ANTI-MYCOBACTERIAL ACTIVITY OF CAPPARIS SPINOSA L. EXTRACT AGAINST CLINICAL ISOLATES OF MYCOBACTERIUM TUBERCULOSIS

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INTRODUCTION

Tuberculosis (TB) is a social disease with the medical features which is described as a barometer of social welfare [1]. TB, the biggest factor causing death among infectious disease in the world [even more than AIDS, malaria and measles] ranks tenth in the global burden of disease 2020 and is expected to maintain its current position or rise to the seventh position [2].

In Africa, many plants are used in traditional medicine to treat TB and other respiratory diseases. The emergence of drug resistance and its conversion into a major threat require an urgent search for anti-TB agents which are effective and safe to consume [3].

TB is a disease caused by a known factor and known epidemiology and its treatment principles are known for about 60 years, and also it is more than a quarter of a century that a short-term treatment regime is used for it; however, some patients are still undiagnosed or not treated appropriately. Unfortunately, the results of these shortcomings and at the same time the increasing incidence of HIV, prepare the conditions for emergence, and spread of multidrug-resistant TB bacillus [2].

Mycobacterium is an aerobics rod-shaped bacillus which does not form spore and gain its energy from simple carbon compounds' oxidation. Moreover, the biochemical activity of bacteria is very slow. They have high chemical agents’ resistance than other bacteria, due to the hydrophobic nature of the mycobacteria surface and their mass growth. Drug-resistant strains pose a serious challenge to global health, in addition to their need for a long-term treatment; they also have side effects. Therefore, the use of herbal remedies as an alternative or complementary for industrial medications gains considerable importance [4].

This Capparaceae have 30 genera and 650 species, and they grow mainly in warm and temperate areas and grow like shrubs. Their bushes are crawling with long life and downy, and they have ramified branches with up to 1.5 m length. This plant has a long root that penetrates soil deeply. Its large flowers have white. Fruits come in different ways such as capsule, pod, and broken shafts and seeds are without albumen content. It has a wide distribution. It grows well in sabulous lands. All parts of the plant such as fruits, roots, bark, leaves, buds, and stems can be used. Its fruit graded by the size in 8 different sizes and export from Iran with the best quality. This plant tolerates 40°C on summer, and due to its mass growth, it has a wide distribution. It grows well in sabulous lands. All parts of the plant such as fruits, roots, bark, leaves, buds, and stems can be used. Its fruit graded by the size in 8 different sizes and export from Iran with the best quality. This plant tolerates 40°C on summer, and due to its

Capparis spinosa L. (Capparaceae; common name: Caper) is a plant from the 1-year herbaceous plant families that grows as shrubs and often covered with glandular. Studies conducted in most parts of the world have been found that most species have anti-bacterial effects and natives consume some species as a traditional medicine. This plant grows in the most parts of the Middle East a wild plant, but not much research has been done on it [5].

Herbal medicines have been used as a non-comprehensive metabolite for pharmaceutical development also as an alternative treatment for human diseases [5]. They contain many active components that have therapeutic value [8]. Iran is a vast country and as a result of its geographical situation on earth, it is a confluence point of several different climates; thus, owns various weather. These climatic conditions can cause rare plant flora on this country. Phytochemical studies on 1700 different plants show that Iran’s plants possess very good quality and quantity [9].

Keywords: Mycobacterium anti-bacterial, Plant Capparis spinosa, Minimum inhibitory concentration.
tendency toward the cold weather, it can tolerate -8°C on winters. It more likes the calcareous soils. Although this plant is cold in nature, it grows even in the dry and nutrient poor soils. Flowers grow on the 1st-year branches and they opened by the sunrise and closed by sunset. Its flowers should be harvesting before unfolding. At the end of season plant’s fruit will split and seeds fall. Capparis has several chemical active compounds but the important one is flavonoid. Roots and flower buds contain perin, saponin, amisin material, aminoglycosides, and substance called kaparynotin. Octadeconoic acid-hexadecanoic acid-octadec-9-enolic acid- 1, 2 - benzene dicarboxylic acid-bis(2-methylpropyl) ester- (Zethylhexylyl) phthalate-2-methoxy-phenol- tetradecanoic acid - dodecanolic acid >2, 9 - 12 - octadecadienoic acid-ethyl ester -1-(1H-pyrrol-2-yf) - ethanone - hexadecanoic acid - methyl ester [11].

The first use of this herb for medicinal purposes assigned to the Sumerians. In the traditional medicine, it is used as support and maintenance, liver stimulant, and it improves the function of the liver. In addition, it is used to reduce bloating, atherosclerosis of the vessels, and as an anti-rheumatic. Moreover, it is reported as a worm excretion factor, diuretic, and nutrition [12].

In traditional medicine, the decoction of the root’s bark is utilized to heal the inflammation of serious fluid in the capillary and relieve arthritis, and gout. In addition, in previous studies, anti-TB property of this pant has been reported. In addition, recent studies confirm its beneficial effect on hysteria and neural states [12].

METHODS

Referring to the suburbs and surrounding villages in Sistan, the Capparis spinosa samples were collected in July 2015. The fruits were then washed and dried in the shade.

Preparation Capparis spinosa fruit extracts

Of the 300 g from powder measured by the ratio of 1-5 was dissolved in methanol. It was then placed on a timer shaker for 48 hrs at 150 rpm, then filtered out by filter paper and poured over the three petri dishes. To accelerate the sublimation of methanol, the samples were incubated at 40°C. After 3 days, the samples were placed in a freeze dryer.

Microorganism preparation

The sample were collected from Sistan and Baluchestan urban and rural health-care centers and sent to test for TB to the Southeast of the country. The obtained samples from medium cultured on Lowenstein-Jensen culture Media. After 6 weeks, the pipes were assessed and it was clear to see that the mycobacterium colonies and the medium were half-covered. A Ziehl-Neelsen test was performed for recognition.

For reference, culture H37RV standard samples were used. Standard samples were collected from the Tehran reference TB. All seven clinical samples were tested to determine the species. First, we had to determine the growth rate. For this purpose, prepared culture medium on Lowenstein-Jensen culture medium and it was evaluated daily for the presence of colonies. The growing temperature was assessed, and for this purpose 5 different plates were prepared and incubated at 44-42-37-33-30-25°C. Samples were examined for colonies on days 3-7-14-21. The colony morphology was assessed and a pigment production test was performed. To do this, both samples were inoculated in 2 different mediums. One was then covered with aluminum foil and incubated for 18 days at 37°C. Next, a nitrate resuscitation test was conducted and the number of colonies of each sample added separately to 0.1% sodium nitrate solution and incubated for 4 hrs in Ben Murray at 37°C then 2 drops of normal hydrochloric acid, 2 drops of sulphuric acid 0.2% and 2 drops the solution N-1-naphthyl diethyl diamine hydrochloride 0.1% added to tube and color examined. Next stage testing niacin, for this test about 1 mL of Sterile Distilled water poured into medium with 50 colonies at least, waiting for 30 minutes and then, 0.5 mL of solution of top on the medium transferred to sterile tube. 0.5 mL of Aniline represented 4% and 0.5 mL of cyanogen bromide reagent added to tube, color examines helped diagnosis.

Preparing Levin-Stein Johnson vitro containing Capparis spinosa fruit extract

All the test equipment was sterilized using autoclave at a temperature of 121°C at 15 minutes and cooled by keeping them at room temperature. To make the egg suspension, first the eggs were cleaned with a brush and placed in a soap solution for 1-5 minutes. Then, these were washed in cold water and placed in 70% alcohol for 15 minutes. After being washed by hand in soap and water, the eggs were broken into a sterile electric mixer container and blended smoothly. The resulting mixture was passed through four layers of typical sterile filter cloth and poured into a 2 L jar. A salt solution was made next by adding malachite green 2% solution. After adding the salt solution and malachite solution to the eggs, they were placed on a hot plate so they could become mixed [10].

To prepare the fruit juice, a concentration of C. spinosa and inoculated that to the under making medium. The weight of the empty plate was calculated. Then, after adding some extract and reassessing the weight of the plate, 30 mg dimethyl sulphoxide was added to dissolve completely on 2.5 mg of extract solvent. The egg solution was divided into 1.5 mL in 6 different containers. Using $C_1 V_1 = C_2 V_2$ formula and adding the extract to a Lowenstein-Jensen solution, 150 mg/mL concentration was gained. Then, 8.6 mL of each concentration was poured into a McCartney tube and the lid tightly closed. This was flocculated diagonally on a coagulator at 85°C for 50 minutes and then, placed in an incubator at 37°C for 24 hrs. After health checking, it was placed in a refrigerator with a temperature of 2-8°C.

Prepare the witness vitro and positive control to determine the possible resistance of microbes

A medium was prepared without any additive for witness purposes. In addition, to ensure that the samples were not resistive, all clinical samples, and reference samples had two witness tubes without the extract and contained anti-mycobacterium. On day two, the mediums were examined for steam and contamination and then, carried to the microbial culture room [3].

Microbial stoke and culture preparation

First, the cabinet was disinfected with 25 mL formaldehyde and 15 g potassium permanganate. Then, the hood was turned on and the culture began 30 minutes later. Eight McCartney tubes with 8-10 sterile Pearl Glasatroom temperature was taken and with disposable sterile lobe, one complete lobe (8-10 colonies) were moved from the medium of the first colonies to the tube and the lid was closed tightly and placed on the shaker for 4-5 minutes. Then, after waiting for 5 minutes, the tube lid was opened. 2cc distilled water was added and the lid was closed again and shaken for 4-5 minutes, then rested for 45 minutes so that the tube contents remain stagnant. The procedure was repeated for each sample and reference sample.

During the next stage, we created a McFarlane. Sulfuric acid 1% (990 mL) + 1% barium chloride (10 mL). Then, with sampler head lifting 0.1 mL of top of liquid, the colony suspension was added to the tube with 1cc distilled water; by adding distilled water, it matched a McFarlane. 500 μl of suspension were picked up with the sampler and added to the 4.5 mL distilled water tube and diluted 10 to the power of minus one. Then, 0.1 mL of the previous solution was added to 9.9 mL distilled water that created 10 to the power of minus two equivalent (3 multiplied by 10 to the power of 6) of bacteria on each mL. Then, 1cc of stock solution was added to the prepared medium, and the cultured tubes were placed in an incubator at 37°C for 4 weeks.

The method used to determine the extract effect on the examined samples was the proportional method. This method is a standard
procedure and is approved by the World Health Organization and Clinical and Laboratory Standards Institute [13]. This method needs 3-4 weeks’ incubation.

RESULTS

Based on species diagnostic tests at the end all of samples were reported as Mycobacterium TB. After adding brown extract to Lowenstein-Jensen medium, medium color changed from green inclined white to dark green.

The statistical examination of this study was carried out using the normality check of the Kolmogorov-Smirnov test, while the Significance level was checked by the Spearman Kendall test.

DISCUSSION

During this research, the methanol extract of C. spinosa was prepared using the maceration method and freeze-dried. 7 clinical samples and one Mycobacterium TB sample were isolated and identified. According to the results, the C. spinosa extract with 25 mg on mL minimum inhibitory concentration (MIC) was effective on 6 clinical samples. On the second sample, with 10 mg on mL MIC of methanol, C. spinosa fruit extract was effective. The H37RV standard strain culture result with 25 mg on mL MIC of methanol C. spinosa fruit extract was effective. It is most probable that the freeze-drying method has the best stability on the known and unknown material of C. spinosa fruit extract, according to the extract methods and their impact on the stability of the effective plant compounds.

This research used the best method to preserve plant properties. According to research conducted Institute of Medicinal Plants (SID), compared with other methods of extraction, such as thermal dryer, spray dryer, and rotary evaporator, using the freeze-dryer method has the most stability of medicine plants' known contents [14]. The other research is considered one of the first studies in Iran on medicine plants used on mycobacterium, who study shallot extract effects considered one of the first studies in Iran on medicine plants used on Mycobacterium. The results were effective on 0.1 mg on mL. Another study assessed the effect of ethanol on the growth of Mycobacterium TB strains. Based on this study, concentrations higher than 5.2% of ethanol prevented growth, but on 0.1-0.5% density has shortened growth time to 3-8 days. Assessing the antimicrobial effects of Salvadora persica water extract at Tehran’s Medical University by Fallah et al with 10 mg on mL was effective [6].

In another study at the University of Spain in 2011, Lea et al. assessed Mozambique’s traditional medicine for phytochemical properties. 15 medicinal plants extracts with hexane-dichloromethane-ethyl acetate - ethanol dissolver, at the same time a decoction of these plants was prepared for traditional use. In the face of two species, Mycobacterium asngmatys H37RV with 1 inhibitory concentration 125 mg on mL was effective [9]. Evaluated the toxicity and anti-inflammatory genes of 10 plants used to cure TB in South Africa. Extraction was performed by 4 dissolvers, petroleum-ether-dichloromethane-ethyl 80% water and using reference samples H37Ra. 6 plants with 1 mg on mL density have suitable activity and water extract known worthless. An assessment of herbal medicines used for curing bacterial infectious diseases in Soudan was carried out by [15]. They assessed 50 extracts from 46 medicinal plants, extracted with hexane-chloroform-ethyl-acetate-n-butane and water. The results show that except for the water extracts, the rest of the extracts with 6.25 mg on mL were effective. A study conducted by [16] in Ghana assessed the treatment of TB with 15 species of plants from 13 different genera. Among these plants, the Amaryllidaceae had the greatest anti-mycobacterium effect. A study in Canada analyzed the anti-mycobacterium effects of Aralia nudicaulis. The plant’s methanol extract and H37RV sample in liquid medium proved to be effective [17].

A survey conducted on the Citrullus colocynthis oil and chloroform and methanol extracts over the Mycobacterium other than TB sensitive and resistive strains, with 16 clinical samples in India by [17], was effective on 62.5 mg on mL over sensitive strains and 125 mg on mL for resistive strains.

In a study assessed the effect of additives monocarboxyl analog of curcumin to prevent the growth of sensitive and resistant anti-TB drug strains. At Atlanta University, Baldwin et al. (2015) [18] carried out a disk diffusion method and liquid measuring on laboratory samples and resistant strain to rifampin. The synergistic effect was confirmed with rifampin. Because of poor bioavailability, it was not appropriate to use curcumin. By Singh et al. (2013) [19] screened for anti-mycobacterial activity of 5 herbal plants using a tube dilution method on M. asngmatys and clinical strains. Prepared Hexan and methanol extracts were effective on 125 and 250 mg/mL.

An overview study was conducted by Gautam et al. (2007) [7] on herbal medicines in India and their anti-mycobacterium effects. By diluted test method macro and micro agar for H37RV sample on middle brook medium and Alamar blue colorimetric system was used and received a wide result. This article reviews the research of 365 plant species between almost 17,500 plant species that show a wide range of anti-Mycobacterium TB. The most interesting part of this research correct confirmation of indigenous medicine and (yvrvdayy) about effective
Table 1: The result MIC of methanol extract of fruits Capparis spinosa on 7 clinical sample and a standard sample and H37Rv

| Extract mg on medium mL | 80 mg | 50 mg | 25 mg | 10 mg | 1 mg | 0.2 mg |
|-------------------------|-------|-------|-------|-------|------|-------|
| First sample            | 0     | 0     | 0     | 0     | 0    | 34.4  |
| Second sample           | 0     | 0     | 0     | 0     | 34.4 | 36.2  |
| Third sample            | 0     | 0     | 0     | 0     | 50   | 86.2  |
| Fourth sample           | 0     | 0     | 0     | 0     | 50   | 86.2  |
| Fifth sample            | 0     | 0     | 0     | 0     | 50   | 50    |
| Sixth sample            | 0     | 0     | 0     | 0     | 50   | 50    |
| Seventh sample          | 0     | 0     | 0     | 0     | 50   | 50    |
| H37Rv                   | 0     | 0     | 0     | 0     | 50   | 50    |

**MIC: Minimum inhibitory concentration**

MIC: Minimum inhibitory concentration, **< 0.001

Table 2: The relation between MIC significantly different from the concentration of the methanol extract of fruits Capparis spinosa marbler clinical sample and a standard sample H37Rv based on 7 compared to control medium

| Statistical analysis | Group | Control |
|----------------------|-------|---------|
| Kendall's tau-b      |       |         |
| Group                |       |         |
| Correlation coefficient | 1.00 | -0.783** |
| Significant (two-tailed) | -   | 0.000   |
| N                    | 56   | 56      |
| Control              |       |         |
| Correlation coefficient | -0.783** | 1.00    |
| Significant (two-tailed) | 0.000 | -       |
| N                    | 56   | 56      |
| Spearman's rho       |       |         |
| Group                |       |         |
| Correlation coefficient | 1.00 | -0.906** |
| Significant (two-tailed) | -   | 0.000   |
| N                    | 56   | 56      |
| Control              |       |         |
| Correlation coefficient | -0.906** | 1.00    |
| Significant (two-tailed) | 0.000 | -       |
| N                    | 56   | 56      |

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