Evaluation of the Epidemiological Factors Responsible for the Appearance of Phytophthora Palmivora and Phytophthora Megakarya, Causal Agent of Black Pods Disease in the Department of Méagi, Southwestern Côte D’ivoire

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Abstract: The cocoa black pod disease is caused by a fungus Phytophthora sp, which comprises two main species (Phytophthora palmivora and Phytophthora megakarya) commonly found in Ivorian cocoa plots and which cause significant damage. Despite numerous studies, this disease continues to spread and even more so Phytophthora megakarya, which did not exist in Côte d'Ivoire, is beginning to spread significantly. This study focused on the evaluation of the epidemiological factors responsible for the progression of Phytophthora megakarya and Phytophthora palmivora in the Department of Méagi, southwestern Côte d’Ivoire. To this end, a prospective survey was conducted in three cocoa plots in Méagi, which collected epidemiological data on “Foliar coverage rate”, “Maintenance level” and the prevalence of black pod. Molecular data on black pod disease isolates were extracted from immature pods with black pods symptoms. A descriptive analysis was first applied to the data to understand their dispersion, followed by a comparative analysis (student’s t-test) to evaluate the relationship between epidemiological parameters and disease prevalence. Finally, a molecular analysis in the laboratory made it possible to characterize the different species of Phytophthora sp. The results of the molecular analysis showed the presence of Phytophthora megakarya at 79% against 21% for Phytophthora palmivora. The results of the student t-test showed that this prevalence of black pod is related to the lack of plot maintenance (P=0.036<0.05) and high leaf coverage of cocoa trees (P=0.003<0.05). However, distribution maps of the different species show that Phytophthora megakarya is the most dominant species on the Méagi site.

Keywords: Cocoa, Maintenance Level, Foliar Coverage, Phytophthora sp.

Introduction

Cocoa tree is a perennial plant originating from the humid tropical forests of America. It belongs to the Malvaceae family (Guyot, 1992), and to the genre Theobroma. It is cultivated for its beans used to make chocolate. Most of the production (70%) comes from Côte d'Ivoire and Ghana (Lass 2004). Cocoa farming in Côte d'Ivoire accounts for 40% of exports and 15% of gross domestic product (ICCO 2015). This crop employs more than 1,000,000 farmers and supports more than 3,000,000 people in the secondary and tertiary sectors of the Ivorian economy. Nevertheless, cocoa farming is subject to high parasitic pressure due to diseases and pests. One of the most important diseases affecting cocoa farms in Côte d’Ivoire is black pod (Kébé et al. 1996; Kébé 1999). Black pod is a fungal disease caused by the genre Phytophthora sp. Phytophthoras are Oomycetes of the family

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Peronosporaceae. Symptoms observed on cocoa trees are generally localized on the fruit (Fig1), on the trunk and sometimes on the leaves. Pods are sensitive at each stage of their development and can be infected at any location on all surfaces. The first symptoms begin with blacking, which later becomes a stain on the pod. This spot spreads rapidly either from top to bottom (Fig1a) or from bottom to top of the infected pod (Fig1b). Water is the main carrier of inoculum. It is under the action of water that sporocysts explode to produce zoospores that are carried by rainwater (Akrofi 2015). Other factors that spread the disease include insects, various animals including rodents and especially humans. Black Pod disease spreads faster when the humidity is high. Indeed, it is favoured by dense shade when the planting density is high or when there is a strong presence of shade trees in the plot with a very large grass cover or proximity to a watercourse (Coffee-Cocoa Council 2015). Two species of *Phytophthora* *sp* are found in cocoa farms in Côte d'Ivoire. These are *Phytophthora palmivora*, which is the most widespread and least aggressive species causing yield losses of around 40%, and *Phytophthora megakarya*, which is very aggressive and less known with damage estimated at nearly 60 to 100% (Flood 2006). In Côte d'Ivoire, black pod has long been attributed to *Phytophthora palmivora*, but in recent decades, *Phytophthora megakarya*, which did not exist in Côte d'Ivoire, has appeared in cocoa plantations (Kébé 1999). The presence of *Phytophthora megakarya* has been reported since 1998 in eastern Côte d'Ivoire, along the entire border with Ghana where significant crop losses have been recorded (CNRA 2007). Since 2006, the first isolates of *Phytophthora megakarya* have been identified in Abengourou and Méagui. This study was extended in 2007 to the Department of Soubré, which is the current cocoa loop. In this Department, two isolates of this *Phytophthora* species have been identified (CNRA 2007). The identification of *Phytophthora megakarya* was confirmed in 2013 in southeastern Côte d'Ivoire towards the border with Ghana where nearly 60 to 80% of crop losses were recorded (Pohe et al. 2013). At the same time, another study was conducted in southwestern Côte d'Ivoire to characterize *Phytophthora spp*, isolates, but the latter study revealed only the presence of the single *palmivora* species (Coulibaly et al. 2013). These results clearly show that the exact distribution of this new species is not well known throughout the national territory. The increasing yield losses observed by producers each year in southwestern Côte d'Ivoire due to black pod could be linked to *Phytophthora megakarya*. To remove the ambiguity, this study is initiated to attest not only to the presence of *Phytophthora megakarya* from molecular analyses, but also to evaluate the epidemiological factors responsible for the development of both *Phytophthora palmivora* and *Phytophthora megakarya* in the Department of Méagui. Specifically, it will first identify the different species of *Phytophthora sp*, then evaluate the epidemiological factors responsible for the appearance of each species of *Phytophthora sp* and finally establish the distribution map of each species of *Phytophthora sp*.

2. Materials and methods

2.1 Study area

This study was conducted in Méagui (5° 24′ 43″ North, 6° 33′ 37″ West), one of the Departmental capitals of the Nawa region, South West Côte d'Ivoire (Fig2). This region was chosen because it constitutes the current cocoa loop, with, for the 2013-2014 campaign, about 320,000 tons, or nearly 20% of national production (Coffee-Cocoa Council 2014). The Department of Méagui is a forest area whose vegetation is essentially dominated by dense forest with deep, permeable and well-drained soil capable of supporting all types of crops, especially cocoa (Ngo et al. 2012). The climate of the Nawa region is of a sub-equatorial type locally called "Attieën climate" (Yao 2009). This climate is characterized by heavy rainfall, which oscillates between 1300 mm and 1600 mm depending on the year and location. The atmospheric humidity is high (90%) with a low annual variation in temperature amplitude (between 26 and 28°C). The climate is characterized by a dry season (December-March) and two rainy seasons (April-June and September-November). These climatic characteristics, marked by high precipitation and a long period of relative humidity, make it a zone with high parasitic pressure.

2.2. Material

The plant material used in this study includes 66 cocoa trees (*Theobroma cacao*) from which 66 pods affected by black pod were collected. The survey took place in three peasant farms. To remove these pods affected by black pod, a pruning shear was used to cut the pod. Once the pods have been removed, they are packaged in newspaper to prevent the rot from spreading, then with adhesive tape, fasteners are made around the pod and labelling has been done with an indelible marker. The Global Positioning System (GPS) coordinates of the cocoa trees sampled were recorded with a Garmin 64s GPS. Survey sheets were completed to compile physical data. Once in the laboratory, the samples of pods affected by black pod were crosscut with a knife following the evolution of the rot and then a scalpel was used to remove small pieces of pod (explants) from the cortex after the crosscut. These decay explants are placed in http://www.ijSciences.com
2.3 Methodology

2.3.1. Prospective survey
This study was carried out through a prospective survey. These surveys were carried out in three peasant farms in Méa-guï. To this end, a survey sheet was prepared containing several parameters that were evaluated in the field. These were date, study area, study site, sample code, tree number, geographical coordinates (longitude in decimal degrees, and latitude in decimal degrees), total number of pods, number of rotten pods, level of maintenance, foliar coverage rate and description of symptoms observed on the pods collected.

2.3.2. Observation
Observations were randomly made on 66 cocoa trees with pods affected by black pod in three peasant plots, i.e. 22 cocoa trees observed per plot. For each tree, observations were made on the total number of pods, as well as the number of rotten or healthy pods from the trunk to the canopy of the test tree. The state of maintenance was also observed at the scale of each plot. The coverage rate was evaluated through the foliar crown of the cocoa tree.

2.3.3. Data collection
Data on epidemiological factors were collected, including the total number of pods, the number of rotten pods, the prevalence of black pod, the level of maintenance of cocoa plots and the cocoa tree coverage rate. Indeed, the total number of pods was obtained by counting all pods on each test cocoa tree. This variable includes both the number of healthy pods and the number of pods affected by black pod. Then, the number of rotten pods was deducted from the total number of pods observed on each test cocoa tree. Each test cocoa tree was the subject of a geographical coordinate survey. These geographical coordinates made it possible to establish the distribution map of the different species of Phytophthora sp.

The prevalence of black pod was determined from the ratio of the total number of pods to the number of pods affected by black pod using the following formula:

\[
P(\text{Pb}) = \frac{\text{Number of rotten pods (TCAB)}}{\text{Total number of pods produced (TCAB)}} \times 100
\]

P (Pb%): prevalence of black pod in percentage.

The prevalence of each Phytophthora sp species was also determined by taking into account the number of infected pods per Phytophthora sp species and the total number of pods produced.

The level of maintenance of the plots has been evaluated. This is a qualitative data with two modalities (maintained or not maintained). This variable relates to regular weeding, draining and pruning of the test plot. The coverage rate of cocoa trees is evaluated at the tree level. It is a qualitative variable with three modalities (low, medium and high). The high coverage rate is characterized by a strong canopy of the tree with a strong shade while the low coverage rate is characterized by a weak foliage of the tree often allowing light to penetrate the undergrowth. At the intersection of these two modalities is the average coverage rate. Molecular data were also collected. These molecular data relate to the black pod isolates collected from the rotten pod samples. Indeed, a sample of pods showing the beginning of rot was collected on each cocoa tree test stock. Once collected, the pod samples are packaged in newsprint to limit the progression of black pod disease. These samples are then numbered and labelled and transported to the laboratory. Once in the laboratory, each pod sample was crosscut to extract an explant from the cortex of the rotten pods following the evolutionary front of the rot. The explants collected are temporarily packaged in eppendorf tubes, containing an agar medium (H2O-Agar) favourable to the development of Phytophthora sp. These different tubes were then labelled according to the identification code since the field sampling. A total of 66 black pods isolate samples were sent to the BGPI laboratory of Cirad Montpellier for molecular analyses.

2.4. Data analysis

2.4.1. Molecular characterization of Phytophthora sp isolates
In the laboratory in Montpellier, the sampled explants were cultured in petri dishes on the "V8" medium, which consists of a 1/10th vegetable juice cocktail, agar at 15g/L and CaCO3 at 3g/L for 4 days in the dark, at 25°C. After 4 days, agar implants containing Phytophthora mycelium were removed from the fungal growth forehead with a scalpel and then transplanted back to the V8 medium under the same conditions described above for 7 days. The resulting mycelium is used for DNA (Deoxyribonucleic Acid) extraction. In the case where the strains are contaminated by other fungi, several transplants on H2O-Agar culture medium at 15g/L are necessary to purify the strain. After purification, the strain is cultured again on the "V8" medium at 25°C for 4 days and then for 7 days in the dark to obtain a mycelium typical of Phytophthora sp. The mycelium obtained after purification is used for DNA
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The identification of the different Phytophthora species was carried out using PCR (Polymerase Chain Reaction) with ITS (Internal Transcribed Spacer) primers specific to each species (P. megakarya and P. palmivora).

2.4.2. Distribution Mapping of Phytophthora sp. species
The results of the molecular analysis were coupled with the geographical coordinates of the cocoa trees from which the samples of rotten pods were taken. The coupling of these coordinates with molecular data made it possible to produce distribution maps of the Phytophthora sp species identified in the Department of Méagui.

2.4.3 Evaluation of epidemiological factors in the progression of black pod disease
The descriptive analysis of the epidemiological data allowed to determine on the one hand the minimum, maximum, average and standard deviation of the quantitative variable represented by the prevalence of black pod over the entire study site and the determination of the numbers and frequency of the qualitative variables relating to the foliar cover rate of cocoa trees and the level of maintenance of the plots surveyed on the other hand. The number and percentage of pods infected by each Phytophthora sp species was also determined. The data from the molecular analysis determined the frequency and distribution maps of each Phytophthora sp species identified at the Méagui site using IBM SPSS statistic.20 software. After checking normality using the Shapiro-Wilk test and variance homogeneity using the Levene test, the student t-test was applied. The student t-test first compared the averages of the "Not maintained" and "Maintained" modalities of the variable maintenance level (NE) of the plot, then the modalities "Medium" and "Strong" of the variable foliar cover rate (TCF) of cocoa trees compared to the prevalence of black pod in cocoa pods. The Chi-square test was used to check the level of dependence between the foliar cover rate and the frequency of Phytophthora sp species identified in Méagui. The Chi-square test was also applied to verify the dependence between the level of maintenance of the plot and the frequency of Phytophthora sp species identified at the Méagui site. Cramer’s Phi and V test was applied to check the magnitude of the link between foliar cover rate and maintenance level in relation to the frequency of Phytophthora sp species identified in Méagui. Boxplots were conducted to visualize this link between prevalence and these considered variables. These statistical tests were all performed at the 5% threshold using the SPSS Statistics 20 software.

3. RESULTS
3.1 Molecular characterization of species
The results of the molecular analysis showed that out of 66 black pods samples analyzed, 52 were positive for P. megakarya, i.e. 78.8% and 14 for P. palmivora, i.e. 21.2%. (Table 1). This result implies an epidemiological prevalence of 10.27% for P. palmivora and an epidemiological prevalence of 28.33% for P. megakarya at the Méagui site. The overall prevalence of the two species is 38.60% over the entire study site (Table 2).

3.2 Distribution maps of the different species of Phytophthora sp
The distribution map of the different species of phytophthora sp (Fig3) shows that P. megakarya is regularly distributed over all the sites of Méagui than P. Palmivora, which has a rather localized distribution. This is justified by a high distribution rate of P. megakarya around 78% compared to a low distribution rate of P. palmivora equivalent to 22%.

3.3 Influences of epidemiological factors on the prevalence of black pod
The result of the descriptive prevalence analysis showed that black pod averaged 43.09% with a standard deviation of ± 19.94% at the Méagui site. This shows that prevalence data are less clustered around the average. The minimum prevalence is 11% and the maximum prevalence is 100%. The description of the leaf coverage rate showed that most of the cocoa trees present at the Méagui site have a high leaf coverage with a frequency of 87.9% compared to 12.1% for moderately covered cocoa trees. The frequency of cocoa trees with low coverage is 0% at the study site. The result of the student t-test showed a highly significant difference (P=0.003<0.05) between the prevalence of cocoa black pods and the leaf cover rate of cocoa trees (Table 3). The boxplots show this link visually (Fig4). The descriptive analysis of the data on the level of maintenance of the plots also showed that 67% of the sampled plots are not maintained, while only 33% of the plots benefit from regular maintenance (Table 4). The means of the modalities of the maintenance level variable (Not maintained: 46.70 ± 20.55 and Maintained: 35.87±16.83) compared to the prevalence of cocoa trees (Table 5). The result of the student t-test showed a significant difference (P=0.036<0.05) between the prevalence of cocoa black pods and the level of maintenance of cocoa fields (Table 5). The boxplots show this link visually (Fig5). The results of the Chi-square test showed no significant relationship (Chi-square = 0.078, ddl=1, P=0.780>0.05) between the frequency of appearance of the two phytophthora species and the different modalities of the level of maintenance of cocoa plots.
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Cramer’s Phi and V test was not significant either (Phi and V to burn=0.034<1, P=0.78>0.05).

4. Discussion

The results of the molecular analysis showed a predominance of Phytophthora megakarya (79%) over Phytophthora palmivora (21%). Indeed, P. megakarya is a very aggressive specie that was not widespread enough in Ivorian cocoa plots. This specie was recently discovered in 1998 in eastern Côte d’Ivoire along the border with Ghana. The increase in P. megakarya has clearly reached almost all cocoa production areas in Côte d’Ivoire. In fact, the damage caused by black pod has long been attributed to P. palmivora, which was more widespread in Côte d’Ivoire and less aggressive. The significant progression of this disease in Méagui could be due to the high proportion of P. megakarya. Some studies carried out by CNRA had shown the presence of P. megakarya in the Department of Méagui since 2006 (CNRA 2007). This result is corroborated with those obtained in this study that shows a strong expansion of P. megakarya. The combined effect of the two Phytophthora species greatly increases the prevalence of the epidemic of black pod disease in the Department of Méagui by 43.09%. These results contradict those obtained by the study by Coulibaly et al (2013) which showed that P. palmivora was the most dominant specie and responsible for the cocoa black pods in southwestern Côte d’Ivoire. Today, this strong spread of P. megakarya in Ivorian cocoa plots, in addition to Swollen Shoot disease, is a real danger to Ivorian cocoa production. The result of the student t-test between prevalence and foliar coverage rate was highly significant (P=0.003<0.05). This significant link shows that the development of black pod in cocoa pods is strongly influenced by the high vegetative cover of cocoa trees. Indeed, the strong foliar coverage of cocoa trees creates shade under cocoa tree, which favours the maintenance of relative humidity. This humid microclimate is a determining factor in the development of fungal diseases, particularly in cocoa black pods disease. In addition, Tarjot (1971), in a study of the impact of various environmental factors on the development of black pod (due to Phytophthora palmivora) in Côte d’Ivoire, showed that high relative humidity increases the sensitivity of the fruit, as well as having a favourable effect on the development of the fungus. In addition, Medeiros (1976) links the evolution of black pods to climatic conditions, particularly relative humidity and rainfall. The low foliar coverage of cocoa trees favours the penetration of light into the undergrowth, thus reducing relative humidity. However, light has been identified as a growth inhibiting factor in different Phytophthora species (Blaha 1983). Efombagn (1999) showed that the in vitro growth of phytophthora sp isolates is influenced by light. He also noted that this growth is slow in continuous light, fast in darkness and average in alternating light (12 hours of light/12 hours of darkness). The results of this study are consistent with those obtained by the Coffee-Cocoa Council (2015), which showed that the development of black pod disease is influenced by the relative humidity associated with high cocoa tree shade. The student Test-t result showed a significant link (P=0.036<0.05) between the prevalence of black pod and the level of maintenance of the plot. This result indicates that the development of black pod can be attributed to the failure to maintain the plots. Indeed, weeding, draining and pruning of the plot are crucial operations in maintaining the cleanliness of the plot. These operations allow the destruction of the nests of certain pests such as insects that are capable of carrying the germs of the fungal pathogen. In addition, the Coffee-Cocoa Council has shown that weeds create a microclimate favourable to the proliferation of cocoa pests. Among these insects, many of them (ants, myriads, spiders...) are able to carry the germ from a rotten pod to a healthy pod. Other insects such as termites and ants, on the other hand, can carry the germ of the disease from the soil to the cocoa tree. While the description of the data relating to the level of maintenance of the plot showed that 67% of the cocoa plots sampled in Méagui are not maintained compared to 33% maintained. The result is confirmed by the studies carried out by Pohe et al (2013), indicating that the failure to maintain plots is linked to the demotivation of farmers who abandon cocoa farming in favour of other crops. This situation could be one of the determining factors in the increase in Phytophthora sp and particularly P. megakarya, which was not very widespread in Ivorian cocoa plots. The Chi-square test and the Phi and V of burning test showed that the foliar coverage rate of cocoa trees was not at the origin of the appearance of one Phytophthora species compared to another. This implies that both the high leaf cover rate is a factor that can favour the appearance of P. megakarya, as well as the appearance of P. palmivora. The same applies to the level of maintenance factor of the plots of land.

5. Conclusion, recommendations and perspectives

At the end of this study, it should be noted that the level of maintenance and the foliar coverage rate are two epidemiological variables favourable to the development of cocoa black pods in Méagui. The high coverage rate of cocoa trees and the lack of maintenance of the plots had a greater influence on the development of cocoa black pods. Indeed, these two factors create a microclimate favourable to the
development of *Phytophthora sp*, which is responsible for black pods disease. Moreover, between the two species identified in Méagui, *Phytophthora megalakarya* grows at a higher proportion (79%) than *Phytophthora palmivora* (21%) thus increasing the prevalence of the epidemic in the Department of Méagui. In view of these results, these recommendations can be observed for the control of black pod. This involves creating an aeration corridor around the farms by destroying the bush over a width of 5 to 10 m around it, maintaining the shade trees at a sufficient distance from each other to allow aeration of the farm, pruning the cocoa trees and carrying out the necessary weeding and finally carrying out the sanitary harvesting to stop the rapid spread of black pod disease in the farms. This requires regular removal of all affected pods as soon as the first typical stain appears. In the future, it would be interesting to extend the study to other sites in order to have a general idea of the prevalence of the *megalakarya* species in south-west Cote d'Ivoire; to carry out a very detailed mapping study in order to establish a risk map in order to better develop control methods for better disease control and for a clearer monitoring of the progression of this very aggressive species, *P. megalakarya*. It would be necessary to identify antagonistic fungi to *Phytophthora sp* in order to carry out biological control.

**Fig.1** Different typical symptoms of black pod disease on pods caused by *Phytophthora palmivora* and *phytophthora megalakarya*. (a): Contamination of the pod from above, (b): Contamination of the pods from below, (c): Lateral contamination of the pods and (d): Shrunken beans, as a result of the *phytophthora sp* attack (Picture taken by Koffi Alain in 2018)

**Fig.2** Map of the Nawa region showing the study site
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**Table 1:** Descriptive statistics of the proportion of each species of *Phytophthora sp*

| Species of *phytophthora sp* | Number | Percentage |
|-----------------------------|--------|------------|
| *P. palmivora*               | 14     | 21.2       |
| *P. megakarya*              | 52     | 78.8       |
| Total                       | 66     | 100        |

**Table 2:** Number of pods infected by each *Phytophthora sp* species in Méagui

| Number of infected pods | Prevalence of *Phytophthora sp* (%) |
|-------------------------|------------------------------------|
| *P. palmivora*           | 141                                | 10.27%                             |
| *P. megakarya*           | 389                                | 28.33%                             |
| Total infected pods      | 530                                | 38.60%                             |
| Total non-infected pods  | 843                                | 61.40%                             |
| Total sampled pods       | 1373                               | 100%                                |

**Table 3:** Comparison table of foliar coverage rate according to the prevalence of black pod

| Prevalence (%) | Foliar coverage rate |
|----------------|----------------------|
|                | Average | Strong |
| Fitolivore     | 8.00    | 58.00  |
| Average        | 24.00   | 45.72  |
| Standard deviation | 10.11  | 19.55  |
| Average standard error | 3.57  | 2.56   |

*P*= 0.003<0.05

**Table 4:** Descriptive statistics on the level of maintenance of the plot

| Level of maintenance of the plots | Number | Percentage (%) |
|-----------------------------------|--------|----------------|
| Not maintained                    | 44     | 66.7           |
| Maintained                        | 22     | 33.3           |
| Total                             | 66     | 100            |

**Fig. 4** Boxplots of coverage rate according to the prevalence of black pod
Table 5: Comparison table of the average of the different groups of the maintenance level of the plot with the prevalence of black pod

| Prevalence(%) | Level of maintenance |
|---------------|----------------------|
|               | Not maintained        |
| N             | 44.00                 |
| Average       | 46.70                 |
| Standard deviation | 20.55               |
| Average standard error | 3.09            |
| Maintained    | 22.00                 |
| Maintained    | 35.86                 |
| Maintained    | 16.83                 |
| Maintained    | 3.58                  |

Fig.5 Maintenance level boxplots according to the prevalence of black pod

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