Thevetia peruviana (Pers.) K. Schum. Potential Antifungal Agent Against Mycosphaerella fijiensis Morelet, Fungi Responsible of Black Leaf Streak Disease (BLSD) of Plantain (Musa spp)

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Abstract

Alternatives to synthetic chemicals are undertaken against phytopathogens. The aim of this work is to evaluate the effect of seed extracts of *Thevetia peruviana* (Pers.) K. Schum. on *Mycosphaerella fijiensis* Morelet, fungus responsible for banana black leaf streak disease. Five extracts of *T. peruviana*, hexane extract (HE), ethyl acetate extract (EAE), acetone extract (AcE), methanol extract (ME) and aqueous extract (AqE), and a fungicide, Azoxystrobin were used. GC-MS of acetone extract was performed. Fifty (50) strains of *M. fijiensis* per sampling site were tested. Three concentrations of extracts 6.25 (C1), 12.5 (C2), and 25 (C3) μl/ml, a negative control (0 μl/ml) and 10 ppm of azoxystrobin were used for the tests. The MIC50 and MIC90 were determined. GC-MS showed chemical compounds with different molecular height such as acids, sugars, and esters. AcE and AqE significantly reduced *M. fijiensis* germ tube growth at C2 and C3 concentrations and with inhibition percentage respectively ranged of 60-90% and 40-80%. The growth levels of the germ tubes were above the strobilurin resistance threshold at Njombe and peasant plantation, ranging from 77.9% to 92.3%. AcE showed the same or superior efficacy as the fungicide used on conidial germination at all tested
concentrations. The MIC$_{50}$ totally reducing mycelial growth and conidial germination was 6.25 μl/ml. *T. peruviana* seeds extracts can be exploited in integrated pests management against *M. fijiensis*.

**Keywords:** *Thevetia peruviana*, *Mycosphaerella fijiensis*, extracts, GC-MS, Inhibition, conidia, germ tube

1. Introduction

Fungi are responsible for almost 60% of crop diseases (Lepoivre, 2003). Their fructifications are the source of primary inoculum in farms. In Cameroon, *Mycosphaerella fijiensis* Morelet, a fungus belonging to Ascomycetes is responsible for the black leaf streak disease (BLSD) of plantain and sweet banana (*Musa* sp). Indeed, banana, which is ranked fourth among agricultural products after rice, wheat and maize, is the most popular fruit on the planet (Lescot, 2006; Lassoudière 2010). This very important sector for Cameroon (first African banana producer), contributes approximately 7% of primary GDP (Gross Domestic Product) and is the second largest national employer after the government and the second largest source of income after timber (Mouliom et al., 1997; Ngando et al., 2006). However, in most banana production areas, black leaf streak disease is the greatest threat (De Lapeyre et al., 2010). This disease caused by *M. fijiensis* affects the photosynthesis of bananas through partial or total drying of the foliar system of the plant (Mouriçhon, 2003). This results in yield losses of up to 100% (Hermento et al., 2010), the reduction of the green lifespan of fruits, making their transport and conservation problematic (Churchill, 2011).

During dry periods, conidial infection may be a very beneficial form of survival for *M. fijiensis*, with the understanding that ascospore infestation is less during these periods (Jacome and Schuh, 1992). Despite much less conidial production, they can cause disease as effectively as ascospores (Stover, 1980; Fouré and Moreau, 1992; Jones 2009; Ngando et al., 2015).

More than 30% of the production of banana devoted for export are to the control of losses due to BLSD over the last 25 years which ranks *M. fijiensis* first among major agricultural pathogens (Abadie et al., 1999; Fullerton, 1994; De Lapeyre et al., 2009; Churchill, 2011).

Several control methods are used against this phytopathogen to reduce its effects on crops. The control of these parasites is accomplished only at the cost of frequent phytosanitary interventions. Farming practices that aim to reduce inoculum potential in the field by eliminating necrotic leaves and turning them upside down against the soil (Abadie et al., 1999; Mouriçhon, 2003). The development of resistant plants against this disease. However, this method is considered expensive and very long for farmers. In addition, fungi populations with a high level of genetic variability are difficult to control, and they can resist any control measure (El Hadrami, 2000). Some crops like bananas do not have resistant varieties, only chemical control methods are available for the farmer (Marin et al., 2003). Chemical control based mainly on the use of synthetic pesticides is the most effective method. This method unfortunately has consequences on the environment, micro-fauna, micro-flora, and human health because of the massive and inappropriate use of these fungicides. In addition, it is
expensive, and it causes the emergence of resistant strains because of the misuse of these chemical pesticides. This phenomenon of resistance to systemic fungicides has become a crucial problem for banana plantations in Cameroon (Lepoivre, 2003; Ngando et al., 2006; El Guilli et al., 2009) and leads to the return to the use of contact fungicides. However, with contact fungicides, the number of sprays per week increase, which is more expensive.

Awareness of the environmental cost of these practices and consumers' fears of the danger that pesticide residues accumulated in plant and fish products may pose to human health are giving rise to a growing interest in other alternatives of control, more efficient and more environmentally friendly.

Among the alternatives, there is biological control which makes use of biological agents and their substances (antagonistic microorganisms) on one hand and the use of plant extracts on the other hand. In the context of integrated pest management, toxic and biodegradable molecules of plant and microbial origin are essential for reducing the harmful effects of plant parasites.

Many plants have been reported in the literature to have pesticidal properties against many plant parasites (Bautista-Banos, 2000; Kassi et al., 2014). The seeds, leaves, fruits and roots of yellow Oleander (Thevetia peruviana) are considered as potential sources of active biological compounds for insecticides (Reed et al., 1982; Ambang et al., 2005), rodenticides (Oji and Okafor, 2000), fungicides (Gata-Goncalves et al., 2003; Ambang et al., 2011; Ngoh Dooh et al., 2014b; Ngoh Dooh et al., 2015), virucides (Tewtrakul et al., 2002) and bactericides (Saxena and Jain, 1990). The general objective of this work is to evaluate the antifungal potential of extracts of Thevetia peruviana against M. fijiensis, agent responsible for black leaf streak disease.

2. Material and Methods

2.1 Chemical Material

The chemical material consisted of the active ingredient of Bankit, azoxystrobin, obtained from CARBAP.

2.2 Obtention of M. fijiensis Conidia

Fifty (50) banana plants, spatially representative of the identified plantation area of the Cavendish AAA variety (dessert banana) in the industrial zone and of the AAB (plantain) plantain subgroup in the peasant zone (Table I), showing typical symptoms of the disease Black streak disease were randomly selected from each zone's plantation. Leaf fragments with stage 2 and stage 3 lesions were collected, wrapped in plastics and transported to the laboratory. Sampling was done on young leaves of the plant not bearing fruit. Hooks were used in areas where banana leaves were very high (Ngando et al., 2006, Nguepjop, 2011, Ngando et al., 2015).
Table I: sampling sites (PHP: Upper Penja Plantation; CDC: Cameroon Development Corporation; Peasant: Mbome)

| Agro-ecological Region | Plantations | Sector |
|------------------------|-------------|--------|
| Littoral               | Peasant     | Mbome  |
|                        | PHP         | Njombe |
| South West             | CDC         | Mussaka|

2.3 Obtention of the Extracts

The fruits were harvested from different parts of Yaounde City and then crushed with a stone. The seeds obtained were dried at room temperature in the Phytopathology laboratory of the Department of Plant Biology of the University of Yaounde I for 3 to 4 weeks. The seeds were crushed using a hand-mill of brand "Victoria". The resulting powder was loaded into cartridges and mounted on the soxhlet. The various extraction solvents hexane (HE), ethyl acetate (EAE), acetone (AcE) and methanol (ME) were each put in turn 48h to 72h (Negrette et al., 1987). The product obtained was concentrated in a rotary evaporator at the evaporation temperature of the corresponding solvent to eliminate the latter. The extract obtained was stored in the refrigerator at 4 °C until use. After this process, the residue was stored and mounted in the apparatus with another solvent.

The aqueous extract (AqE) was obtained by maceration of the powder in sterile distilled water for at least 12 hours (Stoll, 1994). The powder was put into a muslin cloth and soaked in a volume of sterile distilled water necessary to obtain the concentration of the desired stock solution (36 g in 72 ml of water). The extraction yield of each extract was calculated.

Screening of all these extracts was done previously (Table 2).

Table 2. Occurrence of natural products of each respective extract (Ngoh Dooh et al. 2014a).

| Products             | Hexane | Ethyl acetate | Acetone | Methanol | Aqueous |
|----------------------|--------|---------------|---------|----------|---------|
| Essential oils       | T      | -             | +       | +        | +       |
| Saponifiable oils    | +      | +             | +       | +        | T       |
| Coumarines           | _      | +             | +       | +        | +       |
| Alkaloids            | _      | _             | _       | +        | +       |
| Sterols              | _      | +             | +       | +        | +++     |
| Terpenoids           | _      | _             | _       | T        | -       |
| Flavonoids           | _      | _             | _       | _        | _       |
| Anthraquinones       | _      | _             | _       | _        | +       |
| Catechic tannins     | _      | _             | _       | +        | _       |
| Gallic tannins       | _      | _             | _       | _        | _       |
| Saponins             | _      | _             | _       | _        | _       |
| Anthocyanes          | _      | _             | _       | +        | +       |
| Steroid glycosides   | _      | T             | _       | +        | +       |
| Triterpenoid         | _      | _             | T       | _        | _       |
| Glycosides           | _      | _             | T       | T        | +++     |
| Free sugars          | _      | _             | T       | T        | +++     |
| Phenols              | _      | _             | _       | T        | _       |

- Absence of the products, + presence, +++ abundant presence, T presence in traces.
2.4 Preparation of Concentration of Extracts

A stock solution of 500 μl/ml concentration was prepared by mixing 1ml of the extract, 0.25 ml of solvent and 0.75ml of distilled water giving an initial volume of 2 ml.

From the stock solution, the concentrations of 6.25, 12.5 and 25 μl/ml were obtained by taking 0.25 ml, 0.5 ml and 1 ml respectively from the stock solution and adding them to 19.75ml, 19.5ml and 19 ml of culture medium, water agar, i.e., a final volume of 20 ml. (Gata-Gonçalves, 2001). The concentrations of the aqueous extract, 6.25, 12.5 and 25 mg/ml, were obtained by the same method from a stock solution of 500 mg/ml.

2.5 Dilution of Azoxystrobin

A positive control made of the active ingredient of bankit, a fungicide commonly used against leaf streak disease, azoxystrobin, was performed. The required dose is that recommended by the Fungicide Resistance Action Committee (FRAC), International Group of National Associations of Agrochemical Manufacturers for laboratory tests, or 10 ppm (Essis et al., 2010; Ngueujop, 2011; Ngando, et al., 2015). A 10000-ppm stock solution was prepared by mixing 50mg of active ingredient in 5ml of methanol (MeOH) in a tube and all was homogenized in vortex (VMR). Then 100 μl was taken from the stock solution and added to 99.9 ml of water agar medium.

An antibiotic, Chloramphenicol (200mg/l), was added in the medium to prevent bacterial contamination of water agar medium.

The volumes obtained were poured aseptically into Petri dishes of 90mm of diameter under a laminar flow hood the day before the different tests.

2.6 Assessment of the Effects of the Extracts on the Growth of the Germ Tube of M. Fijensis

A well-insulated lesion, susceptible of bearing conidia on its surface is cut with a scalpel from each piece of leaf and following its borders to avoid taking several lesions at the same time. The lower surface of the lesion was applied in the media containing the different treatments so as to leave its footprint. Each lesion was first applied to a test control medium to ensure the effectiveness of the sporulation and then in the treatments. One lesion served as a source of inoculum for the different treatments and for a concentration in the context of this monitoring with the conidia method for each site. The Petri dishes were incubated for 48 hours in the culture room at 25 °C and in continuous light.

After 48 hours, observations were made on 50 strains of M. fijensis for both cultures. The marks of the lesions in the quadrille and numbered Petri dishes were easily visible under the microscope. Readings were made following the number of the lesion using a micrometer (Mouliom et al., 1997; Essis et al., 2010; Nguepjop, 2011).

The first measured criterion is the length (μm) of the inhibited germ tubes which reflects the action of the extracts and fungicide on medium. The measurement of the length of the germ tube was done from the last septum or from the insertion base of the conidial germ tube to its longest end. Three conidia were measured by repetition and the average was selected.
For each site, and for each concentration, an average length of the germ tubes of 50 conidia on control (Lc) and on medium amended with extracts or fungicide (Lf) were calculated. The percentage inhibition of germ tube growth (Ia) was calculated using the formula:

\[ Ia = \frac{Lc - Lf}{Lc} \times 100 \]

(Ngando et al., 2006; Essis et al., 2010)

The growth rate (GT) was subsequently evaluated according to the formula \( GR = 100 - Ia \), to assess the resistance threshold.

The laboratory threshold for reporting resistance was set for Strobilurins (Bankit) at 75% of control (25% sensitivity) according to FRAC recommendations (Brent and Hollomon, 1998; Knight et al., 2002; Essis et al., 2010).

2.7 Assessment of the Effects of Extracts on the Germination of Conidia

The different concentrations that were used in the growth test were maintained.

The Petri dishes were incubated in continuous light for 48 h. About 25 to 30 conidia were counted, some lesions did not ‘spit’ a lot of conidia. The laboratory threshold for declaring resistance was set at 80%, i.e., 20% of the sensitivity (Du Pont, 1983; Smith et al., 1991; Essis et al., 2010).

A spore was considered germinated if the length of the germ tube was equal to or greater than the diameter of the spore.

The percentages of inhibition were then evaluated according to the following formula:

\[ PI = \frac{(A-B)}{A} \times 100 \]

(Leroux et al., 1978).

Avec:  
\( PI = \) percentage of inhibition; \( A = \) number of spores of germinated in the control medium; \( B = \) number of spores germinated in the presence of the extract or fungicide.

2.8 Acetone Extract Analysis (GC-MS)

The acetone extract which exhibiting the highest percentage of inhibition against *M. fijiensis* was analysed by capillary gas chromatography followed by mass spectrometry (GC–MS), using an Autosystem XL gas chromatograph (Agilent GC 7890A) with a vaporisation injector in split mode (1:50) interfaced to a Turbomass Perkin-Elmer mass-spectrometer (Agilent 5975 C TAD VL MSD). The analytical parameters were helium as carrier gas with the column flow rate of 1.21 ml/min. The oven temperature program was 40 °C for 3 min, then increased at 5 °C/min to 180 °C, followed by 15 °C/min to 240 °C and finally to 300 °C at 10 °C/min fractions (isothermal 15 min). A fused-silica capillary column, 30x25 mm i.d.30x32 mm (DB-1; 100% di- 150 99. methylpolysiloxane) was used. The ion source and transfer line temperatures were maintained at 200 and 280 °C, respectively. Electron ionisation mass spectra in the range 40–500 Da were recorded at 70 eV electron energy. The scan time was 1 ms, the multiplier potential 430 V and the source pressure 10 Torr. A computer recorded all data and compounds were identified by comparison with the Wiley’s and Nist libraries spectral data bank. The fraction previously evaporated and re-suspended in dichloromethane.
was analysed twice (1 ml; “hot-needle”) and, for semi-quantitative purposes, the average percentage composition was computed from peak areas normalised without using correction factors.

2.9. Evaluation of Minimum Inhibitory Concentrations (MIC₅₀ and MIC₉₀)

From the linear regression equation between the Neperian logarithms of the abscissa concentrations and the ordinate inhibition percentages, the concentrations reducing growth by 50% (MIC₅₀) and 90% (MIC₉₀) were determined (Dohou et al., 2004).

Only data from germ tube growth were used to calculate MICs, as recommended by the FRAC.

2.10 Statistical Analysis

Statistical analysis of the in vitro observation data and study of correlations was done using the SPSS 16.0 software. The Principal component analyzes (PCA) were performed using the XLSTAT 2007.8.04 software to classify the different extracts with respect to azoxystrobin. The Student Newman Keuls and Duncan tests at the 5% threshold allowed the comparison of the different averages when the differences were significant.

3. Results

3.1 Acetone Extract Chromatographic and Mains Constituents Found

More than 10 peaks were observed in the chromatographic profil of acetone extract (Figure 1). Peak retention time varied from 2.71 to 7.19 mn and peak height ranged from 172006 to 1579559.
Figure 1. GC-MS Chromatographic profile of acetone extract

Many compounds with different molecular weight were obtained from each peak (Table 2). Acid compounds such as decanoid acid, tridecanoid acid and pentadecanoid acid were revealed. Sugar such as maltose, Ethyl beta- d-riboside were obtained. Major compounds were acids (Table 2). Some compounds were not identified.
Table 2. Composition of the constituents found in the acetone extract after GC–MS

| Peak (min) | R.T  | Peak height | Molecular weight | Hypothetical compound name                                      |
|-----------|------|-------------|------------------|-----------------------------------------------------------------|
| 2.71      | 201922 | 88          |                   | 3-Hydroxyphenylacetylene                                         |
|           |       | 175         |                   | 1-Butanamine, N-methyl-N-nitro-                                |
|           |       | 117         |                   | 2-Butenediolyldichloride                                        |
| 3.05      | 254489 | 59          |                   | Ethyl beta.-d-riboside                                          |
|           |       | 56          |                   | Thietane, 2,4-dimethyl-beta.-D-Ribopyranoside, methyl           |
| 3.4       | 172006 | 59          |                   | alpha.-D-Glucopyranoside, methyl                                |
|           |       | 72          |                   | Maltose                                                         |
|           |       | 56          |                   | n-Decanoic acid                                                 |
| 3.78      | 3889785 | 148         |                   | 1,2-Benzenedicarboxylic acid, bi...                             |
|           |       | 56          |                   | 2 Phthalic acid, isobutyl octyl ester                          |
|           |       | 103         |                   | Dibutyl phthalate                                               |
| 4.02      | 3219504 | 72          |                   | Tridecanoic acid                                                |
|           |       | 59          |                   | Pentadecanoic acid                                              |
|           |       | 54          |                   | cis-13-Octadecenoic acid, methyl                                |
| 4.34      | 509524 | 54          |                   | trans-13-Octadecenoic acid, methyl                              |
|           |       | 66          |                   | 13-Octadecenoic acid, methyl ester                             |
|           |       | 68          |                   | cis-13-Octadecenoic acid                                        |
| 4.53      | 6934187 | 54          |                   | 2 cis-Vaccenic acid                                             |
|           |       | 68          |                   | trans-13-Octadecenoic acid                                     |
|           |       | 68          |                   | cis-13-Octadecenoic acid                                        |
| 4.56      | 1963294 | 72          |                   | Octadecanoic acid                                              |
| 5.68      | 457213 | 97          |                   | Glycerol 1-palmitate                                            |
|           |       | 74          |                   | Hexadecanoic acid, 2-hydroxy-1-Undecanedioic acid               |
|           |       | 83          |                   |                                                                |
| 5.86      | 1238475 | 148         |                   | Bis(2-ethylhexyl) phthalate                                    |
|           |       | 166         |                   | Phthalic acid, 2-ethylhexyl isoh...                             |
|           |       | 169         |                   | Phthalic acid, cyclohexyl 2-pent                                |
| 6.32      | 1579559 | 54          |                   | Oleic acid, (2,2-dimethyl-1,3-dioxolan-4-yl) methyl ester       |
|           |       | 128         |                   | 2,3-Difluoroaniline Hydrazinecarboxylic acid, (2-ethoxy-1-methyl-2-oxoethylidene), ethyl ester |
|           |       | 100         |                   |                                                                |
| 6.42      | 310154 | 55          |                   | None identify                                                   |
|           |       | 66          |                   | 9,12-Octadecadienal                                            |
|           |       | 89          |                   | None identify                                                   |
| 6.63      | 4913153 | 67          |                   | 9,17-Octadecadienol, (Z)-                                       |
|           |       | 80          |                   | 9,12-Octadecadienol                                            |
|           |       | 94          |                   | 9,12-Octadecadienoyl chloride                                   |
| 6.65      | 381820 | 67          |                   | 9,17-Octadecadienol, (Z)                                       |
|           |       | 80          |                   | 9,12-Octadecadienol                                            |
|           |       | 68          |                   | cis-13-Octadecenoic acid                                        |
| 7.19      | 1083443 | 67          |                   | 9,17-Octadecadienol, (Z)                                       |
|           |       | 109         |                   | 9,12-Octadecadien-1-ol, (Z,Z)                                  |
|           |       | 81          |                   | 7,10-Hexadecadienoic acid, methyl ester                        |
3.2 Effects of Extracts on the Growth of Germ Tube of M. fijiensis Strains

3.2.1 The Effect of the HE Extract on the Growth of the Germ Tube of M. fijiensis Strains

The HE extract had a very small effect on germ tube growth. No significant difference (P>0.05) were observed between the different concentrations applied and in any of the three study. The highest inhibition percentages recorded were 18.8, 15.0 and 11.5%, respectively at Njombe (PHP), Mussaka and Mbome at the highest concentration C3. The lowest inhibitions were obtained in the peasant plantation (1.1 and 1.2% at C1 and C2 concentrations, respectively. Inhibition of germ tubes by the bankit was highest than that of EH extract. Highest percentage inhibitions recorded were 50.4, 86.3 and 93.4% respectively in Mussaka, PHP, peasant zone ((Figure 2).

Figure 2. Effect of HE on germ tube growth of M. fijiensis strains

For each site the assigned values of the same letter do not differ significantly according to the Duncan test at the 5% threshold

T- : 0µl/ml; C1: 6,25µl/ml; C2: 12,5 µl/ml; C3 : 25µl/ml ; T+: 10 ppm

3.2.2. Effect of EAE on germ tube growth of M. fijiensis strains

The EAE extract was shown to be less effective in inhibiting the elongation of the germ tube of M. fijiensis conidia. An increase in percent inhibition was revealed as a function of concentration with this extract. Thus, at Mussaka 9.0, 32.8 and 54.6% inhibition, respectively of concentrations C1, C2 and C3 were obtained. The highest sensitivity of the conidia to this extract was found at the C3 dose with 48.8, 54.6 and 50.8% inhibition, respectively at Njombe.
(PHP), Mussaka and the peasant zone. At Mussaka, this extract was more effective (P < 0.05) than the bankit, at C₃ concentration (Fig. 6).

![Figure 6. Effect of EAE on germ tube growth of *M. fijensis* strains](image)

For each site, the assigned values of the same letter do not differ significantly according to the Duncan test at 5% threshold.

3.2.3 Effect of AcE on Germ Tube Growth of *M. fijensis* Strains

AcE was very effective in reducing the growth of the germ tube. Inhibition increased with concentrations in all areas. In the Njombe zone (PHP), 62.1, 84.2 and 95.8% inhibition of germ tube growth were obtained with C₁, C₂ and C₃ concentrations, respectively. In the peasant zone, inhibition percentages were 62.8, 91.8 and 97.8%, respectively, at the three concentrations. The AcE extract were more sensitive to germ tube growth compared to the bankit. At Mussaka, all concentrations showed highest inhibition than azoxystrobin (P<0.05). In Njombe and Mbome, only, C₂ and C₃ concentrations inhibited germ tube than the bankit. However, no significant difference was observed between the bankit concentration and the C₂ and C₃ concentrations of AcE at Mbome (P = 0.36), (Fig. 7).

All samples were very sensitive to AcE (Fig. 11c) as the lengths of the germ tube were very short.
Figure 7. Effect of EAc on germ tube growth of *M. fijiensis* strains

For each site the assigned values of the same letter do not differ significantly according to the Duncan test at the 5% threshold

![Bar chart showing effect of EAc on germ tube growth of M. fijiensis strains](chart)

T-: 0µl/ml; C1: 6.25µl/ml; C2: 12.5 µl/ml; C3: 25µl/ml; T+: 10 ppm

3.2.4 Effect of ME on Germ Tube Growth of *M. Fijensis* Strains

Strains from the peasant plantation showed resistance to the methanol extract. The inhibition percentages were 10.9, 8.2 and 25.1%, respectively at C1, C2 and C3 concentrations. A sensitivity to the extract was obtained at the C3 concentration with samples from the two industrial zones, 45.8% in Njombe (PHP) and 43.5% in Mussaka (Fig.8). No concentration matched the effectiveness of azoxystrobin in the areas studied. The lowest inhibition percentages were obtained with the samples from the peasant zone (P> 0.05).
Figure 8. Effect of ME on germ tube growth of *M. fijiensis* strains

For each site, the assigned values of the same letter do not differ significantly according to the Duncan test at the 5% threshold

- **T-**: 0µl/ml; **C1**: 6.25µl/ml; **C2**: 12.5 µl/ml; **C3**: 25µl/ml ; **T+**: 10 ppm

### 3.2.5 Effect of AqE on Germ Tube Growth of *M. fijiensis* Strains

The aqueous extract proved to be sensitive to the development of the germ tube length at all concentrations., the germ tubes growth was reduced compared to the control (Fig. 11b). In the industrial plantation of PHP at Njombe site, the inhibition rates of 40.0, 47.5 and 55.8% were obtained at different concentration, respectively. The sensitivity was very high with the samples from peasant plantation, the percentages of inhibition were 81.4 and 75% at C3 and C1 concentrations. At Mussaka, strains were sensitive at C2 and C3 concentrations than Bankit (P <0.05), the percentages of inhibition were 66.9 and 77.6% against 50.4% for azoxystrobin. No significant difference (P = 0.114) was obtained at Mussaka between C1 and azoxystrobin concentrations (Fig 9).
Figure 9. Effect of AqE on germ tube growth of *M. fijiensis* strains

For each site the assigned values of the same letter do not differ significantly according to the Duncan test at the 5% threshold.

T-: 0 mg/ml; C1: 6.25 mg/ml; C2: 12.5 mg/ml; C3: 25 mg/ml; T+: 10 ppm

The probability correlation circle of 59.78% revealed four distinct groups. A group consisting of HE whose inhibition on germ tube growth was very low. The second group containing EAE, AcE and AqE, which showed an efficiency on germ tube growth. A third group consisting of ME which showed a moderate efficiency on the inhibition of germ tube growth of *M. fijiensis* and finally a group consisting of positive control T + (Fig. 10).
Figure 10. Mapping of extracts and fungicide effective on inhibition of germ tube growth of *M. fijiensis* conidia

Figure 11. Measurement of the germ tube in the various media supplemented with extracts at the C₃ dose: normal conidia on media, control (a). Conidia very sensitive on media with AqE (b) and AcE (c). 1 = germ tube; 2 = conidia

### 3.3 Effect of Different Extracts and Azoxystrobin on the Germination of *M. fijiensis* Conidia

The HE had no effect on the germination of the conidia of *M. fijiensis*. The percentages of inhibition ranged 0-0.5%, 0.2-0.6% and 0.5-8.6% respectively with samples of PHP, Mussaka and peasant plantation. No significant difference was obtained between treatment (P > 0.05%). The lowest value of the percentage inhibition of germination with the bankit was
obtained in Mussaka, 38.6% against 99.5% in the peasant zone. The germination rates obtained with this extract were largely above the resistance threshold of 25% Strobilurins (Table 3).

The EAE extract was less effective on the germination of *M. fijiensis* conidia at all concentrations. With the samples from PHP, the percentages of inhibition obtained were 2.5, 2.5 and 1%, respectively with the doses C₁, C₂ and C₃. At Mussaka the percentages were ranged 0.2, 2.5 and 3.2%. All inhibition percentages obtained with this extract were lower than those obtained with azoxystrobin (Table 3).

The AcE proved to be very effective against the germination of *M. fijiensis* conidia. In industrial plantations, only the concentration of 6.25 μl/ml gave low inhibition percentages, 8.1% and 38.5% in Njombe and Mussaka, respectively. The C₂ and C₃ concentrations were the most effective in these industrial zones, with 64.1% and 99% inhibition respectively at Njombe (PHP) against 85.4% and 96% at Mussaka. The strains of these industrial zones showed a slight resistance to this extract compared to the samples from the peasant plantation where the percentages of inhibition were 54.8, 97.8 and 100% respectively at C₁, C₂ and C₃ concentrations. In Njombe (PHP), the C₃ dose induced 99% inhibition rate while the bankirrate was 81.8% (P <0.05). The same result was obtained at Mussaka, the inhibition percentages obtained were significantly higher and significantly different from those obtained with the bankit (Table 3). However, in the peasant plantation, no significant difference was between concentration of extracts and bankit (P = 0.335). Total inhibition was obtained with samples of peasant zone in the highest concentration, no conidial.

The ME extract was no sensitive to conidial germination. With the samples from the industrial plantation of Njombe (PHP), the inhibition percentages obtained were 17.8, 11.6 and 34.8%, respectively at the three concentrations. In Mussaka the percentages were 0, 0.6 and 34.6% and 13.4, 7.5 and 9.1% in the peasant plantation respectively at C₁, C₂ and C₃. Njombe (PHP) strains were slightly more sensitive to this extract than those of the other two sampling areas. In the three study areas, the bankit showed highest efficacy than this extract (P <0.05) (Table 3).

AqE was effective against germination of conidia from the peasant plantation and the Mussaka industrial area. At C₂ and C₃ there were 42.3 and 62% inhibition in Mussaka against 71.5 and 90% in the peasant plantation. Samples from the peasant zone were therefore the most sensitive to this extract. The synthetic fungicide induced an inhibition of conidial germination up to 99% in the peasant zone and 38.6% in Mussaka. The bankit was less effective than the C₃ concentration with samples from the Mussaka industrial plantation (P <0.05). Conidia from the industrial plantation of Njombe (PHP), showed resistance to AqE. The percentages of inhibition obtained were very low, 10.1, 16.2 and 17.7% respectively at doses C₁, C₂ and C₃ (Table 3).
For each site and each extract, the values assigned to the same letter do not differ significantly at the 5% threshold according to the Duncan test.

|      | PHP  | MUSSAKA | PEASANT |
|------|------|---------|---------|
| HE   |      |         |         |
| T    | 0.00±0 | 0.00±0  | 0.00±0  |
| C₁   | 0.5 a ± 0.3 | 0.2 a ± 0.1 | -0.1 a ± 0.0 |
| C₂   | 0.5 a ± 0.3 | 0.2 a ± 0.1 | -0.1 a ± 0.0 |
| C₃   | 0.5 a ± 0.3 | 0.2 a ± 0.1 | -0.1 a ± 0.0 |
| T+   | 8.5 b ± 9 | 38.6 b ± 7.9 | 99.5 c ± 0.3 |
| AE   |      |         |         |
| T    | 0.00±0 | 0.00±0  | 0.00±0  |
| C₁   | 2.5 a ± 2.2 | 0.2 a ± 0.1 | 7 a ± 1 |
| C₂   | 2.5 a ± 2.2 | 0.2 a ± 0.1 | 7 a ± 1 |
| C₃   | 2.5 a ± 2.2 | 0.2 a ± 0.1 | 7 a ± 1 |
| T+   | 8.5 b ± 9 | 38.6 b ± 7.9 | 99.5 c ± 0.3 |
| AcE  |      |         |         |
| T    | 0.00±0 | 0.00±0  | 0.00±0  |
| C₁   | 8.1 a ± 7.9 | 38.5 a ± 7.1 | 54.8 a ± 5.4 |
| C₂   | 64.1 b ± 4.7 | 85.4 b ± 3.1 | 97.8 b ± 0.8 |
| C₃   | 99 d ± 4.3 | 96 c ± 1.4 | 100 b ± 0.0 |
| T+   | 8.5 c ± 9 | 38.6 a ± 7.9 | 99.5 b ± 0.3 |
| ME   |      |         |         |
| T    | 0.00±0 | 0.00±0  | 0.00±0  |
| C₁   | 17.8 a ± 6.5 | 0 a ± 0.0 | 13.4 a ± 5.7 |
| C₂   | 11.6 a ± 8.3 | 0.6 a ± 0.5 | 7.5 a ± 3.2 |
| C₃   | 34.8 b ± 4.3 | 34.6 b ± 4.4 | 9.1 a ± 3.3 |
| T+   | 8.5 c ± 9 | 38.6 c ± 7.9 | 99.5 b ± 0.3 |
| AqE  |      |         |         |
| T    | 0.00±0 | 0.00±0  | 0.00±0  |
| C₁   | 10.1 a ± 3.5 | 9 a ± 4.3 | 80 b ± 4.3 |
| C₂   | 16.2 b ± 7.5 | 42.3 b ± 3.6 | 71.5 a ± 4.9 |
| C₃   | 17.7 b ± 7.5 | 62 c ± 4.6 | 90 c ± 2.3 |
| T+   | 8.5 c ± 9 | 38.6 b ± 7.9 | 99.5 c ± 0.3 |

The 56.28% probability threshold correlation circle revealed the existence of three distinct groups in the efficiency of the extracts in the inhibition of M. fijiensis conidia germination. A group consisting of HE which was ineffective, another group consisting of effective extracts, AcE, AqE and T+, and a last group for the extracts with low efficiency, ME and EAE (Fig. 12).
Figure 12. Mapping extracts and fungicide effective on the inhibition of conidial germination of *M. fijiensis*

4. Minimal Inhibitory Concentrations MIC$_{50}$ and MIC$_{90}$ of germ tube growth of *M. fijiensis* strains

The lowest MICs were obtained with AcE and AqE. MIC$_{50}$ ranges from 2.0 μl/ml to 3.3 μl/ml for AcE and from 6.0 mg/ml to 15.0 mg/ml for AqE. The highest MICs were obtained with HE. Thus, with EAE, the MIC$_{50}$ values varied from 20.1 to 30 μl/ml respectively for Mussaka and Njombe (PHP) and with the ME from 27.1 to 403.4 μl/ml. However, the MIC$_{50}$ and MIC$_{90}$ of AcE were substantially equal in all the zones (3.3, 2.0 and 3.3 μl/ml) which indicates the same behavior of the extract with respect to all strains regardless of the sampling site (Table 4).
Table 4. CMI<sub>50</sub> and CMI<sub>90</sub> of the growth of the germ tube of \textit{M. fijiensis} with the different extracts tested (µl/ml or mg/ml)

| Sites     | HE CMI<sub>50</sub> | HE CMI<sub>90</sub> | EAE CMI<sub>50</sub> | EAE CMI<sub>90</sub> | AcE CMI<sub>50</sub> | AcE CMI<sub>90</sub> | ME CMI<sub>50</sub> | ME CMI<sub>90</sub> | AqE CMI<sub>50</sub> | AqE CMI<sub>90</sub> |
|-----------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|
| PHP       |                      |                      | 30,0                 | 167,2                | 3,3                  | 14,7                 |                      |                      | 15,0                 | 525,1                |
| MUSSAKA   |                      |                      | 20,1                 | 73,4                 | 2,0                  | 18,0                 |                      |                      | 27,1                 | 74,7                 |
| PEASANT   |                      |                      | 5431 1,210<sup>6</sup> | 24,5                 | 97,7                 | 3,0                  | 18,0                 | 403,4                | 1066                 |                      |

*values no founded (negative correlations)

\section*{5. Discussion}

The present study was based on the extraction of natural substances from the seeds of \textit{T. peruviana} and on the evaluation of the antifungal potential of its extracts on \textit{M. fijiensis}, responsible for the disease of black streaks of banana.

The chromatographic of The AcE has showed many major compounds, its effectiveness could be due to the presence of molecules that act in high doses. Gata-Goncalves et al. (2003) also showed after GC-MS of extract of \textit{T. peruviana}, presence of many compounds

Plant extracts have already shown their effectiveness against the germination of fungal spores. This is the case with the results of Achraf et al. (2012) which found out that the aqueous extract of \textit{Asphodelus tenuifolius} and \textit{Zygophyllum album} were effective in inhibiting spore germination of \textit{Penicillum expansum} with a percentage inhibition of 95.48% and 93.82%. The ethanol extracts of the leaves of \textit{O. gratissimum} and \textit{Aframomum melegueta} prevent the spores of \textit{Fusarium oxysporum} and \textit{Aspergillus niger} from germinating by more than 65%, according to the findings of Okigbo & Ogbonnaya (2006).

The AcE greatly reduced the germ tube elongation of \textit{M. fijiensis} strains, followed by EAq. The AcE was very effective in inhibiting the germination of \textit{M. fijiensis} strains in all banana growing areas studied, being more effective than the chemical fungicide azoxystrobin used. It inhibited more than 90% conidial germination. These results are similar to those of Arciniegas, (2002) and Arciniegas et al. (2002) who showed an \textit{in vitro} test a strong antifungal activity on both colony development and germination of \textit{M. fijiensis} conidia using crude ethanol extracts, amphipolar solvent as well as acetone extract of eight plants such as \textit{Commelina diffusa}, \textit{Momordica charantia}, \textit{Piper hispidum}, \textit{Piper peltatum}, \textit{Sida rhombifolia} and \textit{Syzyzygium aromaticum}.

Furthermore, Paola (2006) obtained a significant reduction in the incidence and severity of black leaf streak disease on plants infected artificially with \textit{M. fijiensis} conidia after being treated with extracts obtained from the maceration of \textit{Momordica charantia} and \textit{Senna reticulata} in water/alcohol mixture. Indeed, the phytochemical analysis performed on \textit{S. reticulata} with which they obtained the best results revealed the presence of a variety of...
secondary metabolites such as polyphenols, coumarins, saponins, triterpenes, flavonoids some of which are found in the extract with acetone (AcE) from *T. peruviana* (Gata-Goncalves et al., 2003; Ngoh Dooh et al., 2014). These extracts are known for their antifungal activity or resistance to the induction on *M. fijiensis* (Riveros et al., 2003; Polanco, 2004).

The growth and germination rates obtained with HE, EAE and ME were significantly higher (77.9% to 92.3%) at the reported Strobilurin resistance threshold which shows the resistance of the strains of these zones to these extracts. Only the C3 doses of EAE and EAq gave lower rates with Mussaka strains. The growth and germination rates obtained for the EAq at doses C1, C2 and C3, despite the effectiveness of this extract did not fall below the resistance threshold of Strobilurins. The growth and germination rates achieved with EAc were below the Strobilurin resistance level at C2 and C3 on growth and in all areas at C2 and C3 with Mbome and Mussaka strains and the C3 dose at the PHP, on the germination, which shows the sensitivity of the strains of these zones to this extract.

With the bankit, the growth level of the germ tube was below the strobilurin resistance threshold in two of the three study areas (Hermento et al., 2010). The lowest rate was obtained in the plantation, 7.6%, followed by the Njombe industrial plantation. An onset of resistance was obtained in Mussaka with azoxystrobin (49.6%).

The bankit whose active ingredient, azoxystrobin, was used in this study was found to be effective in some production areas compared to others. Azoxystrobin, which belongs to the Strobilurins class, is a potent inhibitor of cell respiration, hence its effectiveness against germ tube elongation and germination (both parameters being linked). This result corroborates that of Essis et al. (2010) who obtained an efficacy in the laboratory of azoxystrobin against the germ tube growth of *M. fijiensis* strains from Ivory Coast banana plantations. However, in Mussaka a resistance to this fungicide was detected. This could be explained by the abusive use of this compound which eventually developed resistant strains compared to the peasant zone where there is no use of the bankit. This result is contrary to that of Nguepjop (2011) which indicated a lack of resistant strains in this zone. Ngando et al. (2006) showed resistance phenomena in banana plantations in Cameroon. The low efficiency of azoxystrobin in the peasant plantation denotes the appearance of resistant strains, despite the distance with the industrial zones. This could be explained by the movement of men from one corner to another and who would carry conidia resistant strains to areas where they do not exist (peasant zone).

The MIC50 of the different extracts were determined with the different strains tested. Low variability was observed. The low values and close to the MICs obtained with the acetone extract demonstrate the efficacy and fungicidal properties of this extract in inhibiting the germination of the conidia of the fungus tested. These results are in agreement with those of Doumbouya et al. (2012) who showed that the low MIC values of *Ocimum graticum* extracts inhibit the development of phytopathogenic fungi.

The inhibition percentages obtained were different according to the extracts. This can be explain by the process of obtention the extracts. Okigbo (2005) showed that the levels of
bioactive compounds in plants with antifungal activity could be influenced by many factors, including the plant’s age, harvest time, extraction solvent, and method of extraction. Indeed, for the same species of fungi, an extract preceding or following another could show an effectiveness contrary to its predecessor or its successor. Thus, AcE was effective in inhibiting the germination of all strains studied, unlike methanol which showed no efficacy meanwhile these two solvents are polar. The extraction with acetone would have taken the active ingredient acting on the germination before the application of methanol.

The plant used in this study belongs to the Apocynaceae family. The extracts contain many chemical compounds. These compounds could act on the target organisms in several ways. They could inhibit growth by acting on metabolic functions such as cell division. Others would inhibit respiration by blocking ATP production or inducing plant resistance (Laurent et al., 2003; Lepoiivre 2003; Chwaleka et al., 2006).

6. Conclusion

The AcE and AqE were effective as azoxystrobin, the active ingredient in the bankit, a fungicide commonly used against growth tube length and germination of conidia of *M. fijiensis* responsible of black leaf streak disease. The GC-MS revealed many compounds with antifungal properties.

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