

Original Research Article

Sweet potato (*Ipomea batatas* L.) a new alternative host of two phytoplasmas associated with Coconut Lethal Yellowing disease in Côte d'Ivoire

ABSTRACT

Aims: Coconut Lethal Yellowing Disease has been threatening coconut plantations in Côte d'Ivoire for ten years and has destroyed more than 400 ha. Coconut's destruction, the main host of phytoplasmas associated to the disease, leads to its conservation on reservoir plants grown in association or near from the disease outbreaks. Thus, the search for possible alternative host species was conducted.

Methodology: Surveys were carried out in infected coconut plantations, in order to describe symptoms associated with phytoplasma infections and to collect leaves samples of plant species other than coconut. Symptomatic and non-symptomatic leaves of sweet potato (*Ipomea batatas* L.) were collected for DNA extraction. The extracted DNA was subjected to molecular characterization.

Results: Mosaic and leaf reduction symptoms were observed on sweet potato leaves. From these samples, two phytoplasma strains were associated. Molecular analysis showed the presence of the endemic phytoplasma strain belonging to the 16SrXXII-B group "*Candidatus* Phytoplasma palmicola" and 16SrIV. Sequencing and phylogenetic analysis revealed the presence of a new phytoplasma strain, sharing 99% similarity with 16SrIV group strains "*Candidatus* Phytoplasma palmae".

Conclusion: The detection of these strains confirms sweet potato for the very first time, as an alternative host for coconut lethal yellowing phytoplasmas in Côte d'Ivoire.

Keywords: Sweet potato; coconut Lethal Yellowing Disease; coconut; côte d'Ivoire; 16SrIV; 16SrXXII-B

1. INTRODUCTION

Coconut (*Cocos nucifera* L.) is an economically important crop for many producing countries, including Côte d'Ivoire. Côte d'Ivoire is the 24th country in the world and 5th in Africa [1], with coconut production of 124,810 tons/year [1]. However, in Côte d'Ivoire, coconut plantations have been threatened by lethal yellowing disease for nearly a decade, mainly in the Grand-Lahou locality [2,3,4]. Infected coconut trees show yellow or dried palms along the stipe, which eventually loses its crown [3]. The disease has destroyed more than 400 hectares of producing coconut plantations, causing losses of more than 200 million CFA francs [5]. This has led to a considerable reduction in the economy of the producers. Furthermore, on an environmental aspect, destruction of coconut plantations in Côte d'Ivoire exposes the soil to sunlight and erosion [6].

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Comment [C1]: This sentence need to be rewritten.

The phytoplasma was associated with lethal yellowing disease in the work of [7]. It is a bacteria belonging to the class Mollicutes[8], which lives in both plants and insects [9]. In order to maintain favorable living conditions, phytoplasmas infect a large number of insect vectors and reservoir plants, both cultivated and non-cultivated, especially when the main host tends to disappear as a result of the infection.

Comment [C2]: This sentence need to be rewritten. This is meaningless.

In Côte d'Ivoire, the symptoms of lethal coconut yellowing disease have similarities to those of Cape St Paul disease in Ghana and are associated with phytoplasmas belonging to several groups that are 16SrI, 16SrXXII-B and 16SrXXII-C [10,4] Other studies have also identified phytoplasmas detected in several plant species, namely *Phyllanthusmuellerianus*, *Pennisetumpedicillatum*[9], palm (*Eleaisguineensis*L.), roast palm (*Borassusaethiopicum* L.) [4] and cassava (*Manihotesculenta* L.) [11]. In addition, insect vectors could feed on several plant species while transmitting the phytoplasma to them [12]. This raises the question of the transmission of phytoplasmas to plant species present in the plantations of their preferred or alternative hosts. Recently, symptoms similar to those of coconut lethal yellowing disease have been observed in coconut-producing localities other than the original outbreak in Grand-Lahou. These symptoms were associated with the presence of group XXII-B phytoplasma and the presence of a new phytoplasma infecting coconut belonging to group 16SrIV (MN545965) was even confirmed [13]. As a result, it is necessary to look for alternative host plants that could be involved in the spread of this disease. In Côte d'Ivoire, coconuts are often grown in association with sweet potato plant to meet the food needs of producing families. Coconuts are grown on plots of land devastated by the disease for their tubers and leaves. It is also a source of income to compensate for the losses caused by the fall in production from coconut plantations infected by the lethal yellowing disease. In this context, sweet potato could be an alternative host for phytoplasmas associated with coconut lethal yellowing disease. This study was therefore carried out with the aim of identifying a new alternative host for phytoplasmas associated with coconut lethal yellowing disease in Côte d'Ivoire.

2. MATERIAL AND METHODS

2.1 Collection site

The sweet potato leaves samples were collected in the infected coconut plantations, located in the locality of Grand-Bassam in 2018. The locality of Grand-Bassam is located in Côte d'Ivoire, south-east of the coast, between latitude: 5°12'46" North and longitude: 3°44'35" West.

2.2 Plant material

The plant material consisted of symptomatic and asymptomatic potato leaves. The leaves were collected from plants at the 3 to 5 leaf stage.

2.3 Observation and description of symptoms

In 2 sweet potato plantations in association with coconut plantations infected withcoconut lethal yellowing disease, the general appearance of the sweet potato plants was observed. The symptoms present on the leaves of sweet potato plants were described by considering the different types and forms of symptoms observed.

2.4 Collection of samples

Once the symptoms had been observed and described, samples of symptomatic and asymptomatic sweet potato leaf were collected at random from inside and outside the coconut plantations in Grand-Bassam. Symptomatic and asymptomatic leaf samples were labelled, transported to the laboratory at NANGUI ABROGOUA University (Côte d'Ivoire) and stored at -20 °C.

2.5 Nucleic acid extraction

To confirm the presence of phytoplasmas, total DNA was extracted from all sweet potato leaf samples according to the method of Doyle and Doyle [14]. One hundred mg of fresh symptomatic and asymptomatic plant tissue containing sweet potato leaf midribs were ground into a fine powder in CTAB lysis buffer. The DNA samples were eluted in 25 µL TE. The concentrations and purity of extracted DNA were measured using a NanoDropOne (Thermo Scientific, USA) and stored at -20 °C until use.

2.6 PCR amplifications

The detection of phytoplasmas in sweet potato leaf samples was conducted using PCR reactions. To do this, a 12.5 µL master mixture was used in a direct PCR with the primer pair P1/P7 [15,16], including 2 µL of DNA template, 6.25 µL of GoTaq G2 Green buffer (Promega, USA), 1.25 µL of each primer and 1.75 µL of sterilized distilled water (Promega, USA). Two µL of the products amplified by direct PCR were re-amplified in a nested PCR performed with the primer pair GH813f/AwkaSR [17], in a reaction solution with a total volume of 25 µL containing 12.5 µL of GoTaq G2 Green buffer (Promega, USA), 2.5 µL of each primer and 5.5 µL of sterilized distilled water (Promega, USA). The reaction conditions of the primer pair P1/P7 were as follows: 94 °C for 3 min; 94 °C for 40 s, 56 °C for 40 s and 72 °C for 1 min 40 for 35 cycles in total; 72 °C for 10 min for the final elongation. Nested PCR with the primer pair GH813f/AwkaSR was performed under the same conditions except at 53 °C for the hybridization temperature.

Another direct PCR was performed using the primer pair R16mF2/R16mR1 [18] for amplification of 2 µL of DNA in 12.5 µL master mixture containing 6.25 µL of GoTaq G2 Green buffer (Promega, USA), 1.25 µL of each primer and 1.75 µL of sterilized distilled water (Promega, USA). A 1:30 diluted template generated by R16mF2/R16mR1 primers was used and subjected to nested PCR using the universal primer pair R16mF2n/R16mR2 [18,19]. The master mixture contained 5 µL of diluted product, 25 µL of Go Taq G2 Green buffer (Promega, USA), 5 µL of each primer and 10 µL of nuclease-free water (Promega, USA) for a final volume of 50 µL. The reaction conditions of the primer pairs R16mF2/R16mR1 and R16mF2n/R16mR2 were as follows: 94 °C for 2 min; 94 °C for 1, 50 °C for 2 min, 72 °C for 3 min for 35 cycles in total; 72 °C for 10 min for the final extension. A positive control (phytoplasma DNA from an infected coconut palm) and a negative control (sterilized distilled water) were used in each PCR reaction.

The PCR products were visually detected by 1.5% agarose gel electrophoresis through ethidium bromide staining for 30 min at 80 volts. DNA bands were visualised using a trans-UV illuminator (EBOX VX5, VilberLourmat TM, France).

2.7 Sequencing and phylogenetic classification of the phytoplasma

PCR products using the R16mF2n/R16mR2 primer pair were sequenced by the Eurofins laboratory (France). The consensus nucleotide sequence obtained in the study was assembled using Genious Prime V 2019.1.3 software and saved on GenBank. The nucleotide sequence was compared and analyzed to identify the pathogen strain using the Basic Local Alignment Search Tool (BLAST) from NCBI (<http://www.ncbi.nlm.nih.gov>) [20]. Phylogenetic analysis was performed using MEGA X software [21]. The sequence obtained in this study

was aligned along with the ones published on GenBank or not of phytoplasmas strains using the Clustal X V 2.0 algorithm [22]. Neighbour-joining method with 1000 bootstrap replications was then carried out to evaluate the stability of the phylogenetic tree. The 16S rRNA sequence of *Acholeplasma laidlawii* (M23932.1) was used as an outgroup for the phylogenetic tree construction.

3. RESULTS

3.1 Diversity of symptoms observed on sweet potato leaves

Different types of symptoms were observed on sweet potato leaves collected from plants located inside and around of coconut plantations (Fig. 1). Firstly, there were anatomical changes in terms of a reduction in the size of the apical leaves (A). Secondly, symptoms of color change, with infected leaves showing alternating light green and yellow colors, intermingled over the entire surface, characteristic of the mosaic (B), unlike the green leaves on non-infected control plants (C).

3.2 Amplification of phytoplasma DNA associated with coconut lethal yellowing disease in sweet potato leaves

Amplicons 800 bp in size were obtained with the Gh813f/Awka SR primer (Fig. 2) in 2 samples of sweet potato leaves showing mosaic symptoms and in sweet potato leaves showing leaf reduction symptoms. Amplicons of 1250 bp were obtained with the R16F2n/ R16R2 primer (Fig. 3) in sweet potato leaves showing symptoms of leaf reduction. No amplicons were obtained from the sample of asymptomatic leaves.

3.3 Identification of phytoplasmas associated with coconut lethal yellowing disease in sweet potato leaves

BLAST analysis of homologous gene sequences of 16S rRNA gene sequence of sweet potato leaf reduction isolate obtained in this study and identified on in the Grand-Bassam locality (MN549454.1), shares 99% homology with phytoplasmas in the 16S rRNA IV group "*Candidatus* Phytoplasma palmae", associated with Coconut Lethal Yellowing in Mexico (KX982667.1). Phylogenetic analysis of partial 16S rRNA sequences confirmed the sequence analysis. The phytoplasma associated with the CILY isolate PATBAS identified in Grand-Bassam (MN549454.1) is in the same clade as the phytoplasmas associated with Coconut Lethal Yellowing in Mexico (KX982667.1) and Côte d'Ivoire (MN545965.1) respectively and which belong to the 16S rRNA IV group "*Candidatus* Phytoplasma palmae" (Fig. 4). This sequence obtained on sweet potato (MN549454.1) is in a different clade from that which includes the CILY-associated strains obtained at Grand-Lahou in Côte d'Ivoire on coconut (MN540266), oil palm (KY767914.1), raffia (KY711302.1) and roast palm (KY711392.1).

4. DISCUSSION

Observation of sweet potato (*Ipomea batatas* L.) plants revealed the presence of different types of symptoms and shapes on the leaves present in coconut plantations. These symptoms were reduced leaf size and leaf mosaic. The expression of these symptoms by the sweet potato could be explained by the action of phytoplasmas. The "little leaf" symptom observed on sweet potato leaves was associated with the presence of phytoplasma strains belonging to the 16S rRNA II group. The diversity of symptoms observed on sweet potato leaves is probably due to the action of different phytoplasma strains. These symptoms observed on the leaves could suggest a hormonal imbalance [23]. Indeed, when phytoplasma infects a

plant, it causes several physiological and morphological disorders represented by organ discoloration and deformation [24]. Furthermore, the presence of symptoms on sweet potato plants could be explained by the fact that the sweet potato may be an alternative host for the phytoplasma.

Comment [C3]: Reference(s)?

The present study revealed the presence of phytoplasmas in 4 symptomatic sweet potato leaf samples and not in the asymptomatic ones. This suggests that these phytoplasmas are associated with the various symptoms observed. The 800-bp amplicon obtained with the specific primer pair suggests the presence of the 16SrXXII-B "*Candidatus* Phytoplasma palmicola" phytoplasma associated with coconut lethal yellowing disease [2,9] and which is the most widespread phytoplasma group to date in Côte d'Ivoire as it has been discovered on infected coconut trees in other production localities including Grand-Bassam [13] and is identical to the phytoplasma associated with Cape St Paul Disease (CSPWD) in Ghana [25]. Sequence and phylogeny analyses have also identified another strain associated with leaf reduction in sweet potato (*Ipomea batatas* L.) and belonging to group 16Sr IV "*Candidatus* Phytoplasma palmae". This strain was recently identified and detected in infected coconut trees in Côte d'Ivoire [13]. The presence of this group is a discovery in both coconut and sweet potato in west Africa and more specifically in Côte d'Ivoire, as it is limited to Tanzania and countries in America and the Caribbean [26]. Thus, the presence of group 16Sr IV phytoplasma could be linked to the existence of an insect vector different or identical to the *Nedotepacurta* species suspected of being the vector of the group 16SrXXII-B phytoplasma strain.

The presence of these two strains belonging to the species "*Candidatus* Phytoplasma palmicola" and "*Candidatus* Phytoplasma palmae" in the species sweet potato (*Ipomea batatas* L.), thus indicates the adaptability of these strains to other cultivated species. Indeed, recent studies by Kra et al. [11] have shown that the 16SrXXII-B group strain has been identified in other crops such as cassava (*Manihotesculenta* Crantz) in Côte d'Ivoire, showing that cassava is a reservoir host for this phytoplasma.

Like cassava, sweet potato could constitute an alternative host and a potential reservoir for phytoplasmas associated with Coconut Lethal Yellowing disease in Côte d'Ivoire. However, here is currently no evidence to explain the epidemiological role of sweet potato in the spread of the phytoplasma associated with CILY, although this plant species has been present in coconut plantations infected by this disease. Moreover, diseases caused by phytoplasmas are strongly influenced by the number, abundance and diversity of alternative host plants and insect vectors [27]. Thus, sweet potato could be involved in the spread of these phytoplasmas and in the aggravation of the disease in Côte d'Ivoire.

The results obtained in this study show that sweet potato (*Ipomea batatas* L.) can be infected by both phytoplasmas associated with CILY in Côte d'Ivoire, belonging to the 16SrXXII-B "*Candidatus* Phytoplasma palmicola" and 16SrIV "*Candidatus* Phytoplasma palmae" groups. Studies will be needed to identify the role played by sweet potato in the spread of Coconut Lethal Yellowing phytoplasmas in Côte d'Ivoire.

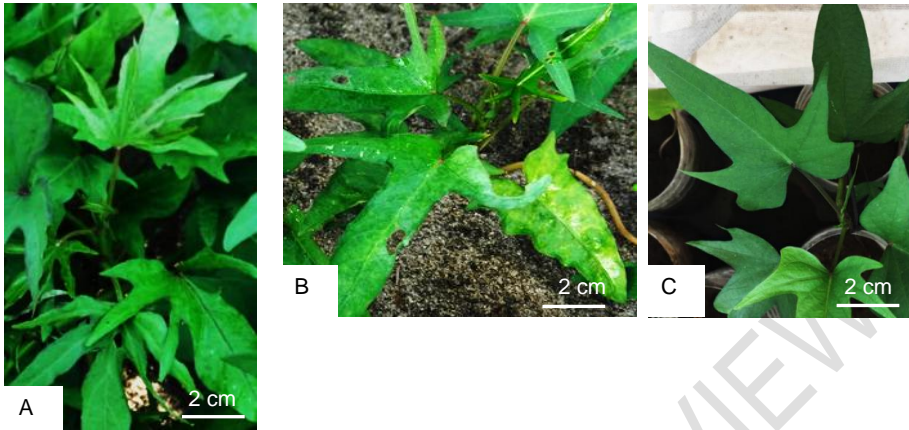


Fig. 1. Sweet potato (*Ipomea batatas* L.) leaves with or without symptoms
 A: Leaves with leaf reduction symptoms; B: Leaves with mosaic symptoms; C: Leaves without symptoms.

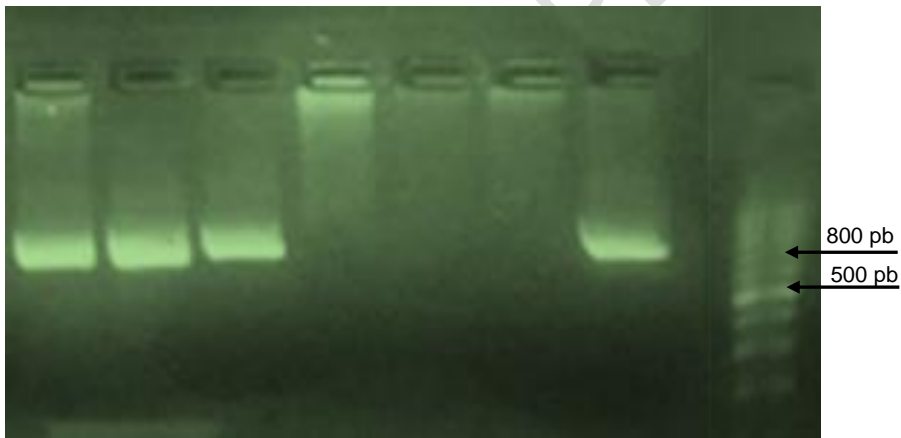


Fig. 2. Agarose gel electrophoretic profile of Nested PCR products of phytoplasma DNA in sweet potato (*Ipomea batatas* L.) leaf samples with the specific primer Gh813F/AwkaSR

M = molecular weight marker (100 bp); 13-14: positive sweet potato leaf samples showing the mosaic symptom; 15: sweet potato leaf samples showing the leaf reduction symptom; 16-17: asymptomatic sweet potato leaf samples; T+: positive control (sample of trunk boring collected from an infected coconut tree); T-: negative control (sterile water).

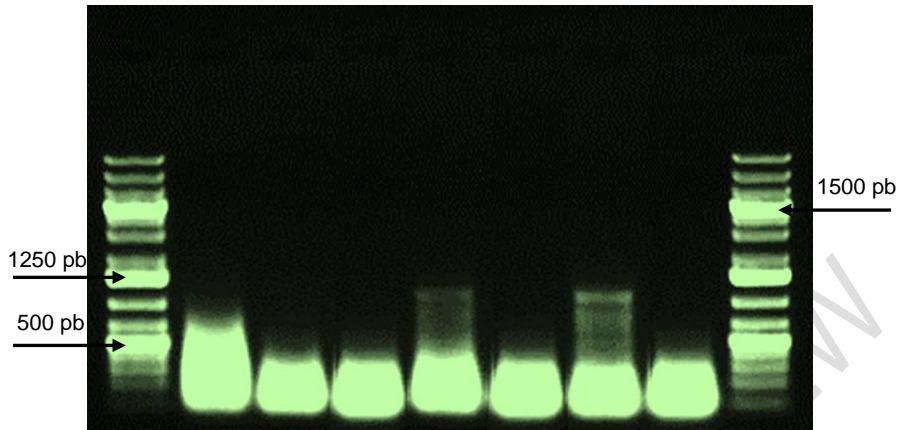
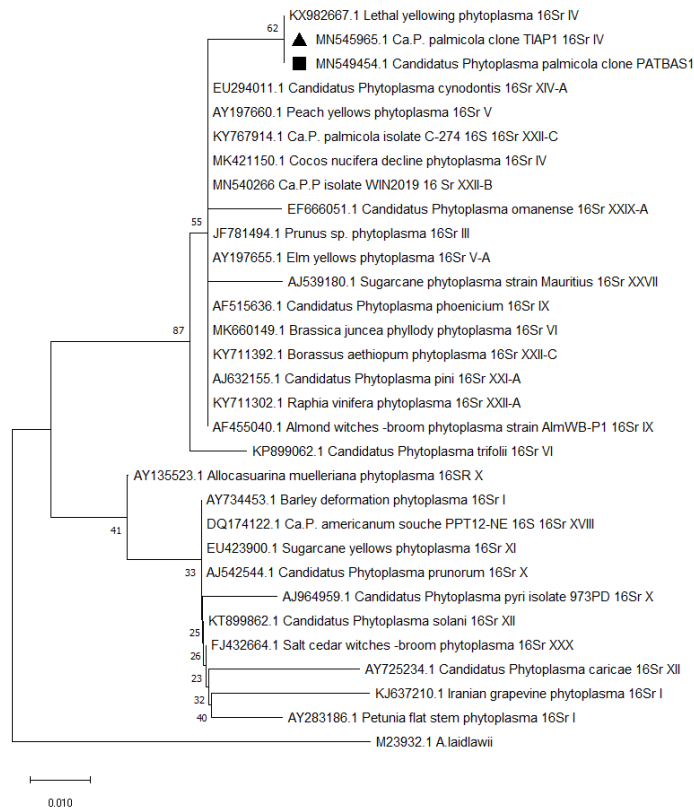


Fig. 3. Agarose gel electrophoretic profile of Nested PCR products of phytoplasma DNA in sweet potato (*Ipomea batatas* L.) leaf samples with the specific primer R16mF2n/ R16mR2

UNDER PEER REVIEW



M = molecular weight marker (1 kb); 1-2: asymptomatic sweetpotato leaf samples; 3 and 5: sweet potato leaf samples showing the mosaic symptom; 4: positive sweet potato leaf sample showing the symptom of leaf reduction; T+: positive control (sample of trunk boring collected from an infected coconut tree); T-: negative control (sterile water).

Fig. 4. Phylogenetic tree based on the R16mF2n/ R16mR2 sequences of the phytoplasma associated with leaf reduction in sweet potato (*Ipomea batatas* L.) and the 16S rRNA gene of phytoplasma reference sequences constructed using the neighbour-joining method with MEGA X

The species *Acholeplasmalaidlawii* was chosen to root the tree. Bootstrap values from 1000 replicates are indicated on the branches. GenBank accession numbers for each sequence are given before the phytoplasma name. ■ The strain obtained in this study ▲ The strain identified in coconut

5. CONCLUSION

Mixed cultivation of sweet potato and infected coconut palms could contribute to an exchange of agents associated with or responsible for infection. The present study reveals for the 1st time that phytoplasmas of groups 16Sr XXII-B and 16Sr IV associated with Coconut Lethal Yellowing disease affect sweet potato in Côte d'Ivoire. Once infected, sweet potato plants show symptoms of leaf deformation and discoloration. However, further studies will be needed to identify the vector and other alternative hosts of these phytoplasmas for better management of Coconut Lethal Yellowing disease in Côte d'Ivoire.

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