Evaluation of antibacterial spectrum and phytochemical analysis of *Laurus nobilis* Leaves extracts

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Abstract

Laurus nobilis is used as medicine and food additives in Saudi Arabia, and also, it could be used to study as antibacterial activity and assessment through agar well diffusion and most susceptible bacteria used was *Pseudomonas aeruginosa*, flowed by *Staphylococcus aureus*. Resistant bacteria mostly were *Bacillus subtilis* followed by MRSA while there was no effect on *Micrococcus luteus*. HPLC analysis found the presence of 11 phenolic compounds isolated from *Laurus nobilis*including Gallic Acid, Catechin, Coffeic Acid, Syringic Acid, Rutin, Coumaric Acid, Naringenin, Quercetin, Ferulic Acid, Vanillin, and Cinnamic Acid. The result showed that obtained found variations from the concentration ratio of ethanolic *Laurus nobilis* leaves extract. HPLC illustrated that Rutin, Gallaic acid, and syringic acid in high concentrations were 418.37, 81.66, and 34.43 $\mu g/g$, respectively and phenolic compounds ranged from 0.21 to 21.42 $\mu g/g$ and the lowest Cinnamic concentration was 0.21 $\mu g/g$.

Keywords: Laurus nobilis, Antibacterial, phytochemical constituents

INTRODUCTION

Resistance microorganism causes diseases that have posed a hazard in human and threatens public health. Many studies started in search of new alternatives to prevent disease ^[1, 2]. Plants consume a rich source of antimicrobial, as also have natural protection products against microbial occurrence [3-^{5]}.Laurel (L.) nobilis is an aromatic and commonly used culinary spice in Western and Asian countries such as Saudi Arabia. Cultivation of such a plant was in the Mediterranean and many warm countries. The aromatic tree is 2 m to 10 m high ^[6]. Leaves and berries are mainly used as sauce and spice aroma ^[7]. Laurus nobilisleaves are used to treat diseases. Essential oils and organic acids in this plant have shown strong antibacterial action against pathogenic strains [7-10]. The leaves have been examined ingredients. The Laurus nobili scomposition and yield are influenced by different factors, including extraction method, plant parts, yield season, growth environment, etc. In the present study, the ethanolic extract of Laurus nobilis leaves was examined using HPLC. This work will help to identify the phenolic compounds, which can be used in beneficial values. This study aimed at evaluating the chemical composition and antibacterial spectrum of Laurus nobilisleaves extract in Saudi Arabia.

MATERIALS AND METHODS

Collection and preparation of samples

Samples of *Laurus nobilis* leaves were collected during May 2019 from Albahahrigon (19°98' 28°N, 41°52' 50°E) southwest Saudi Arabia from cool slopes at 2050 m.a.s.l)

Saudi Arabia, the species status of this plant was fervid at the faculty of Sciences Herbarium, King Abdulaziz University, Jeddah. The plant leaves were transported to the laboratory, washed in running tap water to eliminate dust particles and debris and then washed in distilled water for five minutes.

Preparation of Laurus nobilis leaves extracts

Ten grams of dried *Laurus nobilis*leaves washed to cutting to small pieces (1-2 mm) and adding 100 ml of distilled water and ethanol extract (1:10W/V)/48h and filtration later. Solutions evaporated with pressure (40°C) until dryness and diluted by dimethyl sulfoxide (DMSO) and stored at 20°C ^[11].

Phytochemical screening Phenolic extraction

Phenolic extraction was conducted according to Mattila *et al.* ^[12]. 15 ml of4NNaOHwas added to 0.2L of water extract in a

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How to cite this article: M. Gum Gumjee, N. Evaluation of antibacterial spectrum and phytochemical analysis of *Laurus nobilis* Leaves extracts. Arch PharmaPract. 2020;11(2):145-8.

50-ml Pyrex centrifuge tube purged with nitrogen and shaken in the dark for 2h with a wrist-action shaker. After phenolic acids liberated by alkaline hydrolysis, the samples were acidified with ice-cold HC1 (6N) to decrease pH to 1-2. Then the solution was centrifuged at 3000g and the supernatant was decanted into a 250-ml separator/funnel. The supernatant was extracted with ethyl acetate (3×50 ml) by shaking for 10s and the mixture was allowed to settle for 5min between extractions. Ethyl acetate fractions were collected and pooled. The second supernatant was re-extracted with ethyl acetate (3×50 ml) as before and all ethyl acetate fractions were pooled.

HPLC Analysis

Phenolic was separated according to Shimaduz (Kyoto, Japan) HPLC apparatus (model, LC-4A) equipped with visible/UV detector (model, SPD-2AS) at 280 nm and stainless steel column (25.0cm \times 4.6mm i.d.).

Antibacterial Assay

Three Gram-negativebacteria: *Escherichia coli* (ATCC8739, *Klebsiella pneumonia* (ATCC700603), and *Pseudomonas aeruginosa* (ATCC27853) and four Gram-positive bacteria: *Bacillus subtilis* (ATCC11774); Methicillin-resistant *Staphylococcus aureus* (MRSA) (ATCC977), *S.aureus* (ATCC29213), and *Micrococcus luteus* (ATCC4698) are used to evaluate the efficacy of the extract. The tested strains were subcultured in nutrient agar medium slopes. The stock cultures were stored at 4°C.

Antibacterial Activity

The activity of bacterial strains was determined using the agar well diffusion assay methods according to Holder and Boyce ^[13]. DMSO was used as the negative control and streptomycin and ciprofloxacin (10 mg/disc) were used as the positive controls. Bacterial cultures were incubated at 37°C for 24h. Antimicrobial was determined by measuring the zone of inhibition ^[14].

Statistical Analysis

Means of variable and standard error were accepted using SPSS to detect any significant differences between pathogenic microorganisms and extract type.

RESULTS AND DISCUSSION

Antibacterial influence of the ethanolic extracts of *Laurus* nobilisleaves using disc diffusion against 4 Gram-positive bacteria andthree Gram-negative bacteria at 200 mg/ml, *Laurus nobilis*leaves extract given high activities against the tested organisms. The most susceptible strain was *P. aeruginosa* followed by *S.aureus*, while the most resistant strain was *Bacillus subtilis* followed by MRSA and the diameter of inhibition zones against bacteria was 29.00mm, 28.00mm, 17.00mm, and 18.00mm, respectively (Table 1). This study was consistent with the study conducted by Ramos *et al.* ^[15] in which the antibacterial effect of different solvent extracts and separated ingredients of *Laurus nobilis*were evaluated and showed strong antibacterial action with the ethanol extract against pathogenic bacteria

and foodborne spoilage .Noriakietal^[16] found antimicrobial activity against many pathogens including opportunistic Gram-positive bacteria. In addition, the present results are in line with Al-Hussaini andAdel. [17] who showed that the leave extract of L. nobilis was additionally active against B.subtilis and P. aeruginosa in contract to Gram-negative bacteria. Also, Erturk^[18] described eleven ethanolic extracts of spices for in vitro antibacterial effect against Pseudomonas aeruginosa, Escherichia coli, Staphylococcus aureus, and Bacillus subtilis, which was in line with the current results. Antimicrobial activities of standard antibiotics (Streptomycin (10 µg/disc) and Ciprofloxacin (10µg/disc)) had an inhibitory effect against all the tested bacteria. Ciprofloxacin is more effective than streptomycin (Table 1). Antimicrobial activity of Laurus nobilisleaves in terms of inhibition zone (mm) of the ethanolic extracts were tested at the concentration of 0.5 mg/ml against microorganisms. Laurus nobilis extract leaves has antimicrobial action against P. aeruginosa, S.aureus, Klebsiella pneumonia, and E. coli with inhibition zone of 29, 28, 26, and 24 mm, respectively, which was considerably higher than that of the control (Streptomycin) with the inhibition zones of 22, 19, 25, and 23 mm, respectively and no inhibition zone against Micrococcus strain that was parallel with Shital^[19]. Inhibition diameters against bacteria are shown in Table (1). EL Malti and Amarouch^[20] reported the average inhibition zone of L. nobilisleaves extract ranged from 7mm (Pseudomonas aeruginosa ATCC 27853) to 20mm (L. monocytogenes). The results presented antibacterial use of the bay extract for the treatment of S. aureus infection [21] and on human pathogenic bacteria by disc diffusion method via the average inhibition zone against 9 bacteria strains. L. nobilis effects on bacteria more than that of tetracycline antibiotic. Extract of L. nobilis presented strong anti-bacterial activity similar to the finding of Moghtader and Farahmand^[22].

Table 1. The antibacterial activity of ethanolicextracts of Laurus nobilisextracts against different pathogenic bacteria.

Diameter of the inhibition zone (mm) Mean ± SD

Bacterial strains	Laurus nobilis leaves extracts	Streptomycin (10 µg/disc)	Ciprofloxacin (10µg/disc)
Bacillus subtilis	17.67±0.58	25.00±00.00	34.00±00.00
MRSA	19.00±1.00	20.00±00.00	36.00±00.00
Micrococcus luteus	00.00±00.00	27.00±00.00	46.00±00.00
Staphylococcus aureus	28.00±00.00	19.00±00.00	38.00±00.00
Klebsiella pneumonia	26.00±00.00	25.00±00.00	42.00±00.00

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E. coli	24.67±0.58	23.00±00.00	44.00±00.00
Pseudomonas aeruginosa	29.00±00.00	22.00±00.00	42.00±00.00

Phytochemical Screening of. Laurus nobilis

HPLC analysis found the presence of 11 phenolic compounds isolated from *Laurus nobilis* including Quercetin, Rutin,Syringic Acid, Coffeic Acid, Catechin, Gallic Acid, Coumaric Acid, Vanillin, Ferulic Acid, Naringenin, and Cinnamic Acid). HPLC showed Rutin, Gallaic acid, and syringic acid at high concentrations of 418.37, 81.66, and $34.43\mu g/g$, respectively. Phenolic compounds' concentrations ranged from 0.21 to $21.42 \ \mu g/g$ and Cinnamic Acid had the minimum concentration (0.21 $\mu g/g$) (Table 2).

These results are consistent with the results of Stefano et al. ^[23] who showed that phytochemicals had many flavonoid derivatives. Semipreparative HPLC of laurel leaves found 10 flavonoid O-glycosides, under the best extraction by Muñiz-Márquez et al. ^[24]. Active principles identified showed antimicrobial activity against test organisms parallel to the observations of Gumgumje et al. ^[25], Gumgumjee& Hajar^[26], and Hajar & Gumgumjee^[27]. The extracts from various parts of L. nobilisconsisted of 1, 8 Cineole, sabinene, α-pinene, and p-Cymene and many biological and pharmacological characterization recorded the antibacterial effect of L. nobilis leave extract [28]. These results are consistent with the results by Muñiz-Márqueza et al. [29] in which the phytochemical screening by HPLC analysis determined 4 phenolic compounds in the extract: coumaric, gallic, pyrogallol and resorcinol. Further studies involved in each compound are required. Previous studies established the presence of other phenolic compounds in L. nobilis. Muñiz-Márqueza et al. [30] documented the presence of caffeic, vanillic, and ferulic acids. Lu et al. [31] found rutinin and unknown phenolic acids. Environmental conditions had effects on phenolic compounds in plants ^[32, 33]. Chemical analysis of L. nobilis extract showed the presence of alkaloids, flavonoids, tannins, and essential oil [34]. In addition, a study by Ramling et al. [35], showed that Laurus nobilisleaves produced four nonpolar flavonoids kaempferol-3-O-α-L-(3",4"-di-E-p-coumaroyl)- rhamnoside, kaempferol-3-O-α-L-(2",4"-di-E-pcoumaroyl)-rhamnoside, kaempferol-3-Oα-L-(2"-4"-pcoumaroyl)- rhamnoside, and a new product kaempferol-3-O-α-L-(2",4"-diZ-p-coumaroyl)rhamnoside^[36]. 5 discovered megastigmane glucosides called laurosides A-E were isolated as novel phenolic glucoside from methanolic extract of L. nobilisL. leaves. Kaempferol- 3-rhamnopyranoside, and kaempferol-3, 7- dirhamnopyranoside isolated from Laurus nobilisaqueous and ethanolic extracts ^[37] (Table 2).

Table 2: Chemical composition analysis of Laurusnobilis leave extracts.				
Phenolic compounds	Area	Conc. (µg/ml)		
Gallic Acid	1502.94	81.66		

Catechin93.9021.42Coffeic Acid52.511.33Syringic Acid1308.5934.43Rutin4245.16418.37Coumaric Acid0.000.00Vanillin464.168.87Ferulic Acid178.522.85Naringenin324.7012.14Querectin41.311.28Cinnamic Acid34.380.21				
Coffeic Acid52.511.33Syringic Acid1308.5934.43Rutin4245.16418.37Coumaric Acid0.000.00Vanillin464.168.87Ferulic Acid178.522.85Naringenin324.7012.14Querectin41.311.28Cinnamic Acid34.380.21	Catechin	93.90	21.42	
Syringic Acid 1308.59 34.43 Rutin 4245.16 418.37 Coumaric Acid 0.00 0.00 Vanillin 464.16 8.87 Ferulic Acid 178.52 2.85 Naringenin 324.70 12.14 Querectin 41.31 1.28 Cinnamic Acid 34.38 0.21	Coffeic Acid	52.51	1.33	
Rutin4245.16418.37Coumaric Acid0.000.00Vanillin464.168.87Ferulic Acid178.522.85Naringenin324.7012.14Querectin41.311.28Cinnamic Acid34.380.21	Syringic Acid	1308.59	34.43	
Coumaric Acid 0.00 0.00 Vanillin 464.16 8.87 Ferulic Acid 178.52 2.85 Naringenin 324.70 12.14 Querectin 41.31 1.28 Cinnamic Acid 34.38 0.21	Rutin	4245.16	418.37	
Vanillin 464.16 8.87 Ferulic Acid 178.52 2.85 Naringenin 324.70 12.14 Querectin 41.31 1.28 Cinnamic Acid 34.38 0.21	Coumaric Acid	0.00	0.00	
Ferulic Acid 178.52 2.85 Naringenin 324.70 12.14 Querectin 41.31 1.28 Cinnamic Acid 34.38 0.21	Vanillin	464.16	8.87	
Naringenin 324.70 12.14 Querectin 41.31 1.28 Cinnamic Acid 34.38 0.21	Ferulic Acid	178.52	2.85	
Querectin 41.31 1.28 Cinnamic Acid 34.38 0.21	Naringenin	324.70	12.14	
Cinnamic Acid 34.38 0.21	Querectin	41.31	1.28	
	Cinnamic Acid	34.38	0.21	



CONCLUSION

High differences in volatile components of the ethanolic extract of *Laurus nobilis* leaves are related to diverse geographic origins, growing conditions, periodic conditions, and procedures. Major compounds in *Laurus nobilis* leaves were Gallic acid, catechin, coffeic acid, syringic acid, Rutin, coumaric acid, vannillin, ferulic acid, naringenin, Quercetin, and cinnamic acid. The leaf of *Laurus nobilis*had high pharmacological actions such as antimicrobial and a variety of constituents in that could be responsible for an extensive range of biological activities of the plant.

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