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Phytochemical Screening and Antimicrobial Activity of the Aerial Part of Three Selected Plants

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ABSTRACT

We investigate the antimicrobial activity and phytochemical screening of the aerial parts of Annona senegalensis, *Petiveria alliacea* and Secamone afzelii that have been used as folk medicines. The methanolic extracts were tested on various microorganisms for the antibacterial and antifungal activity using agar well diffusion and poisoned food technique respectively. The length of the inhibition zone was measured in millimetres from the edge of the well to the edge of the inhibition zone. The extracts were assessed in an effort to validate the potential activity of the plants against microbes. The result showed the extracts possess considerable antimicrobial potentials. The phytochemical screening of the plants revealed the presence of alkaloids, flavonoids, tannins, steroids, terpenoids, saponins and anthraquiones. The phytochemicals attributed to the antimicrobials activity of the plants extracts

Keywords: antibacterial, antifungal, Annona senegalensis, Petiveria alliacea, Secamone afzelii

1. Introduction

Plants have been one of the important sources of medicines since the beginning of human civilization. There is a growing demand for plant based medicines, health products, pharmaceuticals, food supplements and cosmetics. Since ancient times, people have been exploring the nature particularly plants in search of new drugs which has resulted in the use of large number of medicinal plants with curative properties to treat various diseases (Verpoorte, 1998). According to WHO survey, 80% populations living in the developing countries rely exclusively on traditional medicine for their primary health care needs of which most involve the use of plant extracts (Sandhya et al., 2006). The studies of plants continue principally for the discovery of novel secondary metabolites or phytochemical which is the non-essential nutrients derived from plants exhibiting a number of protective functions for human beings (Neethu and Neethu, 2016). The medicinal value of plants depends on chemical substance present in them which possess distinct physiological action on the human and animal system. The microorganisms have developed resistance against many antibiotics due to the indiscriminate use of antimicrobial drugs (Ahmad et al., 1998). Antibiotics are sometimes associated with side effects (Cunha, 2001) whereas there are some advantages of using antimicrobial compounds of medicinal plants, such as often fewer side effects, better patient tolerance, relatively less expensive, acceptance due to long history of use and being renewable in nature (Vermani and Garg, 2002). The most important classes of these bioactive constituents of plants are alkaloids, flavonoids, tannins, saponins, terpenoids and phenolic compounds (Hill, 1952). Generally referred to as phytochemicals or metabolites (Okwu, 2005; Iwu MM et al., 1999).

Petiveria alliacea is a plant from the family Phytolaccaceae, known by different names in different countries of Central and South America, the Caribbean and Africa. For hundreds of years it has been used for pain relief, and as an anti-influenza, anti-inflammatory, anti-tumor, anti-bacterial, anti-fungal, anti-hyperlipidemia, and anti-diabetic drug (Tropical Plant Database-Anam, 2011). This plant also grows in Indonesia, but it has not been used extensively. It is traditionally used in Indonesia as an analgesic, anti-inflammatory and for treatment of hemoptysis (Mulyani *et al.*, 2012). Widowati, (2007) and Weniger *et al.*, (1986) reported that *P. Alliacea* can reduce the length of therapy with standard drugs in tuberculosis patients. Several volatile compound like; benzyl-2-hydroxyethyl trisulfide (Szczepanski *et al.*, 1972), cis-3,5-diphenyl-1,2,4-trithiolan (trithiolaniacin), benzaldehyde, benzoic acid, elemental sulfur, and trans-stilbene (Adegosan, 1974), dibenzyltrisulfide (Sousa *et al.*, 1990) and benzaldehyde, benzyl alcohol, cis- and trans-stilbenes, benzyl benzoate, dibenzyldisulfide, and dibenzyltrisulfide (Ayedoun *et al.*, 1998) had been isolated from *P. alliacea*. *P. Alliacea* is also known as skunk weed due to its characteristic odour resulting from the presence of sulfurate compounds (De Sousa *et al.*, 1990). It has been reported that *P. Alliacea* exhibited antirheumatic, anticarcinogenic, antiflu, antitussive, analgesic and antiinflammatory (Villar *et al.*, 1997).

Annona senegalensis is a subtropical plant (Okoli et al., 2010) that has been implicated for the treatment of chest pain, coughs, anaemia, urinary tract infection (Burkill, 1985 and Muanze et al., 1994], cancer treatment

[Durodola *et al.*, 1975 and Fatope *et al.*, 1993], diarrhoea, dysentery (Ekpenda *et al.*, 1998 and Kudi and Myint, 1999), anthritis and rheumatism (Dalziel, 1937 and Audu, 1989). The isolation of monotetrahydrofuran and bistetrahydrofuranacetogenins (Sahpaz *et al.*, 1994) and two cytotoxic monotetrahydrofuranacetogenins (Sahpaz *et al.*, 1996) from this plant are also documented.

Secamone afzelii is a creeping woody climber with pinnately compound leaves. It is often seen as a nuisance to other plants or crops because of its domineering spread wherever it grows (Tavs and Doris, 2012). The root is said to be poisonous but is used, more the less, by the Zulu medicine man as a remedy for spinal disease (Watt and Breyer–Brandwijk, 1962). S. afzelii is used by native people as an anti-inflammatory, anti-bacterial and tonic drug. Latex of S. afzelii is traditionally used in various skin diseases (boils, abscesses and eruptions) Bitter sap of stems and leaves of S.afzelii is used as a stomachic and purgative and diuretic drug. Crushed S.afzelii is used for cooking food for gonorrhoea patient (abbiw, 1990). Hervé et al., (2008) reported the presence of Flavonoids, Saponins, Reducing sugars, coumarines, alkaloids, proteins, tannins, steroids and polyterpenes and Quinones in the methanol extract of leave and stem of S. afzelii. The objective of this study was to evaluate the phytochemical screening and antimicrobial activity of the selected plants

2.0 Materials and Method

2.1 Sample Collection and Preparation

Fresh aerial parts of the plants were collected in their natural habitats. Locations where samples were collected include Akure in Ondo State, Ajilete in Ogun State, Iwo in Osun State and Ibadan in Oyo state. The samples were then air dried in an open ventilated room, to ensure they retain their natural feel and major constituents. After proper drying, they were pulverised into powdery form.

2.2 Extraction Process

700g of each of the dried and pulverized plant material were weighed and poured in 5ltr glass flasks. 2.5ltr N-Hexane was poured in each and left for 24hrs. The N-hexane was used to de-fat the plant materials. The solvent was decanted after 24hrs and 2.5ltr Methanol was thereafter poured in each flask and left for 72hrs. The extracts were taken to MCRL laboratory in University of Ibadan where they were concentrated using Rotary evaporator. The yield was 3.36% for *Anonna senegalensis*, 2.64% for *Petiveria alliacea* and 3.06% for *Secamone afzelli*

2.3 Phytochemical Screening

The phytochemical investigation of the methanol extracts of *Anonna senegalensis*, *Petiveria alliacea and* Secamone afzelli was carried out using standard protocol.

2.4 Test Microorganisms

The microorganisms of choice used for antibacterial activity are *Staphylococcus aureus*, *Bacillus subtilis*, *Sylmonela typhimurium*, *Enterococcus spp*, *Klebsiela aerogenes*, *Escherichia coli*, *Pseudomonas aeruginosa*, *Pseudomonas syringine*, *Xantomona saxonopodis*, *Shigella dysantrioue*, while the microorganism used for antifungal activity are Erwinia herbicola, *Fusariumoxyporium*, *Marcophomina phomoides*, *collectotrichum lindimathiumum*, *Ceratocystis paradoxa and Helminthoporuin toxicum*.

2.5 Antimicrobial Activity

The stock culture of these organisms had already been identified and typed. All bacteria were cultured aerobically at $37^{\circ c}$ for 24hrs on peptone water and antimicrobial testing were carried out on the nutrient agar (NA) plates. All fungi were also grown aerobically at $27^{\circ c}$ of Potato Dextrose Agar (PDA) plates Agar well diffusion methods of Murray *et al* (2000) modified by Olurinola (2004) was employed for this study. 20ml of nutrient agar was dispensed into sterile universal bottles. Then inoculated with 0.2ml of bacteria cultures mixed gently and poured into sterile petri-dishes. After setting, a number 3-cup borer (6mm diameter) was properly sterilized by flaming and used to make three to five uniform cups/wells in each petri dish. A drop of molten nutrient agar was used to seal the base of each cup. The cups/wells were filled with 50ul of the extract concentration and allowed to diffuse for 45minutes. The solvent used for reconstituting these extracts were similarly plated. The plates were incubated at $37^{\circ c}$ for 24hrs for bacteria. The zones of inhibition were measured with Digital Vernier Callipers in mm. The experiment was carried out in triplicate.

Antifungal activities of the samples were carried out by poisoned food technique. 5ml each of the samples was mixed with 20ml of PDA separately before pour plated and allowed to solidify at ambient temperature. A 5mm disc cut from the periphera of 7-day old culture of the test fungi was inoculated in the center of the PDA plates (mixture of PDA and sample). A negative and positive control experiment was set up separately containing distil water and reference antifungal fungicides (Koside). Both plates were incubated aseptically at $27^{\circ c}$ for 72 - 96 hrs. Mycelinl growth of the isolates were measured with the aid of Digital Venier Calipers and the mycelinl growth inhibition (in percentage) was calculated and recorded appropriately by the formula :



$$\frac{dc - dt}{dc} \times 100$$

Where dc = diameter of bacterial/fungal colony in negative control sets

dt = average diameter of bacterial/fungal colony in the set up containing experimental plants

The experiment was carried out in triplicate.

3. Result

3.1 Phytochemical screening

The result of phytochemical screening of the methanolic extract of the selected plant were presented in Table1

3.2 Antibacterial Activity

The result of antibacterial activity omethanolic extract of the selected plants were presented in Table 2 3.3 Antifungal Activity

The result of antifungal activity of methanolic extracts of the selected plants were presented in Table 3

4. Discussion

The yield of the extract was as follows 3.36% for *Anonna senegalensis*, 2.64% for *Petiveria alliacea* and 3.06% for *Secamone afzelli*.

The methanol extracts of *Secamone afzelli, Anonna senegalensis and Petiveria alliacea* showed the presence of alkaloids, flavonoids, tannins, steroids, terpenoids, anthraquinones and saponins. Cardiac glycosides was absent in all the plants

Antibacterial and antifungal activities of methanol extract of *Petiveria alliacea, Secamone afzelii* and *Annona senegalensis* were tested against ten bacteria and five fungi. The results of antibacterial and antifungal activities are presented in table 2 and 3. The extracts recorded promising activities against the tested organisms. The highest activity of *Secamone afzelii* was 9.99 ± 2.00 mm against *Bacillus subtilis* and its lowest activity was 3.63 ± 0.41 mm against *Escherichia coli*. 7.62 ± 0.45 mm was the highest diameter of inhibition recorded by *Petiveria alliacea* and 2.20 ± 1.00 mm was its lowest diameter of inhibition.

For the antifungal activities of the extracts, Secamone afzelii recorded highest activity against Fusarium oxyporium with percentage mycelinl growth of inhibition of 69.35±2.61nm. Annona senegalensis recorded highest activity against Marcophomina phomoides with percentage mycelinl growth of inhibition of 61.77±1.53mm, percentage mycelinl growth of inhibition of 70.86±1.20nm against Ceratocystis paradoxa and 70.31±1.37mm percentage mycelinl growth of inhibition against Collectotrichum lindimutianum. Secamone afzelii recorded percentage mycelinl growth of inhibition against Helminthosporium toxicum with the value of $76.97 \pm 0.93\%$. The antibacterial and antifungal activities can be attributed to the presence of phytochemicals such as flavonoids, tannins, alkaloids, steroids, coumarines, cardiac glycosides, quinines and terpenoids in the plants extract (Rocha and Silva, 1969; Jolad et al., 1984; Adzu et al., 2005; Tavs and Doris, 2012). The antimicrobial properties exhibited by the extracts may be associated with the presence of tannins, saponins and alkaloids found in the plant extracts. A large number of flavonoids have been reported to possess antimicrobial properties (Bastista et al., 1994; Tsuchiya et al., 1996; Boris, 1996; Olowusulu and Ibrahim, 2006; Akinjobi et al., 2006). Tsuchiya et al., (1996) attributed the antimicrobial activities of flavonoids to their ability to complex with extracellular and soluble proteins as well as their ability to complex with bacterial cell walls. They suggested that more lipophylic flavonoids exert antimicrobial activity by disrupting microbial cells (Flora and Folasade, 2008). Herbs that have tannins as their component are astringent in nature and are used for the treatment of gastrointestinal disorders such as diarrhoea and dysentery (Dharmananda, 2003; Asguith and Butler, 1986). Saponins believed to be responsible for numerous pharmacological properties (Estrada et al., 2000) and have been shown to have immense significance as anti hypercholesterol, hypotensive and cardiac depressant properties (Price, 1987). Waterman (1992) reported that alkaloids and flavonoids were useful as antimicrobial, anti-inflammatory and anti-oxidant agents.

5. Conclusion

The result obtained from this study has shown that phytochemical screening of the methanolic extract of *Petiveria alliacea, Secamone afzelii* and *Annona senegalensis* reveal the presence of alkaloids, Flavonoids, saponins, tannins, anthraquinones, terpenoids, and Steroids. The extracts also demonstrated significant antimicrobial and antifungal activities against the tested organisms. Therefore more detailed studies are needed to isolate, characterized and evaluate the active components and the mechanism of action.

REFERENCES

- Adesogan, E.K. (1974), Trithiolaniacin, a novel trithiolan from *Petiveria alliacea*. Journal of the Chemistry Society 21, 906–907
- Ahmad, I. Mahmood, Z & Mohammad, F. (1998), Screening of some Indian medicinal plants for their antimicrobial propreties. *Journal of Ethnopharmacological* 62, 183-193.
- Akujobi C.O., Ogbulie J.N. &Uchegbu U.N., (2006), Antibacterial activities and preliminary phytochemical screening of Vernonia amygdalina and Citrus aurantifolia. *Nigeria Journal of Microbiology* 20 (1), 649-654.
- Asquith T.N. & Butler L.G., (1986), Interaction of condensed tannins with selected proteins. *Phytochemistry*25 (7), 1591-1593.
- Audu, J. (1989), Medicinal herbs and their uses in Bauchi State. The Nigerian Field 54, 157-168.
- Bastista, O., Duarte O., Nascimento, S. &Simones, M.F. (1994),Structure and antimicrobial activity of diferpenes from the root of *Plectranthus hereoensis. Journal of Natural Product* 57, 279-237
- Boris R.P., (1996), Natural Products Research; Perspectives from a major pharmaceutical company. *Journal of Ethnopharamatology* 51, 29-38
- Cunha, B.A. (2001), Antibiotic side effects. Medical Clinics North America 85, 149-185.
- Dalziel, J. M. (1937), The useful plants of West Tropical Africa. Crown overseas agents for the colonies, London, 2-3.
- De Sousa, J.R. Demuer A.J. & Pinheiro J.A. (1990), Dibenzyl trisulphide and trans-N-methyl-α-methoxyproline from *Petiveriaalliacea*, *Phytochemistry* 29, 3653–3655
- Dharmananda S., (2003),Golinuts and the uses of tannins in Chinese Medicine In proceedings of Institute for Traditional Medicine Portland, Oregon, USA
- Durodola, J. I. (1975), Viability and transplanability of developed tumour cells treated in vitro with anti-tumour agent C/M2 isolated from herbal cancer remedy of *Annonasenegalensis*. *PlantaMedica*28, 359
- Ekpendu, T. O. E., Obande, O. D. Anyogo P. O. & Attah, A. D. (1998), Nigerian ethno medicine and medicinal plant flora the Benue experience part 1". International *Journal of Pharmaceutical Research and Development3*, 37–46.
- Estrada, A., Katselis, G.S., Laarveid B. & Bari B., (2000), Isolation and evaluation of immunological adjuvant activities of saponins from *Polygajasenega*.L.Comparative Immunology.Microbial Infectious Disease23, 27-43.
- Fatope, M. O. Ibrahim, H. & Takeda, Y. (1993), "Screening of higher plants reputed as pesticides using the brine shrimp lethality assay. *International Journal of Pharmacognosy* 31, 250–254.
- Flora O. &Folasade D., (2008), Antimicrobial Effect of *Phyllanthusamarus*and*Parquetinanigrescens*on*Salmonella typhi.AfricanJournal of Biomedical Research* 11, 215 219
- Hervé, Z. Charles, K. Anoubilé, B. Janat, M. B. & Yves Alain, B. (2008), Phytochemical screening and determination of flavonoids in *Secamone afzelii*(Asclepiadaceae) extracts; *African Journal of Pure and Applied Chemistry* 2 (8), 80-82
- Hill, A.F. (1952), Economic Botany. A textbook of useful plants and plant products. New York: McGarw-Hill Book Company Inc,
- Iwu, M.M. Duncan, A.R. &Okunji, C.O. (199), New Antimicrobials of plant origin In JanickJ, (Ed).Perspectives in New crops and new uses. ASHS Press: Alexandria, 457-462.
- Kudi, A. C. &Myint, S. H. (1999), Antiviral activity of some Nigerian medicinal plant extracts. Journal of Ethnopharmacology 68, 289–294.
- Muanza, D.N., Kim, B.W., Euler, K.L. & Williams, L. (1994), Antibacterial and antifungal activities of nine medicinal plants from Zaire. *International Journal of Pharmacognosy*, 32, 337-345.
- Mulyani, Y.,Sukandar, E. Y.,Adnyana I. K. & Elfahmi (2012), Petiveria alliacea: New alternative for the treatment of sensitive and multi-resistant Mycobacterium tuberculosis; Journal of Pharmacognosy and Phytotherapy4 (7), 91-95
- Okoli, C. O., Onyeto, C. A., Akpa, B. P., Ezike, A. C., Akah, P. A. & Okoye, T. C. (2010), "Neuropharmacological evaluation of *Annona senegalensis* leaves. *African Journal of Biotechnology* 9 (49), 8435-8444
- Okwu, D.E.(2005), Phytochemicals, Vitamins and Mineral Contents of Two Nigerian Medicinal Plants. *International Journal of Molecular Medicine and Advance Sciences* 1, 375-381

- Olowosulu, A.K. & Ibrahim Y.K.E. (2006), Studies on the antimicrobial screening of Aqueous extracts of five plants used in Folk medicine in Nigeria. *West African Journal of biological science* 3 (5), 21-26
- Price K.R., Johnson T.I. & Fenwick, G.R., (1987), The Chemistry and Biological Significance of Saponins in Food and feeding stuffs. *Critical Reviews in Food Science and Nutrition* 26, 22-48
- Sahpaz, S., González, M.C., Hocquemiller, R., Zafra-Polo, M.C. & Cortes, D. (1996), Annosenegalin and annogalene: two cytotoxic mono-tetrahydrofuran acetogenins from *Annona senegalensis* and *Annonacherimolia.Phytochemistry* 42(1), 103-107
- Sahpaz, S., Laurens, A., Hocquemiller, R., Cavé, A., &Cortés, D. (1994), Senegalène, unenouvelle acétogénineoléfinique mono-tetrahydrofuranique des grainesd' Annona senegalensis. Cannadian Journal of Chemistry 72, 1533-1536
- Sandhya, B., Thomas, S. & Isbael, R. (2006), Complementary and alternative medicines 3, 110-114.
- Souto, X.C.,Bolano, J.C., González, L. & Reigosa, M.J. (2001), Allelopathic effects of tree species on some soil microbial populations and herbaceous plants. *Biology of Plant* 44, 269-275.
- Szczepanski, C.,Zgorzelak, P.& Hoyer, G. A.(1972),Isolierung, Strukturaufkl"arungundSyntheseeinerantimikrobiellwirksamenSubstanzaus Petiveria alliacea L.Arzneimittel Forschung22, 1975–1976
- Tavs, A. A.& Doris, N. O. (2012), Pharmacognostic Evaluation of the Leaves of Secamone afzelii(Schult) K Schum (Asclepiadaceae. Tropical Journal of Pharmaceutical Research 11 (1), 125-131
- Tropical Plant Database–Anamu (*Petiveria alliacea*), Rain tree. Available from <u>http://www.rain-tree.com/anamu.html</u> <u>Accesses 7/2/2017</u>
- Tsuchiya, H.M.S., Miyazaki, T., Fujiwara, S., Taniyaki, S., Ohyama, M., Tanaka, T. &Inuwa M. (1996), Comparative study on the antibacterial activity of bacterial flanones against methcillin resistant *staphylococcu saureus*. *Journal of Ethrophamacology* 50, 27–34
- Umedum,N.L.,Nwajagu, U., Udeozo, I.P.,Anarado, C. E. &Egwuatu C.I. (2014),The Efficacy of *Hyptis Suaveolens*: A Review of Its Nutritional and Medicinal Applications. *European Journal of Medicinal Plants* 4 (6), 661-674
- Vermani, K. &Garg, S. (2002), Herbal Medicines for Sexually Transmitted diseases and AIDS. Journal of Ethnopharmacological 80, 49-66.
- Verpoorte, R. (1998), Chemodiversity and the biological role of Secondary metabolites, some thoughts for selecting plant material for drug development. *Proceedings of the Phytochemical Society of Europe*, Kluwer Publishers 343, 11-24.
- Villar, R., Calleja, J.M. & Morales C. (1997), Screening of 17 Guatemalan medicinal plants for platelet antiaggregant activity. *Phytotherapy Research* 11, 1–5.
- Waterman P. H., (1992), Searching for bioactive compounds various strategies. Journal of National Products 53 (1), 13-22.
- Watt, J.M.&Breyer–Brandwijk, M.G. (1962), The Medicinal and Poisonous Plants of Southern and Eastern Africa. E& S Livingstone Ltd, Edinburgh &London.pp 137-744
- Weniger, B., Rouzier, M., Daguilh, R., Henrys, D., Henrys, J.H., & Anton, R (1986), Popular Medicine off The Central Plateau of Haiti. Journal of Ethnopharmacological 17 (1), 13-30
- Widowati,L.(2007), Khasiatpegagandaripenumpas tuberculosis hinggapeningkatdayaingat.
- Availablehttp://thibbunnabawi.wordpress.com/2007/11/22/khasiat-pegagan-daridaya- ingat penumpas-tbc-sampai-peningkat-

Test		Plant extracts		
		Secamoneafzel	Anonnasenegalen	Petiveriaalli
		li	sis	асеа
Alkaloid	Dragendorff's reagent	+ve	+ve	+ve
	Mayer's reagent	+ve	+ve	+ve
	Wagner's reagent	+ve	+ve	+ve
Flavonoids	Ammonia/H ₂ SO ₄	+ve	+ve	+ve
	Aluminium solution	+ve	+ve	+ve
	Ethyl acetate/Ammonia	+ve	+ve	+ve
Saponins	Frothing	+ve	+ve	+ve
Tannins	Ferric chloride	+ve	+ve	-ve
Anthraquinones	Chloroform/ammonia	+ve	+ve	+ve
Terpenoids	Chloroform/H ₂ SO ₄	+ve	+ve	+ve
cardiac glycosides	Keller- Kiliani test	-ve	-ve	-ve
Steroids	Chloroform/acetic	+ve	+ve	+ve
	anhydride/H ₂ SO ₄			

 Table1: Phytochemical screening of methanol extracts of Secamoneafzelli, Anonna
 senegalensisand

 Petiveriaalliacea
 senegalensisand

Table 2: Antibacterial activity of the methanol extracts of *Secamone afzelli, Anonna senegalensis* and *Petiveria alliacea*

Test organisms	Diameter Zone of Inhibition in mm			
	Secamone afzelii	Petiveria	Annona	Streptomycine
		alliaecea	senegalensis	sulphate
				(control)
Staphylococcus aureus	7.91 ^b ±0.61	$7.62^{\circ}\pm0.45$	$6.58^{abc} \pm 1.27$	23.89 ^a ±4.42
Bacillus subtilis	$9.99^{b} \pm 2.00$	7.53°±0.76	$8.46^{bc} \pm 1.67$	22.13 ^a ±4.99
Sylmonelatyphimurium	8.57 ^b ±0.41	$6.74^{ m bc} \pm 0.87$	$9.35^{bc} \pm 1.15$	23.85 ^a ±4.11
Enterococcus spp	$8.63^{b} \pm 0.81$	$6.87^{bc} \pm 0.58$	$9.80^{ m bc} \pm 0.79$	28.45 ^a ±1.93
Klebsiela aerogenes	$8.90^{b} \pm 1.11$	$6.40^{ m bc} \pm 0.76$	$6.97^{\circ} \pm 0.49$	29.00 ^a ±3.21
Escherichia coli	3.63 ^a ±0.41	$3.77^{ab} \pm 1.21$	$10.10^{a}\pm0.52$	$17.88^{a} \pm 0.65$
Pseudomonas aeruginosa	$8.64^{b}\pm 0.07$	$4.65^{abc}{\pm}0.90$	4.96 ^a ±1.19	28.84 ^a ±1.11
Pseudomonas syringine	3.74 ^a ±0.41	2.20 ^a ±1.00	6.27 ^a ±0.14	29.25 ^a ±0.32
Xantomona saxonopodis	$7.49^{ab} \pm 0.25$	7.13 ^{bc} ±0.43	5.23 ^a ±0.28	29.39ª±0.65
Shigella dysenteriae	$7.09^{ab} \pm 1.23$	$5.07^{abc} \pm 0.13$	5.56 ^a ±0.33	26.43 ^a ±1.23
Erwinia herbicola	$6.13^{ab} \pm 0.49$	5.29 ^{abc} ±0.23	$4.98^{a}\pm0.59$	$21.82^{a} \pm 2.04$

Values are means of duplicate \pm standard error. Column means followed by the same superscript letters are not significantly different at P<0.05.

Table 3: Antifungal activity of methanol extracts of Secamone afzelli, Anonna senegalensis and Petiveria alliacea

Test organisms	Mycelinl Growth Inhibition in Percentage				
	Secamone afzelii	Petiveriaalliae	Annona	Koside	
		сеа	senegalensis	(control)	
Fusarium oxyporium	69.35°±2.61	$67.34^{d} \pm 1.34$	$68.86^{b}\pm 2.10$	$86.89^{b} \pm 0.95$	
Marcophomina phomoides	34.11 ^a ±0.91	$13.88^{a} \pm 0.09$	$61.77^{a} \pm 1.53$	$73.71^{a} \pm 0.90$	
Ceratocystisparadoxa	53.41 ^b ±0.56	$69.60^{d} \pm 0.54$	$70.86^{b} \pm 4.20$	$87.40^{b} \pm 0.33$	
Collectotrichumlindimutian um	$57.52^{b}\pm0.47$	$50.74^{b} \pm 1.05$	70.31 ^b ±1.37	88.64 ^b ±2.00	
Helminthosporiumtoxicum	$76.97^{d} \pm 0.93$	63.37°±0.33	74.59 ^b ±0.56	$90.74^{b}\pm 2.33$	

Values are means of duplicate \pm standard error. Column means followed by the same superscript letters are not significantly different at P<0.05.