

Journal of Applied and Natural Science 11(3): 645-649 (2019) ISSN : 0974-9411 (Print), 2231-5209 (Online) journals.ansfoundation.org

GC-MS analysis of yellow pigmented *Macrococcus* equipercicus isolated from alfalfa rhizosphere soil fields of Coimbatore

Z. Aiysha Thasneem							
Department of Microbiology, 641021, Tamil Nadu, INDIA	Karpagam	Academy	of	Higher	Education,	Coimbatore-	Article Info https://doi.org/10.31018/
K. Aravindh							jans.v11i3.2139
Department of Microbiology,	Karpagam	Academy	of	Higher	Education,	Coimbatore-	Received: July 2, 2019 Revised: August 16, 2019
641021, Tamil Nadu, INDIA							Accepted: August 26, 2019
M. Jeba Malar Fencia		A a a a a a a a a a a	- 4	Linhan		O simb stars	, 1000p104. , 14g401 20, 2010
Department of Microbiology, 641021, Tamil Nadu, INDIA	Karpagam	Academy	01	Higher	Education,	Compatore-	
C. Nitheesh Kumar							How to Cite
Department of Microbiology,	Karnagam		of	Higher	Education	Coimbatore	Thasneem, Z. A. <i>et al.</i>
641021, Tamil Nadu, INDIA	Kaipagam	Academy	01	riighei	Education,	Combatore	(2019). GC-MS analysis of
T. Pavithra							yellow pigmented Macro-
Department of Microbiology,	Karpagam	Academv	of	Hiaher	Education.	Coimbatore-	coccus equipercicus isolat-
641021, Tamil Nadu, INDIA					,		ed from alfalfa rhizosphere
K. Rajkumar							soil fields of Coimbatore. Journal of Applied and
Department of Microbiology,	Karpagam	Academy	of	Higher	Education,	Coimbatore-	Natural Science, 11(3):
641021, Tamil Nadu, INDIA							645- 649 https://
S. Surendran							doi.org/10.31018/
Department of Microbiology,	Karpagam	Academy	of	Higher	Education,	Coimbatore-	jans.v11i3.2139
641021, Tamil Nadu, INDIA							
M. Vidhya							
Department of Microbiology,	Karpagam	Academy	of	Higher	Education,	Coimbatore-	
641021, Tamil Nadu, INDIA							
R. Mahesh	atony South	Travanaa	r 0	Jindu C		raail 620002	
PG and Research Centre of B Kanyakumari District (Tamil Na		Tavanco	ne i		Jilege, Nage	10011-029002,	
S. Ramalakshmi*	add), maia						
Department of Microbiology,	Sri Moogar	nbigai Arts	: ar	nd Scier	nce College	for Women	
Palacode-Hosur Main Road, T							
(Tamilnadu), India	,			. ,		,	
*Corresponding author. E-mail	· arulaksh24	1@omail.co	m				
		i agginaniot	,,,,,				
Abstract	aaaaa imna	mont mior	flor	o which		uido obomicol	
The rhizosphere of plant poss compounds including second							
ment. The microbial flora of alf							
al activity and we found upto							
diversity, this yellow pigmente	ed colony w	as taken f	or f	further s	tudies. Thu	s. the culture	
was molecularly characterized	and identifi	ed for pote	nt k	oioactive	component	s responsible	
for antimicrobial activity. The s							
lites were produced and extra							
The antimicrobial study reveal							
Proteus sp with zone of inhibit							
tion of the isolate by 16S rRN							
cicus with 100 % similarity. If were identified and 13-docos							
found in ethyl acetate extract.							
<i>M.equipercicus</i> isolated from <i>N</i>							
presence of quercetin. Through							
compound producing bacteria							
Konworde: Alfalfa plant	and obros	atography		00 000	otromotry	Maaraaaaa	
Keywords: Alfalfa plant, equipercicus, Medicago sativa			-1118	iss spe	ctrometry,	wacrococcus	
					aaaa daar	reat avatar	n andlives in symbiotic

INTRODUCTION

sesses deep root system andlives in symbiotic association with microbes. The plant has high phy-

Alfalfa plant also known as Medicago sativa pos-

This work is licensed under Attribution-Non Commercial 4.0 International (CC BY-NC 4.0). © 2018: Author (s). Publishing rights @ ANSF.

to pharmacological importance due to the variety of compounds that has been isolated and identified to cure a variety of diseases (atherosclerosis, heart disease, stroke, cancer, diabetes) (Zhang *et al.*, 2006; Bora and Anupam, 2011; Krakowska *et al.*, 2017). They are also cultivated worldwide for high protein and fiber for cows. *Macrococcus* sp belongs to family Staphylococcaceae are gram positive cocci, non-motile and non-spore forming (Becker *et al.*, 2014).

Studies conducted in our lab showed that alfalfa plant rhizosphere soil contains Bacillus horikoshii (Nisha et al., 2019a) and Pantoeaagglomerans (Nisha et al., 2019b) with wide antibacterial activities. Macrococcusluteus and Neisseria sicca has been reported to be isolated from soil of Calotropisprocera and Catharanthusroseus. Their extracts possessed antibacterial activities against pathogens (Arora et al., 2013). Upon preliminary screening of microbes isolated from rhizosphere soil region of alfalfa plant, this isolate was selected for its amylase, cellulase, protease and phosphate solubilization activities. Therefore, the study was aimed to study the antibacterial activity, culture, extract and identify bioactive compounds of functionally diverse organism from rhizosphere soil region of Alfalfa plant (M. sativa) through GC-MS.

MATERIALS AND METHODS

Sample collection: The rhizosphere region soil samples of Alfalfa (*Medicago sativa*) plant were collected from June 2016 to March 2017 at Sulur, Coimbatore, Tamilnadu, India.

Isolation and antibacterial activity: Spread plate method was employed for isolation of microbes from the rhizosphere soil by taking one gram of the collected soil samplesfor serial dilution. About 0.1ml of serially diluted samples $(10^{-1} to 10^{-7})$ was spreaded onto sterile plates and incubated at 37°C for 24-72 hours. The isolated colonies were selected and stored in glycerol stocks until further required. The colonies were characterized by staining and biochemical standard methods. The selected isolates were studied for their antimicrobial activities against bacterial pathogens by well diffusion method (Nisha *et al.*, 2019a).

Production and extraction of the bioactive compounds: The bioactive compounds were extracted from the culture by large scale cultivation of microbes. After growth, the cells are separated by centrifugation and the supernatant is taken for for extraction process. Ethyl acetate was chosen as solvent for solvent to extract the compounds. After extraction, the solvent were evaporated to collect residues and stored and studied for GC-MS (Nisha *et al.*, 2019b).

GC-MS analysis: The microbial extract was further subjected to GC-MS analysis using Thermo MS DSQ II packed DB 35- MS Capillary standard non- polar column. Further the isolated compounds mass spectrums were interpretated by known compounds database NIST.

Molecular identification and phylogenetic analysis: The bacterial genomic DNA isolation were carried out using standard cold spring harbour lab protocol. From the isolated genomic DNA, rRNA genes (1.4 kb length gene)were amplified using the 8 F and 1541 R universal eubacterial primers (5'-AGAGTTTGATCCTGGCTCAG-3' and 5`-AAGGAGGTGATCCAGCCGCA -3` as forward and reverse primers). After amplification, the PCR products were sequenced by big dye terminator cycle sequencing kit (Applied BioSystems, USA) and resolved using Applied Biosystems model 3730XL automated DNA sequencing system (Xcelris Laboratories, India). The phylogeny analyses with multiple closely related sequences were done using MUSCLE 3.7 and PhyML 3.0 aLRT (Nisha et al., 2019a and b).

RESULTS AND DISCUSSION

Antibacterial activity: The plant usually releases a variety of carbon and other important nutrients necessary for microbial growth, which makes the mutualistic relationship between plants and microbes at rhizosphere interface (Bertin et al., 2003). The microbial extract of yellow pigmented colony studied for antibacterial activity against 7 clinical pathogens namely P.aeruginosa, Klebsiella sp, S. aureus, Proteus vulgaricus, S. pneumonia, E. coli, B. cereusrevealed that significant zone of inhibition was against against Streptococcus pneumonia and Proteus sp (18 and 20 mmrespectively) (Fig. 1). Similar to our study results, Janani et al., (2014) studied the pigmented marine bacteria Exiguobacterium sp. showing best antimicrobial and antioxidant activities isolated from different regions of India. The Exiguobacterium sp. showed activity against Shigella, Klebsiellasp and Staphylococcus aureus. Similarly studies by Nisha et al., (2019a, b) has isolated and reported potential isolates with wide antibacterial activity namely Bacillus horikoshii and P.agglomerans from alfalfa plant rhizosphere regions.

Molecular characterization of the isolates: The study results of BLAST showed100 % similarity with *Macrococcus equipercicus* (Fig. 2 and 3) and the gene sequences were submitted to the Gene bank (NCBI, USA) and Genebank ID accession number MK240540 received.

GC-MS Analysis: About twenty five compounds were identified from *Macrococcus equipercicus* extract by using GC-MS (Fig. 4). Table 1 and 2 reveal the compounds molecular formula, weight, structure, mass spectra and compound nature and its activities.

The highest intensity (29.58) with retention time of 36.20 showed 13-Docosenamide compound, has

Thasneem, Z. A. et al. / J. Appl. & Nat. Sci. 11(3): 645- 64) (2019)	
--	----------	--

S.N.	RT	wing GCMS analysis of con Name of the compounds	Molecular formula	Molecular weight	Peak area	Compound structure
1	5.57	Benzene 1,3,5-trimethyl	C ₉ H ₁₂	120	2.71	
2	8.16	Dodecane	$C_{12}H_{26}$	170	3.30	
3	9.18	Memantine	$C_{12}H_{21}N$	179	4.42	
4	10.40	4-Cyano-2H-1- benzothiopyran	$C_{10}H_7NS$	173	5.22	
5	11.29	Tetradecane	$C_{14}H_{30}$	198	2.97	R Anna Crista Management
6	13.25	2-tert-Butyl-4-isopropyl-5- methylphenol	$C_{14}H_{22}O$	206	2.36	
7	13.70	2-tert-Butyl-4-isopropyl-5- methylphenol	$C_{14}H_{22}O$	206	2.36	
8	15.49	Hexadecane	$C_{16}H_{34}$	226	2.23	Name (THE A DESCRIPTION OF A DESCRIPTION
9	15.72	5,8,11-Heptadecatriynoic acid,methyl ester	$C_{18}H_{24}O_2$	272	1.65	
10	19.14	1,4-dioxobicyclononane	$C_7 H_{10} N_2 O_2$	154	2.17	
11	19.59	1-Hexadecanol	$C_{16}H_{34}O$	242	2.05	Terrar Children all Terrar and an exact come
12	22.44	1,4-diaza-2,5-dioxo-3- isobutyl bicyclononane	$C_{11}H_{18}N_2O_2$	210	4.41	
13	23.56	Hexadecanoicacid,ethyl ester	$C_{18}H_{36}O_2$	284	4.47	
14	23.68	Hexadecanoic acid , ethyl ester	$C_{18}H_{36}O_2$	284	4.47	
15	26.07	2-Acetyl-3-ethyl-7- methoxyindole	$C_{22}H_{22}N_2O_3$	362	0.95	
16	27.35	Octadecanoicacid,ethyl ester	$C_{20}H_{40}O_2$	312	3.15	fama calendar and a
17	27.47	Octadecanoic acid ,ethyl ester	$C_{20}H_{40}O_2$	312	3.15	news coloring way of the Color was
18	29.57	Ergotaman-3,6,18-trione	$C_{33}H_{35}N_5O_5$	581	2.83	
19	30.12	Androst-4-en-3-one,17- methoxy,3-methoxime	$C_{21}H_{33}NO_2$	331	3.90	
20	31.76	Lucenin2	$C_{27}H_{30}O_{16}$	610	0.63	
21	32.45	3,17-Dioxo-11-a- hydroxyandrostane-1,4-	$C_{19}H_{24}O_3$	300	1.52	
22	33.92	diene 3,17-Dioxo-11-a- hydroxyandrostane-1,4-	$C_{19}H_{24}O_3$	300	1.52	
23	36.20	diene 13-Docosenamide	$C_{22}H_{43}NO$	337	29.58	
24	37.34	Tetracosa-2,6,14,18,22- pentaene-10,11- diol,2,6,10,15,19,23-	$C_{30}H_{52}O_2$	444	1.35	
25	37.72	hexamethyl QUERCETIN7,3,4- TRIMETHOXY	$C_{18}H_{16}O_7$	344	0.73	

 Table 1. Showing GCMS analysis of compounds obtained from Macrococcus equipercicus extract.

S.N.	RT	Name of the compound	Compound nature	Activity
1	5.57	Benzene 1,3,5-trimethyl	Aromatic hydrocarbon	precursor to styrene
2	8.16	Dodecane	Alkane hydrocarbon	Solvent
3	9.18	Memantine	Amantidine	Treat Alzimer's disease
4	10.40	4-Cyano-2H— benzothiopyran	Aromatic compound	Used in drugs
5	11.29	Tetradecane	Alkane hydrocarbon	Petroleum spirit
6	13.25	2-tert-Butyl-4-isopropyl-5 -metylphenol	Lipophilic organic compound	Food additive
7	13.70	2-tert-Butyl-4-isopropyl-5 -metylphenol	Lipophilic organic compound	Food additive
8	15.49	Hexadecane	Alkane hydrocarbon	Fuel mixture
9	15.72	5,8,11,Heptadecatriynoic acid methyl ester	Acid compounds	Explosive properties
10	19.14	1,4-dioxobicycloninane	Organic compound	Dehydrohalogenation
11	19.59	1-Hexadecanol	Fatty alcohol	Opacifier
12	22.44	1,4-diaza-2,5-dioxo-3- isobutyl bicyclononane	Not reported	Not reported
13	23.56	Hexadecanoic acid ethyl ester	Saturated fatty acid	antimicrobial, antioxidant, antifungal, 5Alpha reductase inhibitor and hypo- cholesterolemic
14	23.68	Hexadecanoic acid ethyl ester	Saturated fatty acid	antimicrobial, antioxidant, antifungal, 5Alpha reductase inhibitor and hypo- cholesterolemic
15	26.07	2-Acetyl-3-ethyl-7- methyoxyindole	Not reported	Transform Harman alkaloids
16	27.35	Octadecanoic acid ethyl ester	Saturated fatty acid	Confers solubility in organic solvent
17	27.47	Octadecanoic acid ethyl ester	Saturated fatty acid	Confers solubility in organic solvent
18	29.57	Ergotaman-3,6,8-trione	Alkaloid	Inhibits vesicular glutamate transporter activity in cow cerebral synaptic vesi- cles
19	30.12	Androst-4-eb-3-one,17- methoxy,3-methoxime	Aromatic compound	Aromatizing enzyme complex of human placenta
20	31.76	Lucenin2	Glycosyl compound	Not reported
21	32.45	3,17-Dioxo-11-a- hydroxydrstane-1,4- diene	Not reported	Not reported
22	33.92	3,17-Dioxo-11-a- hydroxydrstane-1,4- diene	Not reported	Not reported
23	36.20	13-Docosenamide	Amide of docosenoic acid	Reduces mobility and slightly lessened awareness in cerebrospinal fluid of rat and humam
24	37.34	Tetracosa-2,6,14,18,22- pentaene-10,11- diol,2,6,10,15,19,23- hexamethyl	Not reported	Not reported
25	37.72	QUERCETIN 7,3,4- TRIMETHOXY	Flavanoid	Antioxidant, anthelmintic, antimicrobial, antileishmanial, antiplasmodial

Table 2. Activity of compounds identified in Macrococcus equipered	rcicus extract.
--	-----------------

molecular formula is $C_{22}H_{43}NO$ and molecular weight of 337. The compound is amide of docosenoic acid, has been reported for *Ludwigia perennis* antimicrobial activity (Sharmila *et al.*, 2017). The retention time of 4-Cyano-2Hbenzothiopyran from the microbial extract present at 10.40 and its peak area is 5.22, has molecular formula as $C_{10}H_7NS$ and molecular weight is 173 and it is aromatic compound which has the activity in drug usage. The retention time of Androst-4-en-3-one,17-methoxy,3-methoxime is 30.12 and has peak area is 3.90, has molecular formula is $C_{21}H_{33}NO_2$ and molecular weight of 331 and it is an aromatic compound and it has an aromatizing enzyme complex of human placenta. Quercetin 7,3',4'-trimethyl ether is a trimethoxyflavone, derivative of quercetin. The compound

Thasneem, Z. A. et al. / J. Appl. & Nat. Sci. 11(3): 645- 649 (2019)



Fig. 1. Showing antibacterial activity of yellow pigmented colony against Streptococcus pneumonia and Proteus sp.

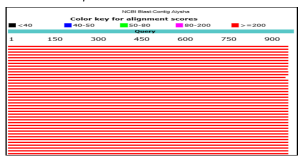
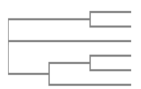


Fig. 2. Showing multiple alignment scores of Macrococcus equipercicus.



NR_036847.1_Macrococcus_brunensis Macrococcus_equipercicus KP345966.1_Macrococcus_sp. DQ279392.1_Micrococcus_sp. AJ576067.1_Macrococcus_equipercicus MG649989.1_Macrococcus_equipercicus

Fig. 3. Phylogenetic tree of Macrococcusequipercicus based on the 16S rRNA gene sequencing.

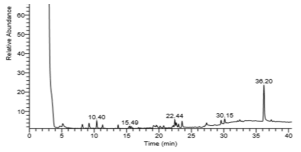


Fig. 4. GCMS spectrum analysis of Macrococcus equipercicus extract.

Quercetin 7,3,4-trimethoxy, a flavonoid has been reported for its activities such as antioxidant, anthelmintic, antimicrobial, antileishmanial, antiplasmodial by Kalpana Devi *et al.* (2016).

Conclusion

In present study, the soil isolate from the rhizosphere region of Alfalfa plant of Coimbatore was molecular characterized and identified with 100 % similarity as *Macrococcus equipercicus*. Highest activity was recorded against two pathogens *Streptococcus pneumonia and Proteus sp* which can be due to presence of 13-docosenamide at retention time of 36.2minutes. *In vitro* and *In vivo* biological studies are further necessary to find new drugs against cancer. Through this study we were able to identify potent antimicrobial compounds such as memantine, quercetin and various esters from medicinally important *Medicago sativa*. Thus, the study provides insight into microflora and its bioactive compounds harbouring alfalfa rhizosphere soil region.

REFERENCES

- Arora, S., Nandi, D., Prasad, N., Rawat, S. and Pandeya, A, (2013). Isolation and characterisation of antibiotic producing microbes present in rhizospheric soil. *International Journal of Scientific & Engineering Research*, 4(9):1157-1166.
- Becker, K., Heilmann, C., Peters, G. (2014). Coagulase-negative Staphylococci. *Clinical Microbiology Revie, ws* 27(4): 870 –926.
- Bertin, C., Yang, X., and Weston, L. A. (2003). The role of root exudates and allelochemicals in the rhizosphere. *Plant and Soil*, 256(1): 67-83.
- Bora, K. S., Anupam Sharma (2011). Phytochemical and pharmacological potential of *Medicago sativa*: A review. *Pharma Biol.*,49(2): 211-220.
- Janani, B., Kiruthika, P., Angayarkanni, J. (2014). Isolation Of Pigmented Marine Bacteria *Exiguobacterium* Sp. From Peninsular Region Of India And A Study On Biological Activity Of Purified Pigment. *International Journal of Scientific & Technology Research*, 3(3): 375-384
- Kalpana Devi, R., Subramani, V., Annamalai, P., Nakulan, Vr., Narayanaperumal, J. (2016). Phytochemical analysis, in vitro antioxidant potential and gas chromatography-mass spectrometry studies of *Dicranopterislinearis. Asian J. Pharm. Clin. Res.*, 9 (2): 220-225.DOI: http://dx.doi.org/10.22159/ ajpcr.2016.v9s2.13636
- Krakowska, A., Ska, K.R., Walczak, J., Kowalkowski, T., Buszewsk, B. (2017). Comparison of Various Extraction Techniques of *Medicago sativa*: Yield, Antioxidant Activity, and Content of Phytochemical Constituents. *J AOAC International*, 100(6): 1681-93.doi: 10.5740/jaoacint.17-0234.
- Nisha, M. Nair, Kanthasamy, R., Mahesh, R., IruthayaKalaiSelvam, S. and Ramalakshmi, S. (2019a). Identification of Antibacterial Compound from *Bacillus horikoshii*, isolated from rhizosphere region of Alfalfa plant. *Journal of Applied Sciences*, 19(2): 140-147.DOI: 10.3923/jas.2019.140.147
- Nisha, M. Nair, Kanthasamy, R., Mahesh, R., IruthayaKalaiSelvam, S. and Ramalakshmi, S. (2019b). Production and Characterization of Antimicrobials from isolate *PantoeaAgglomerans* of Alfalfa Plant Rhizosphere Soil. *Journal of Applied and Natural Science*, 11(2): 267-272.Doi: https:// doi.org/10.31018/jans.v11i2.2031
- 10.Sharmila, M., Rajeswari, M., Jayashree, I. (2017). GC-MS Analysis of Bioactive Compounds in the Whole Plant of Ethanolic Extract of *Ludwigiaperennis* L. Int J Pharm Sci Rev Res 46(1): 124-128.
- 11.Zhang, L., Zhang, D., Feng, K. (2006). Inhibition of refined components of *Medicago sativa* polysaccharides to the activities of reverse transcriptase of HIV and protease of HIV. *Zhongguo Shipin Xuebao*, 6:5-62