

Levels of Biofilm Expression in *Klebsiella pneumoniae* isolates strains exposed to Herbal Drugs

Background: There is continuous rise in antimicrobial resistance globally and factors responsible for this occurrence especially in developing countries are yet to be properly elucidated. Due the financial implications of antimicrobials individuals in developing countries such as Nigeria resort to the consumption of herbal drugs to treat infections.

Aims: To investigate the levels of biofilm expressed in *Klebsiella pneumoniae* isolates pre-treated with herbal drugs.

Methodology: Biofilm assay was performed using 24-well polystyrene plates which mimic the surface for bacterial attachment. Control and clinical isolates strains of *K. pneumoniae* were pre-exposed to different concentrations of herbal solutions (Beta cleanser [Bet], Goko alcoholic bitters [Gab], Goko bitters [Gob], Danko solution [Dan], and Ruzu bitters [Ruz]) (100, 50, 25, 12.5, and 6.25%) in 24-well plate and incubated overnight at 37°C. Cell-to-cell surface attachment of *K. pneumoniae* was recorded by obtaining a photograph of the inoculum in the 24-well plate. Crystal violet method was further used to quantify the level of biofilm attached to the surface of the 24-well plate. Results were analysed using student t-test with Graph pad prism 5.

Results: Cell-to-cell biofilm formation was seen in different drugs used but higher in Bet and Gob. Bet (25%) and Ruz (Ruzu bitter 50%) showed significant level of attached biofilm formed compared to untreated control. This results show that Bet, Gob and Ruz has the ability to induce biofilm in *K. pneumoniae* isolates.

Conclusion: Some herbal drugs could predispose *K. pneumoniae* to enhance its production of biofilm.

Keywords: Biofilm, *Klebsiella pneumoniae*, herbal drug, antimicrobial resistance

1. INTRODUCTION

Since the introduction of antimicrobial agents there have been several observations of the development of antimicrobial resistance in many species of bacteria. The first 'miracle' antibiotics discovered was Penicillin [1]. Resistance to Penicillin was later known to have been caused by Penicillinase, a member of β -lactamases that cleaves the benzylpenicillin. In less than 20 years of the introduction of Penicillin, a rapid increase in the production of penicillinase was observed. This observation was noted for tetracycline, penicillin and macrolide at the end of 1950s. This led to the generation of different strains of microbes, resulting in difficulty in management of infections.

Antimicrobial resistance is a serious health concern that impedes the management and prevention of infections. Different cases of antimicrobial resistance have been seen around the world [2]. Some of these cases of antimicrobial resistance include the emergence of resistant strains of tuberculosis. Globally, this has have been seen or observed in discovered with 4.5 million recent cases of antimicrobial resistance in tuberculosis seen globally in 2012. Other cases of resistance have been observed in other bacteria pathogens such as *Escherichia coli*, *Staphylococcus aureus*, and *Klebsiella pneumoniae*. *E. coli* resistance has now been seen in fluoroquinolone, a widely used antibiotic for the treatment of urinary tract infections. Some isolates of *S. aureus* have shown resistance to first-line drugs. Resistance in *K. pneumoniae* to carbapenem, a last resort antibiotic, is now in all parts of the globe [2].

A number of mechanisms for antibiotics resistance and spread have been discovered. The horizontal gene pool consisting of the mobile genetic elements is responsible for the lateral transfer of genes. This can occur either within individual species or among different species. Multidrug resistance mechanisms occur naturally via erroneously replication or transfer of resistant traits [3]. The force driving this process is the selective force of antimicrobial utilization. This is very notable in hospitals

34 | environments where clear correlation between antimicrobial use and development of resistance can
35 | be seen [4], [5], and [6].

36 | The pathogenesis and outcome of *K. pneumoniae* infection depends on the virulence factors it
37 | produces in the course of the infection. An important virulence factor in this bacterium is the ability to
38 | produces extracellular polysaccharides called biofilm. Bacteria form biofilm in order to successfully
39 | invade and damage the host tissue. Biofilms are surface-attached extracellular polysaccharide matrix.
40 | It could lead to life-threatening bacteremia when formed on medical devices such as catheters [7].
41 | Biofilms pose serious challenges to drug treatment by resisting antimicrobial actions at concentrations
42 | of up to a thousand folds that could easily eliminate free living or planktonic cells. Factors enhancing
43 | biofilm-mediated resistance characteristic include; reduction in the proliferation rate of biofilm [8],
44 | inefficient sequestering off antimicrobial agent within biofilm matrix [9] and presence of "persister"
45 | cells.

46 | The aim of the current study was to examine the hypothesis that exposure of *K. pneumoniae* isolates
47 | to herbal treatments could increase the production of biofilm. The results obtained were compared to
48 | a control conditions (untreated conditions). The data from the biofilm assay demonstrates that pre-
49 | exposure of *K. pneumoniae* strains to some herbal drugs not only results in surface biofilm but also
50 | increases *K. pneumoniae* biofilm attachment to polystyrene plate.

Comment [p1]: These last two sentences are not necessary or suitable here

52 | 2. MATERIALS AND METHODS

54 | 2.1 Collection of Drugs

55 | Locally-made herbal drugs used in this study are Beta cleanser [Bet], Goko alcoholic bitters [Gab],
56 | Goko bitters [Gob], Danko solution [Dan], and Ruzu bitters [Ruz]. They were purchased from Mile 3
57 | market, Port Harcourt, Rivers State, Nigeria.

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59 | 2.2 Determination of the concentrations of the herbal drugs

60 | The concentrations of the herbal antimicrobial solutions were determined by evaporating 1 ml of the
61 | different solutions to dryness in test tubes. The differences in the weight of the test tubes after
62 | dryness were determined. The weight differences obtained were: Goko Alcoholic bitters [Gab] (0.09
63 | g/ml), Ruzu bitters [Ruz] (0.29 g/ml), Beta Cleanser [Bet] (0.09 g/ml), Goko Cleanser Herbal mixture
64 | [Gob] (0.09 g/ml).

65 | 2.3 Collection of Organisms

66 | The laboratory strain also known as control strain of *K. pneumoniae* ATCC 13883 was purchased
67 | from Sigma United Kingdom while the clinical strain was obtained from Lahor Research Laboratory,
68 | Benin, Edo State, Nigeria.

69 | 2.4 Media Preparations

70 | 2.4.1 Culture media Tryptone Soya Agar (TSA) and Tryptone Soya Broth (TSB)

71 | The microbial media used were tryptone soya agar (TSA) and tryptone soya broth (TSB). These were
72 | prepared according to the manufacturer's instructions and autoclaved for 15 minutes at 121°C. TSA
73 | was aseptically poured into sterile Petri dishes and TSB was stored in storage bottles for subsequent
74 | use.

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75 | 2.5 Biofilm attachment assays

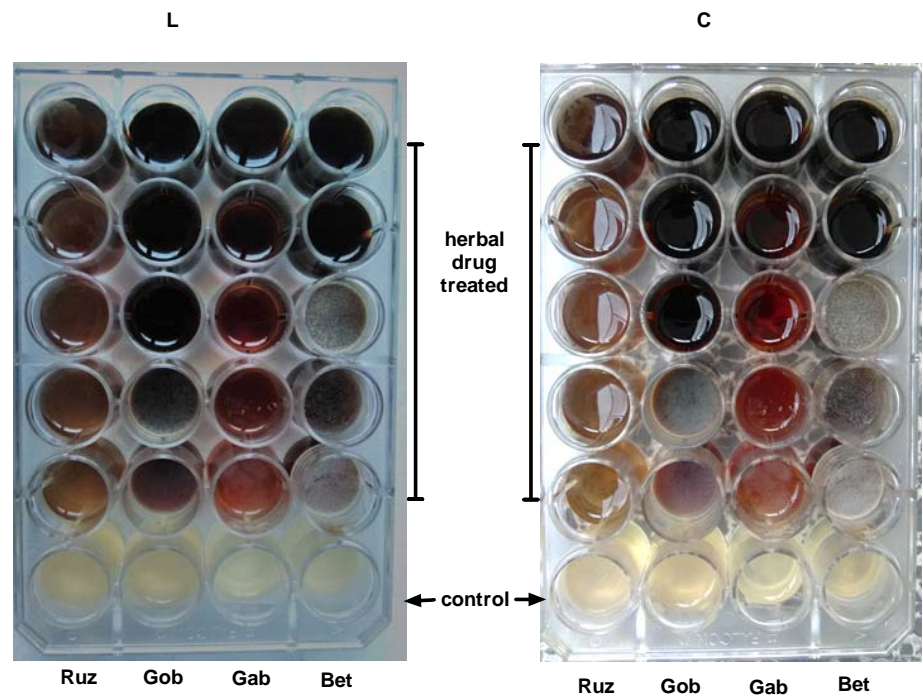
76 | The biofilm assay used in this study is modified from the method used by Lyte et al. [11]. *K.*
77 | *pneumoniae* strains were grown in TSB overnight to log phase (Optical Density 0.5) were diluted to
78 | 1:100 in TSB supplemented with 100%, 50%, 25%, 12.5% and 6.25% of the various original
79 | concentrations of locally-made drugs [Bet, Gab, Gob, and Ruz] stated in section 2.2. A negative

80 control (without herbal drug supplementation) was performed alongside. The cultures (200 µL) were
 81 transferred into a 24-well polystyrene microtiter plate. Wells containing sterile growth medium were
 82 carried out to check for contamination. The plates were incubated at 37°C for 24 and 48 hrs and
 83 photograph of the surface biofilm were taken. The media and loosely adhered bacteria were removed
 84 by vigorously tapping the plate on a tray. Wells were re-washed three times with normal saline to get
 85 rid of any remaining non-adherent bacterial cells and media. Plates were air-dried at about 45°C for 1
 86 hr. Bacteria wells were stained with 1000 µL of 2% crystal violet stain for 15 minutes at room
 87 temperature. After stain was removed, plates were washed twice in normal saline and plates were
 88 dried overnight. Plates were incubated in 1000 µL of 95% ethanol for 10 minutes to solubilise the
 89 crystal violet stains. The attachment of bacterial was quantified by measuring the absorbance of the
 90 crystal violet at 595 nm. The experiment was performed in triplicate on at least three separate
 91 occasions. Data were analysed on Graph Pad Prism 5.0.

93 **3. RESULTS**

94 **3.1 Cell-to-cell attachment**

97 | *K. pneumoniae* isolates produced showed a surface biofilm formation in Gob and Bet in the
 98 laboratory strain but only found in Bet for the Control strain when viewed from the surface (Figure
 99 3.1). No surface biofilm were seen in Gab and Ruz. The two highest concentrations of all the drugs
 100 two did not show any level of surface biofilm induction. The clinical strain showed a higher level of
 101 cell-to-cell aggregation in the Bet compared to the control.



102
 103 | **Figure 3.1. Surface biofilm formation in *K. pneumoniae* isolates exposed to some herbal**
 104 **solutions. Biofilm levels were analysed after 24 hrs of exposure to herbal preparations using**
 105 **spectrophotometer at 595 nM. Beta cleanser [Bet], Goko alcoholic bitters [Gab], Goko bitters [Gob], Danko**
 106 **solution [Dan], and Ruzu bitters [Ruz], L: Laboratory strain, C: Clinical strain.**

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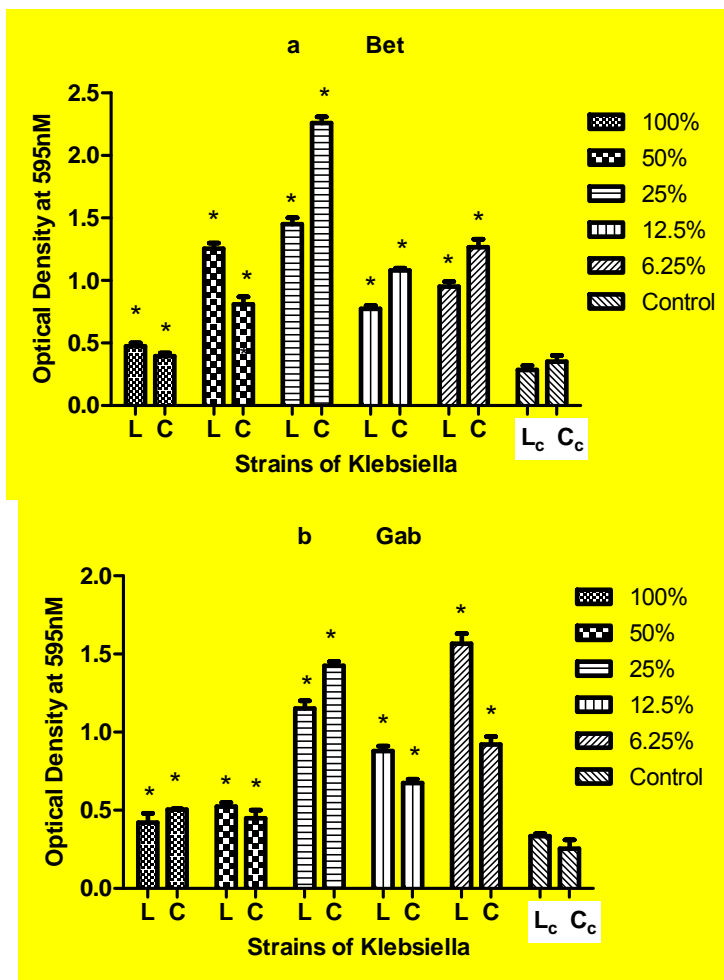
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110 3.2 Biofilm analysis with crystal violet assay

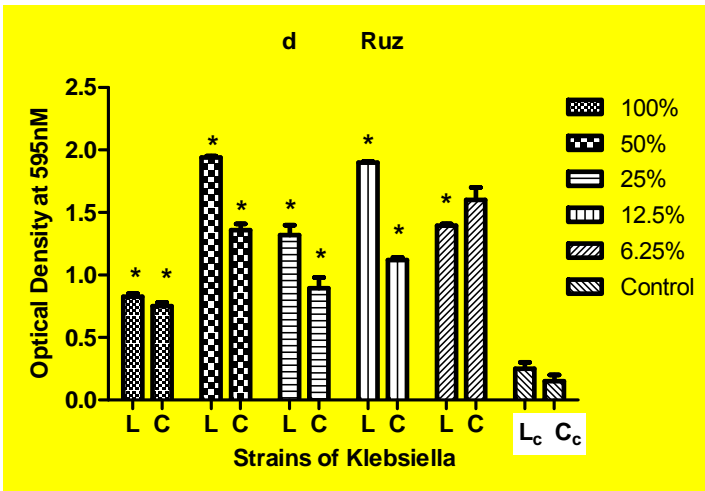
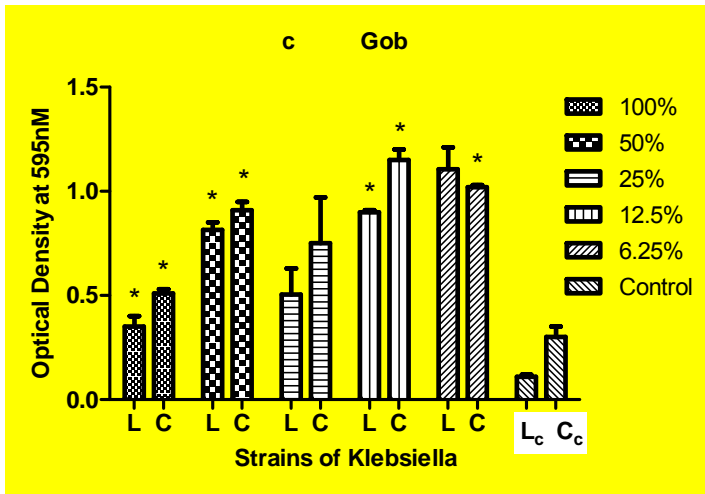
111 Figure 3.2 shows the level of biofilm produced in *Klebsiella pneumoniae* exposed and unexposed. In
112 order to investigate the ability of *K. pneumoniae* to attach to surface of medical devices a modified
113 method of crystal violet biofilm assay was used. The biofilm was detected as optical density measured
114 at 595 nM. In the experiment, all drugs showed higher levels of biofilm induction than the control
115 condition (unexposed). There were similarities in the pattern of biofilm adherence to the polystyrene
116 surface in the different drugs used (Figure 3.2a-d). The unexposed isolates are represented as L_c and
117 C_c. A common trend observed in the experiment is that higher concentrations of the locally-made
118 herbal preparations exhibited reduced level of biofilm production. The lower concentrations of the
119 drug used showed a higher level of biofilm induction. The highest level of biofilm induction is observed
120 in Bet (OD= 2.3), followed by Ruz (OD= 2.0), then Gab (OD= 1.5) and Gob (OD= 1.3). Figure 3.2a
121 and b showed similar pattern of biofilm production: the 25% concentration showed much higher levels
122 of optical densities. Bet (25%) and Ruz (50%) showed significant level of biofilm formed compared to
123 untreated control.

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Figure 3.2. Levels of expression of Biofilm in *K. pneumoniae*. Levels of biofilm formed were measured after 24 hrs incubation with and without herbal drugs at 595 nM. Data plotted above are mean ± standard deviation of three independent experiments performed in triplicates. * Level of significance compared to control not exposed to herbal drugs (L_c and C_c) using p<0.05). Beta cleanser [Bet], Goko alcoholic bitters [Gab], Goko bitters [Gob], Danko solution [Dan], and Ruzu bitters [Ruz], L: Laboratory strain, C: Clinical strain.

4. DISCUSSION

There are two ways biofilm can be formed in bacteria; cell-to-cell aggregation and attachment to surface [10]. The potential of bacteria to resist antibiotics and form biofilm on medical devices is becoming high in hospital-acquired infections [11]. This investigation analysed the level of this virulence factor in *K. pneumoniae* exposed to some common herbal preparations used in Nigeria. The data on the drug resistance mechanism induction by herbal drugs furthers our understanding and appreciation of the possible causes of drug resistance in Nigeria.

The processes in bacterial biofilm formation initially firstly begin by the firstinitial attaching ment to a surface [11]. Findings from other investigations have shown that pathogenic bacteria recognise

147 inotropic drugs and use them to grow and produce biofilm [12] and [11]. However, information is yet
148 available as to whether these **herbal drugs induce biofilm in *Klebsiella*** spp in similar fashion. Hence,
149 the aim of this project was to investigate biofilm levels in *K. pneumoniae* strains response to exposure
150 to herbal drugs. In this investigation, it was shown that concentration of herbal drugs **within the range**
151 **consumed** could markedly increase biofilm levels of *K. pneumoniae* responsible for its ability to persist
152 in the host.

153 Antimicrobial resistance is a growing problem in **infection control and preventioncontrolling**
154 **infection**. Biofilm formation **in *K. pneumoniae*** is an aspect of **its** pathogenicity that enhances the
155 colonization of a host. We demonstrated that herbal drugs most commonly consumed by sick patients
156 (Bet, Gab, Gob and Ruz) all markedly increased *K. pneumoniae* biofilm formation on polystyrene
157 surfaces. This is a crucial discovery as bacterial ability to colonise surfaces such as catheters and
158 other hospital plastic devices is a reason thought to influence patients to acquire pneumonia and
159 other blood related infections [13, 14, 15].

160 Biofilm analysis of herbal drugs induction of biofilm observed in *K. pneumoniae* showed a minimum of
161 two fold increase compared to control (Figure 3.2a) and a maximum of 8-fold increase (Figure 3.2d).
162 A similar study by Freestone et al. [12] demonstrated that *Pseudomonas aeruginosa* **another gram**
163 **negative close** organism **was** also responsible for pneumonia-associated infection showed increase
164 in biofilm level using crystal violet method. Their study showed a minimum of 1.5-fold increase and
165 maximum of 2-fold induction caused by stress factor such as catecholamine. **This is similar to the fold**
166 **increase observed by Freestone et al [12] using catecholamines as a biofilm inducing factor.** This
167 suggests that herbal drug **could be a** stronger inducer of biofilm than catecholamine **in vitro** and
168 **promote the ability of *K. pneumoniae* to cause infection.** Further investigations into the untoward
169 effect of biofilm production such as antibiotic resistance are necessary.

170 A number of people within rural and urban settings in Nigeria consume herbal solutions, some as a
171 way of life while others for the purpose of eliminating infections. **Consequentially, the observations**
172 **from this investigation show the possibility of the effect of consumption of some herbal antimicrobial**
173 **drugs by predisposing herbal drug consumers** to opportunistic infections by enhancing *K. pneumoniae*
174 biofilm formation. **Theirs consumption habit of the herbal drugs by individuals promotes bacteria**
175 **colonization since the bacteria tend to their colonization their survive more survival** in stressful
176 **conditionsituations.** The clinical importance of this *in vitro* investigation is highlighted by the fact that
177 it employed the same herbal **drug** solutions consumed by people in Nigeria together with the low
178 inoculum of bacterial which represents the infectious dosage present during the initial stage of
179 infection [16]. The ***K. pneumoniae* isolates produced biofilm when they were exposed to some herbal**
180 **drugs and this** findings **in** this **current** study further **supports buttress** the observations in previous
181 studies **by Monsi et al** [17, 18] that herbal antimicrobial agents induce resistance, through **suggesting**
182 **that** the production of biofilm_ **could be a mechanism of resistance development employed by**
183 **some herbal drugs.**

184 185 **4. CONCLUSION** 186

187 This study was able to demonstrate for the first time that **in vitro** exposure of *K. pneumoniae* to herbal
188 antimicrobial drugs could **induce biofilm in *K. pneumoniae*.** **However, the** mechanisms behind this
189 biofilm induction are yet to be **determined and warrants further studies,discovered.**
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CONSENT (WHERE EVER APPLICABLE)

This was not applicable in this research.

ETHICAL APPROVAL (WHERE EVER APPLICABLE)

This was not applicable in this research.

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