

Original Research Article

ANTIBIOTIC SUSCEPTIBILITY PATTERN OF BACTERIA IN SACHET WATER, SOLD UYO METROPOLIS, AKWA IBOM STATE

ABSTRACT

Aims: This study was carried out to examine the microbiological quality of sachet drinking water sold in Uyo metropolis, Akwa Ibom State.

Study design: Sachet water was randomly sampled in Uyo Metropolis.

Place and Duration of Study: Department of Microbiology Akwa Ibom State University, Nigeria, between June 2018 and November 2018.

Methodology: Six different brands of sachets water sold and consumed were studied for their physical and microbiological qualities. Thirty (30) sachets water from six (6) different brands, respectively were serially diluted and cultured on Nutrient agar, Eosin Methylene Blue agar, MacConkey agar and Salmonella Shigella agar, while Muller Hinton agar was used for sensitivity test. Pure isolates suspension were standardized with 0.5 McFarland turbidity standard and were subjected to antibiotics susceptibility test using Agar Diffusion method.

Results: The microbial counts ranged from 9.20×10^1 CfU/ml to 1.77×10^2 CfU/ml. Bacterial isolates include; *Klebsiella* sp., *Escherichia* sp., *Staphylococcus* sp., *Salmonella* sp., *Pseudomonas* sp., *Citrobacter* sp., *Bacillus* sp. All the isolates were completely susceptible to Chloramphenicol, Ampiclox, Tarvid and Peflacine. Low percentage of these isolates were resistant to Erythromycin, Gentamycin, Septrin, Ciprofloxacin, Norfloxacin, Chloramphenicol, Levofloxacin and Rifampicin but were highly resistant to Amoxil, Ceporex, Augmentin, Ampicillin, Nalidixic acid and Stretomycin. Some of the sachet water brands from bacteriological standpoints did not meet the World Health Organization Standard.

Conclusion: This study indicted sub-standard packaged waters as a vehicle of spread of antibiotic resistant pathogens, and this poses a high risk to public health. Hence, routine monitoring of producers of sachet water should be enforced.

Keywords: Sachet water, Antibiotics resistance, Public health, water standards.

1. INTRODUCTION

The safety and quality of drinking water has become a public health concern in the world. In Nigeria, high demand of safe drinking water cannot be overemphasized considering the inability of the government to provide adequate pipe-borne water to the populace. Water is

Comment [U1]: Antibiotic sensitivity

Comment [U2]: Wrong calculation.

Comment [U3]: have

Comment [U4]: for

Comment [U5]: change

23 known to be the dwelling place for most bacteria and other microorganisms which cause a
24 variety of waterborne infections [1] and the World Health Organization (WHO) estimated that
25 1.1 billion of the world's population does not have access to safe water. In addition to this,
26 80% of diseases and one-third of deaths in developing countries are due to consumption or
27 drinking of contaminated water [2]. The associated health risks from the consumption of
28 unsafe drinking water vary throughout the world depending on the chemical or
29 microbiological contaminants present in the environment [3]. Many of the bacteria isolated in
30 water distribution systems are opportunistic pathogens. The presence of high numbers of
31 opportunistic pathogens in drinking water is of concern because these microorganisms can
32 lead to infection of certain segments of the population (newborn babies, the sick, and the
33 elderly) [4]. According to the guideline set by the World Health Organisation, quality drinking
34 water must not contain *Escherichia coli* or thermotolerant coliform bacteria, giardia worms,
35 viruses, *Cryptosporidium* spp, *Legionella pneumophila*, *Entamoeba histolytica* and other
36 opportunistic pathogens such as *Clostridium* species, *Klebsiella* species and *Pseudomonas*
37 [2]. The guideline further stated that the water should be tested against the presence of
38 highly virulent pathogens such as *Salmonella typhi*, *Shigella dysenteriae* and *Vibrio cholera*
39 that are responsible for typhoid, bacillary dysentery and cholera diseases, respectively which
40 arises due to the high level of organic decay and fermentation on tropical waters. All these
41 bacteria must not exist in water that are meant for drinking, hence, sources of water for
42 packaged water are subjected to laboratory test by the public analyst in which any of the
43 bacteria must not be found or detected in any 100 ml water sample. "Sachet water is not
44 sterile" according to Linda [3]. Although, sachet water is assumed to be free from certain
45 pathogen during standard treatment processes, but certain organisms are used to confirm
46 the sterility of the water such as coliforms which act as indicator organisms used to assess
47 the safety of water and thus give an idea of the degree of contamination associated with
48 intake of such sachet water [4,5]. Antibiotics is been revolutionized in medicine diversely,
49 saving many lives because it had a major impact on the rate of survival of pathogens from
50 infection. But with this great and remarkable benefit, it is sad to know it is also the bedrock of
51 many other diseases due to their resistance strains. Recently, almost all important bacterial
52 infections are becoming resistant to antibiotics, and these changing patterns caused a
53 demand for new antibacterial agents. Antimicrobial resistance occurs when bacteria adjust
54 or adapt in ways that permits them to stay alive in the presence of antibiotics designed to kill
55 them. Bacteria evolve resistance to these drugs, typically by acquiring chromosomal
56 mutations and multidrug resistant plasmid which has become a public health concern [6,7,8].
57 Antibiotics were formally defined to distinguish their biochemicals which are produced by
58 microorganism from the organic chemicals synthesized in the laboratory. But due to recent
59 development, the distinction between both is no longer meaningful due to the fact that the
60 biochemical structures of many naturally occurring antibiotics are now being synthesized by
61 organic chemist and currently, many antibiotics used in medicals are in the chemically
62 modified forms of the microbial biosynthetic form [9].
63 Antibiotic resistance occurs when the sensitivity of an organism decreases against an
64 antibiotic when compared to officially available breakpoints, usually measured as a decrease
65 in "inhibition zone diameter". The increased use of antibiotics is often associated with
66 increased resistance of bacteria to these chemicals, especially in the hospital setting [10]. A
67 lot of transmissible diseases are waterborne. Many harmful microbial contaminants have
68 been confirmed to be associated with potable water sources. Many people have resorted to
69 patronizing sachet water with the belief that it is pure-hence, fondly called 'pure water'. It is
70 possible that this so-called pure water is not pure after all; hence it may harbour harmful
71 microorganisms as producers of such water may not pay adequate attention to microbial
72 quality. Identification of the major harmful microbial contaminants (*Escherichia coli*,
73 *Salmonella*, *Shigella*, etc.) present in the sachet water is important in assessing its safety.
74 Free from contamination with faecal matter is the most important parameter of water quality
75 because the human faecal matter is generally considered to be a greater risk to human

Comment [U6]: Sentence is too long. Please rephrase.

Comment [U7]:

Comment [U8]: Such as?

Comment [U9]: is

Comment [U10]: rephrase

Comment [U11]: Please explain more.

Comment [U12]: Please rephrase

Comment [U13]: are

Comment [U14]: change

Comment [U15]: major

Comment [U16]: permit

Comment [U17]: change to other words

Comment [U18]: spelling

76 | health as it is more likely to contain human enteric pathogens [11]. There is a need to
77 | constantly access the water quality of water sources available to members of any community
78 | at intervals. This will help monitor or track and prevent the sudden outbreak of waterborne
79 | infections. It is also important to know the antibiotics susceptibility pattern of microorganism
80 | common in our environment in case of any outbreak. This research was borne as a result of
81 | the widespread use of sachet water in Nigeria especially in Akwa Ibom State, conflicting
82 | results on the safety conducted at different locations in the country and lack of data on the
83 | safety of sachet water locally available. This research is aimed at determining the antibiotic
84 | resistant pattern of bacterial isolates obtained from sachet water by testing them against
85 | some of the commonly used antibiotics; taking notes of the antibiotic resistant strains.
86
87

Comment [U19]: chose the best word

Comment [U20]: ?

Comment [U21]: an

88 | 2. MATERIALS AND METHODS

89 | 2.1 Study area

90 | Three major areas in the major city of Uyo in Akwa Ibom State where strategically selected
91 | for this study. The areas comprised of towns where sachet-packaged drinking water is sold
92 | by hawkers. They included: Abak road, Aka road, and Oron road.
93
94

95 | 2.2 Sample collections

96 | A total of thirty (30) sachet water of six different brands was collected randomly from
97 | various parts of Uyo metropolis in Akwa Ibom state and taken to the laboratory for analysis.
98 | The samples were coded as; BC, GO, FD, RS, ML, and CV to reflect the respective brands.
99 | They were collected and transported in clean ice containers and stored at 4.0°C for 30-60
100 | minutes so as to maintain the properties of the samples before the commencement of
101 | analysis. Hygienic and aseptic methods were also observed during sampling of the sachet
102 | water.
103

Comment [U22]: to

Comment [U23]: technique

Comment [U24]: applied.

104 | 2.3 Sterilization

105 | Microbiologically, sterilization is simply any process that eliminates, removes, kills, all forms
106 | of life and any other biological agents (such as bacteria, fungi etc) present in a specific
107 | region. This was achieved through the process of autoclaving (steam under pressure) and
108 | oven drying. Media, water, and other heat stable liquids were sterilized in the autoclaved at
109 | 121°C for 15minutes, while glassware were sterilized at 160°C for at least 2 hours. Also, the
110 | working environment was always kept neat, tidy and sterile by the process of disinfection.
111

Comment [U25]: Unimportant section.

112 | 2.4 Enumeration of microbial load in water sample

113 | 2.4.1 Preparation of the samples

114 | Using aseptic method, six (6) different beakers were labelled according to the 6 different
115 | brands of waters. Five sachets were mixed from each brand to obtain 100ml homogenous
116 | sample in the beaker.
117

118 | 2.4.2 Pour plating method

119 | One millilitre of appropriate dilution was aseptically pipetted into a sterile labelled
120 | petri dish and this was done in duplicates. Appropriate medium (Nutrient agar, Eosin
121 | Methylene Blue, MacConkey agar, Salmonella-Shigella Agar) at 45°C were poured
122 | aseptically into the inoculated petri dishes and swirled gently to mix. They were inversely
123 | incubated at 37°C for 24-48hours. At the end of the incubation period, colonies were counted
124 | and the count for each plate expressed as colony forming units per gram (cfu/mL) of the
125 | sample suspended.
126

Comment [U26]: What dilution?

Comment [U27]: Please rephrase

127 | Nutrient agar (NA) to determine the total viable bacterial Count, Eosin Methylene Blue agar
128 | (EMB) to enumerate *Escherichia coli*, MacConkey agar (MAC) for coliform count and

129 Salmonella-Shigella agar (SSA) for the determination of *Salmonella* and *Shigella* counts.
130 Culture media were prepared according to the respective Manufacturers specification and
131 sterilized in an autoclave at 121°C at 15 psi for 15 minutes.

Comment [U28]: Rewrite the sentences.

Comment [U29]: Delete

132 133 2.4.3 Purification of colonies

134 Using a fresh nutrient agar medium, 24 hours colonies were picked using a sterile wire loop
135 from the plate and streaked on its surface and incubated for 24 hours at 37°C to obtain pure
136 culture. After incubation, discrete growths were observed on the lines of streak. Distinct
137 colony was picked aseptically and cultured on a fresh nutrient agar slant and incubated for
138 24 hours at 37°C and stored in a refrigerator at 4°C. The routine laboratory method of
139 Cruickshank *et al.* [12] was used to characterize different isolates. The isolates were
140 identified using their macroscopic, cultural, physiological and biochemical characteristics.

Comment [U30]: Please rewrite.

Comment [U31]: ?

141 142 2.5 Morphological characterization (Gram's reaction)

143 Gram staining was carried out as described by Olutiola *et al.* [13]. Gram stain is one of the
144 differential stains used to characterize bacteria into two main groups: Gram positive and
145 Gram negative. Gram positive stains blue to purple while Gram negative stains pink to red.
146 The colony of the pure cultures of each bacterial isolates was observed for morphological
147 features using Bergey's Manual of Determinative Bacteriology as a standard for comparison.
148 Cell shape was determined under X100 objective of the light microscope after Gram staining
149 procedure. Bacterial smear (not too thick not too thin) was prepared on the slide using an
150 inoculation loop. This was done by introducing a drop of distilled water on grease-free
151 labelled slide followed by the sample and then smeared, air dried and heat fixed. The slide
152 was flooded with crystal violet staining reagent for about 60 seconds, then washed using a
153 gentle indirect stream of tap water for about 2 seconds. The slide was flooded with a
154 mordant (Lugol's iodine) for 15-30seconds. The slide was decolorized using 70% ethanol for
155 10 seconds and washed off. Lastly, the slide was flooded with 0.5% counter stain (safranin)
156 for 30 seconds, and then washed using indirect stream of tap water and air dried. A drop of
157 immersion oil was dropped on the stained sample and observed under the microscope.

Comment [U32]: delete

Comment [U33]: The pure colonies

Comment [U34]: Wrong magnification

Comment [U35]: Standard procedure. No need include.

158 159 2.6 Biochemical Characterization and Identification of Isolates

160 Pure cultures of bacterial isolates were subjected to various biochemical tests according to
161 standard techniques described by Olutiola *et al.* [13] Biochemical tests carried out include;
162 Catalase test, Coagulase test, Indole test, Oxidase test, Citrate test, Fermentation of
163 glucose, lactose, sucrose, maltose and mannitol [14]. Bacterial isolates were identified [using](#)
164 [according to](#) Bergey's Manual of Determinative Bacteriology [15].

165 166 2.7 Antimicrobial Sensitivity Testing

167 Commercially available antibiotic impregnated 8mm sensitivity discs (Abtek Biological Ltd,
168 UK) ~~was~~ were used to determine the drug sensitivity profile of the isolates. Seventeen
169 different antibiotic discs comprising of Tarivia (OFX), Nalidixic acid (NA), Peflacin (PEF),
170 Gentamycin (CN), Augumentin (AU), Ciproflox (CPX), Septrin (SXT), Ceporek (CEP),
171 Streptomycin (S), Ampicillin (PN) for Gram negative and Levoxin (Lev), Amoxicillin (Amx),
172 Norfloxacin (NB), Chloramphenicol (CH), Erythromycine (E), Ampiclox (APX), Rifampin
173 (RD), Streptomycin (S), Ciproflox (CPX), Gentamycin (CN) for Gram positive organisms .
174 The antimicrobial sensitivity test of each isolate was carried out as described by the Kirby –
175 Bauer disc diffusion method as recommended by the National Committee for Clinical
176 Laboratory Standards [16]

Comment [U36]: antibiotics

177 **Procedures:** The turbidity of the bacterial suspensions was compared with 0.5 Macfarland's
178 standard by inoculating the organism into 10ml peptone water and incubated. The
179 standardized bacterial suspension was then inoculated on to Muller Hinton Agar and left to
180 dry for 10 minutes, before placing the antimicrobial sensitivity discs. After incubation, the

181 diameter of the zone of inhibition were measured and compared with zone diameter of
182 interpretative chart [17,18] to determine the sensitivity of the isolates to antibiotics.

183

184 3. RESULTS

185

186 All the water samples were National Agency for Food and Drug Administration and Control
187 (NAFDAC) approved and had factory addresses on them (Table 1). They were all odourless,
188 colourless and clear in appearance and had no batch number, also non had production and
189 expiration dates meaning that the duration between production and consumption cannot be
190 determined. Only FD contained little particles in it. Lastly, all were the same net volume of 50
191 cl.

192

193 Table 2 shows the Total viable count (TVC) after 48 hours of water samples on different
194 media. All the water samples were contaminated with bacteria. A higher value of TVC on
195 Nutrient agar (NA) was 1.34×10^2 cfu/ml from sample FD, Eosin Methylene Blue agar (EMB)
196 plate was 3.1×10^1 cfu/ml from sample ML, MacConkey agar (MAC) plate was 25cfu/ml from
197 sample ML and on Salmonella Shigella agar (SSA) plate it was 5.0 cfu/ml from sample FD.
198 The highest number of organisms (on all the media) was 1.77×10^2 cfu/ml in FD sachet water
199 and the lowest was 7.6×10^1 cfu/ml in CV sachet water.

200

201 Out of 29 bacteria isolate, 7 distinct isolates were obtained while others were replicates of
202 the seven. *Klebsiella* sp. had the highest frequency showing 7 out of 29 representing
203 24.14%, followed by both *Staphylococcus* sp. and *Pseudomonas* sp. with the frequency of 5
204 out of 29 isolates representing 17.24%. Other bacteria isolated included; *Escherichia* sp.
205 with the frequency of 4 out of 29 representing 13.79%, *Salmonella* sp. and *Citobacter* sp.
206 with frequency of 3 out of 29 representing 10.34% and *Bacillus* sp. with the least frequency
207 2 out 29 representing 6.90% as shown in Figure 1.

208 Six brands of sachet water were analyzed and a total of seven bacterial isolates were
209 identified from the sachet water samples. The isolates were initially differentiated on the
210 basis of the cultural and morphological studies after which they were subjected to various
211 biochemical characterization tests. These tests revealed their probable identity as *Klebsiella*
212 sp., *Escherichia* sp., *Staphylococcus* sp., *Salmonella* sp., *Pseudomonas* sp., *Citrobacter* sp.,
213 *Bacillus* sp.

214 *Klebsiella* sp. was most resistant (70%), followed by *Escherichia* sp and *Salmonella* sp. *E*
215 *Escherichia* sp was resistant to 6 (NA, CN, AU, SXT, S, PN and CEP) out of the 10
216 antibiotics tested against it. Same number of antibiotic resistance was recorded for
217 *Salmonella* sp. (NA, CN, AU, S, PN and CEP). The least resistant gram negative isolate was
218 *Citrobacter* sp. (NA, CPX, S, and PN) and *Pseudomonas* sp. All the Gram's negative
219 isolates were resistant to PN and NA. The Gram's positive organisms were less resistant to
220 all the antibiotics they were exposed to. *Bacillus* sp. was resistant to on ciproflox while
221 *Staphylococcus* sp. was resistant to amoxicillin and Gentamycin (Table 3)

222

223

224

225

226

227

228

229

230

231

Comment [U37]: Redo your result

Comment [U38]: Please rewrite.

Comment [U39]: Please rewrite.

Comment [U40]: Please rewrite.

Comment [U41]: Wrong results.

Table 1: Physical examination of the sampled Sachet water brands sold in Uyo metropolis for compliance. Table pattern according to Dada, 2009.

SAMPLE CODE	NAFDAC	PRODUCTION./ BEST FORE DATE	PRODUCERS' NAME & ADDRESS	COLOUR	APPEAR-ANCE	ODOUR	FLOATING PARTICLES	BATCH NO:	NET VOLUME
BC	+	-	+	-	-	-	None	-	50CL
FD	+	-	+	-	-	-	Few	-	50CL
RS	+	-	+	-	-	-	None	-	50CL
CV	+	-	+	-	-	-	None	-	50CL
ML	+	-	+	-	-	-	None	-	50CL
GO	+	-	+	-	-	-	None	-	50CL

233

234

KEY: +: displayed on sample sachet; -: not displayed on sample sachet

235

236

237

238

239

240

241

242

243

244

245

246

247

248

249

250

251
252
253
254

Table 2: Total viable count (TVC) after 48hours of culturing sachet water samples on different media

Sample/ Media	NA	EMB	MAC	SSA	Total no. of organism (cfu/ml)
BC	110	30	15	0	1.55×10^2
FD	134	29	9	5	1.77×10^2
RS	70	8	14	0	9.20×10^1
CV	25	31	20	0	7.60×10^1
ML	20	45	25	2	9.20×10^1
GO	118	18	12	1	1.48×10^2

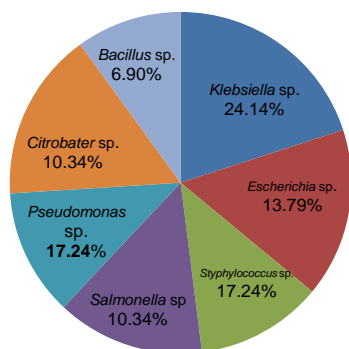
KEYS: NA: Nutrient Agar; EMB: Eosin Methylene blue agar; MAC: MacConkey agar; SSA: Salmonella Shigella Agar

Comment [U42]: How you count the total number of organism? Is wrong to add different cfu from different medium together.

255
256
257
258
259
260
261
262
263
264
265
266
267
268
269
270
271
272
273
274
275
276
277
278
279
280
281
282
283
284
285
286
287
288

UNDER PEER REVIEW

289
290



291
292
293
294
295
296
297
298
299
300

FIG. 1: Percentage frequency of bacteria isolates obtained from sachet water sold in Uyo metropolis

TABLE 3. Antibiotics susceptibility pattern of bacterial isolate from sachet water sold in Uyo metropolis.

S/N	Isolate	Gram Positive Isolates										Gram Negative Isolates										% RESISTANCE	
		AMX	S	NB	CPX	CH	E	LEV	CN	APX	RD	OFX	NA	PEF	CN	AU	CPX	SXT	S	PN	CEP		
1	<i>E. coli</i>											S	R	S	S	R	S	R	R	R	R	R	60
2	<i>K. pneumonia</i>											S	R	S	R	R	R	S	R	R	R	R	70
3	<i>Bacillus sp.</i>	S	R	S	S	S	S	S	S	S	S												10
4	<i>S. aureus</i>	R	S	S	S	S	S	S	R	S	S												20
5	<i>Pseudomonas sp.</i>											S	R	S	S	R	S	S	S	R	R	R	40
6	<i>Citrobacter sp.</i>											S	R	S	S	S	S	R	R	R	S	S	40
7	<i>Salmonella sp.</i>											S	R	S	R	R	S	S	R	R	R	R	60

301
302
303
304
305
306

KEY: Tarivia (OFX), Nalidixic acid (NA), Peflacin (PEF), Gentamycin (CN), Augumentin (AU), Ciproflo (CPX), Septrin (SXT), Ceporek (CEP), Streptomycin(S), Ampicillin(PN) for Gram negative and Levoxin (Lev), Amoxicillin (Amx), Norfloxacin (NB), Chloramphenicol (CH), Erythromycin (E), Ampiclox (APX), Rifampin (RD), Streptomycin (S), Ciproflo (CPX), Gentamycin (CN).

307
308
309
310
311
312
313
314
315
316
317
318
319
320
321
322
323
324
325
326
327
328
329
330
331
332
333
334
335
336
337
338
339
340
341
342
343
344
345
346
347
348
349
350
351
352
353
354
355
356
357
358
359

4. DISCUSSION

This experiment was carried out to determine the microbial quality and the antibiotics resistance pattern among the bacterial isolates from sachet water sold in Uyo with the view of creating public health awareness concerning drinking such water sample. In Nigeria, sachet water is largely taken and they are obtained either from surface or underground sources, and are subjected to various treatment to make it fit for individual consumption, but unfortunately, most of them still fall below the WHO standard from the physical and microbiological analysis [19]. From this analysis, one out of six water samples had particles in it. Meanwhile, all were odourless, colourless, and registered with NAFDAC. Bacteria occurrence was recorded in all the sachet-water samples and the TVC for some were higher than what is acceptable for drinking water (1.0×10^1 cfu/ml) [20].

The presence of pathogenic bacteria (opportunistic pathogens) was recorded which is above the WHO standard for portable water [4]. High occurrence of *Klebsiella* sp. was recorded, followed by *Staphylococcus* sp. Others included *Pseudomonas* sp, *Escherichia* sp., *Salmonella* sp, *Citrobacter* sp. and the least frequent was *Bacillus* sp. Total Viable Count on EMB and MAC for coliform bacteria and the various values obtained for each water sample signified possible faecal contamination. This indicates that the sachet-water samples were contaminated especially with faecal materials, thereby not safe for drinking. Presence of coliforms (*Escherichia* sp. and *Klebsiella* sp. and *Citrobacter* sp.) maybe that some of the water were prepared from shallow and contaminated boreholes. Most of these bacteria are indigenous to aquatic environments [20]. *Salmonella* is also as a result of contaminated water and improper treatment, *Pseudomonas* sp. were also found in water and are considered opportunistic pathogens and *Staphylococcus* sp. isolated from the water samples may have entered the water during packaging or handling since the organism is a normal flora of the human skin [21]. The ingestion of these bacteria with contaminated water constitute public health risks to the immunocompromised members of the population, especially newborn babies, elderly and sick people [22]. The presence of relative heavy load of bacteria in water packaged for drinking purposes has been previously documented in literature [23, 24, 25, 26]. The result of the antibiotics susceptibility testing showed various percentages of antibiotic resistance among the bacterial isolates from packaged water samples. *Escherichia* sp. was highly resistant to six antibiotics and sensitive to only four antibiotics which were; Tarivia (OFX), Gentamycin (CN), Peflacine (PEF) and Ciproflox (CPX). *Klebsiella* sp. was resistant to seven antibiotics and sensitive to Tarivia (OFX), Peflacine (PEF) and Septrin (SXT). *Bacillus* sp. was sensitive to all antibiotics and resistant to only Streptomycin (S). *Staphylococcus* sp. was also highly sensitive to all the antibiotics except Amoxicillin (AMX) and Gentamycin (CN). *Pseudomonas* sp. was also sensitive to most antibiotic except Nalidixic acid (NA), Augumentin (AU), Ampicillin (PN) and Ceporex (CEP). *Citrobacter* sp. was more sensitive to the antibiotics and resistant to only four antibiotics; Nalidixic acid (NA), Septrin (SXT), Streptomycin (S), Ampicillin (PN). *Salmonella* sp. was highly resistant to all the antibiotics except four; Tarivia(OFX), Peflacine(PEF), Ciproflox (CPX) and Septrin (SXT). Generally most of the isolates were resistant to Amoxil, Ceporex, Augmentin, Ampicillin, Nalidixic acid and Stretomycin. The resistance exhibited by *Pseudomonas aeruginosa* and *E. coli* to some of the antibiotics corroborates earlier report from South Eastern Nigeria [27]. The presence of the same type of faecal bacteria in almost all brands shows common source of contamination. It is documented that bacteria harbour series of antibiotic resistant genes which can be transferred to others horizontally [28].

Therefore from observation, a lot of sachet water producers and sellers have emerged making it their major source of income. With this, appropriate health authorities should ensure that producers comply with the government regulations since some of these packaged water may have been produced under unhygienic conditions. Water can be seen as one of the most important, as well as one of the most abundant of those compounds and it is particularly, vital to living organisms [29]. Also, water is like the life wire of the body and

Comment [U43]: Redo your discussion

360 as the basis of life; it is a critical part of human diet. Water constitutes about 90% by weight
361 of the human body [30]. So, water should be treated and the necessary biochemical and
362 microbiological test should be carried out to protect the general public from water-borne
363 disease outbreak.

364

365 5. CONCLUSION

366

367 The study revealed that bacteriological quality of the sachet water brands sold failed to meet
368 the standard for drinking water, even though the bacterial load did not exceed the maximum
369 range of microbial load. However, the bulk of sachet water brands were contaminated by
370 coliform bacteria. It is therefore necessary for sachet water brands to be properly treated and
371 handled to meet the WHO standard for drinking water. To minimise the problem of poor
372 quality of sachet water, government agencies like the NAFDAC and the Environmental
373 Protection Agency should ensure that packaged water manufacturers comply with good
374 manufacturing practices. It is a serious threat to the people of the area if proper
375 measurements are not taken by the concerned authorities. The water sources were
376 contaminated with *Klebsiella* sp., *Escherichia* sp., *Staphylococcus* sp., *Salmonella* sp.,
377 *Pseudomonas* sp., *Citrobacter* sp., and *Bacillus* sp. thus posing a very serious threats to the
378 society. Antibiotic resistance is considered a major problem because many disease causing
379 bacteria are becoming more resistant to the commonly used antibiotics. The bacteria
380 *Klebsiella* sp., *Escherichia* sp., *Citrobacter* sp., showed greater antibiotic resistances. The
381 overuse and misuse of antibiotics can create the conditions for the development of antibiotic
382 resistant bacteria.

383

384 5.1 Recommendation

385 There is need for NAFDAC to intensify efforts in the routine monitoring of activities in the
386 packaged drinking water industry ensuring the safety of sachet drinking water through
387 comprehensive regulatory programs at both the federal and state levels. Also, sample
388 collection and testing of market samples will be a good way of detecting if the water is truly
389 pure as claimed by these producing companies. High emphasis should also be placed on
390 enforcing compliance with Good Manufacturing Practice (GMP) with emphasis on
391 management of raw water source to the consumer product point. Hence, routine monitoring
392 of producers of sachet water should be enforced to ensure adherence to drinking water
393 standard.

394

395

396 COMPETING INTERESTS

397

398 Authors have declared that no competing interest exist.

399

400

401

402 REFERENCES

- 403 1. Spellman FR, Drinan J. The Drinking Water Handbook. Lancaster, Pennsylvania,
404 USA: *Technomic Publishing Company Incorporated*. 2000. Pp. 260.
- 405 2. World Health Organization. Drinking Water Quality Guideline 4th Edition. World
406 Health Organization (WHO), Geneva, Switzerland. 2011;1-28.
- 407 3. Linda OA, Uchenna CO, Moses NI, Chinelo KU, Charles, OE. Microbial Evaluation
408 and Antibiotic Susceptibility Profile of Isolates of Popular Sachet Water Brands Sold
409 in Anambra State. *British Microbiology Research Journal*, 2016;12(4), 1-9.
- 410 4. Bitton G. *Wastewater microbiology*. 3rd Edition. Wiley series in ecological and
411 Applied Microbiology. 2005

- 412 5. Barrell R, Hunter PG. Microbiological Standards for Water and their Relationship to
413 Health Risk. *Communicable Diseases and Public Health*, 2000;3(1): 8-13.
- 414 6. Finch RG, Greenwood D, Norrby SR, Whitley RJ. Antibiotic and chemotherapy: Anti-
415 infective agents and their use in therapy. 8th ed. Edinburgh: *Churchill Livingstone*.
416 2003. 964.
- 417 7. Nichol K, Zhanel GG, Hoban DJ. Molecular epidemiology of penicillin resistant and
418 ciprofloxacin resistant *Streptococcus pneumoniae* in Canada. *Antimicrobial Agents*
419 *and Chemotherapy*. 2003;47:804-808.
- 420 8. Kummerer K. Resistance in the environment. *Journal of Antimicrobial Chemotherapy*
421 2004;54:311–320.
- 422 9. Sharma BC, Rai B. Incidence of multi-drug resistance in *Escherichia coli* Strains
423 isolated from three lakes of tourist attraction (Mirik Lake, Jorepokhari Lake and
424 Nakhapani Lake) of Darjeeling Hills, India. *Indian Journal of Fundamental and*
425 *Applied Life Sciences*. 2012; 2(2):108-114.
- 426 10. Swartz MN. Use of antimicrobial agents and drug resistance. *The New England*
427 *Journal*. 1997; 45-68.
- 428 11. Scott ME, Melton-Celsa AR, O'Brien AD. Mutations in hns reduce the adherence of
429 Shiga toxin-producing *E. coli* 091:H21 strain B2F1 to human colonic epithelial cells
430 and increase the production of hemolysin. *Microbial Pathogenesis*, 2003;34:155–
431 159.
- 432 12. Cruickshank R, Duguid JP, Marmion BP, Swain, RHA. Medical Microbiology,
433 Volume II, 12th edition. Churchill Livingstone, Edinburgh, London and New York.
434 1975.
- 435 13. Olutiola PO, Famurewa O, Sonntag HG. *An Introduction to general microbiology: A*
436 *practical approach*: HeidelbergVerlagsansalt und DruckereiGmbH. Heidelberg.
437 1991; P. 267
- 438 14. Harrigan WF, McCance EM. *Laboratory Methods in Food and Dairy Microbiology*.
439 Academic Press, London, New York, San Francisco, 1976; 452 pp.
- 440 15. Buchanan RE, Gibbon NE. *Bergey's Manual of Determinative Bacteriology*. 9th Edition,
441 Williams and Wilkins Co., Baltimore. 1974
- 442 16. National Committee for Clinical Laboratory Standards. Performance standards for
443 antimicrobial disk susceptibility tests. Approved standard M2-A4. Wayne, Pa:
444 National Committee for Clinical Laboratory Standards; 1990.
- 445 17. CLSI 2009. Performance standards for antimicrobial susceptibility testing; 19th
446 informational supplement M100–S19. Clinical and Laboratory Standards Institute,
447 Wayne, PANational Committee for Clinical Laboratory Standards. Performance
448 standards for antimicrobial disk susceptibility tests. Approved standard.
- 449 18. NCCLS document M2-A5. Wayne, Pa: National Committee for Clinical Laboratory
450 Standards; 1993
- 451 19. WHO 2003
- 452 20. Berger, P.S., Oshiro, R.K. Source water protection: Microbiology of source water.,
453 *In: Encyclopedia of Environmental Microbiology*. G.Bitton, Editor-in-chief, Wiley
454 InterScience, New York. 2002; Pp2967-2978
- 455 21. Ollos PJ, Huck PM, Slawson, RM. Factors Affecting Biofilm Accumulation in Model
456 Distribution Systems. *Journal of American Water Works Association*, 2003; 95: 87–
457 97.
- 458 22. LeChevallier MW, Seidler RJ, Evans TM. Enumeration and characterization
459 of standard plate count bacteria in chlorinated and raw water supplies. *Applied*
460 *Environmental Microbiology* 1980; 40:922–930.
- 461 23. Onifade AK, Ilori RM. Microbiological Analysis of Sachet Water Vended in Ondo
462 State, Nigeria. *Environmental Research Journal*, 2008; 2: 107-110.
- 463 24. Oladipo IC, Onyenika IC, Adebisi AO. Microbial analysis of some vended sachet
464 water in Ogbomosho, Nigeria. *African Journal of Food Science*, 2009; 3(12): 406-412.

- 465 25. Oyedeji O, Olutiola PO, Moninuola MA. Microbiological quality of packaged drinking
466 water brands marketed in Ibadan metropolis and Ile-Ife city in South Western
467 Nigeria. *African Journal of Microbiology*. 2010; 4: 96-102.
- 468 26. Onilude AA, Adesina FC, Oluboyede OA and Adeyemi BI. Microbiological
469 quality of sachet packaged water vended in three local governments of Oyo State,
470 Nigeria. *African Journal of Environmental Microbiology*. 2013; 4(9): 195-200.
- 471 27. Nwachukwu E, Emeruem CM. Presence of antibiotic resistant bacteria in sachet
472 water produced and sold in the eastern Nigeria. *Research Journal of Microbiology*,
473 2007; 2(10):782-786
- 474 28. Piddock, L. J. Clinically relevant chromosomally encoded multidrug resistance
475 efflux pumps in bacteria. *Clinical Microbiology Review*, 2006; **19**, 382–402.
- 476 29. Tortora JG, Funke RB, Case LC. Microbiology An introduction. *Media update of 7*
477 *Edition including bibliography and index publisher*. Daryl Fox. 2002; 258-260.
- 478 30. William C, Sonzogoni P, Standridge J, Bussen M. Madison Preservation and
479 survival of *Escherichia coli* in well water sample. *Wisconsin State Laboratory of*
480 *Hygiene, University of Wisconsin*. 2002; 4-10.
- 481
- 482