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**Isolation, purification and partial characterization of thermostable serine alkaline protease from a newly isolated *Bacillus thuringiensis*-SH-II-1A**

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In the present study, the isolation, purification and partial characterization of thermostable serine alkaline protease produced from *Bacillus thuringiensis* SH-II-1A was reported. The culture was isolated from soil of slaughter house waste and identified further from ribosomal sequence. The crude enzyme was purified by ammonium sulfate precipitation, dialysis and Sephadex G-200 gel permeation chromatography up to 17.04 fold with recovery of 8.47%. Relative molecular weight (67 kDa) of purified enzyme was observed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). Maximum production of enzyme and cell biomass was observed for 48 h of incubation period at 45°C. Strong activity of enzyme was observed at pH 10 to 11; also stability of up to 2 and 20 h incubation at the same pH range confirms alkaline protease. Optimum temperature recorded for protease activity was 45°C, and 100% thermal stability up to 350 min of incubation was recorded. Among different natural substrates tried, casein was found as ideal substrate. Enzyme activity was strongly enhanced by metal ions like Ca²⁺, Mg²⁺ and Mn²⁺ whereas, 100% enzyme activity was inhibited by phenylmethylsulphonyl fluoride (PMSF), and up to 92% inhibition by diisopropyl fluorophosphates (DFP) confirmed serine protease. Detergent compatibility of the enzyme was studied in the presence of 10 mM CaCl₂ and 1 M glycine at 45°C. This indicates 80 to 100% stability for a period of 0.5 to 2.5 h incubation. Improved washing performance and removal of blood stains from the cotton cloth was observed when detergent Surf excel was used with enzyme. Overall, the observed properties of isolated protease conclude its commercial application in detergent and leather industries.

**Key words:** Thermostable serine alkaline protease, *Bacillus thuringiensis*, purification, partial characterization.

**INTRODUCTION**

Among the large number of microbial enzymes, proteases occupy a pivotal position owing to their wide applications. The current estimated value of the worldwide sales of microbial enzymes is $ 1 billion; proteases alone account for about 60% of the total worldwide sales (Immaculate and Jamila, 2014) and were the first enzymes to be produced in bulk (Beg et al., 2003; Eilaiah et al., 2003). Alkaline proteases are a physiologically and commercially important group of enzymes used primarily as detergent additives. They play...
a specific catalytic role in the hydrolysis of proteins. This enzyme accounts for 40% of the total worldwide enzyme sales. It is expected to be an upward trend in the use of alkaline proteases in the future (Adinarayana et al., 2003; Moreira et al., 2003). Proteases are used in laundry detergents for over 50 years to facilitate release of proteinaceous materials in stains (blood and milk) and account for approximately 25% of total worldwide sales of enzymes (Ahmed et al., 2011). Thermostable alkaline proteases are having application in detergent industry as additive (Lagzian et al., 2012).

In the present study, the authors produced protease enzyme from Bacillus thuringensis SH-II-1A (Foda et al., 2013). The bacillus strain was obtained from natural source of slaughter house waste drainage soil of different locations (Joachim et al., 2008). Maximum growth of culture and production of enzyme was observed at 48 h of incubation period. Isolation and purification of the enzyme was carried out by sequential steps of ammonium sulphate precipitation, DEAE-cellulose dialysis and Sephadex G-200 gel permeation chromatography. Isolated protease enzyme was identified as serine type after maximum inhibition of enzyme activity by phenyl methyl sulfonyl fluoride (PMSF) and di-isopropyl fluoro phosphate (DFP). Maximum hydrolysis of casein as a substrate and maximum stability was observed at pH 10 to 11, which supports alkaline type protease. Proteolytic efficiency of the enzyme was observed at 45°C, which supports thermal stability of enzyme at higher temperature. Activity of enzyme was enhanced by the use of metal ions like calcium (CaCl₂) supporting cofactor requirement. Enzyme has shown greater compatibility and stability with commercial detergent formulations like surf excel up to 1.5 h of incubation. Improved washing performance and removal of blood stains from the cotton cloth was observed upon incubation of enzyme along with detergent powder. Molecular weight of the purified enzyme was up to 68 kDa estimated with SDS-PAGE and supported by gelatin zymogram activity.

As it was desirable to search for new proteases with novel properties from as many different sources as possible, the present enzyme could become one of the ideal sources to fulfill all the required properties. Overall, the isolated protease enzyme could be considered as an additive for commercial detergent formulations. Commercially, it would have many industrial applications like hair removal property in leather industry, as a laundry additive in detergent industry, and breakdown of X-ray films to remove trapped silver particles. It would have therapeutic applications as a digestive aid, fibrinolytic agent, dissolution of blood clots and treatment of atherosclerosis in the future scope.

MATERIALS AND METHODS

The following chemicals were obtained from Sigma-Aldrich, St. Louis, MO (USA): Casein, Sephadex G-200, bovine serum albumin (BSA), phenylmethylsulfonyl fluoride (PMSF), β-mercaptoethanol (β-ME), DEAE-Cellulose equilibrium dialysis Harvard apparatus. Heavy metal ions, surfactants, organic solvents and chemicals for electrophoresis were purchased from Merck, Germany. Molecular mass markers (20 to 205 kDa) were obtained from Biotech-Genetix Pvt Ltd, Bangalore, India. Commercial detergent powder and electrophoresis Vertical Slab Gel System were from Asian Scientific Instruments Ltd. All other chemicals used such as Ariel, Tide, Surf, Surf excel, Nirma and Rin were purchased from local market of high analytical grade.

Isolation and cultivation of microorganisms

The soil samples were collected in the form of soil from the waste drainage areas of slaughter house at six different locations of Pune district region (MS) India, and were diluted in sterile saline solution (Palsaniya et al., 2012). The clear supernatant liquid of the diluted samples was plated onto skim milk agar plates containing peptone (0.1% wt/vol), NaCl (0.5% wt/vol), agar (2.0% wt/vol) and skim milk (10% vol/vol). Plates were incubated at 37°C for 24 h. A clear zone of skim milk hydrolysis gave an indication of protease producing organisms (Shata, 2005; Shieh, 2009). Depending on the maximum zone of hydrolysis, strain SH-II-1A was selected for further experimental studies. The pure isolated proteolytic strain was further subjected to identification of nucleotide sequence at National Centre for Cell Science (NCCS), Pune (MS), India. It was identified as B. thuringensis and it was designated as B. thuringensis SH-II-1A used in further study. Production of protease from B. thuringensis SH-II-1A was carried out in a medium containing the following components: glucose 1% (wt/vol); yeast extract 0.5% (wt/vol); tryptone, 0.5% (wt/vol); casein, 1.0% (wt/vol); soluble starch, 1.0% (wt/vol); NaNO₂ 1.0% (wt/vol); CaCl₂, 1.0% (wt/vol); NH₄Cl 1.0% (wt/vol); it was maintained at 45°C for 72 h in a shaker incubator (150 rpm). The pH of the medium was preadjusted with 0.1 N NaOH or 0.1 N HCl. After the completion of fermentation, the whole fermentation broth was centrifuged at 5000 x g at 4°C for 15 min, and the clear supernatant was recovered. The crude enzyme supernatant was subjected to purification and further studies (Amrita et al., 2012; Akolkar 2009).

Enzyme purification

Ammonium sulphate precipitation

As described above, fermentation was carried out for 72 h at 45°C and clear supernatant was separated from the cells by centrifuge at 5000 xg for 15 min. Further supernatant was fractionated by precipitation with ammonium sulfate between 40 and 60% of saturation. All subsequent steps were carried out at 5 to 8°C. The protein mass obtained after precipitation was resuspended again in 0.1 M Tris-HCl buffer, pH 10.0, and dialyzed using diethyl amino ethyl (DEAE) cellulose membrane against the buffer having same composition (Asker et al., 2013).

SephadexG-200 gel permeation chromatography

Sephadex G-200 (5 gm) gel was added to 0.1 M Tris-HCl (pH 10.0) and allowed to swell overnight; and column (1.5 x 65 cm) (Sigma-Aldrich, St Louis, MO) was packed. The column (1.5 x 65 cm) was equilibrated with 0.1 M Tris-HCl buffer and pH 10.0. The protein pellet obtained after saturation with ammonium sulphate between 50 and 70% was dissolved in 0.1 M Tris-HCl buffer and pH 10.0, and a total of 5 mL of protein precipitate was loaded onto a column of SephadeG-200 (1.5 x 65 cm). The column was equilibrated
with Tris-HCl buffer and pH 10.0. The process of elution was started at a flow rate of 1 mL/min with a 1:1 volume gradient flow from 0.1 to 1 M NaCl in the same buffer. Protein concentration from each fraction was determined by Lowry method using bovine serum albumin as standard. Such fractions were pooled, dialyzed and concentrated to be used for further studies (Asker et al., 2013).

**Molecular mass determination**

Molecular mass of the purified enzyme fractions showing greater activity was determined on sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) according to the method of Laemmli (1970) and Ali et al. (2014) with 15% acrylamide concentration. Approximately, 10 µg of purified protein was loaded on 15% SDS-PAGE with standard molecular mass markers into two side wells of the gel. The dragging process of loaded protein was started at initial current of 15 mA and then constant current of 30 mA. After electrophoresis, the gel was stained with Co-omassie Brilliant Blue R-250 (CBB R-250) and destained in glacial acetic acid to visualize protein bands.

**Determination of proteolytic activity (caseinase determination)**

Caseinase activity was assessed by the modified procedure (Tsuchida et al., 1986) using 2% casein substrate dissolved in 0.2 M carbonate buffer pH 10. Casein solution (0.5 ml) with an equal volume of suitably diluted enzyme solution was incubated at 37°C. After 15 min, the reaction was terminated by addition of 1 ml of 10% trichloroacetic acid (TCA). The reaction mixture was centrifuged and 5 ml of 0.44 M Na,CO₃ was added to the supernatant. Addition of two fold diluted (1 mL) Folin Ciocalteu reagent to the mixture resulted in blue color. After 45 min at room temperature, the reaction mixture was subjected to centrifugation (Remi, 4000 rpm, 4°C). The intensity of color developed in the supernatant mixture was measured at 660 nm against a reagent blank prepared in the same manner. Tyrosine served as the reference standard and optical density of colored solution was measured by UV spectrophotometer (Shimadzu, Japan).

**Protein assay**

Protein was measured by the method of Lowry et al. (1951) with bovine serum albumin (BSA) as the standard. The concentration of protein during purification studies was calculated from the absorbance at 280 nm (Singhal et al., 2012).

**Effect of pH on enzyme activity and stability**

The activity of the enzyme (10 µl) was measured at different pH values in the presence of 10 mM CaCl₂. The pH was adjusted using the following buffers (0.05 M): phosphate buffer (pH 6.0 to 7.0), Tris-HCl buffer (pH 8.0) and glycine-NaOH buffer (pH 9.0 to 12.0). Reaction mixtures containing 0.5% casein (1 mL) were incubated at 45°C for 30 min. The reaction was stopped by addition of 0.5% Tri chloro acetic acid (TCA). Absorbance of TCA soluble peptides was measured at 280 nm. The purified enzyme was diluted in different relevant buffers (pH 6.0 to 12.0) and incubated at 45°C for 2 and 20 h for checking of enzyme stability. The relative activity at each exposure was measured as per assay procedure (Siala et al., 2009).

**Effect of temperature on enzyme activity and stability**

The activity of the enzyme was determined by incubating the reaction mixture at different temperatures ranging from 35 to 65°C in the presence of 10 mM CaCl₂ for 30 min. To determine the enzyme stability with changes in temperature, purified enzyme was incubated at different temperatures (35, 40, 45, 50, 55, 60 and 65°C) in the presence of 10 mM CaCl₂ and relative protease activities were assayed at standard assay conditions (Bhunia et al., 2012).

**Effect of protease inhibitors on enzyme activity**

The effect of different protease inhibitors on purified protease activity was studied using phenyl methyl sulfonyl fluoride (PMSF) (1 mM), ethylene diamine tetra acetic acid (EDTA) (5 mM), p-chloromercuric benzoate (p-CMB) (5 mM), lodoacacetate (5 mM), D-isopropyl fluoro phosphate (DFP) 5 mM and β-mercaptoethanol (β-ME) (5 mM) (Sigma et al., 1975). The reaction mixture was prepared by pre-incubating the purified enzyme with inhibitors without substrate for 10 min at 45°C. The protease assay was performed by addition of casein as a substrate for 30 min. Protease activity obtained without inhibitor was considered as 100% (Usharani et al., 2010).

**Effect of various metal ions on enzyme activity**

Effect of monovalent (Na⁺ and K⁺), divalent (Ca²⁺, Mg²⁺, Mn²⁺, Co²⁺, Cd²⁺, Zn²⁺ and Hg²⁺) and trivalent (Al³⁺, Fe³⁺) metal ions on enzyme activity at a concentration of 5, 10 and 15 mM was investigated by using casein as substrate (Sigma et al., 1975). The reaction mixture was prepared by pre-incubating the purified enzyme with metal ions at each concentration for 10 min at 45°C and the proteolytic activity was determined for 1 h by the above mentioned method. Enzyme activity in the absence of metal ions was considered as 100% (George et al., 2012).

**Hydrolysis of protein substrates**

Protease activity with different protein substrates including BSA, casein, egg albumin and gelatin was assayed by mixing 100 ng of the enzyme and 200 µL of assay buffer containing the protein substrates (2 mg/mL). After incubation at 45°C for 30 min, the reaction was stopped by adding 200 µL of 10% (wt/vol) trichloro acetic acid (TCA) and allowed to stand at room temperature for 10 min. The undigested protein was removed by centrifugation and peptides released were assayed. The specific protease activity towards casein as a substrate was considered as a control (Iqbal et al., 2011).

**Detergent stability**

The compatibility of SH-II-1A protease with local laundry detergents was studied in the presence of 10 mM CaCl₂ and 1 M glycine. Detergents used were Nirma (Nirma Chemical, India); Henko (Henkel Spic, India); Surf, Surf Excel, Super Wheel, Rin (Hindustan Lever Ltd, India); and Ariel (Procter and Gamble, India). The detergents were diluted in distilled water (0.7% wt/vol) and incubated with protease for 3 h at 45°C, and the residual activity was determined. The enzyme activity of a control sample (without any detergent) was taken as 100% (Ire et al., 2011).

**Washing test with protease preparation**

Application of protease (5000 U/mL) as a detergent additive was studied on white cotton cloth pieces (4 × 4 cm) stained with blood.
The stained cloth pieces were taken in separate flasks. The following sets were prepared and studied: 1. Flask with distilled water (100 mL) + stained cloth (cloth stained with blood); 2. Flask with distilled water (100 mL) + stained cloth (cloth stained with blood) + 1 mL Wheel detergent (7 mg/mL); 3. Flask with distilled water (100 mL) + stained cloth (cloth stained with blood) + 1 mL Wheel detergent (7 mg/mL) + 2 mL enzyme solution.

The above flasks were incubated at 45°C for 15 min. After incubation, cloth pieces were taken out, rinsed with water, and dried. Visual examination of cloth pieces exhibited the effect of enzyme in removal of stains. Untreated cloth pieces stained with blood were taken as control for comparison (Ram et al., 2012).

### RESULTS AND DISCUSSION

**Identification of bacterial culture**

The most significant characteristics of the isolate SH-II-1A are summarized: these are vegetative cells and characterized as large stout rods that are straight or slightly curved with rounded ends (Gopala, 2011). They usually occur in pairs or short chains. It is Gram positive, non-capsulated and motile with peritrichous flagella. Classification of *B. thuringiensis* strains has been accomplished by H serotyping the immunological reaction to the bacterial flagellar antigen (Ash et al., 2011). The *hag* gene encodes flagellin, which is responsible for eliciting the immunological reaction in H serotyping. This showed that the isolate belongs to the class of *Bacillus* sp.

**Purification of extracellular protease of *B. thuringiensis* SH-II-1A**

In *B. thuringiensis*, maximum growth and maximum enzyme production was observed at 48 h (Figure 1). Fermentation was carried out at 45°C and clear supernatant was separated from the cells by centrifuging at 5000 ×*g* for 15 min. Further supernatant was fractionated by precipitation with ammonium sulfate between 40 and 60% of saturation. All subsequent steps were carried out at 5 to 8°C. The protein mass obtained after precipitation was resuspended in 0.1 M Tris-HCl buffer, pH 10; it was dialyzed using diethyl amino ethyl (DEAE) cellulose membrane against the buffer having same composition in order to achieve desired purity of enzyme.

The protein pellet obtained after dialysis using DEAE membrane was loaded onto a column of Sephadex G-200 (1.5 × 65 cm) equilibrated with Tris-HCl buffer, pH 10. The elution profile of gel filtration chromatography is shown in Figure 2. Elution profile of column was monitored and it was observed that the protease enzyme was eluted in the form of well resolved single peak, showing casein hydrolysis (caseinase) activity coinciding with a single protein peak at a NaCl concentration of 0.6 M. Eluted fractions sequence numbers (25 to 30) were observed with high protease activity. The summary of purification steps involved for alkaline protease is presented in Table 1 (Zhou et al., 2009; Fakhfakh et al., 2010).

When the ammonium sulphate precipitation and purified protease was analyzed by SDS-PAGE, seven bands were observed in the presence of ammonium sulphate precipitation (Figure 3); while purified protease showed a single band on SDS-PAGE, indicating a homogenous preparation. The molecular weight of the protease was determined by comparison of the migration distances of standard markers protein. The molecular mass standards were bovine serum albumin (67 kDa), ovalbumin (45 kDa), carbonic anhydrase (30 kDa), trypsinogen (24 kDa) and α-lactalbumin (14 kDa) on SDS-PAGE. The molecular mass was determined by extrapolation from a linear semi logarithmic plot of

![Figure 1. Maximum growth of culture and enzyme production.](image-url)
Figure 2. Elution profile of protease SH-II-1A by Sephadex G-200 column chromatography.

Table 1. Summary of purification steps for isolated enzyme.

<table>
<thead>
<tr>
<th>Purification step</th>
<th>Total enzyme activity (U)</th>
<th>Total protein (mg)</th>
<th>Specific activity (U/mg)</th>
<th>Purification folds</th>
<th>Recovery (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude enzyme</td>
<td>105000</td>
<td>6025</td>
<td>17.4</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>Ammonium sulphate ppt. and dialysis</td>
<td>90500</td>
<td>4000</td>
<td>22.62</td>
<td>1.29</td>
<td>86.19</td>
</tr>
<tr>
<td>Sephadex G-200</td>
<td>8900</td>
<td>30</td>
<td>296.66</td>
<td>17.04</td>
<td>8.47</td>
</tr>
</tbody>
</table>

Figure 3. SDS-PAGE and Gelatin zymogram for enzyme. Lane A, Molecular mass marker; Lane B, crude enzyme; Lane C, purified enzyme; Lane D, standard BSA; Lane E and G, gelatin zymogram for standard BSA; Lane F, gelatin zymogram for purified enzyme. The molecular mass markers are: myosin 205 kDa, phosphorylaseB-97 kDa, bovine serum albumin-68 kDa, ovalbumin-43 kDa, carbonic anhydrase-29 kDa, tyrosine-20 kDa.
Characterization of purified enzyme

For the determination of the pH optimum, phosphate (pH 6 to 7), Tris-HCl (pH 8), and glycine-NaOH (pH 9 to 12) buffers were used in the absence and presence of 10 mM CaCl₂. The highest protease activity was found to be at pH 10 to 11 using glycine-NaOH buffer (Figure 4). These findings are in line with several earlier reports showing pH optima of 10 to 10.5 for protease from *Bacillus* sp., *Thermus aquaticus*, *Xanthomonas maltophilia* and *Vibrio metschnikovii*. The important detergent enzymes, subtilisin Carlsberg and subtilisin Novo or BPN (Dhandapani et al., 1994) also showed maximum activity at pH 10.5. The stability of enzyme was checked at various pH 6 to 12. 100% stability was achieved at pH 9 to 11 for a period of 2 and 20 h incubation (Figure 5).

The activity of the crude and purified enzyme was determined at different temperatures ranging from 35 to 65°C in the absence and presence of 10 mM CaCl₂. The optimum temperature recorded was at 45°C for protease activity. The enzyme activity gradually declined at
temperatures beyond 50°C (Figure 6). A similar type of result was observed by other investigators where a maximum temperature of 55°C was recorded for an alkaline protease from *Bacillus stearothermophilus* AP-4 and 60°C for a protease (Jellouli et al., 2009) derived from *Bacillus* sp B21-2. The thermal stability of the purified protease was tested at different temperatures of 45, 50 and 55°C for different periods (50 to 350 min) in the presence of 10 mM CaCl₂. The enzyme was almost 100% stable at 45°C even after 350 min of incubation (Figure 7). Previous reports on thermostability have shown half-lives of >200 min at 50°C and 2 to 22 min at 60°C for heat stable serine proteases, and half-lives of 3.4 and 2.4 min at 50°C have been recorded for subtilisin Carlsberg and subtilisin BPN, respectively.

**Effect of protease inhibitors on enzyme activity**

Inhibition studies primarily give an insight into the nature of an enzyme, its cofactor requirements, and the nature of the active center. The effect of different inhibitors on the enzyme activity of the purified protease was studied. Among the inhibitors tested (at 5 mM concentration), PMSF was able to inhibit the protease completely, while DFP exhibited 94% inhibition. In this regard, PMSF sulphonates, the essential serine residue in the active site of the protease and has been reported to result in the complete loss of enzyme activity. Relative enzyme activity by different inhibitors is described in Figure 8. Our findings are similar to those of Tsuchida et al. (1986) where the protease was completely inhibited by PMSF.
This indicated that it is a serine alkaline protease. Slight inhibition was observed with other inhibitors like iodoacetate, p-CMB and β-ME.

**Effect of various metal ions on enzyme activity**

Most of the metal ions tested had a stimulatory effect (Ca$^{2+}$, Mg$^{2+}$ and Mn$^{2+}$) or a slight inhibitory effect (other ions) on enzyme activity. Some of the metal ions such as Ca, Mg$^{2+}$ and Mn$^{2+}$ increased and stabilized the protease activity of the enzyme; this is possible because of the activation by the metal ions. These cations also have been reported to increase the thermal stability of other *Bacillus* alkaline proteases. These results suggest that concerned metal ions apparently protected the enzyme against thermal denaturation and played a vital role in maintaining the active confirmation of the enzyme at high temperatures. Other metal ions such as Zn$^{2+}$, Cu$^{2+}$, Hg$^{2+}$, Co$^{2+}$, N$^{3+}$, Cd$^{2+}$, Al$^{3+}$ and Fe$^{3+}$ did not show any appreciable effect on enzyme activity. Relative enzyme activity is described in Figure 9.
Hydrolysis of protein substrates

When assayed with native proteins as substrates, the protease showed a high level of hydrolytic activity against casein and poor to moderate hydrolysis of BSA and egg albumin; although the hydrolysis was hardly observed with gelatin (Figure 10) (Loops et al., 2003).

Compatibility with detergents

Besides pH, a good detergent protease is expected to be stable in the presence of commercial detergents. Protease from B. thuringiensis SH-II-1A showed stability and compatibility with a wide range of commercial detergents at 45°C in the presence of CaCl$_2$ and glycine (Annamalai et al., 2013). High activity alkaline protease was reported from Conidiobolus coronatus showing compatibility at 50°C, in the presence of 25 mM CaCl$_2$, with a variety of commercial detergents. Also, 16% activity was reported in Revel, 11.4% activity in Aerial and 6.6% activity in Wheel. Comparing these results, the Bacillus subtilis PE-11 enzyme was significantly more stable in commercial detergents. The compatibility of alkaline protease was studied with Surf excel in the presence of 10 mM CaCl$_2$ and 1 M glycine for different periods (0.5 to 2.5 h) at 45°C. The enzyme retained about 82% activity after 1.5 h in the presence of surf excel at 45°C and was almost inactivated after 2.5 h in the absence of any stabilizer (Figure 11). However, the addition of CaCl$_2$ (10 mM) and glycine (1 M), individually
and in combination, was very effective in improving the stability, where it retained 60% activity even after 2.5 h. As the protease produced by our isolate *B. thuringinsis* SH-II-1A was stable over a wide range of pH values and temperatures and also showed compatibility with various commercial detergents tested in the presence of CaCl₂ and glycine, it was used as an additive in detergent to check the contribution of the enzyme in improving the washing performance of the detergent. The supplementation of the enzyme preparation in detergent (that is, Surf excel) could significantly improve the cleansing of the blood stains (Figure 12).

**Conclusion**

The alkaline protease isolated from *B. thuringinsis* SH-II-1A is a thermostable serine protease. It is stable at alkaline pH, at high temperatures, and in the presence of commercial detergents and is compatible with commercial and local detergents. These properties indicate the possibilities for use of the protease in the detergent industry. This enzyme can be exploited commercially.

**CONFLICT OF INTERESTS**

The authors have not declared any conflict of interests.

**Abbreviations**

NCCS, National Centre for Cell Sciences; DEAE, di ethyl amino ethyl; CBB, co-ommassie brilliant blue; SDS-PAGE, sodium dodecyl sulphate poly-acrylamide gel electrophoresis; TCA, tri-chloro acetic acid; UV, ultra violet; BSA, bovine serum albumin; p-CMB, para chloro mercuric benzoate; PMSF, phenyl methyl sulfonyl fluoride; β-ME, β-mercapto ethanol; DFP, di isopropyl fluoro phosphate.

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Cucurbit yellow stunting disorder virus and Watermelon chlorotic stunt virus induced gene silencing in tobacco plants

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Cucurbit yellow stunting disorder virus (CYSDV) and Watermelon chlorotic stunt virus (WmCSV) are the most widespread and damaging viruses to cucurbits in the Middle East. CYSDV and WmCSV are cucurbit-infecting bipartite whitefly-transmitted viruses. Post-transcriptional gene silencing (PTGS) is a universal mechanism by which plants are able to systemically switch off the expression of targeted genes via the reduction of steady-state levels of specific RNAs. PTGS was used in this study to control the two viruses. In this study, the efficiency of the dsRNA for the ability to trigger resistance against the CYSDV and WmCSV was investigated. Three regions of three genes of CYSDV genome were selected; the coat protein gene (CP), heat shock gene (Hsp70) and ORF3, while the two regions of two genes of WmCSV genome were selected; CP gene and rep gene. Bioassay, dot-blot hybridization and polymerase chain reaction (PCR) methods were capable to evaluate the resistance against viruses. Clear symptoms on tobacco plants took two to three weeks to appear and all non-infiltrating tobacco plants (positive control) showed viral symptoms after inoculation. Most of the agro-infiltrating sense/antisense constructs did not yield symptoms of the viruses. Dot-blot hybridization, showed that negative hybridization was obtained with infiltrating tobacco plants with prepared constructs compared to those non-infiltrating tobacco plants used as the control. Only one out of five gave positive signals with the construct pasCYSDV-Hsp70. Using PCR, positive reactions of the expected size of 500 bp fragment with WmCSV and 800 bp with CYSDV were obtained with the infiltrating tobacco plants with sense constructs, which pointed out the existence of viral genome in challenging tobacco plants. Infiltrating tobacco plants with sense/antisense constructs gave negative PCR pointed out the lack of the viral genome.

Key words: Cucurbit yellow stunting disorder virus (CYSDV), watermelon chlorotic stunt virus (WmCSV), Post-transcriptional gene silencing (PTGS), coat protein (CP), Hsp70, ORF3, Rep, dot-blot, hybridization.

INTRODUCTION

Cucurbit crops are widely grown in the Middle East and consumed in large quantities in the traditional diet. The warm climate permits the nearly year-round development of insect vectors such as the whiteflies, which are the main vector for the viruses that infect cucurbits. The most widespread and damaging viruses are members of the
genus Begomovirus (*Geminiviridae*) and Crinivirus (*Closteroviridae*). The most important time of the year for whitelyf infestation of cucurbit crops in Saudi Arabia is the autumn (Raja Al-Zahrani, 2010). During 2001, two new viral diseases were detected in cucurbits: watermelon (*Citrusillus lanatus*), squash (*Cucurbita pepo*) and cucumber (*Cucumis sativus L.*) in the Middle East region. Sequence analysis revealed that the diseases were caused by the native *Watermelon chlorotic stunt virus* and *Squash leaf curl virus* (Abdel-Salam et al., 2006; Abudy et al., 2010; Ali-Shtayeh et al., 2014; Idris et al., 2006). *WmCSV* was first identified in Yemen (Jones et al., 1988) and after that, in the following years, it was reported through the Middle East and also in North Africa as the following: Sudan (Khey-Pour et al., 2000), Jordan (Al-Musa et al., 2011), Oman (Khan et al., 2012) and Palestine (Ali-Shtayeh et al., 2014). The symptoms caused by WmCSV are chlorotic mottling, vein yellowing, stunting and severe decrease of yield, generally in watermelon (*Citrusillus species*) crops. WmCSV host range covers the most cucurbits including squash, cucumber, melons, and pumpkin.

The first detection of CYSDV in the Middle East was reported in the United Arab Emirates in 1982 (Hassan and Duffus, 1991). In subsequent years, it was identified throughout the Mediterranean region, in Europe, and in North America (Abou-Jawdah et al., 2000; Brown et al., 2007; Kuo et al., 2007; Papayiannis et al., 2009; Sweiss et al., 2007). CYSDV is a bunched crinivirus, which has had a main negative impact on the production of cucumbers in several regions of the world, a phenomenon increased by the absence of effective preventive arrangements against the whitefly (Owen et al., 2016). WmCSV and CYSDV infect all cultivated cucurbits, some legumes (alfalfa, bean), and malvaceous species. CYSDV causes severe symptoms in cucurbit plants, green spots on the oldest leaves, severe interveinal chlorosis, and spot appears between 14 and 22 days’ post-inoculation and severe symptoms including leaf rolling, complete leaf lamina yellowing, brittleness, severe stunting of young leaves and a drastic reduction of yield (Célix et al., 1996).

One of the primary aspects of virus-plant host interactions is derived from plant natural defenses through RNA silencing (Ding and Voinnet, 2007; Ruiz-Ferrer and Voinnet, 2009). Through infection, plants process viral double-stranded RNAs (dsRNAs) into small interfering RNAs (siRNAs) and use the siRNAs to guide specific antiviral silencing activities (Li et al., 2015). siRNAs guide members of the Argonaute protein family to specific mRNAs for cleavage, a process referred to as post-transcriptional gene silencing (PTGS), which is a universal defense mechanism against RNA viruses and transcripts produced by DNA viruses such as geminiviruses (Li et al., 2015). The lack of natural sources of resistance to WmCSV and CYSDV and the need to reduce pesticide application motivates us to develop multiple virus resistance through the development of transgenic cucurbit plants. This study was conducted to improve the socioeconomic status of the Saudi Arabia farmers and the local economy by the accurate diagnosis of cucurbit virus diseases and development of transgenic plant resistant to those viruses adapted to local conditions. In the present study, infiltrating tobacco plants with sense/antisense constructs gave negative PCR which pointed out the lack of the viral genome while the infiltrating tobacco plants with sense constructs pointed out the existence of viral genome in challenging tobacco plants.

**MATERIALS AND METHODS**

The present work was conducted at Pest and Plant Diseases Unit (PPDU), College of Agriculture and Food Sciences, King Faisal University, Saudi Arabia.

**Extraction of total nucleic acids from cucurbit tissues**

The total RNA (for CYSDV) was extracted from infected cucurbit plants using the RNeasy® Plant Mini Kit (Qiagen, Germany) according to the instructions of the manufacturer while the total DNA (for WmCSV) was extracted from infected cucurbit plants using the DNeasy® Plant Mini Kit (Qiagen, Germany) according to the instructions of the manufacturer.

**Design and synthesis of oligonucleotide primers**

Specific primers were designed for parts (selected fragments) of the coat protein gene (CP) of CYSDV (176 bp), heat shock gene (Hsp70) of CYSDV (232 bp), ORF3 of CYSDV (198 bp), CP gene of WmCSV (160 bp) and rep gene of WmCSV (230 bp) (Table 1) according to the sequences obtained from GenBank. The sense primers are CYcpr, Hsp702f, CYorf3f, Wmcp2 and Wmrep1 with Ascl and Xbal sites (underlined), respectively at the 5’ end (Table 1). The antisense primers are CYcpr, Hsp702r, CYorf3r, Wmcp2 and Wmrep2 with Swal and BamHI sites (underlined) respectively, at the 5’ end (Table 1). The restriction sites were inserted at the 5’ end of the prepared primers to ease the cloning of the selected fragments into the binary vector.

**One-step reverse transcriptase-polymerase chain reaction (RT-PCR) for the selected fragments of CYSDV**

Platinum Quantitative RT-PCR Thermo Script One Step System experiment. The RT reaction started with incubation for 30 min at 50°C, then denaturation for 5 min at 95°C. Thirty-five cycles of PCR started with denaturation for 1 min at 95°C, primer annealing for 1 min at 55°C, and extension for 1 min at 72°C with a final extension

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for 7 min at 72°C.

**PCR for the selected fragments of WmCSV**

PCR cycle parameters were as follows: 2 min at 94°C, after that, 35 cycles of 1 min at 94°C, 2 min at 55°C and 2 min at 72°C, then the final extension for 10 min at 72°C.

**Cloning and sequencing of the amplified fragments of CYSDV and WmCSV**

PCR products of CYSDV and WmCSV pGEM®-Easy vector (Promega, USA) were ligated together. The insertion of recombinant plasmids in strain DH5α of E. coli occurred according to the instructions of the manufacturer. The Wizard® Plus SV Miniprep DNA Purification System (Promega, USA) was applied for the isolation of DNA from different white colonies. Then the digestion of DNA was done using EcoRI and electrophoresed on agarose gels. The orientation of these inserts in the plasmid was determined using two different primers for sequencing, the first one was T7 forward primer in the promoter region (5’ TAATACGACTCACTATAGGG ‘3) and the second was SP6 reverse in promoter region (5’ATTTAGGTGACACTATAG ‘3). The clones having inserts of accurate sizes were selected for sequencing using an automated instrument for DNA sequencing (ABI 377XL). Sequencing Analysis Software, ABI™ version 3.0 was used to analyze the data. All sequencing was carried out in Macrogen Company, Seoul, South Korea.

**Sub-cloning of the selected fragments into a binary vector**

The selected fragments for gene silencing (CYSDV-Cp, CYSDV-Hsp70, CYSDV-ORF3, WmCSV-CP and WmCSV-rep) were re-amplified from the clones into pGEM-Teasy and sub-cloned into the binary vector pFGC5941 (kindly provided from the University of Arizona, USA) in both orientations once to give the sense orientation by restriction enzymes Ascl and SwaI and another one to give the antisense orientation with XbaI and BamHI departing the intron Chalcon Synthase (ChS) in the middle (Rezk et al., 2006; Soliman et al., 2008).

**Preparation of sense, antisense and sense/antisense constructs**

The sense constructs were prepared through the digestion of the DNA plasmids (pCYSDV-Cp, pCYSDV-Hsp70, pCYSDV-ORF3, pWmCSV-CP and pWmCSV-rep) with Ascl (Thermo Fisher Scientific, USA) and SwaI (Thermo Fisher Scientific, USA) as the following: 50 U of SwaI and 10 µl of 10X buffer O were used to digest 50 µl of each DNA plasmid and dhO2 were added to reach 100 µl. The tubes were incubated at 30°C for 4 h and 50 U of Ascl and 10 µl of 10X buffer Tango™ were added, and incubated at 37°C for 4 h. The antisense constructs were prepared through the digestion of the plasmids with XbaI and BamHI as described in antisense preparation and the tubes were incubated at 37°C in water bath for 4 h. The digested DNAs were electrophoresed in 2% agarose gel. A clean sharp scalpel was used to cut out the digested bands for gene cleaning. The restriction enzymes (the same enzymes) were used to digest the binary vector as described in each case of sense and antisense constructs. At the same time, the prepared sense constructs (psCYSDV-CP, psCYSDV-Hsp70, psCYSDV-ORF3, psWmCSV-CP and psWmCSV-rep) were digested in another reaction with XbaI and BamHI. The digested vector (100 ng) was added with 15 ng of the digested DNA of the selected fragment to a clean tube (both of them were digested with the same restriction enzymes). 1 U of T4 DNA ligase, 1 µl of 10X ligase buffer and final volume of the 10 µl of the ligation mixture was obtained using nuclease-free water, and finally incubated overnight at 4°C.

**Transformation of the constructs into E. coli competent cells**

Competent cells E. coli (strain DH5α) were thawed on ice for 30 min and 100 µl was transferred to a pre-chilled 5 ml tube and 5 µl of ligated DNA was added and incubated on ice for another 30 min. Cells were heat-shocked for 2 min at 42°C by immersing the tube into a water bath for 2 min and then incubated on ice for 2 min. One milliliter of Luria-Bertani (LB) medium was added and cells grown for 1 h at 37°C in shaking water bath. Transformed cells were plated on the surface of prepared plates (LB solid medium with 50 µg/ml kanamycin) for overnight at 37°C and bacterial colonies containing recombinant plasmids were selected for further characterization. The enzyme digestion for the DNA minipreps of the selected colonies of constructs (sense, anti-sense, sense/anti-sense) were carried out using the restriction enzymes SwaI and Ascl (sense), BamHI and XbaI (anti-sense) and Ascl and XbaI (sense/anti-sense) as described earlier.

**Nucleotide sequence analysis**

Two primers were designed in the binary vector pFGC5941; one in the p35S promoter location (pFGC35SF, 5’AGATGGACCCCCACCCAGGAGGAGG3’) and the other in the OCS location (pFGC-OSCR, 5’AGGATCTGAGCTACATGCTAGG3’).

**Transformation into Agrobacterium tumefaciens LBA4404**

One microgram of plasmid DNA was added to the cells. Freezing the cells in liquid nitrogen was performed. The cells were thawed by incubating the tubes in a 37°C water bath for 5 min. 1 ml of YEP medium was added to the tube and incubated at 28°C for 2 to 4 h with gentle shaking. This period allowed the bacteria to express the antibiotic resistance genes. The tubes were centrifuged for 30 s in a table centrifuge. The supernatant solution was discarded and the cells were resuspended in 0.1 ml YEP medium. The cells were spread on a YEP agar plate containing 50 µg/ml kanamycin and 50 µg/ml streptomycin. The plate was incubated at 28°C. Transformed colonies should appear in 2 to 3 days.

**Gene delivery using the syringe-spotting technique (SST)**

After A. tumefaciens LBA4404 has been transformed with vectors harboring different constructs using chemical transformation, 5 ml culture was grown overnight at 28°C in the appropriate antibiotic selection medium. The next day, the culture was inoculated into a 50 ml LB medium containing the selective antibiotics. The culture was grown overnight in a 28°C shaker. Agrobacterium cells were harvested and resuspended in infiltration media [10 mM MgCl2, 10 mM MES (2-N-morpholino ethane sulfonic acid), and 20 µM acetosyringone (4’-hydroxy-3,5-dimethoxyacetophenone)], adjusted
to OD_{600} of 2.0 and left at room temperature for 3 h. The prepared constructs (pasCYSDV-CP, pasCYSDV-Hsp70, pasCYSDV-ORF3, pasWmCSV-CP, and pasWmCSV-rep) in sense and sense/antisense direction were used to inoculate all experimental plants in this study. Tobacco (Nicotiana benthamiana) plants were infiltrated at the stage of seedlings in 2 to 3 leaves (each treatment contains 10 plants) with Agrobacterium containing both prepared constructs and the empty binary vector without construct (as a control) using a 5 ml syringe with no needle (Johansen and Carrington, 2001; Abhary, 2003). Fifteen days after the infiltration of the constructs, the challenge with infection viruses was carried out using the syringe spotted technique with infectious clones of WmCSV and CYSDV.

### Evaluation of the resistance triggered against viral infection

The agro-infiltrating plants of N. benthamiana with different constructs were inoculated with viral infectious clones and the development of the disease symptoms was observed. In addition, RNA for CYSDV and DNA for WmCSV were extracted as described earlier; after that, PCR and non-radioactive hybridization methods were applied to detect the two viruses in the challenged plants.

### RESULTS

#### PCR and RT-PCR analysis of the selected fragments

The designed specific primers CYcpf/CYCpr, Hsp702f/Hsp702r and CYorf3f/CYorf3r (Table 1), were used successfully for the amplification of the selected fragments of the coat protein gene (CP), heat shock 70 (Hsp70) and ORF3, respectively. In addition, the designed specific primers Wmcp1/Wmcp2 and Wmrep1/Wmrep2 (Table 1) were used successfully for the amplification of the selected fragments of coat protein gene (CP) and rep gene. The selected fragments of CYSDV were 176 bp of CP, 232 bp of Hsp70 and 198 bp of ORF3. While the selected fragments of WmCSV were 160 bp of CP and 230 bp of rep gene (Figure 1).

### Cloning and sequencing of the selected fragments

The recombinant plasmids were digested with EcoRI restriction endonuclease to test for inserts of 176, 232 and 198 bp of CYSDV and 230 and 160 bp of WmCSV. One recombinant plasmid with inserts pCYSDV-CP, pCYSDV-Hsp70, pCYSDV-ORF3, pWmCSV-CP and pWmCSV-rep were selected for sequencing and sub-cloning in the binary vector pFGC5941.

### Nucleotide sequence analysis of the selected fragments

Sequences of the PCR amplified fragments in the recombinant (pCYSDV-CP, pCYSDV-Hsp70, pCYSDV-ORF3, pWmCSV-CP and pWmCSV-rep) were completed. The sequences obtained with T7 forward, SP6 reverse primers were aligned with CYSDV, and WmCSV sequences available in the GenBank.

### Sub-cloning of the selected fragments into the binary vector

Sub-cloning of the selected fragments into the binary vector were done in the two orientations once by the Ascl and Swal (sense orientation) and second time with Xbal and BamHI (antisense orientation) departing the Chalcon Synthase (ChaS) Intron in the middle and finally the third

---

### Table 1. Nucleotide sequences of the prepared primers to amplify the selected fragments of CYSDV and WmCSV.

<table>
<thead>
<tr>
<th>Viruses' name</th>
<th>Primers' name</th>
<th>Nucleotide sequences (5’---------------------------3’)</th>
<th>Restriction enzymes</th>
<th>Size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CYSDV</td>
<td>CYcpf</td>
<td>ATATTCTAGAGGCCGCGCACCACACACTCATGCGACGG</td>
<td>XbaI, Ascl</td>
<td>176</td>
</tr>
<tr>
<td></td>
<td>CYcpr</td>
<td>ATATGGATCCATTTAAATAGTTCTAGGCCTGCGCG</td>
<td>BamHI, Swal</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CYorf3f</td>
<td>ATATTCTAGAGGCCGCGCACCACACACTCATGCGACGG</td>
<td>XbaI, Ascl</td>
<td>198</td>
</tr>
<tr>
<td></td>
<td>CYorf3r</td>
<td>ATATGGATCCATTTAAATAGTTCTAGGCCTGCGCG</td>
<td>BamHI, Swal</td>
<td></td>
</tr>
<tr>
<td>Hsp702f</td>
<td></td>
<td>ATATTCTAGAGGCCGCGCACCACACACTCATGCGACGG</td>
<td>XbaI, Ascl</td>
<td>232</td>
</tr>
<tr>
<td>Hsp702r</td>
<td></td>
<td>ATATGGATCCATTTAAATAGTTCTAGGCCTGCGCG</td>
<td>BamHI, Swal</td>
<td></td>
</tr>
<tr>
<td>WmCSV</td>
<td>Wmcp1</td>
<td>ATATTCTAGAGGCCGCGCACCACACACTCATGCGACGG</td>
<td>XbaI, Ascl</td>
<td>160</td>
</tr>
<tr>
<td></td>
<td>Wmcp2</td>
<td>ATATGGATCCATTTAAATAGTTCTAGGCCTGCGCG</td>
<td>BamHI, Swal</td>
<td></td>
</tr>
<tr>
<td>Wmrep1</td>
<td></td>
<td>ATATTCTAGAGGCCGCGCACCACACACTCATGCGACGG</td>
<td>XbaI, Ascl</td>
<td>230</td>
</tr>
<tr>
<td>Wmrep2</td>
<td></td>
<td>ATATGGATCCATTTAAATAGTTCTAGGCCTGCGCG</td>
<td>BamHI, Swal</td>
<td></td>
</tr>
</tbody>
</table>
construct of sense/antisense was prepared.

Nucleotide sequence analysis

The selected clones of the sense/antisense constructs were prepared for sequencing as described earlier. Nucleotide sequencing of the silencing fragments from the sense/anti-sense constructs were completed and the sequences were the same as the sequence obtained earlier.

PCR for the recombinant plasmids

PCR was done on the recombinant plasmids obtained from the Alkaline Lysis Miniprep method as described earlier. Using the previous profile of PCR, it was found that the transformation was carried out successfully and the bands appeared at their expected sizes.

Transformation into Agrobacterium

The constructs pasCYSDV-CP, pasCYSDV-Hsp70, pasCYSDV-ORF3, pasWmCSV-CP and pasWmCSV-rep in sense and anti-sense direction and binary vector pFGC5941 without construct as a control were transformed into A. tumefaciens LBA4404. Tobacco (N. benthamiana) plants were transformed using Syringe-Spotting Technique.

Evaluation of the transgenes

Screening for viral symptoms

The monitoring of symptoms appearance, after inoculation with viral infectious clones, was done daily to record the severity of symptoms. Clear symptoms on tobacco plants (N. benthamiana) took two to three weeks to appear (Table 2). All non-infiltrating tobacco plants (positive control) showed viral symptoms after inoculation. Most of the agro-infiltrating sense/antisense constructs did not yield symptoms of the viruses.

Screening with dot-blot hybridization

Tobacco plants inoculated with viral infectious clones were used as sources for DNA in dot-blot hybridization for the detection of viral genome in challenging plants. Total RNA and total DNA were extracted from new leaves using the methods mentioned earlier and blotted into nylon membranes. Hybridization was carried out using probes prepared to detect the viral genome in infiltrating plants with the different constructs. Results presented in Figure 2 showed that negative hybridization was obtained with infiltrating tobacco plants compared to those of non-infiltrating plants used as control. Only one out of five gave positive signals with the construct pasCYSDV-Hsp70.

Screening with RT-PCR and PCR

RT-PCR and PCR were carried out on the extracted RNA and DNA, respectively for the detection of the viral genome in challenging plants, after 15 days of inoculation with infectious clones of CYSDV and WmCSV. RT-PCR was applied using CYSDV-CP1 and CYSDV-CP2 primers to detect CYSDV. While, PCR was done using WmF2 and WmR2 primers to detect WmCSV. Positive reaction of the expected size, 500 bp of WmCSV and 800 bp of CYSDV were obtained with the infiltrating plants in case
Table 2. Viral symptoms severity after challenging with viral infectious clones showed by the number of infected tobacco plants per the number of infiltrating tobacco plants.

<table>
<thead>
<tr>
<th>Construct’s name</th>
<th>No symptoms</th>
<th>Mild symptoms</th>
<th>Severe symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td>pasCYSDV-CP</td>
<td>5/5</td>
<td>0/5</td>
<td>0/5</td>
</tr>
<tr>
<td>pasCYSDV-Hsp70</td>
<td>2/5</td>
<td>2/5</td>
<td>1/5</td>
</tr>
<tr>
<td>pasCYSDV-ORF3</td>
<td>4/5</td>
<td>1/5</td>
<td>0/5</td>
</tr>
<tr>
<td>pasWmCSV-CP</td>
<td>5/5</td>
<td>0/5</td>
<td>0/5</td>
</tr>
<tr>
<td>pasWmCSV-rep</td>
<td>5/5</td>
<td>0/5</td>
<td>0/5</td>
</tr>
<tr>
<td>pFGC5941</td>
<td>0/5</td>
<td>0/5</td>
<td>5/5</td>
</tr>
<tr>
<td>Untreated plants</td>
<td>0/5</td>
<td>0/5</td>
<td>5/5</td>
</tr>
</tbody>
</table>

Figure 2. Non-radioactive labeling system dot-blot hybridization for virus detection in the treated plants. 1, pasWmCSV-CP; 2, psWmCSV-CP; 3, pasWmCSV-rep; 4, psWmCSV-rep; 5, pasCYSDV-CP; 6, psCYSDV-CP; 7, pasCYSDV-ORF3; 8, psCYSDV-ORF3; 9, pasCYSDV-Hsp70; 10, psCYSDV-Hsp70; 11, Untreated plants as control.

of sense constructs (ps) which proved the existence of the genome of both viruses in challenging plants (Figures 3 and 4). Infiltrating plants with sense/antisense (pas) constructs gave negative reactions indicating the lack of the viral genome. The obtained results indicate that the pasWmCSV-CP and pasCYSDV-CP constructs succeeded in preventing viral replication and infection in infiltrating plants through the siRNA-mediated resistance.

DISCUSSION

The application of new techniques is becoming a widely used strategy to control diseases caused by plant viruses. Post-transcriptional silencing (PTGS) of viral genes using co-suppression or anti-sense constructs performs a little ratio of silenced individuals (Wesley et al., 2001). There are two hallmarks in PTGS; first, silencing of target miRNAs occurs in the cytoplasm. Secondly, small interfering RNA (siRNA) molecules (21-25 nt) are generated from the silenced target mRNAs. Based on differences in their biogenesis, types of small RNAs (miRNA and siRNA) have been identified (Carthew and Sontheimer, 2009). The second type (siRNA) was used as a tool to have resistance cucurbit plants for CYSDV and WmCSV. The A. tumefaciens-mediated transient expression system is a multilateral system to quickly introduce genes into plant tissue. This system enables gene expression within a short period of time and without the demand for regenerating transgenic plants. The Agrobacterium mediated expression system has also been used effectively as a way to deliver RNA silencing suppressors and inducers into transgenic plants that express a silencing reporter gene (Johansen and Carrington, 2001).

Data from this study demonstrate clearly the effectiveness of the gene silencing application technique to trigger resistance against CYSDV and WmCSV. Three conserved, un-translated sequences of CYSDV genome were selected; the coat protein gene (CP), heat shock gene (Hsp70) and ORF3. While two conserved, un-translated sequences of WmCSV genome were selected;
Figure 3. PCR for DNA extracted from different agro-infiltrating plants after 15 days of inoculation with infectious clone, using the primers WmF2 and WmR2. (1) A, plants treated with psWmCSV-CP; B, plants treated with pasWmCSV-CP. (2) A, plants treated with psWmCSV-rep; B, plants treated with pasWmCSV-rep.

Figure 4. RT-PCR for RNA extracted from different agro-infiltrating plants after 15 days of inoculation with infectious clone, using the primers CYSDV-CP1 and CYSDV-CP1. (1) A, plants treated with psCYSDV-CP; B, plants treated with pasCYSDV-CP. (2) A, plants treated with psCYSDV-ORF3; B, pasCYSDV-ORF3. (3) A, psCYSDV-Hsp70; B, pasCYSDV-Hsp70.

CP gene and rep gene. In this study, the Syringe-Spotting Technique has been adapted to deliver different constructs into tobacco (N. benthamiana) cells. The high efficiency of this technique to convey the constructs may be attributed to the presence of large amount meristematic cells in small leaf tissues of newly evolved
seedlings, which supply the most suitable host-cells for the DNA to be expressed, and the bacterium to be linked (Soliman et al., 2008). Although the transformed plant cells are limited to the spotted region, RNA silencing signals could be systemically transmitted, as supposed by Eckardt (2002). Andrieu et al. (2012) mentioned that, the results of their work indicate that the method is efficient at inducing gene silencing in the agro-infected leaf area.

The transfer of low amounts of siRNA, probably occurring passively through the symplastic pathway from the agro-infected area, seemed sufficient to trigger degradation of target transcripts in the adjacent tissues. Data presented in this study shows that all transformed tobacco plants with empty vector and subsequent inoculated with CYSDV and WmCSV showed viral symptoms after 2 to 3 weeks of inoculation. These results suggest that the binary vector was single-handedly incapable to trigger resistance against CYSDV and WmCSV. In a similar way, all transformed tobacco plants with sense construct were oversensitive (susceptible) to the infection with CYSDV and WmCSV. These results suggest that, the transcription of the sense silencing trigger resulted in a mRNA that lacked the coding for any protein and did not recombine to the viral mRNA during the viral infection cycle.

Plants that had been transformed with antisense construct were all susceptible to the infection with CYSDV and WmCSV. However, these plants developed disease symptoms after two weeks of inoculation with CYSDV and WmCSV. This suggests that the progression rate of disease in these plants was lower than that with other treatments. High level of resistance could be induced in tobacco plants against CYSDV and WmCSV using sense/antisense construct in transformation. No disease symptoms could be observed in plants two and three weeks after inoculation and these plants remained symptomless until the experiment has been terminated (3 weeks after inoculation with CYSDV and WmCSV). These results clearly showed that, dsRNA folding could be produced by the sense/antisense orientation through the complementarities between the sense and the antisense orientations of the silencing triggers after the splicing of the ChaS intron post-transcriptionally. As previously described in the literature review, the dsRNA would be chopped into small temporal RNAs that can guide the host cell RISC protein to degrade the complementary sequence, which is the viral mRNA, in a sequence specific manner. This might explain the reason why plants transformed with sense/antisense construct did not show any disease symptoms during the experiment. The sequence homology of the silencing trigger might guide the RISC protein in tobacco cells to degrade the CYSDV and WmCSV messenger RNA before expressing the replicas and the coat protein, preventing the virus from initiating the replication cycle. Previous studies showed that the 21-23 nucleotides RNA known as siRNA is the main factor of gene silencing (Voinnet, 2002). The siRNA is derived from dsRNA by the action of an RNase III-like enzyme (Dicer). In another study, Llave (2002) showed that the small RNAs, extracted from inflorescent tissues, were accumulated in tissue-specific manner.

Conclusion

Data presented in this study provide evidence that the gene silencing (siRNA) technology could be used as significant virus-control measure. As indicated, high rate of plants was resistant to CYSDV and WmCSV when they were transformed with sense/antisense construct.

CONFLICT OF INTERESTS

The authors have not declared any conflict of interests.

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**Full Length Research Paper**

**Tomato yellow leaf curl virus: Diagnosis and metabolites**

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The existence of Tomato yellow leaf curl virus (TYLCV) was figured out in different locations in Al-Ahsaa of Saudi Arabia. Polymerase chain reaction (PCR) results of samples collected showed that TYLCV existed in all locations. Using AVcore and ACcore primers, begomoviruses family were detected in symptomatic tomato plants and by using TYv2664 and TYc138 (specific primers for the detection of TYLCV), the results proved that the samples were infected with TYLCV. The lipid-soluble fraction of healthy and infected tomato leaves extract was compared using gas chromatography techniques. A total of 46 compounds were identified in both healthy and virus-infected leaf tissues; among which 37 metabolites were common between both samples and increased or decreased in concentration due to the virus attack. Nevertheless, eight compounds were exclusively detected in the infected samples with only one compound consumed and thus recognized only in the healthy samples. The classifications and roles of the identified metabolites were discussed from the point of view of plant defense mechanisms or virus resistance against plant defense.

**Key words:** Tomato yellow leaf curl virus (TYLCV), begomoviruses, Polymerase chain reaction (PCR), gas chromatography.

**INTRODUCTION**

Tomato (\textit{Solanum lycopersicum}, L.) is economically important in Saudi Arabia and is one of the most important vegetable crops in the world. It is considered as one of the most popular and widely grown vegetable crops worldwide with the area harvested in Saudi Arabia being 14,902 ha in 2016 (FAOSTAT, 2016). The production of tomatoes in Saudi Arabia in 2016 was 503,217 tonnes with most of that production (60%) grown in greenhouses (FAOSTAT, 2016).

Begomoviruses have one (monopartite) or two (bipartite) genomic components, denominated DNA-A and DNA-B, and are transmitted in a persistent manner.
Figure 1. Healthy (A) and TYLCV-infected (B) tomato leaves. Picture on the left represent the healthy leaves and picture on the right represent infected leaves by TYLCV. The plant samples were taken from tomato fields and green houses in Al-Ahsaa, Eastern Province, Saudi Arabia.

by whiteflies of the species complex Bemisia tabaci Gennadius (Hemiptera: Aleyrodidae) to dicotyledonous plants (Fernandes et al., 2010). Tomato yellow leaf curl virus (TYLCV) is one of the most important harmful and invasive members of the genus begomovirus (family Geminiviridae), which is widespread over the world associated with tomato yellow leaf curl disease (Barboza et al., 2013). TYLCV is transmitted by whiteflies and can spread rapidly; it is also not transmitted through seed or by mechanical transmission. Severe symptoms such as leaf curling, stunting, and yellowing showed on TYLCV-infected tomato plants, which cause serious production loss in tomato cultivation (Kil et al., 2016; Papayiannis et al., 2010). In addition to tomato, other cultivated plants including pepper (Capsicum species), common bean (Phaseolus vulgaris), cucurbit (Cucumis species) and eustoma (Eustoma grandiflora) have been reported to be TYLCV hosts (Anfoka et al., 2009; Kil et al., 2016). Polymerase chain reaction (PCR) was applied to detect and establish provisional identity of begomoviruses through amplification of 575 bp fragment of the begomoviral coat protein gene (CP), referred to as the ‘core’ region of the CP gene (core CP). The core CP fragment contains conserved and unique regions, and was hypothesized to constitute a sequence useful for begomovirus classification (Brown et al., 2001).

Metabolomics is used nowadays as a high potential tool for understanding different metabolic changes in many biological systems and its applications have been recognized in the quality control validations and natural products research (Dai et al., 2010). However, the use of metabolomics in investigating interactions between different organisms is until now infrequent. For example, a metabolic profile for Catharanthus roseus leaves infected with phytoplasma has been determined (Choi et al., 2004) and the aromatic metabolite profiles of Arabidopsis thaliana infected by Pythium sylvaticum has been investigated (Bednarek et al., 2005).

To the knowledge of the authors, the metabolites that resulted from the interaction between the Tomato yellow leaf curl virus and its host the tomato plant (S. lycopersicum) has never been investigated. The main aim of this study was to explore the type of phytochemicals newly produced, increased or decreased in concentration during the attack of TYLCV on the tomato plant leaves. The study could ascribe to the better knowledge of the plant-virus chemical connection.

MATERIALS AND METHODS

Plant material

Healthy tomato (S. lycopersicum, family Solanaceae) leaves samples (HTL) were identified and collected from the local fields and greenhouses in Al-Ahsaa, Eastern province, Saudi Arabia. Meanwhile, TYLCV naturally-infected tomato leaves samples (ITL) showing the typical symptoms of TYLCV infection (severe stunting, yellowing, curling of leaves and chlorosis on leaves) were collected from same field greenhouses (Figure 1). All plants were identified by experts and taxonomists in the College of Agriculture and Food Sciences, King Faisal University, Saudi Arabia with specimens deposited to the herbarium of the college.

Primers for begomoviruses and TYLCV

Two sets of primers were used in this study to identify the TYLCV infection (Table 1). The first set of primers were AVcore and ACCore used for the detection of begomoviruses and the second set of primers were TYv2664 and TYc138 used for the detection of TYLCV (Table 1).
Table 1. The sequences of the primers used to detect the Begomoviruses and TYLCV.

<table>
<thead>
<tr>
<th>Viruses</th>
<th>Primers’ Name</th>
<th>Nucleotide Sequences</th>
<th>Size (bp)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Begomoviruses</td>
<td>AVcore</td>
<td>5’- GCCHATRTAYAG RAAGCCNAGRAT -3’</td>
<td>575</td>
<td>Brown et al. (2001)</td>
</tr>
<tr>
<td></td>
<td>ACcore</td>
<td>5’- GGRRTDGARCCATGHGATACANGCC -3’</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TYLCV</td>
<td>TYv2664</td>
<td>5’- ATTGACCAAGATTTTACCTTACCC -3’</td>
<td>316</td>
<td>Anfoka et al. (2005)</td>
</tr>
<tr>
<td></td>
<td>TYc138</td>
<td>5’- AAGTGCGGTCCACATATTGCAAGAC -3’</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Extraction of total DNA from plant tissues**

Total DNA was isolated from the infected tomato plants using DNeasy® Plant Mini Kit obtained from QIAGEN as manufacturer’s instruction.

**Polymerase chain reaction (PCR)**

The extracted DNA was used as a template for PCR using set of primers as shown in Table 1. AVcore and ACcore primers were used as degenerate primers for begomoviruses group to amplify 575 bp while TYv2664 and TYc138 primers were used to amplify 316 bp of IR of TYLCV. PCR reactions were optimized for 25 µl and the final concentrations of reaction components were: 25 µM deoxynucleotide triphosphate (dNTPs), 2.5 µM of 10X PCR buffer, 2.5 mM MgCl₂, 5 units Taq DNA polymerase, 1 µl of 10 µM of each complementary and viral-sense primers and 3 µl of DNA were used as target templates. PCR cycle parameters for AVcore and AC core primers were as follows: one cycle at 94°C for 2 min; 35 cycles at 94°C for 1 min, 55°C for 2 min, and 72°C for 2 min, followed by one cycle at 72°C for 10 min. PCR cycle parameters for TYv2664 and TYc138 primers were as follows: one cycle at 94°C for 5 min; 30 cycles at 94°C for 1 min, 62°C for 1 min, and 72°C for 1 min, followed by one cycle at 94°C for 1 min, 56°C for 1 min, and 72°C for 10 min. Five microliters aliquots of PCR products were analyzed on 1% agarose gels in 0.5X TBE buffer.

**Plant sample preparation for analysis**

Three hundred grams of the healthy and TYLCV-infected tomato leaves were isolated directly from the field to make the HTL and ITL samples, respectively. The samples were immediately placed into liquid nitrogen for preservation and enzyme deactivation and then pulverized into a powdered form and kept in -20°C until further analysis.

**Metabolites extraction**

The powdered samples were extracted by n-hexane (HPLC-grade, Fisher Chemicals) using a Soxhlet apparatus for 3 h (20 cycles, each) according to Shah and Alagawadi (2011). The n-hexane extracts were evaporated under reduced pressure to yield different residues. Twelve plants were used to give six HTL and ITL samples.

**Metabolites isolation and identification**

The n-Hexane extracts were investigated using gas-chromatography-mass spectrometry (GC-MS) for qualitative analysis and gas chromatography-flame ionization detection (GC/FID) quantitative analysis. The GC conditions involved the use of Shimadzu-QP-2010 machine equipped with a capillary column (DB-5 ms 30 m x 0.25 mm I.D., 0.25 µm). The chromatograph was programmed for an initial temperature of 50°C for 2 min followed by a 5°C/min temperature ramp to 280°C. The final temperature was maintained for 4 min. Injector and detector temperatures are maintained at 250 and 280°C, respectively. The initial head pressure of the carrier gas (He) was 90 kPa and a split injection system (ratio 1:20) was used. In GC/MS, the capillary column was directly coupled to a quadrupole mass spectrometer (Shimadzu model QP2010S), the ionization mode was electron impact (EI) and ionization energy was 70 eV.

**Components identification and percentage area calculation**

Different separated compounds were identified using Kovat’s Retention indices (RI) calculated with respect to a set of co-injected homologous series of saturated hydrocarbon standards (C8 to C40, Sigma, UK). Compounds were identified by comparing their spectral data and RI with Wiley Registry of Mass Spectral Data 9th edition/NIST Mass Spectral Library (2011), and literature data (Adams, 2007). Some of the compounds were identified using authentic samples and those compounds are marked in Table 2. Calculations of peak percentage areas, based on FID response, are as follow:

\[
\text{Percent area of peak} = \left( \frac{\text{The FID peak area}}{\text{the sum of all the FID peaks areas}} \right) \times 100
\]

Most of non-identified components are present as traces with relative abundances of less than 0.1%. The most important constituents identified in the n-hexane fractions analyzed are listed in Table 2. The percent area ratio was calculated for each component and displayed in Table 2. This ratio indicates that this component increased in concentration due to the virus attack (that is, the ratio will be more than 1) or decreased due to the attack (the ratio will be less than 1).

**Statistical analysis**

Six samples were used for both HTL and ITL (n=6), respectively and each sample was injected in triplicate. Quantitative values are expressed as mean ± standard error of mean of percentage areas and significance difference was determined using unpaired student-sample-t-test performed using SPSS statistical package version (SPSS for Windows, Version 11.5, SPSS Inc., Chicago, IL). P<0.05 was considered significant.

**RESULTS**

**Detection of TYLCV**

DNA of the expected sizes, 575 and 316 bp of
Table 2. Comparison between the n-hexan extract constituents of healthy tomato leaves (HTL) and infected tomato leaves (ITL) TYLCV. The major 46 components the n-hexan fraction were compared. The percentages of infected (ITL) to healthy (HTL) fraction of the same compound of both n-hexan fractions are in the ITL/HTL ratio column. The Quantitative values are expressed as mean ± SEM of six independent infected and healthy leaf samples (n=6).

<table>
<thead>
<tr>
<th>S/N</th>
<th>Compound name</th>
<th>Rt (min)</th>
<th>RI</th>
<th>Area percentage</th>
<th>Significance difference</th>
<th>ITL/HTL ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>HTL</td>
<td></td>
<td>ITL</td>
</tr>
<tr>
<td>1</td>
<td>Octane</td>
<td>2.444</td>
<td>801</td>
<td>0.250 ± 0.014</td>
<td>Yes</td>
<td>3.327</td>
</tr>
<tr>
<td>2</td>
<td>1,3,5-Bisabolatrien-7-ol</td>
<td>21.151</td>
<td>1604</td>
<td>0.117 ± 0.014</td>
<td>Present in ITL only</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>E-Bisabol-11-ol</td>
<td>23.701</td>
<td>1666</td>
<td>0.057 ± 0.004</td>
<td>Present in ITL only</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Palmitic acid**</td>
<td>24.268</td>
<td>1680</td>
<td>0.116 ± 0.007</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Z2,6E-Famesol</td>
<td>26.012</td>
<td>1723</td>
<td>0.162 ± 0.009</td>
<td>Yes</td>
<td>1.289</td>
</tr>
<tr>
<td>6</td>
<td>E,6E-Farnesal</td>
<td>26.268</td>
<td>1729</td>
<td>0.131 ± 0.018</td>
<td>Yes</td>
<td>2.711</td>
</tr>
<tr>
<td>7</td>
<td>E-β-Santalol</td>
<td>26.493</td>
<td>1735</td>
<td>0.309 ± 0.015</td>
<td>Present in HTL only</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>6R-7R-bisabolone</td>
<td>27.609</td>
<td>1762</td>
<td>0.383 ± 0.025</td>
<td>Yes</td>
<td>1.499</td>
</tr>
<tr>
<td>9</td>
<td>β-Bisabolinalen</td>
<td>27.771</td>
<td>1766</td>
<td>0.241 ± 0.029</td>
<td>Present in ITL only</td>
<td></td>
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<tr>
<td>10</td>
<td>2E,6E-Methyl famesoate</td>
<td>28.38</td>
<td>1781</td>
<td>8.379 ± 0.521</td>
<td>Yes</td>
<td>0.853</td>
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<tr>
<td>11</td>
<td>Z-β-Santalol acetate</td>
<td>30.042</td>
<td>1818</td>
<td>0.095 ± 0.011</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>Phytol</td>
<td>30.953</td>
<td>1837</td>
<td>1.645 ± 0.179</td>
<td>No</td>
<td>1.207</td>
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<tr>
<td>13</td>
<td>Linolenic acid methyl ester**</td>
<td>31.275</td>
<td>1843</td>
<td>3.786 ± 0.168</td>
<td>Yes</td>
<td>2.449</td>
</tr>
<tr>
<td>14</td>
<td>Bisabolatrien-1-ol-4-one</td>
<td>31.377</td>
<td>1845</td>
<td>1.643 ± 0.149</td>
<td>No</td>
<td>1.135</td>
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<tr>
<td>15</td>
<td>2E,6E-Famesyl acetate</td>
<td>31.452</td>
<td>1847</td>
<td>1.072 ± 0.087</td>
<td>No</td>
<td>1.159</td>
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<td>16</td>
<td>Z.Z-Famesyl acetone</td>
<td>31.649</td>
<td>1851</td>
<td>1.176 ± 0.245</td>
<td>Yes</td>
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<td>17</td>
<td>Stearic acid**</td>
<td>31.857</td>
<td>1855</td>
<td>2.288 ± 0.196</td>
<td>Yes</td>
<td>1.446</td>
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<tr>
<td>18</td>
<td>E-β-Santalol acetate</td>
<td>32.243</td>
<td>1863</td>
<td>Trace</td>
<td>Present in ITL only</td>
<td></td>
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<tr>
<td>19</td>
<td>5Z,9E-Famesyl acetone</td>
<td>33.903</td>
<td>1897</td>
<td>Trace</td>
<td>Present in ITL only</td>
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<td>20</td>
<td>cis-9-Hexadecenal</td>
<td>34.775</td>
<td>1915</td>
<td>0.095 ± 0.012</td>
<td>No</td>
<td>0.905</td>
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<td>21</td>
<td>Isophytol</td>
<td>36.602</td>
<td>1952</td>
<td>0.504 ± 0.024</td>
<td>Present in ITL only</td>
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<tr>
<td>22</td>
<td>E-β-ionone</td>
<td>37.08</td>
<td>1962</td>
<td>1.415 ± 0.092</td>
<td>Yes</td>
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<td>23</td>
<td>Stearylvialdehyde</td>
<td>37.518</td>
<td>1971</td>
<td>0.546 ± 0.013</td>
<td>Yes</td>
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<td>24</td>
<td>Kaur-15-ene</td>
<td>38.994</td>
<td>2001</td>
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<td>Yes</td>
<td>0.578</td>
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<td>25</td>
<td>Palmitaldehyde</td>
<td>39.657</td>
<td>2025</td>
<td>1.567 ± 0.145</td>
<td>Yes</td>
<td>1.616</td>
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<td>26</td>
<td>Kaurene</td>
<td>40.108</td>
<td>2041</td>
<td>1.234 ± 0.098</td>
<td>Yes</td>
<td>1.772</td>
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<td>27</td>
<td>6Z,10E-Pseudo phytol</td>
<td>40.292</td>
<td>2048</td>
<td>Trace</td>
<td>Present in ITL only</td>
<td></td>
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<tr>
<td>28</td>
<td>6E,10E-Pseudo phytol</td>
<td>40.642</td>
<td>2060</td>
<td>Trace</td>
<td>Present in ITL only</td>
<td></td>
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<tr>
<td>29</td>
<td>Humulene epoxide</td>
<td>41.524</td>
<td>2092</td>
<td>0.067 ± 0.084</td>
<td>Yes</td>
<td>Present in ITL only</td>
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<tr>
<td>30</td>
<td>Methyl linoleate</td>
<td>41.796</td>
<td>2102</td>
<td>0.680 ± 0.009</td>
<td>No</td>
<td>0.999</td>
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<td>31</td>
<td>Squalene</td>
<td>41.9</td>
<td>2105</td>
<td>0.602 ± 0.041</td>
<td>Yes</td>
<td>0.852</td>
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<td>32</td>
<td>Stearic acid methyl ester**</td>
<td>42.262</td>
<td>2118</td>
<td>1.094 ± 0.112</td>
<td>Yes</td>
<td>1.444</td>
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<td>33</td>
<td>linoleic acid**</td>
<td>42.683</td>
<td>2134</td>
<td>0.999 ± 0.128</td>
<td>No</td>
<td>0.837</td>
</tr>
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<td>34</td>
<td>Oleic acid**</td>
<td>42.857</td>
<td>2140</td>
<td>0.486 ± 0.039</td>
<td>Present in ITL only</td>
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<td>35</td>
<td>Abieta-(8(14),13(15)-diene</td>
<td>43.244</td>
<td>2154</td>
<td>4.473 ± 0.418</td>
<td>Yes</td>
<td>0.743</td>
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<tr>
<td>36</td>
<td>Phytol acetate</td>
<td>44.951</td>
<td>2215</td>
<td>1.808 ± 0.147</td>
<td>Yes</td>
<td>0.829</td>
</tr>
<tr>
<td>37</td>
<td>3-β-Stigmast-5-en-3-ol**</td>
<td>46.242</td>
<td>2261</td>
<td>3.894 ± 0.221</td>
<td>Yes</td>
<td>1.850</td>
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<tr>
<td>38</td>
<td>Dehydro abietal</td>
<td>46.474</td>
<td>2270</td>
<td>0.812 ± 0.061</td>
<td>No</td>
<td>0.962</td>
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<tr>
<td>39</td>
<td>β-Stitsterol**</td>
<td>47.138</td>
<td>2293</td>
<td>20.256 ± 1.387</td>
<td>Yes</td>
<td>0.790</td>
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<tr>
<td>40</td>
<td>Abietal</td>
<td>47.459</td>
<td>2305</td>
<td>0.629 ± 0.025</td>
<td>Yes</td>
<td>Present in ITL only</td>
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<tr>
<td>41</td>
<td>Campesterol</td>
<td>48.646</td>
<td>2347</td>
<td>0.970 ± 0.087</td>
<td>Yes</td>
<td>Present in ITL only</td>
</tr>
<tr>
<td>42</td>
<td>Methyl dehydro abietate</td>
<td>48.939</td>
<td>2358</td>
<td>3.171 ± 0.278</td>
<td>Yes</td>
<td>0.793</td>
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<td>43</td>
<td>4-epi-Abietal</td>
<td>49.646</td>
<td>2383</td>
<td>3.346 ± 0.391</td>
<td>No</td>
<td>1.041</td>
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<td>44</td>
<td>Stigmast-4-en-3-one</td>
<td>51.642</td>
<td>2566</td>
<td>2.282 ± 0.222</td>
<td>Yes</td>
<td>0.971</td>
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<tr>
<td>45</td>
<td>neo-Abietol</td>
<td>51.997</td>
<td>2605</td>
<td>0.566 ± 0.054</td>
<td>Yes</td>
<td>Present in ITL only</td>
</tr>
<tr>
<td>46</td>
<td>Lupeol**</td>
<td>52.918</td>
<td>2704</td>
<td>6.604 ± 0.428</td>
<td>No</td>
<td>0.909</td>
</tr>
</tbody>
</table>
Table 2. Contd.

<table>
<thead>
<tr>
<th>Total fraction percentage area</th>
<th>HTL 80.400</th>
<th>ITL 82.640</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sesquiterpenes</td>
<td>24.701</td>
<td>25.945</td>
</tr>
<tr>
<td>Fatty acids</td>
<td>9.118</td>
<td>11.656</td>
</tr>
<tr>
<td>Sterols and triterpenes</td>
<td>38.068</td>
<td>34.607</td>
</tr>
<tr>
<td>Abietic acid derivatives</td>
<td>10.938</td>
<td>11.154</td>
</tr>
<tr>
<td>Bisaboline derivatives</td>
<td>1.703</td>
<td>2.440</td>
</tr>
<tr>
<td>Phytol derivatives</td>
<td>3.544</td>
<td>5.155</td>
</tr>
</tbody>
</table>

Rt, Retention time; RI, retention index. Trace: concentration less than 0.1%. *Significance difference is between H and I percentage area for each component, determined using unpaired student-sample-t-test (p<0.05). ** Identified using standards compounds. #The ITL/HTL ratio indicate if this component increased in concentration due to TYLCV attack (implying the ratio is more than 1) or decreased due to the attack (implying the percentage is less than 1).

Figure 2. Agarose gel electrophoresis of PCR products using the degenerate primers AVcore/ACcore. M, 100 bp DNA ladder (Promega); from 1 to 6, six tomato samples showed symptoms of TYLCV infection; 7, healthy tomato sample; 8, negative control.

Figure 3. Agarose gel electrophoresis of PCR products using the primers TYv2664/TYc138. M, 100 bp DNA ladder (Promega); from 1 to 3, three tomato samples showed symptoms of TYLCV infection and gave positive reaction with degenerate primers AVcore/ACcore.

begomoviruses and TYLCV, respectively, were amplified from symptomatic tomato plants using the primer pairs AVcore/ACcore (Figure 2) and TYv2664/TYc138 (Figure 3), respectively.

Analysis of n-hexane fraction

This study was performed to compare the lipid-soluble metabolic pool of compound in healthy tomato leaves to that of the TYLCV-infected leaves, which allows the identification of the newly synthesized metabolites, or those, which differ in concentration because of the virus attack.

The gas chromatographic analysis of the n-hexane fraction (Figure 4 and Table 2) resulted in the separation of 84 components, 46 of which were identified, representing 80.40 and 82.64% of the total fraction contents of the HTL and ITL sample, respectively. Thirty-seven common compounds were identified between the two extracts; however eight compounds were produced uniquely in the infected tissue extracts and a compound was identified in the healthy samples only indicating its total consumption during the virus infection process. Similarly, five compounds were found in trace in the healthy tissues indicating that 14 compounds were more or less produced due to the virus infection (Table 2). The concentrations of the 12 compounds increased in the infected tissues in relation to the healthy ones. Nevertheless, ten common compounds decreased in concentration when the tissue was infected and both cases can be recognized from ITL/HTL ratio in Table 2. The change in concentration in ten common compounds was considered insignificant, and thus those compounds are considered to have no change in concentration due to the virus infection.
Figure 4. Aligned and expanded GC-MS chromatograms of the n-hexane fraction extracted from healthy Tomato leaves (HTL, lower in black) and TYLCV-infected leaves (ITL, upper in pink). To follow up the retention time on the chromatograms (from 1 to 55 min), follow the numbers from 1 to 4.
DISCUSSION

In the present study, the detection of begomoviruses in tomato was done and the results agreed with that of Alhudaib et al. (2014) and Rezk (2016) who used the degenerate primers of AVcore and ACcore to detect the begomoviruses in infected tomato samples in Saudi Arabia. Also, a leaf curl disease with symptoms typical of begomoviruses was observed in bean (P. vulgaris) at the Main Research Farm of the Indian Institute of Pulses Research, Kanpur, India (Kamaal et al., 2015) whereas Herrera-Vásquez et al. (2016) detected begomoviruses in commercial tomato plots using both production systems (open field and greenhouse) in Panama with different degenerate primers.

Just et al. (2014) stated that, imported tomato fruits infected with Tomato yellow leaf curl virus (TYLCV) were identified on the market in northern Europe using paper-based FTA Classic Cards (Whatman), PCR and partial DNA sequence analysis.

Impact of the metabolites pool changes due to the virus attack

Analysis of Table 2 discloses many classes of lipid-soluble components in the healthy and infected tissues of tomato leaves with different ratios. Sterols and triterpenes are the main class of compounds, which can be identified in the metabolites of both healthy and virus-infected leaves (Table 2). Although there was total decline in sterols and triterpenes concentration, 3-β-Stigmaster-5-en-3-ol increased by 1.85 folds due to the virus infection. The concentration of β-Sitosterol represents nearly 25% of the total lipid-soluble fraction and it declined in the virus-infected leaves to reach 0.7 of its amount in healthy tissues. Accumulation of stigmasterol is a characteristic for plants during pathogens infection (Griebel and Zeier, 2010). Stigmasterol is chemically produced from β-sitosterol through C22 desaturation and this can explain the consumption of β-sitosterol and the increase in stigmastanol concentration due to the virus attack. Campesterol is produced uniquely in the virus-infected tomato leaves tissues. Campesterol are the precursor of steroidal phytohormones called brassinosteroids (Schaller, 2003), which are of vital role in plant defense mechanism against any pathogen attack (Choudhary et al., 2012). The increase in Campesterol concentration due to the virus attack might indicate an effect applied by the virus to weaken the plant defense through prevention of the production of brassinosteroids. On the other hand, a low ratio of Campesterol to sitosterol is needed for high plant cell membrane integrity and functionality (Schaeffer et al., 2001), though this ratio was affected by the production of Campesterol in the virus infected tissues which could lead to interruption and weakness of cell member, that is, the curling effect.

Sesquiterpenes are another class of compounds which are found in high ratio in the lipid soluble metabolic pool of tomato leaves. Although the total sesquiterpene compounds concentration insignificantly changed due to the virus infection, many individual components were produced exclusively or showed meaningful increase in concentration due to infection. Sesquiterpenes in tomato leaves metabolic pool can be divided into three main classes; Fernesols, bisabolines and abietic acid derivatives. Fernesols represent around 11.6% of the total lipid-soluble fraction of tomato leaves and although this whole ratio did not change due to the virus infection, many compounds were individually increased or decreased. The most abundant fernesol-type sesquiterpene in tomato leaf is methyl farnesoate, which represents 9.82% in healthy tissue, and this percentage decreased to 8.37 in the infected tissues. Fernesols are insect hormones (Nagaraju, 2007) and prevent fungal mycelia development with slight anti-fungal properties (Hornby et al., 2001). Bisabolene-type compound are another type of component that belong to the sesquiterpenes pool. Bisabolenes are recognized as sexual pheromones (Brézot et al., 1994; Lu and Teal, 2001) and thus they, together with Fernesols could be emitted by the plant to attract insects (War et al., 2012) as a way to resist the virus attack. Abietic acid derivatives is another class of sesquiterpenes, which is represented in the total lipid-soluble fraction tomato leaves extract by nearly 11%. This ratio did not change significantly between the infected and non-infected leave tissues. Abietic acid and its derivatives are diterpenes which are known for their role in plant defense mechanisms and are recognized for their tissue healing properties and pathogen trapping capabilities (Costa et al., 2016).

Phytols are acyclic diterpene alcohols, which decreased due to the virus attack on the plant. Although phytol and phytol acetate concentration decreased, isophytol was produced uniquely in the attacked tissues. The production of isophytol can explain the decrease in concentrations in phytol and its acetate. The role of isophytol as a production in the virus infected tissues is not clear and needs further investigation. Fatty acids have been identified in both the HTL and ITL samples with 9.11 and 11.6%, respectively. The main fatty acids found in both extracts was stearic acids and its derivative; stearylaldehyde and its methyl ester. The concentration of stearylaldehyde has dramatically decreased in ITL tissues to reach 0.2% of its original concentration in HTL. However, the concentration of stearic acid and its methyl ester has increased by 1.4% for both compounds due to the virus attack and this could explain the decline in stearylaldehyde concentration.

Conclusion

TYLCV has been identified in local area of Al-Ahsaa region, Eastern province of Saudi Arabia using means of
specific PCR primers. The fat-soluble metabolites resulting from the virus attack on the tomato plants has been revealed using means of GC/MS and quantified using GC/FID. Forty-six compounds were separated in both healthy and virus-infected leaf tissues, among which eight compounds were exclusively detected in the infected samples and only one compound was consumed and thus recognized only in the healthy samples.

CONFLICT OF INTERESTS

The authors have not declared any conflict of interests.

ACKNOWLEDGEMENT

The authors are thankful to Dr. Mohamed S. Al-Saikhan, the Supervisor of Central Labs, College of Agricultural and Food Sciences, King Faisal University for assistance during scientific experiments.

REFERENCES


Full Length Research Paper

Polymerase chain reaction amplification of 16S rDNA from three nosocomial bacterial isolates in Kaduna State, Northern Nigeria

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A wide variety of opportunistic pathogens has been detected in hospital surfaces. Medical center surfaces can serve as reservoirs of pathogenic bacteria. Among this pathogens, Pseudomonas species are one of the leading causes of nosocomial infections, frequently found in hospital environments. Polymerase chain amplification system remains one of the best methods for the rapid detection of low numbers of pathogenic bacteria with core references to nucleic Acid content. However, there is a limited researches focusing on these techniques to examine the molecular content of nosocomial bacteria. The present investigation examines types and strains of bacteria present in indoor air of wards, fomites and surgical tools of three prominent hospitals namely, Ahmadu Bello University Teaching Hospital Zaria, Hajiya Gambo Sawaba Hospital Zaria City and Barau Dikko Teaching Hospital Kaduna using standard methods. Preliminary Grams reaction and biochemical characterization was done according to standard methods, DNA extraction precede PCR amplification, probable organisms include Pseudomonas aeruginosa, Corynebacterium sp., P. aeruginosa, Bacillus sp., Klebsiella pneumonia, Neisseria sp., Staphylococcus aureus and Staphylococcus epidermidis. Out of all the isolates that were of public health concern, Neisseria sp., S. aureus and P. aeruginosa are the most prevailing isolates. A strain of P. aeruginosa was observed to give a DNA sequence. P. aeruginosa was the bacteria isolates sequenced and it showed 100% similarity having the id query: 86603, when blast using National Center for Biotechnology Information (NCBI). In general, patterns were specific at either the genus level or the species level. This research has been able to show that PCR is a promising fast method for the identification of nosocomial microorganisms.

Key words: Polymerase chain reaction, nosocomial bacterial, hospital and DNA.

INTRODUCTION

Nosocomial infection is one of the most common complications within health care facilities, certain studies have reported outbreaks resulting from contaminated hospital environments, it is called nosocomial infection if it develops 72 h after admission to the hospital (Akihiro et al., 2017). These infections are more dangerous than
other infections because they are caused by bacteria with a high resistance to antibiotics.

Health care facilities provide an environment conducive to exposure and transmission of bacteria; infections caused by these bacteria are an important cause of increased morbidity, mortality and health care costs worldwide (Daniel and Michael, 2015). Nowadays, although modern antibiotics have improved; still sometimes the treatment is difficult and causes morbidity and mortality to patients. Many outbreaks of nosocomial infections have come from reservoirs of pathogens in the inanimate hospital surfaces. The contribution of the environment surfaces remains an important factor in nosocomial infection. It has been reported that majority of the patient acquire this infection while staying in hospital (Ananthan et al., 2011).

The major causative agents in the Western world comprise Gram-positive Staphylococcus aureus, Enterococcus spp., Streptococcus spp., Gram-negative Escherichia coli, Enterobacter spp., Proteus mirabilis, Klebsiella spp. In Africa, P. aeruginosa are in increasing proportions due to intensive care treatment (Souza et al., 2015). Pseudomonas sp. are considered opportunistic pathogens that causes opportunistic infection, they are commonly found in nature (soils, water, plants and animals) and water treatment systems, thus demonstrating their adaptation to environments with low nutrient concentration and over a large temperature range (between 4 and 42°C) (Frickmann et al., 2013). Conventional examination of a bacterial infection mainly relies on culture-based techniques. These cultivations usually yield diagnostic results in days or in some cases up to a week after sampling. Furthermore, cultivation of bacteria is not always successful under laboratory conditions (Kerremans et al., 2008; Yoshimura et al., 2011). Such failures may occur due to unsuitable culturing conditions and methods for the bacterial species under study. Alternatively, the particular patient under investigation may have received antimicrobial therapy before sampling.

Molecular methods based on nucleic acid amplification and hybridization aim to circumvent these problems and hasten diagnostic procedures. In such methods, the pathogen is simultaneously detected and identified, which results in more rapid diagnosis than those obtained by conventional culturing methods and obviates the need for additional culture tests (Akihiro et al., 2017). The purpose of this research was to use molecular techniques as alternative methods for strains level identification of nosocomial bacterial found in indoor air of wards, fomites and surgical tools of three prominent hospitals namely, Ahmadu Bello University Teaching Hospital Zaria, Hajiyam Gambo Sawaba Hospital Zaria City and Barau Dikko Teaching Hospital Kaduna, Northern Nigeria.

MATERIALS AND METHODS

Synopsis of the study location

Kaduna with location Coordinates 10°20’N 7°45’E, and 10.333°N 7.750°E usually referred to as Kaduna state to distinguish it from the city of Kaduna, is a state in Northwest Nigeria. It is the capital of Kaduna with a population of over 6,066,562 (2006 Census). The selected hospitals are the three major government hospitals in the state and the justification for selection of these hospitals was based on high patronization.

Isolation and molecular characterization of bacteria

Bacteria were isolated from indoor air of wards, fomites and surgical tools of Ahmadu Bello University Teaching Hospital Zaria, Hajiyam Gambo Sawaba Hospital Zaria City and Barau Dikko Teaching Hospital Kaduna using standard methods described by Kerremans et al. (2008) and Cheesbrough (2006). The bacteria from bed sheets, pillow cases, nurses’ desk and surgical tools were isolated using sterile swab sticks while those from the air in surgical wards and main wards were isolated using exposed plate technique.

Media preparation

All the media used in this research work (nutrient agar, nutrient broth, mannitol salt agar and blood agar) were prepared according to manufacturer’s instructions.

Sample collection

Sampling sterile swab sticks were immediately introduced into nutrient broth and taken to Kaduna State University Microbiology Laboratory for incubation and further bacteriological analysis. In the exposed plate technique, the agar plates were opened and exposed to the indoor air of the wards for about 5 min. After 5 min of exposure the plates were covered again and taken to Microbiology laboratory for incubation. Both the agar plates and the broth cultures were incubated at 37°C for 24 h. After incubation, the bacterial colonies that showed positive growth were sub-cultured, gram stained and viewed under the microscope using oil immersion x100 objectives lens (Kerremans et al., 2008).

Biochemical characterization and identification of the nosocomial isolate

Bacteria isolates extracted were characterized and identified after studying their Gram reaction as well as cell micro morphology. Other tests performed were spore formation, motility and catalase production, citrate utilization, fermentative utilization of glucose, indole production, methyl red- Voges Proskauer reaction, urease and coagulate production, starch hydrolysis, production of H2S from triple sugar iron (TSI) agar and sugar fermentation. The test were carried out according to the methods described (Cheesbrough, 2006; Adeoye, 2007; Agwung-Fobellah and Kemajou, 2007; Ochei

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Molecular characterization of species of bacteria

The bacteria species characterized using biochemical methods were subjected to strain level identification using molecular approach. These involve extracting the DNA using standard protocols, amplifying the DNA using PCR, sequencing the gene of interest using next generation sequencing (NGS).

DNA extraction using phenol chloroform method

DNA was extracted using phenol chloroform standard method as described by Psfidi et al. (2010). 200 μl of the bacterial cells were added in a 1.5 ml Eppendorf tube, 400 μl of lysis buffer (Tris–HCL) and 10 μl of proteinase k was added to the 1.5 ml tube. The tube was vortexed and placed on heat block at 65°C for 1 h while vortexing at interval. 400 μl of phenol chloroform was added and vortexed briefly, it was centrifuged for 10 min at 13,000 rpm to separate the phases. The upper layer was carefully removed with a micro pipette, 400 μl of chloroform was added and vortexed, and was then centrifuged for 5 min at 13,000 rpm. The upper layer was carefully removed, 1000 μl of absolute ethanol and 40 μl of 3 M sodium acetate and mixed by inverting the tube several times.

The tubes containing pure DNA were incubated at -20°C overnight. The tubes were centrifuged for 5 min at 14,000 rpm using cool centrifuge, the upper layer was carefully removed using micro pipette, 400 μl of 70% ethanol was added to the precipitate, the tubes were centrifuged for 5 min at 14,000 rpm using cold centrifuge, the upper layer was carefully removed using micro pipette in order to remove all traces of ethanol, the DNA was then allowed to dry by leaving the tubes open for 20 min at room temperature and 50 μl of sterile water was added, vortexed and kept at -20°C.

PCR amplification of 16S rDNA using conventional PCR

The extracted DNA from different species of bacteria was quantified using Nano drop. Thereafter, the quantified genomic DNA was placed in a tube containing master mix and primer and the tube was spun for 30 s and introduced into a thermocycler. The thermocycler was operated based on initial denaturation (95°C for 5 min), 25 cycles of denaturation (94°C for 1 min), annealing (52°C for 1 min), extension (72°C for 1 min and final extension (72°C for 7 min) according to Psfidi et al. (2010). 16S rDNA forward sequence 5’GGACTACGGGTATCTAAAT 3’ and reverse sequence 3’AGAGTTTGATCCAGG 5’ after the PCR in the thermocycler, the amplified PCR products of expected size 789 bp were confirmed by agarose gel electrophoresis system. The amplified PCR products was run in 1.5% agarose gel electrophoresis stained with ethidium bromide and was viewed under ultra violet (UV) light (Smith et al., 2003; Black and Foarde, 2007).

Sequencing of the gene

The amplified genes of interest were documented, cut and freeze dried. The freeze dried samples of DNA were sent to Macrogen U.S.A laboratory for sequencing. The genomic sequence data was appropriately analyzed using bioinformatics tools, sequence identification was performed using NCBI Basic Local Alignment Search Tool (BLAST) algorithm, similar sequences were downloaded and aligned while phylogenetic tree was drawn with MEGA 7 software (Kumar et al., 2016).

RESULTS

A total number of 66 bacteria were isolated from three hospitals in Kaduna State namely; Ahmadu Bello University Teaching Hospital, Zaria, Hajiya Gambo Sawaba Hospital, Zaria City and Barau Dikko Teaching Hospital, Kaduna (Table 1). Twenty-two (22) samples were collected from each of the hospitals for preliminary antibiotic resistance potency. Observation shows that only 20 of the bacterial isolates were recorded as multi drugs resistant which includes six (6) from Ahmadu Bello University Teaching Hospital, Zaria, eight (8) from Hajiya Gambo Sawaba Hospital, Zaria City and six (6) from Barau Dikko Teaching Hospital, Kaduna. The morphological and biochemical characterization of nosocomial bacterial isolates from hospital environments was presented in Table 1. Investigation revealed that S. aureus occurred as the highest predominant bacterial isolate with the percentage occurrence of 10 (50%) followed by P. aeruginosa with 4 (20%) and Bacillus sp. had 2 (10%), Corynebacterium sp., Klebsiella pneumoniae, Neisseria sp. and Staphylococcus epidermidis all recorded 1 (5%) appearances (Table 1).

DISCUSSION

A number of morphological and biochemical parameters have been used to facilitate the determining of the identities of nosocomial bacteria and other hospital acquired infections (Ateba and Mbewe, 2011). Even though selective and differential media has been used to aid the identification of the bacterial species, yet the sensitivity of these protocols might not be very reproducible between laboratories. Generally, selective and differential media rely on some structural or metabolic property of the species that is preferentially selected. It is highly recommended that they should be combined with confirmatory biochemical and morphological tests.

Table 1 show the morphological and biochemical characteristics of the 20 isolated multidrug resistant nosocomial bacterial (MDRNB). The present study (morphological observation) revealed the cell shapes of 40.00% of the overall isolated microorganisms possessed rod shape under the microscope while 60.00% were found to be cocci in shapes. This is a good characteristic of some organisms but not enough to categorize, hence, all isolates were subjected to grams staining test. About 25.00% of the isolated microorganism from analyzed samples stained pink to red, indicating them to be Gram negative organisms, while 75.00% of the overall isolates stained blue to purple by retaining the crystal violet dye indicating them to be Gram positive organisms (Berger et al., 1994).
Table 1. Morphological and biochemical features of the selected examined Multi-Drugs Resistant (MDRI) isolate.

<table>
<thead>
<tr>
<th>Isolate code</th>
<th>Cell shape</th>
<th>Gram reaction</th>
<th>Catalase</th>
<th>TSI reaction</th>
<th>SIM reaction</th>
<th>C-U</th>
<th>MR</th>
<th>VP</th>
<th>GL</th>
<th>MAL</th>
<th>MNT</th>
<th>SU</th>
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<th>OF OR H L</th>
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<th>Probable organisms</th>
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<td>NCNCNC</td>
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<td>NC</td>
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<td>OX</td>
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<td>+</td>
<td>YGY+</td>
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<td>Staphylococcus aureus</td>
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CC, Cocci shape; SR, short rod; MLR, medium long rod; LR, long rod; YG, acid and gas production; Y, acid production only; TSI, triple sugar ions; SIM, sulphide,indole,motility tests; CU, citrate utilization test; MR/VP, methyl red and Voges, Proskauer test; GL, glucose; NC, No change; F, fermentative; OX, oxidative; I, indole production; M, motility; H2S, hydrogen sulphide production; MAL, maltose; MNT, mannitol; SU, sucrose; LAC, lactose; nitrate reduction test; OF, oxidation,fermentation Test; HSP, hospital; ABU, Ahmadu Bello University; HGS, Hajiya Gambo Sawaba; BDT, Barau Dikko Teaching Hospital Kaduna.

PCR amplification of 16S rDNA from the three highest prevailing nosocomial bacterial isolates in Kaduna State

Organisms of concern in this study are P. aeruginosa, S. aureus and Neisseria sp. Figure 1 shows the amplified 16S rDNA gene with band size of 789 bp bands from Neisseria sp., P. aeruginosa and S. aureus obtained from some selected hospitals in Kaduna State. To test whether we can discriminate between these isolates, Neisseria sp., P. aeruginosa and S. aureus strains were examined for their electrophoretic mobility patterns in PCR amplification (Figure 1). Although the observed differences were small, PCR amplification was capable of distinguishing all the isolates from one and other (Lee et al., 2009). For the Pseudomonas species, the observed electrophoretic mobility patterns showed light differences between Staphylococcus and Neisseria sp. (Figure 1). Because of the closed mobility relativity of all these on electrophoresis gel, these data suggest species-specific patterns for identification. To evaluate the applicability of PCR amplification as a general tool for the identification of bacteria, the amplified 16S rDNA products from the three nosocomial bacterial isolates were sequenced according to Cole et al. (2009) and the result obtained is shown in Figure 2. Out of the three bacterial isolates analyzed, only one which is P. aeruginosa gave a DNA sequence. P. aeruginosa was the bacterial isolates sequenced and it showed 100% similarity.
Figure 1. PCR amplification of 16S rDNA gene in three nosocomial bacterial isolates from some selected hospitals in Kaduna State. Lane 1, DNA marker; lane 2, Neisseria sp.; lane 3, P. aeruginosa; lane 4, S. aureus; lane 5, negative control.

Figure 2. Sequence of bacteria isolate (Pseudomonas aeruginosa).

having the id query: 86603, when using BLAST National Center for Biotechnology Information (NCBI). The red bullet indicates the sequenced P. aeruginosa (Figure 3). In general, patterns were specific at either the genus level or the species level.

However, we opined that in order to overcome lane-to-lane and gel-to-gel differences, PCR resolution was improved by the addition of an internal DNA marker for each sample lane. This makes it feasible to precisely compare relative migration times. This research has been able to show that PCR is a promising fast method for the identification of microorganisms. Also, the use of PCR with universal primers and Single Strand Conformational Polymorphism (SSCP) patterns as an identification method can generally be applied to a wide range of nosocomial bacteria without the need of a large panel of probes.

In order to intensify our research, we added an option for molecular identification of methicillin resistant S. species by including the methicillin resistance gene mecA in the assay. The identification was based on PCR amplification as shown in Figure 4. mecA gene was amplified in S. aureus which showed no visible band size. The presence of coagulase negative staphylococcal DNA other than that associated with S. epidermidis was then detected by genus-specific probes.

Conclusion

Molecular and genotypic characterization of bacteria is advantageous when compared to biochemical and phenotypic methods of characterization and identification of nosocomial bacterial. The latter require a prolonged cultivation period for the suspected bacteria and pure bacterial cultures for various biochemical assays. It is therefore concluded that broad-range PCR amplification with subsequent hybridization on a microarray is a rapid diagnostic tool in identifying causative agents of bacterial infections in various specimens from normally sterile site of the hospital environment to medical and surgical tool surfaces.
CONFLICT OF INTERESTS

The authors have not declared any conflict of interests.

ACKNOWLEDGEMENT

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The nitrogen-fixing *Frankia* significantly increases growth, uprooting resistance and root tensile strength of *Alnus formosana*

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Restoration of *Alnus formosana* (Burk.) Makino on landslide areas is important for agroforestry, forestry and soil erosion control in Taiwan. To ensure successful reforestation, *A. formosana* seedlings have to develop strong root system for nutrient and water acquisition as well as anchorage. Inoculating of *A. formosana* with symbiotic nitrogen-fixing actinobacteria *Frankia* may help mitigate drought and nutrient deficiencies on landslide sites. However, the effects of *Frankia* inoculation on growth, root architecture and mechanical properties of *A. formosana* seedlings are not well understood. In this research, a *Frankia* strain AF1 was isolated from actinorhizal nodules of local *A. formosana* and recognized as *Frankia* species, and its influences on growth performance and root mechanical properties of *A. formosana* seedlings were examined and analyzed. The results showed that the inoculated seedlings had significantly larger height and root biomass, longer root length, and more root tip number than that of the non-inoculated controls. Consistently, the inoculated seedlings had statistically significant higher uprooting resistance, root tensile resistance force and tensile strength than the controls. The results reveal that this native *Frankia* strain can promote growth performance, root system architecture, anchorage ability and root tensile strength of *A. formosana*.

**Key words:** Actinorhiza, Betulaceae, inoculation, root morphology, tensile strength, uprooting resistance.

INTRODUCTION

Landslide hazards and their large negative impacts on human lives, economies and infrastructure have become a growing challenge globally (Dai et al., 2002). Due to the fragile geology, steep terrain and torrential rains brought by typhoons, landslides have become one of the most severe disasters in Taiwan. Vegetation landslide engineering has become increasingly important for landslide prevention and rehabilitation in recent years (Chen et al., 2014). In general, trees and forests can play an important role in preventing and rehabilitating landslides. Actinorhizal trees are pioneer species and can improve tree growth and survival on degraded landslide soils (Diagne et al., 2013). *Alnus formosana* (*Alnus formosana* (Burk.) Makino), belonging to the family Betulaceae, is a native nitrogen-fixing actinorhizal woody species, widely distributed throughout the island of...
Taiwan in landslide sites (Liao, 1996). It has high potential for agroforestry, forestry, lumber production and landslide restoration. It can establish symbiosis with nitrogen-fixing *Frankia* and form actinorhizal root nodules in which *Frankia* provides fixed nitrogen to the host plant for growth and development (Lee, 1986; Lin, 1992). Several previous studies have demonstrated that *Frankia* can improve the establishment and growth of *Alnus* in degraded lands (Lefrançois et al., 2010; Santi et al., 2013; Bissonnette et al., 2014; Põlme et al., 2014). Inoculation with *Frankia* significantly increases seedling growth, biomass and root nodules of *Alnus crispa* and *Alnus sieboldiana* (Yamanaka et al., 2005; Quoreshi et al., 2007).

Morphological types of tree root system architecture have been classified into heart system, plate system and taproot system (Stokes and Mattheck, 1996). Stokes et al. (2009) indicated that taproot length, amount of lateral roots and root architecture affects uprooting resistance of plants. Orfanoudakis et al. (2010) showed that inoculation of *Alnus glutinosa* with *Gigaspora rosea* and *Frankia* improves ramous root. Past studies on root morphological characteristics and biomechanical properties of *Alnus* species were focused on *Alnus subcordata* and *Alnus viridis*. In *A. viridis*, the maximum root area ratio (RAR) values were located in the upper 30 cm soil layer and the maximum rooting depth was about 1 m, whereas the root tensile strength decreased with diameter (Bischetti et al., 2005). In *A. subcordata*, the root density, root number and RAR decreased with increasing depth and the maximum rooting depth was 1 m, while the root tensile strength decreased with increasing root diameter (Naghdi et al., 2013). However, there were few studies that tried to investigate the effects of inoculation of *Frankia* on growth, root morphological characteristics and mechanical properties of *A. formosana*. Therefore, the purposes of this research were: (1) to isolate the *Frankia* strain from *A. formosana*, and (2) to examine the influences of *Frankia* inoculation on growth performance, uprooting resistance and tensile strength of *A. formosana* seedlings in order to provide strategy for landslide prevention and erosion control practices. This study focused on the application of nitrogen-fixing *Frankia* to alder seedling production in order to improve seedling growth performance and enhance root mechanical properties, which is important in prevention of landslide hazards.

**MATERIALS AND METHODS**

**Sample collection**

An elite tree of *A. formosana* was selected from the natural forest stand located at Fengliu Township, Chiayi County, Taiwan (219417.59E, 2599775.50N, TWD 97) in October 2014. Actinorhizal nodules were gathered from roots at 6 to 30 cm deep in soil, kept in sealed plastic bags and transported to laboratory in a cold box for *Frankia* strain isolation (Lin, 1992). Cones were also collected from upper crown of the same tree. Cones were sun-dried in trays for seed release. Seeds were extracted, cleaned and freeze-stored in polyethylene bags at -20°C.

**Frankia strain isolation and purification**

Actinorhizal root nodules were washed in distilled water to remove soil particles. Single nodule lobes were cut 2 mm from the tip with scalpel, ultrasonically cleaned, surface-sterilized with 75% ethanol for 10 min, 15% NaOCl for 10 min, and 30% hydrogen peroxide for 10 min. The endosymbiont *Frankia* in nodule lobes were isolated aseptically on QMOD medium (Lalonde and Calvert, 1979).

**DNA extraction, sequencing, and gene sequence similarity analysis**

*Frankia* genomic DNA was extracted with Puregene DNA Purification Kit (QIAGEN, Pleasanton, CA, USA), and subsequently subjected to 1.2% agarose gel electrophoresis. The primers used for PCR of rDNA were primer FGPL2054 (5’-CCGGGTTCCTCCCATTCCG-3’) and primer FGPS998e (5’-GGG GTC CTG AGG GGC T-3’) (Daniel et al., 1999). Then, the amplified samples were analyzed by gel electrophoresis and the particular PCR products were sequenced. The DNA sequences were submitted to NCBI to access GenBank for sequence similarity analysis of the *Frankia* rDNA gene sequences.

**Seedling preparation**

Seeds of *A. formosana* were surface cleaned with tap water, sterilized 2 times with 10% NaOCl solution for 5 min and washed with sterile water, and then germinated in autoclaved peat moss and vermiculite mixtures (1:1, v/v) in October 2015. The wooden boxes (b x w x h, 30 cm x 30 cm x 60 cm) were used for transplanting. The boxes were sterilized with 10% NaOCl solution, and the sandy loam soils collected from the same natural forest stand were autoclaved and then fumigated with 200 g Basamid fumigant per cubic meter of soil, and the soil surface was sealed with polyethylene sheets for 14 days to prevent the toxic gas from escaping. Then, the boxes were filled with the sterilized soils. When seedlings attained a height of 5 cm, they were individually transplanted to the boxes, and watered regularly. Twenty-eight seedlings in boxes were arranged randomly into two individual plastic houses.

**Inoculum preparation**

The isolated *Frankia* strain was successively cultured twice for 30 days. Inocula were prepared by concentrating *Frankia* cultures in sterile 1 ml tubes (15,000 rpm for 10 min at 4°C) using a Hitachi centrifuge (HIMAG Centrifuge CR 15T, Rotor RT15A, Tokyo, Japan). The pellets were then homogenized with glass tissue homogenizer (Wheaton 25802a, Millville, NJ, USA) in sterile BAP medium and sonicated for 1 min using an ultrasonicator (Biologics 150VT, Manassas, VA, USA) on ice, and used as inoculum. The protein concentration of the homogenized inoculum was assayed with the Bradford method (Bradford, 1976). The protein concentration of the *Frankia* inoculum was 2.8±0.02 µg ml⁻¹.

**Inoculation test**

Four weeks after transplanting, 14 plants in one plastic house were inoculated with the isolated *Frankia* strain. A 5 ml suspension of the
strain (protein concentration 2.8±0.02 µg ml⁻¹) was dripped into five small holes near the seedling. The process was duplicated after fourteen days for ensuring high rate of root colonization. Another 14 non-inoculated control seedlings were treated with sterilized water. The boxes of inoculated and control seedlings were placed individually in two separate plastic houses. The seedlings were grown at 26±4°C during daytime and 18±5°C at night time, with 60 to 80% relative humidity, and 1000±200 µmoles photon m⁻² s⁻¹ photosynthetic photon flux density during daytime. Eight months later, the seedlings were sampled for measurements of growth performance, root morphological characteristics, uprooting resistance and tensile strength.

Plant growth performance and root morphological characteristics

After 8 months of cultivation, seven inoculated and seven non-inoculated control plants were randomly chosen, separately. The height and stem-base diameter of seedlings were measured with ruler and caliper. The root systems were carefully excavated by hand with trowel (Böhm, 1979). The root length and root numbers were recorded. Images of seedling roots were captured for analysis of root architecture and morphological characteristics. The root morphological characteristics analysis was conducted using a WinRHIZOPro analysis system (Regent Instruments, Quebec, QC, Canada) (Bouma et al., 2000). However, root volume was evaluated with water displacement technique due to large quantity of roots (Pang et al., 2011). Biomass of leaf, root, stem and root nodule was estimated by drying in a hot air oven at 75°C until a constant weight was obtained. Root functional characteristics, that is, root mass density (g dm⁻³), root length density (m dm⁻³), tissue mass density (g cm⁻³), specific root length (m g⁻¹), and root to shoot ratio were computed (Stokes et al., 2009; Burylo et al., 2012; Gould et al., 2016). Live roots were also collected from sampled seedlings and prepared for subsequent tensile testing.

Vertical uprooting test

After 8 months of cultivation, 7 inoculated and 7 non-inoculated control seedlings were randomly sampled for vertical uprooting test, respectively. The soil material was categorized as sandy loam soil (containing 65.2% sand, 27.4% silt and 7.4% clay). At first, seedling base diameter were recorded. The seedling stem base diameter of seedlings were measured with digital caliper (accuracy of 0.01 mm). The local strain was recognized as AF1. Molecular analysis was conducted using a WinRHIZOPro analysis system (Regent Instruments, Quebec, QC, Canada) (Bouma et al., 2000). However, root volume was evaluated with water displacement technique due to large quantity of roots (Pang et al., 2011). Biomass of leaf, root, stem and root nodule was estimated by drying in a hot air oven at 75°C until a constant weight was obtained. Root functional characteristics, that is, root mass density (g dm⁻³), root length density (m dm⁻³), tissue mass density (g cm⁻³), specific root length (m g⁻¹), and root to shoot ratio were computed (Stokes et al., 2009; Burylo et al., 2012; Gould et al., 2016). Live roots were also collected from sampled seedlings and prepared for subsequent tensile testing.

Root tensile test

After root excavation, live roots of different diameter classes (0 to 1, 1 to 2, 2 to 5, and 5 to 10 mm) at a depth of 30 cm below the soil surface were collected randomly from the sampled seedlings, respectively. The roots were kept in separate sealed plastic bags to prevent drying of root tissues (De Bates et al., 2008) and transported with a cold box (Bischetti et al., 2005). Root samples were then immersed in a 15% ethanol solution at 4°C in order to conserve root tensile strength (Bischetti et al., 2009). Tensile tests were performed in the laboratory using a tensile-testing machine (U-Soft USPT-003, U-Soft Technology Co., Taipei, Taiwan). The load cell (Kyowa LCN-A, Tokyo, Japan; sensor resolution 0.1 N, maximum force 500 N) was connected to a loading recorder and control unit. The data of tensile force and displacement were compiled on a portable computer. A total of 110 root segments were randomly sampled from the inoculated seedlings. Another 110 root segments were also randomly collected from the non-inoculated seedlings. All root segments were washed and cut to 60 mm in length, and the root segments were clamped with sand paper during testing to prevent slippage. Then, the root segments were pulled vertically at a constant speed of 4.7 mm min⁻¹ until the resisting force dropped sharply. The root tensile strength (Tₛ, MPa) was calculated using the following formula (De Baets et al., 2008; Osman et al., 2011; Zhang et al., 2012):

\[
T_s = \frac{4F_{\text{max}}}{\pi d^2}
\]

where \(F_{\text{max}}\) is the maximum force (N) at rupture and \(d\) is the mean root segment diameter (mm) measured at three points, that is, near the upper clamp, halfway and near the bottom clamp, using a digital caliper (accuracy of 0.01 mm).

Data analysis

T-test in SPSS 22.0 software (Chicago, IL, USA) was used for analyzing variations in growth performance and morphological characteristics data between inoculated and non-inoculated control seedlings. The relationships between uprooting resistance, root tensile strength, tensile strength and morphological characteristics were evaluated using Microsoft Excel regression analysis.

RESULTS

Actinobacterial strain isolation and sequencing

The actinobacterial strain was isolated and purified on QMOD medium and classified as AF1. Molecular analysis showed that the rDNA gene sequence of the isolated AF1 has 100% similarity to that of Frankia species genus (Figure 1). The local strain was recognized as Frankia spp. AF1. Inoculation test showed that the isolated Frankia strain can induce nodule development in the roots of A. formosana (Figure 2).

Growth performance

Results of statistical analysis showed that Frankia inoculation significantly promoted growth performance characteristics of A. formosana seedlings. Generally, seedlings inoculated with Frankia developed significantly larger height (50%), tap root length (60%), root biomass (60%), and shoot biomass (46%) than the non-inoculated ones (Table 1).

Root system architecture

Results of the investigation revealed that inoculated A.
Figure 1. Partial sequence of strain AF1 rDNA gene (query), compared to that of Frankia spp. genus (subject).

Figure 2. Actinorhizal nodules formed on the roots of A. formosana inoculated with Frankia strain (bar = 3 cm).
Table 1. Growth performance of A. formosana seedlings inoculated and non-inoculated with Frankia after 8 months of cultivation.

<table>
<thead>
<tr>
<th>Frankia inoculation</th>
<th>Height (cm)</th>
<th>Stem-base diameter (mm)</th>
<th>Taproot length (cm)</th>
<th>Root biomass (g)</th>
<th>Shoot biomass (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inoculated</td>
<td>218.8±5.5a</td>
<td>21.9±3.0a</td>
<td>43.0±7.5a</td>
<td>68.6±11.1a</td>
<td>162.6±26.8a</td>
</tr>
<tr>
<td>Non-inoculated</td>
<td>145.2±4.3b</td>
<td>18.2±2.4a</td>
<td>26.4±3.1b</td>
<td>42.8±14.0b</td>
<td>111.2±20.8b</td>
</tr>
</tbody>
</table>

All values are the mean ± standard error of 7 replicates. Values in the same column followed by different superscript letters significantly differ at 5% significant level.

Figure 3. Root morphologies of A. formosana seedlings after 8 months of cultivation: (a) inoculated with Frankia strain; (b) non-inoculated.

*formosana* plants developed larger root systems than the non-inoculated ones (Figure 3). The taproots of inoculated plants grew to 43 cm deep. In addition, the inoculated plants developed about 85% of the root matrix in the top 40 cm soil, and its lateral roots extended profusely, with many nodules (Figure 3a). Conversely, the taproots of non-inoculated ones grew only to 20 cm deep. Also, the non-inoculated plants developed about 90% of the root matrix in the top 30 cm soil, and its lateral roots grew sparsely, without any nodules (Figure 3b). The types of root architecture of inoculated and non-inoculated A. formosana seedlings were categorized to the heart root system according to Stokes and Mattheck (1996).

WinRHIZO analysis of root morphological characteristics revealed that Frankia inoculation significantly influenced all morphological characteristics. Generally, the inoculated plants developed larger total...
Table 2. Root morphological characteristics of *A. formosana* seedlings inoculated and non-inoculated with *Frankia* after 8 months of cultivation.

<table>
<thead>
<tr>
<th>Frankia inoculation</th>
<th>Total root length (cm)</th>
<th>Root surface area (cm²)</th>
<th>Root volume (cm³)</th>
<th>Root tip number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inoculated</td>
<td>5217.4±379.1a</td>
<td>3120.8±162.0a</td>
<td>498.3±53.9a</td>
<td>6570±265a</td>
</tr>
<tr>
<td>Non-inoculated</td>
<td>3254.2±585.0b</td>
<td>1939.9±153.2b</td>
<td>322.3±16.6b</td>
<td>3823±229b</td>
</tr>
</tbody>
</table>

All values are the mean ± standard error of 7 replicates. Values in the same column followed by different superscript letters significantly differ at 5% significant level.

Table 3. Root functional characteristics of *A. formosana* seedlings inoculated and non-inoculated with *Frankia* after 8 months of cultivation.

<table>
<thead>
<tr>
<th>Frankia inoculation</th>
<th>Root density (kg m⁻³)</th>
<th>Root length density (km m⁻³)</th>
<th>Root tissue density (g cm⁻³)</th>
<th>Specific root length (mg⁻¹)</th>
<th>Root shoot ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inoculated</td>
<td>1.27±0.21a</td>
<td>0.97±0.26a</td>
<td>0.14±0.02a</td>
<td>0.76±0.14a</td>
<td>0.42±0.09a</td>
</tr>
<tr>
<td>Non-inoculated</td>
<td>0.79±0.19b</td>
<td>0.58±0.30b</td>
<td>0.12±0.02b</td>
<td>0.85±0.16b</td>
<td>0.39±0.07a</td>
</tr>
</tbody>
</table>

All values are the mean ± standard error of 7 replicates. Values in the same column followed by different superscript letters significantly differ at 5% significant level.

Table 4. Uprooting resistances of inoculated and non-inoculated *A. formosana* seedlings after 8 months of cultivation.

<table>
<thead>
<tr>
<th>Frankia inoculation</th>
<th>Uprooting resistance (kN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inoculated</td>
<td>1.09±0.40a</td>
</tr>
<tr>
<td>Non-inoculated</td>
<td>0.60±0.17b</td>
</tr>
</tbody>
</table>

All values are the mean ± standard error of 7 replicates. Values in the same column followed by different superscript letters significantly differ at 5% significant level.

In this study, seven uprooting tests were conducted to examine the uprooting resistance of the inoculated and non-inoculated seedlings. The results revealed that the average maximum uprooting resistance of the inoculated seedlings (1.09±0.40 kN) was significantly higher than the non-inoculated ones (0.60±0.17 kN) (Table 4). The uprooting resisting force increased with displacement up to the peak and then decreased sharply as the roots broke (Figure 4). Regression analysis exhibited the significant linear positive correlations between the maximum uprooting resistance force and morphological characteristics, that is, tree height, stem-base diameter, taproot length, root biomass, and shoot biomass (Figures 5, 6, 7, 8, and 9).

**Root tensile strength**

In total, 220 tests were performed to investigate the root tensile strength of the inoculated and non-inoculated *A. formosana* seedlings. Among them, 86 root tensile tests of the inoculated seedlings and 102 root tensile tests of the non-inoculated controls were successful. The average root tensile resistance force of inoculated seedlings (81.1±19.0 N) was significantly higher than that of the controls (68.0±23.5 N). The mean value of root tensile strength of inoculated seedlings (17.45±3.36 MPa) was significantly higher than that of the non-inoculated controls (11.42±1.83 MPa). On the other hand, the average root diameter of non-inoculated seedlings (2.88±1.28 mm) was significantly higher than the inoculated ones (2.42±1.20 mm) (Table 5). Regression analysis revealed that root tensile resistance force increased with increasing root diameter in accordance with a positive power function correlation (Figure 10), whereas the root tensile strength decreased with increasing root diameter in accordance with a negative logarithmic function correlation (Figure 11). Furthermore, the maximum root tensile resistance and tensile strength of inoculated seedlings were significantly higher than that of the non-inoculated controls.

**DISCUSSION**

The results showed that the native symbiotic
Figure 4. Uprooting resistance force-displacement curves for *A. formosana* seedlings inoculated (—) and non-inoculated (—) with *Frankia*.

Figure 5. Maximum uprooting resistance force-tree height relationship for *A. formosana* seedlings inoculated (—) and non-inoculated (—) with *Frankia*. N = 7.
Figure 6. Maximum uprooting resistance force-stem base diameter relationship for *A. formosana* seedlings inoculated (—) and non-inoculated (- -) with *Frankia*. N = 7.

Figure 7. Uprooting resistance force-taproot length relationship for *A. formosana* seedlings inoculated (—) and non-inoculated (- -) with *Frankia*. N = 7.
Figure 8. Maximum uprooting resistance force-root biomass relationship for *A. formosana* seedlings inoculated (—) and non-inoculated (---) with *Frankia*. N = 7.

Figure 9. Maximum uprooting resistance force-shoot biomass relationship for *A. formosana* seedlings inoculated (—) and non-inoculated (---) with *Frankia*. N = 7.
Table 5. Average root diameter, tensile resistance force and tensile strength of inoculated and non-inoculated A. formosana seedlings after 8 months of cultivation.

<table>
<thead>
<tr>
<th>Frankia inoculation</th>
<th>Average root diameter (mm)</th>
<th>Tensile resistance force (N)</th>
<th>Tensile strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inoculated</td>
<td>2.42±1.20</td>
<td>81.1±19.0</td>
<td>17.45±3.36</td>
</tr>
<tr>
<td>Non-inoculated</td>
<td>2.88±1.28</td>
<td>68.0±23.5</td>
<td>11.42±1.83</td>
</tr>
</tbody>
</table>

All values are the mean ± standard error of 86 root segments and 102 root segments of inoculated and non-inoculated A. formosana, respectively. Values in the same column followed by different superscript letters significantly differ at 5% significant level.

Figure 10. Root tensile resistance force-root diameter relationship for A. formosana seedlings inoculated (●—●) and non-inoculated (○—○) with Frankia.

An actinobacterium strain in nodules of A. formosana was isolated and recognized as Frankia by 16S rDNA gene similarity analysis. Inoculation test showed that this Frankia strain can induce actinorhizal nodule development in the roots of A. formosana seedlings. A number of studies indicated that Frankia strains associate with Alnus spp. (that is, Alnus acuminata, A. crispa, A. glutinosa, Alnus nepalensis, Alnus rubra and Alnus sieboldiana, respectively) (Benson, 1982; Carlson and Dawson, 1985; Hooker and Wheeler, 1987; Vendan et al., 1999; Carú et al., 2000; Oliveira et al., 2005; Yamanaka et al., 2005). Faure-Raynaud et al. (1991) analyzed the diversity of Frankia strains isolated from single nodules of A. glutinosa and showed no divergence among strains isolated from the same nodule. McEwan et al. (2015) also demonstrated that a single ribotype of Frankia is the major bacterium in single lobe from a nodule of A. glutinosa.

The research revealed that A. formosana seedlings inoculated with the native Frankia had significantly higher growth performance than the non-inoculated ones. Many previous studies have shown that inoculation with Frankia strains could significantly increase growth and development of alder seedlings (Prat, 1989; Wheeler et al., 1991; Kendall et al., 2003; Martin et al., 2003; Schrader and Graves, 2008; Bissonnette et al., 2014; Yamanaka et al., 2009). Lumini et al. (1994) showed that inoculation of selected Frankia strains and arbuscular mycorrhizal fungi along with sterilized media developed significantly higher shoot biomass than the non-inoculated controls. Quoreshi et al. (2007) demonstrated that A. crispa inoculated with Frankia had significantly
higher biomass, nodule lobes, and nodule weight than the controls. Moreover, Vendan et al. (1999) showed that Nepalese alder (A. nepalensis) plants inoculated with Frankia have a higher shoot length, root length, and biomass, while the Frankia strain AVC-II exhibited better infectivity and productivity of Nepalese alder than other strains tested. They also clearly indicated the potentiality of utilizing the Alnus-Frankia specificity for higher productivity through effective symbiosis. Schrader and Graves (2008) demonstrated that alder seedlings inoculated with species-specific Frankia strain grew larger, and gained more biomass than the cross-inoculated ones. Consistently, the study also demonstrated the positive effect of local Frankia strain on growth performance of A. formosana seedlings.

The root system architecture of A. formosana seedlings inoculated and non-inoculated with Frankia were similar to the heart root system (Stokes and Mattheck, 1996). The seedlings inoculated with Frankia had deeper taproot and more profuse roots than the non-inoculated ones. The inoculated seedlings had significantly higher total root length, root surface area, root volume, and root tip number than the controls. Moreover, seedlings inoculated with Frankia also had significantly higher root density and root length density than the non-inoculated ones. Wheeler et al. (1979) demonstrated that alder inoculated with Frankia developed more lateral root primordia than the non-inoculated ones, suggesting that Frankia can induce lateral root formation. De Bates et al. (2006) indicated that root density is a pertinent parameter to estimate the erosion-reducing efficacy. Stokes et al. (2009) also showed that higher root length density increases the uprooting resistance of plants.

This study demonstrates that A. formosana seedlings inoculated with Frankia have significantly higher uprooting resistance than the controls, indicating a higher anchorage capability in roots of the inoculated seedlings. Also, there were strong linear positive correlations between uprooting resistance and tree height, stem-base diameter, taproot length, root biomass and shoot biomass. Additionally, the inoculated A. formosana seedlings have longer taproot and more profuse lateral roots than the non-inoculated ones. Hence, inoculation with Frankia significantly augmented the numbers of lateral roots, which consequently stimulate seedling anchorage capability and uprooting resistance. Stokes et al. (2005) also indicated that heart root system is more resistant to uprooting than taproot system and plate root system.

The results of root tensile tests showed that the root tensile resistance force increased with increasing root diameter, whereas the root tensile strength decreased with increasing diameter. The findings are congruent with many other studies (Bischetti et al., 2005; Genet et al., 2005; De Baets et al., 2007; Normaniza et al., 2011; Nyambane et al., 2011; Abdi et al., 2014; Mohammed and
Normaniza, 2014; Capilleri et al., 2016). Root chemical components, such as cellulose, lignin, hemicellulose and holocellulose, are closely related to root mechanical properties. Genet et al. (2005) found that root cellulose content increased with increasing root diameter and increasing tensile strength in both Pinus pinaster and Castanea sativa. Several studies have shown that the root cellulose content increases and lignin content decreases with an increase in root diameter, and decrease in tensile strength (Lv et al., 2013; Zhang et al., 2014; Yang et al., 2016). In addition, the results from the present study also demonstrated that the average root tensile resistance force and tensile strength of inoculated seedlings were significantly higher than that of the non-inoculated controls. This suggests that inoculation with Frankia promotes root tensile resistance force and tensile strength of A. formosana seedlings.

Conclusions

Collectively, results of the present study clearly show that the native Frankia strain significantly enhances growth performance, root system architecture, uprooting resistance and root tensile strength of A. formosana seedlings. The findings of this study are of great importance in the application of Frankia in alder seedling production for landslide reforestation and soil conservation practices. This is the first report to demonstrate that inoculation with Frankia can significantly enhance growth, root system architecture, anchorage capability and root tensile strength of A. formosana seedlings. Additionally, further studies on the diversity of native Frankia strains and its symbiotic compatibility with alder species are needed for seedling production. Also, researches on the effects of alder roots on hillslope stability and erosion control are useful in ecological restoration and soil conservation in landslide areas.

CONFLICT OF INTERESTS

The authors have not declared any conflict of interests.

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REFERENCES

Alnus rubra


Characterization of biodiesel obtained from atemoya (Annona squamosa × A. cherimola) seed oil

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Biodiesel is derived from renewable sources, such as vegetable oils, by means of a transesterification process in which triacylglycerols are transformed into smaller molecules of esters of fatty acids and glycerol. The transesterification reactions of ‘Gefner’ atemoya (Annona squamosa × A. cherimola) seed oil extracted by pressing (physical) and solvent (chemical) processes were studied, with analysis of the methyl esters produced. The reactions were monitored using gas chromatography coupled to mass spectrometry (GC-MS), as well as by hydrogen nuclear magnetic resonance spectroscopy (¹H-NMR). The methyl esters formed during the transesterification reaction with methanol were determined for each oil. The major methyl esters (16:0, 18:0, 18:1 and 18:2) formed during 50 min of reaction were similar to those reported in the literature for other biodiesels; the peak areas and retention times were also similar. No changes in signal intensity over time were observed for the oils obtained by the two extraction methods. It was also noted that the extraction method had no influence on the types of methyl esters formed during biodiesel production.

Key words: Annona, oil extraction, transesterification.

INTRODUCTION

Biodiesel consists of mono-alkyl esters of long-chain fatty acids derived from renewable sources such as vegetable oils, obtained by a transesterification process in which triglycerides are transformed into smaller molecules of fatty acid esters and glycerol. Its use is intended to replace fossil fuels in diesel engines. It has promising potential, not only for its important contribution to reducing environmental pollution, but also for the generation of renewable energy as a replacement for fossil diesel and other petroleum products (Pinto et al., 2005).

In 2013, Brazil was the world’s second largest biodiesel
consumer, only behind the United States, which had a demand of 5.2 million m³. In terms of production, the USA is the global leader, with production of 5.1 million m³ in 2013, followed by Germany and Brazil, with production of 3.6 and 3.0 million m³ of biodiesel, respectively (Agência Nacional do Petróleo (ANP), 2013).

In 2004, the Brazilian government launched the National Program for Biodiesel Production (PNPB). Biodiesel can be used to partially or totally replace mineral diesel for light vehicles, trucks, tractors, and generators. In Brazil, the biodiesel mixture has been regulated by law since 2008. At first, the mandatory use was 2%, and it has been progressively increased to 5% (Kohlhepp, 2010).

Biodiesel is registered by the United States Environmental Protection Agency as a fuel and as an additive for fuels (Ferrari et al., 2005). After transesterification, biodiesel can be used neat at 100% (B100) or at proportions of 5% upwards in mixtures whose use is intended to replace fossil fuels in diesel cycle engines, without any need for modification of the engine. Various vegetable oils have been successfully tested in transesterifications with methanol or ethanol for the production of biodiesel. The seeds of peanuts, sunflowers, and soybeans, with oil contents of 41.3, 60.2, and 24.5 g 100 g⁻¹, respectively, are widely used for biodiesel production (Constantino et al., 2014). Oils extracted from different fruits have also been explored for biodiesel production (Adekanle et al., 2016; Alexandre et al., 2015), offering non-conventional sources of this biofuel.

Atemoya is an interspecific hybrid of cherimoya (Annona cherimola) and sugar-apple (Annona squamosa). It was introduced to Brazil in the 1980s and is mainly grown in the south and southeast of the country. In the 1990s, the ‘Gefner’ hybrid variety was successfully introduced in the northeast of Brazil. The cultivated area now exceeds 1,500 hectares, spread over the States of São Paulo and Paraná, as well as the northeast region (Braga-Sobrinho, 2014).

Atemoya seeds represent around 8.4% of the weight of the fruit and have potential as a source of biodiesel, since the lipid content is 27.3 g 100 g⁻¹ (Cruz et al., 2013). This content is close to that of other seeds such as soybeans, which are widely used for biofuel. The use of atemoya seeds to produce biodiesel can add value to the fruit.

The objective of this study was to analyze methyl esters produced during the transesterification reaction of the oil from ‘Gefner’ atemoya seeds, obtained by physical (pressing) and chemical (solvent) extraction.

**MATERIALS AND METHODS**

The atemoya was obtained during the 2010/2011 agricultural cycle in an orchard situated in the municipality of Jaíba, in northern Minas Gerais State, Brazil (14°33′-15°28′S, 43°29′-44°06′W, altitude of 500 m). The fruits were harvested at the appropriate stage of maturity and transported overland to Universidade Federal de Lavras. In the laboratory, the fruits were selected considering size, maturity, and absence of defects. Each replicate employed 82 fruits, totaling 902 fruits.

The seeds were separated and washed with distilled water, weighed, and dried in a forced-air circulation oven at 60 to 65°C until they reached humidity lower than 6%. The seeds were then vacuum-packed in plastic bags and stored at around -10°C in a cold chamber until oil extraction (AOAC, 2012).

**Oil extractions**

Oil extractions were performed by pressing (physical) and solvent (chemical) methods, as described by Cruz et al. (2015). Oil pressing was performed in a continuous expeller press, while chemical extraction employed a Soxhlet extractor with hexane as solvent at 68°C. Humidity determination was performed by dehydration of the oil until constant weight in an oven at 105°C, (Lutz, 2008).

**Biodiesel production**

Transesterification reactions were performed for 40 min at 50°C in a jacketed reactor, to which 200 ml of vegetable oil and 50 ml of methanol were added. This mixture was heated to a temperature of 50°C under mechanical stirring for 20 min. After this time, 6 ml of sodium methoxide (30%) were added, maintaining the temperature and stirring for 40 min. The solution was then transferred to a separation funnel for separation of the phases (biodiesel and glycerin) (Silva, 2005). Aliquots were removed at 0, 10, 20, 30, 40, and 50 min of reaction. Subsequently, 1 ml of each of the six aliquots was treated with 5 ml of chloroform, 0.5 ml of sulfuric acid, and 10 ml of saturated sodium chloride solution. The organic phase obtained was dried with magnesium sulfate, the solvent was removed in a rotary evaporator, and the product was dried with a flow of nitrogen gas. The samples obtained were analyzed using gas chromatography-mass spectrometry (GC-MS), as well as by hydrogen nuclear magnetic resonance spectroscopy (¹H-NMR). For the GC-MS analyses, the samples were resuspended in 0.1 ml of hexane.

**Chromatographic analysis**

The samples were analyzed using a gas chromatograph coupled to a GC-MS QP2010 Plus mass spectrometer (Shimadzu, Japan) equipped with an AOC-5000 autosampler for liquids and gases (Shimadzu, Japan). A 30 m × 0.25 mm × 0.25 μm RTX-5MS column (5% phenyl to 95% dimethylsiloxane) was used for separation and identification of the compounds. The injector was operated at 220°C in split mode, with a split ratio of 1:20. The carrier gas used was He 5.0, at a flow rate of 1.18 ml min⁻¹. The oven temperature was programmed from 60 to 240°C, with a heating ramp of 5°C min⁻¹; and then from 240 to 270°C, with a heating ramp of 10°C min⁻¹, followed by a final hold at 270°C for 7 min. An electron impact mass spectrometer (70 eV) was used in scan mode (45 to 500 Da), with solvent cutting at 3.5 min. The detector interface and ion source temperatures were kept at 240°C and 200°C, respectively. The compounds were identified by comparing the mass spectra with library spectra (Wiley 8 and FFNSC 1.2 libraries).

**Nuclear magnetic resonance analysis**

The ¹H-NMR analyses employed an EFT-60 spectrometer (Anasazi Instruments, Indianapolis, USA), with one-dimensional spectra acquired for the biodiesel samples obtained by both methods.
Table 1. Methyl esters obtained from transesterification of the seed oil from 'Gefner' atemoya (% peak area), using two oil extraction methods: physical (P) and chemical (C).

<table>
<thead>
<tr>
<th>Methyl esters of fatty acids</th>
<th>Number of carbons from the fatty acid: unsaturation number</th>
<th>Retention time (1st identification)</th>
<th>Reaction time (min)</th>
<th>P</th>
<th>C</th>
<th>P</th>
<th>C</th>
<th>P</th>
<th>C</th>
<th>P</th>
<th>C</th>
<th>P</th>
<th>C</th>
<th>P</th>
<th>C</th>
<th>P</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methyl octadecanoate</td>
<td>18:0</td>
<td>24.45</td>
<td>1</td>
<td>25.46</td>
<td>24.65</td>
<td>19.95</td>
<td>20.74</td>
<td>18.50</td>
<td>21.31</td>
<td>20.95</td>
<td>19.96</td>
<td>22.85</td>
<td>21.86</td>
<td>20.64</td>
<td>23.42</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl nonanoate</td>
<td>9:0</td>
<td>26.57</td>
<td>10</td>
<td>0.09</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.02</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Methyl eicosenoate</td>
<td>20:0</td>
<td>28.10</td>
<td>20</td>
<td>0.07</td>
<td>8.95</td>
<td>0.08</td>
<td>0.12</td>
<td>3.5</td>
<td>3.05</td>
<td>2.83</td>
<td>0.03</td>
<td>-</td>
<td>-</td>
<td>0.06</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl (9z)-9-octadecenoate</td>
<td>18:1</td>
<td>30.09</td>
<td>25</td>
<td>41.95</td>
<td>24.31</td>
<td>56.62</td>
<td>29.96</td>
<td>32.89</td>
<td>27.95</td>
<td>29.2</td>
<td>29.82</td>
<td>25.06</td>
<td>26.13</td>
<td>28.69</td>
<td>25.26</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl heptadecanoate</td>
<td>17:0</td>
<td>30.53</td>
<td>29</td>
<td>0.64</td>
<td>1.77</td>
<td>0.66</td>
<td>0.47</td>
<td>0.39</td>
<td>0.64</td>
<td>0.62</td>
<td>0.44</td>
<td>0.87</td>
<td>0.84</td>
<td>0.44</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl nonadecanoate</td>
<td>19:0</td>
<td>34.16</td>
<td>32</td>
<td>0.14</td>
<td>0.61</td>
<td>0.14</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.19</td>
<td>0.14</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl eicosanoate</td>
<td>21:0</td>
<td>35.86</td>
<td>34</td>
<td>3.25</td>
<td>-</td>
<td>-</td>
<td>2.92</td>
<td>1.76</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.58</td>
<td>3.78</td>
<td>2.96</td>
<td>6.25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl docosanoate</td>
<td>10:0</td>
<td>38.62</td>
<td>37</td>
<td>0.56</td>
<td>1.64</td>
<td>0.48</td>
<td>0.49</td>
<td>0.25</td>
<td>0.47</td>
<td>0.51</td>
<td>0.48</td>
<td>0.79</td>
<td>0.58</td>
<td>0.51</td>
<td>0.98</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl tetracosanoate</td>
<td>24:0</td>
<td>41.04</td>
<td>41</td>
<td>0.34</td>
<td>0.9</td>
<td>0.31</td>
<td>-</td>
<td>0.16</td>
<td>0.34</td>
<td>0.32</td>
<td>0.31</td>
<td>0.47</td>
<td>0.29</td>
<td>0.3</td>
<td>0.51</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl tetradecanoate</td>
<td>14:0</td>
<td>24.45</td>
<td>23</td>
<td>-</td>
<td>0.49</td>
<td>0.19</td>
<td>-</td>
<td>0.06</td>
<td>0.24</td>
<td>0.19</td>
<td>0.16</td>
<td>0.3</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl (9z)-9-hexadecenoate</td>
<td>16:1</td>
<td>28.19</td>
<td>22</td>
<td>-</td>
<td>1.31</td>
<td>0.3</td>
<td>0.45</td>
<td>0.3</td>
<td>0.94</td>
<td>-</td>
<td>0.36</td>
<td>1.56</td>
<td>0.94</td>
<td>0.5</td>
<td>1.65</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl (9z,12z)-9,12-octadecadienoate</td>
<td>18:2</td>
<td>31.86</td>
<td>20</td>
<td>-</td>
<td>3.73</td>
<td>-</td>
<td>23.08</td>
<td>24.86</td>
<td>21.12</td>
<td>22.04</td>
<td>23</td>
<td>17.72</td>
<td>21.03</td>
<td>22.81</td>
<td>15.32</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl (11e)-11-eicosenoate</td>
<td>20:1</td>
<td>35.45</td>
<td>22</td>
<td>-</td>
<td>1.53</td>
<td>-</td>
<td>0.14</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.44</td>
<td>0.84</td>
<td>0.59</td>
<td>0.42</td>
<td>1.15</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Previously treated biodiesel samples (0.1 ml) were dissolved in 0.1 ml of chloroform deuterated with 99.8% deuterium (CDCl₃), in 5 mm NMR tubes. Tetramethylsilane (0.1 ml) was used as an internal reference standard.

RESULTS AND DISCUSSION

The yield of the transesterification reaction for the oil obtained by pressing of atemoya seeds was 89% methyl esters and 11% glycerin. The biodiesel yield for the chemical extraction was 91% methyl esters and 9% glycerin. These values differed from the results obtained for soybean biodiesel by Ferrari et al. (2005), who reported 57.26% ethyl esters, 22.29% glycerin, 10.04% recovered ethanol, and 10.41% losses. The concentration of glycerin in the biodiesel obtained from atemoya seeds was approximately half that obtained from soybeans, so the concentration of biodiesel was much higher, and losses were not observed.

Table 1 shows the profile of methyl esters for the transesterification reaction of the oil from 'Gefner' atemoya seeds. The major esters found in the atemoya biodiesel were 16:0, 18:0, 18:1, and 18:2.

The ester area percentages obtained for the physical and chemical extractions, after a reaction time of 50 min, at which the biodiesel was decanted, were 22.2 and 24.3 (16:0), 20.64 and 23.42 (18:0), 28.7 and 25.26 (18:1), and 22.81 and 15.32 (18:2), respectively. Comparison of the results obtained here with the findings of Urioste et al. (2008) revealed the presence of one additional ester (18:2).

In chromatographic, determination of the esters of fatty acids was obtained in transesterification reactions of babassu oil with ethanol, propanol, and butanol. Urioste et al. (2008) reported the following areas (%) for ethyl, propyl, and butyl esters, respectively: 24.82, 22.82, and 23.52 (16:0); 21.56, 22.14, and 23.28 (18:0); and 24.60, 22.50, and 24.56 (18:1). It appears that there was virtually no difference between the areas of the three ethyl, propyl and butyl esters formed in babassu biodiesel.

In their study with soybean oil, Ferrari et al. (2005) observed the following areas (%) for ethyl esters of fatty acids: 11.29 (16:0); 21.56, 22.14, and 23.28 (18:0); and 24.60, 22.50, and 24.56 (18:1). It appears that there was virtually no difference between the areas of the three ethyl, propyl and butyl esters formed in babassu biodiesel.
15.32%, respectively, which were smaller than the value obtained for soybean oil. On the other hand, the 18:0 and 18:1 esters of atemoya oil had greater areas, compared to the soybean oil, with values of 20.64% (pressed) and 23.42% (solvent extracted), and 28.69% (pressed) and 25.26% (solvent extracted), respectively. These data show that the predominant esters in current biodiesels are 16:0, 18:0, 18:1, 18:2, and 18:3.

Marques et al. (2010) emphasized the importance of considering instrumental precision when evaluating major peak areas, in order to increase the accuracy of determination of esters of fatty acids formed during transesterification reactions. Barbosa et al. (2010) reported the following areas (%) for ethyl esters of soybean seed oil: 16.0 (16:0), 2.4 (18:0), 23.5 (18:1), and 51.2 (18:2). In the present work, greater areas were obtained for 16:0, 18:0, and 18:1 esters in the atemoya oil biodiesel obtained using both forms of oil extraction (press and solvent), while the value obtained for the 18:2 ester was smaller. Benito et al. (2014) studied the biodiesel potential of Annona diversifolia seed oil and reported the following areas (%) for methyl esters: 16.4 (16:0), 5.22 (18:0), 70.4 (18:1), and 7.97 (18:2). The areas obtained for the 16:0, 18:0, and 18:2 methyl esters were greater for the decanted biodiesel from atemoya oil extracted by both techniques (physical and chemical), while the area was smaller for the 18:1 ester.

Methanol is the main alcohol used in transesterification in many countries (Pinto et al., 2005). In Brazil, several research groups and small producers use the methyl pathway for the production of biodiesel, because methanol is more reactive, while ethanol causes greater dispersion of glycerin in the biodiesel, making separation difficult (Lôbo et al., 2009). A reaction time of 30min was required for the formation of methyl esters in the biodiesel, similar to the duration of 25min reported by Urioste et al. (2008) for biodiesel from Babassu, where the major esters formed were 16:0, 18:0, and 18:1.

Encinar et al. (2002) observed that the transesterification reaction was very fast, with conversion into ethyl esters close to the maximum value after only 5 to 10min of reaction, and stabilization at a maximum value after 20 to 30min. These values were similar to those found for the formation of biodiesel from atemoya oil, which occurred after 20 to 30min of reaction. However, in the study of Ferrari et al. (2005), chromatographic monitoring of the products formed after various reaction times showed that a time of 5min was sufficient for the conversion of neutral and dried oil into ester. The conversion of fatty acids into methyl esters in the atemoya oil occurred between 5 and 10min of reaction, stabilizing at a maximum value after 20 to 30min. The physical extraction (pressing) of atemoya seed oil is economically advantageous and provides a high oil extraction efficiency of 88.9g 100 g–1 (dry mass basis) (Cruz et al., 2015).

Furthermore, in comparison with chemical extraction using solvent, a disadvantage of the latter has greater oxidation of the extracted oil. The profile of the methyl esters identified by GC-MS was confirmed by 1H-NMR. Figures 1 and 2 show the major chemical shifts characterizing the esters, formed after 50 min of reaction. No changes in signal strength or in the hydrogen chemical shifts with time were observed after the
Figure 2. $^1$H-NMR spectrum of methyl esters obtained in the transesterification of the ‘Gefner’ atemoya seed oil extracted chemically (using solvent), at 50 min reaction time.

**Table 2.** Chemical shifts (ppm) obtained by $^1$H-NMR (60 MHz) of biodiesel in CDCl$_3$ at 25°C (physical extraction).

<table>
<thead>
<tr>
<th>$^3$H</th>
<th>Atemoya biodiesel</th>
<th>Paiva et al. (2010)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH$_3$</td>
<td>0.89</td>
<td>0.90</td>
</tr>
<tr>
<td>CH$_2$</td>
<td>1.27</td>
<td>1.2 – 2.5</td>
</tr>
<tr>
<td>CH$_2$ (α-carbonyl)</td>
<td>2.40</td>
<td>2.1 – 2.5</td>
</tr>
<tr>
<td>CH$_3$-O</td>
<td>3.68</td>
<td>3.5 – 4.8</td>
</tr>
<tr>
<td>CH$_2$ (vinylic)</td>
<td>5.99</td>
<td>4.5 – 7.0</td>
</tr>
</tbody>
</table>

**Table 3.** Chemical shifts (ppm) obtained by $^1$H-NMR (60 MHz) of biodiesel in CDCl$_3$ at 25°C (chemical extraction).

<table>
<thead>
<tr>
<th>$^3$H</th>
<th>Atemoya biodiesel</th>
<th>Paiva et al. (2010)</th>
</tr>
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<tbody>
<tr>
<td>CH$_3$</td>
<td>0.89</td>
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</tr>
<tr>
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<td>1.27</td>
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<tr>
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<td>3.5 – 4.8</td>
</tr>
<tr>
<td>CH$_2$ (vinylic)</td>
<td>5.99</td>
<td>4.5 – 7.0</td>
</tr>
</tbody>
</table>

The method used for oil extraction had no influence on the types of methyl esters formed during biodiesel production. The oil from atemoya seeds has potential for use as biodiesel, with advantages including addition of value to the fruit and a use for unwanted biomass (the seeds) that might otherwise be treated as waste.

**CONFLICT OF INTERESTS**

The authors have not declared any conflict of interests.

**ACKNOWLEDGMENTS**

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