

Analysis of Lower Leg Ulcer Pathogens using 16S rRNA Gene Based Method

Abstract

The study was carried out to determine the bacterial pathogens associated with lower leg ulcers in Ebonyi State from July, 2016 to July of 2017, using wound swabs from eligible patients with lower leg ulcers. The swabs were processed and analysed using standard microbiological methods, isolated microbial pathogens were identified by employing standard biochemical test, microbial identification tests and standard molecular methods for DNA extraction. Pressure ulcers 450 (37.2 %) was found to be the most commonly infected, closely followed by diabetic foot ulcers 300 (24.8%) and non-healing surgical ulcers 210 (17.4%). *Staphylococcus aureus* strain ATCC 12600 (12.0%, 6.7% and 16.7%) was the most predominant in venous, diabetic and non-healing surgical ulcers respectively, while pressure ulcer had *Pseudomonas aeruginosa* strain M37351 (8.2%). Out of 1500 specimens examined, 1210 (80.7%) showed positive microbial growth, while 290 (19.3%) were not infected. Age group of 31-40 years had the highest prevalence rate of 20.7%, followed by 41-50 years (20.5%) while the least was 0-10 years (0.1%). The males were mostly affected than females. This study has revealed a high index of microbial involvement in lower leg ulcer in Ebonyi State. We recommend a multidisciplinary approach to leg ulcer management and specific intervention strategies, not only to treat but also to reduce and subsequently prevent their spread in rural communities. This results and findings will hopefully help to create awareness on the imperative to improve the quality of the treatment regime employed. Thus, each health institution should carry out a survey to determine the common microbial wound pathogens among their patients.

Key words: lower leg ulcer, bacteria, 16S rRNA and pathogens

Introduction

The term ulcer refers to a serious, long-lasting wound, and is described as chronic if it does not show any healing tendency within six weeks. Ulcers (wounds) may arise post-operatively, following farm injuries, scratches, hoe cuts, thorns cuts, burns, or in association with certain medical conditions such as diabetes mellitus, haemoglobinopathy, lower extremities arterial disease, vasculitis, ulcerative skin diseases and malignancies [1].

Leg ulcer is known as lower limb ulcer, is defined as a defect in the skin below the level of knee persisting for more than six weeks and shows no tendency to heal after three or more months [2]. Ulceration of the lower legs is a relatively common condition amongst adults, and it causes pain and social distress [3]. Ulcers of skin can result in complete loss of the epidermis and often portions of the dermis and even the subcutaneous fat [4]. The incidence of ulceration is rising as a result of the aging population and increased risk factors for atherosclerotic occlusion such as smoking, obesity, and diabetes. In the course of a lifetime, almost 10% of the population will develop a chronic wound, with a wound-related mortality rate of 2.5%. [5] reported that inappropriate treatment of acute traumatic wounds was the most common cause of the chronic wound. A study carried out by [6] showed that the principal etiology (67%) of ulceration is trauma or traumatic wounds compounded by infection, while diabetic ulcers, venous ulcers, and pressure ulcers accounted for 4.9%, 6.5%, and 9.2%, respectively. The majority of these wounds were seen in farmers and other agricultural workers [6, 7]. The current spread of multi-drug resistant bacteria pathogens has added a new dimension to the problem of wound infections. This is particularly worse in resource-poor countries where the sale of antibiotics is under poor control [8]. Leg ulcers are debilitating and they greatly reduce the patients' quality of life.

In this study, we employed 16S ribosomal RNA gene based method to investigate pathogens associated with lower leg ulcers in Ebonyi State, Nigeria. Through this approach, we identified the predominant bacterial species present within the ulcer types and assessed whether differences in patient demographics affected the composition of the microbes in these samples.

48 **Materials and Methods**

49 **Study Area**

50 The study was carried out at the Federal Teaching Hospital Abakaliki (FETHA) and private hospitals in
51 the three Senatorial Districts of Ebonyi State.

52 **Study Design**

53 A total of 1500 specimens were collected from patients of lower leg ulcers in the three Senatorial Districts
54 of Ebonyi State.

55 **Ethical Consideration**

56 Prior to the commencement of the study, ethical clearance was sought from the Federal Teaching
57 Hospital Abakaliki (FETHA) ethical committees. In addition, letters requesting for collaboration was written
58 to the management of all the private hospitals from which specimens were collected and consent was
59 obtained from parents or close relatives.

60 **Collection of Specimens**

61 The specimens were collected with sterile swab sticks in accordance with standard routine procedure. If
62 delay is unavoidable (more than 2 hours), specimens were placed in Stuarts transport medium and
63 refrigerated immediately [1]. Specimens were analyzed on the same day of collection. A structurally
64 designed questionnaire was used for obtaining information concerning each patient. Specimens were
65 completely and properly labeled as well as from the hospital records. Patients on antimicrobial therapy
66 within 72hr of presentation were included in the study.

67 **Culture of Specimens on Media**

68 Cultures of the specimens were made on nutrient agar, Mannitol Salt Agar, EMB, blood and MacConkey
69 agar for the isolation of the bacterial. After incubation at 37°C for 24hrs, the plates were observed;
70 carefully examined and distinct growths were sub-cultured on fresh medium for purity. The bacterial
71 isolates were identified using their cultural identification, morphological and other biochemical
72 characteristics as described by Bergey's Manual of Determinative Bacteriology [9,10].
73

74 **Identification of isolates by DNA sequencing**

75 The Methods employed were the following.

76 1: Culture on Nutrient Broth

2: DNA extraction: ZR D3024 Quick-gDNA™ MiniPrep (50 Preps)

77 3: PCR amplification and DNA sequencing by Sanger sequencing method

78 4: Blasting analysis: CLC BIO AND NCBI BLAST ONLINE

79 **SEQUENCING: Sanger sequencing**

80 **Primer that was used:** Bacteria: 16S, 27-F

81 **Molecular Analysis:**

82 DNA Extraction was performed at the Anaerobe Laboratory, Molecular Biology and biotechnology
83 Division, Nigerian Institute of Medical Research, Yaba, Lagos. Methodology was based on PCR and
84 metagenomics analysis. While sequencing analysis was done at Inqaba Biotechnology Pty South Africa.

85 **DNA extraction ZR D3024 Quick-gDNA™ MiniPrep (50 Preps)**

86 DNA extraction was from a 24 hours growth of microbial isolates in BHI broth harvested by centrifugation
87 at 14, 000 x g for 10 minutes. The cells was washed three times in 1 ml of ultra-pure water by centrifuging
88 at 12,000 rpm for 5 min. DNA extraction and purification was done using ZR Fungal/Bacterial DNA
89 MiniPrep™50 Preps Model D6005 (Zymo Research, California, USA). 50-100 mg of bacterial cells was
90 resuspended in 200µl of sterile water. This was transferred into a ZR Bashing Bead™ Lysis Tube. Exactly
91 750µl lysis solution was added to the tube. The bead containing the solution was secured in a bead

92 beater fitted with a 2 ml tube holder assembly and process at maximum speed for 5 minutes. The ZR
93 BashingBead™ Lysis Tube was centrifuged in a microcentrifuge at 10,000 x g for 1 minute. 400µl of the
94 supernatant was pipetted into a Zymo-Spin™ IV Spin Filter in a Collection Tube and centrifuged at 7,000
95 x g for 1 minute. This was followed by the addition of 1,200µl of Bacterial DNA Binding Buffer into the
96 filtrate in the Collection Tube. After this, 800µl of the mixture was transferred into a Zymo-Spin™ IIC
97 Column in a Collection Tube and centrifuge at 10,000 x g for 1 minute.
98 The flow through was discarded from the Collection Tube and the process was repeated to obtain the
99 remaining products. The 200µl DNA Pre-Wash Buffer was added into the Zymo-Spin™ IIC Column in a
100 new Collection Tube and centrifuge at 10,000 x g for 1 minute. This was followed by the addition of 500µl
101 Bacterial DNA Wash Buffer into the Zymo-Spin™ IIC Column and centrifuged at 10,000 x g for 1 minute.
102 The Zymo-Spin™ IIC Column was transferred into a clean 1.5 ml micro-centrifuge tube and 100µl of DNA
103 Elution Buffer was then added directly to the column matrix. This was centrifuged at 10,000 x g for 30
104 seconds to elute the DNA. The Ultra-pure resulting filtrate (DNA) obtained was used as a template during
105 the assay. This was transported in ice to the laboratory for sequencing.

106 **Gel Electrophoresis of DNA**

107 For the electrophoresis, 0.8% agarose was prepared by weighing 0.8 g of agarose powder with a
108 weighing balance. The powder was mixed with 100ml of electrophoresis buffer and then heated in a
109 microwave oven until completely melted. Ethidium bromide was added to the gel at a final concentration
110 0.5ug/ml to facilitate visualization of DNA after electrophoresis. After cooling the solution to about 60°C, it
111 was poured into a casting tray with a comb placed across its rim to form wells. The gel was allowed to set
112 for 30 minutes and the comb was removed. 20µl of the DNA samples was then loaded into the wells after
113 mixing with 2µl of bromophenol blue. A DNA molecular weight marker was loaded into one of the wells.
114 The plastic tray with the gel was inserted horizontally into the electrophoresis chamber and covered with
115 buffer. 70V was applied for 1hr 30 min. The distance DNA migrated in the gel was judged by visually
116 monitoring migration of the tracking dyes. The DNA was visualized by placing the gel in an ultraviolet
117 transilluminator in a photo documentation system (Clinix Japan, Model 1570). The size of the visible
118 bands obtained was calculated by matching that of the isolates with the standard bands produced by
119 HIND III marker.

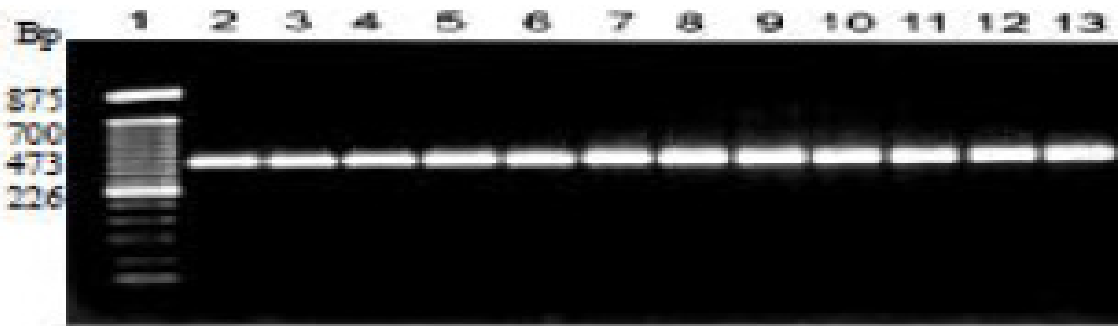
120 **PCR amplification and DNA sequencing by Sanger sequencing method and Blast analysis (CLC 121 BIO and NCBI blast online)**

122 DNA sequencing was performed by Sanger (dideoxy) Sequencing Technique to determine the nucleotide
123 sequence of the specific microorganism isolated using automated PCR cycle- Sanger Sequencer™
124 3730/3730XL DNA Analyzers from Applied Biosystems [11,12]. This result was obtained as nucleotides.
125 Sequence analysis from resultant nucleotides base-pairs was performed by BLAST analysis by using
126 CLO Bio software and by direct blasting on <http://blast.ncbi.nlm.nih.gov>.
127 For every set of isolate, a read was BLASTED and the resultant top hits for every BLAST result showing
128 species name was used to name the specific organism

129 **Results**

130 The bacteria associated with lower leg ulcers of patients in Ebonyi State were studied.

131 Figure 1 depicted the PCR amplification product of the 16S rRNA gene using the primer 27F (5'AGA GTT
132 TGA TCC TGG CTC AG-3'). This was separated on a 1.5% agarose gel electrophoresis. The DNA bands
133 were visualized after ethidium bromide staining using 100bp DNA ladder as DNA molecular weight
134 standard.



135
 136 Fig.1: Amplification of PCR products from isolates.
 137 Lane 1 was 100 bp DNA ladder. Lanes 2–13 were PCR amplified from different isolates.

138 The gene sequences of the various bacterial isolates are presented in Tables 1-13. The result shows the
 139 nucleotide sequence of the isolated bacteria based on 16S rRNA. The blasting of the sequence results
 140 was done using the online blast software at <http://blast.ncbi.nlm.nih.gov/blast.cgi>. The result for every set
 141 of isolate was taken from the top hit of the blast showing species name and the strain number. Following
 142 this, the bacteria identified were *Staphylococcus aureus* strain ATCC 12600, *Enterococcus faecalis* strain
 143 ATCC 19433, *Proteus mirabilis* strain ATCC 29906, *Proteus vulgaris* strain ATCC 29905, *Klebsiella*
 144 *aerogenes* strain KCTC 2190, *Escherichia coli* strain U 5/41, *Serratia marcescens* strain NBRC 102204,
 145 *Pseudomonas aeruginosa* strain M37351, *Streptococcus pyogenes* KS030, *Enterobacter cloacae* strain
 146 NBRC 13535, *Peptostreptococcus* strain *anaerobius* DSM 2949, *Clostridium tetani* strain E88 and
 147 *Corynebacterium ulcerans* strain 0102.

148 Table 1: Nucleotide sequence of the isolate code LU10 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe/strain code	GeneBank/Accession number
U5	LU 10	AGGATGAACGCTGGCGGCGTGCCTAATACATGCAAGTCG AGCGAACGGACGAGAAGCTTGCTTCTCTGATGTTAGCGG CGGACGGGTGAGTAACACGTGGATAACCTACCTATAAGA CTGGGATAACTTCGGGAAACCGGAGCTAATACCGGATAA TATTTTGAACCGCATGGTTCAAAAGTGAAAGACGGTCTTG CTGTCACTTATAGATGGATCCGCGCTGCATTAGCTAGTTG GTAAGGTAACGGCTTACCAAGGCAACGATGCATAGCCGA CCTGAGAGGGTGATCGGCCACACTGGAAGTGAACACG GTCCAGACTCCTACGGGAGGCAGCAGTAGGGAATCTTCC GCAATGGGCGAAAGCCTGACGGAGCAACGCCGCGTGAG TGATGAAGG	<i>Staphylococcus aureus</i> strain ATCC 12600, 16S rRNA gene	NR 115606.1

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 150 Table 2: Nucleotide sequence of the isolate code LU14 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank/Accession number
U9	LU14	GACGAACGCTGGCGGCGTGCCTAATACATGCAAGTCGAA CGCTTCTTTCTCCCGAGTGCTTCACTCAATTGGAAAGA GGAGTGGCGGACGGGTGAGTAACACGTGGGTAACCTAC CCATCAGAGGGGATAACACTTGGAAACAGGTGCTAATA CCGCATAACAGTTTATGCCGCATGGCATAAGAGTGAAAG GCGCTTTCGGGTGTGCGCTGATGGATGGACCCGCGGTGC ATTAGCTAGTTGGTGAAGTAACGGCTCACCAAGGCCACG ATGCATAGCCGACCTGAGAGGGTGTGATCGGCCACACTGG GACTGAGACACGGCCAGACTCCTACGGGAGGCAGCAG TAGGGAATCTTCGGCAATGGACGAAAGTCTGACCGAGCA	<i>Enterococcus faecalis</i> strain ATCC 19433, 16S rRNA gene	NR 115765.1

ACGCCGCGTGAGTGAAGAAGGTTTTCGGATCGTAAACT
 CTGTTGTTAGAGAAGAACAAGGACGTTAGTAACTGAACGT
 CCCCTGACGGTATCTAACCAGAAAGCCACGGC

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152 Table 3: Nucleotide sequence of the isolate code LU15 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank/ Accession number
U10	LU15	TGATCCTGGCTCAGATTGAACGCTGGCGGCAGGCCTAAC ACATGCAAGTCGAGCGGTAACAGGAGAAAGCTTGCTTTC TTGCTGACGAGCGGCGGACGGGTGAGTAATGTATGGGG ATCTGCCCGATAGAGGGGGATAACTACTGGAAACGGTGG CTAATACCGCATAATGTCTACGGACCAAAGCAGGGGGCTC TTCGGACCTTGCACTATCGGATGAACCCATATGGGATTAG CTAGTAGGTGGGGTAAAGGCTCACCTAGGCGACGATCTC TAGCTGGTCTGAGAGGATGATCAGCCACACTGGGACTGA GACACGGCCCAGACTCCTACGGGAGGCAGCAGTGGGGA ATATTGCACAATGGGCGCAAGCCTGATGCAGCCATGCCG CGTGTATGAAGAAGGCCTTAGGGTTGTAAAGTACTTTTCAG CGGGGAGGAAGGTGATAAGG	<i>Proteus mirabilis</i> strain ATCC 29906, 16S rRNA gene	NR 114419.1

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154 Table 4: Nucleotide sequence of the isolate code LU16 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank/ Accession number
U11	LU16	AAATTGAAGAGTTTGATCATGGCTCAGATTGAACGCTGGC GGCAGGCCTAACACATGCAAGTCGAGCGGTAGCACAGA GAGCTTGCTCTCGGGTGACGAGCGGCGGACGGGTGAGT AATGTCTGGGAAACTGCCTGATGGAGGGGGATAACTACT GGAAACGGTAGCTAATACCGCATAACGTCGCAAGACCAA AGTGGGGGACCTTCGGGCCTCATGCCATCAGATGTGCC AGATGGGATTAGCTAGTAGGTGGGGTAATGGCTCACCTA GGCGACGATCCCTAGCTGGTCTGAGAG	<i>Klebsiella aerogenes</i> strain KCTC 2190, 16S rRNA	NR 102493.2

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156 Table 5: Nucleotide sequence of the isolate code LU17 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank/ Accession number
U12	LU17	TCAGATTGAACGCTGGCGGCAGGCCTAACACATGCAATC GAGCGGTAACAGAAGAAAGCTTGCTTTCTTGCTGACGAG CGGCGGACGGGTGAGTAATGTATGGGGATCTGCCCGAT AGAGGGGGATAACTACTGGAACGGTGGCTAATACCGCA TGACGTCTACGGACCAAAGCAGGGGCTCTTCGGACCTTG CGCTATCGGATGAACCCATATGGGATTAGCTAGTAGGTG AGGTAATGGCTCACCTAGGCAACGATCTCTAGCTGGTCT GAGAGGATGATCAGCCACACTGGGACTGAGACACGGCC CAGACTCCTACGGGAGGCAGCAGTGGGGAATATTGCACA ATGGGCGCAAGCCTGATGCAGCCATGCCGCGTGTATGAA GAAGGCCTTAGGGTTGTAAAGTACTTTTCAGCGGGGAGGA AGGTGATAAAGTTAATACCTTT	<i>Proteus vulgaris</i> strain ATCC 29905, 16S rRNA gene	NR 115878.1

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158 Table 6: Nucleotide sequence of the isolate code LU18 based on 16S ribosomal RNA

Specimen	Isolati	Gene sequence	Microbe /strain	GeneBank/
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n code	on code	Gene sequence	Microbe /strain code	Accession number
U13	LU18	AGTTTGATCATGGCTCAGATTGAACGCTGGCGGCAGGCC TAACACATGCAAGTCGAACGGTAACAGGAAGCAGCTTGC TGCTTTGCTGACGAGTGGCGGACGGGTGAGTAATGTCTG GGAAACTGCCTGATGGAGGGGGATAACTACTGGAAACGG TAGCTAATACCGCATAACGTCGCAAGCACAAAGAGGGGG ACCTTAGGGCCTCTTGCCATCGGATGTGCCCAGATGGGA TTAGCTAGTAGGTGGGGTAACGGCTCACCTAGGCGACGA TCCCTAGCTGGTCTGAGAGGATGACCAGCAACTGGAA CTGAGACACGGTCCAGACTCCTACGGGAGGCAGCAGTG GGGAATATTGCACAATGGGCGCAAGCCTGATGCAGCCAT GCNGCGTGTATGAAGAAGGCCTTCGGGTTGTAAAGTACT TTCAG	<i>Escherichia coli</i> strain U 5/41, 16S rRNA gene.	NR 024570.1

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Table 7: Nucleotide sequence of the isolate code LU19 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank / Accession number
U14	LU19	ATTGAACGCTGGCGGCAGGCTTAACACATGCAAGTCGAG CGGTAGCACAGGGGAGCTTGCTCCCTGGGTGACGAGCG GCGGACGGGTGAGTAATGTCTGGGAACTGCCTGATGGA GGGGGATAACTACTGGAAACGGTAGCTAATACCGCATAA CGTCGCAAGACCAAAGAGGGGGACCTTCGGGCCTCTTG CCATCAGATGTGCCCAGATGGGATTAGCTAGTAGGTGGG GTAATGGCTCACCTAGGCGACGATCCCTAGCTGGTCTGA GAGGATGACCAGCCACACTGGAAGTGAAGACACGGTCCA GACTCCTACGGGAGGCAGCAGTGGGGAATATTGCACAAT GGGCGCAAGCCTGATGCAGCCATGCCGCGTGTGTGAAG AAGGCCTTCGGGTTGTAAAGCACTTTCAGCGAGGAGGAA GGTGGTGAGCTTAATACGTTCAATTGACGTTACTCGC AGAAGAAGCACCGGCTAACTCCGTGCCAGCAGCCG	<i>Serratia marcescens</i> strain NBRC 102204, 16S rRNA gene	NR 114043.1

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Table 8: Nucleotide sequence of the isolate code LU20 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank / Accession number
U15	LU20	TTCCTTGAGCGACAGCGCCGCGATGATGGCTTTCAGGAG TTCATCCACGGAGGTTTCCTCTTGGGTGGGGTAAAGGCC AAAGGTGGCCCGCGCTCGGCGGGCGAGCGGCTCCATGC CATCGATGGCGGGTCTTTGGTATGGCGCCCATCAGAAT CGACAGGACCGTGCCGTCCGGGGCGTAGCTGAAGACCG GCGAGAAGTAGAGGTAATCACCCTCCTCGATCATCCGTG CGGCGCGGGCGGTGTATTTCGACACGGCCCCACAGGCCG GAGCCTTCGCGCCATTGAAATCGAGGAAGCGGCCGGC AGCAGGCGCCGGCTGGCCGTTTTCTCTTTCTGAGGGTC TGGTGCTCATAGTCCAGGACAGGCGGTGTCTTGCGTGCC CTGGCCCCGTC	<i>Pseudomonas aeruginosa</i> strain M37351, 16S rRNA gene	CP 008863.1

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Table 9: Nucleotide sequence of the isolate code LU22 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank / Accession number
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				Accession number
U17	LU22	TTGTTGATATTCTGTTTTTCTTTTTAGTTTTCCACATAAA AAATAGTTGAAAACAATAGCGGTGTCACCTTAAAATGACT TTCCACAGGTTGTGGAGAACCCAAATTAACAGTGTTAAT TTATTTTCCACAGATTGTGGAAAACTAACTATTATCCATT GCTCTGTGGAAAAGTAAGTATGTTATGGTAGAATAGTTC TAGAATTATCCACAAGAAGGAACCTAGTATGACTGAAAAT GAACAAATTTTTTGAACAGGGTCTTGAATTAGCTCAGA GTCAATTAAACAGGCAACTTATGAATTTTTGTTCATGAT GCCCGTCTATTAAGGTCGATAAGCATATTGCAACTATTT ACTTAGATCAAATGAAAGAACTCTTTTGGGAAAAAATCTT AAAGATGTTATTCTTACTGCTGGTTTTGAAGTTTATAACGC TCAAATTTCT	<i>Streptococcus pyogenes</i> MGAS315, 16S rRNA gene	AE 014074.1

166 Table 10: Nucleotide sequence of the isolate code LU23 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank/ Accession number
U18	LU23	ATTGAACGCTGGCGGCAGGCCTAACACATGCAAGTCGAA CGGTAGCACAGAGAGCTTGTCTCGGGTGACGAGTGGC GGACGGGTGAGTAATGTCTGGGAACTGCCTGATGGAG GGGATAACTACTGGAAACGGTAGCTAATACCGCATAAN GTGCAAGACCAAAGAGGGGGACCTTCGGGCCTTTGC CATCAGATGTGCCAGATGGGATTAGCTAGTAGGTGGGG TAACGGCTCACCTAGGCGACGATCCCTAGCTGGTCTGAG AGGATGACCAGCCACACTGGAAGTACGACACGGTCCAGA CTCCTACGGGAGGCAGCAGTGGGGAATATTGCACAATGG GCGCAAGCCTGATGCAGCCATGCCGCGTGTATGAAGAAG GCCTTCGGGTGTAAAGTACTTTTACGCGGGGAGGAAGGT GTTGTGGTTAATAACCGCAGCAAT	<i>Enterobacter cloacae</i> strain NBRC 13535, 16S ribosomal RNA gene	NR 113615.1

167 Table 11: Nucleotide sequence of the isolate code LU24 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank / Accession number
U19	LU24	GACTTCTCACAGTTGTTGTTACTAGGTTTTCCCTCTGTG AGACCTCACGGGATAAGTCGTACGTCCTTCCCTCGTCTACC TGCCTAATTTACTTACATAAGTTACGTTTGCCTTTTGGACT TCGTGACTTTGGGCCCACTCATCCTTTATGTAAGCCTTTA TATTAGTTTTCTGTACGTCAGGCTACGATTTTGTCTATTGCT TCTTCTCGCCATCACCTCACGGTGATAACCTTGCAAGTTG CTATGAGGTTTCGTGCGCAACTACGCCCTACGTGGACTTT CACCACAGACTGACGGCATGCCCGTCATACCCAAAAAAA GAAGCCCACAGTATATAACTGAGGGCTTGTATCTTCTATT ATTTTATTAATAAACTCGGCTGCCTTTACCTTTGAAGGTA TAAATTTACCTATAACACCACCTATGATAGCTGGTACTAC CCAGTTGAATCCTAGGAATCCAAGTGGTAGGTTGTTTACA GGTGCTAGTGATGGTATACTTGCTGATAATAGTGTAAGAG CACTTGTTATCACAGTAAATAGTACAGACACCTTGTAGA	<i>Peptostreptococcus</i> strain <i>anaerobius</i> DSM 2949	KB 906605.1

169 Table 12: Nucleotide sequence of the isolate code LU25 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank/ Accession number

				n number
U20	LU25	CAGAGCTTTGGAATAAGATAGGTGATAATAAAAATACATA TAGTGACTTATTAGAGCTGTTTAATAAAGTAGGAGAAAA TATTCAGATACAATAAAAAAGACTATTTGGGTTTATAATAG AGAATTTTTTATATACTAATATGACAGAAATAACTGAAAA TACAAAAATGAATATTTAACATTGAAAAAGAGGAATTTA AGGGTAAAATAAGTTTATAAATAAAAATAATTTGAATTTAT ATAAACAAAAATAAAAACCTCTTAAAGGCTGAAAGTTTCG CGGCTTTACCTTTAAGAGTTATAAGAGTACCCTATATAGA AAAATTCTACATAGGAGAACCTTTATTTACTTAAAGACATTT TAATATATTTATTGTATATAGTCAATAGTAAAAAGTTCTTCT ATAACTTTGGGTACAAGATAGGAGGACTTTATTTGTTAGC AAAACAATTAAGCTTATATGATTATATAGACAATACATTA AAAACCTTAGTAACCTGTGAAGTGGAAATAAAGGCAGAGG AAGATAACATACAAAAGGCAACACTT	<i>Clostridium tetani</i> strain E88	AF 528097.1

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172 Table 13: Nucleotide sequence of the isolate code LU26 based on 16S ribosomal RNA

Specimen code	Isolation code	Gene sequence	Microbe /strain code	GeneBank/Accession number
U21	LU26	TGTGGATAACTCTCAAGTTTATGTGCGAACTTTCCACATA GAATTGAATTTACGCAGGTTAAAAGCAAATTCCTTCACAG TGAGCGTTATCCACAAGGTTGTGAAATAACTGTGGATAAC TTTTCACACACCCAGTGTAGGAAGTTATCCACAGTTGTG GAAAACCTCTGTGGAATACGCGGTCACAGCCCCAAAGCGT TGTGAACAACCTCGGTGAAATCCCCGTGGACAGTGAATA ACATTTTTTACCAGCCGGATGGGGAAATCTCGTTGCGTTG GACATTATTTTTGGGGCTAGTGAGCGCTGCTGATAGGGC ACTGACCCGCGAATGCACTGCCATATCAGCATGAAAATTA GTTGTTGTTTCATTTCGCACGAGGTAAAGGAAATACGAAGTG TCGGAGACTCCATCGACATGGAACGAGCGGTGGCAGGA AGTTACTAACGAGCTGCTGTCACAGTCTCAGGACCCAGA TAGTGGTATTTCCATTACTCGCCAGCAAAGTGCATACCTG CGATTGGTAAAGCCAGTCGCGTTTGTAGAGGGTATTGCA GTTTTAAGCGTCCCTCACGCCGAGCGAAAAAAGAGATT GAAACTACGCT	<i>Corynebacterium ulcerans</i> strain 0102	AP 012284.1

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176 As shown in Table 14, the ulcer type designated as pressure ulcers, which amounted to 450 (37.2 %)
 177 were found to be the most commonly infected. All acute soft tissue wounds such as road traffic accidents,
 178 lacerations, domestic violence, burn sites and gunshot injuries were classified under pressure ulcer. This
 179 was closely followed by diabetic foot ulcers, 300 (24.8%). Non-healing surgical ulcers 210 (17.4%) were
 180 the least frequent.

Table 14: Prevalence of lower ulcer types in Ebonyi State.

Category of patients	No examined	No. infected (%)	No. uninfected (%)
Venous leg ulcer	300	250 (20.7)	50 (17.2)
pressure ulcer	600	450 (37.2)	150 (51.7)
Diabetic foot ulcer	350	300 (24.8)	50 (17.2)
Non-healing surgical ulcer	250	210 (17.4)	40 (13.8)

Total 1500 1210 (80.6) 290 (19.3)

181
 182 The distribution of microorganisms isolated from lower leg ulcer patients according to the ulcer type
 183 showed that *Staphylococcus aureus* (12.0%) was the most predominant in venous leg ulcer, followed
 184 *Pseudomonas aeruginosa* which had 8.0%, whereas *Peptostreptococcus* (1.2%) was the least in
 185 occurrence. In pressure ulcer, the most predominant microbe was *Pseudomonas aeruginosa* (8.2%),
 186 followed by *Escherichia coli* (7.6%). Diabetic leg ulcer had *Staphylococcus aureus* (6.7%) and
 187 *Escherichia coli* strain 6.7%) as the highest in occurrence, followed by *Pseudomonas aeruginosa* (6.3%),
 188 as the least was *Clostridium tetani* strain (2.0%). The non-healing surgical ulcer had *Staphylococcus*
 189 *aureus* (16.7%) and *Streptococcus pyogenes* (16.7%) as the most occurring microbes, followed by 15.2%
 190 for *Pseudomonas aeruginosa*, *Streptococcus pyogenes* and *Corynebacterium ulcerans* respectively
 191 (Table 15).

Table 15: Occurrence of microorganisms isolated from lower leg ulcer patients according to ulcer type

Ulcer type	<i>Staphylococcus aureus</i>	<i>Staphylococcus aureus</i>	<i>Staphylococcus saprophyticus</i>	<i>Staphylococcus epidermidis</i>	<i>Enterococcus faecalis</i>	<i>Proteus mirabilis</i>	<i>Klebsiella aerogenes</i>	<i>Proteus vulgaris</i>	<i>Escherichia coli</i>	<i>Serratia marcescens</i>	<i>Pseudomonas aeruginosa</i>	Total
Venous	30(12.0)	20(8.0)	10(4.0)	10(4.0)	5(2.0)	10(4.0)	15(6.0)	13(5.2)	12(4.8)	6(2.4)	20(8.0)	250(20.7)
Pressure	31(6.9)	29(6.4)	14(3.1)	16(3.6)	13(2.9)	25(5.6)	26(5.8)	22(4.9)	34(7.6)	14(3.1)	37(8.2)	450(37.2)
Diabetic	20(6.7)	17(5.7)	12(4.0)	11(3.7)	10(3.3)	16(5.3)	16(5.3)	14(4.7)	20(6.7)	10(3.3)	19(6.3)	300(24.8)
Non healing surgical	35(16.7)	9(4.3)	2(1.0)	0(0.0)*	0(0.0)*	3(1.4)	10(4.8)	1(0.5)	2(1.0)	0(0.0)*	32(15.2)	210(17.4)
Total	116(9.6)	75(6.2)	38(3.1)	37(3.1)	28(2.3)	54(4.5)	67(5.5)	50(4.1)	68(5.6)	30(2.5)	108(8.9)	1210(100)

192 p ≥ 0.05: Ulcer type differed significantly except for values with asterisks.

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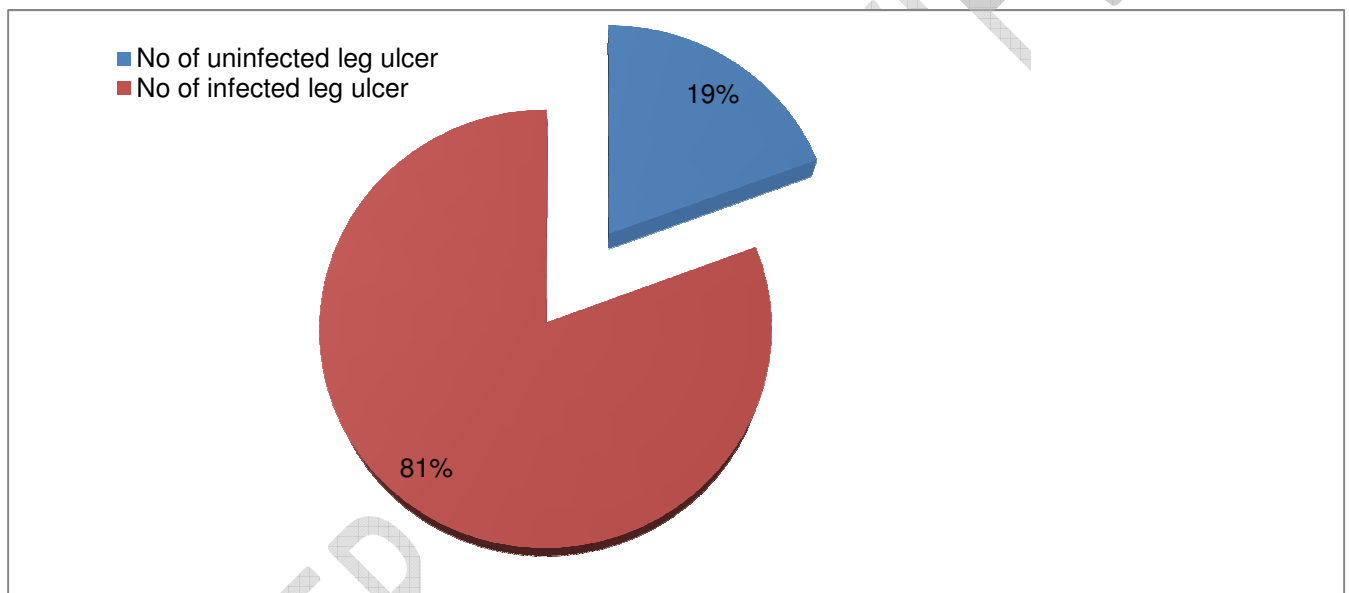
Table 15 conti.: Occurrence of microorganisms isolated from lower leg ulcer patients according to ulcer type

Ulcer type	<i>Streptococcus pyogenes</i>	<i>Streptococcus pyogenes</i>	<i>Enterobacter cloacae</i>	<i>Peptostreptococcus</i>	<i>Clostridium tetani</i>	<i>Corynebacterium ulcerans</i>	<i>Klebsiella pneumoniae</i>	Total
Venous	15(6.0)	15(6.0)	6(1.2)	6(1.2)	5(2.0)	13(5.2)	10(4.0)	250(20.7)

Pressure	24(5.3)	32(7.1)	11(2.4)	16(3.6)	14(3.1)	24(5.3)	13(2.9)	450(37.2)
Diabetic	15(5.0)	17(5.7)	12(4.0)	12(4.0)	6(2.0)	17(5.7)	11(3.7)	300(24.8)
Non healing surgical	35(16.7)	32(15.2)	0(0.0)*	5(2.4)	0(0.0)*	32(15.2)	10(4.8)	210(17.4)
Total	89(7.4)	96(7.9)	29(2.4)	39(3.2)	25(2.1)	86(7.1)	44(3.6)	1210(100)

194 $p \geq 0.05$: Ulcer type was significantly different except for values with asterisk.

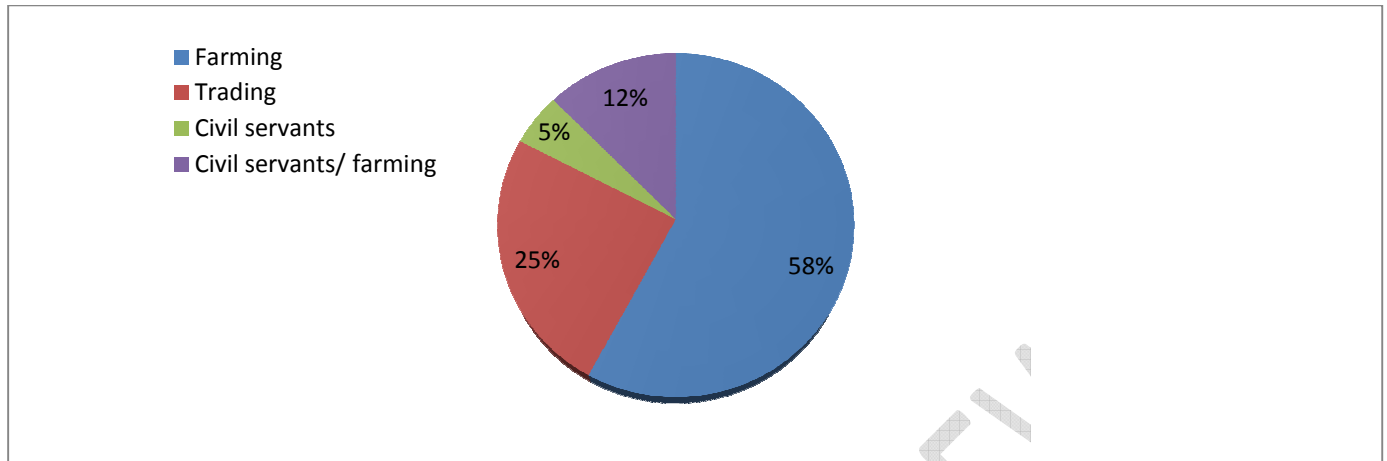
195 Out of 1500 specimens examined, 1210 (80.7%) showed positive microbial association, while 290
 196 (19.3%) did not produce any growth due to the general purpose media employed.



197

198 Figure 2: Prevalence of lower leg ulcers in Ebonyi State.

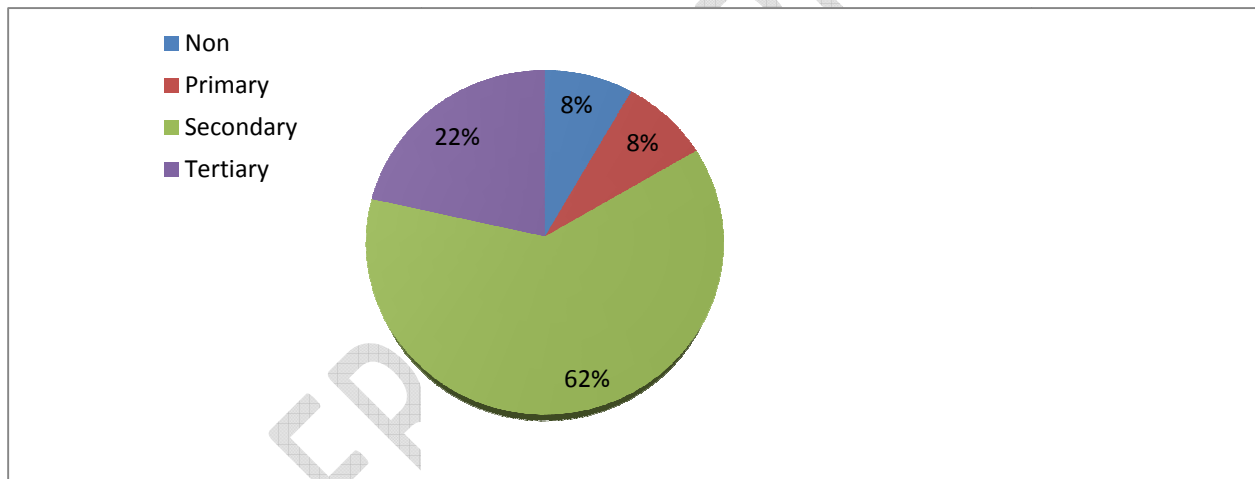
199 The subjects in Ebonyi State are involved in active farming, 58% prevalent rate of leg ulcer was recorded
 200 in this occupation. The infection was less prevalent among civil servants.



201

202 Figure 3: Prevalence of lower leg ulcer in relation to occupation in Ebonyi State.

203 Prevalence of lower leg ulcer in relation to educational level of patients in Ebonyi State showed that the
 204 highest rate was seen among the secondary level (62%), followed by tertiary education (22%), while the
 205 least was 8% for primary and non-educated respectively.



206

207 Figure 4: Prevalence of lower leg ulcer in relation to educational level of patients in Ebonyi State

208 The age and gender prevalence of patients of lower leg ulcer is shown in Table 16. The age group of 41-
 209 50 years (29.5%) was the most infected among females, while 51-60 years was mostly infected among
 210 males.

Table 16: Age and gender prevalence of lower leg ulcers in Ebonyi State.

Age group (years)	No. of specimen examined	No infected (%)	No. uninfected (%)	Male (%)	Female (%)
0-10	5	1(0.1)*	4(1.4)	1(0.1)*	0(0.0)*
11-20	29	12(1.0)	17(5.9)	7(0.9)	5(1.2)
21-30	250	196(16.2)	54(18.6)	99(12.5)	97(23.1)
31-40	300	251(20.7)	49(16.9)	150(19.0)	101(24.1)
41-50	357	248(20.5)	109(37.6)	124(15.7)	124(29.5)
51-60	250	211(17.4)	39(13.5)	161(20.4)	50(11.9)
61-70	180	165(13.6)	15(5.2)	135(17.1)	30(7.1)
71-80	75	72(6.0)	3(1.0)	62(7.9)	10(2.4)

≥ 81	54	54(4.5)	0(0.0) *	51(6.5)	3(0.7)
Total	1500	1210(80.7)	290(19.3)	790(65.3)	420(34.7)

211 p ≥ 0.05: Number of infected and uninfected among age groups and gender were significantly different
212 except for values with asterisks.

213 Discussion

214 Modern molecular tools such as 16S rRNA gene-based sequencing provide powerful means to define
215 chronic wound bacteria. We found that leg ulcers supported complex bacterial communities comprised of
216 a wide-range of bacterial taxa including fastidious anaerobic bacteria that were not observed using
217 culture-based methods. The bacterial communities characterized in this study were similar to those
218 reported by other groups using 16S rRNA gene-based methods [13,14]. Of the numerous organisms that
219 colonize chronic wounds, wound care experts believe *Staphylococcus aureus*, *Pseudomonas aeruginosa*,
220 beta-hemolytic streptococcus, and anaerobes are the most likely bacterial causes of delayed healing and
221 infection. Of these organisms, *S. aureus* is most commonly isolated from chronic wounds with the others
222 occurring at relatively low rates.
223 *S. aureus* is a known pathogen with an extensive array of virulence factors including proteases and
224 toxins. As with most bacteria, these factors are primarily expressed at higher densities to enable the
225 organism to further colonize, and subsequently invade surrounding tissues. Such factors are rarely
226 expressed at lower densities where adherence and survival are paramount. Wound contaminants are
227 likely to originate from the environment (exogenous microorganisms in the air or those introduced by
228 traumatic injury), the surrounding skin (involving members of the normal skin microflora such as
229 *Staphylococcus epidermidis*, micrococci, skin diphtheroids, and propionibacteria), and endogenous
230 sources involving mucous membranes, primarily the gastrointestinal, oropharyngeal and genitourinary
231 mucosae [13].

232 The distribution of microorganisms isolated from lower leg ulcer patients in relation to the ulcer type
233 showed that *Staphylococcus aureus* (12.0%) was the most predominant in venous leg ulcer, followed by
234 *Staphylococcus aureus* subsp. anaerobius and *Pseudomonas aeruginosa* which had 8.0% respectively,
235 whereas *Peptostreptococcus* (1.2%) was the least in occurrence. This is consistent with similar studies in
236 Nigeria which reported that *Staphylococcus aureus* was the predominant pathogen in wound ulcers [15],
237 but differed from the studies of [16] in Okolobiri, Bayelsa, Nigeria and [17] in Sagamu, Nigeria, who
238 reported *P. aeruginosa* and *Klebsiella* respectively as the most common pathogens. Although the
239 presence of *P. aeruginosa* among subjects may be attributed to contamination of wounds with
240 contaminated medical devices, it also shows local variability in wound management procedures.

241 It was observed that pressure ulcers 450 (37.2 %) was the most commonly infected, and these include
242 acute soft tissue infections such as road traffic accidents, lacerations, domestic violence, burn sites and
243 gunshot injuries. This was closely followed by diabetic foot ulcer 300 (24.8%). Infections of non-healing
244 surgical ulcers 210 (17.4%) were the least frequent. This report deferred from the result of [18], who
245 observed that surgical site infection ranked highest among wound infections. [18] further attributed the
246 claim that patients are likely to undergo surgical operations and more likely to have breaks in their local
247 defence system. The predominant microbe in pressure ulcers was *Pseudomonas aeruginosa* (8.2%),
248 followed by *Escherichia coli* (7.6%). Similar findings had also been reported by [19]. In wound sepsis,
249 *Pseudomonas aeruginosa* was also the most prevalent infectious organism caused by incision or fluid
250 collection under the skin surface. This finding differed from that obtained by [20]. The susceptibility of
251 burn wound to opportunistic colonization by bacteria and fungi results from several factors, including the
252 presence of coagulated proteins, the absence of blood-borne immune factors, and the avascularity of the
253 burn wound [21]. Multiple studies had examined the microflora that affects these wounds; both superficial
254 and deep tissue cultures had been used, and the results are fairly consistent. The organisms isolated by
255 culture were *S. aureus*, *S. epidermidis*, and *Streptococcus* spp. [22,23]. Other bacteria that consistently
256 appeared include *Proteus mirabilis*, *P. aeruginosa*, and *Propionibacterium* spp.

257 Diabetic leg ulcers had *Staphylococcus aureus* (6.7%) and *Escherichia coli* (6.7%) as the highest in
258 occurrence, followed by *Pseudomonas aeruginosa* (6.3%), while the least was *Clostridium tetani* (2.0%).

259 [24] reported *S. aureus* as the most prevalent isolate in diabetic foot ulcers, together with other aerobes
260 such as *S. epidermidis*, *Streptococcus* spp., *P. aeruginosa*, *Enterococcus* spp., and coliform bacteria. In
261 addition, increased plasma glucose stimulates the growth of Gram-positive organisms. High glucose
262 levels have been associated with an increased risk of wound infection in both humans and animal
263 models, and hyperglycemia has been demonstrated to be associated with Gram-positive septicemia [25].

264 The non-healing surgical ulcers had the most occurring microbes as *Staphylococcus aureus* and
265 *Streptococcus pyogenes* (16.7%), followed by 15.2% for *Pseudomonas aeruginosa*, *Streptococcus*
266 *pyogenes* and *Corynebacterium ulcerans* respectively. This finding is similar to that reported by [26] who
267 found that 21.3%, 19.0% and 10.9% were *S. aureus*, *E. coli*, *P. mirabilis* and *P. aeruginosa*, respectively.
268 This research finding indicates that the presence of enteric organisms in fresh wounds or at operation
269 probably leads to their subsequent sepsis. These findings, therefore, imply that both enteric and non
270 organisms are important determinants of healing traumatic and other in surgical wounds. The incidence of
271 enteric bacteria also confirms the observation that most wound infections arising from abdominal
272 procedures are mainly acquired from the patient's own faecal flora [27]. However, various traumatic and
273 surgical wounds are potentially heavily contaminated with exogenous and endogenous aerobic and
274 anaerobic bacteria derived from the disruption of mucosal surfaces [28].

275 The number of specimens collected from male patients with leg ulcer infections were much higher than
276 those from female patients (900 males compared to 600 females) and the proportions with infection in
277 each gender class were 790 (65.3%) for males and 420 (34.7%) for females. There was significant
278 correlation ($r = 0.12$) between gender and contracting wound infection. A similar result was also reported
279 in India, the difference in the number of males to females with wound infection might have been due to
280 the social behavior where males were given superiority to the females, and if contacted disease were
281 immediately taken to hospitals in comparison to delays in bringing females for treatment [19]. Age
282 prevalence in relation to number of microbial isolates among patients showed that 31-40 years had the
283 highest rate, followed by 41-50 years while the least was 0-10 years. The proportion of adults with wound
284 infection was much higher than children, and there was a moderate correlation ($r = 0.43$) between age
285 and contracting wound infection. This might have been due to the fact that more adults are involved in
286 farming, laboring jobs and more exposed to occupational hazards and therefore likely to have more
287 wounds and injuries which then provide the sites for microbial infections.

288 Conclusion

289 This work discovered a high index of bacterial involvement in lower leg ulcers in Ebonyi State, and the
290 isolates had their highest occurrence in males except for *Escherichia coli* which occurred mostly in
291 females. *Staphylococcus aureus* and *Pseudomonas aeruginosa* had the high prevalence rate. In
292 addition, the findings of this study might also guide policy makers to implement specific intervention
293 strategies to reduce the bacterial infections and their transmission.

294

295 Conflict of interest

296

297 **Competing interests:** None declared

298

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UNDER PEER REVIEW