

REVIEW OF ANTIBIOTICS USAGE IN THE MANAGEMENT OF OPEN

FRACTURE IN A NIGERIAN HOSPITAL

ABSTRACT

BACKGROUND. Antibiotics are adjuncts in the management of open fractures, and microbial characteristic of open fractures will guide the use of antibiotics. With changing pattern in microbial colonization of wounds, the need to review antibiotic usage in hospitals becomes imperative. The study aimed to evaluate the antibiotic protocol of managing open fractures at the Accident and Emergency department, with the advent of new antibiotics introduced into the hospital drug formulary.

MATERIALS AND METHODS. This study is a hospital-based prospective evaluation of the antibiotic sensitivity of cultured microorganisms from the patients with open fractures presenting between January 2013 and December 2013 in the Accident and Emergency Department, of a tertiary hospital in Nigeria. Swabs of superficial and deeper parts of the wound were taken at the presentation of the patients before wound debridement and commencement of antibiotics. Other two samples and biopsies were taken at the deeper parts of the wound on the 3rd and 7th day of admission. Culture and Sensitivity pattern of isolates were determined for positive cultures using antibiotics impregnated disks. Descriptive and inferential statistics of the findings are presented.

RESULT. One hundred and thirty patients were recruited for the study, a sterile swab was taken from their wounds at presentation, but 81 patients completed the study. Forty patients discharged themselves against medical advice and while nine patients were referred to other hospitals. *Staphylococcus aureus* and *Clostridium perfringens* was the most common aerobic and anaerobic isolates respectively. The aerobic isolates and anaerobes were susceptible to ceftriaxone, ciprofloxacin, co-Amoxyclav, gentamycin, and cefotaxime and metronidazole respectively.

CONCLUSION. The antibiotic sensitivity pattern in the emergency department of the Hospital has changed not significantly as previously reported about 12 years earlier. Therefore, the hospital antibiotic protocol in the treatment of open fractures in the Accident and Emergency department should be retained.

Keywords: Open fracture. Antibiotics sensitivity, Antibiotic usage, Ibadan, Nigeria

Introduction

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The choice of antibiotics in the treatment of open fractures as an adjunct to debridement and wound care is determined by established microbial characteristics of open fractures in the locality or empirically using combination therapy to cover most of the available organisms such as Gram-positive and Gram-negative aerobes as well as the anaerobes. The trend of microbial infections and antibiotic sensitivity pattern in the hospital where this particular study was undertaken had been established by a previous study [1] The choice of antibiotics in the treatment of infections is determined by the potential bacterial contamination based on historical or research documented patterns for each locality [2]. On account of their findings, Wilkins and Patzakakis recommended the use of a combination of cephalosporins, penicillins and aminoglycosides in open fractures depending on the severity of the wound and extent of contamination [3]. However, Alonge et al. in Ibadan Nigeria, found that pefloxacin, ciprofloxacin and ceftriaxone were the antibiotics which exhibited relatively higher sensitivity to the micro-organisms isolated [1], which is in agreement with the findings in other studies [4] [5] [6] [7][8].

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An open fracture can be defined as a break in the structural continuity of a bone in which the fracture hematoma communicates through the soft tissue with epithelial lining including skin and mucosal lining. It is relatively common especially in developing countries and accounts for a third of all trauma referrals [1]. In one study, Forty-eight percent of fractures were open fractures with a preponderance for males and a predilection for the tibia and the forearm bones [9]. Open fractures usually result from high energy trauma such as motor vehicle crashes, falls from height, gunshot injury, assault and machine injury [5] and are prone to contamination and infection[4]. Open fractures have been classified into three major types (I, II, III) and type III has been further sub-classified into three groups, based on the mechanism of injury, the degree of soft tissue damage, the configuration of the fracture and the level of contamination [2] [10]

Decades of research correlating the Gustilo-Anderson types and the risks of infection have helped refine surgical protocols, change in antibiotic prescriptions, and in defining the appropriate timing for interventions including debridement, modalities of fracture fixation, and soft tissue coverage [11][12][13][14][15][16]. Infections in open fractures often develop after six hours of injury if adequate surgical treatment is not carried out along with the administration of appropriate antibiotics early enough after the injury. Deep fracture site infections could lead to complications of chronic osteomyelitis, nonunion and sometimes limb loss. Apart from the exposure of the fractured bone, numerous predisposing factors which influence the development of infection include shock from blood loss, hypoxia and the degree of comminution [17]. Majority of infections in open fractures are caused by *Staphylococci* species especially *Staphylococcus aureus* and coagulase-negative *Staphylococci*, gram-negative bacilli which include *Acinetobacter spp*, *Escherichia coli*, *Pseudomonas spp*, *Klebsiella spp* and *Proteus spp* amongst others[4][14][17]. However, Alonge et al. in 2002 established that *E coli* was the most prevalent single isolate while *Staphylococcus aureus* was the most prevalent microbial isolate in poly-microbial infections [1].

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While antibiotics have been established as an essential adjunct in the treatment of open fractures, resistance to available antimicrobial drugs is an established and ever-growing challenge in clinical practice. Such resistance can result from two mutually non-exclusive phenomena: mutations in house-keeping structural or regulatory genes and the horizontal acquisition of foreign genetic information [18]. Outbreaks of infections due to *Klebsiella* pneumonia harboring plasmid-encoded cephalosporinases and the spread of this resistance mechanism to bacterial species naturally susceptible to cephamycins have been reported [19]. An infection engrafted on a biomaterial (thick, adherent biofilm) responds poorly to antimicrobial therapy and usually is not cured until the biomaterial is removed. Bacterial isolates may not be entirely representative of the microbial components of the biofilm because the coherent properties of the adherent biofilms that are found on surfaces in these infections may prevent genuinely representative organisms from detaching in sufficient numbers to be detected entirely and consistently by simple sampling and routine culture techniques. Therefore, antimicrobials that are chosen from the culture results may not be effective against all of the bacterial species in these biofilm infections [20].

The rapid spread of antimicrobial resistance in a wide variety of bacteria is mainly due to the location of antimicrobial resistance genes on mobile genetic elements such as plasmids and transposons [21]. Globally, *Enterobacter* isolates resistant to expanded-spectrum cephalosporin is becoming a matter of concern for the possibility of transmitting antimicrobial resistance from one microorganism to another [22].

This study aimed to review the antibiotic treatment protocol for open fractures in the A&E of a tertiary hospital in Nigeria with the view for recommendations for possible change in practice.

MATERIALS AND METHODS

This study is a hospital-based prospective evaluation of antimicrobial pattern and antibiotics sensitivity pattern in open fractures presenting in the Accident and Emergency Department of the University College Hospital, Ibadan from January 2013 to December 2013.

Proforma for the study was completed for all patients seen in the Accident and Emergency department of the hospital with open fracture after obtaining securing informed consent from the included patients. Patients with an open fracture who had wound debridement and antibiotics before presenting at the Accident and Emergency of the University College Hospital, Ibadan were excluded.

Poly-traumatized patients with concomitant open fractures were resuscitated and treated using the advanced trauma life support (ATLS) protocol. The associated wounds with open fractures were inspected, and clinical photographs obtained to record the injury at presentation. Four sterile wound swabs, (superficial aerobic and anaerobic, deep aerobic and anaerobic) were collected from the superficial and deep parts of open fracture wounds using the Levine's technique. The swabs of the wounds were obtained aseptically before wound

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debridement and antibiotics were commenced within 30 minutes of patient's arrival at the Accident and Emergency Department. Two other samples and biopsies were taken at the deeper parts of the wounds on the 3rd and 7th day of admission. Samples were collected into sterile Stuarts transport medium, and sterile Robertson cooked meat medium for aerobic and anaerobic organisms respectively. The samples were labelled "S" for superficial swab samples, "D" for deep swab samples, "BS" and "BD" for superficial and deep biopsy samples with the patient's research number on the laboratory request form and also on the bottle. All samples arrived the laboratory within 30 minutes to 3 hours of collection. The samples were stored at room temperature in a cupboard for less than 6 hours until ready for analysis. Microscopy, culture and sensitivity patterns of the samples to various antibiotics (penicillin, cephalosporin, quinolone, aminoglycoside, clindamycin, sulphonamides and trimethoprim, and metronidazole) were carried out. The samples for aerobic cultures were plated out on sterile Sheep blood agar and MacConkey agar aseptically and incubated at 37⁰C for 24 hours. The direct Gram staining of the swabs was carried out, and the slides examined to identify the presence of organisms and pus cells. After 24 hours of incubation, the plates were analyzed for the growth of the bacteria and gram staining of the bacteria colonies were carried out.

The confirmatory test of all the isolated gram-negative bacilli was based on the use of API 20 E while the gram-positive cocci were based on the use of control organisms for coagulase test. Sensitivity testing was carried out using the disc diffusion technique (Bauer Kirby method), where the Mueller Hinton agar was seeded with the confirmed bacteria, and the observed zone of inhibition around the antibiotic discs was measured and compared with the controlled organism. It was recorded as sensitive if the observed area was greater or equal to the zone of the controlled microorganisms and resistant if less than the observed zone of the standard organisms. The anaerobic samples were inoculated aseptically into a sterile Sheep blood agar and MacConkey agar within five minutes of sample collection. The inoculated plates were incubated in the anaerobic gas chamber containing anaerobic catalytic agent, Anaero Gen kit and anaerobic control kit (Oxoid Ltd of United Kingdom). Strict anaerobic control bacteria and strict aerobic bacteria were also included as an added quality control. The anaerobic organisms were left in the chamber to incubate at 37⁰C for three days to isolate the fast-growing anaerobes which are mostly contaminants while the late growing anaerobes were further incubated for ten days and these are the bacteria of medical importance.

RESULTS

Eighty-one of the 130 patients recruited completed the study with superficial and deep swab samples taken from all patients on the first day and the second and third swab and biopsy samples taken on the third and seventh day of admission. Forty patients took their discharges against medical advice while nine patients were referred to other hospitals of their choice. Eight of the open fractures were excluded based on the study exclusion criteria. There were 93 (71.5%) male and 37 (28.5%) female patients as shown in figure 1 while figure 2 represents open fractures in different regions of the body with the tibia and fibula constituting 78 (60%) of the cases while the femur accounted for 19 (14.6%). Gustilo and Anderson type

130 3B [23] was the most common grade of open fracture 48 (36.9%), while type 3A occurred in
131 43 (33.1%) as presented in figure 3. The microbial culture shows that *Staphylococcus aureus*
132 and *Clostridium perfringens* were the predominant aerobic and anaerobic isolates.
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Figure 1: Showing the sex distribution

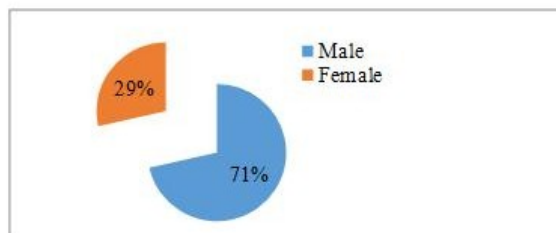
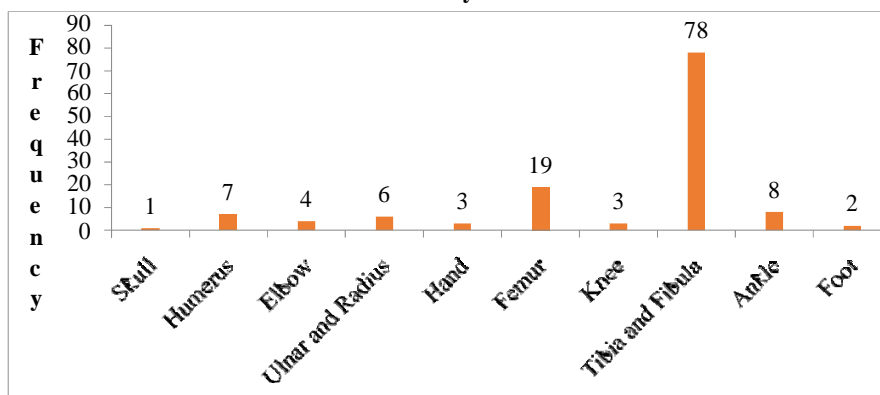
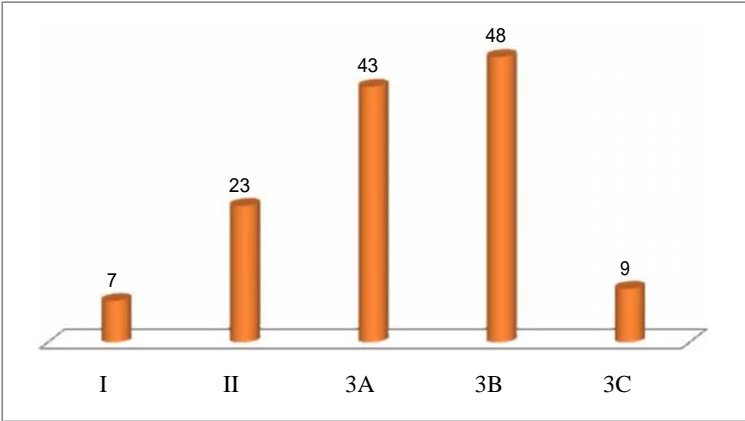


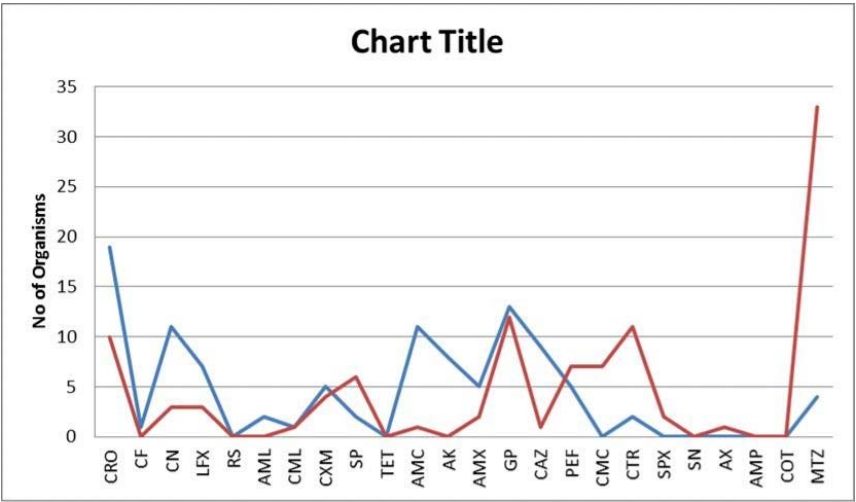
Figure 2: Shows open fracture in the various regions of the body



147 **Figure 3: Shows the grades of open fracture**



150 **Figure 4: Antibiotic sensitivity pattern for aerobes (blue) and anaerobes (red)**



152 **Abbreviations**

153 CRO – cephalosporin, CF – cefazolin, CN – gentamycin, LFX – levofloxacin, RS –rosoxacin,
154 AML –amoxycillin, CLM – clindamycin, CXM – cefuroxime, SP – sparfloxacin, TET –
155 tetracycline, AMC – co-Amoxyclav, AMX – amoxycillin, GP – ciprofloxacin, CAZ –
156 ceftazidime, PEF- pefloxacin, CTR – cefotaxime, SPX – sparfloxacin, SN-sulphonamides,
157 AX – amoxycillin, AMP – ampicillin, MTZ – metronidazole and COT–cotrimoxazole.

159 The antibiotic sensitivity pattern are shown in figure 4 and tables 1 and 2. Ciprofloxacin
160 (GP), cephalosporin (CRO), co-amoxiclav (AMC) and gentamycin (CN) were the drugs most

aerobic organisms were sensitive to, while anaerobic microorganisms were highly sensitive to cefotaxime (CTR), and metronidazole (MTZ).

Table 1. Aerobic Organism sensitivity

Organism	Antibiotics									
.	CRO	CN	LFX	CXM	AMG	AMX	GP	CAZ	CTR	MTZ
SA	5	4	1	2	3	3	4	0	0	0
EC	0	1	2	0	1	0	1	2	0	0
KS	3	1	0	1	5	1	4	0	0	0
PsA	2	1	1	0	0	0	2	1	0	0

Key: *SA* – *Staphylococcus aureus*, *EC* – *Escherichia coli*, *KS* – *klebsiella spp*, and *PsA* - *Pseudomonas auregenosa*

Table 2. Anaerobic Organism sensitivity

Organism	Antibiotics									
	CRO	CN	LFX	CXM	AMG	AMX	GP	CAZ	CTR	MTZ
CP	3	0	2	2	0	0	3	1	3	20
BS	0	0	0	0	0	0	0	0	0	5
CT	1	0	1	0	0	0	1	0	2	9
AI	4	1	0	2	0	0	4	2	1	0

Key: *CP* – *Clostridium perfringens*, *CT* – *Clostridium tetani*, *BS* – *Bacteroides spp* and *AI* – *Actinomyces israelii*.

169 Discussion

170

171 The hospital antibiotic protocol in the Accident and Emergency Department of the hospital,
 172 for the treatment of open fractures, has been a combination of ceftriaxone, quinolones
 173 (ciprofloxacin) and metronidazole-based on findings of Alonge et al. in 2002. The role of
 174 early wound debridement and antibiotic administration is recognized as necessary in the
 175 management of open fractures in the hospital. Appropriate antibiotic(s) are administered
 176 according to the established hospital protocol following the identified historical and
 177 sensitivity pattern of wound swabs [24]. The current hospital antibiotic protocol was guided
 178 by an earlier study that confirmed *Escherichia coli* as the most common single gram-negative
 179 aerobic isolate sensitive to ceftriaxone, quinolones, but since anaerobic organisms were not
 180 cultured the inclusion of metronidazole in the hospital antibiotic protocol was based on
 181 evidence from other practices. The result of the earlier study in the center was at variance to
 182 the findings in this study which showed that *Staphylococcus aureus* and *Clostridium*
 183 *perfringens* as the most common single aerobic and anaerobic isolates respectively. The
 184 predominant aerobic gram-positive organism (*Staphylococcus aureus*) was sensitive to
 185 ceftriaxone (CRO), Gentamycin (CN), co-amoxiclav (AMC), cefuroxime (CXM) and
 186 amoxycillin (AMX) while the aerobic gram-negative organisms (*Escherichia coli* and
 187 *Klebsiella spp*) were sensitive to ceftriaxone, amoxycillin, levofloxacin and ceftazidime. The
 188 antibiotic sensitivity pattern was similar to the findings by Alonge et al. 2002 and other
 189 studies [1][4][5]. Also, anaerobes were significantly sensitive to metronidazole (MTZ) and
 190 moderately sensitive to ceftriaxone, levofloxacin, cefuroxime, ciprofloxacin and cefotaxime
 191 (CTR), affirming the inclusion of metronidazole in the hospital antibiotic protocol. Since the
 192 antibiotic sensitivity pattern from this study is in keeping with findings of an earlier study
 193 which results guided the hospital antibiotic protocol, the hospital antibiotic protocol should
 194 therefore be retained.

195

196 The organisms cultured in this study showed high resistance to ampicillin (AMP),
 197 cotrimoxazole (COT), sulphonamides (SN), clindamycin (CML), rosoxacin (RS),
 198 amoxycillin, cefazolin (CF), and tetracycline (TET). The aerobic gram-positive organisms
 199 were resistance to ceftazidime (CAZ), cefotaxime (CTR) and metronidazole while the aerobic
 200 gram-negative microorganisms were resistance to cefotaxime, metronidazole, amoxycillin,
 201 cefuroxime). The anaerobic organisms also showed significant resistance to co-amoxyclav,
 202 amoxycillin, gentamycin and Ceftazidime. These findings are comparable to a similar study
 203 in another African hospital by Sitali and colleagues in 2017 [25].

204

205 Apart from antibiotic sensitivity and microbial patterns, the hospital antibiotic protocol is also
 206 influenced by the cost and availability of the drugs. In the centre where this study was
 207 undertaken as well as in most hospitals in the region, availability of some of the antibiotics
 208 can be challenging. Even when the drugs are available, affordability often becomes another
 209 challenge as the majority of persons that in the region lives below the WHO poverty line
 210 [26]. The use of generic forms of these antibiotics, therefore, the norm in the region.

The value of antibiotics in the treatment of open fractures has been established, but this does not substitute for proper wound debridement and adequate skeletal stabilization as an essential aspect of open fracture management. The choice of antibiotic should be guided by the knowledge of possible contaminating organisms at presentation, but subsequent infections are most likely multiple organisms which should be covered by choice of antibiotics. Evidence-based guidelines for prophylactic antibiotic use in open fractures recommend short-course, narrow-spectrum antibiotics for Gustilo Grade I or II open fractures and broader gram-negative coverage for Grade III open fractures [27].

It is worth noting that cultured isolates from a wound especially in the presence of biomaterials and biofilms may not be truly representative of the actual organisms causing infections. Since an infection engrafted on a biomaterial (thick, adherent biofilm) responds poorly to antimicrobial therapy and usually is not cured until the biomaterial is removed, the reliance on only antibiotics without appropriate debridement of dead tissue should be with caution. Antimicrobials that are chosen from the swab culture results may not be effective against all of the bacterial species in these biofilm infections [27]. Incidentally, it takes some time before biofilms developed. Since the cultures in this study were all done within seven days of admission, the identified sensitivity patterns may not be entirely reflective of the antibiotic sensitivity and resistance in open fractures with chronic wounds where there is an existence of biofilms.

CONCLUSION

The hospital antibiotic protocol which recommends the combination of ceftriaxone, quinolones, gentamycin, co-amoxycylav and metronidazole in treating open fractures in the Accident and Emergency department, was based on their sensitivity to cultured microbial organisms in the hospital. The existing microbial and antibiotic sensitivity patterns had not changed significantly over the preceding 12 years when the protocol was established as such there is no reason for a change in the current practice.

REFERENCES

1. Alonge TO, Ogunlade SO, Salawu SA, Fashina AN. Microbial isolates in open fractures seen in the accident and emergency unit of a teaching hospital in a developing country. *West African J Med*. 2002 Oct-Dec;21(4):302-4
2. Gustilo RB, Anderson JT. Prevention of infection in the treatment of one thousand and twenty-five open fractures of long bones: retrospective and prospective analyses. *J Bone Joint Surg. Am*. 1976;58:453-8.
3. Levine NS, Lindberg B, Mason AD Jr, Pruitt BA Jr. The quantitative swab culture and smear: a quick method for determining the number of viable aerobic bacteria on open wounds. *J Trauma* 1976;16: 89-94.

- 250 4. Ikem IC, Oginni LM, Bamgboye EA, Ako-Nai AK, Onipade AO. The bacteriology of open
251 fractures in Ile-Ife, Nigeria. *Niger J Med*. 2004; 13: 359-65.
- 252 5. Bowler P, Duerden B, Armstrong D. Wound microbiology and associated approaches to
253 wound management. *Clinical Microbiology Reviews* 2001; 14: 2, 244-269.
- 254 6. Wilkins J, Patzakis M. Choice and duration of antibiotics in open fractures. *Ortho Clin N*
255 *Amer*. 1991; 22(3): 433-437.
- 256 7. Akinyoola AL, Ako-Nai AK, Dosumu O, Aboderin AO, Kassim OO. Microbial isolates in
257 early swabs of open musculoskeletal injuries. *Niger Postgrad Med J*. 2006 Sep; 13(3): 176-
258 81.
- 259 8. Alonge TO, Salawu SA, Adebisi AT, Fashina AN. The choice of antibiotic in open
260 fractures in a teaching hospital in a developing country. *Int J Clin Pract*. 2002 Jun; 56(5):
261 353 – 6.
- 262 9. Ifesanya AO, Alonge TO, Ogunlade SO, Omololu AB, Nottidge TE, Ayorinde RO.
263 Changing Trends in the pattern of tibial fractures in Nigeria: A review of 70 cases.
264 *J.Orthopaedics* 2008;5(2)e4.
- 265 10. Gustilo RB, Mendoza RM, Williams DN. Problems in the management of type III
266 (severe) open fractures: A new classification of type III open fractures. *J Trauma* 1984;24:
267 742-746.
- 268 11. Paul H. Kim MD, Seth S. Leopold MD. Gustilo-Anderson classification: *Clin Orthop*
269 *Relat Res* 2012;470: 3270 – 3274.
- 270 12. Zalavras CG, Marcus RE, Levin LS, Patzakis MJ. Management of open fractures and
271 subsequent complications. *J. Bone Joint Surg. Am*. 2007;89: 884-95.
- 272 13. Okike K, Bhattacharyya T. Trends in the management of open fractures. A critical
273 analysis. *J Bone Joint Surg. Am*. 2006;88: 2739-48.
- 274 14. Patzakis MJ, Wilkins J. Factors influencing infection rate in open fracture wounds.
275 *ClinOrthopRelat Res*. 1989;243: 36 – 40.
- 276 15. Ostermann PA, Seligson D, Henry SL. Local antibiotic therapy for severe open fractures.
277 A review of 1085 consecutive cases. *J Bone Joint Surg Br*. 1995; 77: 93 – 7.
- 278 16. Gosselin RA, Roberts I, Gillespie WJ. Antibiotics for preventing infection in open limb
279 fractures. *Cochrane Database Syst Rev*. 2004; (1) CD003764.
- 280 17. Khosravi AD, Ahmadi F, Salmanzadeh S, Dashtbozorg A, AbasiMontazeri E. Study of
281 bacteria isolated from orthopedic implant infections and their antimicrobial susceptibility
282 pattern. *Res J Microbiol*. 2009; 4: 158-63.
- 283 18. Courvalin P. Antimicrobial Drug Resistance: “Prediction Is Very Difficult, Especially
284 about the Future.” *Emerg Infect Dis*. 2005;11: 1503-6.
- 285 19. Garazzino S, Aprato A, Maiello A, Masse A, Biasibetti A, De Rosa FG, Di Per G.
286 Osteomyelitis Caused by *Enterobacter cancerogenus* Infection following a Traumatic
287 Injury: Case Report and Review of the Literature. *Journal of Clinical Microbiology*
288 2005; 43(3):1459 – 1461.

- 289 20. Gristina AG, Costerton JW. Bacterial adherence to biomaterials and tissue. The
290 significance of its role in clinical sepsis. *J Bone Joint Surg Am.* 1985; 67(2):264-73.
- 291 21. Gebreselassie S. Pattern of isolates of common gram positive bacterial pathogens and
292 their susceptibilities to antimicrobial agents in Jimma Hospital. *Ethiop Med J.* 2002;
293 40(2):115-27
- 294 22. Yishak A., Wamisho BL. Microbial susceptibility of bacteria isolated from open fracture
295 wounds presenting to the Err of black-lion hospital, Addis Ababa University, Ethiopia
296 *African Journal of Microbiology Research* 2009; 3(12): 939-951.
- 297 23. Gustilo RB, Mendoza RM, Williams DN. Problems in the management of type III
298 (severe) open fractures: A new classification of type III open fractures. *J Trauma* 1984; 24:
299 742-746.
- 300 24. Purghele F, Badea R, Ciuvica R, Anastasiu A. The use of antibiotics in traumatology and
301 Orthopaedic surgery. *Journal of clinical medicine*, 2006; 1: 58-65.
- 302 25. Sitati FC, Mosi PO, Mwangi JC. Early Bacterial Cultures from Open Fractures -
303 Differences Before and After Debridement. *Ann Afr Surg.* 2017;14(2):66-70.
- 304 26. <https://www.indexmundi.com/g/g.aspx>. Accessed: 12th December, 2017.
- 305 27. Rodriguez L, Jung HS, Goulet JA, Cicalo A, Machado-Aranda DA, Napolitano LM.
306 Evidence-based protocol for prophylactic antibiotics in open fractures: Improved antibiotic
307 stewardship with no increase in infection rates. *Journal of Trauma and Acute Care Surgery.*
308 2014; 77(3):400-408.