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Bouillon, Marc; Bertocco, Katia; Bischoff, Laura; Buri, Michelle; Davies, Lucy Rebecca; Wilkinson, Elizabeth Jane; Lahmann, Martina

European Journal of Organic Chemistry

DOI:
10.1002/ejoc.202001317

Published: 31/12/2020

Peer reviewed version

Cyswllt i'r cyhoeddiad / Link to publication

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Accepted Article

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**Authors:** Marc Etienne Bouillon, Katia Bertocco, Laura Bischoff, Michelle Buri, Lucy Rebecca Davies, Elizabeth Jane Wilkinson, and Martina Lahmann

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**To be cited as:** Eur. J. Org. Chem. 10.1002/ejoc.202001317

**Link to VoR:** https://doi.org/10.1002/ejoc.202001317
Synthesis of Anemoclemosides A and B, two Saponins Isolated from Anemoclema glaucifolium


Abstract: Steroidal and triterpenoid saponins are attractive for their wide-ranging pharmacological properties. The triterpenoid saponins Anemoclemoside A and B are root constituents of the Chinese folk medicinal plant Anemoclema glaucifolium (Ranunculaceae). Both compounds feature an unusual cyclic acetal linkage to the carbohydrate L-arabinose in its open chain form rather than the typical glycosidic bond present in normal saponins. The straightforward and scalable syntheses of both saponins starting from L-arabinose as well as L-lyxose and L-rhamnose are described.

Saponins are a diverse group of natural products widely distributed in the plant kingdom as well as some marine organisms. Their structure is characterized by comprising a triterpene or steroid aglycone and one or more carbohydrate side chains.[1] Both steroidal and triterpenoid saponins form an interesting class of compounds due to their pharmacological and biological properties and novel synthetic approaches are frequently reported.[2,3,4,5] In 1995, Yamasaki et al. reported the isolation and characterization of two unusual triterpenoid saponins from the roots of Anemoclema glaucifolium (Ranunculaceae), a Chinese folk medicinal plant growing at altitudes between 1600 to 3000m in the Yangtse River valley region of China.[6] Accordingly named Anemoclemosides A and B, these saponins feature an unprecedented cyclic acetal linkage of the triterpene hederagenin to the carbohydrate L-arabinose in its open chain form rather than the typical glycosidic bond present in usual saponins. About 20 years later, Anemoclemoside A was also identified by Kirmizizgil and co-workers to be a constituent in the aerial parts of Cephalaria elazigensis var. purpurea, a perennial medicinal herb belonging to the Dipsacaceae family, widely distributed in southwestern Anatolia, Turkey.[7] Anemoclemosides A and B are structural isomers of the hederagenin glycosides δ- and α-hederin, two saponins first isolated and characterized from the leaves of common English Ivy (Hedera helix, Araliaceae) by Hostettmann.[8] They were later also found in a multitude of other unrelated plant sources. The glycosidic linkage of saponins is considered as an important feature for their pharmacological and biological properties. However, it is susceptible to enzymatic cleavage under physiological conditions, and loss of activity due to loss of the glycon part has been reported.[9] The unusual open chain linkage of the Anemoclemosides might not be recognized by common glycosidases and consequently enhance their stability against enzymatic cleavage. To further investigate this feature, larger quantities of this type of saponins were required. Hence, here we present a straightforward and scalable approach for the synthesis of Anemoclemoside B as well as the first synthesis of Anemoclemoside A.

A first synthesis of Anemoclemoside B was reported by Yu et al. in 2005.[10] Their synthesis included the formation of the cyclic acetal glycosidic linkage between hederagenin and its disaccharide side chain via a TMS triflate promoted condensation reaction at -78 °C. Given the optimal geometry of the diol moiety of hederagenin to form chair-shaped cyclic acetals and ketals under acidic catalysis as observed with simple carbonyl compounds like benzaldehyde, we expected the acetal coupling to proceed in the presence of an appropriate Bronsted acid catalyst like PTSA at rt. As a proof of concept, we tested this on the synthesis of Anemoclemoside A with the simpler L-arabinosylidene acetal.
The synthesis of Anemoclemoside A started with the preparation of 2,3,4,5-tetra-O-benzyl-L-arabinose (3) as the glycon building block (Scheme 1). Commercially available L-arabinose was first converted into its open-chain diethyl dithioacetal 1 with ethanethiol in conc. hydrochloric acid followed by O-benzylation with NaH, BnBr, and TBAI. Cleavage of the thioacetal in 2 with mercury(II) chloride in the presence of mercury(II) oxide then provided the fully benzylated L-arabinose aldehyde 3 in 61% yield over the three steps. Like Yu and co-workers,10 we chose the benzyl ester of hederagenin (BnHed) as triterpene building block and coupling partner for the acetal condensation, since unprotected hederagenin is poorly soluble in the reaction solvent dichloromethane. The requisite hederagenin was obtained in multi-gram quantities via the hydrolysis of ivy saponins which, in turn, had been isolated from an ethanolic ivy fruit extract.11 Following a procedure by Eldrige et al.,12 hederagenin was then treated with benzyl bromide in DMF in the presence of potassium carbonate to afford BnHed in 87% yield. For the construction of the cyclic acetal bond, aldehyde 3 and a slight excess (1.5 equiv.) of BnHed were treated with a catalytic amount of PTSA at ambient temperature to give the coupling product 4 in 91% yield. According to NMR, a single diastereoisomer with the sugar chain in the thermodynamically favored equatorial position was obtained. In a test reaction, Yu et al. observed the formation of an L-arabinosylidene acetal at -78 °C with the tetroal moiety in the axial position. However, they were able to equilibrate to the favored equatorial acetal in the presence of a catalytic amount of TMS triflate at room temperature.10 The hydrogenolytic cleavage of all benzyl groups over Pearlman catalyst concluded the synthesis, eventually providing Anemoclemoside A in 46% overall yield over 6 steps from L-arabinose and hederagenin (Scheme 1). As anticipated, the double bond between C-12 and C-13 in the triterpene skeleton was not affected during the deprotection.15

Next, we turned to the more challenging synthesis of Anemoclemoside B. As glycosyl donor, 2,3,4,5-tetra-O-acetyl-L-rhamnopyranosyl trichloroacetimidate (5) was synthesised over three steps according to standard procedures (Scheme 2).13 The glycosyl acceptor required an 1,3,4,5-O-protected L-arabinose building block with a free OH group at C-2 to be linked to the L-rhamnose donor.

While the thioacetal was convenient as aldehyde precursor in the synthesis of Anemoclemoside A, it is not compatible with the conditions applied in glycosylation reactions with trihaloacetimidates, as Yu and co-workers already demonstrated.10 Thus, we decided to use orthogonally protected 3,4,5-tri-O-benzyl-1-O-TBDDS-L-arabinol (11) as acceptor and to regenerate the aldehyde functionality later in the synthesis. Yet, since the conversion of L-arabinose into this building block requires a 10-step synthesis with a total yield of about 20%, as reported for the 5-O-acetylated 3,4-di-O-benzyl-1-O-TBDDS-L-arabinol used in the first Anemoclemoside B synthesis,10 we decided to investigate the preparation of 11 starting from L-lyxose instead (Scheme 3).
The synthesis of Anemoclomoside B started with the conversion of L-lyxose into its open-chain diethyl dithioacetal 6 in 81% yield. The following regioselective introduction of the 4,5-O-isopropylidene acetal, however, proved to be challenging. According to Redlich et al., the kinetically controlled reaction of thioacetal 6 with acetone in the presence of 2 N HCl at 0–5°C should give the desired 4,5-O-isopropylidene acetal within a couple of hours as the major product. Yet, these conditions preferentially produced the thermodynamically favored 3,4-O-isopropylidene acetal as the major product accompanied by the desired 4,5-derivative and other acetals in minor quantities. Consequently, we performed a series of experiments to find suitable reaction conditions to produce the desired 4,5-acetonide as the main product. Lowering the reaction temperature to 0°C had only an effect on the reaction rate but no significant effect on the regioselectivity. Strong acid catalysts (PTSA, CSA) gave complex product mixtures that were separable by column chromatography but produced the desired 4,5-acetonide only in moderate yields (max. 40%). Side products included the 3,4- and 3,5-O-monooisopropylidene acetals. Similar results were obtained when neat acetone or acetone in combination with 2,2-dimethoxypropane or 2-methoxypropene were used. However, when a solution of thioacetal 6 was treated with weakly acidic PPTS in neat 2,2-dimethoxypropane for 2 hours, the desired 4,5-O-isopropylidene acetal 7 was obtained in 70% yield. Longer reaction times led to...
a decrease of the 4,5-acetonide in favor of the 3,4- and 3,5-acetonides.

The conversion of 7 into the O-benzylated acetonide 10 was initially planned via a three-step sequence, i.e. hydrolysis of the thiocetal group and reduction of the intermediate aldehyde followed by perbenzylolation 4,5-O-isopropylidene-L-lyxitol to yield acetonide 10. However, all attempts to cleave the thiocetal under various conditions resulted in the complete decomposition of the starting material. Therefore, we decided to benzylation the two remaining free hydroxyl groups in 7 prior to the thiocetal cleavage. Thiocetal derivatives of pentoses like 8 are known to give the corresponding protected aldehydes in hydrolysis reactions in good yields. Thus, diol 7 was benzyl protected under standard conditions affording the fully protected L-lyxose derivative 8 in 92% yield. Initial hydrolysis attempts with HgCl$_2$/HgO and NBS resulted in partly concommitant cleavage of the acetonide moiety. Nevertheless, when the hydrolysis of thiocetal 8 was performed with NBS/AgNO$_3$ buffered with 2,6-lutidine, alcohol 9 was obtained in 98% yield after reduction of the intermediate aldehyde with NaBH$_4$. Benzylation of the primary alcohol then yielded the desired acetonide 10 in 97%.

The remaining synthesis steps proceeded smoothly. Hydrolysis of the isopropylidene acetal with dilute aqueous sulfuric acid at 90 °C followed by selective TBDDS protection of the primary alcohol eventually gave the requisite L-arabinol acceptor 11 in 43% yield over 8 steps. L-Arabinol 11 was then coupled with the L-rhamnose donor 5 in the presence of TMS triflate as Lewis acid catalyst affording disaccharide 12 in a yield of 91% as a single diastereoisomer. Silyl ether 12 was subsequently transformed into the required aldehyde 14 in two steps. First, the TBDDS group was cleaved with TBAF in the presence of excess acetic acid as buffer to prevent acetate migration followed by oxidation of alcohol 13 with TEMPO/BAIB to yield the desired aldehyde 14 in 74% over two steps. The coupling of disaccharide aldehyde 14 with BnHed was performed under the same conditions as applied in the synthesis of Anemoclemoside A affording the fully protected Anemoclemoside B precursor 15 in 94% yield. A two-step deprotection sequence concluded the synthesis of Anemoclemoside B. First, the acetate groups of coupling product 15 were saponified with dilute aqueous NaOH keeping the benzyl ester intact. The crude semi-deprotected saponin was then debenzylated over Pearlman catalyst with excess hydrogen yielding the desired saponin in 91%. Spectroscopic data of the synthetic saponins matched the data reported for the natural products by Yamasaki et al. In summary, the saponins Anemoclemoside A and B were synthesised in overall yields of 46% and 18% over 6 and 18 steps, respectively. A mild and facile method was employed to construct the characteristic cyclic acetal glycosidic linkage in both saponins via the PTSA catalyzed condensation of benzyl hederagenate with the respective saccaride side chains. 2.3 g of Anemoclemside A and 7.5 g of Anemoclemoside B were produced in straight forward procedures demonstrating the scalability of this approach.

Acknowledgements

The authors gratefully acknowledge the financial support by the BEACON initiative, Innovate UK, Bangor University, and the Erasmus+ program (M. Buri).

Keywords: Saponins • Anemoclemosides • Glycosylation • Hederagenin glycosides • Cyclic acetal glycosidic linkage


The straightforward and scalable syntheses of the triterpenoid saponins Anemoclemoside A and B are described. Both compounds feature an unusual cyclic acetal linkage to the carbohydrate L-arabinose in its open chain form. A mild and facile method was employed to construct the characteristic cyclic acetal glycosidic linkage in both saponins via the PTSA catalyzed condensation of benzyl hederagenate with the respective saccharide side chains.