# Communications

#### **Total Synthesis**

DOI: 10.1002/anie.200602569

### Total Synthesis of (+)-Phyllantidine\*\*

Cheryl A. Carson and Michael A. Kerr\*

Dedicated to Professor K. C. Nicolaou on the occasion of his 60th birthday

There exists a small group of alkaloids isolated from the *Euporbiaceae* family of plants known as the securinega alkaloids (Figure 1).<sup>[1]</sup> These compounds have an indolizidine

[\*] C. A. Carson, Dr. M. A. Kerr Department of Chemistry The University of Western Ontario London, ON, N6A5B7 (Canada) Fax: (+1)519-661-3022 E-mail: makerr@uwo.ca

[\*\*] We thank the Natural Sciences and Engineering Research Council (NSERC) of Canada and Boehringer Ingelheim Canada for funding. We are grateful to D. Hairsine for performing MS analyses. C.A.C. is the recipient of an NSERC CGSM postgraduate scholarship.

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Figure 1. The securinega alkaloids including (+)-phyllantidine (5).

core imbedded within an azabicyclo [3.2.1] ring system. This is fused to a butenolide moiety forming a rather interesting and structurally complex molecular framework. Securinine (1) and its C2 epimer allosecurinine (2) are constituents of Securinega suffruticosa,<sup>[2]</sup> and the antipodal compounds virosecurinine (3) and viroallosecurinine (4) are found in Securinega virosa.<sup>[3]</sup> While these compounds show interesting activity in the central nervous system (CNS) in the form of antagonism of the  $\gamma$ -aminobutyric acid (GABA) receptor,<sup>[4]</sup> the synthetic chemist is drawn to the compact and complex architecture of the compounds. Indeed several syntheses of the securinine series of compounds have been reported.<sup>[5]</sup> Of interest to us are not the indolizidines 1-4 but a related and much rarer alkaloid phyllantidine 5 (isolated from Phyllanthus discoides and Seurinega suffruticosa)<sup>[6]</sup> and its enantiomer (from *Breynia coronata*).<sup>[7]</sup>

To date, no syntheses of phyllantidine (or *ent*-phyllantidine) have been reported, although phyllantidine is available through the peroxide (or peracid) oxidation of virosecurinine.<sup>[8]</sup> This proceeds via the *N*-oxide which undergoes a Meisenheimer rearrangement yielding phyllantidine. Since the synthetic routes to the securinine alkaloids are quite complex and lengthy, this route to phyllantidine is less than appealing. Herein, we present a convenient and direct synthesis of (+)-phyllantidine.

Perhaps the most significant structural feature of phyllantidine is the tetrahydro-1,2-oxazine ring. This heterocyclic motif is uncommon in natural products and is found in FR-900482 (and a few related compounds)<sup>[9]</sup>. This structural feature also makes phyllantidine an elusive target since there are few ways to directly prepare tetrahydro-1,2-oxazines. Recently, however, we reported that nitrones (either as isolated compounds or generated in situ) react smoothly with 1,1-cyclopropane diesters under the influence of Lewis acids to form tetrahydro-1,2-oxazines in what we have termed a "homo-1,3-dipolar cycloaddition".<sup>[10]</sup> The reactions are diastereoselective and yield only 3,6-*cis* adducts. Both the substitution pattern and relative stereochemistry of the adducts from this cycloaddition would fulfill the requirements of a practical synthetic route to phyllantidine (Scheme 1).



Scheme 1. Retrosynthesis of phyllantidine. RCM = ring-closing olefin metathesis.

Initially, we envisioned a cycloaddition between a cyclic nitrone 11 and cyclopropane 8 as a rapid access to an advanced bicyclic intermediate such as 12 [Eq. (1)]. However, we have not been able to obtain adducts from the cyclo-



addition reaction using nitrones such as 11 as the substrate.

Our synthesis of phyllantidine commenced with the threecomponent coupling of hydroxylamine 13, aldehyde 14, and cyclopropane  $\mathbf{8}^{[11]}$  under the influence of catalytic ytterbium triflate hydrate to give the tetrahydro-1,2-oxazine 15 in 86% yield as a 12:1 mixture of diastereomers, in which the major product bore the required cis relationship between the oxazine vinyl substituent and the alkyl chain (Scheme 2). Interestingly, we occasionally saw this slight loss of diastereochemical integrity when the cycloaddition reaction mixture was heated to reflux. However, perhaps more interesting and surprising is that there was approximately 10% erosion of the absolute stereochemistry, with the cis adduct isolated as a 90:10 mixture of enantiomers. This has mechanistic implications and a detailed study of the stereochemistry of these cycloadditions is underway. Krapcho decarboxylation proceeded smoothly to produce 16 as a 1:1 mixture of diastereomers in 85% yield. Treatment of 16 (mixture of epimers) with KHMDS to generate the potassium enolate followed by treatment with the Davis oxaziridine<sup>[12]</sup> gave the hydroxy esters 17 and 18 as an inseparable mixture of diastereomers (1:3 in favor of the desired isomer; Scheme 2).

We were delighted to find that our expectations of a diastereoselective hydroxylation were born out, since this

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**Scheme 2.** Synthesis of advanced intermediate **23**. a) Ytterbium(III) trifluoromethanesulfonate hydrate (5 mol%), MS (4 Å), toluene, heated at reflux (86%); b) LiCl (5 equiv), DMSO, H<sub>2</sub>O, 160 °C (85%); c) KHMDS, Davis oxaziridine, THF, -78 °C (80% as a 3:1 diasteromeric mixture of **18** and **17**); d) LAH, THF, 0°C (97%); e) 2,2-dimethoxypropane, *p*-toluenesulfonic acid, DMSO (74%); f) 6 N HCl, THF (93% based on **20**); g) IBX, DMSO (64%); h) CH<sub>2</sub>=CHMgBr, THF, (76%); i) IBX, DMSO (79%). PMB = *p*-methoxybenzyl, Ts = toluene-4-sulfonyl, LAH = lithium aluminum hydride, KHMDS = potassium bis(trimethylsilyl)amide, IBX = *o*-iodoxybenzoic acid.

transformation was the greatest concern to us at the outset of the project. Our rationale for the predicted selectivity is shown in Scheme 3. The enolate derived from 16 may be envisaged as the chair conformers **A** or **B**. Whether one invokes **A** or **B**, approach of the electrophilic oxidant should be from the top face (bold arrow in Scheme 3) to avoid either a 1,3-diaxial interaction with the vinyl group in **A** or a 1,2interaction with the alkyl chain in **B**. It is worth noting that in a simpler model system, in which both the substituents on the nitrogen atom and the alkyl chain were replaced by phenyl moieties, there was complete selectivity for the desired isomer.

Reduction of this mixture with LAH afforded the diols (in 97% yield), which upon derivatization gave the acetonides **19** 



*Scheme 3.* Model for selective hydroxylation showing steric interactions between oxidant and vinyl group in **A** and between oxidant and alkyl chain in **B**.

and **20** as a 1:3 mixture as expected (Scheme 3). This mixture was amenable to simple separation by flash column chromatography giving, after acetonide removal (in 93 % yield), **21** as a single isomer. Oxidation of the primary hydroxy group gave an aldehyde **22** (in 64 % yield), which was treated with vinylmagnesium bromide to give a diastereomeric mixture of the allylic alcohols in 76 % yield. Oxidation gave the enone **23**, which we anticipated would be an appropriate substrate for RCM.

While the sequence of reactions in Scheme 2 certainly produced **23** in a relatively efficient way, this sequence of protection through formation of acetonides, separation, and deprotection seemed rather clumsy. This procedure was necessary at the time to better understand the reaction sequence. In addition, separation of the diols resulting from the reduction of mixture **17/18** was not possible in a preparative manner.

In order to circumvent the tedious protection/deprotection sequence, the diastereomeric mixture of 17/18 (ratio 1:3) was subjected to a sequence of four reactions (Scheme 4), namely, reduction using LAH, oxidation of the primary alcohol to an aldehyde using IBX, addition of vinylmagnesium bromide, and finally oxidation of the allylic alcohol to give the vinyl ketone 23. At this juncture, the unwanted diastereomer was readily removed by flash column chromatography. Enone 23 proved to be a suitable substrate for RCM with Grubbs second-generation catalyst. In the event, ring closure to give enone 24 proceeded in 74% yield. The Oacetyl derivative of 24 was explored as a substrate for aldoltype ring closure to form the butenolide, yet all attempts to affect this cyclization failed. Fortunately, an alternative acylation<sup>[13]</sup> using diethylphosphonoacetic acid afforded substrate 25 (in 71% yield), which after an intramolecular Horner-Emmons reaction, was converted into the desired 26 in quantitative yield.<sup>[14]</sup> The natural product **5** was ultimately

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5: (+)-phyllantidine

**Scheme 4.** Synthesis of (+)-phyllantidine. a) LAH, THF, 0°C; b) IBX, DMSO; c)  $CH_2$ =CHMgBr, THF; d) IBX, DMSO (20% overall yield of one diastereomer from a mixture of **17** and **18**); e) second-generation Grubbs catalyst (20 mol%),  $CH_2Cl_2$ , heated at reflux (74%); f) dieth-ylphosphonoacetic acid, DCC,  $CH_2Cl_2$  (71%); g)  $K_2CO_3$ , [18]crown-6, toluene (quant.); h) DDQ/CH<sub>2</sub>Cl<sub>2</sub>, H<sub>2</sub>O (98%); j) Ph<sub>3</sub>P, DIAD, toluene (98%). Mes = mesityl = 2,4,6-trimethylphenyl, DCC = 1,3-dicyclohexyl-carbodiimide, DDQ = 2,3-dichloro-4,6-dicyano-1,4-benzoquinone, DIA-D = diisopropylazodicarboxylate.

secured through oxidative removal of both the *p*-methoxybenzyl groups and ring closure under Mitsunobu conditions. This method of piperidine formation is unusual for C–N bond formation, and we were pleasantly surprised at its efficiency. However, there is ample precedent for such transformations.<sup>[15]</sup> The spectral data for synthetic (+)-phyllantidine correspond to those reported in the literature, including the sign of optical rotation. Analysis by HPLC on a chiral stationary phase indicated no trace of the (–) isomer.

In summary, we have succeeded in preparing for the first time the structurally unusual and demanding alkaloid phyllantidine using a homo [3+2] dipolar cycloaddition. The overall yield of the natural product is around 6% over 12 synthetic operations from the cycloaddition. Efforts to adapt this protocol (through N–O bond reduction and ring closure to form a pyrrolidine) to other *securinega* alkaloids are in progress.

Received: June 27, 2006 Published online: September 13, 2006

**Keywords:** alkaloids  $\cdot$  cycloaddition  $\cdot$  nitrone  $\cdot$  oxazine  $\cdot$  total synthesis

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