Multi-Drug Resistance Pattern of *Staphylococcus aureus* from Pediatric ward, General Hospital, Ikot-Ekpaw, Mkpat Enin LGA, Akwa-Ibom State, Nigeria

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Received date: August 08, 2019; Accepted date: February 21, 2020; Published date: February 27, 2020

Citation: Ibanga IA, Akan OD, Uyanga FZ, Mantu EC, Asuquo M (2020) Multi-Drug Resistance Pattern of *Staphylococcus aureus* from Paediatric ward, General Hospital, Ikot-Ekpaw, Mkpat Enin LGA, Akwa-Ibom State, Nigeria. Arch Clin Microbiol Vol. 11 No. 1:105.

**Abstract**

The evaluation of the Multi-drug resistance pattern of *Staphylococcus aureus* from a pediatric ward, in Akwa-Ibom State was conducted using standard clinical microbiological procedures.

Of the 100 samples from skin, wound, ear, throat and nose swabs, 28 isolates were confirmed as *S. aureus* and were subjected to a range of selected commercially available antibiotics like: amoxicillin, ampiclox, chloramphenicol, ciprofloxacin, erythromycin, gentamicin, levofloxacin, norfloxacin, rifampicin and streptomycin, to evaluate their susceptibilities. The wound swabs gave the highest isolate percentage yield (32%) followed by skin swabs (29%). While susceptibility results showed that amoxicillin and ampiclox were more resisted by the isolates, while ciprofloxacin, levofloxacin and norfloxacin were more effective against the isolates.

The MAR indices showed that 85.7% of the isolates had confirmed multi-drug resistance status, with 60.7% of the isolates having resistance for between four or more the tested antimicrobials. MAR indices revealed that 96.4% of the isolates had 0.3, indicating that the resistance resulted from isolates that adapted to the tested drugs due to some form of abuse. Restricted use of these drugs would help curtail the high resistance currently experienced amongst microorganisms.

**Keywords:** Multi-drug resistance; *Staphylococcus aureus*; Multiple antibiotics resistance index; Susceptibility test; Nosocomial infections

**Introduction**

Microbes do manifest themselves in three ways, through substance spoilage, fermentation of organic and inorganic matters and causation of ailments. Different microorganisms, with their different mode of aetiology, causing different types of ailments, will require different methods and capable drugs for treatments. Continuous deployment of antimicrobial drugs in treating microbial infections has led to the emergence of resistance amongst various strains of microorganisms [1,2]. MDR literally means ‘being resistant to more than one antimicrobial agent’, although a standardized definition has not yet been agreed upon by the medical community. There are currently other definitions that are used to characterize patterns multidrug resistance.

The most practical definition used for Gram-positive and Gram-negative bacteria is ‘resistance to three or more antimicrobial classes according to Magiorakos et al. [3]. MDR could also be defined as the insensitivity or resistance of a microorganism to administered antimicrobial medicines (which are structurally unrelated and have different molecular targets) despite earlier sensitivity to the same medicines [4]. According to Nikaido [5], multidrug resistance in bacteria cells come about by their accumulation, on resistance (R) plasmids or transposons, of genes, with each coding for resistance to a specific agent, and/or by the action of multidrug efflux pumps, each of which can pump out more than one drug type. This MDR abilities lead to ineffective ailment treatment, resulting in its persistence, infection’s spread and high cost [2,6].

The hospital environment have been said to be an active reservoir for infectious microorganisms, being the meeting point for people with diverse disease etiological agents and susceptible individuals [7-9]. Nikaido [5] mentioned that *Staphylococcus aureus* has a known nosocomial, multi-drug-resistant strain referred to as the methicillin-resistant *Staphylococcus aureus* (MRSA).

MRSA was initially controlled but currently is also resistant to other antimicrobials like the aminoglycosides, chloramphenicol, lincomamides, macrolides and tetracycline [1]. This study was conducted as part of evidence to buttress the efficacy of *Staphylococcus* infections in young children and the scale to which MDR pathogens are becoming threats to
the health of the younger generation amongst the Mkpat Enin, Akwa-Ibom State populace.

Materials and Methods

Study facility, group and sample collection

The study facility is in a growing town and services a couple of adjoining communities, with a number of established institutions, stable commerce and ever growing population. The General Hospital, Ikot-Ekpaw, in Mkpat Enin LGA, Akwa-Ibom State, South-South Nigeria, has Out-patient department, Post-natal ward, Pediatric ward and servicing laboratories. The study focused on children aged 1-15 years.

A hundred (100) sterile swabs samples from skin, nostrils, wound, throat and ear was sourced over a period of three months. Once gotten, the swabs were labelled, placed in an ice pack and taken to the Microbiology laboratory, Akwa-Ibom State University.

Sample analysis

The sterile swab sticks were depth into peptone water and incubated at 37°C for 24 hours. A loop full from each sample was streaked on separate Mannitol salt agar (MSA) plates and incubated at 37°C for 24 hours. Discrete golden yellow colonies were sub-cultured, purified and preserved. Only Gram positive cocci bacterial colonies were further tested for catalase and coagulase.

Antibiotics susceptibility test

Confirmed Staphylococcus aureus isolates were tested for their susceptibilities to various selected commercial antibiotic drugs like Ciprofloxacin, Erythromycin, Levofloxacin, Gentamicin, Ampiclox, Rifampicin, Amoxicillin, Streptomycin, Norflxacin and Chloramphenicol. Overnight cultures using Kirby-Bauer method [10] were inoculated on Mueller-Hinton agar (Oxoid, Uk), cultures adjusted to 0.5 McFarland standard. After pre-diffusion, the plates were inoculated at 37°C for 24 hours.

Diameter of zones of inhibition (IZDs) produced by the antibiotics were measured and recorded in millimeter. Thereafter, the Multiple antibiotics resistance (MAR) index was determine for each isolate using a formula MAR = x/y, where x is the number of antibiotics to which test isolate displayed resistance and y is the total number of antibiotics to which the test organism has been evaluated for sensitivity [11,12].

Results

Result for the prevalence of S. aureus is as shown in Table 1. The result table shows that of the 28 confirmed, wound samples had the highest number, which was followed by samples from children skin swabs. The least number of confirmed Staphylococcus aureus isolates were from the ear swabs.

The susceptibility pattern of the 28 confirmed test isolates to the selected commercially available drugs (amoxicillin, ampiclox, chloramphenicol, ciprofloxacin, erythromycin, gentamicin, levofloxacin, norflxacin, rifampicin and streptomycin) is as shown in Figure 1.

Ciprofloxacin was the most effective drugs against the test organism, followed by Levofloxacin and Norflxacin. Confirmed Staphylococcus aureus isolates had high resistance for Amoxicillin, closely followed by their resistance for Ampiclox.

Table 1 Prevalence of Staphylococcus aureus from clinical samples.

| Type of Specimen | Sample Size | Number of S. aureus isolated | Total percentage |
|------------------|-------------|----------------------------|-----------------|
| Skin             | 20          | 8                          | 29              |
| Nose             | 20          | 5                          | 18              |
| Wound            | 20          | 9                          | 32              |
| Throat           | 20          | 4                          | 14              |
| Ear              | 20          | 2                          | 7               |

Figure 1 Antibiotics susceptibility pattern of Staphylococcus strains from clinical sample.

Table 2 has the multiple antibiotics resistance (MAR) index result which shows that 85.7% of the confirmed test isolates were multi-drug resistant (showing resistance to three or more classes of antibiotics). Only 14.3% of the isolates showed resistance to only two classes of antibiotics.

Results from this study show that 60.7% of test isolates had resistance for four or more antibiotics drugs. MAR index value for 96.4% of the test isolates reveal cases of source drug abuse.
### Table 2 Antibiotic Resistance Pattern and MAR index of *Staphylococcus aureus*.

| S/N | Isolate code | Antibiotic Resistant Pattern | MARI | Antibiotic Resistant Class |
|-----|--------------|-------------------------------|------|---------------------------|
| 1   | S9           | RD, S                         | 0.2  | RIF, AMG                  |
| 2   | S8           | APX, AMX, CH                  | 0.3  | PEN, CHL                  |
| 3   | N3           | LEV, APX, AMX                 | 0.3  | QUI, PEN                  |
| 4   | W1           | CN, AMX, CH                   | 0.3  | AMG, PEN, CHL             |
| 5   | W9           | APX, AMX, S                   | 0.3  | PEN, AMG                  |
| 6   | S10          | CN, APX, RD, AMX              | 0.4  | AMG, PEN, RIF             |
| 7   | T6           | E, APX, AMX, NB               | 0.4  | MAC, PEN, QUI             |
| 8   | T8           | CPX, E, APX, AMX              | 0.4  | QUI, MAC, PEN             |
| 9   | S2           | E, APX, RD, AMX, S            | 0.5  | MAC, PEN, RIF, AMG        |
| 10  | S3           | E, CN, APX, AMX, S            | 0.5  | MAC, AMG, PEN             |
| 11  | N1           | E, CN, APX, AMX, CH           | 0.5  | MAC, AMG, PEN, CHL        |
| 12  | N2           | CN, APX, AMX, S, CH           | 0.5  | AMG, PEN, CHL             |
| 13  | W2           | E, LEV, APX, AMX, S           | 0.5  | MAC, QUI, PEN, AMG        |
| 14  | W5           | E, APX, AMX, S, CH            | 0.5  | MAC, PEN, AMG, CHL        |
| 15  | W6           | CN, APX, AMX, S, CH           | 0.5  | AMG, PEN, CHL             |
| 16  | W7           | APX, AMX, S, NB, CH           | 0.5  | PEN, AMG, QUI, CHL        |
| 17  | W10          | E, LEV, APX, AMX, S           | 0.5  | MAC, QUI, PEN, AMG        |
| 18  | T3           | E, APX, AMX, S, CH            | 0.5  | MAC, PEN, AMG, CHL        |
| 19  | E5           | E, CN, APX, RD, AMX           | 0.5  | MAC, AMG, PEN, RIF        |
| 20  | S1           | E, CN, APX, RD, AMX, S        | 0.6  | MAC, AMG, PEN, RIF        |
| 21  | S5           | E, LEV, APX, RD, AMX, S       | 0.6  | MAC, QUI, PEN, RIF, AMG   |
| 22  | S6           | CN, APX, RD, AMX, S, CH       | 0.6  | AMG, PEN, RIF, CHL        |
| 23  | N5           | CPX, CN, APX, AMX, S, CH      | 0.6  | QUI, AMG, PEN, CHL        |
| 24  | W8           | E, CN, APX, AMX, S, NB        | 0.6  | MAC, AMG, PEN, QUI        |
| 25  | N8           | E, CN, APX, RD, AMX, S, NB    | 0.7  | MAC, AMG, PEN, RIF, QUI   |
| 26  | W3           | LEV, CN, APX, RD, AMX, S, NB  | 0.7  | QUI, AMG, PEN, RIF        |
| 27  | T2           | E, CN, APX, RD, AMX, S, CH    | 0.7  | MAC, AMG, PEN, RIF, CHL   |
| 28  | E9           | CPX, E, CN, APX, AMX, S, CH   | 0.7  | QUI, MAC, AMG, PEN, CHL   |

KEY: CPX: Ciprofloxacin; E: Erythromycin; LEV: Levofloxacin; CN: Gentamicin; APX: Ampiclox; RD: Rifampicin; AMX: Amoxicillin; S: Streptomycin; NB: Norfloxacin; CH: Chloramphenicol; MARI: Multiple Antibiotic Resistance Index; RIF: Rifamycins; AMG: Aminoglycosides; PEN: Penicillins; CHL: Chloramphenicol; MAC: Macrolides; QUI: Quinolones

### Discussions and Conclusion

Data for isolate occurrence and confirmation showed that there were more confirmed *Staphylococcus* isolates from wound swab-samples, followed by skin sample-swabs. Parta et al. [13] also recorded very high *Staphylococcus* number from wound swabs. This high isolate-numerical is suggestive of the exposed nature of the sampling points. This is supported by findings presented by Nimmo et al. [14], who found more *Staphylococcus* isolates on exposed body surfaces than the internal parts. While uncovered wounds have sticky surfaces and the skin is continuously exposed, it is therefore easy for such high microbial numbers to be recorded.

Susceptibility data showed that all the confirmed and tested isolates resisted two or more antimicrobials. Qureshi et al. [15] also isolated MRSA that resisted multiple anti-microbials from hospital specimens. This study’s result showed a higher MDR percentage than the “nearly half” proportion reported by Nimmo et al. [14].

More than 96.4% of the MAR indices were 0.3 from this study evaluation. This assertion is indicative that resistance to these multiple drugs come from over exposure of the isolates.
to drugs, making them adapt or resistant to them with recurrent treatments. The high case of MDR amongst microorganisms can drastically be reduced by mere restricting the indiscriminate and readily availability of these drugs over the counter [14].

Many studies have shown that prolonged stays in hospitals increases the risk for colonization or infection with MDR Staphylococcus aureus. This reflects an inherent risk in acquiring MDR organisms through environmental contaminations and hospital stay conditions. With infant patients, another potential transmission route is through infected staff members handling and this calls for special care [16,17].

The isolation of MDR Staphylococcus aureus from infant patients is a call for the proper implementation of contact precautions during hospitalization especially in developing countries [16,18].

**Ethical Consideration**

The study was approved by the ministry of health, Akwa Ibom State. Permission was obtained from General Administration of the General hospital prior to collecting any data. Participants’ privacy and confidentiality have been assured (no names have been used, only numbers were used) and all data and results have been handled and treated confidentially.

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