The isolation of isoagathenediol: a new tricyclic diterpene from the lipids of *Rhodospirillum rubrum*

Jo-Anne Chuck† and Kevin D. Barrow

Author for correspondence: Jo-Anne Chuck. Tel: +45 33 27 5229. Fax: +45 33 27 4766.
e-mail: chuck@biobase.dk

INTRODUCTION

Rhodospirillaceae are photosynthetic non-sulphur bacteria which are found in fresh water, marine and hypersaline environments such as lakes, marshes and paddy fields (Imhoff & Trüper, 1989). Members of this family are thought to be unique amongst contemporary organisms as they contain the genetic material for all or most modes of energy metabolism (Youvan & Bylina, 1989). For example, *Rhodospirillum rubrum* is able to grow as a phototrophic or photoautotrophic under anaerobic conditions in the light, as well as being capable of microaerophilic or aerobic growth in the dark (Trüper & Imhoff, 1989).

Terpenoid compounds play important roles in the photosynthetic processes of *R. rubrum*. The light harvesting bacteriochlorophyll *a* from this organism has mainly geranylgeraniol, and to a minor extent phytol, esterified to the porphyrin (Jones, 1978). This is in comparison with most other species which possess phyto! and farnesol (Brockman & Knobloch, 1972). Carotenoids of the spirilloxanthin series (Schmidt, 1978) and membrane fluidity modulators (hopanoids) are also found as lipid constituents (Rohmer et al., 1984).

This investigation reports the isolation of a new class of diterpene, isoagathenediol, in the lipids of this organism. This is also the first report of the isolation of this compound as a natural product. The structure of isoagathenediol is such that it could function as a membrane fluidity modulator in a similar manner to the hopanoids. Its isolation as a prokaryotic lipid indicates the origin of some of the diterpene biomarkers which are detected in oil and shale sediments (Anders & Robinson, 1971; Chicarelli et al., 1988).

METHODS

Organism. *Rhodospirillum rubrum* Van Neils strain S1 (UNSW 009500) was obtained from the School of Microbiology University of NSW, Kensington, Australia. The organism was cultured anaerobically in the medium of Ormerod et al. (1961) in four 20 l carboys at 30°C under the illumination of tungsten lamps for 10 d. The cells were harvested by centrifugation (5000 g, 10 min at 5°C), washed and freeze-dried (90 g dry wt).

Isolation of isoagathenediol monoacetate. A lipid fraction obtained by refluxing the cells in CHCl₃/CH₃OH (2:1, v/v) for 2 × 1 h and the evaporated nitrates were acetylated using pyridine/acetic anhydride (1:1, v/v) for 18 h at room temperature. The evaporated extract (18 g) was dissolved in CHCl₃ and separated by flash chromatography (Still et al., 1978) using a column of Merck 60H silica gel (14 × 8 cm i.d.). The column was equilibrated with hexane and fractions (200 ml) of the following solvents were passed sequentially through the column under the suction of a water pump and collected separately: hexane, hexane/ethyl acetate (95:5, v/v), hexane/ethyl acetate (9:1, v/v), hexane/ethyl acetate (4:1, v/v), hexane/ethyl acetate (3:1, v/v), hexane/ethyl acetate (7:3, v/v), hexane/ethyl acetate...
(1:1, v/v), ethyl acetate and CH$_2$OH. Thin layer chromatography (TLC) was carried out using Polygram SIL-HR/UV254 plates (Macherey-Nagel) developed in paper-lined tanks using toluene/ethyl acetate (95:5, v/v) and visualized by spraying with 50% (v/v) H$_2$SO$_4$ and charring on a hot plate (diterpene acetate R$_f$ 0.3).

Fractions 5 and 6 were pooled and further purified using a rotary Chromatotron (Model 7924T, Professional Technology Pty Ltd) and a 2 mm silica gel (Merck, 60 PF$_{38}$) rotor using toluene as a solvent (50 ml). Fractions (4 ml) were collected and analysed by TLC. Fractions 9 and 10 were further purified by HPLC using a Whatman Partisil M9 10/50 normal phase column with 50% (v/v) H$_2$SO$_4$, a Waters Differential Refractometer R401 detector and a solvent system of toluene/ethyl acetate (6:4, v/v) at a flow rate of 12 ml min$^{-1}$ (diterpene acetate, retention time 15.5 min). A Gold Pak EXSIL 100, 5 μm silica gel column (Actiavan) using a solvent system of toluene/ethyl acetate (8:2, v/v, 1 ml min$^{-1}$, retention time 4 min) was used as a final purification step. The compound (16 mg) was recrystallized from hot acetonitrile and purity assessed by gas chromatography using a Hewlett Packard 5890 GC with a 15 m SE-30 capillary column (Alltech), a H$_2$ flow rate of 12 ml min$^{-1}$, an oven temperature which increased from 200 to 280 °C at 10 °C min$^{-1}$ and flame ionization detection (FID) (retention time 7.2 min).

**Preparation of isoagathenediol.** Isoagathenediol monoacetate (4 mg) was deacetylated by dissolving in CH$_3$OH (0.5 ml) and adding 2 M NaOH (0.5 ml). The reaction was carried out at 30 °C for 2 h and then acidified with 1 M HCl. After extraction with CHCl$_3$ (4 x 1.5 ml), the reaction products were analysed by normal phase TLC (isoagathenediol, R$_f$ 0.1). Final purification of the diol was by HPLC in the previously described system using the Gold Pak 5 μm silica column (retention time of the diol 9 min).

**Physical analyses.** Infra red spectra were recorded in CHCl$_3$ on a Perkin–Elmer 580B infra red spectrometer. Melting points were carried out using a Buchi type S melting point apparatus. Optical rotations were determined in CHCl$_3$ using a Perkin–Elmer 141 polarimeter.

Chemical ionization (CI) spectra using methane as the ionization gas were recorded using a Finnigan Model 3200 mass spectrometer. Isoagathenediol monoacetate CI positive ion spectrum (direct insertion): m/z: 351 (2%), 333 (20%), 317 (6%), 281 (5%), 273 (65%), 263 (7%), 257 (12%), 207 (30%), 205 (15%), 171 (13%), 163 (15%), 149 (65%), 61 (100%). Electron impact ionization (EI) mass spectra were recorded using a VG AUTOSPEC-Q mass spectrometer operating at a filament emission energy of 70 eV and a source temperature of 180 °C. Isoagathenediol monoacetate EI positive ion spectrum (direct insertion): m/z: 290 (12%), 275 (7%), 205 (10%), 191 (43%), 137 (6%), 123 (15%), 95 (17%), 69 (37%).

$^1$H and $^{13}$C NMR spectra were recorded on a Bruker AM 500 spectrometer operating at 500 MHz and 125 MHz, respectively. Spectra were recorded on samples containing CDC$_3$ and chemical shifts were expressed in p.p.m. downfield from Me$_4$Si. In the case of the $^{13}$C spectra, the reference was CDC$_3$, with the centre of the triplet resonating at 77.0 p.p.m. relative to Me$_4$Si. $^1$H–$^1$H homonuclear shift correlated (COSY) spectra were recorded using 90° and 45° pulses of 12 and 6 μs, respectively, and a recycle time of 2 s. Homonuclear Hartman Hahn transfer (HOHAHA) spin locked experiments were run using the MLEV-17 sequence (Bax & Davis, 1985) with mixing times of 65–70 ms and 90° and 180° pulse widths of 35 and 70 μs, respectively, and a recycle delay of 2 s. Distortionless enhancement by polarization transfer (DEPT) spectra were run with the standard Bruker program. Carbon–proton correlations (heteronuclear shift correlated 2D) were carried out using the sequence of Wilde & Bolton (1984). The 90° $^1$H pulse width was 11 μs and the carbon 90° and 180° pulses were 5-5 and 11 μs, respectively. A heteronuclear J-resolved 2D NMR spectrum was obtained with 90° and 180° carbon pulses of 5-5 and 11 μs, respectively. A long range carbon–proton correlation experiment was carried out using the sequence of Reynolds et al. (1985), a 2 s recycle time and delays of 0:011 s and 0:22 s.

**RESULTS**

Isoagathenediol monoacetate [melting point 159 °C (corrected), literature 159–161 °C; Imamura & Ruveda, 1980; Fig. 1] was isolated from the acetylated lipids of *R. rubrum* (16 mg per 90 g dry wt of cells). The compound could not be detected by TLC of the crude acetylated lipid extract of the organism. Partial purification of the extract...
Table 1. $^{13}$C and $^1$H NMR assignments for isoagathenediol monoacetate

The numbering of the carbon atoms is shown in Fig. 1.

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using flash chromatography allowed the diterpene to be detected. The compound gave a red/purple colour on the plate after spraying with H$_2$SO$_4$ and charring and this was quite distinct from the purple colour that the hopanoids and squalene develop after similar treatment. The compound did not absorb light at 254 nm.

Isoagathenediol monoacetate was detected in the crude acetylated lipid extract of the organism by capillary GC using FID. The same extract, when chromatographed before acetylation, had no peaks in the chromatogram corresponding to the compound (results not shown), thus the diterpene must exist in the lipids of the organism in the unacetylated form. The structure of the compound was determined by extensive use of NMR spectroscopy and high resolution mass spectrometry.

NMR analyses

The proton NMR spectrum of isoagathenediol monoacetate indicated that the compound contained six quaternary methyl groups. A resonance at 1.20 p.p.m. was consistent with a methyl group close to an electron withdrawing group. The resonance at 2.08 p.p.m. indicated the presence of a methyl group of an acetate moiety. The compound also contained two methylene protons resulting in resonances at lower field centred at 4.33 p.p.m. The other proton resonances in the spectrum occurred in the chemical shift range of 0.8–2.4 p.p.m. (Table 1).

The $^{13}$C spectrum of isoagathenediol monoacetate depicted 22 carbon resonances and, analogous to the $^{13}$C spectra of other acetylated terpenes (Nishizawa et al., 1986), contained resonances at 171.3 p.p.m. and 21.3 p.p.m. These were the carbonyl and methyl carbon resonances of the acetoxy group added to the compound during derivatization. Due to the similarities of the spectrum with that of other cyclic terpenes, it appeared that the compound had a typical terpenoid A/B ring system (Wehrli & Wirthlin, 1976).

DEPT experiments indicated that the unacetylated compound had eight CH$_3$, five methyl groups, four quaternary carbons and three CH groups. This was consistent with the results obtained from a 2D $J$-resolved heteronuclear experiment which also indicated that the coupling constants were all in the usual range of 125–150 Hz and thus excluded the existence of strained or small rings in the molecule (Wehrli & Wirthlin, 1976). Assignment of the proton spectrum of the acetate derivative was carried out using the results of $^{13}$C–$^1$H correlation experiments and on the basis of the carbon assignments by Nishizawa et al. (1986) for unacetylated compounds (Table 1).

A COSY spectrum of the molecule indicated that the two lower field protons at 4.29 and 4.37 p.p.m. were coupled to each other and to a proton at 1.55 p.p.m. These protons were assigned to C-15 and were coupled to H-14. A long range coupling was also seen from the two protons at C-15 to the methyl protons of the acetoxy group. The results of a HOHAHA experiment showed three spin systems in the ring proton region of the spectrum. With the results of the HOHAHA experiment and the DEPT experiments, seven of the CH$_3$ groups were assigned to the three spin systems and the eighth to the carbon bearing the acetoxy group. Two of the CH groups were also involved in the spin systems of the ring protons while the third CH group (C-14), which was surrounded by quaternary ring carbons, showed coupling to the CH$_3$ (C-15) bearing the acetoxy group. The long range $^{13}$C–$^1$H correlation experiment clearly confirmed the structure of the molecule around carbons 13, 14 and 15 and allowed for the interconnection of the spin systems detected in the HOHAHA experiment.

Mass spectrometry and other analyses

High resolution mass spectrometry was carried out on both the acetylated and unacetylated diol giving rise to very similar mass spectra. No molecular ion was evident in the spectra of the molecules using EI fragmentation; however, a pseudomolecular ion of low intensity ($m/\text{x} 351$, [M + 1]$^+$, 2 %) was evident in the CI spectra of the acetate derivative.

By EI, both compounds had a similar mass spectrum as the monoacetate derivative lost the acetate moiety and then both compounds lost water, presumably at the position of the tertiary hydroxyl group. This gave the ion $m/\text{x} 290$ (12 % high resolution mass measurement found $m/\text{x} 290$-2610; calculated for C$_{19}$H$_{28}$O ($\text{M-CCH}_3\text{COOH})^+$ $m/\text{x} 290$-2610) and the skeleton fragmented to give the ion $m/\text{x} 191$ (43 %) in high abundance.
An infra red spectrum of the diterpene monoacetate showed absorbance bands at 3597 and 1735 cm\(^{-1}\) which indicated the presence of a tertiary hydroxyl group and an acetate group in the molecule. On deacylation, the carbonyl band was missing and implied that the molecule contained two hydroxyl groups, of which only one was acetylated under the experimental conditions used.

**Determination of complete configuration of isoagathenediol**

The compound isolated in this investigation was identified as the enantiomer of a compound synthesized by Imamura & Ruveda (1980). Debromoisoaplysin 20-monoacetate (Fig. 1) had an identical melting point and \(^1\)H NMR spectrum to the compound isolated in this investigation. The optical rotation of isoagathenediol monoacetate ([\(\alpha\)]\(_D\) = 3-7; \(a = 0.004)\) though the same in magnitude, was laevorotatory rather than dextrorotary. From the NMR spectrum of the synthetic compound 13-epi-debromoisoaplysin 20-monoacetate (Fig. 1), the stereochemistry of the hydroxyl group compared with the carbon at C-15 was determined. The 1D \(^1\)H NMR spectrum of this compound had the resonance of the methyl group at C-13 at 1-02 p.p.m. compared with the diterpene isolated in this investigation where the methyl group was at 1-20 p.p.m.

Nishizawa et al. (1986) also synthesized the diol (spongial-13\(\alpha\),15-diol) as a racemic mixture. They synthesized various isomers of the compound with isomeric methyl groups and hydroxyl groups in ring C. The comparison of the \(^{13}\)C spectrum of the diterpene isolated in this investigation with the spectra of the analogue allowed for the determination of the stereochemistry of isoagathenediol (13\(\alpha\) OH, 14 \(\alpha\) H).

**DISCUSSION**

Isoagathenediol is a new microbial lipid and is not a previously known natural product. It has, however, been chemically synthesized (Nishizawa et al., 1986; Vlad et al., 1986) and named according to Vlad et al. (1986) based on the nomenclature of isoagatholactone. Previous to this study, the only known microbial diterpenes were the carotenoid intermediate, geranylgeraniol pyrophosphate (Langworthy, 1982), a chemically uncharacterized gibberellin from *Azotobacter vinelandii* (Lee et al., 1970) and verrucosan-2\(\beta\)-ol from *Chloroflexus aurantiacus* (Hefter et al., 1993).

Five other prokaryotes, including those known to contain hopanoids, were screened for the presence of the diterpene by GC of crude acetylated lipid extracts (results not shown). In acetylated extracts derived from *Rhodopsseudomonas palustris*, *Alicyclacoccus acidocaldarius*, *Zymomonas mobilis*, *Rhodobacter capsulatus* and *Rhodopsseudomonas viridis* the diterpene could not be detected (< 10 \(\mu\)g per 15 mg of lipid extract). This may be due to the composition of bacteriochlorophyll \(a\) of *R. rubrum* having geranylgeraniol as a constituent (Jones, 1978). It is likely that this molecule or geranylgeraniol pyrophosphate would be the substrate for the cyclization reaction biosynthesizing the diterpene.

The cyclase giving rise to this diterpene appears to be distinct from those required for hopanoid and tetrahymanol biosynthesis. The substrate folding for the active site of the enzyme would require an all pre-chair folding in preparation for an 'all Markovnikov' reaction (Ourisson & Rohmer, 1982). Renoux & Rohmer (1986) have demonstrated the cyclization of hexaprenol methyl ether to an extended tricyclic product in a cell-free system of *Tetrabymena pyriformis*. It was assumed that the squalene cyclase was responsible for the synthesis and this was supported by the fact that the cyclization of both the enantiomers of squalene epoxide was demonstrated by the preparation indicating a lack of substrate specificity. The stereochemistry of these products at the side chain of the molecule was different to the diterpene isolated in this investigation. It was assumed that the substrate had adopted a chair, chair, boat conformation in the active site of the cyclase resulting in an axial side chain (Renoux & Rohmer, 1986). This suggests that isoagathenediol is cyclized by a different enzyme to the squalene cyclases.

As terpenoid compounds are very recalcitrant they are often detected in sediment samples such as oil or shale and are used to determine the source and age of organic matter making up the sediment. Geochemical analysis of Irati shale has shown that \(C_{20}\) diterpene isomers 13\(\beta\) H, 14\(\alpha\) H and 13\(\alpha\) H, 14\(\alpha\) H are abundant, with approximately equal quantities of the isomers (Chicarelli et al., 1988). This would be consistent with the stereochemistry of isoagathenediol (13\(\alpha\) OH, 14 \(\alpha\) H) which, via the maturation process of the sediment, would rapidly dehydrate at the position of the tertiary hydroxyl group. Thus the stereochemistry of the methyl group at C-13 would be lost and the double bond between C-12 and C-13 could be saturated above or below the plane of the ring to give the stereochirnistry of the two predominate isomers. The generation of isomers of sterols in sediments via the hydrogenation of ring double bonds is well documented (Mackenzie et al., 1982). This investigation again reinforces the importance of the characterization of lipid components of organic sediments to gain knowledge about existing lipids.

Isoagathenediol is a planar compound with the hydroxyl groups at one end of the molecule. According to Ourisson & Rohmer (1982), for molecules to optimize the mechanical properties of a membrane, the compound must have a hydrophilic head and a lipophilic body. To enable close packing with the acyl chains of lipids, the molecule must be approximately 6 Å thick and 6 Å wide. Isoagathenediol fulfills this criterion as it is approximately 6 Å high, 6 Å wide and 11 Å long (dimensions determined using a scaled model). Like the hopanoids, this compound could be incorporated into the membranes of *R. rubrum* in a sterol-like manner and function as a membrane-reinforcing molecule. This has yet to be confirmed by biophysical analyses.

Prior to this study, structurally related tricyclic diterpenes had not been isolated from a biological source even...
though they were readily detected in sediment samples and so it was assumed that the molecules were derived from unknown or ancient organisms (Ourisson & Rohmer, 1982; Brassell et al., 1983). If the evolutionary model of Rohmer et al. (1979) is accepted, with membrane stability maintained by carotenoids and tetraterpenes prior to the evolution of a cyclase, the enzyme would be likely to use a carotenoid or tetraterpene precursor as a substrate. Isoagathenediol may thus be more primitive than hopanoids as its biosynthesis does not require the synthesis of squalene but an intermediate from carotenoid biosynthesis, geranylgeraniol. It is likely that the diterpene, however, acts as a less effective membrane stabilizer due to the reduced length of its hydrophobic region and thus evolution has selected for the squalene cyclase in most bacteria.

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REFERENCES


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