Dr. Vijay

ISSN 2277 - 5730 AN INTERNATIONAL MULTIDISCIPLINARY QUARTERLY RESEARCH JOURNAL

AJANTA

Volume - XII Issue - I

January - March - 2023

ENGLISH PART II / MARATHI

Peer Reviewed Refereed and UGC Listed Journal Journal No. 40776



IMPACT FACTOR / INDEXING 2020 - 6.306 www.sjifactor.com

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SOURCE SOURCE STATE STATE

S. No.	Title & Author	Page No.
1	Perception of undergraduates towards transit advertising	1-12
	Ms. Subhashini Iyengar	
	Dr. Pooja Ramchandani	
2/	Paraben Derivatives as Future Potential Drug	13-21
	Vijay D. Gangan	13-22
	Uttam Yadav	
	Kundan Mewad	
	Yogita Purohit	
	Neha Rastogi	
	Priyanka A. Save	
	Anamika Singh	
	Krishnalekha Nadar	
3/	To Study Investment Patterns and Customer Attitudes in Relation	22-32
	to Mutual Funds	23-33
	Ms. Abeda Shaikh	
	Dr.Vaishali Kothiya	
4	The Dynamics of Sustainable Development: Social change through	33-38
	Teacher Education Programme	
	Dr. Shaheen Parveen	
5	Impact of Technology on Different Leadership Styles to Support Distanced	39-45
	Working: A Study of India's Service Sector in Telangana State	
	Linga Reddy Katipalli	
6	Challenges of Human Resource Management in Current Scenario	46-50
	Ms. Saniya Shiurkar U.	
	Dr. Padmapani Bhagwan Sawai	
7	Production of Cost Effective, Biodegradable, Disposable Sanitary	51-58
	Napkins using Banana Fibres	52-58
	Dr. Vijay D. Gangan	
	Dr. Uttam Yadav	
	Yogita Purohit	
	Neha Rastogi	
	Krushnal Devlekar	
	Mrs. Priyanka A. Save	



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SOUNTENTS OF ENGLISH PART - II <</p>

S. No.	Title & Author	Page No.
8	Traditional Accounting and Modern Accounting system in India with	59-667
	Special Reference to Frauds after 2000-01	
	Shaikh Mumtaz	
9	Digital Transformation in the Economy is a Boon!	68-74
2	Dr. Prakash Laxmanrao Dompale	
10/	A Study on Students' Attitudes towards Online Shopping Platforms	75-81
10	Amit Chhotelal Gupta	
	Urmila Arjun Chauhan	
11/	Statistics and Blood Glucose Monitoring	21-86
<i>''</i>	Krishnalekha Nadar	82-86
	Vijay D. Gangan	
	Priyanka A. Save	
	Sukanya Sawant	
12//	Subliminal Advertising: Utilizing Deception to Influence	87-92
Del	Consumption Habits	
•	Mr. Frank Remedios	
13	Higher Initiative to Learn: Essential Quality for	93-96
10	Entrepreneurship Development	
	Prof. Himanshul Panchal	
14	Game Changer: The Role of Sports in Achieving Sustainable National	97-101
	Development	9
	Dr. Surayakant Pandurand Dhonde	

2. Paraben Derivatives as Future Potential Drug

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Abstract

Parabens are class of chemicals widely used as preservatives in the cosmetic and pharmaceutical industries. They are attractive preservatives in many types of formulas. These compounds and their salts are used primarily for their bacterial and fungicidal properties. They are also used as food additives. Their analogues also possess various biological activities which prompted us to synthesize few more analogues for their future application as bioactive molecules. All the synthesized compounds were characterized by 1HNMR and mass spectral data and screened for their antibacterial activity against Gram + ve and Gram – ve cultures. Few of them are showing promising antibacterial activity.

Keywords: Parabens, antibacterial, fungicidal, 1HNR, elemental analysis.

Introduction

Phenolic phytochemicals are known to exhibit anti-inflammatory, antioxidant, anticarcinogenic, antidiabetic, antiatherosclerosis and immunomodulatory activities in animals1, 2. These are mostly polyphenols known as secondary plant metabolites3 present in plant and trees. One of such compound is 4-hydroxy benzoic acid which is used as antifungal, antimutagenic, antisickling, esterogenic4 and antimicrobial5 agent. It is primarily known as the basis for the preparation of its esters, known as parabens, which are used as preservatives in cosmetics. Parabens are used for their bactericidal and fungicidal properties. They can be found in shampoos, commercial moisturizers, shaving gels, personal lubricants, topical / parenteral

pharmaceuticals, spray tanning solution, makeup and toothpaste. They are also used as food additives. In the present study, we are converting 4-hydroxy benzoic acid to methyl and ethyl paraben using conventional method and their further diversification to ester derivatives. Since methyl and ethyl paraben are naturally occurring active compounds having antioxidant and antimicrobial properties, we decided to make a library of compounds using various permutation and combinations to come up with novel ether and ester derivatives of methyl and ethyl paraben using conventional methods. The objective of this study is to condense two molecules of the same disease domain to produce more potent candidate in the same disease domain or to condense two molecules of different disease domain to produce mixed variety of those disease domain or to have drug candidate with entirely different disease domain. In the present work, we are converting 4-hydroxy benzoic acid to methyl and ethyl paraben which in turn furl converted to ether and hybrid derivatives respectively using conventional methods.

Results and Discussion

Preparation of methyl and ethyl paraben

They were prepared by refluxing 4-hydroxy benzoic acid with methanol / ethanol using sulphuric acid as a catalyst for 8 hrs. The progress of the reaction is monitored by TLC for the completion of reaction.

Work up

The reaction mixture concentrated under reduced pressure to minimum and to that 200 ml of dichloromethane + 200 ml of water was added. The aqueous layer was extracted successively with dichloromethane (2 x 100 ml). The total organic layer was washed with water (200 ml), brine (100 ml) and concentrated to yield methyl and paraben respectively which can directly used for further diversification. The general yields were 95 - 98 %

General method for the preparation of compounds (I and II)

These were prepared by following general method as depicted below.

To a stirred solution of [A] (1 eq.) in 30 ml acetone was added [B] (2.5 eq.) and stirring continued at 400 C for the next 30 min. for complete formation of K-salt. To this compound [C] (2 eq.) was added and stirring continued at 45-50° C for the next 8 hrs. The progress of the reaction is monitored by TLC for the completion of this reaction.

Work Up

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The reaction mixture filtered through Buchner funnel, wash the cake with 25 ml acetone. The total organic layer was concentrated to minimum, preadsorbed on silica gel and purified by silica gel (100 - 200 mesh) column chromatography with increase in concentration of ethyl acetate in petroleum ether. The general yields ranges between 60-70 %.

Reaction Scheme:

Ethyl 4-hydroxybenzoate

Ethyl paraben ether derivatives

Compound No.	R
I	Heptyl
II	3-Methyl butyl

Taking Isoeugenol as general example, the probable mechanism for ethers can be given as follows.

1-butoxy-2-methoxy-4-[(E)-prop-1-enyl]benzene

Compound I Ethyl 4 – heptoxybenzoate.

1H NMR (CDCl3, 400 MHz) δ ppm : 0.88 (t, J = 4.7 Hz, 3H, terminal –CH3 from heptoxy moiety), 1.2 – 1.5 (m, 8H, 4 x –CH2, methylenes from n- heptyl moiety), 1.37 (t, J = 6.9 H, 3H, from –OCH2CH3 group) , 1.7 – 2.0 (m, 2H, 1x –CH2 from n-heptyl moiety), 3.99 (t, J = 6.6 Hz, 2H, 1 x –OCH2 from n-heptyl moiety), 4.34 (q, J = 7.7 Hz, 2H, -OCH2 from -OCH2CH3 group), 6.89 (d, J = 8.7 Hz, 2H, ArH), 7.97 (d, J = 8.7 Hz, 2H, ArH) Molecular Formula C16H24O3 Pure viscous mass (0.954 gms, 60 %). Anal. Calcd. for C16H24O3 : C 72.69, H 9.15, O 18.16 Found C 72.66 , H 9.12, O18.19

Compound II Ethyl 4- isopentyloxybenzoate

1H NMR(CDCl3, 400 MHz) δ ppm- 0.98 (d, J = 6.6 Hz, 2 x -CH3, 6H), 1.37 (t, J = 6.9Hz, 3H from –OCH2CH3 group), 1.7 (m, 2H, -CH2 from 3-Methyl butane moiety), 1.85 (1), 11, CH(CH3)2), 4.0 (t, J = 6.6Hz, 2H, 1 x -OCH2 from 3-Methyl butane moiety), 4.34 (q, J = 7.7 Hz, 2H, -OCH2 from ethyl paraben moiety), 6.89 (d, J = 8.7 Hz, 2H, ArH), 7.07 (d, J = 8.7 Hz, 2H, ArH). Molecular Formula C14H2OO3. Pure viscous mass (0.894 gms, 63 %). Anal. Calcd .for C14H2OO3 : C 71.16, H 8.53, O 20.31 Found C 71.13, H 8.50, O20.34

General method for the preparation of **compounds** (III - V) These were prepared by following general method as depicted below.

To a stirred solution of Ethyl paraben (1 eq.) in dichloromethane (30 ml) was added pyridine (2.5 eq.) and cool the reaction in ice bath at 15° C. Clear solution of reaction mixture was obtained. To this, was added Acetyl chloride/ benzoyl chloride / substituted benzoyl chloride (2 eq.) at 15 - 20° C and stirred, allowed to attain the room temperature and stirring was continued for the next 24 hrs. (TLC). The organic layer was concentrated under reduced pressure to minimum, preadsorbed on the silica gel and purified by column chromatography (SiO₂, 100-200 mesh) with increase in concentration of Ethyl acetate in petroleum ether to yield pure compound. The purified compounds were unambiguously characterized by ¹HNMR and mass spectroscopy. The general yields of these reactions were ranges between 60 - 80 %.

The most significant features of this methodology are (a) good accessibility of the reagents and its stability (b) a stoichiometric amount of reagent can be used by direct weighing, avoiding excess (c) no evolution of hazardous vapors during the reaction (d) the total elimination of the use of toxic organic solvents (e) a simple experimental procedure (g) good control over the outcome of the reaction by varying the amount of reagent. The aforesaid protocol thus provides

an improved procedure for the synthesis of useful paraben derivatives having important pharmaceutical, agricultural and other physicochemical properties.

Reaction Scheme:

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Ethyl 4-hydroxybenzoate

Ethyl paraben ester derivatives

Compound No.	R
III	Acetyl
IV	Benzoyl
V	4-Chloro benzoyl

The probable mechanism for esters can be given as follows.

Probable mechanism for Esters:

Compound III

Ethyl 4 – acetoxybenzoate

1H NMR (CDCl3, 400 MHz) δ ppm: 1.3 (t, J = 7.7 Hz, 3H, from - OCH2CH3), 2.31 (S, 3H, from >COCH3 group), 4.37 (q, J = 7, 14 Hz, 2H, from - OCH2CH3), 7.16 (d, J = 8.4 Hz, 2H, ArH), 8.07 (d, J = 8.4 Hz, ArH). Molecular Formula C11H12O4. Pure viscous mass (0.90 gms, 72 %). Anal. Calcd. for C11H12O4 : C 63.45, H 5.81, O 30.74 Found C 63.41, H 5.78, O 30.78

Compound IV

Ethyl 4 - benzoyloxybenzoate

1HNMR (CDCl3, 400 MHz) δppm: 1.4 (t, J = 4.7 Hz, 3H, -CH3 from –OCH2CH3 group), 4.34 (q, J=7.7 Hz, 2H,-OCH2 from Etyl paraben moiety), 7.2 - 8.4 (m, 9H, Ar-H). Molecular Formula C16H14O4. Off white solid (0.975 gms, 64 %); Melting range 65 – 680C; Anal. Calcd .for C16H14O4: C 71.10, H 5.22, O 23.68 Found C 71.06, H 5.19, O 2

Compound V

Ethyl 4-(4-chlorobenzoyl) oxybenzoate

1H NMR (CDCl3, 400 MHz) δppm: 1.4 (t, J = 4.7 Hz, 3H, -CH3 from –OCH2CH3 group), 4.34 (q, J = 7.7 Hz, 2H, OCH2 from ethyl paraben moiety), 7.0 - 8.7 (m, 8H, ArH). Molecular Formula C16H13ClO4. Off white solid (1.34 gms, 74 %). Melting range 100 - 1040C Anal. Calcd .for C16H13ClO4: C 63.06, H 4.30, O 21.00 Cl 11.63 Found C 63.10, H 4.26, O 21.03

General method for the preparation of compounds (VI to IX):- These were prepared by the general method as mentioned for ethyl paraben.

Reaction Scheme:

Methyl-4-hydroxybenzoate

Methyl paraben ester derivatives

Compound No.	R
VI	Acetyl
VII	Benzoyl
VIII	4-Chloro benzoyl
IX	2-Chloro Benzoyl
11.1	2 Cinoro Benzo.

The general mechanism for ester is same as depicted for ethyl paraben.

Compound VI

H

H3

H).

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Methyl 4-acetoxybenzoate

1HNMR (CDCl3, 400 MHz) δppm: 2.3 (s, 3H, >CO-CH3), 3.9 (s, 3H, -COOCH3), 7.16 (d, J = 8.4 Hz, 2H, ArH), 8.07 (d, J = 8.4 Hz, ArH). Molecular Formula C10H10O4; White solid (1.297 gms, 68 %). Melting range 70 - 750C. Anal.Calcd .for C10H10O4: C 61.85, H 5.19,O 32.96 Found C 61.81, H 5.16, O 32.99

Compound VII

Methyl 4-benzoyloxybenzoate

1HNMR (CDCl3, 400 MHz) δ ppm: 3.9 (s, 3H, from -COOCH3 group), 7.22 (d, J = 8.4 Hz, 2H, ArH), 8.07(d, J = 8.4 Hz, ArH from methyl paraben moiety), 7.2 - 8.3 (m, 5H, ArH from Benzoyl Moiety). Molecular Formula C15H12O4; White solid (1.09 gms, 65 %). Melting range 106 - 1100C Anal. Calcd .for C15H12O4 C 70.31, H 4.72, O 24.97. Found C 70.34, H 4.69, O 24.95.

Compound VIII

Methyl 4-(4-chlorobenzoyl)oxybezoate

1H NMR (CDCl3, 400 MHz) δppm: 3.9(s, 3H,-OCH3 from –COOCH3 group), 7.22(d, J = 8.4 Hz, 2H, ArH), 8.1 (d, J = 8.4 Hz, 2H, ArH from Methyl paraben moiety), 7.2 - 8.2 (m, 4H, ArH from 4-Chloro benzoyl moiety). Molecular Formula C15H11ClO4. Off white solid (1.297 gms, 68 %). Melting range 126 - 1300C. Anal. Calcd .for C15H11ClO4: C 61.98, H 3.81, O 22.02, Cl 12.20 Found C 61.95, H 3.78, O 22.05

Compound IX

(4- methoxycarbonyl phenyl)-2-chlorobenzoate

1HNMR (CDCl3, 400 MHz) δppm : 3.9 (s, 3H,-OCH3 from -COOCH3 group), 7.24 (d, J = 8.4 Hz, 2H, ArH from Methyl paraben moiety), 8.1 (d, J = 8.4 Hz, H, ArH from methyl paraben moiety), 7.2 - 8.2 (m, 4H, ArH from 2-Chloro benzoyl moiety). Molecular Formula

C15H11ClO4. White solid (1.297 gms, 68 %). Melting range 67 - 700C Anal. Calcd for C15H11ClO4 C 61.98,H 3.81, O 22.02 Cl 12.20 Found C 61.95, H 3.78, O 22.03

Chromatographic system

Column chromatography: For column chromatography 100 – 200 mesh Acme grade silica gel is used. The crude reaction mixture is concentrated under reduced pressure to yield crude mass which is preadsorbed on silica gel and purified by column chromatography with increase in concentration of Ethyl acetate in Petroleum ether. The fractions having similar 'rf' values were pooled together, concentrated and subjected for characterization using various spectroscopic techniques.

Thin layer chromatography TLC plates were prepared using silica gel G (ACME, BOMBAY). Pet. ether: EtOAc (85: 15) was used as the solvent system.

Radial chromatography The circular glass plates of thickness 1 mm, were prepared by using silica gel (PF254, E. MERCK, 50 g) in cold distilled water (105 ml). For elution, gradually increasing concentrations of EtOAc in pet ether were employed.

Biological Activity

Antibacterial activity using agar diffusion method 6 Concentration 100 µm

The synthesized molecules were screened for their antibacterial activity using agar diffusion method at 100 µm concentration against Gram positive (Staphylococcus aureus) and Gram negative (Escherichia coli) bacterial species qualitatively. The results of the antibacterial activities are summarized in Table 1.

Table 1: Antibacterial Activity Results

	Antibacterial Activity		
Compound	Against Gram - ve bacteria species	Against Gram +ve bacterial species (Staphylococcus	
No.	(Escherichia coli)	aureus)	
Methyl paraben	+	+	
Ethyl paraben	-	-	
Ampiciline	+	+	
	No. Methyl paraben Ethyl paraben	Against Gram - ve bacteria Compound species No. (Escherichia coli) Methyl paraben + Ethyl paraben -	

	(Std. Drug)		
4	I	-	-
4	II	+	-
5	III	-	-
6	IV	-	-
7	V	-	+
8	VI	-	
9	VII	-	-
10	VIII	-	+
11	IX	-	-

The above results shows that the base molecule Methyl paraben has anti-bacterial activity against both the bacterial culture. Its derivative *viz*. V and VIII were active against *Staphylococcus aureus* (Gram + ve bacteria) and II was active against *Escherichia coli* (Gram - ve bacteria) respectively. Thus, 3-methyl butyl and 4-chloro benzoyl derivatives were potential antibacterial candidates. In depth analysis of these compounds through structure activity relationship studies would provide further insight and can be an interesting topic of future studies.

The structural diversity and the pronounced biological activities encountered in the paraben derivatives suggests that this class of compounds is worthy for further studies that may lead to derivatives by using combinatorial chemistry approach is an alternative strategy to new therapeutic discovery. In other words the generation of diverse benzofuran derivatives develop new therapeutic molecules that might result in candidates having better activity.

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Output No.: May and (West), Dist. Thane - 401

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