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To cite this article: Alina Owczarek, Renata Kaczmarek, Barbara Mikołajczyk, Ewa Wasilewska, Dariusz Korczyński, Janina Baraniak, Maria Koziółkiewicz, Wojciech J. Stec & Charles Brenner (2003) Stereochemical Analysis of Diastereomeric 1,3-bis(Adenosine-5'-O-phosphorothioyl)glycerols, *Nucleosides, Nucleotides and Nucleic Acids*, 22:5-8, 797-799, DOI: [10.1081/NCN-120022637](https://doi.org/10.1081/NCN-120022637)

To link to this article: <https://doi.org/10.1081/NCN-120022637>



Published online: 31 Aug 2006.



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Stereochemical Analysis of Diastereomeric 1,3-bis(Adenosine-5'-O-phosphorothioyl)glycerols

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INTRODUCTION

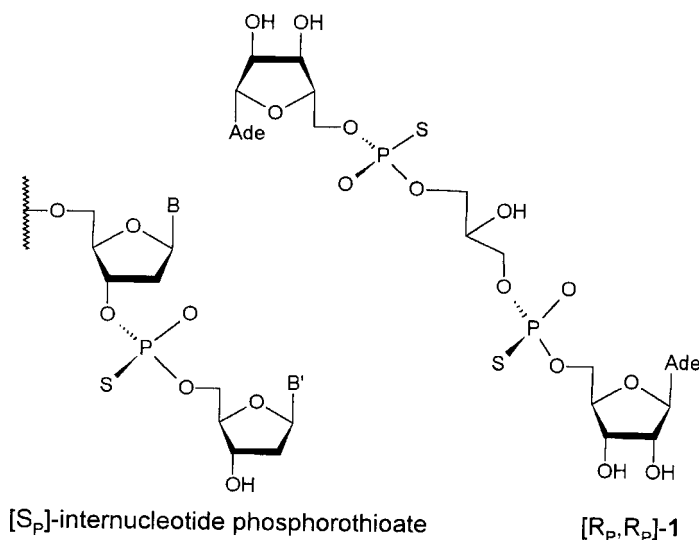
We have published recently^[1] that 1,3-bis(adenosine-5'-phosphorothioyl)glycerol (**1**), prepared according to the oxathiaphospholane methodology^[2] possesses stronger inhibitory activity towards FHIT protein, as compared with all known inhibitors.^[3] Fhit protein is the Ap₃A hydrolase which binds and cleaves diadenosine polyphosphates and acts as a tumor suppressor.^[4] Loss of Fhit protein is among the earliest known events in the development of a variety of the most common and lethal human malignancies. Function of Fhit in tumor suppression does not require diadenosine polyphosphates cleavage but correlates with the ability to form enzyme-substrate complexes. If Fhit-substrate complexes promote tumor suppression by stimulating a pro-apoptotic effector,^[5] then Fhit inhibitors, that resemble natural substrates, may promote or antagonize Fhit function, depending on their features, in Fhit + cells.



RESULTS AND DISCUSSION

In this communication we present preliminary studies concerning stereochemical analysis of compound **1**, prepared^[2] as the mixture of eight diastereoisomers (2³, two chiral centers at phosphorus and one chiral/pseudochiral center at C-2 atoms of glycerol moiety). Since interaction of **1** with proteins may be *a priori* considered as stereodependent event, attempts at separation of **1** into individual diastereoisomers by means of RP-HPLC technique were undertaken. These efforts were partially successful, and only one isomer assigned as **1a** was isolated as pure specimen; inhibitory effect of this individual isomer towards Fhit is under investigation. Independently, experiments towards assignment of absolute configuration at P atom in **1a** were performed. Diastereoisomer **1a** appeared to be resistant towards *snake venom phosphodiesterase* (*svPDE*) which is known to hydrolyse stereoselectively internucleotide 3',5'-phosphorothioates of R_P-configuration.^[6,7]

Topological analysis indicates (Sch. 1) that *svPDE*-resistant S_P-dinucleoside 3',5'-phosphorothioate, due to Cahn-Ingold-Prelog rules, corresponds to *svPDE*-resistant R_P,R_P-isomer of **1**. Such assignment is validated by observation, that diastereoisomeric mixture of **1** under treatment with *svPDE* undergoes degradation, rendering intact **1a**. From the resulting mixture besides two compounds corresponding to mono-(adenosine-5'-O-phosphorothioyl)glycerol of R_C,R_P- and S_C,R_P-configuration, adenosine-5'-O-phosphorothioate has been isolated (HPLC coinjection, MALDI-TOF MS analysis). Compound **1** (as the mixture of all possible diastereoisomers) was digested with *svPDE* in the buffer containing [¹⁸O] water, and resulting adenosine-5'-O-[¹⁸O]phosphorothioate was isolated by means of RP-HPLC. Its stereochemical analysis was performed according to the methodology developed recently in this laboratory.^[8] Taking into account that *svPDE* cleaves internucleotide



Scheme 1.

phosphorothioate of R_P-configuration with retention (involvement of covalent enzyme-substrate complex^[9]) we were able to prove that undigested by *sv*PDE isomer **1a** possesses R_P-configuration (data not shown). Studies upon inhibitory activity of **1a** towards Fhit protein are in progress and results will be published in due course.

ACKNOWLEDGMENT

This work was financially assisted by the State Committee for Scientific Research (KBN), grant no. Z-KBN K005/T09 (to W.J.S.).

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