**Anti-Inflammatory And Antimicrobial Activities Of Powered And Ethanaol Extracts Of Scent Leaf (*Ocimumgratissimum*) And Curry Leaf (*Murrayakoenigii*)**

Onyeyilim Ebuka Leonard, Urama Festus Chinonso

Department of pure and Industrial Chemistry, University of Nigeria, Nsukka (41001)

ebuka.onyeyilim@unn.edu.ng

**Abstract:** The antimicrobial activities of ethanol extract and anti-inflammatory activities of powdered extract of *Ocimum gratissimum L* (scent leaves) and *Murraya koenigii* (curry leaves) were investigated. The tested organisms were *Staphylococcus aureus, Bacillus subtilis, Salmonella typhi and Candid albicans, Aspergillus niger* for bacteria and fungi respectively in the antimicrobial test. The antimicrobial test was determined using agar-well dilution method. The scent leaf extracts was active against all the microbes except *Staphylococcus aureus* and *Salmonella typhi*. The powdered extract of scent leaf and curry leaf showed immense anti-inflammmatory activities, although the extract of curry leaf showed higher anti-inflammatory activities close to that of indomethacin which acted as the standard drugs. The minimum inhibitory concentration (MIC) result showed that the ethanolic extracts of scent leaf has more antibacterial activities than that of curry leaf, which showed more antifungal activities.

[Onyeyilim Ebuka Leonard, Urama Festus Chinonso. **Anti-Inflammatory And Antimicrobial Activities Of Powered And Ethanaol Extracts Of Scent Leaf (*Ocimumgratissimum*) And Curry Leaf (*Murrayakoenigii*).** *N Y Sci J* 2020;13(3):58-63]. ISSN 1554-0200 (print); ISSN 2375-723X (online). <http://www.sciencepub.net/newyork>. 5. doi:[10.7537/marsnys130320.05](http://www.dx.doi.org/10.7537/marsnys130320.05).

**Key words**: *Ocimum gratissimum, Murraya koenigii,* antimicrobial, anti-inflammatory, MIC

**Introduction**

Infectious diseases are nowadays the second major cause of death worldwide and the third leading cause of death in developed countries (Khan, *et al.,* 2008). Also, microorganism resistance to multiple antimicrobial agents has become a serious problem (Reambiah, *et al.,* 2014 and Varshney *et al.,* 2009). Scent leaf (*Ocimumgratissimum*) is a well-known medicinal plant which has received a great deal of attention over the past few decades around the world (Disegha, *et al*., 2014).

Curry leaves (*Murrayakoenigii*) also are a popular leaf-spice used in very small quantities for their distinct aroma due to the presence of volatile oil and their ability to improve digestion. These leaves are widely used in Asian and African cuisines for flavouring foods. The leaves have a slightly pungent, bitter and feebly acidic taste, and they retain their flavour and other qualities even after drying (Suman, *et al.,* 2014). The Seeds and leaves yield an essential oil containing eugenol, it is used widely in perfumes, cosmetics, pharmaceuticals and confectionary industries (Bhandari, 2012). Extracts of Ocimumgratissimum having strong antibacterial and antioxidant properties are widely used for medicinal purposes. The administration of aqueous extract of both the curry and olive leaves has shown a significant decrease in the blood glucose level in STZ-induced diabetic rats (Maha, *et al.*, 2013). Extracts from *Ocimumgratissimum.* Plants have shown strong inhibitory effects on HIV-1reverse transcriptase and platelets aggregation (Das, *et al*., 2012; Halliwell and Gutteridge 1990).

This work therefore, is intended to extract the chemical constituent of the leaves of *Ocimumgratissimum* and *Murrayakoenigii,* using ethanol as a solvent for a portion and obtaining the solid extract of another portion with the view to authenticating the antimicrobial potentials and anti-inflammatory activity of these extracts against some harmful pathogenic bacteria and inflammation.

**Materials And Methods**

**Preparation of Ethanol Extracts for Antimicrobial test**

The antimicrobial test was carried out by the method adopted by Harika, *et al.,* 2010 and Mohar, *et al.,* 2011. The fresh leaves of *Murrayakoenigii* and *Ocimumgratissimum* were washed with tap water, dried in the lab under ambient temperature and powdered by a mechanical grinder. The dried powders were then extracted with ethanol to give their extracts respectively. The plant leaves about 10gms each was sequentially extracted with ethanol (150ml) by maceration process for 5 days at room temperature. The obtained extracts were filtered by using whatmann

no.1 filter paper. The extracts were concentrated using a rotary evaporator at 60°Cevaporated to obtain syrupy solution weighing 19.8g and19.2g for curry leaves and Scent leaves respectively. This was put in an air-tight container and stored in a refrigerator for further analysis.

**Preparation of Solid Extracts for Anti-inflammatory Test**

This test was carried out using the method reported by Winter, *et al.,* 1962 The fresh leaves of *Murrayakoenigii* and *Ocimumgratissimum* L. were washed with tap water, dried in the lab under ambient temperature and powdered by a mechanical grinder. The powdered plant materials was filtered to obtain the extract, which was stored in an air-tight container and stored for further analysis.

**Evaluation of Antimicrobial Activity**

The antimicrobial sensitivity testing of both crude and prepared concentrations of the plant extracts was determined using the agar-well dilution method.

**Test Micro-organisms**

Six standard clinical isolates used in this work were obtained from laboratory stock organisms in the Department of Pharmaceutical Microbiology, University of Nigeria, Nsukka. They include four bacteria, -*Staphylococcus aureus, Escherichia coli, Bacillus subtilis* and *Salmonella typhi* and two Fungi, - *Candida albicans, and Aspergillusniger.* These micro-organism are selected because they have been implicated to be among the organisms causing enteric diseases/infections and easily develop resistance to commonly prescribed antibiotics.

**Preparation and sterilization of materials**

All glass wares used were first soaked in detergent solution for about 30 minutes, then washed and rinsed with clan water and allowed to dry. The test tubes were plugged with cotton wool, while the pipettes and petri dishes were packed in a canister. All these were sterilized using hot air oven at 160°C for one hour.

**Preparation of Culture Media**

**Nutrient agar**: A 28g of nutrient agar powder was dissolved in 1000ml of distilled water, and was allowed soaking for 10mins. The agar suspension was brought to melt by boiling in a water bath. A 20 ml aliquot of the molten nutrient agar was dispensed into a bijou bottles, cocked, and sterilized in an autoclave at 121oc for 15mins. The sterile molten nutrient agar was stored at 42oc until use.

**Potato Dextrose Agar (PDA)**: A 47g of PDA powder was dissolved in 1000ml of distilled water, and was allowed soaking for 10mins. The agar suspension was brought to melt by boiling in a water bath. A 20ml aliquot of the molten PDA was dispensed into a bijou bottles, cocked, and sterilized in an autoclave at 121oC for 15mins. The sterile molten potato dextrose agar was stored at 42oC until use.

**Preparation of Test Micro-organism**

Four standard laboratory isolates; *S. aureus, E. coli, B. subtilis and S. typhi* and two fungi; A. *niger* and *C. albicans* (standard laboratory isolates), were all obtained from Pharmaceutical Microbiology Laboratory, UNN.

The pure samples of each test micro-organism were transferred into a sterile petri dish with the aid of sterile inoculating loop, and incubated at 37oC for 24 hours for bacteria and at 25oC for 48 hours for fungi.

**Standardization of the test organism suspension**:

The micro-organisms were standardized using 0.5 MacFaland turbid equivalent.

**Preparation of Sterile Water:**

A 200ml of distilled water was dispensed in conical flask and the flask was plugged with cotton and foil. The water was sterilized in an autoclave at 121oC for 15 minutes.

**Preparation of the different concentration of the extract used:**

A 5mg/ml (5000µg/ml) stock concentration of the crude extract was prepared by dissolving 10g of the extract in 2ml of 50% DMSO. 1.0mg/ml, 0.9mg/ml, 0.8mg/ml, 0.7mg/ml, 0.6mg/ml, 0.5mg/ml, 0.4mg/ml, 0.3mg/ml, 0.2mg/ml, 0.1mg/ml, concentrations were obtained using C1V1=C2V2 formula.

**Control test (standard)**:

The standard antibiotics used were Ciprofloxacin and Fluconazole.

**Experimental:**

A 4.0ml of crude extract suspension of stock concentration 50mg/ml was transferred to the sterile Petri dish, a 16.0ml volume of double strength sterile molten agar was transferred to the same plate to mix uniformly thus, 1mg/ml concentration was obtained. The other concentrations 0.9g/ml, 0.8mg/ml, 0.7mg/ml, 0.6mg/ml, 0.5mg/ml, 0.4mg/ml, 0.3mg/ml, 0.2mg/ml, 0.1mg/ml, were obtained using the same C1V1=C2V2 formula. The molten agar plates with different concentrations of the extract were allowed to gel. The plates were divided into seven equal parts with permanent marker. The test microorganisms were streaked on the segments, and labeled. The culture plates were incubated in inverted position at 37oc for 24hours, and at 25oc for 48hours. After the due period of incubation, the plates were observed for sensitivity and resistivity of the organisms to the agents, and the observation was recorded. The plates were further incubated for another 24hour at 37oc, and 48hours at 25oc to determine whether the activity was bactriostatic or bacteriocidal. The observation was also recorded.

**3.5 Evaluation of Anti-inflammatory Activity**

Albino rats were divided into four groups of six animals each. Oedema was produced by injecting 0.2 ml of a solution of 1% carrageenan in the hind paw. The rats were injected intraperitoneally with 1ml suspension. Group I served as toxicant control, received carrageenan (0.1 ml of 1% solution), group II served as standard, received indomethacin (10 mg/kg, p.o), group III and IV were treated with 100 mg / kg & 100 mg/kg of solid extract of scent leaf and curry leaf respectively by oral route. Note that after the carrageenan was injected into the rats, they were left for about 48 hours to allow the carrageenan to circulate proportionally in the rats causing the inflammation to spread rapidly.

One hour after drug or test compound (extract) administration, 0.1 ml. of 1% carrageenan in distilled water was injected into the sub plantar region of right hind paws of all groups. The paw oedema volume was measured with the help of plethysmometer at zero hour (immediately after injecting carrageenan). The same procedure was repeated at 1, 2, 3 and 4 hour after carrageenan injection. The difference between the initial and subsequent reading gave the actual ooedema volume. Reduction in paw volume compared to the control animals was considered as anti-inflammatory response (Winter, *et al.,* 1962)

Percentage Inhibition = [Vc - Vt/Vc] x 100

Where, Vt = mean paw volume of test group & Vc = mean paw volume of control group.

**Control Test (Standard):**

The standard anti-inflammatory drug used in this study is Indomethacin.

**Results And Discussions**

**Results of the Antimicrobial Activities of the Ethanol extract of Scent leaf and Curry leaf**

The results obtained from the antimicrobial activities of Scent Leaf (*Ocimumgratissimum L*.) and Curry Leaf (*Murrayakoenigii*) are as presented below;

**Results of the Antimicrobial activities of Scent leaf (*Ocimumgratissimum L*.)**

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Concentration (mg/ml)** | **Staphylococcusaureus** | **Bacillus subtilis** | **Escherichia coli** | **Salmonelatyphi** | **Candida albicans** | **Aspergillusniger** |
| 1.0 | - | - | - | - | - | - |
| 0.9 | + | - | + | - | + | + |
| 0.8 | + | + | + | + | + | + |
| 0.7 | + | + | + | + | + | + |
| 0.6 | + | + | + | + | + | + |
| 0.5 | + | + | + | + | + | + |
| 0.4 | + | + | + | + | + | + |
| 0.3 | + | + | + | + | + | + |
| 0.2 | + | + | + | + | + | + |
| 0.1 | + | + | + | + | + | + |
| Ciprofloxacin 30µg/ml | - | - | - | - | + | + |
| Fluconazole 30µg/ml | + | + | + | + | - | - |

**Key:** The sign **(-)** indicates that there is inhibition, The sign **(+)** indicates that there is no inhibition

**Results of the Antimicrobial activities of Curry leaf (*Murrayakoenigii*).**

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Concentration (mg/ml)** | **Staphylococcusaureus** | **Bacillussubtilis** | **Escherichia coli** | **Salmonelatyphi** | **Candida albicans** | **Aspergillusniger** |
| 1.0 | + | - | - | + | - | - |
| 0.9 | + | - | + | + | - | + |
| 0.8 | + | +  | + | + | - | + |
| 0.7 | + | + | + | + | + | + |
| 0.6 | + | + | + | + | + | + |
| 0.5 | + | + | + | + | + | + |
| 0.4 | + | + | + | + | + | + |
| 0.3 | + | + | + | + | + | + |
| 0.2 | + | + | + | + | + | + |
| 0.1 | + | + | + | + | + | + |
| Ciprofloxacin 30µg/ml | - | - | - | - | + | + |
| Fluconazole 30µg/ml | + | + | + | + | - | - |

**Key;** The sign **(-)** indicates that there is inhibition, The sign **(+)** indicates that there is no inhibition

**Determination of Minimum Inhibitory Concentration (MIC)**

The minimum inhibitory concentration (MIC) is defined as the lowest concentration of the extract inhibiting the visible growth of any micro-organism permanently. It is best determined by incorporating various concentrations of the extracts into the culture media, observing and measuring the zone of inhibition.

The minimum inhibitory concentration of the two leaf extracts on the microorganisms is given in the tables below:

**Minimum Inhibitory Concentration of Ethanolic extract of Scent leaf on the test micro-organisms.**

|  |  |
| --- | --- |
| **Test Micro-organism** | **Minimum Inhibitory Concentration (MIC) (mg/ml)** |
| Staphylococcus aureus | 1.0 |
| Bacillus subtilis | 0.9 |
| Escherichia coli | 1.0 |
| Salmonella typhi | 0.9 |
| Candida albicans | 1.0 |
| Aspergillusniger | 1.0 |

**Minimum Inhibitory Concentration of Ethanolic extract of Curry leaf on the test micro-organisms.**

|  |  |
| --- | --- |
| **Test Micro-organism** | **Minimum Inhibitory Concentration (MIC)****(mg/ml)** |
| Staphylococcus aureus | N/A |
| Bacillus subtilis | 0.9 |
| Escherichia coli | 1.0 |
| Salmonella typhi | N/A |
| Candida albicans | 0.8 |
| Aspergillusniger | 1.0  |

**Key:** N/A in this context means no activity.

**Results of the Anti-inflammatory Activities of Solid extracts of Scent leaf and Curry leaf**

|  |
| --- |
| **Percentage Inhibition Of Oedema Formation** |
| Groups | 1h  | 2h | 3h | 4h |
| I | 1.27  | 1.36  | 1.49  | 1.53  |
| II | 19.4  | 34.1  | 45.3  | 55.7  |
| III | 12.2  | 26.8  | 36.8  | 43.7  |
| IV | 17.8  | 29.7  | 39.7  | 51.2  |

**Note;** h: in the table means hour

* Group I is the toxicant control, they received carrageenan but were not treated.
* Group II is the standard control, they were treated with a standard drug, indomethacin
* Group III was treated with 100mg/kg of solid extract of Scent leaf
* Group IV was treated with 100mg/kg of solid extract of Curry leaf

**Discussion of results**

Currently, in addition to antibiotics and chemically synthesized drugs, the trend to look out for alternative medicine in nature is increasing as the natural resources are less toxic and less deleterious to the overall health of human beings. In this study curry leaves and scent leaves have displayed immense potential as that natural alternative, especially with their antimicrobial anti-inflammatory activities. Table 4.1 and 4.2 gives the antimicrobial activities of *Ocimumgratissimum L*. and *Murrayakoenigii* leaf extracts against different micro-organisms at different concentrations.

The six micro-organism used in this research were sensitive to the leaf extract of scent leaf but it showed higher activity against the bacterial strains; *Bacillussubtilis* and *Salmonelatyphi* at a concentration of 0.9mg/ml (typhoid causing agent) where it showed activity even at a lower concentration. These activities was found to be more profound than the antibiotics used in my study, alluding to the excellent medicinal properties of scent leaves without any side effects that antibiotics may produce in humans, so they can be employed for treating typhoid and other bacterial infections. There was no significant effect on other bacterial and fungi. The antimicrobial effect on the later bacterial and fungi species could have been limited by contamination in the laboratory procedures. The observation suggests that the ethanolic extracts of Scent leaf has a higher antibacterial activity than antifungal activity as can be clearly seen from table 4.3. which shows the minimum inhibitory concentration (MIC) of Scent leaf.

On the other hand, not all of the six micro-organisms showed sensitivity to the ethanolic leaf extracts of curry leaves. It had more activity on the bacterial strain; *Bacillus subtilis* at a concentration of 0.9mg/ml and a more profound activity on the fungal specie; *Candida albicans* at a concentration of 0.8mg/ml, it showed no activity on two bacterial strains; *Staphylococcus aureus and Salmonelatyphi* as can be seen above. These limitations in activities could have been due to contamination in the laboratory procedures. It showed sensitivity on the two fungal strains used, this observation suggests that the ethanolic extracts of curry leaf has a higher antifungal activity than antibacterial activities as can be seen clearly from table 4.4 above which shows the minimum inhibitory concentration (MIC) of Curry leaf. This indicates that it can serve as a good alternative for treating any form of fungi infections with little or no side effects as compared to the usual antibiotics.

Generally, from this study it can be seen that the ethanolic leaf extracts of Scent leaf and Curry leaf have very good antimicrobial activities against microbes and can be employed in place of antibiotics for the treatment of drug resistant pathogenic strains that do not respond to the usual line of treatment. It can also be deduced from this study that the ethanolic leaf extracts of scent leaves showed a higher antibacterial activity than that of curry leaves which showed a higher antifungal activity when compared.

The standard antibiotics used, Ciprofloxacin was effective against all the bacterial strains at standard concentration (30µg/ml), while Fluconazole was effective against all the fungal species at a standard concentration (30µg/ml).

For the anti-inflammatory activities of curry leaf and scent leaf, in this work it was observed that after the drugs were administered to the groups, they had different but similar activities. After the measurement of the paw oedema volume using the plethysmometer in 1h, 2h, 3h and 4h intervals and was used to calculate the percentage inhibition, it was observed that the solid extract of scent leaf and curry leaf showed immense activities on the paw oedema. Curry leaf (Group IV) showed a higher activity on the inflammation as its activity was closer to that of the standard drug indomethacin at the 1h, 2h, 3h and 4h intervals there was no much difference compared to the activity of Scent leaf (Group III) at this intervals, this is well illustrated in table 4.5.

**Conclusion**

*Murrayakoenigii* (Curry leaves) extracts have demonstrated antibacterial effects particularly on *Bacillus subtlis* and *Escherichia coli*, it has also shown antifungal effects particularly on *Candidaalbicans* and mildly on *Aspergillusniger* as compared to the antibiotics such as ciprofloxacin and fluconazole used in this study. The ethanolic extracts of curry leaves were found to be effective against all tested microorganism except *Staphylococcus aureus* and *Salmonelatyphi*.

For *Ocimumgratissimum L*. (Scent leaves), the results of these particular study suggests that its extracts demonstrated antibacterial effects strongly on *Bacillus subtilis* and *Salmonelatyphi*, and mildly on *Staphylococcus aureus* and *Escherichia coli*, it also showed moderate antifungal effects on *Candida albicans* and *Aspergillusniger* as compared to the antibiotics used in this study. The ethanolic extracts of Scent leaves were found to be effective against all tested microorganism.

The solid extracts of Scent leaf and Curry leaf showed high anti-inflammatory activities especially that of curry leaf (Group IV) which showed an activity that is almost equal to that of the standard drug used in this study (indomethacin). Therefore the solid extracts of scent leaf and curry leaf can be comfortably used as an alternative drug to address cases of inflammation, due to their less toxicity, less side effects and high activities against any form of inflammation.

*Murrayakoenigii* and *Ocimumgratissimum L*. has the bioactive potential and they may be the best natural alternative to antibiotic and anti-inflammatory therapy for tested microbes and paw oedema. Therefore, curry leaves and Scent leaves could be effectively used as a natural remedy in everyday meal, for the prevention of bacterial, fungal and inflammatory infections. Indeed these phenomenal plants may serve as a useful resource in the food industry and clinical medicine. This study gives way for further attention and research to identify the active compounds responsible for these plants biological activities.

**References**

1. Bhandari, P.R. (2012). Curry leaf (*Murrayakoenigii*) or Cure leaf: Review of its curative properties. *J. Med. Nutr. Nutraceut*, 1: 92-97.
2. Das, B.N and Biswas, B.K. (2012). Antibacterial and cytotoxic activities of the leaf extract of *Murraya Koenigii*. *Int. J. Life Sc. Bt & Pharm. Res.,* 1(3): 59-63.
3. Disegha, G.C, and Onuegbu Izionworu V. (2014). Antifungal Activities of Curry Leaf (Murraya Koengii) Extract on Some Selected Fungi. *Chemistry and Materials Research,* 6(11): 2224-3224.
4. Halliwell, B, Gutteridge J.M.C. (1990). Role of free radicals and catalytic metal ions in human disease, *An overview. Methods Enzyme molecule*, 186:1-85.
5. Harika, V.C., Padmavathi, P., Sambasiva Rao K.R.S., Phani, R.S.C. (2010). In vitro antimicrobial activity of leaf powder. *Int. J. Res. Pharm. Biomed. Sci.,* 2(1): 128-131.
6. Khan S. A, Lamba H. S, Alam O, Mohd I. (2008) Synthesis and antihyperglycemic activity of [2-(substituted phenyl)-3-[{4-(1-naphthyl)- 1,3-thiazol-2-yl}amino]-4- oxo-1,3-thiazolidin-5-yl] Acetic Acid. *Asian J Chem* 20: 4987-4993.
7. Maha El Amin, Promy Virk, Mai Abdel Rahman El Obeid, Zainab M Al Marhoon, Zainab Korany Hassan, et al. (2013). Antidiabetic effect of Murrayakoenigii and Oleaeuropaea leaf extracts on streptozotocin induced diabetic rats. *Pak. J. Pharm. Sci.,* 2(26): 359-365.
8. Mohar, S. A., Sanjay, K., Hotam, Singh. C., Ravindra M Thakka Santosh K Verma, Chandrabhan Seniya. (2011).The efficacy of Murrayakoenigii leaf extract on some bacterial and a fungal strain by disc diffusion method*. J. Chem. Pharm. Res*., 3(5): 697-704.
9. Reambiah, L.K., Ndong, J.C., Massoua, P.M.M., Mezegue, S., Elisee-Ndam, M., Mintsa Ndong. A., Siawaya. J.F.D. (2014). *Int. J. infect. Dis*. 29, 48-53.
10. Suman Singh A1, P.K. Omre B and Sandhya Madan Mohan C. (2014). Curry leave (Murrayakoenigii) a miracle plant. *Ind. J. Sci. Res.,* 4 (1): 46-52.
11. Varshney, V., Mishra, N.N., Shukla. P.K., Sahu. D.P. (2009). *Bioorg Med. Chem. Lett.*
12. Winter, C., Risley, E.A., Nuss, G.W. (1962). Carragennian induced ooedema in hind paw of the rats for anti-inflammatory drugs. *Proceedings of the Society for Experimental Biology and Medicine.*; 3:544-547.

3/7/2020