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Genetic diversity of antibiotic and plant extract resistant *Staphylococcus aureus* isolated from hospitalized patients in Ekiti State, Nigeria

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This study aimed to determine the genetic diversity of antibiotics and plants extract resistant *Staphylococcus aureus* using molecular technique. A total of 106 human clinical samples were collected from patients in three different hospitals within Ekiti State. Microbiological and molecular analyses were performed using standard methods. Antimicrobial susceptibility test with common antibiotics and plant leaves extracts was carried out using disk and agar well diffusion methods. Urine samples from male patients recorded high percentage of *S. aureus* (57.1%) as compared to the female patients (43.1%). Percentage of *S. aureus* recorded from the ear swab samples in male patients (58.3%) was also high as compared to the female (41.7%). From the nose swab samples, female patients recorded 60% as compared to their male counterpart (40%). Equal percentage (50%) of wound infection caused by *S. aureus* was observed in both male and female patients. Of the eight antibiotics used in this study, the *S. aureus* isolated were susceptible to ofloxacin (25 to 36 mm), gentamycin (16 to 23 mm) and erythromycin (11 to 25 mm). All the isolates confirmed resistance to ceftaxidime and cloxacillin. Nine isolates were susceptible to cefuroxime with zones of inhibition that ranged from 10 to 25 mm, while 11 were susceptible to ceftriaxone with zone of inhibition between 11 and 20 mm. Only one isolate was sensitive to augmentin (amoxicillin/clavulanate) with zone of inhibition of 20 mm. Out of three plant extracts used in this study, aqueous leaf extract of *Terminalia catappa* demonstrated highest antibacterial activity on the test isolates with zone of inhibition ranging from 11 to 36 mm followed by *Mangifera indica* with inhibition ranging from 11 to 32 mm, while least inhibition ranging from 11 to 20 mm was exhibited by *Acalypha wikesiana*. Random amplified polymorphic DNA (RAPD) proved to be useful as genetic markers in determining genetic diversity among antibiotic and plant extract resistant *S. aureus*.

**Key words:** *Acalypha wikesiana*, multidrug-resistant strains, *Staphylococcus aureus*, genetic diversity.

INTRODUCTION

Modern medicine is no doubt faced with an unprecedented threat posed by multidrug-resistant bacteria (Levy and Marshall, 2004). It has been advised that a multidisciplinary approach is required to tackle the

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emergence and evolution of drug resistance microorganisms (Martinez and Baquero, 2002). In their study, Baquero et al. (2003) averred that the multidisciplinary approach will be a challenge given that the trend of antibiotic resistance appears as a reflection of the characteristic of a dynamic, highly complex, and self-organizing system evolving at an instance of complex situations. New molecular technologies promise to offer adequate and effective tools in the fight against antibiotic resistance strains of microorganisms. Advance molecular approaches have also been reported to be effective containment of multidrug-resistant strains, given the accurate detection of bacterial species from different environment. Currently, the main advantage of molecular resistance testing is the rapid turn-around time and improved accuracy, both of which are relevant particularly in cases of life-threatening diseases such as meningitis and sepsis, for which rapid detection, identification and resistance testing is important (Johansen et al., 2003). It is noteworthy to add that, rapid polymerase chain reaction (PCR)-based assays for resistance testing have been used as excellent complementary tools in the laboratory for the detection of methicillin-resistant Staphylococcus aureus (MRSA) (Warren et al., 2004).

Medicinal plants as alternative therapeutic agent have recently attracted the attention of the biological scientific communities. Use of medicinal plants involves the isolation and identification of secondary metabolites produced by plants and is used as active principles in medicinal preparations. Application of various herbal medicinal preparations for the treatment of S. aureus associated illnesses in Nigeria has been reported (Akinyemi et al., 2005). Volumes of reports have demonstrated the antimicrobial properties of several of these plants against multi-drug resistant bacteria, particularly methicillin resistant S. aureus (Ankri and Mirelman, 1999; Adesino et al., 2011; Khan et al., 2016).

Methicillin resistant S. aureus (MRSA) is a bacterium responsible for several difficult-to-treat infections in humans. MRSA is any strain of S. aureus that has developed through the horizontal gene transfer and natural selection, multi-resistance to beta-lactam antibiotics, which include the penicillins (methylcillin, dicloxacillin, nafcillin, oxacillin, etc.) and the cephalosporins. Understanding the S. aureus lineages therefore, is crucial in controlling nosocomial infections. Recent studies on S. aureus in Nigeria have revealed an escalating burden of MRSA. However, the S. aureus genotypes circulating among hospitalized patients in hospitals found in Ekiti State are not known, hence this study.

MATERIALS AND METHODS

Sampling

In this research, a total of 106 human clinical samples were collected and transported within 2 h of collection to the Microbiology Research Laboratory in the Department of Biological Sciences, Afe Babalola University, Ado-Ekiti for microbiological analysis. The samples used in this research were urine, wound, nose and ear swabs obtained from patients hospitalized at different wards of Ekiti State University Teaching Hospital (EKSUTH), State Specialist Hospital Ikere-Ekiti, and General Hospital Iyin-Ekiti in Ekiti State Nigeria. Samples were obtained from both male and female of different ages. From men, 33 urine samples, 14 ear swabs, four nose swabs and seven wound swabs were obtained. On the other hand, a total of 45 samples were obtained from women comprising 25 urine samples, ten ear swabs, six nose swabs and seven wound swabs. Each urine sample was aseptically collected into a sterile sampling bottle, while ear, nose and wound swabs were collected using a sterile swab stick.

Plant sample collection and preparation

Fresh plant leaves of Terminalia catappa (Almond), Mangifera indica (Mango) and Acalypha wikesiana (Acalypha) were obtained within the premises of Afe Babalola University and identified in the Department of Botany, University of Ibadan, Nigeria. Voucher specimens for T. catappa (UIH 22567), M. indica (UIH 22568) and A. wikesiana (UIH 22569) were deposited at University of Ibadan Herbarium of the Department of Botany. The leaves were brought to the laboratory, stripped from their stems, washed with distilled water and crushed into a fine infusion using a laboratory mortar and pestle. The solution was filtered with No. 1 Whatman filter paper and finally passed through membrane filter to obtain sterile extract. These were labelled and kept at 4°C in the refrigerator for use. A clean filter was used to obtain the extracts respectively into three well labelled beakers.

Qualitative analysis of phytochemical constituent

Simple standard chemical tests as previously described by Harborne (1973) and Chethana et al. (2013) were employed for detecting the presence of some phytochemical components such as saponins, tannins, alkaloids, steroids, phenol, terpenoid, flavonoid and glycosides in the plant extract.

Isolation and identification of test organisms

The isolates gotten from the various sites were identified using standard microbiological method as described by Cheesbrough (2000).

Antibiotic sensitivity testing

Antimicrobial susceptibility of the isolates was determined by the Kirby Bauer disc diffusion method according to CLSI guidelines (2012) using the following antibiotics: ofloxacin (OFL), augmentin (AUG), ceftaxidime (CAZ), cefuroxime (CRX), gentamycin (GENT), ceftriaxone (CTR), erythromycin (ERY) and cloxacillin (CXC). Young culture of the isolate was suspended in a sterile saline solution (0.85% NaCl). The suspension was adjusted to match 0.5 McFarland turbidity standards. The antibiotics discs were aseptically placed on Mueller Hinton agar surface using sterile forceps and incubate at 37°C for 24 to 48 h. Effectiveness of the antibiotics was shown by clear zones around them which were measured and interpreted as either resistance, intermediate, or susceptible.

Plant extract inhibitory test

Agar well diffusion method was used as previously described
DNA isolation and purification of bacterial genomic DNA

Sterile dry swabs were used for streaking clinical samples onto sterile nutrient agar plates (CONDA). Inoculated plates were incubated at 37°C for 48 h. Stock cultures of the pure isolates were prepared and refrigerated at 4°C to be kept for further analysis. Extraction and purification of genomic DNA of the bacterial isolates was carried out using the ZR Fungal/Bacterial DNA MiniPrep™ (Zymo). The procedure employed was as described by the manufacturer.

RAPD PCR analysis

RAPD PCR analysis used in this study was performed as previously described by Onasanya et al. (2003). DNA primers tested were purchased from Operon Technologies (Alemada, California, USA) with each being 10 nucleotides long. Two separate concentrations of DNA, 24 and 96 ng per reaction were used to test reproducibility as well as eliminate irregular amplification products from the analysis. Thirteen primers were screened for their ability to amplify the DNA of the S. aureus isolate. Six out of the 13 screened primers were found useful because they showed polymorphism. These were used for amplification of DNA of the 20 isolates. Amplification were performed in 25-μl reaction mixture consisting of genomic DNA, IX reaction buffer, (Promega), 100 μM each of dATP, dCTP, dGTP and dTTP, 0.2 μM Operon random primer, 2.5 μM MgCl₂ and 0.2 μl (1 unit) of Taq polymerase (Boehringer, Germany). A single primer was used in each reaction. Amplification was performed in a thermowell microtiter plate (Costa Corporation) using a Perkin Elmer programmable Thermal Controller model 9600. The cycling program was set as follows: initial denaturation at 94°C for 3 min followed by 45 cycles each at 94°C for 1 min, 40°C for 1 min., 72°C for 2 min and final extension at 72°C for 7 min. Amplified DNA products were analyzed by electrophoresis in 1.4% agarose gel using TAE buffer (45 Mm Tris-acetate, 1 Mm EDTA, pH 8.0) at 100 V. Gels were visualized by staining with ethidium bromide solution (0.5 μg/ml) and banding patterns were observed using the Photogel documentation system.

Phylogenetic analysis

Positions of unequivocally scorable RAPD bands were transformed into a binary character matrix (to indicate the presence or absence of a band at a particular position). Pair-wise distance matrices were compiled by the Numerical Taxonomy System (NTSYS) 2.0 software (Rohlf, 2000) using a simple matching coefficient for estimation by means of the Jaccard’s coefficient to construct a similarity matrix (Ivchenko and Honov, 1998). Cluster analysis and dendrogram were produced on the basis of the unweighted pair group method arithmetic (UPGMA) (Sneath and Sokal, 1973; Jako et al., 2009) as the banding profiles of the six RAPD primers represent a novel approach for rapid screening of the multidrug resistant.

Statistical analysis

Data generated were expressed in mean ± standard error mean (SEM). One-way analysis of variance (ANOVA) followed by Student T-test (Walson, 1989; AOAC 1990) was used for this research.

RESULTS AND DISCUSSION

All samples analyzed for the presence of *Staphylococcus* on mannitol salt agar and nutrient agar media showed obvious bacterial growths. Out of 106 collected samples from both male and female patients of different age brackets admitted to the three hospitals within Ekiti State, Ekiti State University Teaching Hospital (EKSUTH) had the highest number of patient colonized with *S. aureus*.

From urine samples, high percentage of *S. aureus* in male patients (57.1%) as compared to their female counterpart (43.1%) was recorded. Percentage of *S. aureus* recorded from the ear swab samples in male patients (58.3%) was also high as compared to the female (41.7%). From the nose swab samples, female patients recorded 60% as compared to their male counterpart (40%). Equal percentage (50%) of wound infection caused by *S. aureus* was observed in both male and female patients.

Further to this study, 20 representatives of the *S. aureus* isolates were picked for multidrug resistance pattern to antibiotics and plant leaves aqueous extracts of which 9 were obtained from patients’ urine, three from the noses, four from the ears and four from wounds. These isolates were code-numbered accordingly. All the isolates were completely resistant to CAZ and CXC as there was no zone of inhibition observed. Nine isolates were observed to be sensitive to CRX with zones of inhibition ranging from 10 to 25 mm, while 11 isolates were sensitive to CTR with range between 11 and 20 mm. Meanwhile, only one isolate was sensitive to AUG 20 mm zone of inhibition (Figure 1). However, the 20 *S. aureus* isolates used in the study were all susceptible to OFL with clear zones of inhibition ranging from 24 to 35 mm followed by GEN 16 to 23 mm and ERY 11 to 25 mm.

This apparently high level of susceptibility of the isolates to OFL suggest that ofloxacin (OFL) could be a drug of choice for treating infections caused by *S. aureus* in the study area, especially at the present time, when *S. aureus* strains are resistant to other commonly used antibiotics as reported in many literatures. This finding is consistent with previous reports of Amadi et al. (2007) and Chigbu and Ezeronye (2003). Also, 100% sensitivity of *S. aureus* isolates to ofloxacin (OFL) was reported by Obi et al. (1996). In addition, high inhibition of OFL against *S. aureus* isolates was also highlighted by Chalita et al. (2004). However, low sensitivity was reported among most strains of methicillin resistant *S. aureus* (MRSA) isolated from patients with ocular infections (Uwaezuoke and Aririatu, 2004). *S. aureus* susceptibility to gentamycin (GEN) in this present study compares favourably with reports published in the South-western part of Nigeria, by Paul et al. (1982) and Ndip et al. (1997).

On the other hand, the sensitivity pattern of 20 *S. aureus* isolates to leaves extract shows that *T. catappa*
Figure 1. Susceptibility pattern of *S. aureus* to antibiotics. MSW = Male surgical ward; FSW = female surgical ward; MMW = male medical ward; FMW = female medical ward; SOP = surgical outpatient; ENT = ear nose and throat, B, A, W (5, 4, 1, 8, 3, 6, 10 respectively if bed no.; ES = ear swab; NS = nose swab; WS = wound swab. OFL = Ofloxacin; AUG = augmentin; CAZ = ceftaxidime; CRX = cefuroxime; GENT = gentamycin; CTR = ceftriaxone; ERY = erythromycin; CXC = cloxacillin.

demonstrated the highest level of effectiveness against 18 isolates which included isolates gotten from urine, ear, nose and wounds with zone of inhibition that ranges between 16 to 36 mm, with 2 isolates resistant to the plant. Although, 19 isolates were inhibited by *M. indica*, their zone of inhibition ranged between 11 and 32 mm, with only one isolate resistant and 14 out of the 20 isolates showed resistance to the aqueous leaf extract of *A. wikesiena*. Susceptibility of 6 isolates to *A. wikesiena* ranged from 11 to 20 mm zones of inhibition (*Figure 2*). The highest antimicrobial activity was observed with *T. catappa* which had better effect on 18 isolates with wider zones of inhibition than all the commercial antibiotics used in this study. Also, aqueous leaf extract of *M. indica* demonstrated higher antimicrobial effect on 19 isolates and its inhibitory activity was better than all the antibiotics used except OFL from which a wider zone of inhibition was observed. This indicates that the extracts of the leaf are as effective as the commercially prepared antibiotics thus confirming its potency in the treatment of *S. aureus*-mediated infections as well as gastroenteritis and skin dermatitis. Moreover, since the leaf extracts exhibited inhibitory activity on the test organisms, the antibacterial activity appears to be broad spectrum (Oluduro et al., 2011). The phytochemicals detected in the leaves aqueous extracts of *T. catappa*, *M. indica* and *A. wikesiena* in this study corresponds to earlier studies carried out with other plants by Onah et al. (2002), Akpan and Usoh (2004) and Oyewole et al. (2008).

The quantity of tannin present in *T. catappa* was significantly different from the quantity of tannin present in *A. wikesiena* but not significantly different from the value obtained in *M. indica* (Table 1). Furthermore, flavonoids content in *T. catappa* shows no significant difference from that in *M. indica*. No significant difference existed in the phenol quantity between *T. catappa*, *M. indica* and *A. wikesiena*. Also, the quantity of saponin in *T. catappa* was not significantly different from the values of *M. indica* and *A. wikesiena* (Table 1).

In this study, it was observed that all isolates from patients' samples receiving treatment from EKSUTH expressed similar pattern in their resistance to the antibiotic and plant extract used. Genetic fingerprinting of the 20 *S. aureus* isolates from urine, ear, nose and wound swabs of patients receiving treatment from 3 hospitals within Ekiti State was compared using RAPD-PCR markers. Of the 13 arbitrary primers tested, six primers (Table 2) showed a good level of polymorphism and reproducibility. The amplification reaction of 6 oligonucleotide primers yielded 135 bands of which 121 were polymorphic with sizes ranging between 200 and 3,000 bp (*Figure 3a to f*). 121 polymorphic RAPD
Figure 2. Antibacterial effect of *T. catappa*, *M. indica* and *A. wikesienia* leaves aqueous extracts on the twenty *S. aureus* isolates. MSW= male surgical ward; FSW= female surgical ward; MMW= male medical ward; FMW= female medical ward; SOP= surgical outpatient; ENT= ear nose and throat; B, A; W (5, 4, 1, 8, 3, 6 and 10 respectively) = Bed no.; ES= ear swab; NS= nose swab; WS= wound swab.

Table 1. Quantity of phytochemicals from the aqueous leaf extracts (mg/g).

<table>
<thead>
<tr>
<th>Samples</th>
<th>Tannin</th>
<th>Flavonoids</th>
<th>Phenol</th>
<th>Saponin</th>
<th>Alkaloids</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>T. catappa</em></td>
<td>0.04 ±0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.17 ±0.02&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.03 ±0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.60 ±0.02&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.63±0.02&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><em>M. indica</em></td>
<td>0.03±0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.17±0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.03 ±0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.31 ±0.36&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.43±0.40&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><em>A. wikesienia</em></td>
<td>0.06±0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>-</td>
<td>0.04 ±0.02&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.26 ±0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.52±0.01&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>S. V. (α = 0.05)&lt;sup&gt;(p = 0.002)&lt;/sup&gt;</td>
<td>0.000</td>
<td>0.868</td>
<td>0.096</td>
<td>0.089</td>
<td></td>
</tr>
</tbody>
</table>

Values are mean ± standard deviations of triplicate determinations. Values with different superscript per column are statistically significant (p<0.05).

markers were used to construct the phylogenetic relationship dendrogram among 20 *S. aureus* isolates into different group’s genotypes and sub groups depending on their reaction with antibiotics and plants extract used in the study.

From the reaction of 8 antibiotics on the 20 *S. aureus* isolates, two different reactive groups emerged which consists of antibiotics reaction group 1 (ARg1) and antibiotics reaction group 2 (ARg2) at 75% similarity (Figure 4a). ARg1 yielded 3 subgroups which include ARg1a, ARg1b and ARg1c. ARg1a consisted of six isolates (1-MSWB5U, 4-MMWB1iU, 11-ENTN2, 13-
Table 2. Oligonucleotide primers showing levels of genetic diversity among S. aureus isolates using the RAPD-PCR analysis.

<table>
<thead>
<tr>
<th>Operon primer code</th>
<th>Primer sequence 5' – 3'</th>
<th>Total number of fragments</th>
<th>Monomorphic bands</th>
<th>Polymorphic bands</th>
</tr>
</thead>
<tbody>
<tr>
<td>OPT06</td>
<td>CAAGGGCAGA</td>
<td>25</td>
<td>3</td>
<td>22</td>
</tr>
<tr>
<td>OPT14</td>
<td>AATGCCGCAG</td>
<td>28</td>
<td>2</td>
<td>26</td>
</tr>
<tr>
<td>OPT01</td>
<td>GGGCCACTCA</td>
<td>22</td>
<td>2</td>
<td>20</td>
</tr>
<tr>
<td>OPB04</td>
<td>GGACTGGAGT</td>
<td>20</td>
<td>4</td>
<td>16</td>
</tr>
<tr>
<td>OPH08</td>
<td>GAAACACCCC</td>
<td>22</td>
<td>1</td>
<td>21</td>
</tr>
<tr>
<td>OPT12</td>
<td>GGGTGTGTAG</td>
<td>18</td>
<td>2</td>
<td>16</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td><strong>135</strong></td>
<td><strong>14</strong></td>
<td><strong>121</strong></td>
</tr>
</tbody>
</table>

Figure 3a. Band patterns obtained after RAPD amplification of the 20 S. aureus isolates with the oligonucleotide OPT06. Lane M, DNA molecular size marker 1 kbp; Lane 1 – 20, 20 S. aureus isolates (ENTE4iES, 17-FMWB3i and 14-MSWB4iiiES). The reactions of these isolates with five antibiotics were identical. ARg1a consisted of four isolates (2-FSWB4U, 5-MMWB8iiiU, 18-MSWB6iiWS and 19-MSWB10WS) that were also identical in their reaction with 5 antibiotics. ARg1c had only 1 isolate (16-SOPB3i). ARg2 yielded only 2 subgroups consisting of ARg2a and ARg2b. ARg2a consisted of 7 isolates (3-FSWB4iiU, 6-MMWB8iiU, 20-MMWB1iES, 15-ENTE4iES, 12-FMWB4iNS, 9-SOPB2ii and 8-FMWA1iiU). These isolates show similar pattern of activities in their reaction with the 5 antibiotics. ARg1a consisted of two isolates (7-MMWB13iiiU and 10-FMWWA4iNS). Both isolates in ARg2b were identical in their reaction with the five antibiotics.

Figure 4b presents cluster E that shows the reaction of 3 plant leaf extracts (T. cattapa, M. indica and A. wikesiena) on 20 S. aureus isolates. At 55% coefficient of similarity, 2 major groups were formed, namely Gg1 and Gg2. Gg1 developed two subgroups namely Gg1a and Gg1b. Gg1a which consists of 12 isolates (1-MSWB5U, 2-FSWB4U, 3-FSWB4iiU, 4-MMWB1iU, 17-FMWB3i, 18-MSWB6iiWS, 14-MSWB4iiiES, 16-SOPB3i, 10-FMWWA4iNS, 9-SOPB2ii, 8-FMWA1iiU and 5-MMWB8iiiU). These isolates reacted in the same way to the aqueous leaf extracts. Gg1b consisted of six isolates (6-MMWB8iiU, 7-MMWB13iiiU, 20-MMWB1iES, 19-FMWB10WS, 13-ENTE4iES and 15-ENTE14i) which also reacted in the same way with the extracts. Gg2 had two isolates (11-ENT N2 and 12-FMWB4iNS) which reacted differently to the extracts.

Figure 4c presents cluster G which consists of two major group genotypes (Gg) represented as Gg1 and Gg2 at 10.5% similarity. Gg1 yielded 2 subgroups namely Gg1a and Gg1b at 11% similarity. Gg1a further gave rise to two subgroups consisting of Gg1ai and
Gg1aii; Gg1b gave rise also to two subgroups Gg1bi and Gg1bii as with 11.5% similarity. Gg2 did not yield any obvious subgroups. However, the (Gg) cluster formed three distinct groups; the first group consisted of four isolates (1, 16, 15 and 19), the second group consisted of four isolates (2, 6, 7 and 3) while the third group consisted of 9 isolates (8, 9, 17, 10, 11, 12, 18, 13 and 14). From the first group, it was observed that only the group genotype of isolate 1 was resistant to the antibiotics, making the group an antibiotic resistance emergent group. Group genotypes of the isolates in group 2 revealed antibiotics susceptibility because no resistance was found in the isolates.

Molecular research studies indicate that resistance is a complex system, resulting from multiple interacting mechanisms at the genomic, regulatory and expression levels. If the complementary effort of both the culture-dependent and molecular technique as evidenced in this study is anything to go by, molecular assays for resistance testing will not replace conventional culture-based antibiotic susceptibility testing in the immediate future (Fluit et al., 2001).
**Figure 3d.** Band patterns obtained after RAPD amplification of the 20 *S. aureus* isolates with the oligonucleotide OPT01. Lane M, DNA molecular size marker 1 kbp; Lane 1 – 20, 20 *S. aureus*.

**Figure 3e.** Band patterns obtained after RAPD amplification of the 20 *S. aureus* isolates with the oligonucleotide OPT12. Lane M, DNA molecular size marker 1 kbp, Lane 1 – 20, 20 *S. aureus*.

**Figure 3f.** Band patterns obtained after RAPD amplification of the 20 *S. aureus* isolates with the oligonucleotide OPT14. Lane M, DNA molecular size marker 1 kbp, Lane 1 – 20, 20 *S. aureus*. 
Figure 4a. Cluster A showing the reaction of twenty *S. aureus* isolates against five antibiotics.

Figure 4b. Cluster E showing the reaction of 20 *S. aureus* isolates against three plant leaf extracts.
According to Adesida et al. (2007), identification of antibiotic resistant organism using molecular techniques such as 16s rRNA, random amplification of polymorphic DNA (RAPD) and plasmid profile causing infectious processes proves to be very essential for effective antimicrobial and supportive therapy.

Conclusion

By virtue of this analysis using molecular method (RAPD- PCR), it was possible to group isolates of similar genotypes with the same resistance behaviour to antibiotics and plants extract. Although major bands from RAPD reactions are highly reproducible, challenges like minor bands proves difficult to repeat due to random priming nature of PCR reaction and potential confounding effects associated with co-migration with other markers.

CONFLICT OF INTERESTS

The authors have not declared any conflict of interests.

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