The Importance of using Arbutus Pavarii Plant as a Medicinal Herbal Plant and its Benefits in our Daily Life

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Abstract: Arbutus pavarii Pamp. (Ericaceae) is one of the endemic species in Libya. Samples of the non-ripened and ripened fruits of Arbutus pavarii Pamp. Were collected from its natural habitat (El-Gabel El-Akhdar, Libya). Fruit characters as fruit size, weight, fruit mass and seeds per fruit were estimated. The vitamin content of A, C and E was estimated using different methods. The total phenolic and flavonoids content of the fruits was determined spectrophotometrically. The antioxidant and antiviral activity of the fruits was determined using different methods. The results were discussed according to the nutritional value of the fruits. The human impacts and their effects on plant vegetation and biodiversity became a field of major interest in the last few years. many of human activities occurred in AL-fjabal AL-akhdar area as result of increase of development activities and growth of population. The present investigation was carried out to study the Important of Arbutus pavarii Pamp. (Ericaceae) As A herbaceous plant useful in medicinal uses, as well as many benefits in use and national economic wealth. The paper presents the results of phytochemical screening, antioxidant, antimicrobial and anti-proliferative activity studies of Arbutus pavarii (A. pavarii) plant. The whole aerial part of the plant was collected during the spring season (2020), identified and extracted with methanol, chloroform and n-Hexane. Phytochemical screening was carried out using standard procedures. Antioxidant activity was done using 2, 2-Diphenyl-1-picrylhydrazyl (DPPH) assay. Antimicrobial evaluation was performed using agar well diffusion method. The anti-proliferative activity was evaluated on two breast adenocarcinoma and lung cancer cells. Phytochemical screening showed the presence of flavonoids, tannins, glycosides, simple phenolic, free reducing sugars, terpenes and sterols in the plant extract.

Keywords: Arbutus Pavarii, Antioxidant Activity, Antimicrobial Activity, Anti-Proliferative Activity.

1. INTRODUCTION

Arbutus pavarii Pamp. (Ericaceae) is one of the endemic species in Libya and it distributes naturally as wild plant in El-Gabel El-Akhdar area, which characterized with Mediterranean climatic conditions (Elshatshat 2009; Elshatshat et al 2009; and Elabidi and Elshatshat 2017). It is evergreen shrub or small tree, 1.5 to 3 m tall with reddish brown peeling bark. The flowering season appear from late October to February and the flowers are a good source of nectar for bees. Because of its nutritional and medicinal value, A. pavarii Pamp. honey is widely used for folk medicinal purposes (El abidi and Elshatshat 2017), in addition, other honey types which collected from other plant species (Elshatshat and Elsilini 2006). In addition, the antioxidant properties and activities (Isbilir, et al. 2012; Mendes, et al. 2011). Because of its endemism and lack of information about the fruits of Arbutus pavarii Pamp., this work was conducted to shed some light on the fruit characters and the characterization
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of the antioxidant composition at different fruit ripening stages. Plant biodiversity is reported that one of important genetic resources of many species growing spontaneously around the Mediterranean basin (Louhaichi et al, 2011). Libya is one of Mediterranean basin country and a native of many plant species that model of biodiversity (El-Darier and El-Mogaspi, 2009). Plants that have economic importance such as Haplophyllum tuberculatum (Forsk) and Strawberry trees (Arbutus pavarrii) which are located in different environments (coastal, mountainous and desert) in Libya (El-Darier and El-Mogaspi, 2009). There are a number of factors for the difficulty of germination and growth of these plants naturally in the wild land. Drop in rate of rainfall annually it is the most important environmental factor which has made the wild lands drier and decreased significantly seeds germination (Elmaghrabi at al. 2017). In addition, overgrazing and the use of lumber as firewood and also expansion of new farms, which contributed of deterioration sharply of edible and medical wild plant resources which led these species to endanger (Elmaghrabi et al 2017). The genus Arbutus belongs to the Vaccinioideae subfamily which includes evergreen shrub-like woody taxa with laurel-like and sclerophyllous leaves of the Ericaceae family (Torres et al., 2002). There were about six species of Arbutus grows spontaneously around the Mediterranean basin. The species is drought tolerant and able to regenerate following forestry fires making it quite interesting for forestation programs in Mediterranean regions. Fruits are used to make jellies and a spirit which represents the main income for owners (Torres et al., 2002). Arbutus pavarrii species it is only located in the coast of Libya especially at El-Jabal El-Akhdar Region in the Mediterranean regions (El-Darier and El-Mogaspi, 2009). Haplophyllum tuberculatum (Forsk) juss that belongs to the Rutaceae family (El-Naggar et al., 2014). Haplophyllum tuberculatum is a herbal plant is, simple leaves, reciprocal, heterogeneous white, small yellow flowers (Puricelli et al., 2002). The whole plant is being used in pharmaceutical product with the exception of the roots. The essential oil of Haplophyllum tuberculatum was reported by (Al-Rehaily et al, 2014) which prepared by hydrodistillation of the fresh flowering aerial parts of the plant collected from wild types. The oil was subsequently analyzed by GC and GC-MS. Thirty-seven compounds, accounting for 96.4 % of the oil composition were identified in this study (Al-Rehaily et al, 2014). The antimicrobial and activity of the essential oil was also evaluated against various human pathogens, where a relatively low inhibitory range was observed. Because these species (Haplophyllum tuberculatum and Arbutus pavarrii) which have good economic value and, the scarcity of plant biomass available in the natural habitat, we set up non-conventional methods for plant propagation from nodal stem segments and, at the same time, we established cell cultures of the plant (Elmaghrabi et al, 2017).

2. MATERIALS AND METHODS

2.1. Materials

2.1.1. Plant Material

The samples of Arbutus Pavarii. Collected from the EL-Gabal EL-Akhdar area (Asulntah area) and then choose three random plants were next to some includes (Sample1= plant intact naturally grow, sample2= plant grow almost normal and sample 3= No plant grows naturally). All samples were collected from the same location.

2.1.2. Chemicals

1. 1-Diphenylpicrylhydrazyl (DPPH’), methanol, Ethanol and acetone were supplied by Sigma and Merck company. Ascorbic acid, Folin-Ciocalteu reagent, ferric chloride, potassium ferricyanide, monobasic dihydrogen phosphate, dibasic monohydrogen phosphate, trichloro acetic acid, sodium carbonate, anhydrous sodium sulfate and pyrogallol were obtained from the biochemistry laboratory of Chemistry department-Benghazi University.

3. METHODS

3.1. Extraction of Essential Oil from Leaves of Arbutus Pavarii. (Sample 1, Sample 2 and Sample 3)

The dry powdered leaves of Juniperus phoenicea (500g) were subjected to hydro distillation using Clevenger apparatus. The isolation of volatile oils was completed within 6 hours (Clevenger, 1928).

3.1.1. Store Essential Oils

The oil samples were stored at 7°C in dark air-tight containers after drying over anhydrous sodium sulfate and filtered before injecting to GC-MS analysis.
3.2. Oil Analysis

The oil samples extracted from leaves of *Arbutus Pavarii*. Were subjected to the following tests:

3.2.1. Gas Chromatography/Mass Spectra

Thermo Scientific, Trace GC Ultra & ISQ Single Quadruple MS, DB-5 bonded-phase fused-silica capillary column was used in for GC/MS analysis of essential oils. This experiment has been conducted in the central laboratory at Cairo University-Egypt.

3.2.2. Antioxidant Activities Assays and Quantitative Analysis

All these experimental assays have been conducted in biochemistry laboratory at Benghazi University.

**Total Phenolic Content (TPC):**

Total concentration of phenolic compound in all oil extracts obtained from leaves of *Arbutus Pavarii*. Was estimated using the colorimetric method based on Folin-Ciocalteu reagent (Sawsan et al. 2010). 0.05 ml of the oils at different concentrations "100,200,300,400,500 µg/ml" were mixed separately with 0.05 ml of Folin-Ciocalteu reagent. Then 0.5 ml of 15% sodium carbonate solution was added to the mixture and then the adjusted to 1 ml with 0.4 ml of distilled water. The reaction was allowed to stand for 10 min, after which the absorbance was recorded at 725 nm by UV-visible spectrophotometer. Quantification was done with respect to standard calibration curve of Pyrogallol the results were expressed as pyrogallol "µg/ml" Fig (6). Estimation of the phenolic compounds was carried out in triplicates. The results were mean values (standard deviations).

**Table1. Total phenolic content (TPC) of Pyrogallol**

| Concentration of Pyrogallol (µg/ml) | Mean (Standard Deviation) |
|-----------------------------------|---------------------------|
| 100                               | 0.410 (  0.032)           |
| 200                               | 0.799 (0.0220)            |
| 300                               | 1.333 ( 0.0045)           |
| 400                               | 1.828 ( 0.0117)           |
| 500                               | 2.105 ( 0.0225)           |

**Total Flavonoids Content (TFC)**

Aluminum chloride colorimetric method was used for determination of total flavonoids (Chang et al., 2002). 2 ml of different concentration "100, 200, 300, 400, 500 µg/ml " of oil extracts mixed with 0.1ml of 10% aluminum chloride, 0.1 ml of 1 M potassium acetate and 2.8 ml of distilled water. It kept at room temperature for 30 min; the absorbance of the reaction mixture was measured at 415 nm with a UV-visible spectrophotometer. The calibration curve was obtained by preparing different quercetin solutions in methanol at concentrations "100 to 500 µg/ml" Fig (7).

**Table2. Total flavonoids content of quercetin**

| Concentration of quercetin (µg/ml) | Mean (Standard Deviation) |
|------------------------------------|---------------------------|
| 100                                | 0.279 ( 0.092)            |
| 200                                | 0.560 (0.035)             |
| 300                                | 0.834 ( 0.003)            |
| 400                                | 1.066 (0.009)             |
| 500                                | 1.300 ( 0.006)            |
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Reducing Power Assay (RPA)

The reducing power was determined according to (Naznin and Hasan, 2009). 2ml of the all oil extracts with different concentrations "100,200,300,400,500µg/ml" were mixed with 2.5 ml phosphate buffer (0.2 M, pH 6.6) and 2.5 ml potassium ferricyanide then mixture was incubated in water bath at 50 C° for 20 minutes and 2.5 ml of trichloroacetic acid was added to the mixture which was then centrifuged at 3000 rpm for 10 minutes. Finally, 2.5 ml of the supernatant was mixed with 2.5 ml of distilled water and 1 ml FeCl₃ substances, which have reduction potential react with potassium ferricyanide (Fe³⁺) to form potassium ferricyanide (Fe²⁺), which then reacts with ferric chloride to form ferric ferrous complex that has an absorption maximum at 700nm by UV-Visible spectrophotometer. Quantification was done with respect to standard calibration curve of ascorbic acid the results were expressed as ascorbic acid "µg/ml" Fig (8).

Potassium ferricyanide + ferric chloride → antioxidant potassium ferricyanide + ferrous chloride.

Table 3. Reducing power assay of vitamin C

| Concentration of vitamin C "µg/ml" | Mean (Standard Deviation) |
|-----------------------------------|---------------------------|
| 100                               | 0.201 (0.0280)            |
| 200                               | 0.495 (0.0350)            |
| 300                               | 0.697 (0.0087)            |
| 400                               | 0.992 (0.0727)            |
| 500                               | 1.201 (0.0305)            |

DPPH Free Radical Scavenging Activity (RSA)

The antioxidant activity of all oil extracts were measured in terms of hydrogen donating or radical-scavenging ability using the stable DPPH˙ method as modified by (Potapovich and Kostyuk 2003). The reaction mixture containing 2 ml of all extracts at different concentrations "100,200,300,400,500µg/ml" and 2ml of DPPH˙ (0.2mM) was vigorously shaken and incubated in darkness at room temperature for 30 minutes. When the DPPH˙ reacted with an antioxidant compound in oil that can donate hydrogen, it was reduced and resulting in decrease of absorbance at 517nm using UV-visible spectrophotometer, and the mean values were obtained from triplicate experiments. The
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percentage of the remaining DPPH’ was plotted against the sample concentration. A lower value indicates greater antioxidant activity. Radical scavenging activity was expressed as percent ages of inhibition and was calculated using the following formula:

\[
\% \text{DPPH 'RSA'} = \left( \frac{\text{Abs. of Control} - \text{Abs. of Sample}}{\text{Abs. of Control}} \right) \times 100
\]

3.3. Growth Characteristics of J. Phoenicea

The study samples were divided into lots different tree, depending on the appearance exterior of the tree with three replicates in each tree were made to find out the quantitative and morphological differences.

3.4. Estimation of Photosynthetic Pigments

The photosynthetic pigments were extracted from a known fresh weight of leaves in 85% aqueous acetone to a certain concentration for spectrophotometric measurements. The photosynthetic pigments (chlorophyll a, b and carotenoids) were determined by spectrophotometric method as described by Metzner et al. (1965). The pigments extract was measured against a blank of pure 85% aqueous acetone at three wavelengths of 452.5, 644 and 663 nm. Taking into consideration the dilution factor, it was possible to determine the concentration of pigment fractions (chl. a, b and carotenoides) as mg/ml using following equations:

\[
\text{Chlorophyll a} = 10.3 E_{663} - 0.918 E_{644} = \text{mg/ml}
\]

\[
\text{Chlorophyll b} = 19.7 E_{644} - 3.87 E_{663} = \text{mg/ml}
\]

\[
\text{Charotenoids} = 4.2 E_{452.5} = \text{mg/ml}
\]

Finally, these pigment fractions were calculated as mg/g fresh matter

3.5. The Estimation of Microbial Number

3.5.1. Nutrient Agar medium

23 g of nutrient agar medium were suspened in 1-liter ml distilled water and then heat to boiling to dissolve the medium completely. Sterilized by autoclaving at 15 lbs pressure (121°C) for 15 min.

3.5.2. Pure Culture

A single bacterial colony from each mixed culture was taken by loop, streaked on nutrient agar medium, and incubated at 37.5 °C for three days.

3.5.3. Gram Stain

Differential stains (Gram stain) was carried out to confirm that the bacteria were gram negative and rode shaped. Smears from single colony bacteria grown on EMB at 37 °C and 44.5°C were prepared and covered by primary stain (Crystal violet) for 1 min and then Mordant (Iodine) for 1 min to made CV-I complex after that decolorized agent (70% ethanol) was washed the slide to remove the CV-I complex in the thin layer of peptidoglycan of gram negative bacteria. Wall. Drops of counter stain (safranin) were added for a half minute.was The slide then washed in The distilled water and dried by filter paper after that examined by microscope using an oil immersion lens. In summary, gram-positive cells retain the dye and remain purple, Gram-negative cells did not retain the dye; they were colorless until counterstained with a red dye (pink cells).

3.5.4. Endospore Staining

A film was made on a clean slide by emulsifying part of a colony in loopful of distilled water. The film was then air-dried and stained with malachite green for 4-5 min, using a flame. The smear was rinsed rapidly with water and stained with safranin solution for 30 sec. The slide was washed with water and allowed to dry. On microscopic examination the endospores appeared green and the cells were pink (Abualdahab and Gorani, 1983).
4. RESULTS

4.1. Antioxidant Evaluation of Essential Oils Extracted from Leaves of Arbutus Pavarii Pamp. (Ericaceae).

The antioxidant activities of essential oils extracted of three samples from leaves of Arbutus pavarii Pamp. (Ericaceae) growing in Al-Jabal Al Akhdar were evaluated by:

4.1.1. Total Phenolic Content (TPC):

Figure (9) show the total phenolic content found in essential oils where the essential oil of sample number (1) contained high total phenolic content, the results expressed according to pyrogallol as phenolic compound in figure (6).

4.1.2. Total Flavonoids Content (TFC):

The results obtained in this study as shown in figure (10) indicated that the essential oil of healthy sample contain high amounts of flavonoids compounds as compared with the quercetin in figure (7) which was used as standard.

4.1.3. Reducing Power Assay (RPA):

As shown in figure (11) the reducing power assay of essential oil of sample 1 and sample 2 exhibit higher reducing activity than the ascorbic acid.

4.1.4. The DPPH' Radical Scavenging Activity:

The result of the DPPH' radical scavenging activity of essential oil are shown in figure (12), this result compared with the well-known antioxidant ascorbic acid were the percent of the inhibition is 97% at 500 µg/ml of the essential oil of sample 1 and 92% at 500 µg/ml of the essential oil of sample 2, while the percent of the inhibition is 38% at 500 µg/ml of the essential oil sample 3.

Table 4. Total phenolic content (TPC) of essential oils extracted of three samples from leaves Arbutus pavarii (Sample1= plant intact naturally grow, sample2= plant growth almost normal, sample 3= No plant grows naturally) and % variation from the corresponding control (pyrogallol as standard)

| concentration "µg/ml" | Mean phenolic content (Standard Deviation) |
|-----------------------|------------------------------------------|
|                       | Pyrogallol | Sample1 | % var. | Sample2 | % var. | Sample3 | % var. |
| 100 (1)               | 0.410 (0.032) | 0.711 (0.03) | 73.4↑ | 0.699 (0.05)↑ | 70.5↑ | 0.025 (0.0162)↑ | 97.5↓ |
| 200                   | 0.799 (0.0220) | 0.799 (0.01)↑ | Zero | 0.782 (0.01)↑ | 2.13↓ | 0.085 (0.020)↑ | 89.4↓ |
| 300                   | 1.333 (0.0045) | 1.435 (0.02)↑ | 7.65↑ | 0.988 (0.002)↑ | 31.2↓ | 0.222 (0.022)↑ | 84.5↓ |
| 400                   | 1.828 (0.0117) | 1.78 (0.01)↑ | 2.63↑ | 1.56 (0.016)↑ | 12.4↓ | 0.754 (0.036)↑ | 57.6↓ |
| 500                   | 2.105 (0.0225) | 2.69 (0.07)↑ | 27.8↑ | 2.21 (0.009)↑ | 17.8↓ | 1.061 (0.037)↑ | 60.6↓ |

† Insignificant difference from the corresponding control at P > 0.1
* Significant difference from the corresponding control at P < 0.05
** Highly sig. difference from the corresponding control at P < 0.01
*** Very highly sig. difference from the corresponding control at P < 0.001
↓ Decrease
↑ Increase

Fig9. Total phenolic content (TPC) of essential oils extracted from leaves Arbutus pavarii and Pyrogallol as phenolic compound.
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Table 5. Total flavonoids content essential oils extracted of three samples from leaves Arbutus pavarii (Sample1= plant intact naturally grow, sample2= plant growth almost normal , sample 3= No plant grows naturally) and % variation from the corresponding control (quercetin as standard).

| Concentration " µg/ml" | Mean phenolic content (Standard Deviation) | % var. | % var. | % var. | % var. |
|------------------------|------------------------------------------|--------|--------|--------|--------|
|                        | quercetin | Sample1 | Sample2 | Sample3 | Sample3 |
| 100                    | 0.279 (0.092) | 0.297 (0.032)† | 0.276 (0.002)† | 1.07↓ | 1.07↓ |
| 200                    | 0.560 (0.035) | 0.497 (0.011)† | 0.375 (0.001)* | 30.1↓ | 1.095 (0.045)** |
| 300                    | 0.834 (0.003) | 0.664 (0.002)† | 0.609 (0.008)† | 27.0↓ | 28.0↓ |
| 400                    | 1.066 (0.009) | 0.820 (0.090)† | 0.869 (0.003)† | 18.4↓ | 20.4↓ |
| 500                    | 1.300 (0.006) | 1.690 (0.011)* | 1.340 (0.001)† | 3.08↑ | 3.34↑ |

† Insignificant difference from the corresponding control at P > 0.1
* Significant difference from the corresponding control at P < 0.05
** Highly sig. difference from the corresponding control at P < 0.01
*** Very highly sig. difference from the corresponding control at P < 0.001
↓ Decrease
↑ Increase

Fig 10. Total flavonoids content of essential oils extracted from leaves Arbutus pavarii and quercetin as flavonoid compound.

Table 6. Reducing power assay essential oils extracted of three samples from leaves Arbutus pavarii (Sample1= plant intact normally grow, sample2= plant growth almost normal , sample 3= No plant grows naturally) and % variation from the corresponding control (vitamin C).

| Concentration " µg/ml" | Mean phenolic content (Standard Deviation) | % var. | % var. | % var. | % var. |
|------------------------|------------------------------------------|--------|--------|--------|--------|
|                        | vitamin C | Sample1 | Sample2 | Sample3 | Sample3 |
| 100                    | 0.201 (0.0280) | 0.503 (0.0448)** | 0.564 (0.07)** | 180.6↑ | 0.044 (0.006)** |
| 200                    | 0.495 (0.0350) | 0.875 (0.0965)** | 0.871 (0.0)*** | 75.9↑ | 0.104 (0.002)** |
| 300                    | 0.697 (0.0087) | 1.293 (0.0471)** | 1.321 (0.003)*** | 89.5↑ | 0.432 (0.005)** |
| 400                    | 0.992 (0.0727) | 1.563 (0.0266)*** | 1.899 (0.01)***** | 91.4↑ | 0.902 (0.016)*** |
| 500                    | 1.201 (0.0305) | 2.339 (0.0401)***** | 2.486 (0.01)***** | 107.0↑ | 1.052 (2.965)*** |

† Insignificant difference from the corresponding control at P > 0.1
* Significant difference from the corresponding control at P < 0.05
** Highly sig. difference from the corresponding control at P < 0.01
*** Very highly sig. difference from the corresponding control at P < 0.001
↓ Decrease
↑ Increase
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Fig 11. Reducing power assay of methanol, ethanol, and acetone extracted from leaves Arbutus pavarii and vitamin C.

Table 7. DPPH radical scavenging activity of vitamin C, and essential oils extracted of three samples from leaves Arbutus pavarii according to % inhibition.

| Concentration (µg/ml) | vitamin C | Sample 1 | Sample 2 | Sample 3 |
|-----------------------|-----------|----------|----------|----------|
|                       |           | % var.   | % var.   | % var.   |
| 100                   | 92.3%     | 57% *    | 38.2%    | 39% *    |
|                       |           |          | 57.7%    | 59% *    |
| 200                   | 93.8%     | 79% †    | 15.8%    | 75% †    |
|                       |           |          | 37.1%    | 21.1%    |
| 300                   | 95.1%     | 85% †    | 10.6%    | 79% †    |
|                       |           |          | 38%      | 27% †    |
| 400                   | 95.8%     | 89% †    | 7.1%     | 87% †    |
|                       |           |          | 9.19%    | 32% †    |
| 500                   | 96.7%     | 97% †    | 0.31%    | 90% †    |
|                       |           |          | 6.93%    | 38% **   |

† Insignificant difference from the corresponding control at P > 0.1
* Significant difference from the corresponding control at P < 0.05
** Very highly sig. difference from the corresponding control at P < 0.001
↓ Decrease
↑ Increase

Fig 12. DPPH radical scavenging activity of vitamin C and essential oils extracted from leaves Arbutus pavarii according to % inhibition.

4.2. The GC-MS Analysis of the Essential Oils Extracted from Leaves Arbutus Pavarii:

The retention time of the different compounds in the essential oils of Arbutus pavarii are presented in Table 8. Thirty-four volatile compounds, representing 88.29% of the total oil composition in sample 1, were identified in the leaves oils (Table 8). Monoterpenes hydrocarbons were found to be the major group of compounds. The most abundant component found in the leaf oil was α-pinene (20.85%) followed by Germacrene D (16.49%), while α-pinene and Germacrene D in sample 2 and sample 3 were (12.45%, 3.09%) and (9.19%, 1.05%) respectively. Surprisingly, there were decrease of these compounds in the sample diseased (table 8).

Table 8. The main components of essential oils of Arbutus pavarii L. leaves collected from Al-Jabal Al Akhdar area (A sulnta)

| Constituents | sample 1 | Sample 2 | Sample 3 |
|--------------|----------|----------|----------|
| α-pinene     | 20.85    | 12.45    | 3.09     | 4.16     |
β–myrcene & 1.36 & 0.23 & - & 5.06 
Terpinolene & 0.07 & - & - & 5.26 
β–phellandrene & 3.84 & 1.79 & 0.02 & 5.91 
Trans-Caryophyllene & 5.44 & 5.00 & 2.087 & 14.66 
4,7,10-Cycloundecatriene,1,1,4,8-tetramethyl-cis,cis,cis & 5.31 & 4.13 & 0.12 & 15.37 
Germacrene D & 16.49 & 9.19 & 1.054 & 16.05 
Germacrene B & 2.73 & 1.72 & - & 17.46 
α – cedrol & 0.36 & - & - & 18.27 
+ alpha-longipinene & 2.98 & 2.082 & 0.471 & 18.08 
Naphthalene,1,2,3,4,4a,5,6,8a-octahydro-7-methyl-4-methylene-1- (1-methylthyl)-(1.alpha,4a.alpha.,8a,alpha) & 5.22 & 2.011 & 0.57 & 16.23 
Camphene & 0.29 & - & - & 4.32 
β-pinene & 0.49 & - & - & 4.81 
1-phellandrene & 0.39 & - & - & 5.33 
delta.3-carene & 0.40 & - & - & 5.46 
α – terpinolen & 0.23 & - & - & 7.10 
Terpinolene & 5.13 & 5.13 & 2.093 & 13.11 
Linalool & 0.47 & - & - & 7.37 
Citronellol & 0.63 & - & - & 10.30 
α-cubebene & 0.39 & - & - & 13.60 
β-bourbonene & 0.22 & - & - & 13.80 
β-elemene & 0.93 & - & - & 13.94 
Widdrene & 1.24 & 0.53 & - & 14.83 
α-gurjunene & 0.35 & - & - & 16.40 
Zingiberene & 0.43 & - & - & 16.92 
delta-cadinene & 5.63 & 5.00 & 3.73 & 16.81 
α-muurolene & 0.43 & - & - & 17.01 
α-calacorene & 0.24 & - & - & 17.10 
Elemol & 0.61 & - & - & 17.28 
caryophyllene oxide & 2.06 & 1.890 & 0.281 & 17.95 
Fonenol & 0.36 & - & - & 18.58 
α-ylangene & 1.33 & 0.072 & - & 19.03 
β- seline & 0.30 & 0.30 & 0.30 & 19.54 
vulgarol B & 1.09 & 0.873 & - & 19.88 

*Rt: Retention time obtained by chromatogram.

Sample1= plant intact naturally grow, sample2= plant growth almost normal and sample 3= No plant grows naturally.

The Results of Morphological and Growth Characteristics are Given in Table (9). From this Table, the Sample Number (1) Reflected the Highest Mean Value of Plant Height and Stem Diameter. They were 222.67 Cm and 16.42 Cm, Respectively.

Table 9. Growth in terms of plant height (cm), stem diameter (cm) and leaf dry weight (mg) of Arbutus pavarii in EL-Gabal EL-Ahkdar area.

| Sample Name      | Plant height (cm) | Stem diameter (cm) | Leaf dry weight (mg) |
|------------------|-------------------|--------------------|---------------------|
|                  | Mean (SD)         | Mean (SD)          | Mean (SD)           |
| Healthy sample   | 222.67 (21.35)    | 16.42 (5.90)       | 3.0234 (0.19)       |
| Semi healthy sample | 216.55 (17.93)    | 11.22 (3.14)       | 3.2058 (0.23)       |
| Sample sick      | 214.23 (23.47)    | 9.89 (2.05)        | 3.8240 (0.11)       |
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The Results Obtained in this Study as Shown in Table(10) Indicate that the Healthy Sample Contain High Amount of Chlorophyll A, Chlorophyll B and Carotenoids as Compared with the Semi Healthy Sample and Sample Disease.

Table10. Estimation of photosynthetic pigments of Arbutus pavarii leaves in EL-Gabal EL-Ahkdar area.

| Sample          | Chlorophyll A (mg/ml) | Chlorophyll B (mg/ml) | Carotenoids (mg/ml) |
|-----------------|-----------------------|-----------------------|---------------------|
| Healthy sample  | 5.7708                | 4.894                 | 6.323               |
| Semi healthy sample | 5.670                | 4.721                 | 5.126               |
| Sample sick     | 4.346                 | 2.915                 | 3.503               |

Fig13. Estimation of photosynthetic pigments of Arbutus pavarii leaves in EL-Gabal EL-Ahkdar area

As shown in Fig (14) decreased the number of bacteria in the sub-branches of Semi-healthy sample but increased the number of bacteria in diseased sample. The number of bacteria in the sample healthy at the main branch of the highest less than the bottom, but in the semi-healthy sample, the number of bacteria from the top down and more of them there are more in the center, while in diseased sample, the number of bacteria is too much and this shown in Fig (15).

In addition, the number of bacteria in lichens the number of bacteria in three samples and an equal number was more than 300, as shown in the Fig (16).

Through our results was observed that the number of bacteria in the green leaves of the sample healthy sample more than semi-healthy sample, as shown in Fig (17), while the number of bacteria in the brown leaves in all samples were large Fig (18).

Fig (19) indicated that the number of bacteria taken of selected locations of Arbutus pavarii tree in an environment actinomycetes were a few specific sites, but the number of bacteria is high in lichens and green leaves and soil.

Fig14. The number of bacteria on the sub-branches in Arbutus pavarii.
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Figure 15. The number of bacteria on the main branch in Arbutus pavarii.

Figure 16. The number of bacteria on lichens in Arbutus pavarii plant.

Figure 17. The number of bacteria on green leaves in Arbutus pavarii plant.

Figure 18. The number of bacteria on brown leaves in Arbutus pavarii plant.
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Figure 19. The number of bacteria and total Actinomycetes mass

Figure 20. The number of bacterial colonies in soil samples

Figure 21. The number of bacteria in the Nutrient Agar media taken by Wipes.

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