Antimycobacterial activity and low cytotoxicity of leaf extracts of some African Anacardiaceae tree species.

Short Title: Antimycobacterial and cytotoxicity of Anacardiaceae tree species

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Treatment of tuberculosis is a challenge due to multi and extremely drug resistant strains of Mycobacterium tuberculosis. Plant species contain antimicrobial compounds that may lead to new anti-TB drugs. Previous screening of some tree species from the Anacardiaceae family revealed the presence of antimicrobial activity, justifying further investigations. Leaf extracts of 15 Anacardiaceae tree species were screened for antimycobacterial activity using a twofold serial microdilution assay against the pathogenic Mycobacterium bovis and multidrug resistant M. tuberculosis and rapidly growing mycobacteria, Mycobacterium smegmatis, Mycobacterium fortuitum and Mycobacterium aurum. The vaccine strain, M. bovis and an avirulent strain, H37Ra M. tuberculosis, were also used. Cytotoxicity was assessed using a colorimetric assay against Vero kidney, human hepatoma and murine macrophage cells. Four out of 15 crude
acetone extracts showed significant antimycobacterial activity with MIC varying from 50 to 100µg/mL. *Searsia undulata* had the highest activity against most mycobacteria, followed by *Protorhus longifolia*. *M. fortuitum* was the strongest predictor of activity against MDR-TB (correlation coefficient=0.65). Bioautography against *M. aurum* and *M. fortuitum* worked well as indicators of the Rf values of active compounds yielding strong zones of inhibition. The leaf extracts of *S. undulata* and *P. longifolia* had more than 10 different antimycobacterial compounds and had low cytotoxicity with LC50 values above 100µg/ml.

**Keywords**: Anacardiaceae; MDR-*M. tuberculosis*; antimycobacterial; cytotoxicity; selectivity
1 Background

Tuberculosis occurs in humans, cattle, wild animals and many other domesticated species. The *Mycobacterium tuberculosis* complex (MTC) comprises *M. tuberculosis*, *M. bovis*, *M. africanum*, *M. canetti*, *M. pinnipedii*, *M. caprae*, *M. microti*, *M. mungi*, Dassie bacillus, Oryx bacillus and the attenuated *M. bovis* Bacille-Calmette-Guerin (BCG) vaccine strain. With the exception of BCG, these species are pathogenic and can cause tuberculosis (TB) in mammalian hosts (Kabongo-Kayoka *et al.*, 2015).

Despite the fact that TB can be cured, this disease remains a worldwide public health threat due to the emergence of multi and extremely drug resistant strains and subsequent poor response to current antibiotics. One third of the world’s population is infected with *Mycobacterium tuberculosis* (MTB) and is hence at risk of developing active TB. In the order of 9.6 million people every year develop TB and 1.5 million die from the disease. This includes 0.4 million TB deaths among HIV positive people (WHO, 2015). From the 9.6 million, only 12% were HIV positive and it was also estimated that 480 000 cases of multidrug-resistant TB (MDR-TB) have occurred in 2014. This confirms that TB alone is still a major challenge.

There is currently no effective vaccine to protect against TB. In humans, control of the disease relies heavily on detecting infectious cases and treating them for at least six months with a combination of antibiotics. The course of antibiotics involves the administration of first-line anti-TB drugs, which are combinations of isoniazid (INH), rifampicin (RIF), pyrazinamide and ethambutol for 2 months to kill the rapidly growing bacteria. The treatment is continued in the next 4 months with a combination of INH and RIF because of their sterilizing activity to eliminate bacilli which are dormant in the macrophages, or slow growers (Rivers and Mancera, 2008). Toxicity associated with these first-line drugs and the long duration of treatment has led
to low patient compliance, giving rise to drug-resistant strains. Multidrug-resistant mycobacteria
are resistant to INH and RIF whereas extremely drug-resistant mycobacteria are resistant to
second-line drugs such as fluoroquinolones and to at least one injectable drug in addition to INH
and RIF. Hence, there is a need to find new anti-tubercular agents with novel modes of actions
(Chinsembu, 2016).

It is most likely that the emergence of resistant *Mycobacterium* isolates will continue in the
future, exhausting the current range of effective antibiotics available. Therefore, new classes of
anti-TB agents are urgently needed, and research projects seeking for alternative therapeutics
have been encouraged. Natural resources have been explored to develop new antibiotics.
Substances that can either inhibit the growth of pathogens or kill them while having little or no
toxicity to host cells are considered good candidates for developing new antimicrobial drugs
(Nguta *et al*., 2016).

Medicinal plants are used in many parts of the world to treat different ailments including TB-
related symptoms such as chest complaints, respiratory ailments, fever and coughing (McGaw *et
al*., 2008). Relevant reviews of traditional medicinal use of plants in Africa are available
(Hutchings *et al*., 1996; Kuete, 2010; Maroyi, 2013; Nguta *et al*., 2016; Stark *et al*., 2013; Van
Wyk *et al*., 1997).

South Africa possesses the richest temperate flora in the world, with an estimated 24 000 species
and intraspecific taxa in 368 families—more than 10% of the world’s vascular plant flora
(Germishuizen and Meyer, 2003). There is much potential for discovery of structurally
interesting metabolites with activity against *Mycobacterium* species from these plants. The
existence of a strong cultural reliance on traditional medicines within South African communities
justifies the importance of studying their safety and efficacy (Fennell *et al*., 2004).
In a study conducted on Indian medicinal plants, Gautam et al., (2007) reported that with many of the plant species examined, there was a strong positive correlation between antimycobacterial activity and ethnomedicinal use for TB and TB-related diseases. Although this provides support for investigating plants customarily used in other cultures to treat symptoms relating to TB, other plants tested for antimycobacterial activity using indicator organisms such as *M. smegmatis* and *M. bovis* BCG should also be investigated (McGaw et al., 2008). Testing plant material not only against ATCC strains but against current isolates is crucial as these microorganisms are part of the changing environment harbouring emerging resistant strains.

The Anacardiaceae, cashew or Sumac is a tropical family that comprises flowering plant bearing fruits that are drupes and in some cases producing urushiol which is an irritant substance (Mabberley, 1997). The Anacardiaceae comprises several genera of economic importance namely Cashew (*Anacardium occidentale*), Mango (*Mangifera indica*), Poison Ivy (*Toxicodendron radican, Smodingium argutum*), Sumac (*Searsia species*), Smoke tree (*Cotinus coggygria, Cotinus obovatus*), Marula (*Sclerocarya birrea*), Yellow mombin (*Spondias mombin, Spondias purpurea var. lutea*), Cuachalalate (*Amphipterygium adstringens*) and Pistacia (*Pistacia lentiscus, Pistacia integrrima*). This family which belongs to the order of sapindales is represented by 77 genera and 600 species mostly distributed in tropical, subtropical, and temperate areas (Mabberley, 1997). Very few plant species of the Anacardiaceae family have been investigated for antimycobacterial activity. One of the objectives of this study was to investigate *in vitro* antimycobacterial activities as well as cytotoxicity of selected plant species from the Anacardiaceae family. These plants were selected mainly based on a preliminary broad antimicrobial screening (Pauw and Eloff, 2014).
A summary on ethnobotanical use of the selected plant species from the Anacardiaceae family is presented in Table 1. Eight out of the 15 plants namely *Lannea discolor*, *Loxostylis alata*, *Ozoroa mucronata*, *Ozoroa paniculosa*, *Protorhus longifolia*, *Searsia chirindensis*, *Searsia lancea* and *Sclerocarya birrea* have antimicrobial properties mainly against Gram-negative and Gram-positive organisms. Three species (*Searsia chirindensis*, *Searsia undulata* and *Smodingium argutum*) have good anti-inflammatory activity. It would be useful to identify plant species with both antimycobacterial and anti-inflammatory efficacy as acute and chronic inflammations are induced during the course of tuberculosis (Volpe et al., 2006).

2 Materials and Methods

2.1 Source of plant materials and extraction

Leaves of the 15 plant species were collected in November 2009 from the Lowveld National Botanical Gardens in Nelspruit, the University of Pretoria Botanical Gardens and the Pretoria National Botanical Garden, South Africa in terms of a signed material transfer agreement. Leaves were kept in labelled bags with full identification and collection dates. Voucher specimens were kept in the Lowveld National Botanical Gardens (Nelspruit), HGWJ Schweickerdt Herbarium of the University of Pretoria and Phytomedicine Programme, University of Pretoria (Table 1). The leaves were air-dried at room temperature suspended in bags of net material in a ventilated room, ground to a fine powder in a Macsalab Mill (Model 200 LAB Eriez®), Bramley, Johannesburg, South Africa) and stored in closed glass containers away from light until needed. Three grams of each plant were extracted in 30 mL of acetone, (technical grade, Merck, Darmstadt, Germany) in a 50 mL polypropylene centrifuge tube. Acetone was used as the extracting solvent due to its ability to extract a wide range of compounds and low toxicity on different microorganisms (Eloff, 1998a). The tube was
vigorously shaken for 30 min on an orbital shaker, then centrifuged at 4000 × g for 10 min and the supernatant was filtered using Whatman No.1, 110mm diameter filter paper, before being transferred into pre-weighed glass containers (Eloff, 1998a). The extraction was repeated thrice on the same plant material and the solvent was removed by evaporation under a stream of air in a fume hood at room temperature to produce the dried extract. Extracts were made up to a concentration of 10 mg/mL in acetone for use in relevant tests.

Insert table 1

2.2 Antimycobacterial activity

The antimycobacterial activity of the acetone crude extracts of leaf powder was determined by using bioautography and the two-fold serial microdilution method. The pathogenic and rapidly growing mycobacteria were test microorganisms used for antimycobacterial activity.

2.2.1 Pathogenic mycobacteria

2.2.1.1 *Mycobacterium tuberculosis* multidrug resistant isolates (MDR-TB)

Two clinical isolates of multidrug resistant *Mycobacterium tuberculosis* (MDR-TB) were used. Both isolates were obtained from patients admitted to the MDR-TB ward at Tshepong hospital in Klerksdorp, North West province of South Africa in December 2012. The patients were a 34 year old female and an 18 year old male. Samples of sputum were submitted to the National Health Laboratory Services (NHLS) in Pretoria for culture in liquid medium and PCR/Line Probe Assay. Both isolates were resistant to isoniazid and rifampicin.
2.2.1.2 *Mycobacterium bovis*

*M. bovis* cultures were isolated from field samples of cattle from herds that tested positive on tuberculosis skin test during the period January 2009 to January 2011 (Kabongo-Kayoka *et al*., 2015).

2.2.2 Rapidly growing mycobacteria

The *Mycobacterium bovis* BCG (Pasteur strain P1172) was obtained from Tuberculosis laboratory at Onderstepoort Veterinary Institute and the *Mycobacterium tuberculosis* H37Ra which is routinely used as reference material at NHLS was obtained from the American Type Culture Collection (ATCC) number 25177. Other reference strains used included a *Mycobacterium aurum* strain from the National Collection of Type cultures (NCTC), a culture collection of Public Health England, *Mycobacterium aurum* (NCTC 10437), *Mycobacterium smegmatis* (ATCC 1441) and *M. fortuitum* (ATCC 6841).

2.2.3 Maintenance of cultures

Fresh cultures were used in the relevant assays. The pathogenic isolates of *Mycobacterium* spp. were kept at room temperature on Lowenstein-Jensen (LJ) slants supplemented with glycerol or pyruvate in the case of the *M. bovis* BCG cultures and were used within a month. Prior to each assay, cultures were reactivated in liquid medium, Middlebrook 7H9, using MGIT 960 tubes which were incubated at 37°C in the BACTEC MGIT 960 instrument, in which they were automatically monitored each hour for fluorescence development for 42 days or until a positive signal developed. Bacterial suspensions from MGIT tubes were then subcultured on solid medium LJ slants with pyruvate for *M. bovis* and LJ slants with glycerol for *M. tuberculosis*. Löwenstein Jensen tubes were then incubated in a walk-in incubator at 37 °C for 4 to 6 weeks. A stained Ziehl Neelsen smear was made from the sediment of the MGIT tube and the slant of LJ
medium. Reference cultures of *M. avium* (ATCC 25291), *M. bovis* BCG and *M. tuberculosis* H37Ra (ATCC 25177) were used as positive controls.

The rapidly growing *Mycobacterium* species were maintained on Middlebrook 7H10 Agar prepared according to the manufacturer instructions (Merck) and kept in a fridge at 4°C until used within a month. *Mycobacterium aurum* and *M. fortuitum* were revived using Middlebrook 7H9 broth with glycerol and tween and supplemented with OADC (Oleic acid Albumin Dextrose Catalase) whereas *M. smegmatis* was revived using MH broth (Merck). Bacterial suspensions were inoculated on MH agar plates for *M. smegmatis* and the rest on Middlebrook 7H9 Agar. All agar plates were then incubated at 37 °C until colony growth was visible within 24 to 72 hours depending on the growth rate of the specific mycobacterial species. The final density of $1 \times 10^5$ CFU/mL used was confirmed after spreading an inoculum of 100 μL of tenfold serial dilutions of the bacterial suspensions at McFarland No 1 onto Middlebrook 7H10 (M7H10) agar plates incubated at 37°C and counting colonies.

2.2.4 Minimum inhibitory concentration (MIC) determination

The MIC values were determined using the serial microplate method developed by **Eloff (1998b)** and slightly modified for mycobacteria by **McGaw et al. (2008)**. The MIC value was the lowest concentration of the tested extract or sample inhibiting the growth of the *Mycobacterium* species under investigation. Mycobacterial suspensions were prepared from a pure culture of fresh colonies from solid medium and suspended in Middlebrook 7H9 (M7H9) liquid medium supplemented with 10% OADC. These colonies were transferred into a sterile screw capped tube containing 3 ml of M7H9 broth and homogenized by placing the tube on a Vortex mixer for 5 minutes. After the larger particles had settled, the mycobacterial suspension was adjusted to McFarland no.1 turbidity standard by adding more broth (Lall and Meyer, 1999).
The assay was performed using sterile 96-well microplates with rounded bottoms. The sample to be tested (plant extract, drug, fraction) was prepared at a concentration of 10 mg/ml before serial dilution. One hundred µl of M7H9 broth or MH broth was added to all the wells then 100 µl of the sample to be tested were added in three adjacent column wells in the first row whereas other wells included standard drugs positive controls to assess bacterial growth, solvent control, broth to be inoculated and clean broth for sterility check. Two samples of crude extract and one standard drug were run per plate. A two fold serial dilution was carried out leaving 100 µl of different concentrations of diluted tested samples in each well starting with a concentration of 2.5 mg/ml in the first wells. Then 100 µl of the test bacterial suspension were added to all the wells except column 12 (negative control) containing pure broth to make sure that there was no contamination of the broth. Each dose was run in triplicate and the experiment repeated at least six times. The standard antibiotics (reference drugs) included ciprofloxacin, rifampicin, isoniazid and streptomycin starting with a concentration of 100µg/ml. The microplates were covered and sealed in plastic bags, placed in humid chambers to minimize the evaporation of the culture medium and incubated at 37°C for 24 hours (M. smegmatis), 48 hours (M. fortuitum) and 72 hours (M. aurum). For the pathogenic mycobacteria, microplates were incubated for a period of 7 to 15 days.

At the end of incubation, a volume of 40 µl of 0.2 mg/ml of iodonitrotetrazolium chloride (INT) was added to each well, plates were incubated for 30 minutes or longer at 37°C and the development of colour observed. A coloured red-purple formazan or pink color indicated the reduction of INT by metabolizing organisms whereas a yellow color or decrease in color indicated the inhibition of bacterial growth (Eloff, 1998b). If the colour development was not strong enough for slow growing organisms, plates were incubated much longer and monitored.
The correlation assessing the relationship between the MIC values of the 7 mycobacterial species used (Table 3) was calculated using Microsoft Excel 2010 software (Microsoft, Redmond, WA, USA).

2.3 Thin Layer Chromatography (TLC) analysis and Bioautography

Ten microliter from 10 mg/mL of relevant extracts (containing 100 μg of extract) was loaded on aluminium-backed TLC plates (Merck silica gel F254) in lines of about 1 cm wide. The TLC plates were eluted in three different mobile solvent systems that included ethyl acetate/methanol/water (EMW) 10:1.35:1, chloroform/ethyl acetate/formic acid (CEF) 10:8:2 and benzene/ethanol/ammonia (BEA) 18:2:0.2 (Kotzé and Eloff, 2002). Chromatograms were examined under ultraviolet light at wavelengths of 254 and 366 nm to visualize UV active compounds that fluoresce and might not be visible after vanillin spray; the plates were then sprayed with a solution made of 0.1 g vanillin in 28 ml of methanol and 1 ml of sulphuric acid (vanillin–Methanol-sulphuric acid) and heated at 110°C with a heat gun for colour development and visualization of UV inactive compounds present in the plant extract. More than 10 compounds of different polarities were observed. These compounds were very close to each other in both the solvent systems BEA and CEF. In general, the active plants had different activities against different Mycobacteria. Active compounds included both polar and non-polar compounds based on the Rf values determined in bioautography. In the non-polar solvent BEA, the non-polar compounds travelled some distance up the plate with clear zone of inhibition on plates sprayed with different Mycobacteria species whereas in the polar solvent system EMW, non-polar compounds were pushed towards the solvent front and polar compounds off the baseline. The CEF solvent system led to a better separation of some compounds (B in Figure 1).
For bioautography, only the rapidly growing organisms were used, namely *M. aurum*, *M. fortuitum* and *M. smegmatis*, because the pathogenic mycobacteria present a high biohazard risk if handled in a spray format. Aerosols of pathogenic species increase risk of possible contamination of the environment and human exposure. Duplicate chromatograms prepared as described above were left uncovered in a dark place and air dried until solvents were completely evaporated for at least 24 hours or overnight. A saturated bacterial culture was centrifuged at 3000 x g for 10 min and the pellet resuspended in fresh appropriate liquid growth medium and prepared at McFarland No 1. The TLC plates were then sprayed with active bacterial suspension (18 to 72 hour old cultures depending on the growth rate of the particular *Mycobacterium* species) until completely moist using a glass spray gun linked to a vacuum pump. The moist plates were incubated at 37°C in a humidified chamber under 100% relative humidity for 24 hours for *Mycobacterium smegmatis*, 48 hours for *M. fortuitum* and 72 hours for *M. aurum*. The plates were then sprayed with 2 mg/mL of INT (Sigma, Johannesburg, South Africa) in distilled water and incubated for a further 30 min to few hours until the development of clear zones against a purple-red background. The emergence of purple-red colour resulting from the reduction of INT into formazan indicated the presence of viable microorganisms whereas clear zones against the purple-red background indicated inhibition of mycobacterial growth indicating the Rf values of antimycobacterial compounds separated on the thin layer chromatography plates (Hamburger and Cordell, 1987).

2.4 *In vitro* cytotoxicity assay and selectivity index

The cytotoxicity of crude acetone extracts from plants with good MIC values and clear zones of inhibition on bioautography was determined using a tetrazolium-based colorimetric assay, 3-(4, 5-dimethylthiazol)-2, 5-diphenyl tetrazolium bromide (MTT assay) described by (Mosmann,
1983) and slightly modified by (McGaw et al., 2007). Vero African Monkey kidney cells were obtained from the cell culture collection of the Department of Tropical Diseases (University of Pretoria), cancer liver cells (hepatoma C3A) purchased from the American Type Culture Collection (ATCC), Manassas, USA, CRL-10741 and murine macrophage (RAW 264.7) cells purchased from ATCC, ATCC®TIB71™ were used. Cells were cultured in Minimal Essential Medium Eagle with L-Glutamine (EMEM) (BioWhittaker, Lonza, Belgium) supplemented with 0.1% gentamicin (Genta50) (Virbac, Centurion, South Africa) and 5% foetal calf serum (Highveld Biological, Sandton, Modderfontein, South Africa) for Vero cells and 5% sodium pyruvate (Adcock Ingram, Midrand, South Africa) and 10% of foetal calf serum were added for C3A cells. For C3A and Vero cells, a 5 day old confluent culture in 75 cm² flasks were harvested and centrifuged at 200×g for 2 min, and the pellet resuspended in growth medium to a density of 0.1 ×10⁶ cells/mL whereas RAW 264.7 cells were mechanically scraped, plated at a density of 4 × 10⁵ cells/mL onto 96-well plates containing 100 μL of Dulbecco’s Modified Eagle Medium (DMEM) (BioWhittaker, Lonza, Belgium) and incubated in a 37°C, 5% CO₂ incubator for 24 h prior to exposure. The following procedure was the same for all the different types of cells. A sterile 96-well microplate was used; cell suspension (100 μL) was added into each well of columns 2–11. Columns 1 and 12 were used as blanks with 200 μl of growth medium to minimize the “edge effect” and maintain humidity. The plates were incubated for 24 h at 37°C in a 5% CO₂ incubator and cells were observed until they reached the exponential phase of growth. The cells were then exposed to different concentrations of the extract sample as well as to Doxorubicin Hydrochloride (Adriblastina CSV, Pfizer, Johannesburg, South Africa) as positive control and acetone as negative control. From a concentration of 100 mg/ml of the extract sample, different concentrations using growth medium as a diluent were prepared,
namely 1, 0.75, 0.5, 0.25, 0.1, 0.075, 0.05 and 0.025 mg/ml. The microplates were then incubated for 48 h. After this period of incubation, the medium was removed from the wells containing the cells using a multichannel pipette attached to a pump via a plastic tube; cells were rinsed with 200 µl phosphate buffered saline (PBS, Sigma-Aldrich, Johannesburg, South Africa) and replaced with 200 µL of fresh medium. Then thirty (30 µl) of a 5 mg/ml solution of 3-(4, 5-dimethylthiazol)-2, 5-diphenyl tetrazolium bromide (Sigma-Aldrich, Johannesburg, South Africa) in PBS was added to each well. The plates were incubated for a further 4 h at 37°C in the CO₂ incubator. After incubation, the medium was carefully removed using the same multichannel pipette connected to a pump without disturbing the MTT formazan crystals in each well. Then 50 µl of dimethyl sulfoxide (DMSO) was added to each well to dissolve the MTT crystals absorbance values were read immediately using a microplate reader (BioTek Synergy HT, Analytical and Diagnostic Products, Johannesburg, South Africa) at a wavelength of 570 nm. Columns 1 and 12 were used as blank. The different concentrations were tested in quadruplicate and the assay was repeated at least three times. The LC₅₀ values were calculated as the concentration of plants extracts where 50% of cells were still viable compared to untreated cells in columns 2 and 11. The selectivity index of the active plants was calculated using LC₅₀ values divided by MIC values (LC₅₀/MIC).

2.5 Statistical analysis

All experiments were conducted in triplicate and values expressed as mean ± standard deviation. For the MIC values, re-evaluation of the growth inhibition was conducted where a wide variation within the three experiments was noticed. The relationship between the different mycobacterial MIC values was calculated using the Pearson’s correlation coefficient (r).

3 Results and Discussion
3.1 Acetone extract plant yield

The extraction yield of the plant species ranged from 0.8 to 18.8% (Table 2). The highest percentage was observed with Searsia magaliesmontana (18.8%) followed by Searsia undulata (12.5%) and Protorhus longifolia (8.3%). Heeria argentea had the lowest yield of 0.8%. The extraction yield and bioactivities of extracts using different extractants vary strongly (Kotzé and Eloff, 2002), hence the choice of acetone in this study.

Insert table 2

3.2 Biological activity of extracts

Bioautography of extracts using two (M. aurum and M. fortuitum) out of the three rapidly growing mycobacteria led to strong, clear zones of inhibition. This also corresponded with low MIC values obtained ranging from 0.05 to 0.1 mg/ml.

Extracts with MIC values \( \leq 0.1 \text{mg/mL} \) were considered to have significant activity; moderate activity was between 0.1 and 0.625 \( \text{mg/mL} \) and weak or poor activity (resistant) with MICs > 0.625 \( \text{mg/mL} \) (Kuete, 2010). All 15 plant extracts had good to moderate antimycobacterial activity against rapidly growing and pathogenic mycobacteria ranging from 0.05 to 0.590 mg/mL. Searsia undulata extracts had significant antimycobacterial activity with the lowest MIC value of 0.07 mg/mL for M. aurum and M. fortuitum followed by M. smegmatis with MIC of 0.09 mg/mL. Protorhus longifolia extracts also had significant antimycobacterial activity against all three non-tuberculous mycobacteria with MIC values of 0.11 mg/mL for both M. aurum and M. fortuitum and 0.07 mg/mL for M. smegmatis. Searsia lancea, Sclerocarya birrea and Harpehyllum caffrum had moderate activity of 0.420 mg/mL, 0.520 mg/mL and 0.590 mg/mL against M. aurum respectively whereas the same plant species had moderate activity with low MIC values of 0.21 and 0.11 mg/mL; 0.13 and 0.21 mg/mL and 0.21 and 0.11 mg/mL against M.
*M. fortuitum* and *M. smegmatis* respectively. *Searsia undulata* had significant activity against all mycobacteria including *M. bovis* and MDR-TB isolates with MIC ranging from 0.05 to 0.11 mg/mL, and the lowest value of 0.05 mg/ml against *M. tuberculosis* ATCC strain H37Ra. Positive correlation was observed between the pathogenic isolates, *M. bovis* and *MDR-TB*, with *M. fortuitum* with (r) coefficient correlation values of 0.62 and 0.65 respectively whereas negative correlation with value of -0.005 was observed with *M. aurum*. Good correlation was observed between ATCC strain, H37Ra and rapidly growing mycobacteria with values of 0.92, 0.87 and 0.37 for *M. fortuitum*, *M. aurum* and *M. smegmatis* respectively and the highest coefficient correlation value of 0.98 between *MDR-TB* isolate with the pathogenic *M. bovis* isolate (Table 3).

**Insert table 3**

The total activity value of an extract from a plant indicates which species could be the candidates for organic production of active extracts for use by communities because it takes into account not only the MIC of the extract but also the yield of the extract. By dividing the extraction yield in mg/g by the MIC in mg/ml the total activity is calculated in ml/g (Eloff, 2000). This provides the volume in ml that the extract from 1 g of dry material can be diluted and still kills the pathogen. *Searsia undulata* had the lowest MIC values (significant activity) with the highest % yield and highest total activity of 5 357 mL/g on *M. fortuitum* and 2885 mL/g on MDR-TB followed by *Protorhus longifolia* showing the highest total activity of 3557 mL/g on *M. smegmatis* and 2264 mL/g on MDR-TB. The results of total activity using rapidly growing mycobacteria are not shown due to limitation on number of tables. These two plants are the best candidates for further investigation due to their low cytotoxicity and more than 30 antimycobacterial compounds observed using bioautography. In addition, (Mosa *et al.*, 2014)
reported that *Protorhus longifolia* had antibacterial activity towards resistant clinical isolates of
Gram negative (*Pseudomonas aeruginosa*) and Gram positive (*Staphylococcus aureus*) whereas
*Searsia undulata* has been used to treat chest cold (Hutchings *et al.*, 1996). Specific
antimicrobials targeting Gram negative and Gram positive bacteria are useful but this does not
eliminate the need for broader spectrum antimicrobials. However, other plants with moderate
activity are worthy of investigation as studies have shown that there is synergy among different
compounds within an extract and once fractionated, different fractions/compounds might be
more active than the crude extract (Awouafack *et al.*, 2013).
3.3 Bioautography and thin layer chromatography analysis

Although all selected plant species had significant to moderate activity against all *Mycobacterium* species tested, plants showing at least MIC below 0.1 mg/mL or close to 0.1 mg/mL for any of the mycobacteria tested were selected to be investigated by bioautography. Five plants were selected namely *Heeria argentea*, *Lannea discolor*, *Protorhus longifolia*, *Searsia undulata* and *Sclerocarya birrea*. Only three plant species, *Harpephyllum caffrum*, *Protorhus longifolia* and *Searsia undulata*, had one or more clear zones of inhibition after spraying with *M. aurum* or *M. fortuitum*. *Searsia undulata* was the only species to show clear zones of inhibition with both *M. aurum* and *M. fortuitum* with more compounds visible with *M. aurum*. Using bioautography, *M. aurum* was the best indicator of antimycobacterial activity followed by *M. fortuitum*. Low MIC value might not always correlate with visualization of clear zones of inhibition due to the nature of separated compounds on the TLC plate, which may be volatile or experience loss of activity due to the separation of compounds that were acting in synergy in the crude extract. The crude extract of *Searsia undulata* had several active compounds against *M. aurum* – potentially more than ten compounds as some compounds were very close to each other in terms of Rf values ranging from 0.05 to 0.98 in the different solvent systems BEA, CEF and EMW (results not shown). Based on the MIC values and bioautogram patterns, the same plant extract have shown different activities towards different mycobacteria species. *M. smegmatis* also had some low MIC values with few or no visible zones of inhibition. The BEA and CEF solvent systems gave the best separation of compounds at this initial stage (Figure 1).

*Insert Figure 1*
3.4 Cytotoxicity assay

Most drugs or other substances are metabolised in the liver and excreted by the kidney. Mycobacterial species causing tuberculosis multiply and hide in macrophages. Therefore, three cell lines were selected to represent these organs namely Vero monkey kidney, human liver hepatoma (C3A) and mouse macrophage cells (RAW 264.7). Cancer cells are abnormal cells, and substances showing cytotoxicity to C3A cells and low cytotoxicity to the Vero cells may also be good candidates for further investigation as anticancer preparations. Cell-based in vitro toxicity assays are performed at an early stage of the drug development process in order to remove high-risk materials. Plants extracts showing sensitivity to cell lines with LC$_{50}$ values >0.1 mg/mL are considered not cytotoxic in terms of searching for anticancer compounds (Kuete, 2010). Crude extracts (Heeria argentea, Lannea discolor, Protorhus longifolia, Searsia undulata and Sclerocarya birrea) with good antimycobacterial activity (from significant to moderate) and those that showed visible zones of inhibition on bioautograms were selected for cytotoxicity testing. All crude extracts tested showed low toxicity against all three cell lines (C3A, Vero cells and RAW (264.7) except Searsia undulata that had moderate toxicity to C3A cells with LC$_{50}$ of 0.034 mg/mL. This deserves further investigation as source of anticancer substance as C3A cells are cancer cells with abnormal metabolic activity. The $S$. undulata extract had LC$_{50}$ of 0.50 and 0.12 mg/mL on Vero cells and RAW cells respectively with good selectivity indexes of 7.08 on Vero cells and more than 1 on RAW cells for non-tuberculous and MDR-TB mycobacteria. Protorhus longifolia had an LC$_{50}$ of 0.62 mg/mL on C3A cells, 0.88 mg/mL on Vero cells and >1 mg/mL for RAW cells with highest selectivity index on Vero cells of 12.6 for $M$. smegmatis followed by 8.02 for $M$. aurum, $M$. fortuitum and MDR-TB. Plant extracts with SI values less than1 mean that the extracts are relatively less toxic to the bacteria.
and more toxic to the mammalian cells (Eloff, 2000). Therefore, extracts with SI >1 may be relatively safer to use in vivo (not accounting for pharmacokinetic parameters) as they are less toxic to mammalian cells but more toxic to the pathogens. The selectivity indexes, especially of *P. longifolia* with SI = 12.6 could be considered as very promising as a good therapeutic index for a remedy or drug should be ≥10 (Caamal-Fuentes *et al.*, 2011). It is also worth noting that efficacy “in vitro” might differ from efficacy ‘in vivo” due to different parameters influencing pharmacodynamics and pharmacokinetic of drugs administered by different routes to humans and animals. In this study, *Protorhus longifolia* and *Searsia undulata* had the highest selectivity index against the three rapidly growing mycobacteria and pathogenic mycobacteria and the highest total activity. These rapidly growing species had the highest total activity of 5357 mL/g for *S. undulata* and 3557 mL/g for *P. longifolia* which indicates the volume to which the extract from 1 g can be diluted and still inhibit the growth of mycobacteria. It is also worth noting that the leaves of *S. undulata* are chewed by Khoi people to treat chest cold, indicating a low toxicity. The mode of administration and interaction with other factors in” vivo” must be considered when assessing toxicity.

**Insert Table 4**

**4. Conclusion**

This is the first report on antimycobacterial activity of these tree species from the Anacardiaceae family in Africa. Acetone leaf crude extracts of all the selected plants have shown significant to moderate antimycobacterial activity. *Harpephyllum caffrum, Lannea discolor, Protorhus longifolia, Searsia undulata* and *Sclerocarya birrea* susbsp. *caffra* had good antimycobacterial activity and low cytotoxicity. Some rapidly growing mycobacteria may be used as indicators for detecting good antimycobacterial activity of plant extracts against pathogenic mycobacteria
based on the correlation of MIC values. The best indicators of antimycobacterial activity on
MIC and bioautography were *M. aurum* followed by *M. fortuitum*. *P. longifolia* and *S. undulata*
had good antimycobacterial activity against all mycobacterial species. Low cytotoxicity was
observed on Vero cells and RAW cells. *Searsia undulata* was cytotoxic to hepatoma cells and
could be investigated further as a source of potential anticancer substances. Adding to results
obtained from the bioautography, MIC values and cytotoxicity as well as ethnobotanical use, *P.
longifolia* and *S. undulata* had good potential antimycobacterial activity worthy of further
investigation. The ethnobotanical use of chewing *S. undulata* leaves by Khoi people to treat
chest colds is interesting and the antimicrobial activity of leaf extracts of this species supports
this use although further study is necessary to validate this supposition. Isolation and
characterization of compounds from these highly promising species are underway.

**Acknowledgments**

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Africa (UNISA) Reference number: CAES/087 and the Department of Agriculture, Forestry and
Fisheries under section 20. UNISA and National Research Foundation (NRF grant No: 66141
and 86458) provided funding to support this study.

**Conflict of interest**

The authors have declared no conflict of interest.


Eloff JN. 1998a. Which extractant should be used for the screening and isolation of antimicrobial components from plants? *J Ethnopharmacol* 60: 1–8.


Kotzé M, Eloff JN. 2002. Extraction of antibacterial compounds from *Combretum*


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<th>Voucher number</th>
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<td>PRU2583</td>
<td>Bark, leaves and roots</td>
<td>Purifying blood, skin problems scarifications around sprains and fractures and leg paralysis due to poison</td>
<td>Polyphenolic and flavonoids/anticonvulsant</td>
<td>Hutchings et al., 1996; Van Wyk et al., 1997</td>
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<td>Stem Bark</td>
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*: not reported, Eng: English, Afr: Afrikaans, Z: Zulu, Tsw: Tswana
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<td>CIP (µg/ml)</td>
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<td>0.013 ± 0.006</td>
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<td>0.013 ± 0.006</td>
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<td>Strep (µg/ml)</td>
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<td>&gt;100</td>
<td>0.33 ± 0.11</td>
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### Table 3 Pearson's correlation coefficient (r) between MIC values of tested Mycobacteria

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<th>Mf</th>
<th>Ms</th>
<th>H37Ra</th>
<th>Mb</th>
<th>MDR-TB</th>
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Ma, Mycobacterium aurum; Mf, Mycobacterium fortuitum; Ms, Mycobacterium smegmatis; H37Ra, Mycobacterium tuberculosis ATCC strain 25177; Mb, Mycobacterium bovis; MDR-TB, Multidrug resistant Mycobacterium tuberculosis.

The bolded values represent the correlation between rapidly growing Mycobacteria and pathogenic Mycobacteria.
Table 4 Cytotoxicity (LC<sub>50</sub> in mg/mL) of extracts and selectivity index against C3A liver cells, Vero kidney cells and RAW 264.7 murine macrophage cells

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<th>SI Vero cells</th>
<th>SI RAW cells</th>
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<td>MF</td>
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<td>0.488</td>
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<td>Sclerocarya birrea</td>
<td>0.169</td>
<td>0.326</td>
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</table>
Figure 1

PL (A)              PL (B)                   HC (A)                  HC (B)                SU (A)                 SU (B)

PL: *Protorhus longifolia*, HC: *Harpephyllum caffrum*, SU: *Searsia undulata*

Figure 4.3  Bioautogram (A) of the three plants extracts with low MIC values and thin layer chromatography plates eluted in chloroform/ethyl acetate/formic (CEF) solvent system sprayed with vanillin sulphuric acid (B) showing varied chemical constituents. Bioautogram of the screened plants against *Mycobacterium aurum*