INTRODUCTION
Tuberculosis (TB) is a chronic bacterial infection caused by Mycobacterium tuberculosis. It is easily transmitted from one person to another through the air by droplet nuclei (Moulding, 1988). Tuberculosis remains a leading cause of death in the world from a single infectious agent. It is estimated that one-third of the world’s population is infected with the tubercle bacillus and about 80% of individuals diagnosed with the disease every year live in the 22 most populous countries such as China, India and Pakistan (Dye, 2006). Medicinal plants have enjoyed use in virtually all cultures as sources of medicines (Plotkin, 1991). The history of the use of medicinal plants for their therapeutic purposes probably dates back to the origin of man. Fossil records date human use of plants as medicines to at least the middle Paleolithic age, some 60,000 years ago (Solecki, 1977). Ancient texts of India and China contain exhaustive depictions of the use of a variety of plant-derived medications (Ahmed et al., 2006).

In Africa, Asia and Latin America, traditional medicine and medicinal plants have continued to play a very important role in their health care delivery systems. According to the World Health Organization (WHO 2003a), it is estimated that about 80% of the population in these developing countries rely on traditional medicine for their primary health care needs. There has also been an increase on the reliance of the use of medicinal plants by the population in the industrialized societies, as herbal remedies have become more popular in the treatment of minor ailments and also on account of the increasing costs of personal health maintenance (Horeau and Dasilva, 1999). In 2001, it was estimated that approximately 25% of prescription drugs originated from plants, 121 active compounds were then in use (Houghton, 2001; Rates, 2001).

However, the difficulty in the use of medicinal plants is that they are used without any standardization. This makes it difficult to document and institute a system of verification or assessment of the efficacy of the treatment. Thus, the local herbal medicine practitioner is quick to profess the efficacy of these remedies but stops short of providing a sound scientific basis and explanation for the remedy and more often than not thrives on the long, continuous and sometimes uneventful use of the remedy for treatment. Plants may contain constituents that can be used to treat diseases such as infections, inflammatory conditions and cardiovascular diseases, but the scientific information on most of these medicinal plants in use are lacking. Therefore, as part of the efforts to promote the use of medicinal plants either as an alternative or an adjunct to conventional medicine, it is necessary for scientists to carry out investigations into herbal medicines. This will help bridge the gap between conventional and herbal medicines. For a long
time, medicinal plants and herbs were used intensively in folkloric medicine for treatment of various diseases. Among the plant species that show relevant significant importance include Calotropis procera and Garcinia kola. On the part of Calotropis procera, the leaf is traditionally considered for the treatment of tuberculosis and it is a flowering plant in the dogbane family, Apocynaceae, that is native to North Africa, Western Africa, South Asia, and China. The green globes are hollow but the flesh contains a toxic milky sap that is extremely bitter and turns into a gluey coating resistant to soap. Common names for the plant include Sodom apple, King's crown, rubber bush, or rubber tree. The name apple of Sodom is derived from Hebrew; the milky sap contains a complex mixture of chemicals, some of which are steroidal heart poisons known as "cardiac aglycones". These belong to the same chemical family as similar chemicals found in foxgloves (Digitalis purpurea).

Garcinia kola Bitter cola is another plant species used in the treatment of tuberculosis, the plant is essentially found in some parts of Nigeria. It belongs to the family “Guittifferea". Among the Yorubas, it is called ‘Orogbo’, the Igbos calls it ‘Agbilu’, ‘Adi’ or ‘Aki ilu’ while to Hausas, this very popular nut is known as ‘Namijin Goro.’ It is a wonderful agricultural product with a wide range of applications in natural and orthodox medicine. Bitter cola is potent, with antibiotic properties, which could be effective in the treatment of many ailments and infections. Bitter cola has lots of health benefits, like the treatment of cough, sneezing, cold, diarrhea, tuberculosis, bacterial infection and fever. "It improves lung function by expanding the alveolar ducts and sac in the lungs thereby improving and strengthening the fibers in the lung tissue" (De Caluwé et al., 2009).

Study Area
The study was conducted at Infectious Diseases Hospital (I.D.H), located at France road, Kano, state, Northern Nigeria. The state is within the savannah zones of the country, latitude 11°30’N and longitude 8°30’ E. It has an elevation of about 525 meters above the mean sea level.

Collection, Identification and Preparation of the plant materials.
The plants used in this study were selected from the list of plants used by local herbalist in the preparation of various medicaments and their history in the folkloric medicine, used for curing tuberculosis. Two plants i.e. seed of Garcinia kola (Bitter kola) and leaves of Calotropis procera (Sodom apple) were collected. They were properly identified at the Department of Plant Biology, Bayero University Kano with the following herbarium accession number
A. Calotropis procera (Sodom apple) BUKHAN 0132
B. Garcinia kola (Bitter kola) BUKHAN 0433
C. Calotropis procera
The leaf of plants Calotropis procera (Sodom apple) was collected from the old campus of Bayero University Kano which is located between latitude 13° 21’ N, 42° E in Gwale local government area of Kano state Northern Nigeria. The leaf of the growing plant collected, was thoroughly washed under running tap water and rinsed with distilled water and finally shade dried. The dried leaves were crushed into powder form using motor and pestle as described by Fatope et al.(1999).

Powdered samples were then stored into air- tight container until used.

Garcinia kola
The Garcinia kola was bought from the popular Mariri ‘Yangoro Market of Kumbotso local government area, of Kano state. It was peeled and washed using running tap water and then rinsed with distilled water, later the seeds were crushed using mortar and pestle, spread under shade till it was dried and then kept in a container until used.

Extraction and Preparation of the Plant Material
Two hundred grams each of the powdered plant sample was extracted exhaustively with 95% methanol. The percolation processes was carried out on each of the fine powdered plant sample. The processed plant material was placed in closed vessels, followed by addition of 95% methanol and then later allowed to stand for fourteen days with regular shaking occasionally under room temperature. The extracts were then filtered through Whatman filter paper, No.3. Solid residue was pressed to recover much solution, liquid strained off. The percolation process was repeated three times, the second and third residue were allowed to stand until it is evaporated before adding to the first residue, concentrated extract of C. Procera yielded (8.1g) and G. kola (19.1g) and was referred to as Crude Methanol Extract (CME). Filtrates were labelled and stored respectively until used.

Preliminary Phytochemical Screening
The screening methods were carried out using the procedure described by Harbone (1984), Sofowora (1986), Mukhtar and Okafor (2002). The following active constituents were tested for: - alkaloids, tannins, flavonoids, cyanogenic glycosides, anthaquinone, saponins, anthrocyanosides (anthrocyanin pigment) and reducing sugar compounds.

Test for Terpenes:
Salwoski test:
0.3g of the extract was dissolved in 1ml chloroform followed by addition of 1ml of concentrated H₂SO₄ down the side of the test tube to form two phases. Formation of red or yellow coloration indicated the presence of sterols (Silva et. al., 1998).

Test for flavonoids:
Sodium hydroxide test:
0.3g of the extract was dissolved in water and filtered. To this, 2ml of 10% aqueous sodium hydroxide was added to produce a yellow coloration. A change of color from yellow to colorless on addition of dilute hydrochloric acid was an indication of the presence of flavonoids (Trease and Evans, 2002).

Test for Alkaloids:
0.5g of the extract was stirred with 5ml of 1% aqueous HCl on water bath and then filtered, 1ml each of the filtrate was taken in two test tubes. To the first portion few drops of Dragendorff’s reagent were added to observe the presence of orange-red precipitate while to the second portion 1ml of Mayer’s reagent was added. Formation of a blue colored precipitation indicated the presences of alkaloids (Sofowora, 1993).

Test for Tannins:
About 0.5g of the extract was stirred with 10ml of distilled water and then filtered. Few drops of 1% ferric chloride solution were added to 2ml of the filtrate, formation of a blue-black, green or blue-green precipitate indicated the presence of tannins (Trease and Evans, 2002).

**Test for Saponins:**
About 0.5g of the extract was shaken with water in a test tube. Frothing which persist for 15 minutes indicated the presence of saponins (Silva et al., 1998).

**Test for Anthraquinones**
0.5g of extract was shaken with 10ml of benzene, the content was filtered and 5ml of 10% ammonia solution was added to the filtrate the mixture was shaken. Presence of a pink, red or violet color in the ammoniacal layer (lower phase) indicated the presence of free anthraquinones (Trease and Evans, 2002).

**Test for glycosides**

**Keller-killiani Test**
The extract was reduced to dryness after which 5mg was dissolved in 2ml chloroform. Tetraoxosulphate (VI) acid was added to form a layer and the purple color at the inter-phase indicated the presence of glycosides (Trease and Evans, 2002).

**Test for carbohydrates**

**Fehling’s Test**
Fehling A and Fehling B reagents were mixed together and few drops of extracts were added and boiled. A brick red colored precipitate of cuprous oxide forms, confirmed the presence of carbohydrates (Trease and Evans, 2002).

**Test for reducing sugars**
One ml each of Fehling’s solutions I and II was added to 2 ml of the aqueous solution of the extract. The mixture was heated in a boiling water bath for about 2 – 5 min. The production of a brick red precipitate indicated the presence of reducing sugars(Silva et al., 1998).

**Test for steroids**
Salkowski method was used to test for steroids. About 0.5 g of the extract was dissolved in 3 ml of CHCl3 and filtered. To the filtrate concentrated H2SO4 was added to form a lower layer. A reddish brown color was taken as positive for steroid ring(Silva et al., 1998).

**Test for Amino Acid (Ninhydrin test)**
Two drops of freshly prepared 0.2% ninhydrin reagent was added to 0.3g of the extract and heated. Development of blue color indicated the presence of Amino acids(Trease and Evans, 2002).

**Test for proteins (Biuret test)**
0.3g of the extract was added to 1 ml of 40% sodium hydroxide and 2 drops of 1% copper sulfate solutions were added. A violet color indicated the presence of proteins(Trease and Evans, 2002)

**Test for resins**
Two (2g) of the methanic extract was dissolved in 10ml of acetic anhydride. A drop of concentrated sulphuric acid was added. Appearance of purple color, which rapidly changed to violet, was indicative of the presence of resins (Trease and Evans, 2002).

**Test Organism**
Clinical isolate of mycobacterium tuberculosis were used in the study and it was collected from infectious diseases hospital (I.D.H) Kano.

**Inoculums preparation**
A sterilized wire loop was used to scrape the colony of *Mycobacterium* and it was dipped into 5ml normal saline solution and gently shaken. The turbidity of the actively growing bacterial suspension was adjusted to match the turbidity standard of 0.5 Macfarland unit, it was prepared by mixing 0.5ml of 1.75(w/v) barium chloride dehydrate with 99.5ml of 1% (v/v) sulphuric acid. The grown suspension was used for further testing.

**Preparation of stock solution**
The stock solution of plant extract was prepared in a screwed capped bijou bottle different concentration ranging from (250,500,1000 and2000µg) of the extract was dissolved in 5ml Dimethly sulphoxide (D.M.S.O) and the solution was kept at room temperature until used.

**Preparation of Culture Media**
About 6.2g of Lowenstein-Jensen (LJ) powdered medium was measured and poured into a conical flask containing 100ml of sterile distilled water. It was then shaken until the media dissolved evenly into the solution. The conical flask was then covered with foil paper to avoid impurities from air. Thereafter, it was taken into an autoclave for between45-50min. 2ml of glycerol was added into the conical flask containing the LJ medium, aseptically a fresh egg was smashed into a beaker and stirred gently until it was evenly mixed. The egg mixture was poured into the conical flask containing the media, while continuously shaking until mixed evenly. The LJ solution was then poured into Petri-dish and allowed to cool and hardened. Later the Petri-dishes were sealed up with tape to prevent air passage and kept at room temperature until used, as it was described by manufacturer.

**Preparation of Sensitivity Discs**
Sterilized paper disc of No 1 Whatmann filter paper was punched using puncher to an approximately 6.0mm in diameter and were placed in a screwed capped bijou bottle and sterilized in dry heat oven at 140°C for 1hour and the discs were allowed to cool 500,1000,2000,5000 and 10,000µg/ml of each plant extract concentration was dissolved in Dimethly sulphoxide (D.M.S.O) solution. The sterilized paper discs were soaked in each concentration of plant extract sample respectively and allowed to stand until it soaked all the solution. They were kept at room temperature until used. Standard Rifampicine antibiotic disc was used as control.

**Bioassay Procedure**
The antimycobacterial activities of the extract were determined by the Kirby-Bauer (1966) agar diffusion method according to NCCLS standard (2000). Lowenstein Johnson medium were used for the antimycobacterial test under septic condition. 15ml of the LJ medium were dispensed into pre-sterilized Petri-dish.
A sterile wire loop was dipped into the prepared inoculum of *Mycobacterium tuberculosis* isolate and then it was gently streaked on the Petri-dish turning the plate to an angle 360° until the entire surface of the media was covered. Using sterile forceps aseptically the sterile extract discs were placed on the surface of each inoculated media and they were pressed down gently to ensure contact with the agar surface. The discs were spaced about 20mm to avoid overlapping ring of the zone of inhibitions and the control disc of rifampicin was placed at the center of the Petri-dish. It was then covered and incubated. Readings was taken on the 7th day after inoculation and subsequently once a week for five weeks.

**Minimum Inhibitory Concentration (MIC)**

Minimum Inhibitory Concentration (MIC) is the lowest concentration of the plant extract that inhibits the visible growth of *Mycobacterium*. This was done using agar well diffusion method as described by Kirby-bauer (1966) and demonstrated by Arzai (2002). From the prepared inoculum wire loop was used to streak on the surface of the Lowenstein Johnson medium. Using a sterilized cock borer a hole of about 4mm was punched into the media leaving a space in between each hole. About 0.5ml of each prepared extract concentration was pipette and poured into respective hole. Readings were observed on each Petri-dish and recorded.

**RESULTS**

**Phytochemical studies**

Phytochemical studies were conducted on methanolic extract of the two plant species. Result indicated that the extract had different characteristics in terms of color, odor and texture. *G. kola* yielded more in weight (19.1g).

| Plant sample | Plant part | Initial weight (g) | Final weight (g) | Color | Odor         | Texture |
|--------------|------------|--------------------|------------------|-------|--------------|---------|
| *C. procera* | Leaf       | 200                | 8.1              | Green | Repulsive    | Oily    |
| *G. kola*    | Seeds      | 200                | 19.1             | Black | Repulsive    | Oily    |

From the phytochemical characteristics of the crude methanol extract of the two plants, tannins and flavanoids were found to be present among in both plants, while resins were absent. Furthermore there was variation in the occurrence of the phytochemicals among the plant as *G. kola* had saponins while *C. procera* had glycosides and steroids.

**Table 2: Phytochemical characteristics of the plant extracts**

| Phytochemical     | *C. procera* | *G. kola* |
|-------------------|--------------|-----------|
| Carbohydrates     | +            | -         |
| Tanins            | +            | +         |
| Terpenoids        | -            | -         |
| Flavanoids        | +            | +         |
| Saponins          | -            | +         |
| Alkaloids         | -            | +         |
| Glycosides        | +            | -         |
| Anthroquinone     | -            | +         |
| Proteins and amino acids | +     | -         |
| Steroids          | +            | -         |
| Reducing sugar    | -            | +         |
| Resin             | -            | -         |

Key (+) Detected  (-) Not Detected

From the results in table 3 below, it could be seen that *Calotropis procera*, *Garcinia kola*, and Rifampicin had a mean diameter (zone of inhibition) of 11.38mm, 6.2mm, and 17.8mm respectively.

**Table 3: Diameter of zone of inhibition (mean±std) of the plant extracts**
From the results presented in the table 4 below it could be seen that *Garcinia kola* fell within a subset 1 with a zone inhibition 6.20mm. *Calotropis procera* fell under subset 2 with a zone of inhibition having a diameter of 11.40mm and the Rifampicin which is the control has a diameter of 17.80mm around the zone of inhibition. The study further revealed that *Garcinia kola* had lower effect on the *Mycobacterium tuberculosis* compared to *Calotropis procera*. Both plant extracts (*Garcinia kola* and *Calotropis procera*) were not as effective as Rifampicin.

Table 4: Scheffé Multiple Comparison Test of the plant extract on inhibition of *Mycobacterium tuberculosis* growth.

| Zone of Inhibition (mm) | Treatment                  | N  | Subset for alpha = 0.05 |
|------------------------|----------------------------|----|------------------------|
|                        | *Garcinia kola*             | 25 | 1                      |
|                        | *Calotropis procera*        | 25 | 2                      |
|                        | Rifampicin                 | 25 | 3                      |

Means for groups in homogeneous subsets are displayed

From the results in the table 5 below, the *Calotropis procera* concentrations at 500, 1000, 2000, 5000 and 10000µg/ml had mean diameters of 7.06mm, 8.92mm, 11.96mm, 11.96mm and 16.98mm respectively.

Table 5: Effect of *Calotropis procera* leaf extract on inhibition of *Mycobacterium tuberculosis* growth.

| Concentration   | N  | Mean± std.     |
|-----------------|----|----------------|
| 500 µg/ml       | 5  | 7.06± 2.01817  |
| 1000 µg/ml      | 5  | 8.92± 4.21746  |
| 2000µg/ml       | 5  | 11.96± 3.78259 |
| 5000 µg/ml      | 5  | 11.96± 3.78259 |
| 10000µg/ml      | 5  | 16.98± 5.06429 |

From the results presented in the table 6 below it could be seen that the 500µg/ml, 1000 µg/ml, 2000 µg/ml, 5000 µg/ml had fallen within a subset group 1 having a diameter (zone of inhibition) of 7.06mm, 8.92mm, 11.96mm and 11.96mm respectively and they had similar effect on the *Mycobacterium tuberculosis* while concentration level of 10000 µg/ml had a diameter of 16.98mm had fallen under the subset group 2. The analysis therefore reveals that the *Calotropis procera* concentration level of 10000µg/ml is the most effective and therefore the analysis accepted the alternative hypothesis and rejects the null hypothesis.

Table 6: Multiple comparison test for the *Calotropis procera* concentration levels on inhibition of *Mycobacterium tuberculosis* growth.

| Concentration of *Calotropis procera* | N  | Subset for alpha = 0.05 |
|--------------------------------------|----|------------------------|
| 500µg/ml                             | 5  | 7.06                    |
| 1000µg/ml                            | 5  | 8.92                    |
| 2000µg/ml                            | 5  | 11.96                   |
| 5000µg/ml                            | 5  | 11.96                   |
| 10000µg/ml                           | 5  | 16.98                   |

Means for groups in homogeneous subsets are displayed

The result indicated that *C. procera* leaf extract was effective across the concentration used but was not effective at 250µg/ml,
and *G. kola* seed extract showed inhibition at 2000µg/ml and 1000µg/ml but growth was not inhibited at 500µg/ml and 250µg/ml.

### Table 7: Minimum inhibitory concentration of C.M.E of plant sample on inhibition of *Mycobacterium tuberculosis* growth.

| Plant sample | Extract concentration (µg/ml) |
|--------------|-------------------------------|
|              | 2000 | 1000 | 500 | 250 |
| *C. procera* | +    | +    | +   |    |
| *G. kola*    | +    | +    |     |    |

Key: (+) Growth inhibited, (-) Growth not inhibited

### DISCUSSION

Phytochemical analysis is very useful in the evaluation of bioactive and biochemical components of seeds and other parts of plants. The results of the preliminary phytochemical screening carried out on the crude methanolic extracts of *Calotropis procera* and *Garcinia kola* revealed the presence of tannins and flavonoids in both plant samples. This agrees with the work of (Musa et al., 2000) which also found the same phytochemical in *C. procera* leaf. Carbohydrate, proteins and amino acids occurred only in *Calotropis procera* while alkaloids, anthroquinones and reducing sugar were only present in *G. kola*. (Kawo et al., 2009) also reported similar phytochemicals during extraction of *C. procera* leaf and latex. The phytochemical constituents may be associated with the antimicrobial activities of the plant (should be the other way round. In any case, you have not stated your own result before discussing it). Tannins and flavonoids have been reported to possess antimicrobial activities (Cowan, 1999). Thus the antimicobacterial activity of tannins may be related to their ability to inactivate microbial adhesion of enzymes (Cowan, 1999). Tannins have been reported to inhibit growth of microorganisms by precipitating microbial protein and making nutritional proteins unavailable to them (Ogunleye and Ibitoye, 1986). The result obtained in the experiments agreed with an earlier report of Aliyu (2006) who recorded similar results for the extract of *Garcinia kola*. He further reported that the invaluable pharmacological potential of *Garcinia kola* such as antioxidant, antibacterial, antiviral, antifungal and anti-inflammatory properties may be attributed to the presence of these bioactive compounds. The presence of these components in this species is an indication that it may have some medicinal potential. This is due to the fact that each of the components identified has one therapeutic usage or another. For instance, plants rich in saponins have immune boosting and anti-inflammatory properties (Kenner and Requena, 1996). Similarly, tannins have been reported to have antibacterial potential due to their basic character that allows them to react with proteins to form stable water soluble compounds thereby killing bacteria by directly damaging its cell membrane (Elmarie and Johan, 2001). The antibacterial activities of alkaloids and flavonoids have been reported by a number of authors (Hassan et al., 2005; Alierio et al., 2008; Yesmin et al., 2008). Elsewhere in Democratic Republic of Congo similar observations have been made on plants employed for traditional medicines, which were known to contain the above mentioned bioactive components (Otshudi et al., 2000).

The ant tubercular activity of methanolic plant extracts of (*C. procera, G. kola*) and rifampicine (control) tested against the isolate (*M. tuberculosis*) has a mean diameter (zone of inhibition) of 11.38mm, 6.2mm, and 17.8mm respectively, as it was presented in table 3, however, it could be seen that between the two plant samples, *C. procera* had a higher mean value (zone of inhibition). This indicated that the plants have different activities on *Mycobacterium tuberculosis* isolate.

From the analysis done above it was found out that *C. procera* is having the highest degree of influence among the plant extract. It is of paramount importance to find out which of the concentrations will be most effective in the treatment of the *Mycobacterium tuberculosis* isolate which led us to further analysis of variance in respect concentration which ranges from (500, 1000, 2000, 5000, 10,000µg/ml).

From the descriptive statistics on the concentration of *C. procera* (table 6) it will be seen that 500, 1000, 2000, 5000 and 10,000 µg/ml had mean diameter (zone of inhibition) of 7.06mm, 8.92mm, 11.96mm, 11.96mm and 16.98mm respectively. Analysis of variance revealed a significant variation among the concentration levels at 1% probability. This led to accept the alternate hypothesis that says at least one of the concentrations had more significant effect in the treatment of *Mycobacterium tuberculosis* isolate than the other, and this was found to be 10,000µg/ml with the highest diameter (zone of inhibition) of 16.98mm. The result agrees with the work of Johnson et al. (2000) also reported that plant used to treat T. B. in Mbarara district is *G. kola* which contains some of the chemical constituent. Moreover Gill (1992) also stated that *C. procera* has some ameliorative effect on nasophaningeal and pulmonary diseases.

For the minimum inhibitory concentration the result indicated that *C. procera* leaf extract was effective across the concentration used but was not effective at 250µg/ml. *G. kola* seed extract showed inhibition at 2000µg/ml and 1000µg/ml but growth was not inhibited at 500µg/ml and 250µg/ml.

### CONCLUSION

It shows that the phytochemical analysis of the plant samples contain tannins, flavonoids, proteins and amino acids to mention but few and the crude methanolic extract of the two plant samples possess some antimicrobial activities with *C. procera* having the highest degree of activity due to the higher
zone of inhibition followed by *G. kola* with moderate activity among the plant extract. However all of them were not as effective as the control (Rifampicin). This provides the basis for the use of the plant in traditional medicine. However to evaluate the potential effectiveness in human beings, further studies are needed.

REFERENCES
Ahmed, M., Amin, S., Islam, M., Takahashi, M., Okuyama, E., and Hossain, C.F. (2006) Analgesic Principle from *Abutilon indicum*. Pharmacie 55:314

Aliero, A.A., B.L. Aliero, and U. Buhari (2008). Preliminary Phytochemical and Antibacterial screening of *Scadoxus multiflorus*, *International Journal of Pure and Applied Sciences*. 2(4); 13-17

Arzai, H.A. (2002). Preparation of Wet disk for Sensitivity Testing (Undergraduate Lecture notes) Unpublished, Bayero University, Kano.

Ayele W.Y., Svastova P., Roubal P., Bartos M., Pavlik I. (2008): *Mycobacterium avium* Sub-species para-tuberculosis cultured from locally and commercially pasteurized cow’s milk in the Czech Republic. *Applied and Environmental Microbiology*. 71, 1210–1214.

Aliyu, S.B. (2006). *Common Ethnomedicinal plants of the semi arid regions of West Africa their description and phytochemicals*, Triumph Publishing Company Limited Kano, Nigeria. Pp 193-196

Cowan, M.M. (1999). Plants Products as Antimicrobial agents. *Clinical Microbiology Review* 12: 564-582

De Caluwè, E., Halamová, K., and Van Damme, P. (2009). Baobab (*Adansonia digitata* L.): A review of traditional uses, phytochemistry and pharmacology. In: Rodolfo, H., Simon, J.E., Ho, C.-T. (Eds.), *African natural plant products: new discoveries and challenges in chemistry and quality*. Oxford University Press, USA, pp. 51–84.

Dye, C. (2006). Global epidemiology of tuberculosis *Lancet* 367,938–940.

Elmarie, V.W. and Jonah, C.P. (2001). Purification and identification of active antibacterial component in *Carpobrotus edulis*. *Journal of Ethnopharmacology*. 76: 87-91

Fatope M. O Ibrahim H., and Takeda Y. (1999) screening of higher plants reported as pesticides using brine shrimps assay. *International Journal of Pharmacognosy volume3 number1* Pp 250-260.

Hassan, S. W., Umar, R. A., Ebbo, A. A., and Matazu, I.K. (2005). Phytochemical, antibacterial and toxicity studies of *Parkinsonia aculeate* L. (Fabaceae), *Nigerian Journal of Biochemistry and Molecular Biology*. 20(2): 89-97

Harbene, J. B. (1984). Phytochemical Methods, *In: A guide to modern techniques of plant analysis*. Fakenhan Press, Britain 162 – 167.

Hoareau, L. and DaSilva, E. J. (1999). Medicinal plants: a re-emerging health aid. *Electronic Journal of Biotechnology*. 2, 56-70.

Houghton, P. J. (2001). Old yet new pharmaceuticals from plants, *Journal of Chemical Education*. 78(2), 175-184.

Johnson E, Long NH, Diwan VK, and Winkvist A. (2000). Gender and tuberculosis control: perspectives on health seeking behavior among men and women in Vietnam. *Health Policy* 52:33–51.

Kawo, A. H., Mustapha, A., Abdullahi, B. A., Rogo, L. D., Gaiya, Z. A. and Kumurya, A. S. (2009). Phytochemical Properties and Antibacterial Activities of the leaf and latex extracts of *Calotropsis procera*. *Bayero Journal of Pure and Applied Sciences*, 2(1):34-40

Kenner D and Requena Y (1996) Botanical medicine, a European professional perspectives paradigm publication Brookline Massachusetts p7-12

Kirby – Bauer (1966). Antimicrobial Sensitivity Testing by Agar Diffusion Meth. *African Journal of Clinical Pathology*, 44: 493.

Moulding, T., (1988). Pathogenesis, Pathophysiology and Immunology, In: Schloss-berg, D. (Ed.), *Tuberculosis*, 2nd edition. Springer-Verlag, NewYork, pp.33–22.

Musa, K.Y., A. Ahmed, H. Ibrahim, G. Arowosiaye and O.S. Olonitola (2000): Phytochemical antimiocrobes studies of *Acrypha racemosa*. *Nigerian Journal of Natural Products and Medicine* 4:67-69.

Mukhtar, M.D. and Okafor A.C. (2002). Bioactive Evaluation of Ethanol Extracts of Leaf and Stem – Bark of Guiera Senegalensis. *Nigerian Journal of Research and Production* 1 (1) Pp. 114 – 121.

Ogunleye B and Ibitoye E, (1986) *Medicinal plants in tropical West Africa*. Cambridge University Press, London

Otshudi, A. L., Foriers, A., Vercryse, A., VanZeebroeck, A. and Lawwers, S. (2000). In vitro antibacterial activities of six medicinal plants traditionally used for the treatment of dysentery and diarrhea in Democratic Republic of Congo (DRC), *Phytochemistry* 7(2):167-172.

Plotkin, M. J. (1991). *Traditional Knowledge of Medicinal Plants the search for new jungle medicine*. Cambridge University Press, Cambridge UK. 245-246.

Rates, S. M. K.(2001). Review: Plants as a source of drugs. *Toxicon science directs*, 39, 603-613.

Silva, G.L., I. Lee and K.A. Douglas, 1998. Special problems with extraction of plants In: *Cannell, J.P.R. (ed.). Natural Products Isolation*. Humana press publishers, New Jersey (USA). Pp 356-358.
Sofowora, A. (1986). Retrospect and Prospects. In: The state of medical plants Research in Nigeria. University of Ile press, Ile – Ife Nigeria 1 – 10.

Solecki, R. (1977), A Neanderthal flower burial in Northern Iraq. Science, 19, 880-881.

Trease and Evans, (2002), "Carbohydrates in Pharmacognosy", 5th edition, Ed. W.C. Evans, Harcourt Publisher Limited, London, Pp. 193

World Health Organization (WHO), (2003a). The promotion and Development of traditional Medicine, Technical Report Series, 622

APPENDICES

From table 1 below, it could be seen that F-value was found to be significant at 1% probability. This implies that the plant extracts selected differed in their effect on Mycobacterium tuberculosis and the research therefore accepted the alternative hypothesis and rejects the null hypothesis that says the plant extracts (treatment) have the same effect in the treatment of Mycobacterium tuberculosis.

Table 2: Analysis of Variance (ANOVA)

| Zone of Inhibition | Sum of Squares | Df  | Mean Square | F        | Sig. |
|--------------------|---------------|-----|-------------|----------|------|
| Between Groups     | 283.734       | 4   | 70.933      | 4.659    | 0.008 |
| Within Groups      | 304.492       | 20  | 15.225      |          |      |
| Total              | 588.226       | 24  |             |          |      |

** - Significant at 1%

From the results in the table 2 it could be seen that the F-value which measure the level of dispersion or variation between or among the concentration levels was found to be significant at 1% probability and the research therefore has to accept the alternative hypothesis that says at least one of the concentration is having more significant effect in the treatment of Mycobacterium tuberculosis than others. The results show that 10000µg/ml having a highest diameter of 16.98mm will have more significant effect than the other concentration.

Table 2: Anova of Calotropis procera Concentration Levels on inhibition of Mycobacterium tuberculosis growth.

| Diameter (mm) | Sum of Squares | Df  | Mean Square | F        | Sig. |
|---------------|---------------|-----|-------------|----------|------|
| Between Groups | 2517,416      | 3   | 839.139     | 23.369   | 0.00**|
| Within Groups  | 3447.211      | 96  | 35.908      |          |      |
| Total          | 5964.628      | 99  |             |          |      |

Least significant difference was found to be 14.13 and from the table 8 below of the mean value of Calotropis procera Concentration Levels on inhibition of Mycobacterium tuberculosis growth showed that all the calculated means were lower than the LSD value which indicated that there was no significance difference between concentration levels hence the null hypothesis was rejected.

Table 8: Least significant difference (LSD) table of Calotropis procera Concentration Levels on inhibition of Mycobacterium tuberculosis growth.

| Difference between Mean | Greater than (>) or Less than (<) LSD | Null hypothesis |
|-------------------------|---------------------------------------|-----------------|
| X1–X2                   | 7.06 – 8.92                           | 1.86            | H0, Rejected |
| X1–X3                   | 7.06 – 11.96                          | 4.90            | H0, Rejected |
| X1–X4                   | 7.06 – 11.96                          | 4.90            | H0, Rejected |
| X1–X5                   | 7.06 – 16.98                          | 9.92            | H0, Rejected |
| X2–X3                   | 8.92 – 11.96                          | 3.04            | H0, Rejected |
| X2–X4                   | 8.92 – 11.96                          | 3.04            | H0, Rejected |
| X2–X5                   | 8.92 – 16.98                          | 8.06            | H0, Rejected |
| X3–X4                   | 11.96 – 11.96                         | 0.00            | H0, Rejected |
| X3–X5                   | 11.96 – 16.98                         | 5.02            | H0, Rejected |
| X4–X5                   | 11.96 – 16.98                         | 5.02            | H0, Rejected |

Key: X (Mean) H0 (Null hypothesis)

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