. Epoxidation of the enone **Q** under the conditions of H₂O₂ (30%)/NaOH at rt provided the desired epoxide **4** (93%),¹² which was followed by acetylation with Ac₂O and consequent reduction with NaBH₄ in MeOH at –45 °C to yield α -hydroxy epoxide **5** in 75% yield over two steps (Table 1). Treatment of **5** with BF₃·Et₂O in dry DCM at 0 °C to rt for 30 min gave a mixture of epoxide-rearranged products **Q1** (37%), **Q2** (33%), and **6** (21%) (Table 1, entry 1). We propose that the plausible mechanism of epoxide rearrangement was an oxonium-promoted Wagner–Meerwein rearrangement¹³ to form **Q2**, with further release of acetaldehyde to give **Q1** (Path a, Scheme 3). By contrast, a 3-*exo* rearrangement¹⁴ followed by semipinacol rearrangement^{13a} under acidic conditions gave compound **6** (Path b, Scheme 3). In comparison, when THF was used as solvent, **6** was obtained as the major product in 70% yield (Table 1, entry 2). Interestingly, treatment of **5** with TFA in DCM at 0 °C to rt resulted in the formation of **Q1** as the major product in 68% yield (Table 1, entry 3). Additionally, when THF was again used as a solvent in the presence of TFA, compound **Q2** was the major product, with a 74% yield (Table 1, entry 4). The structure of **Q2** was unequivocally determined by X-ray crystallographic analysis. Moreover, **Q2** could be obtained as a single product with an 82% yield in NaIO₄/H₅IO₆ and THF/H₂O (1:1, v/v) (Table 1, entry 5).

Next, the reaction-discovery strategy was tested on ring A. Schmalz et al.¹⁵ reported a cyclopropyl-methyl cation rearrangement method for tricyclo[4.4.1.0^{5,10}]undecane **I**, which easily

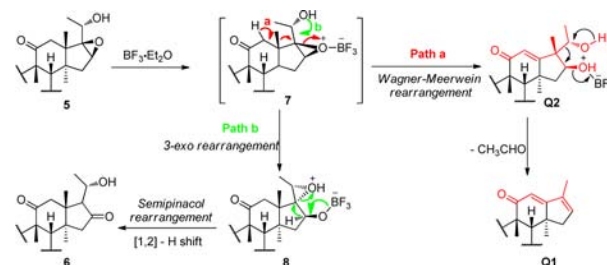
Table 1. Conditions and Optimization of Epoxide Rearrangement



entry	conditions	solvent	yield (%) ^d		
			Q1	Q2	6
1 ^a	BF ₃ ·Et ₂ O, 0 °C–rt, 30 min	DCM	37	33	21
2 ^a	BF ₃ ·Et ₂ O, 0 °C–rt, 30 min	THF	12	15	70
3 ^b	TFA, 0 °C–rt, 30 min	DCM	68	17	11
4 ^b	TFA, 0 °C–rt, 30 min	THF	16	74	8
5 ^c	NaIO ₄ /H ₅ IO ₆ , rt, 72 h	THF/H ₂ O (1:1, v/v)	nd ^e	82	nd

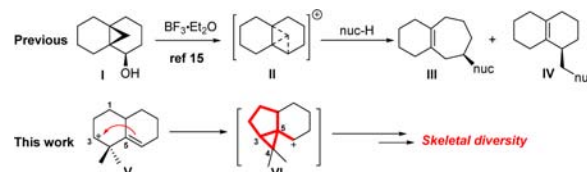
^aReaction conditions: **Q** (20 mg, 0.04 mmol); BF₃·Et₂O (6 μ L, 0.048 mmol); corresponding solvent (3 mL) in table; 0 °C to rt. ^bReaction conditions: **Q** (20 mg, 0.04 mmol); TFA (9 μ L, 0.12 mmol); corresponding solvent (3 mL) in table; 0 °C to rt. ^cReaction conditions: **Q** (20 mg, 0.04 mmol); H₅IO₆ (45 mg, 0.2 mmol); NaIO₄ (43 mg, 0.2 mmol); corresponding solvent (3 mL) in table; rt. ^dYield of the isolated product after column chromatography. ^eNot detected.

Scheme 3. Proposed Mechanisms for the Formation of **Q1**, **Q2**, and **6**



formed the bicyclic 6/7- and 6/6-fused compounds **III** and **IV** (Scheme 4). We envisioned the acid-promoted formation of the

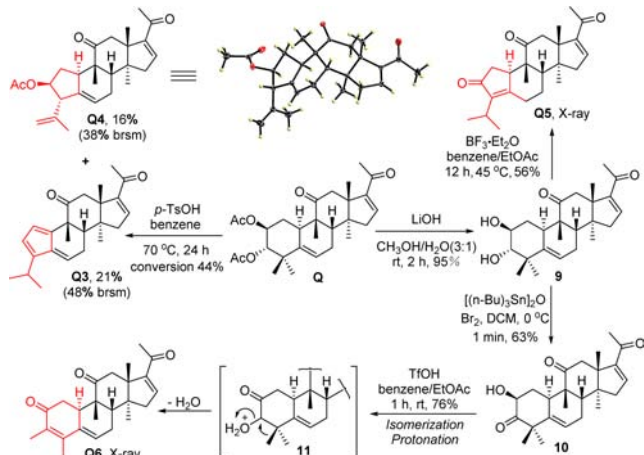
Scheme 4. Rearrangement of Cyclopropyl Methyl Cation



C3 cation, which could be further trapped by C5 to generate the cyclopropyl-methyl cation intermediate **VI**. This intermediate could likely access the ring-A-contracted skeleton, resulting in further skeletal diversity. As expected, when the intermediate **Q** was treated with *p*-TsOH in benzene at 70 °C for 24 h, the ring-

contracted products **Q3** and **Q4** were isolated in 21% (48% brsm) and 16% (38% brsm) yields, respectively (Scheme 5).

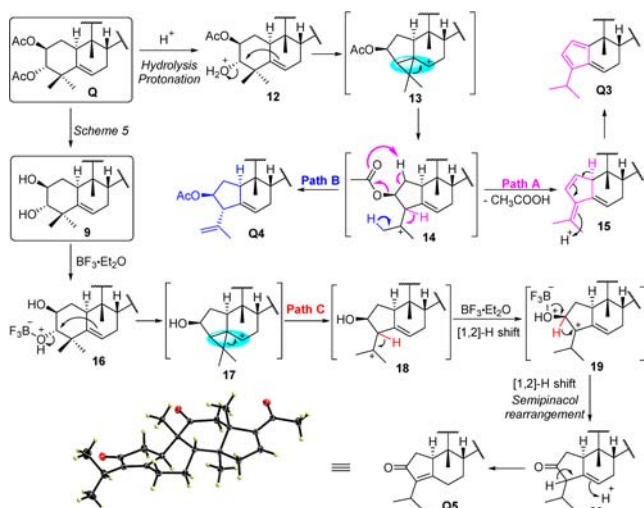
Scheme 5. Skeletal Diversity Based on Ring A Rearrangement



To further investigate the acid-promoted rearrangement of ring A, we prepared deacetylated substrate **9** in 95% yield. Regioselective oxidation of **9** using bis(tri-*n*-butyl-tin)oxide in the presence of bromine at 0 °C resulted in the α -hydroxyl ketone **10** in 63% yield.¹⁶ Interestingly, when compound **9** was treated with $\text{BF}_3 \cdot \text{Et}_2\text{O}$ in benzene/EtOAc (50:1, v/v) for 12 h, the cyclopentenone **Q5** was obtained in 56% yield. To our surprise, when compound **10** was treated with TfOH in benzene/EtOAc (50:1, v/v) at rt, a methyl [1,2]-shifted product **Q6** was obtained in 76% yield.

We propose that the transformation from **Q** to products **Q3**, **Q4**, and **Q5** was initiated by the formation of cyclopropyl-methyl cations **13** and **17**. The plausible pathway is shown in Scheme 6.

Scheme 6. Proposed Mechanisms for the Formation of **Q3**, **Q4**, and **Q5**

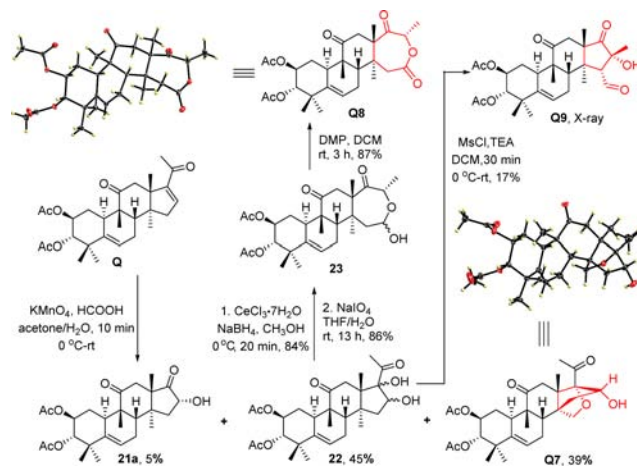


Hydrolysis and protonation of the intermediate **Q** under acidic conditions led to the formation of the cyclopropyl methyl cation **13**. The cation **13** underwent a ring-opening reaction to give the more stable tertiary cation **14**. Deprotonation and elimination of acetic acid from cation **14** gave the intermediate **15**, and the subsequent isomerization of **15** resulted in **Q3** (Path A), whereas

deprotonation of the methyl group yielded **Q4** (Path B). For cation **17**, the reaction pathway likely involved two [1,2]-H shifts and subsequent isomerization to give **Q5** (Path C).

To access a high level of skeletal diversity, we exploited the α,β -unsaturated ketone at ring D of the intermediate **Q** (Scheme 7).

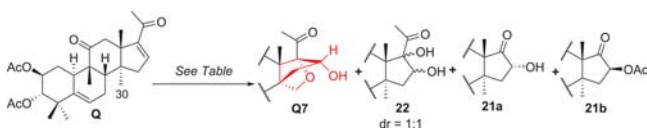
Scheme 7. Skeletal Diversity via Remodeling Ring D



The regioselective dihydroxylation of the C16–C17 double bond was first carried out in the presence of KMnO_4 (3 equiv)/HCOOH (5 equiv) in acetone/ H_2O (4:1, v/v), which resulted in the expected dihydroxylated product **22** (45%, dr = 1:1, determined by ^1H NMR) and the C17–C20 cleavage product **21a** (5%). The reaction also furnished the C30-methyl-oxidized product **Q7** (39%) with a newly formed oxabicyclo[2.2.1]-heptane motif at ring D. To the best of our knowledge, site-selective aliphatic C–H oxidations of inert angular methyl C–H bonds using KMnO_4 for the construction of strained and transannulated tetrahydrofuran ring systems have rarely been reported, although such reactions are extremely interesting. To further explore the origins of the selectivity for C–H functionalization in this transformation and to improve the yield of **Q7**, a series of reaction conditions were examined (Table 2). Finally, we obtained **Q7** in a higher yield of 54% by adjusting the ratio of acetone and water (Table 2, entry 4). The stereochemistry of the product **Q7** was further clarified by X-ray crystallographic analysis. Subsequent treatment of compound **22** with $\text{NaBH}_4/\text{CeCl}_3 \cdot 7\text{H}_2\text{O}$ in MeOH was followed by treatment with NaIO_4 to give the ring-expanded hemiacetal **23** in 72% yield over two steps. Oxidation of the resulting hemiacetal **23** with Dess–Martin periodinane provided lactone **Q8** in 87% yield, the structure of which was also confirmed by X-ray crystallographic analysis. In addition, when compound **22** was treated with MsCl/TEA in DCM at 0 °C to rt for 30 min, the rearrangement product **Q9** was isolated in 17% yield. Thus, synthesis of a diverse range of compounds was achieved; investigations of their biological activities are now underway and will be reported in due course.

In summary, we have accomplished a facile chemical approach to a natural-product-like molecular library with complex diversity by using the common synthon **Q** derived from hemslecin A. Systematic modulation of the architecture of hemslecin A was achieved via oxonium-promoted semipinacol rearrangement, Wagner–Meerwein rearrangement, and a cyclopropyl-methyl cation rearrangement. Most importantly, this work led to the discovery of a Mn-mediated site-selective aliphatic C–H

Table 2. Condition Screening for C(sp³)-H Oxidation of the C30-Methyl Group



entry ^a	conditions	solvent	yield (%) ^b		
			Q7	21a/21b	22
1	KMnO ₄ , HCOOH	acetone/H ₂ O (4:1, v/v)	39	5/nd ^c	45
2	KMnO ₄ , HCOOH	acetone/H ₂ O (2:1, v/v)	40	25/nd	35
3	KMnO ₄ , HCOOH	acetone/H ₂ O (1:1, v/v)	42	15/9	12
4	KMnO ₄ , HCOOH	acetone/H ₂ O (1:2, v/v)	54	nd/25	nd
5 ^d	KMnO ₄ , HCOOH	18-crown-6 (1 equiv)/acetone	37	nd/nd	51
6	KMnO ₄ , HCOOH	CH ₃ CN/H ₂ O (1:2, v/v)	36	24/17	11

^aUnless otherwise noted, all reactions were carried out with **Q** (50 mg, 0.11 mmol), KMnO₄ (49 mg, 0.33 mmol), and HCOOH (21 μ L, 0.55 mmol) in the corresponding solvent (4 mL) with the corresponding ratio listed in the table at 0 °C to rt. ^bYield of the isolated product after column chromatography. ^cNot detected. ^dReaction conditions: **Q** (50 mg, 0.11 mmol), HCOOH (21 μ L, 0.55 mmol), 18-crown-6 (24 mg, 0.11 mmol), KMnO₄ (49 mg, 0.33 mmol) in acetone (4 mL) at 0 °C to rt.

oxidation of inert angular methyl groups, which could easily lead to a synthetically challenging oxabicyclo[2.2.1]heptane motif. These rearrangement reactions with Mn-mediated C-H oxidation could be used as a reaction tool box to remodel other related triterpene scaffolds and to access novel chemotypes and pharmacological tools.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.orglett.6b01654.

Detailed experimental procedures; characterization data; copy of ¹H and ¹³C NMR spectra for new compounds; and X-ray crystallographic data for compounds **5**, **6**, **Q2**, **Q4**, **Q5**, **Q6**, **Q7**, **Q8**, and **Q9** (PDF)

Crystallographic data for compound **5** (CIF)

Crystallographic data for compound **6** (CIF)

Crystallographic data for compound **Q2** (CIF)

Crystallographic data for compound **Q4** (CIF)

Crystallographic data for compound **Q5** (CIF)

Crystallographic data for compound **Q6** (CIF)

Crystallographic data for compound **Q7** (CIF)

Crystallographic data for compound **Q8** (CIF)

Crystallographic data for compound **Q9** (CIF)

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Notes

The authors declare no competing financial interest.

■ ACKNOWLEDGMENTS

This project was supported by the National Natural Science Foundation of China (U1502223, 21402212).

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