

1 Isolation, Identification, and Antibigram of Colistin- 2 Resistant *Acinetobacter baumannii* from Rivers in and 3 around Kathmandu Valley

4 Abstract

5 Background: *Acinetobacter baumannii*, an opportunistic Gram-negative pathogen, poses an
6 escalating threat in clinical settings due to the rise of multidrug-resistant infections. Despite its
7 clinical significance, there exists a considerable gap in understanding its environmental
8 dissemination.

9 Aims and Objectives: The primary objective is to examine the distribution of *A. baumannii* and its
10 antibiotic resistance in river ecosystems. Specifically, we aim to identify strains resistant to
11 Colistin, a last-resort antibiotic, and elucidate the susceptibility patterns to other antibiotics.

12 Materials and Methods: Water samples from 10 rivers were collected and subjected to analysis
13 using Leeds *Acinetobacter* Agar Base and a series of biochemical tests. Antibiotic susceptibility
14 testing, focusing on Colistin resistance, was performed using standard procedures.

15 Results: Out of the 284 isolated strains, 14 (4.9%) exhibited resistance to Colistin, while
16 demonstrating varying susceptibility patterns to other antibiotics. Notably, Gentamycin showed
17 effectiveness against resistant strains (14.28%), while Ceftazidime resistance was complete.
18 Colistin-sensitive strains displayed high susceptibility to Ciprofloxacin (84.44%) and lower
19 susceptibility to Chloramphenicol (53.33%). Carbapenem susceptibility was observed across all
20 isolates.

21 Conclusion: The study underscores a concerning environmental presence of multidrug-resistant *A.*
22 *baumannii* in rivers around Kathmandu Valley, with Sundarijal being the exception. The findings
23 emphasize the necessity of scrutinizing environmental reservoirs for pathogen spread, advocating
24 for heightened awareness of potential health implications beyond clinical settings. Urgent attention
25 is needed to comprehend and counteract the emergence and dissemination of antibiotic resistance,
26 necessitating comprehensive strategies and continued surveillance

28 Keywords: *Acinetobacter baumannii*, antibiotic resistance, Colistin, ESKAPE, Carbapenem,
29 Kathmandu Valley.

30

31 Introduction

32 Antimicrobial resistance presents a global health challenge, with rising bacterial infections defying
33 conventional antibiotics (1). Environmental monitoring reveals the pervasive presence of
34 pharmaceuticals in soil and water, raising concerns about the emergence of antibiotic-resistant
35 bacterial strains over time (2). While antibiotic resistance in clinical settings is acknowledged, the
36 environmental impact has been largely overlooked. The use of antibiotics and the spread of
37 antibiotic resistance in clinical settings is a well-recognized problem, but antibiotics and
38 antibiotic resistance as environmental problems and pollutants have largely been overlooked. As
39 a result, the increasing incidence of resistance to a wide range of antibiotic agents by a variety of
40 organisms outside clinical settings is a major concern (3).

41

42 Similarly, in the case of *A. baumannii* too, hospital-acquired infections have increased dramatically
43 worldwide (4). Difficulties caused by this pathogen in the hospital setting are exacerbated by its
44 ability to form biofilms on abiotic or biotic surfaces and to cope with different environmental
45 conditions, including desiccation and disinfectants (5). Non-susceptibility to commonly used
46 antimicrobials has also been observed, with Carbapenem resistance becoming a global problem
47 since 2000 (4).

48

49 *A. baumannii*, recognized as one of the six 'superbugs' by the Infectious Diseases Society of
50 America (IDSA) (6), is a Gram-negative, non-fermentative coccobacillus within the
51 Moraxellaceae family. Gaining prominence as a nosocomial pathogen in recent years (7), *A.*
52 *baumannii*'s adaptability to antibacterial agents in its environment is facilitated by mutation,
53 selection, and genetic exchange mechanisms (2). Studies have demonstrated that wastewater is one
54 of the highly selective environments and that it contributes to the high rates of resistant bacteria
55 that are being discharged in the natural environment, including the river systems (8).

56

57 Untreated hospital waste containing residual antibiotics at low concentrations emerges as a
58 significant contributor to antibiotic resistance development. Such locales become pivotal hotspots
59 for horizontal gene transfer (HGT) of antibiotic-resistant genes, fostering the emergence of
60 resistant pathogens (9). This intricate interplay underscores the urgent need for comprehensive
61 strategies to mitigate the environmental spread of antibiotic resistance associated with *A.*
62 *baumannii* (6).

63
64 The basic principle underlying wastewater management is the stringent control of discharging
65 hazardous liquids into sewers without prior treatment, ensuring the prevention of introducing
66 pathogenic organisms into the environment. However, in Nepal, only Kathmandu Valley boasts a
67 sewerage network, accessible to a mere 15% of households (10). Therefore, even if the hospitals
68 discharge their healthcare liquid waste into the sewerage system, it is mixed with the sewage and
69 gets into surface water without proper treatment (11).

70
71 Untreated hospital effluents pose a significant risk, releasing concentrated forms of infectious
72 agents and antibiotic-resistant microbes into communities, leading to waterborne diseases such as
73 cholera, typhoid fever, dysentery, and gastroenteritis. Environmental compartments, including
74 wastewater, surface water, groundwater, sediments, and soils, have revealed the presence of
75 antibiotics, disinfectants, and bacteria resistant to them (8). Prolonged exposure of microorganisms
76 to low concentrations of antibiotics in wastewater and surface water enhances the potential for the
77 development of antibiotic resistance in these organisms (11). Addressing this issue is imperative
78 for safeguarding public health and environmental well-being.

79
80 The recent development of antibiotic resistance clearly demonstrates the urgent need for global
81 surveillance data that can inform clinicians, public health experts, policymakers and
82 pharmaceutical companies about the dynamic spread of antibiotic-resistant pathogens in a
83 geographically explicit and timely manner (12). However, the antibiotic susceptibility pattern of
84 bacterial isolates in much of the developing world is unknown. Susceptibility testing cannot be
85 done readily because equipment, personnel, and consumables are scarce and expensive (13).
86 Therefore, the purpose of the present study was to examine the presence of MDR *A. baumannii* in
87 river water across Kathmandu Valley.

88

89 The escalating challenge of antibiotic resistance underscores the critical need for global
90 surveillance data, essential for guiding clinicians, public health experts, policymakers, and
91 pharmaceutical companies in understanding the dynamic spread of antibiotic-resistant pathogens
92 (14). Unfortunately, the antibiotic susceptibility pattern of bacterial isolates in much of the
93 developing world is unknown (13). Considering these challenges, the present study aims to
94 investigate the prevalence of multidrug-resistant *A. baumannii* in river water across the Kathmandu
95 Valley. This research contributes to addressing the information gap in antibiotic resistance patterns
96 in resource-constrained settings.

97 Methods

98 Sampling site

99 Ten different sites across Kathmandu Valley (A to J) were selected to carry out this study, namely:

100 Site A: Chobar (Bagmati)

101 Site B: Pashupati (Bagmati)

102 Site C: Balkhu (Balkhu)

103 Site D: Jadibuti (Hanumante)

104 Site E: Tilganga (Bagmati)

105 Site F: Teku (Bishnumati)

106 Site G: Balkumari (Manohara)

107 Site H: Chunikhel (Dhobikhola)

108 Site I: Sundarijal (Bagmati)

109 Site J: Tripureshor (Tukucha)

110

111 These sites were chosen for their geographical significance. Site I is located upstream and offers a
112 baseline for natural water quality before urban influence. Sites C, D, F, G and J are near hospital
113 effluents. Sites A, B, E, and H are influenced by urban runoff.

114 Study period

115 The study was carried out in the Microbiology Laboratory of St. Xavier's College,

116 Maitighar, Kathmandu from February 2021 to December 2021.

117 Sample size

118 A total number of 30 water samples were collected from 10 different sites from different rivers
119 around Kathmandu Valley using the grab sample method. The grab sample method involves taking
120 water samples at a specific point in time (7:00 am), providing a representative sample of the water's
121 composition at that moment. This approach is particularly useful when studying short-term
122 variations and assessing the immediate impact of environmental factors on water quality. Each
123 sample was collected weekly over a period of five months from March 2nd to August 16th, 2021,
124 allowing for a comprehensive examination of temporal variations in water quality.

125 Sample collection

126 River water samples for bacteriological analysis were collected in sterile BOD bottles. During
127 sample collection, the bottle caps were opened aseptically, and the bottles were lowered into the
128 water with their mouth directed against the water current. The water samples were transported in
129 an ice box to the laboratory and processed within two hours of sample collection.

130 *A. baumannii* isolation and identification

131 *A. baumannii* was isolated through the spread plate technique. 0.1 ml of sample was pipetted out
132 from 10⁻³ dilution series onto the centre of the surface of prepared Acinetobacter Agar Base (with
133 added Leeds Acinetobacter Selective Supplement) from HiMedia Laboratories Pvt. Ltd, Mumbai,
134 India. An L-shaped glass spreader was dipped into alcohol. The glass spreader was flamed over a
135 Bunsen burner. The sample was spread evenly over the surface of the agar using the sterile glass
136 spreader, carefully rotating the Petri plate underneath at the same time. The plate was incubated at
137 42°C and growth of the organism was observed after 24 hours.

138

139 Microscopic examination was done after the incubation period and the isolated *A. baumannii* were
140 subcultured on MacConkey agar (MA) and Nutrient agar (NA) and incubated at 42°C for 24 hours.
141 Further biochemical tests (Oxidase, Catalase, Methyl red, Voges-Proskauer test, Citrate utilization
142 test, Indole production test, Triple sugar ion agar test, and Urease test) were performed on all

143 isolates. All the media were) from HiMedia Laboratories Pvt. Ltd, Mumbai, India. The obtained
144 data were interpreted following the Clinical Laboratory Standards Institute (CLSI) (15).

145 Screening for colistin-resistant *A. baumannii*

146 Colistin (4 mg/ml) infused Nutrient agar plates were prepared and a well diffusion method was
147 used to screen colistin-resistant isolates of *A. baumannii*. 0.1 ml of bacterial inoculum prepared by
148 suspending a single colony from overnight agar plates in nutrient broth to the final turbidity of a
149 0.5 McFarland standard was added to the well. After diffusion, the plates were incubated at 37°C
150 for 24 hours, and growth was observed.

151 Antibiogram of isolates

152 Colistin-resistant *A. baumannii* were further tested against a series of antibiotics to study their drug
153 resistance profile. Kirby-Bauer's disc diffusion method was used for the antibiotic susceptibility
154 test.

155 Detection of carbapenemase production

156 The modified carbapenem inactivation method was used for phenotypic detection of
157 carbapenemase production in the colistin-resistant *A. baumannii* isolated. The isolates were
158 subcultured on Blood Agar (from HiMedia Laboratories Pvt. Ltd, Mumbai, India) and incubated
159 at 42°C for 24 hours. A single colony from each isolate was emulsified in 2 ml Tryptic Soy Broth
160 (TSB) (from HiMedia Laboratories Pvt. Ltd, Mumbai, India). Meropenem disk was immersed in
161 the suspension and was incubated at 37°C for four hours. A 0.1 ml of *E. coli* ATCC 25922 was
162 then prepared by suspending a single colony from overnight agar plates in nutrient broth to the
163 final turbidity of a 0.5 McFarland standard and was carpet cultured on Mueller Hinton Agar
164 (MHA) plates. Meropenem disks were removed from TSB and placed on the plates. Another fresh
165 meropenem disk was placed as control. The MHA plates were incubated at 37°C for 24 hours and
166 observed for a zone of inhibition.

167 Preservation of *A. baumannii*

168 *A. baumannii* isolates in pure culture, after performing the biochemical and antimicrobial
169 susceptibility testing, were preserved in Tryptic Soy broth (TSB) containing 20% Glycerol and
170 kept at -70°C until further tests were required.

171 Quality control

172 Laboratory equipment like incubators, refrigerators, autoclaves, and hot-air ovens were regularly
173 monitored for their efficiency. The temperature of the incubator and refrigerator were monitored
174 twice a day. Reagents and media were regularly monitored for their expiry date and proper storage
175 condition. After media preparation, they were properly labelled with the preparation date. The
176 quality of the media prepared was checked by subjecting one plate of each batch to sterility and
177 performance testing.

178

179 A purity plate was used to ensure that the inoculation used for biochemical tests was pure culture
180 and to ensure that the biochemical tests were performed in an aseptic condition. Thus, while
181 performing biochemical tests, the same inoculums were subcultured in the respective medium and
182 incubated. The media were then checked for the appearance of pure growth of organisms. The
183 development of pure culture in the medium would confirm the purity of the inoculum.

184

185 Antibiotic susceptibility tests were performed by maintaining the thickness of Mueller Hinton agar
186 at 4mm and pH at 7.2-7.4. Similarly, antibiotic discs containing the correct amount as indicated
187 were used. MHA and the antibiotic discs were checked for their lot numbers, manufacturing dates,
188 expiry dates, and storage conditions. For the standardization of the Kirby Bauer test and for
189 performance testing of antibiotics and MHA, the control strain of *E. coli* ATCC 25922 was tested.

190 Statistical analysis

191 The data obtained were entered into MS Excel and analyzed using Statistical Package for Social
192 Science (SPSS) software (Version 21.0).

193

194 **Result**

195 **Culture result of the samples**

196 The samples were cultured on Leeds Acinetobacter Agar Base at 42°C. Out of the 30 samples, 27
197 samples showed growth for *A. baumannii*, whereas three samples from Site I showed no growth
198 on the media.

199 **pH**

200 The pH was noted every time the samples were collected. Each sample's maximum and minimum
201 pH are given below, which was found to be within the range of WHO guidelines and was between
202 6.1 to 8.4. There were no huge differences in pH among the 10 sites.

Site	pH	
	Maximum	Minimum
A	7.1	6.8
B	7.0	6.7
C	6.9	6.8
D	7.1	6.8
E	6.5	6.4
F	8.1	7.8
G	7.1	7.0
H	7.4	7.2
I	7.4	7.2
J	6.6	6.4

205

206

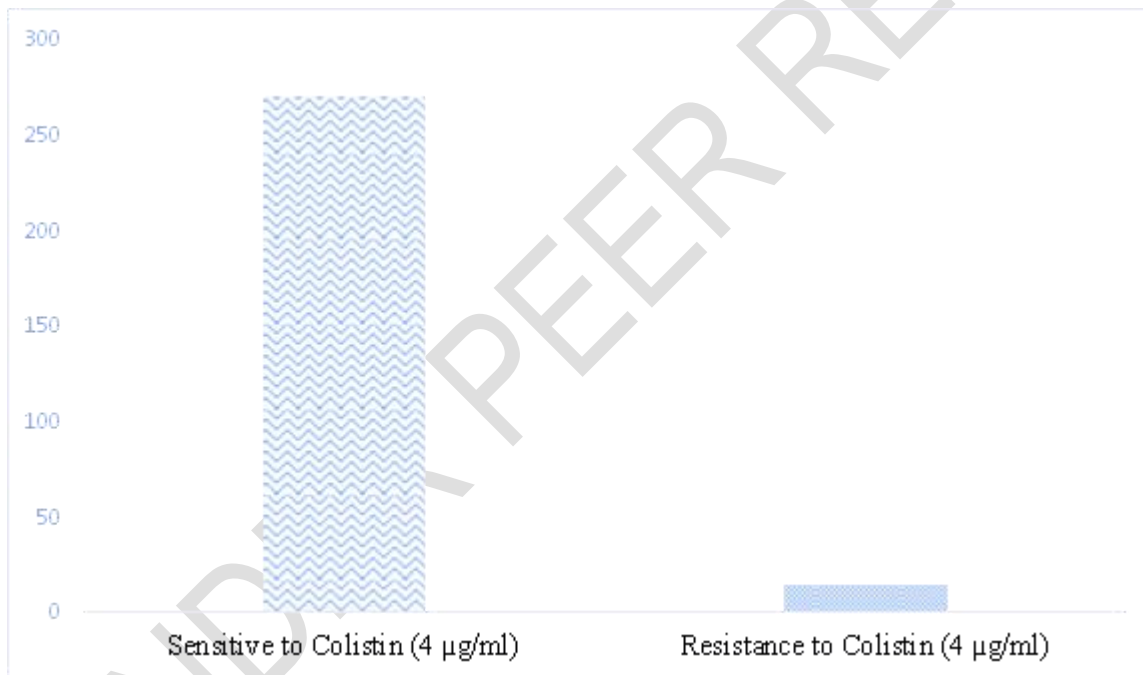
Table 1: pH of the river water samples

207 Morphological and biochemical characteristics of *A. baumannii*

208 From the 27 samples, 284 isolates of *A. baumannii* were isolated using the Acinetobacter agar
209 base. All the isolates were subjected to Gram staining, followed by biochemical tests. The isolates
210 were Gram-negative, non-motile coccobacillus which showed growth on Simmon's citrate and
211 MacConkey agar at 37 °C.

212 Screening of colistin-resistant *A. baumannii*

213 The susceptibility breakpoint of colistin is 4 µg/ml for *A. baumannii* 17978 (CLSI 2022). All 284
214 isolates of *A. baumannii* were grown on MHA with 4 µg/ml colistin. Out of them, 14 (4.9%)
215 isolates showed growth on the media and 270 (95.1%) showed negative growth. The isolates that
216 grew on MHA incorporated with 4 µg/ml were screened as colistin-resistant.



217

218

Figure 1: Colistin-resistance in *A. baumannii* isolates

219 Antibiotic susceptibility pattern of colistin-resistant *A. baumannii*

220 The 14 colistin-resistant *A. baumannii* were further subjected to different types of antibiotics from
221 different groups, namely chloramphenicol, gentamycin, ceftazidime, nalidixic acid, ciprofloxacin,
222 cefotaxime, by Kirby-Bauer disk diffusion method following CLSI guidelines. The highest

223 susceptibility was found towards gentamycin (14.28%) and the least susceptible towards
 224 ceftazidime (0.00%).

Antibiotics	Disc content (µg)	Sensitive	Intermediate	Resistance	Total isolates
Cefotaxime (CTX)	30	1 (7.14%)	0 (0.00%)	13 (92.86%)	14
Ceftazidime (CAZ)	30	0 (0.00%)	0 (0.00%)	100 (100.00%)	14
Chloramphenicol (C)	30	1 (7.14%)	0 (0.00%)	13 (92.86%)	14
Ciprofloxacin (CIP)	5	1 (7.14%)	0 (0.00%)	13 (92.86%)	14
Gentamycin (GEN)	10	2 (14.28%)	1 (7.14%)	11 (78.57%)	14
Nalidixic Acid (NA)	30	1 (7.14%)	NA	13 (92.86%)	14

225

226 Table 2: Antibiotic susceptibility pattern of in *A. baumannii* isolates

227 Screening of carbapenem-resistant *A. baumannii* isolates

228 The 14 colistin-resistant *A. baumannii* were initially subjected to meropenem by Kirby-Bauer disk
 229 diffusion methods following CLSI guidelines to screen out meropenem-resistant isolates i.e. those
 230 isolates whose zone of inhibition were less than 20 mm. Four of the isolates were resistant to
 231 Meropenem.

Antibiotics	Total	Sensitive	Resistance
	isolates		
Meropenem (MRP)	14	10 (71.43%)	4 (28.57%)

233

234

Table 3: Screening of Carbapenem resistance in Colistin-resistant *A. baumannii* isolates235 **Confirmation of carbapenem-resistant *A. baumannii* isolates**

236 Four isolates that were screened as carbapenem-resistant were further subjected to MCIM
 237 (Modified Carbapenem Inactivation Method--procedure) for confirmation of carbapenem
 238 resistance. All four of them were sensitive towards meropenem after the MCIM test.

239 **Antibiotic susceptibility pattern of colistin-sensitive *A. baumannii***

240 The 270 remaining isolates that did not show resistance to colistin were further subjected to
 241 different types of antibiotics from different groups, namely chloramphenicol, gentamycin,
 242 nalidixic acid, cefotaxime, ciprofloxacin and ceftazidime by Kirby-Bauer disk diffusion method
 243 following CLSI guidelines. The highest susceptibility was found towards ciprofloxacin (84.44%)
 244 and the least susceptible towards chloramphenicol (53.33%).

245

Antibiotics	Disc content (µg)	Sensitive	Intermediate	Resistance	Total isolates
Cefotaxime (CTX)	30	174 (64.44%)	42 (15.56%)	54 (20.00%)	270
Ceftazidime (CAZ)	30	210 (77.78%)	48 (17.79%)	12 (4.42%)	270

Chloramphenicol	30	144	54	72	270
(C)		(53.33%)	(20.00%)	(26.67%)	
Ciprofloxacin	5	228	24	18	270
(CIP)		(84.44%)	(8.89%)	(6.67%)	
Gentamycin	10	225	24	21	270
(GEN)		(83.31%)	(8.89%)	(7.80%)	
Nalidixic Acid	30	210	NA	60	270
(NA)		(77.78%)		(22.22%)	

246

247

Table 4: Antibiotic susceptibility pattern of Colistin-sensitive *A. baumannii* isolates

248 Screening of carbapenem resistance among the *A. baumannii* colistin-sensitive
249 isolates

250 The 270 colistin-sensitive *A. baumannii* were initially subjected to Meropenem by the Kirby-Bauer
251 disk diffusion method following CLSI guidelines to screen out those isolates whose zone of
252 inhibition was less than 20 mm. All the isolates were sensitive to Meropenem.

253 The multidrug resistance profile of *A. baumannii* isolates

254 Multidrug-resistant *A. baumannii* is defined as resistant to at least three classes of antimicrobial
255 agents—penicillins, cephalosporins, fluoroquinolones, and aminoglycosides. Of the 284 *A.*
256 *baumannii* isolates, 158 (57.04%) were found to be multidrug resistant. 73 were resistant to
257 penicillins + quinolones + cephalosporins, 53 were resistant to penicillins + quinolones +
258 aminoglycosides, 32 were resistant to penicillins + quinolones + aminoglycosides +
259 cephalosporins, and none were resistant to penicillins + quinolones + aminoglycosides +
260 cephalosporins + carbapenem.

261

262 Relation between MDR and colistin-resistant isolates

263 All 14 initially screened colistin-resistant *A. baumannii* isolates were found to be multidrug-
264 resistant (MDR) as well. The chi-square test for independence was conducted to examine the
265 association between multidrug resistance (MDR) and colistin resistance in the *A. baumannii*
266 isolates. The contingency table presented a chi-square statistic of $\chi^2=90.03$ with 1 degree of
267 freedom. The test revealed a statistically significant association between MDR and colistin
268 resistance ($\chi^2(1) = 90.03, p<0.05$). The observed distribution significantly deviated from what
269 would be expected under the assumption of independence. This suggests that the likelihood of
270 being colistin resistant is associated with the presence of multidrug resistance in the studied *A.*
271 *baumannii* isolates.

272

MDR	Susceptibility to Colistin		Total	p-value
	Colistin resistant	Colistin sensitive		
Positive	14	144	158	
Negative	0	126	126	<0.05
Total	14	270	284	

273

274

Table 5: Relation between MDR and colistin-resistant *A. baumannii* isolates

275

276 Discussion

277 *A. baumannii* in hospital settings is not a new finding; however, the propagation of viable *A.*
278 *baumannii* in the natural environment is a pressing public health challenge. From soil to water, the
279 antibiotic-resistant bacteria may also transmit and colonize a new home/habitat elsewhere.

280

281 We undertook this research due to the growing evidence of multidrug-resistant bacteria in river
282 water, as demonstrated by numerous studies, including those conducted in South Asian countries.
283 For instance, Lamba et al., in 2017, investigated the release of carbapenem-resistant pathogens in
284 Delhi, India (16). Additionally, a parallel study in Kathmandu by Thakali et al. (2021) focused on
285 the release of antibiotic-resistance genes from hospitals (17). The existing body of research
286 highlighted the urgent need to further explore and understand the dynamics of antibiotic resistance
287 in river water, particularly in the context of the Kathmandu Valley.

288
289 In this study, river water samples (pH 6.4-8.1) yielded 284 *A. baumannii* isolates as they showed
290 positive growth at 42°C on Acinetobacter Agar Base (with added Leeds Acinetobacter Selective
291 Supplement). Confirmatory tests identified isolates as Gram-negative, non-motile coccobacilli,
292 with specific growth characteristics. Because the study focused on isolating colistin-resistant *A.*
293 *baumannii* and further investigating their antimicrobial patterns, all 284 isolates were subjected to
294 4 µg/ml Colistin. Out of them, 14 isolates showed positive growth against it. Jovicic et al, in March
295 2021, observed a similar result in Zagreb, where clinically relevant isolates of *A. baumannii* were
296 recovered from hospital wastewater and wastewater treatment plants, seven of which were found
297 to be colistin-resistant (18). The 14 colistin-resistant isolates of *A. baumannii* also showed
298 similarity to the clinical isolate with which it was compared in the pattern of susceptibility to
299 different antibiotics, namely chloramphenicol, gentamicin, ceftazidime, nalidixic acid,
300 ciprofloxacin, cefotaxime, ceftazidime/clavulanic acid, and cefotaxime/clavulanic acid. The
301 highest susceptibility was found towards gentamycin (14.28%). The βlactam antibiotic,
302 cefotaxime, used in combination with clavulanic acid, showed the least sensitivity (0.00%) towards
303 all 14 isolates.

304
305 According to a 2012 study done by Espinal et al, the β-lactam antibiotic is the recommended
306 antibiotic treatment for Acinetobacter infections (5). However, it did not show promising results
307 against the isolated *A. baumannii* in this study. This is a point to be noted that within a decade, the
308 bacteria have evolved to show resistance towards even the most recommended antibiotics as well.
309 Also, a combination of antibiotic-resistant mechanisms could be one of the likely reasons for a
310 wide range of resistance to different classes of antibiotics by *A. baumannii*. The 14 isolates were
311 subjected to meropenem too. First, the screening process was carried out where the isolates were

312 tested against meropenem by the Kirby-Bauer disk diffusion method following CLSI guidelines
313 to screen out those isolates whose zone of inhibition was less than 20 mm. From this process, four
314 isolates were initially screened out. These four isolates were further subjected to MCIM (Modified
315 Carbapenem Inactivation Method) out of which all were susceptible to Carbapenem (meropenem),
316 which is not 42 unexpected as the prevalence of carbapenem-resistant *A. baumannii* is highly
317 unlikely in environment settings (19).

318
319 The remaining 270 isolates that were susceptible to colistin were further tested against different
320 classes of antibiotics to understand their antibiograms. The highest susceptibility was found
321 towards ciprofloxacin (84.44%) and least susceptible towards chloramphenicol (53.33%). After
322 this, the same 270 Colistin-sensitive *A. baumannii* were initially subjected to meropenem by the
323 Kirby-Bauer disk diffusion method following CLSI guidelines to screen out those isolates whose
324 zone of inhibition was less than 20 mm. All 270 of the isolates showed susceptibility towards
325 meropenem, so further MCIM test was not carried out. The colistin-resistant and susceptible
326 isolates were further studied to understand their multidrug-resistant patterns.

327
328 Multidrug-resistant *A. baumannii* are those that are resistant to at least three classes of
329 antimicrobial agents—penicillins, cephalosporins, fluoroquinolones, and aminoglycosides. Of the
330 284 *A. baumannii* isolates, 158 (57.04%) were multidrug resistant. 73 isolates were resistant to
331 penicillins, quinolones and cephalosporins, 53 were resistant to penicillins, quinolones and
332 aminoglycosides, 32 were resistant to penicillins, quinolones, aminoglycosides and
333 cephalosporins. All 14 colistin-resistant *A. baumannii* that were initially screened were also found
334 to be multidrug-resistant and thus were classified as extensively drug-resistant (XDR). A similar
335 study was done at a university hospital in Kathmandu, where 122 (49.6%) *A. baumannii* isolates
336 out of 246 were MDR *A. baumannii*, with the majority being resistant to fluoroquinolones,
337 aminoglycosides and carbapenems. However, they were not resistant to colistin (20). In a similar
338 study in China, nine MDR *A. baumannii* were recovered from wastewater followed by disinfection
339 (chlorination), afterwards out of 9 only one remained MDR but those strains were not related to
340 clinical isolates (21).

341

342 Another study conducted in Croatia also recovered viable MDR *A. baumannii* which was also not
343 related to clinical isolates from municipal wastewater in Zagreb, Croatia, both before and after
344 passage through the secondary wastewater treatment process (22). However, despite the above
345 observations concerning the ubiquity of *A. baumannii*, there is a lack of clear evidence about the
346 relationship between hospital settings and the natural environment in the propagation of this
347 increasingly important pathogen (9). In this study, *A. baumannii* showed excellent survival in river
348 water. These isolates disseminated in the environment could represent the source of serious
349 community-acquired infections. The fact that *A. baumannii* is responsible for uncountable
350 hospital-acquired infections worldwide and has recently become one of the top most important
351 healthcare-associated infections in hospitals is a matter of discussion, but their presence in the
352 environment, that too in river water indicates the occurrence of horizontal gene transfer (HGT).

353

354 In the rivers in Kathmandu, especially in central city areas where wastewater is directly dumped
355 into the rivers, human-associated and environmental bacteria are mixed together and exposed to
356 many substances, including various antimicrobial compounds, which in turn increase the HGT
357 (17). This can also be backed by the fact that no *A. baumannii* was isolated from the water collected
358 from Sundarijal, which is considered the starting point of the Bagmati River. The river's
359 downstream and midstream areas, however, flow through settlement areas where hospitals are
360 present. Not only that, the only functioning wastewater treatment plant in Kathmandu Valley also
361 is in the midstream site. With more than a hundred hospitals and a treatment plant that lacks a
362 chlorine disinfection phase, which is a critical step in removing antimicrobial-resistant genes in
363 bacteria, it is inevitable for multidrug-resistant bacteria, even those found mainly in the hospital
364 environment, to be present in river water (23). However, a study conducted in Spain by Rodriguez-
365 Mozaz et al in 2015 showed that even after hospital wastewater treatment in proper wastewater
366 treatment plants, antimicrobial-resistant genes still spread in the receiving river sources (14). This
367 further backs the claim that hospitals and even wastewater treatment plants are potential sources
368 of multidrug-resistant *A. baumannii* in the rivers in Kathmandu Valley.

369

370 The findings of this study highlight the presence of colistin-resistant *A. baumannii* in the
371 environmental reservoirs of Kathmandu Valley, underscoring a critical aspect of Nepal's
372 environmental and public health landscape. This discovery fills a notable gap in the current

373 understanding of antibiotic resistance dynamics in the region and emphasizes the need for a holistic
374 approach to combatting the spread of multidrug-resistant pathogens. The study also highlights the
375 potential risks associated with the environmental presence of colistin-resistant *A. baumannii*,
376 particularly in a country with limited healthcare resources. The elevated prevalence of antibiotic
377 resistance in environmental water sources raises concerns about its transmission to humans and
378 animals, posing significant challenges for public health.

379
380 To address these concerns, we propose a series of measures for monitoring, control, and
381 prevention. Enhanced surveillance of water sources, coupled with the development of
382 antimicrobial stewardship programs, can play a pivotal role in curbing the dissemination of
383 resistant strains. Moreover, public awareness campaigns and educational initiatives are crucial in
384 fostering a proactive community response to mitigate the risks associated with environmental
385 antibiotic resistance.

386 Conclusion

387 The findings from this study strengthen the urgent need not just to develop new classes of
388 antibiotics against highly resistant bacteria like *A. baumannii*, but also to strictly monitor the
389 haphazard use of antibiotics.

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