Development of Antitumor Compounds Using the Chiral Benzaldehyde

メタデータ	言語: jpn
	出版者:
	公開日: 2022-05-16
	キーワード (Ja):
	キーワード (En):
	作成者: Hanaoka, Miyoji
	メールアドレス:
	所属:
URL	https://doi.org/10.24517/00065990
	This work is licensed under a Creative Commons

This work is licensed under a Creative Commons Attribution-NonCommercial-ShareAlike 3.0 International License.



1999 Fiscal Year Final Research Report Summary

Development of Antitumor Compounds Using the Chiral Benzaldehyde

Research Project

Project/Area Number
09470484
Research Category
Grant-in-Aid for Scientific Research (B)
Allocation Type
Single-year Grants
Section
一般
Research Field
Chemical pharmacy
Research Institution
Kanazawa University
Principal Investigator
HANAOKA Miyoji Faculty of Pharmaceutical Sciences, Kanazawa University Professor, 薬学部, 教授 (80028844)
Co-Investigator(Kenkyū-buntansha)
KATAOKA Osamu Faculty of Pharmaceutical Sciences, Kanazawa University Research Associate, 薬学部, 助手 (40303292) MUKAI Chisato Graduate School of Natural Science and Technology, Kanazawa University Professor, 大学院・自然科学研究科, 教授 (70143914)
Project Period (FY)
1997 – 1999
Keywords
chiral benzaldehyde / styryllactone / chiral aldol reaction / stereoselective reaction / chiral synthesis / lactone structure / antitumor compound

Research Abstract

1. Chirai aldol reaction of the chiral benzaldehyde with silyl ketene acetals afforded anti-aldol products with highly stereoselectivity. Thus, chirality of the chiral benzaldehyde was effectively transformed to a chiral carbon chain.

2. Aldol reaction of the chiral benzaldehyde with a titanium enolate derived form a thioester gave stereoselectively an anti-aldol product.

3. Common synthetic intermediate having four continuous asymmetric carbons was synthesized in highly stereoselective manner and highly optical yield from the chiral benzaldehyde through twice aldol reactions. Total synthesis of styryllactones having a 5-membered lactone, goniofufurone, goniobutenolide A and B was achieved.

4. Conversion of a 5-membered lactone to a 6-membered lactone through a lactol was developed. According to this method, styryllactones having a 6membered lactone, goniodiol, goniotriol, 8-acetylgoniotriol, altholactone, and 9-deoxygoniopypyrone were synthesized.

5. Chiral total synthesis of goniofupyrone was succeeded. This synthesis established the structure of goniofupyrone including absolute stereochemistry and revised the previously proposed structure.

6. The structure of gonioheptolide A was revised by its synthesis from goniofupyrone. The proposed 8-membered lactone structure was found to be incorrect.

7. Amino acid part of AI-77B, an antiulcerogenic antibiotic, and AI-77B analogue were synthesized from the chiral benzaldehyde.

8. 1,3-Dipolar cycloaddition of chiral nitrons, derived form the chiral benzaldehyde, with olefins gave stereoselectively cis-3,5-disubstituted isoxazolidines.

Research Products (6 results)

			All	Other	
	All Publications		ıs (6 ı	(6 results)	
[Publications] Chisato Mukai: "New Approach to AI-77B : Stereoselective Conversion of a Potential Precursor of the Amino Acid Side Chain"J. Chem. Soc., Perkin Trans. 1. 913-917 (1997)					
[Publications] Chisato Mukai: "Stereoselective Syntheses of (+)-Goniotriol, (+)-8-Acetylgoniotriol, (+)-Goniodiol, (+)-9-Deoxygoniopypyrone, (+)- Altholactone and (-)-Goniofupyrone"J. Org. Chem 62. 6619-6626 (1997)				~	
[Publications] Chisato Mukai: "Revised Structure of Gonioheptolide A"Chem. Pharm. Bull 47. 131-132 (1999)				~	
[Publications] MUKAI, Chisato: "New Approach to AI-77B : Stereoselective conversion of a Potential Precursor of the Amino Acid Side Chain"J. Chem. So Perkin Trans 1 · 6. 913-917 (1997)			n. Soc	···, 🗸	
[Publications] MUKAI, Chisato: "Stereoselective Syntheses of (+)-Goniotriol, (+)-8-Acetyl-goniotriol, (+)-Goniodiol, (+)-9-Deoxygoniopypyrone, (+)- Altholactone and (-)-Goniofupyrone"J. Org. Chem 62 · 19. 6619-6626 (1997))-	~	
[Publications] MUKAI, Chisato: "Revised Structure of Gonioheptolide A"Chem. Pharm. Bull 47 • 1. 131-132 (1999)				~	

URL: https://kaken.nii.ac.jp/report/KAKENHI-PROJECT-09470484/094704841999kenkyu_seika_hokoku_

Published: 2001-10-22