Effect of Plant-Products Fumigation on Air-Borne Microbes

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Authors’ contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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ABSTRACT

Background and Objective: Purity of the ambient air is essential for our health and well-being. One of the main factors influencing air quality is the presence of microbes. Fumigation of herbs has been recommended in Unani medicine to purify the air. Hence, the present study was aimed to evaluate the effect of selected Unani herbs fumigation on air-borne microbes.

Methods: In this study, the effect of fumigation with Unani Medicinal herbs powder on air-borne microbes was assessed using differences in total colony counts of microbes in pre and post fumigation samples. Microbial load in the air was quantified using the passive open-air petri plate method. Formalin and potassium permanganate served as positive control, while tamarind wood charcoal fumigation served as negative control.

Results: Fumigation with Unani medicinal herbs powder at a dose of 45 grams was found to be the most effective in reducing the microbial load of the air. Significant reduction in aerial microbial colonies was observed with fumigation at 30 and 45 gms in the fumigated areas (P<0.05).

Conclusion: It can be inferred from the findings of the present study that the test drugs fumigation efficiently reduces the air-borne microbes, hence may be recommended for air disinfectant. However,
various factors that were not considered in this study, such as the effect of temperature and humidity on disinfection efficiency of fumigants should be addressed in further studies.

Keywords: Unani medicine; fumigation; air disinfection; air-borne microbes.

1. INTRODUCTION

On average, a person inhales approximately 14,000 litres of air per day. Therefore, clean air is a basic requirement of life and is essential to human health and well-being [1]. Since people spend the majority of their time indoors, indoor air quality has a significant impact on our health and lives [2]. Recognizing the importance of clean air in health promotion, several editions of air quality guidelines were issued by World Health Organization (WHO). In 2006, global update on air quality guidelines issued by W.H.O. highlighted the significant impact of indoor air pollution on health [3].

One of the most common factors affecting indoor air quality is the presence of bioaerosols. Bioaerosols are particulate matter usually associated with compounds of biological origin such as bacterial cells and cellular fragments, fungal spores and fungal hyphae, viruses, and by-products of microbial metabolism [4]. These are considered to be responsible for approximately 5 to 34% of indoor air pollution and are produced by a combination of natural and anthropogenic activities. Some of the microbial aerosols produced during these activities may be infectious. Exposure to these microbial aerosols can result in a variety of adverse health effects such as infectious diseases, acute toxic effects, allergies, and cancer [5,6].

Therefore, reducing microbial aerosols in indoor environments is essential for the prevention of various airborne diseases. Various methods such as air filters, chemical fumigants are used to reduce microbial aerosols [7]. However, in developing countries, chemical fumigation such as hydrogen peroxide, formalin is frequently used for air disinfection, which is associated with various toxic side effects [8]. Hence, there is a necessity to find natural and safe alternative method of air disinfection.

Fumigation of plant-based products has been described by Unani scholars as a method to purify the indoor air of residential places [9-14]. The air purification method described in Unani literature corresponds to environmental disinfection procedures of modern science. However, the efficacy of fumigation using these plant products in improving the microbiological quality of air has not been scientifically validated.

Therefore, the present study was carried out to validate the traditional fumigation method using some Unani medicinal herbs to improve the microbiological quality of air in relation to formalin, so that ancient knowledge of Unani scholars can be unfolded to practical application. Herbs used in this study were selected from a renowned Unani treatise, Kitâbal-Mansûri, authored by Zakariya Râzî in 9th century A.D. concerning their availability and potential.

2. MATERIALS AND METHODS

2.1 Study Design and Setting

The in-situ experiment was designed to compare the efficacy of traditional fumigation of Unani medicinal herbs with formalin plus potassium permanganate fumigation in improving the microbiological quality of indoor air. The efficacy of fumigation on aerial microbes' population was assessed through differences in total colony counts (CFU) of microbes in pre and post samples. The experiment was conducted in three different locations of the National Institute of Unani Medicine campus in the months of September to December 2020. Prior to the start of the study, air samples were collected from various locations and the microbial load in the air was estimated; Places with the highest microbial load were chosen for the study. The dimensions of the three selected locations were 1200, 900, and 700 cu. ft. respectively.

2.2 Air Sampling

Passive open-air petri plate exposure method was used to quantify the microbial load in the air. Air sampling was done before fumigation and at periodic intervals, after fumigation (1hr, 4hrs and 12 hrs). For sampling pre-cultured, sterilized petri-plates were placed in the desired locations according to the 1/1/1 scheme (1 m above the floor, about 1m away from walls, for 1 hour) after closing the doors of the rooms [15]. To isolate the aerobic and anaerobic bacterial colonies, Luria Bertani (LB) medium and for fungi, potato...
The aerial microbial population has been presented in the form of colony-forming units (CFU). Reduction in microbial colonies due to fumigation has been presented in the form of a percentage reduction. For intragroup comparison, Friedman test for repeated-measures was used to determine the significant effect of each fumigant in reducing the microbial colonies. For intergroup comparison, the kruskal-wallis test was used. P-value less than 0.05 was considered significant.

Fumigation with Unani formulation at 30 gms ($P=0.058$) and 45 gms ($P=0.01$) dosage resulted in a significant reduction in microbial colonies in area 1 (1200 cu. ft. room). Intergroup comparison revealed a significant effect on microbial colonies at 4 and 12 hours after fumigation ($P=0.057$) (Table 1). In area 2 (900 cu. ft. room), the only fumigation of Unani formulation at 45 gms resulted in a significant reduction in microbial colonies ($P=0.01$); however, on the intergroup comparison, the effect was found to be insignificant ($P>0.05$) (Table 2). Fumigation with Unani formulation at 30 gms ($P=0.056$) and 45 gms ($P=0.01$) dosage resulted in a significant reduction in microbial colonies in area 3 (700 cu. ft. room). The effect was found to be significant at 1 hour ($P=0.051$) and 4 hours ($P=0.055$) after fumigation in an intergroup comparison (Table 3).
Table 1. Comparison of the effect of different fumigants on aerial microbial population (CFU) in Area 1 [1200 cubic feet]

| Fumigants                        | Sampling Time Before Fumigation | After Fumigation | Significance |
|----------------------------------|--------------------------------|------------------|--------------|
|                                  | CFU | 1 hrs | 4 hrs | 12 hrs | CFU | % | CFU | % | CFU | % |
| Tamarind Wood Charcoal*          | 22  | 22    | 0     | 24     | 9   | 23  | 4.5 |
| Unani Formulation (5gms)         | 53  | 50    | 5.66  | 50     | 5.66| 52  | 1.88|
| Unani Formulation (10gms)        | 61  | 48    | 21.3  | 49     | 19.6| 57  | 6.55|
| Unani Formulation (15gms)        | 38  | 24    | 36.8  | 29     | 23.6| 33  | 13.16|
| Unani Formulation (30gms)        | 35  | 11    | 68.57 | 14     | 60  | 18  | 48.57|
| Unani Formulation (45gms)        | 58  | 11    | 81.03 | 15     | 74.14| 18 | 68.97|
| Formalin and potassium permanganate | 29  | 22    | 24.13 | 24     | 17.24| 9  | 68.97|

Significance

|        | P=0.819 | P=0.135 | P=0.156 | P=0.156 |

*Increase in microbial colony counts observed. % indicates reduction in aerial microbial colonies. *P< 0.05 considered significant

CFU=Colony Forming units.

Table 2. Comparison of the effect of different fumigants on aerial microbial population (CFU) in Area 2 [900 cubic feet]

| Fumigants                        | Sampling Time Before Fumigation | After Fumigation | Significance |
|----------------------------------|--------------------------------|------------------|--------------|
|                                  | CFU | 1 hrs | 4 hrs | 12 hrs | CFU | % | CFU | % | CFU | % |
| Tamarind Wood Charcoal*          | 21  | 21    | 0     | 23     | 9.5 | 23  | 0   |
| Unani Formulation (5gms)         | 53  | 52    | 1.8   | 51     | 3.7 | 53  | 0   |
| Unani Formulation (10gms)        | 57  | 50    | 12.2  | 49     | 14.03| 54  | 5.2  |
| Unani Formulation (15gms)        | 42  | 24    | 42.8  | 26     | 38.09| 32  | 23.8 |
| Unani Formulation (30gms)        | 28  | 7     | 75    | 11     | 60.71| 14  | 50  |
| Unani Formulation (45gms)        | 60  | 10    | 83.3  | 12     | 80  | 15  | 75  |
| Formalin and potassium permanganate | 34  | 27    | 20.5  | 23     | 32.23| 10  | 70.5 |

Significance

|        | P=0.061 | P=0.061 | P=0.061 | P=0.061 |

*Increase in microbial colony counts observed. % indicates reduction in aerial microbial colonies. *P< 0.05 considered significant

CFU=Colony Forming units.

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Table 3. Comparison of the effect of different fumigants on aerial microbial population (CFU) in Area 3 [700 cubic feet]

| Fumigants                        | Before Fumigation CFU | After Fumigation 1 hrs CFU | % | After Fumigation 4 hrs CFU | % | After Fumigation 12 hrs CFU | % | Significance |
|----------------------------------|-----------------------|----------------------------|---|-----------------------------|---|-----------------------------|---|--------------|
| Tamarind Wood Charcoal*          | 29                    | 28                         | 3.4| 28                         | 3.4| 30                         | +3.4 | P=0.112      |
| Unani Formulation (5gms)         | 52                    | 49                         | 5.7| 51                         | 1.9| 52                         | 0   | P=0.145      |
| Unani Formulation (10gms)        | 59                    | 49                         | 16.9| 49                         | 16.9| 54                         | 8.4  | P=0.112      |
| Unani Formulation (15gms)        | 38                    | 22                         | 42.1| 28                         | 26.3| 33                         | 13.1 | P=0.120      |
| Unani Formulation (30gms)        | 35                    | 11                         | 68.5| 14                         | 60  | 18                         | 48.5 | P=0.056*     |
| Unani Formulation (45gms)        | 54                    | 11                         | 79.6| 16                         | 70.37| 18                         | 66.66 | P=0.001*     |
| Formalin and potassium permanganate | 29                    | 24                         | 17.2| 23                         | 20.6| 9                          | 68.9  | P=0.120      |

Significance: P=0.266, P=0.051*, P=0.055*, P=0.06

*CFU=Colony Forming units. Increase in microbial colony counts observed. % indicates reduction in aerial microbial colonies. *P< 0.05 considered significant.
4. DISCUSSION

The present study compared the effect of Unani Medicines fumigation on reducing the microbial load in ambient air in relation to positive (formalin plus potassium permanganate) and negative (tamarind wood charcoal) controls; in order to validate their air disinfection efficiency and reveal the potential of these plant-products fumigation as an alternative to conventional chemical fumigants.

The results of this study showed that fumigation of the negative control had no effect on the population of aerial bacteria and fungi, whereas the effect of test formulation fumigation was dose-dependent, with its effect increasing as the dose of the test formulation was increased. Test formulation at 45gms dose had the greatest effect on bacterial and fungal colonies. The effect of fumigation with formalin and potassium permanganate was found to be maximum after 12 hours of fumigation; whereas its effect was found to be very little at 1 and 4 hours of fumigation.

The results of other studies are not directly comparable with our study because, to the best of our knowledge, this is the first study that examines the effect of Unani Medicines fumigation in improving the microbiological quality of air. Although, some AYUSH researchers [15-16,20-22] have attempted to validate the use of fumigation with herbal products as an air disinfectant; however, the drugs they have used were different from our test formulation.

Bisht et al. [20] reported that among the various tested herbs, 15gms of Cedrus deodara resulted in around 96% reduction in airborne bacteria after 45 minutes of fumigation [20]. Nautiyal et al. [15] and Samanth et al. [21] demonstrated that 500gms havansamagri (mixture of more than 50 odiferous and medicinal plants) reduced the airborne bacteria by 94% and 95% respectively within 1 hour of fumigation. Bhatwalkar et al. [22] demonstrated the use of 3gms powder of four herbal drugs reduced airborne bacteria by nearly 60-70% after 12 hrs of fumigation. Whereas in the present study, fumigation with the test formulation powder reduced airborne bacteria and fungi colonies by 75% and 81.5 percent respectively, in much larger settings (1200, 900, and 700 cu. ft. vs 594 cu. ft.) than used by Bisht et al., at a much lower dose (45 gmsvs 500gms) than used by Nautiyal et al. and Samanth et al., and in a shorter period of time than Bhatwalkar et al. (1 hr after fumigation vs 12 hours).

From the Unani medicine point of view, the effectiveness of the test formulation in reducing aerial microbial population may be attributed to the cold and dry temperament, as well as the daf‘i-ta‘affun (antiseptic) and mānī‘i-‘ufūnat (disinfectant) properties [23,24]. According to the Unani medicine, hot and moist air, particularly humid air, provides a favourable environment for microbial growth. As a result, measures such as increasing the dryness of the air should be taken to inhibit microbial growth. All the components of the test formulation are dry (Yābis) and some are cold (bārid) in temperament, both of which are contrary to microbial growth [25]. While, according to modern medicine, the presence of antibacterial phytochemicals in these plants may explain their efficacy in air disinfection. These plants are rich in a wide variety of bioactive compounds such as terpenoids, alkaloids, flavonoids, tannins and phenolic compounds, which have been found in-vitro to have antimicrobial properties [26].

Boswellic acids (BAs), which are pentacyclic triterpenoids, are the bioactive phytoconstituents of Boswellia that are considered to be responsible for its antimicrobial properties [27]. Mansumbinone, 3, 4-seco-mansumbinoic acid, β-elemene, and T-cadinol are four antimicrobial terpene compounds isolated from Commiphora; among these, 3, 4-seco-mansumbinoic acid has the most potent antimicrobial activity [28]. The presence of alcoholic compound elemol in Costus has been attributed to its antimicrobial activities [29]. The active antimicrobial compounds of Liquidambar are phenolics and terpenes in nature. The major terpenes identified in its essential oil are terpinen-4-ol, α-terpinol, sabine and γ-terpinene [30]. Phytochemical studies revealed the presence of several phenylpropanoids and sesquiterpenoids including α- and β- santalol in the sandal. The antibacterial activity of α- and β-santalol has been demonstrated in various studies [31]. The presence of alkaloids and saponins may be attributed to the antimicrobial activity of Aquilaria [32].

Formalin as a comparative disinfectant was also used by Bisht et al. [20] in their study and reported that its fumigation completely destroys aerogenic microbes within 15 minutes of fumigation. However, the present study found that fumigation with formalin and potassium
permanganate had the greatest effect on aerial microbes after 12 hours of fumigation. Our findings are contrary to the findings of Bisht et al. [20]; variation in findings could be due to the differences in fumigants dose and fumigation method. While this finding is consistent with formalin fumigation standards, which state that the results of formalin fumigation can be better achieved after 12-24 hours of fumigation.

Three types of colonies, yellow, white, and orange were seen in the air samples of the study area. On sequencing, it was found that the bacterial species representing the yellow, white, and orange colonies are *Neomicrococcus lactis*, *Micrococcus lylae*, and *Kocuria rosea* respectively. This finding is in line of previous studies findings. Kooklen et al. reported that two thirds of the environmental isolates in studies were microcoss [33]. According to Gorny et al. [34], the majority of bacteria found in indoor air are *Micrococcus*, *Staphylococcus*, and *Pseudomonas* [34]. However, only the Micrococcus species of bacteria were isolated in this study. The passive air sampling method and the study settings may be the major factors that can be attributed to this finding. Because only 5% of the total bacterial species present in the air can be cultured using the passive sampling method [35]. In the present study, air samples were collected from areas with very low human occupancy, and the literature indicates that human occupancy is the most important factor influencing the total number and community structure of bioaerosols in the indoor environment [36].

On analyzing the effect of fumigation on these bacterial colonies, it was found that fumigation with test formulation at 45 gms dose had the highest effect. The bacterial species isolated from the air sample belong to the micrococcus genus. These organisms are generally of low virulence and consired to be harmless commensals of skin and oropharynx but may cause opportunistic infections in immunocompromised individuals [37]. Therefore, it is assumed that researchers have rarely used these organisms in antibacterial studies. However, the effects of test formulation drugs on other gram-positive airborne pathogens have been extensively researched. Hence, on that basis, it can be said that the test formulation have acted on these bacteriae in the same way as it does on other gram-positive bacterial population [38,39].

The data of the current study clearly indicated that fumigation of Unani medicinal herbs is more effective than conventional chemical fumigants in improving the microbiological quality of air by reducing the total colony count of microbes (CFU). The findings of this study not only validate the claim of Unani scholars about the efficiency of the test formulation in air purification but also add an Unani formulation to the list of potential herbal air disinfectants that can be used for air disinfection after further researches.

5. CONCLUSION

It can be concluded from the findings of the present study that fumigation with Unani medicinal herbs (Káfür, Kundur, Mlásá‘la, Murr, Qusţ, Şandal and ‘Ud) is effective in reducing the microbial load of air. The effect of fumigation with these medicinal herbs powder was found to be dose-dependent with the maximum effect occurring at a dose of 45 gms. However the effect of certain factors such as temperature and humidity on disinfection efficiency was not considered. Hence, the authors recommended that this ancient concept should be further evaluated in the light of modern medical science and can be utilized for air disinfection if found suitable.

CONSENT

It is not applicable.

ETHICAL APPROVAL

It is not applicable.

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

1. Martin WJ 2nd, Glass RI, Balbus JM, Collins FS. A major environmental cause of death. Science. 2011;334(6053): 180-81.
2. Gall ET, Carter EM, Earnest CM, Stephens B. Indoor air pollution in developing countries: research and implementation needs for improvements in global public health. Am J Public Health 2013; 103(4): e67-e72. DOI:10.2105/AJPH.2012.300955

3. WHO Guidelines for Indoor Air Quality: Selected Pollutants. Geneva: World Health Organization; 2010. Available from: https://www.ncbi.nlm.nih.gov/books/NBK138700/

4. Cincinelli A, Martellini T. Indoor Air Quality and Health. Int J Environ Res Public Health 2017; 14(11):1286. DOI:10.3390/ijerph14111286

5. Kim KH, Kabir E, Jahan SA. Airborne bioaerosols and their impact on human health. J Environ Sci (China). 2018:67:23-35. DOI: 10.1016/j.jes.2017.08.027

6. Stetzenbach LD. Airborne infectious microorganisms. Encyclopedia of Microbiology3rd Edition.Amsterdam: Academic Press. 2009:175-182. Available:https://doi.org/10.1016/B978-012373944-5.00177-2.

7. Bolashikov ZD, Melikov AK. Methods for air cleaning and protection of building occupants from airborne pathogens. Build Environ. 2009;44(7):1378-1385.

8. Puck TT. The mechanism of aerial disinfection by glycols and other chemical agents: I. Demonstration that the germicidal action occurs through the agency of the vapor phase. J Exp Med. 1947;85(6):729-739. DOI: 10.1084/jem.85.6.729.

9. Sina I. Al-QānūnFiTibb. Vol. 1. & V. New Delhi: IdārāKitābuahShifā; 2010:102-110, 1205-08.

10. Baghdādī HI. Kitāb al-MukhtarätFiTibb. Part-4. New Delhi: Central Council for Research in Unani Medicine. 2007:237-38.

11. Rāżī Z. Kitābāl-Mansūrī. New Delhi: Central Council for Research in Unani Medicine. 1991:175-79.

12. Rāżī Z. Kitāb al-Murshid. New Delhi: Tāraqqī Urdu Beoro. 2000:34-37.

13. Rāżī Z. Kitāb al-Hawi. Vol. 15. New Delhi: Central council for Research in Unani Medicine. 2008:151-55.

14. Rushd I. Kitāb-al-Kulliyat. Lahore: Maktaba Daniyal. 2017;368-370.

15. Samantha TU, Jha SG, Sinha V, Patel S, Desai KJ. Effect of smoke from medicinal herbs on the nosocomial infections in ENT outpatient department. Indian J Otol. 2018; 24 (1):9-12.

16. Kumari N, Shashirekha. Antimicrobial action of dhupana (fumigation with herbs) with respect to air borne microbes in indoor environment of central hospital. Int. J. Res. Ayurveda Pharm. 2016;7(5):48-55.

17. Anthrax in Humans and Animals. 4th edition. Geneva: World Health Organization; 2008. Annex 3, Disinfection, decontamination, fumigation, incineration. Available:https://www.ncbi.nlm.nih.gov/books/NBK310477/

18. Zhang LP. The ratio of a formalin and potassium permanganate preparation in a sealed container and its bacteriocidal effect. Zhonghua Hu Li ZaZhi. 1985 Feb;20(1):29-30.

19. Scarlett CM, Mathewson GK. Terminal disinfection of calf houses by formaldehyde fumigation. Vet Rec. 1977; 101(1):7-10. DOI: 10.1136/vr.101.1.7.

20. Bisht LS, Brindavanam NB, Kimothic P. Comparative study of herbal agents used for fumigation in relations to formulation. Anc Sci Life. 1988;8(2):125-32.

21. Nautiyal CS, Chauhan PS, Nene YL. Medicinal smoke reduces airborne bacteria. J Ethnopharmacol. 2007;114(3): 446-51. DOI: 10.1016/j.jep.2007.08.038.

22. Bhatwal NK, Shukla P, Srivastava RK, Mondal R, Anupam R. Validation of environmental disinfection efficiency of traditional Ayurvedic fumigation practices. J Ayurveda Integr Med. 2019;10(3):203-206. DOI: 10.1016/j.jaim.2019.05.002

23. Khan MA,Muht-i-Azam. Vol. 3 & 4. New Delhi: Central council for Research in Unani Medicine. 2013:27, 303, 463, 558, 621, 711, 820.

24. Ghanī N. Khazāin’alAdviya. New Delhi: IdārāKitābuahShifā: YNM: 932, 955, 981, 999, 1069,1229.

25. Itrat M, Khan TN, Riaz Z, Zulkifle M. Epidemic containment measures in Unani medicine and their contemporary relevance. J Indian Sys Medicine. 2020; 8(2):84-90.

26. Elisha IL, Botha FS, McGaw LJ, Eloff JN. The antibacterial activity of extracts of nine plant species with good activity against Escherichia coli against five other bacteria
and cytotoxicity of extracts. BMC Complement Altern Med. 2017;17(1):133. DOI: 10.1186/s12906-017-1645-z.

27. Raja AF, Ali F, Khan IA, Shawl AS, Arora DS, Shah BA, Taneja SC. Antistaphylococcal and biofilm inhibitory activities of acetyl-11-keto-boswellic acid from Boswelliaserrata. Bio Med Microbiol. 2011;11:54. DOI: 10.1186/1471-2180-11-54.

28. Rahman MM, Garvey M, Piddock LJ, Gibbons S. Antibacterial terpenes from the oleo-resin of Commiphoramolmol (Engl.). Phytother Res. 2008;22(10):1356-60.

29. Pandey MM, Rastogi S, Rawat AK. Saussureacostus: botanical, chemical and pharmacological review of an ayurvedic medicinal plant. J Ethnopharmacol. 2007;110(3):379-90.

30. Okmen G, Turkcan O, Ceylan O, Gork G. The antimicrobial activity of Liquidambar orientalis mill. against food pathogens and antioxidant capacity of leaf extracts. Afr J Tradit Complement Altern Med. 2014;11(5):28-32.

31. Kumar R, Anjum N, Tripathi YC. Phytochemistry and pharmacology of santalum album: a review. World J Pharm Res 2015; 4(10):1842-1876.

32. Dash M, Patra JK, Panda PP. Phytochemical and antimicrobial screening of extracts of Aquilariaagallocha Roxb. Afr J Biotechnol 2008; 7(20):3531-3534.

33. Kookan JM, Fox KF, Fox A. Characterization of micrococcus strains isolated from indoor air. Mol Cell Probes 2012; 26(1):1-5. DOI: 10.1016/j.mcp.2011.09.003

34. Gorný R, Dutkiewicz J, Krysińska-Traczyk E. Size distribution of bacterial and fungal bioaerosols in indoor air. Ann Agric Environ Med 1999;6(2):105-113.

35. Napoli C, Marcotrigiano V, Montagna MT. Air sampling procedures to evaluate microbial contamination: a comparison between active and passive methods in operating theatres. BMC Public Health. 2012; 12:594. DOI: 10.1186/1471-2458-12-594

36. Nazaroff WW. Indoor bioaerosol dynamics. Indoor Air. 2016;26(1):61-78. DOI:10.1111/ina.12174

37. Baron S, editor. Medical microbiology. 4th edition. Galveston (TX): University of Texas Medical Branch at Galveston; 1996. Available:https://www.ncbi.nlm.nih.gov/books/NBK8147/

38. Agnihotri S, Tamrakar K. Antibacterial activity of aqueous methanolic extract of leaf and fruit extract of Santalum album. Int J Sci Res. 2017;7(8):524-25.

39. Canli K, Yetgin A, Akata I, Altuner EM. In vitro antimicrobial screening of aquilariaagallocha roots. Afr J Tradit Complement Altern Med. 2016;13(5):178-81.

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