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Tunable Cyclization Strategy for the Synthesis of

Zizaene-, allo-Cedrane-, seco-Kaurane-, and seco-

Atesane-Type Skeletons

Qianqian Yang, Wenjing Ma, Gaopeng Wang, Wenli Bao, Xiaoshu Dong, Xuefeng Liang, Lizhi Zhu,*and Chi-Sing Lee*

Laboratory of Chemical Genomics, School of Chemical Biology and Biotechnology, Peking University Shenzhen Graduate School, Shenzhen University Town, Xili, Shenzhen 518055, China

ABSTRACT: A versatile Lewis acid-mediated cyclization strategy has been developed for selectively establishing zizaene-, *allo*-cedrane-, *seco*-kaurane-, and *seco*- atesane-type skeletons. The zizaene- and *seco*- atesane-type skeletons can be obtained in a cascade manner, which involves Diels-Alder reaction of cyclic enones with bis- silyloxy dienes and carbocyclization of yne-enolates through Lewis acid dependent 5-or 6- *exo-dig* modes. This cyclization strategy was also employed for the core synthesisof tashironin.

Bicyclo[3.2.1] octanes (I) and bicyclo[2.2.2] octanes (II) are common bridged ring systems found in natural products (Figure 1a). When they are fused with a 5- or 6-membered ring, I will become a zizaene- 2 or seco-kaurane-type 3 skeleton, respectively, and II will become an allo-cedrane- 4 or seco- atesane-type 3 skeleton, respectively. These substructures are common motifs in the core structures of a variety of bioactive natural products. Our group has initiated a long-term projectfor developing a tunable cascade cyclization strategy for these bridged ring systems using dual-mode Lewis acids, which could provide two different modes of activation for cyclization via σ - and π -complexation. As shown in Figure 1b, we have previously reported a dual-mode Lewis acid induced Diels- Alder (DA)/carbocyclization cascade cyclization of enone III with diene IV for establishing the seco-kaurane-type skeleton V. However, this cascade cyclization strategy can only provide the 5-exo-dig cyclization products V, and the DAcycloaddition of diene IV with 5-membered enone VI (n = 1) under Lewis acid conditions has been reported to be difficult. Instead of using Danishefsky's base-mediated "sequential Michael addition protocol", we decided to employ more electron-rich bis-silyloxy diene VII for cyclization with enone VI to overcome the limitation of our previous strategy (Figure 1c). By tuning the reaction conditions, the DA intermediate VIII could undergo either 5-exo-

dig carbocyclization and lead to zizaene- and seco-kaurane-type skeletons X or 6-exo-dig carbocyclization and give allo-cedrane- and seco-atesane-type skeletons IX. Moreover, the cyclization products bear a hydroxyl at the ring junction, which could be useful for the synthesis of natural products, such as steviol glucosides, gibberellic acid, tashironin, and platencin SL3.

Figure 1. (a) Skeletons containing bicyclo[3.2.1]octanes (I) and bicyclo[2.2.2]octane (II). (b) Our previous work. (c) The goal of thiswork.

To study the selectivity of the carbocyclization, bis-silyloxy diene (±)-3 was prepared by DA cycloaddition between enone 1 and 2,3-bis(*tert*-butyldimethylsiloxy)-1,3-butadiene (2) usingMe₂AlCl in CH₂Cl₂. However, (±)-3 underwent intramolecular Mukaiyama aldol reaction at rt and gave (±)-4 as the major side product. This side reaction was suppressed by quenching the reaction at -20 °C. With (±)-3 in hand, the effects of Lewis acids on the selectivity of the carbocyclization were studied. As shown in Table 1, dual-mode Lewis acids, such as InCl₃, InBr₃, and FeCl₃, led to decomposition of the substrates (Table 1, entries 1–3). ZnCl₂ gave 43% of the cyclized products with a (±)-5a/(±)-5b ratio of 1:9 (Table 1, entry 4). Interestingly, only (±)-5b was obtained with an extended reaction time (20 h) (Table 1, entry 5), indicating Pinaol-type rearrangement of (±)-5a to (±)-5b occurred under the effectof Lewis acid. Finally, the formation of (±)-5b was optimized by using ZnBr₂ in CH₂Cl₂ at rt for 20 h (Table 1, entry 6), while ZnI₂ gave only 50% of the cyclized products with a ratio(±)-5a/(±)-5b at 1:2.8 (Table 1, entry 7). The *seco*-kaurane-type skeleton of (±)-5b was determined unambiguously by X- ray crystallography.¹²

Table 1. Effects of Lewis Acids on Cyclization of (\pm) -3^a

(=/ -	type skeleton type ske			
entry	Lewis acid (equiv)/solvent	temp (°C) yield ^b (%)) (±)-5a/5b ^e
1	InCl ₃ (1)/CH ₂ Cl ₂	rt		
2	InBr ₃ (1)/CH ₂ Cl ₂	rt		
3	FeCl ₃ (1)/CH ₂ Cl ₂	rt		
4	ZnCl ₂ (1)/CH ₂ Cl ₂	rt	43	1:9
5	ZnCl ₂ (1)/CH ₂ Cl ₂	rt ^d	46	(\pm) -5b only
6	ZnBr ₂ (1)/CH ₂ Cl ₂	rt ^d	98	(\pm) -5b only
7	ZnI ₂ (1)/CH ₂ Cl ₂	rt <u>e</u>	50	1:2.8
8	AuCl (1)/CH3CN	rt	82	1.3:1
9	AuCl (1)/CH3CN	- 40) 59	1.4:1
10	AuCl (1)/CH ₂ Cl ₂	rt	46	3:1
11	AuCl (1)/CH ₂ Cl ₂	- 78	3 28	3:1
12	PPh ₃ AuCl (0.2)/toluene	0 or :	rt	
13	AuCl ₃ (0.2)/CH ₂ Cl ₂	0 or :	rt	
14	Cu(OTf)2 (1)/CH2Cl2	0 or :	rt	
15	Pd(OAc) ₂ (1)/THF	0 or :	rt	
16	PdCl ₂ (1)/CH ₃ CN	0 or :	rt	
17	PtCl ₂ (1)/THF	0 or :	rt	
18	PtCl ₂ (1)/CH ₃ CN	rt		
19	PtCl ₂ (1)/CH ₃ CN	0	10	(\pm) -5a only
20	PtCl ₂ (1)/CH ₂ Cl ₂	0	65	3.6:1
21	PtCl ₂ (0.5)/CH ₂ Cl ₂	0	62	5.1:1
22	PtCl ₂ (0.2)/CH ₂ Cl ₂	0	60	5:1
23	PtCl ₂ (0.5)/CH ₂ Cl ₂	- 78	3 72	7.8:1

 $^{^{}d}$ The general procedures were followed (time = 30 min). b Isolated yield (%) after silica gel column chromatography. c The product ratios were determined by 1 H NMR signals of the exocyclic alkenes. d Reaction time = 20 h. e Reaction time = 40 h.

After studying the effects of dual-mode Lewis acids, a variety of π -Lewis acids were investigated. Using AuCl³ in CH₂CN gave 82% yield of the cyclized products with slightly higher selectivity for (±)-5a (Table 1, entry 8). Lowering the reaction temperature to -40 °C led to a lower yield (59%) with a similar product ratio (Table 1, entry 9). Switching the solventto CH₂Cl₂ gave only 46% yield of the cyclized products with a higher selectivity for (±)-5a (3:1) (Table 1, entry 10). An attempt at increasing the product ratio by lowering the reaction temperature to -78 °C led to a much lower yield (28%) with a similar level of selectivity (Table 1, entry 11). PPh₃AuCl, ¹³ AuCl₃, ¹⁴ Cu(OTf)₂, ¹⁵ Pd(OAc)₂, and PdCl ¹⁶ did not provide any cyclized product at 0 °C or rt (Table 1, entries 12–16). PtCl₂¹⁷ in THF led to hydrolysis of (±)-3 and resulted in amixture of α-silyloxy ketones (Table 1, entry 17). Using PtCl2in CH2CN also led to decomposition of the substrate at rt(Table 1, entry 18) but surprisingly afforded (±)-5a in 10% yield at 0 °C (Table 1, entry 19). This encouraging result prompted us to study the effects of PtCl2 under different conditions. Switching the solvent to CH₂Cl₂ afforded the cyclized products in 65% yield with a ratio (\pm)-5a/(\pm)-5b at 3.6:1 (Table 1, entry 20). After studying the effects of catalyst loading and reaction temperature, the optimal results was obtained by using 0.5 equiv of PtCl₂ in CH₂Cl₂ at -78 °C, which afforded 72% yield of the cyclized products in favor of the seco-atesane-type skeleton of (\pm) -5a (7.8:1) (entry 23). The results of this study indicated that π -Lewis acids generally favored the 5-exo-dig cyclizations and afforded the seco-kaurane- type skeleton preferentially.

Based on the results of the above study, ZnBr2 could be a suitable dual-mode Lewis acid for the cascade cyclization between enone 1 and diene 2. After a survey of different reaction conditions, the cascade cyclization was optimized by using 1 equiv of ZnBr2 in CH2Cl2 at rt, which afforded 93% yield of (±)-5b in a single operation (Table 2, entry 1). With the above optimal conditions in hand, a series of dienes and enones with different steric hindrances were investigated. Cascade cyclization of enone 1 with diene 6 led to only 35% yield of the cyclized products (Table 2, entry 2) due to rapid hydrolysis of diene 6. The selectivity for (±)-7b is low, and equilibration of (\pm) -7a to (\pm) -7b became unfavorable after hydrolysis of the TMS ethers of the cyclized products. The more bulky diene 8 gave a reasonable good yield of the cyclized product (±)-9b, but the equilibration of (\pm) -9a to (\pm) -9b is slow (Table 2, entry 3). These results indicated that diene 2 has the optimal size for balancing the stability and reactivity in the cascade cyclization. Cascade cyclization of enone 1 with cyclicdiene 10 afforded 79% yield of (±)-11a (the structure was determined by X-ray crystallography)¹² as the only product (Table 2, entry 4). However, diene 12 gave only the Mukaiyama Michael side product 13 (Table 2, entry 5) due to its steric hindrance. Cascade cyclization of (±)-enone 14 with diene 2 gave 79% yield of (±)-15b as a single diastereomer (Table 2, entry 6), suggesting the possibility for developing an diastereoslective version via substrate-control. For synthesis of the zizaene- and allo-cedrane-type skeletons, 5membered enone 16 was employed for the cascade cyclization with diene 2, which resulted in 71% of a 4:1 mixture of (\pm) -17a(zizaene) and (\pm) -17b (allo-cedrane) (Table 2, entry 7). Interestingly, the ratio of the cyclized products remains unchanged upon heating with long reaction time probably due to the ring strain of the cyclized products. Introducing an ethyl group to the alkyne terminus (enone 18) led to excellent selectivity for the zizaene-type-skeleton (±)-19a as a single isomer with moderate yields (Table 2, entry 8).

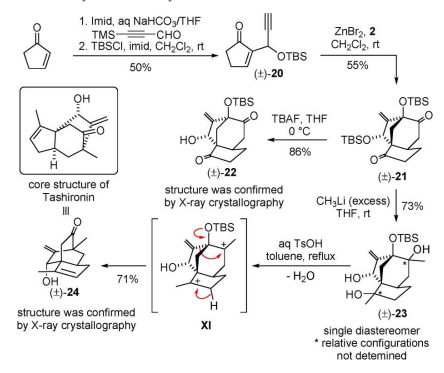
Table 2. Substrate Scope of the Cascade Cyclization^a

entry	substra	tes	pr	oduct(s)	yield (%) ^b	product ratio (a/b) ^c
1	R	OR	or or	OR	93	(±) -5b only
2 3	1, R = H 1, R = H 1, R = H	2, R = TBS 6, R = TMS 8, R = TIPS	(\pm) -5a, R = TBS (\pm) -7a, R = H (\pm) -9a, R = TIPS	(±)- 5b , R = TBS (±)- 7b , R = H (±)- 9a , R = TIPS	35 78	1:2 1:16
4	1, R = H	OTBS OTBS	OTBS O (±)-11a ¹²	-	79	(±)- 11a only
5	1, R = H	OTBS OTBS	13	Отвѕ	35^d	single diastereomer ^d
6	(±) -14 , R = OTBS	OR OR OR OR OR	-	твsо (±)- 15 b	79	(\pm)-15b only single diastereomer ^d
7	16	2, R = TBS	OTBS (±)-17a	(±)-17b ¹²	71	4:1
8	18	2 , R = TBS	OH O (±)-19a	-	45	(±)-19a only single isomer

^aThe general procedures were followed. ^bIsolated yield (%) after silica gel column chromatography. ^cThe product ratios were determined by the ¹HNMR signals of the exocyclic alkenes. ^aThe relative configurations of the OTBS group were not determined. ^cThe geometry of the alkene was notdetermined.

To demonstrate the utility of the cascade cyclization, a model toward the synthesis of tashironin18 was studied. As shown in Scheme 1, Morita—Baylis—Hillman reaction of cyclopentenone with 3-(trimethylsilyl)propynal followed by silylation gave (±)-20, which underwent cascade cyclization with diene 2 using ZnBr2 and afforded good yields of cyclized product (±)-21 diastereoselectively. The structure of (±)-21 was determined by X-ray crystallography of (±)-22,12 which was obtained from by treating (±)-21 with TBAF at 0 °C. Addition of excessive MeLi gave (±)-23 as a single diastereomer. The relative configurations of the tertiary alcohols were not determined since the stereogenic centers will be destroyed in the subsequent step. Upon treatment of aq TsOH in refluxing toluene,19 the carbocation (XI) generated in situ underwent elimination and Pinol-type rearragement, which led to the allo- cedrane-type skeleton of (±)-24.12 The model compound (±)-24 contains the core structure of tashironin, which provides a foundation for developing a total synthesis for this natural product and related compounds based on this cascade cyclization strategy.

Scheme 1. Model Study toward the Synthesis of Tashironin



In summary, we have extensively studied the effects of Lewisacids on the selectivity of the carbocyclization of the DAintermediate (±)-3 and developed a tunable cyclization strategyfor establishing the zizaene-, *allo*-cedrane-, *seco*-kaurane-, and *seco*-atesane-type skeletons. Upon treatment of ZnBr₂/CH₂Cl₂ at rt, the 6-membered enone 1 and diene 2 underwent cascade cyclization and afforded the 6-*exo-dig* cyclized product (bearingthe *seco*-atesane-type skeleton) selectively in 93% yield. The 5-*exo-dig* cyclized product (bearing the *seco*-kaurane-type skeleton) can be obtained via carbocyclization of the DA intermediate (±)-3 using PtCl₂ at -78 °C (72%, 7.8:1). On the other hand, the cascade cyclization between 5-membered enone(±)-20 and diene 2 using ZnBr₂ favors the 5-*exo-dig* cyclizationproduct (bearing the zizaene-type skeleton), which was converted to the *allo*-cedrane-type skeleton of (±)-24 (the core of tashironin) with two steps in a model study. Development of an asymmetric version using chiral Lewis acids and a total synthesis of tashironin and related natural products using this cascade cyclization strategy are ongoing in our laboratory.

AUTHOR INFORMATION

Corresponding Authors

*E-mail: lzzhu86@pku.edu.cn.

*E-mail: lizc@pkusz.edu.cn.

ORCID

Chi-Sing Lee: 0000-0002-3564-8224

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

We gratefully thank the NSFC (National Natural Science Foundation of China, Nos. 21502002 and 21572006), Chinese Postdoctoral Science Foundation (2015M580910), the SSTIC (Shenzhen Science and Technology Innovation Committee, JSGG20160301103446375, JCYJ20160330095659560, and JCYJ20150626111042525), and Peking University Shenzhen Graduate School for the *fi*nancial support. Special acknowledgement is made to Dr. Wesley Tingkwok Chan (Hong KongPolytechnic University) for the X-ray crystallography of compounds (±)-5b, (±)-11a, (±)-17b, (±)-22, and (±)-24.

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