Total Synthesis of (–)-Blepharocalyxin D and Analogues

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An efficient strategy for the total synthesis of (–)-blepharocalyxin D and an analogue is described. The key step involves an acid-mediated cascade process in which reaction of methyl 3,3-dimethoxypropanoate with γ , δ -unsaturated alcohols possessing diastereotopic styrenyl groups gives *trans*-fused bicyclic lactones with the creation of two rings and four stereocenters in one pot.

Kadota and co-workers have reported the isolation of a family of related polyphenolic compounds from extracts of the seeds of *Alpinia blepharocalyx*.¹ Their structures were determined originally using spectroscopic methods. More recent synthetic studies by Rychnovsky and co-workers have led to structural reassignment of some of these compounds.² We were drawn to one particular diarylheptanoid blepharocalyxin D (1), which was isolated in very small quantities from seeds of A. blepharocalyx. Blepharocalyxin D exhibits potent antiproliferative activity against murine colon 26-L5 carcinoma and human HT-1080 fibrosarcoma cells.³ It has an unusual structure assembled upon a trans-2,8-dioxabicyclo[4.4.0]decane adorned by four side chains each in an equatorial position (Figure 1). We now describe a flexible strategy for the synthesis of trans-2,8-dioxabicyclo[4.4.0]decanones in which two rings and four stereocenters are created in a one-pot cascade process and culminates in the total synthesis of blepharocalyxin D and an analogue of the natural product.

Lee and co-workers have reported the only previous total synthesis of blepharocalyxin D in which they employed two separate Prins cyclizations to construct each of



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the oxane rings.⁴ However, introduction of the C-5 equatorial side chain proved challenging. An interesting Prinspinacol reaction gave predominantly an unwanted axial aldehyde at C-5 which was epimerized prior to a Julia chain extension to the required styrenyl group. Our goal was to develop a new synthetic strategy to blepharocalyxin D in which both rings of the *trans*-fused bicyclic framework would be generated in one-pot with all side chains equatorial. The approach was to be versatile to provide analogues of potential biological interest.

Our retrosynthetic analysis of 1 is shown in Scheme 1. The C-9 side chain would be introduced via a Grignard addition to lactone 2 followed by reduction of the resultant lactol. An advantage of using the lactone is that various

⁽¹⁾ Tezuka, Y.; Gewali, M. B.; Ali, M. S.; Banskota, A. H.; Kadota, S. J. Nat. Prod. 2001, 64, 208.

^{(2) (}a) Tian, X.; Jaber, J. J.; Rychnovsky, S. D. J. Org. Chem. 2006, 71, 3176. (b) Tian, X.; Rychnovsky, S. D. Org. Lett. 2007, 9, 4955.

^{(3) (}a) Ali, M. S.; Tezuka, Y.; Banskota, A. H.; Kadota, S. J. Nat. *Prod.* **2001**, *64*, 491. (b) Tezuka, Y.; Ali, M. S.; Banskota, A. H.; Kadota, S. *Tetrahedron Lett.* **2000**, *41*, 5903.

^{(4) (}a) Ko, H. M.; Lee, D. G.; Kim, M. A.; Kim, H. J.; Park, J.; Lah, M. S.; Lee, E. *Org. Lett.* **2007**, *9*, 141. (b) Ko, H. M.; Lee, D. G.; Kim, M. A.; Kim, H. J.; Park, J.; Lah, M. S.; Lee, E. *Tetrahedron* **2007**, *63*, 5797.

Grignard reagents could be used to prepare analogues of the natural product.

Scheme 1. Retrosynthesis of Blepharocalyxin D



The *trans*-2,8-dioxabicyclo[4.4.0]decanone (2) would be assembled from γ , δ -unsaturated alcohol 3 in which the double bond is placed one position further away from the hydroxyl group than in homoallylic alcohols and derivatives commonly used in Prins cyclizations.⁵ The success of this strategy relied upon reaction of 3 (to be prepared from (*S*)-homoallylic alcohol 4) with an electrophile to generate an intermediate oxycarbenium I with diastereotopic styrenyl groups. Cyclization was predicted to proceed to give stabilized carbocation II with the first of the oxane rings with an equatorial substituent at C-5. Intramolecular trapping of the resultant carbocation with an ester would generate the second heterocycle giving lactone 2 with an equatorial substituent at C-7.⁶

To establish conditions for the proposed cyclization, first the known⁷ γ , δ -unsaturated alcohol **5** was converted to enol ether **6** using methyl propiolate and catalytic quinuclidine. Treatment of **6** with TMSOTf gave bicyclic lactone **7** in 69% yield (Scheme 2).⁸ The structure was elucidated by NMR spectroscopy and confirmed by X-ray crystallography.





The second approach to 7 involved the direct reaction of γ , δ -unsaturated alcohol **5** with commercially available methyl 3,3-dimethoxypropanoate **8** giving, after optimization, lactone **7** in 67% yield. These studies established an approach for the direct conversion of γ , δ -unsaturated alcohols to *trans* fused bicyclic lactones with equatorial groups at both C-3 and C-7.

For the proposed synthesis of blepharocalyxin D (Scheme 2), a γ , δ -unsaturated dienol **3** was required as the substrate for the key cyclization to generate the bicyclic framework with an equatorial styrenyl group at C-5. Following the studies outlined in Scheme 2, we selected dienol **16** with phenyl groups as the initial target.

Dienol **16** was prepared from the known⁹ (*S*)-homoallylic alcohol **9** as shown in Scheme 3. Following protection of the secondary alcohol as TBS ether **10**, the double bond was oxidatively cleaved with $OsO_4/NaIO_4$ to give aldehyde **11**. A Horner–Wadsworth–Emmons chain extension of **11** using phosphonate **12** and K₂CO₃ gave (*E*)- α , β -unsaturated ketone **13** with excellent stereocontrol. The final carbon– carbon bond was formed via a rhodium-mediated 1,4addition of styrenyl boronic acid to enone **13** under the conditions reported by Hiyashi¹⁰ to give ketone **14** in 94% yield as a 1:1 mixture of diastereomers. The lack of stereocontrol was not a problem as this newly created stereocenter would be destroyed later in the synthesis as the target, dienol **16**, has two identical (*E*)-phenylethenyl side chains.

We envisaged that the second double bond of **15** could be readily generated from ketone **14** via reduction to an alcohol followed by elimination. While reduction of ketone **14** with NaBH₄ proceeded smoothly to give the expected benzylic alcohol as a mixture of isomers, the elimination step proved problematic via either the corresponding acetate or mesylate. However, the rarely used xanthate formation/Chugaev elimination in this instance worked well. Thus, conversion of the alcohol to a xanthate using NaH/CS₂/MeI followed by refluxing with NaHCO₃ in

⁽⁵⁾ For reviews on Prins cyclizations, see: (a) Olier, C.; Kaafarani, M.; Gastaldi, S.; Bertrand, M. P. *Tetrahedron* **2010**, *66*, 413. (b) Crane, E. A.; Scheidt, K. A. *Angew. Chem., Int. Ed.* **2010**, *49*, 8316.

⁽⁶⁾ For examples of the use of intramolecular trapping using tethered oxygen nucleophiles, see: (a) Fráter, G.; Muller, U.; Kraft, P. Helv. Chim. Acta 2004, 87, 2750. (b) Elsworth, J. D.; Willis, C. L. Chem. Commun. 2008, 13, 1587. (c) Yadav, J. S.; Chakravarthy, P. P.; Borkar, P.; Reddy, B. V. S.; Sarma, A. V. S. Tetrahedron Lett. 2009, 50, 5998. (d) Reddy, B. V. S.; Borkar, P.; Yadav, J. S.; Reddy, P. P.; Kunwar, A. C.; Sridhar, B.; Grée, R. Org. Biomol. Chem. 2012, 10, 1349. (e) Reddy, B. V. S.; Sarma, B. Org. Biomol. Chem. 2012, 10, 6562.

⁽⁷⁾ Bunt, A. J.; Bailey, C. D.; Cons, B. D.; Edwards, S. J.; Elsworth, J. D.; Pheko, T.; Willis, C. L. *Angew. Chem., Int. Ed.* **2012**, *51*, 3901.

⁽⁸⁾ Previous examples of Prins reactions using enol ethers include:
(a) Hart, D. J.; Bennett, C. E. *Org. Lett.* 2003, *5*, 1499. (b) Barry, C. S.; Bushby, N.; Harding, J. R.; Willis, C. L. *Org. Lett.* 2005, *7*, 2683. (c) Yang, Y.; Jia, P.; Liu, S.; Yu, W. *Chin. J. Chem.* 2012, *30*, 1439.

⁽⁹⁾ Keck, G. E.; Krishnamurthy, D. Org. Synth. 1998, 75, 12.

⁽¹⁰⁾ Takaya, Y.; Ogasawara, M.; Hayashi, T.; Sakai, M.; Miyaura, N. J. Am. Chem. Soc. **1998**, 120, 5579.

⁽¹¹⁾ Chugaev, L. Chem. Ber. 1899, 32, 3332. For selected examples of the Chugaev elimination in synthesis, see: (a) Meulemans, T. M.; Stork, G. A.; Macaev, F. Z.; Jansen, B. J. M.; deGroot, A. J. J. Org. Chem. 1999, 64, 91178. (b) Padwa, A.; Zhang, H. J. Org. Chem. 2007, 72, 2570. (c) Nicolaou, K. C.; Ortiz, A.; Zhang, H.; Dagneau, P.; Lanver, A.; Jennings, M. P.; Arseniyadis, S.; Faraoni, R.; Lizos, D. E. J. Am. Chem. Soc. 2010, 132, 7149.

xylene to effect a Chugaev elimination¹¹ gave the required alkene **15** in 68% overall yield from ketone **14** and with excellent *E*-selectivity. Silyl ether **15** was deprotected using 2% HCl in ethanol to give the required γ , δ -unsaturated alcohol **16**.



With dienol **16** in hand, the key cyclization was carried out with acetal **8** and TMSOTf in CH_2Cl_2 at -30 °C giving bicyclic lactone **17** with the creation of four new stereocenters (Scheme 4). The presence of the three equatorial substituents and *trans* ring junction was determined from the ¹H NMR spectrum which showed the expected *trans* diaxial couplings for each of the methine protons. It was evident that there was excellent stereocontrol at C-5 with cyclization occurring to give exclusively the equatorial styrenyl side-chain.

With five out of the six asymmetric centers of the blepharocalyxin D framework intact, the final carboncarbon bond-forming reaction introduced the side-chain at C-9. Treatment of **17** with 2-(*p*-methoxyphenyl)ethylmagnesium bromide followed by reduction of the resultant lactol with TMSOTf, Et₃SiH gave **18** in 48% yield over the two steps. The NMR data were consistent with the side chains all being in equatorial positions and was confirmed by X-ray crystallography. It is interesting to note the π stacking between the 7-phenyl ring and the styrenyl side chain at C-5 in the crystal structure.

Turning to the synthesis of blepharocalyxin D, the first challenge was to prepare a dienol with appropriate *para*substituted phenyl groups. In their total synthesis, Lee and Scheme 4. Completing the Synthesis Blepharocalyxin D Analogue 18 and Crystal Structure of 18



co-workers⁴ had used a *p*-methoxyphenyl group which was deprotected using LiSPr/HMPA. However, the unsaturated *p*-methoxyphenyl derivative **19** has an activated electron-donating aromatic ring and proved unstable to the TMSOTf-promoted cyclization conditions (Scheme 5).⁷ In contrast, the acid-mediated reaction of methyl 3,3-dimethoxypropanoate **8** and unsaturated alcohol **20** with the electron-withdrawing *p*-phenylsulfonyl group gave bicyclic lactone **21** in 93% yield. Hence the use of phenylsulfonyl esters was selected for the total synthesis of blepharocalyxin D.





For the natural product, a similar synthetic strategy was used to dienol intermediate **29** as for the analogue **16** (Scheme 3).¹² The chain extension of aldehyde **24** to **26** was achieved in 91% yield using a Wittig reaction with

(12) In this case, the synthetic route began with the novel (S)-homoallylic alcohol **22** which was prepared in 98% yield via a Nokami crotyl transfer reaction.



Nokami, J.; Ohga, M.; Nakamoto, H.; Matsubara, T.; Hussain, I.; Kataoka, K. J. Am. Chem. Soc. 2001, 123, 9168.

stabilized ylide 25; this was a significant improvement on the HWE reaction used to prepare enone 13 (65% yield). The rhodium-mediated 1,4-addition to enone 26 using the novel boronic acid (prepared from vinyl boronic acid MIDA boronate and *p*-sulfonylstyrene¹³) gave ketone 27 which was converted to the required dienol 29 using the same reduction/elimination sequence as for analogue 16.

Reaction of dienol **29** with acetal **8** and TMSOTf gave bicyclic lactone **30** as a single diastereomer with the creation of the two rings and four stereocenters in 75% yield (Scheme 6). The C-9 side chain was introduced via a Grignard addition/reduction protocol giving **31** with the fully assembled carbon framework of our target. Finally deprotection of the phenolic groups with LiSPr/HMPA gave (–)-blepharocalyxin D (1) in 85% yield. Interestingly Lee and co-workers⁴ reported that their synthetic blepharocalyxin D exhibited variable specific rotation values in methanol (ranging from -77.1 to -89.9 depending on concentration), our synthetic sample [α]_D -79.2 (*c* 0.23, MeOH) was in accord with their data.¹⁵

In conclusion, a new synthetic strategy to (-)-blepharocalyxin D and an analogue has been developed. The approach proceeds via γ , δ -unsaturated dienols **16** and **29** prepared using an efficient rhodium-mediated conjugate addition to enones **13** and **26** followed by reduction of the resultant ketones and xanthate formation/Chugaev elimination to the required (*E*)-double bond. The key step involves reaction of the dienols with

(13) Initial attempts to synthesize boronic acid including the hydroboration of the corresponding alkyne and lithiation of the corresponding β -halostyrene followed by quenching with triisopropylborane were unsuccessful. Instead the required reagent was synthesized in 92% yield from vinyl boronic acid MIDA boronate and *p*-sulfonylstyrene using conditions reported by Burke.¹⁴



G2 = Grubbs 2nd generation catalyst

Scheme 6. Completing the Synthesis of 1



methyl 3,3-dimethoxypropanoate **8** to give bicyclic lactones **17** and **30** via a one-pot cascade to generate two rings and four stereocenters. A Grignard addition/ reduction protocol followed by deprotection gave blepharocalyxin D. The bicyclic lactones are flexible intermediates for use as building blocks in organic synthesis and have potential widespread value.

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Supporting Information Available. Preparation and characterization of the compounds described in this paper. This material is available free of charge via the Internet at http://pubs.acs.org

(14) Uno, B. E.; Gillis, E. P.; Burke, M. D. Tetrahedron 2009, 65, 3130.

(15) The optical rotation for the natural product reported by Kadota and co-workers³ was $[\alpha]_D$ +18.5 (*c* 0.025, MeOH), but Lee reports that the NMR spectra of the natural product contained impurities.⁴

The authors declare no competing financial interest.