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Bacterial community dynamics and associated genes in hydrocarbon contaminated soil during bioremediation using brewery spent grain during bioremediation using brewery spent grain

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- 13
- 14 Original Research

15 Key words

4 5 6

7 8

- 16 Bacterial community profiling
- 17 *alk*B, *cat*A, *xylE* catabolic genes
- 18 Diesel metabolites
- 19 Total Petroleum Hydrocarbons
- 20 Brewery spent grain
- 21 Bioremediation

23 Abstract

25 Brewery spent grain (BSG) has previously been exploited in bioremediation. However, detailed knowledge of 26 the associated bacterial community dynamics and changes in relevant metabolites and genes over time, is 27 limited. This study investigated the bioremediation of diesel contaminated soil amended with BSG. We observed 28 the complete degradation of three Total Petroleum Hydrocarbon (TPH C10 - C28) fractions in amended 29 treatments as compared to one fraction in the unamended, natural attenuation treatments. The biodegradation rate 30 constant was higher in amended treatments (0.1021 k) than in unamended (0.059 k) and bacterial CFUs 31 increased significantly in amended treatments. The degradation compounds observed fitted into the elucidated diesel degradation pathways and quantitative PCR results showed that the gene copy numbers of all three 32 33 associated degradation genes, alkB, catA and xylE, were significantly higher in amended treatments. High-34 throughput sequencing of 16S rRNA gene amplicons showed that amendment with BSG enriched autochthonous 35 hydrocarbon degraders. Also, community shifts of the genera Acinetobacter and Pseudomonas correlated with 36 the abundance of catabolic genes and degradation compounds observed. This study showed that these two genera are present in BSG and thus, may be associated with the enhanced biodegradation observed in amended 37

treatments. The results suggest that the combined evaluation of TPH, microbiological, metabolite and genetic analysis provides a useful wholistic approach to assessing bioremediation.

41 Data summary and availability

The DNA sequencing data generated and analysed during the current study is publicly available and has been deposited with NCBI Sequence Read Archive (SRA) with BioProject accession number PRJNA861128.
 https://www.ncbi.nlm.nih.gov/sra/PRJNA861128

47Impact Statement

49 This report summarises findings relating to enhancing the bioremediation of diesel fuel using brewery spent grain as an amendment, and the associated bacterial populations, during incubation in an organic soil. The data 50 51 provides information on the biodegradation of hydrocarbons, relates the presence of signature metabolites in the 52 degradation pathway to specific hydrocarbon degradation genes and changes in the microbial population. The 53 use of various amendments to enhance bioremediation has received increased interest in recent years and is 54 recognised as having value in industrial practice. This is enhanced when routine amendment materials such as 55 spent grain is used, rather than expensive waste treatments. Our data is novel in using the same treatment 56 samples for a combination of biochemical analysis of hydrocarbons and their degradation metabolites with the 57 genetic identification of genes and microbial communities to illuminate correlations at each time point during the 58 degradation. Intrinsic BSG bacterial populations have also been identified and profiled.

59 Declaration – This work is based on the corresponding author's publicly available doctoral thesis, which is not
 60 considered a prior publication.

61 Introduction

Environmental contamination by petrogenic hydrocarbons is ubiquitous and diesel fuel is one of the most
common pollutants of this type (Tellechea et al. 2017; Aziz et al. 2020). It is introduced to the environment
mainly by spillage during transportation and storage, thereby contaminating water and soil (Yergeau et al. 2012;
Tellechea et al. 2017). The impacts of diesel pollution go beyond environmental degradation to include health
risks for humans and other organisms due to the toxic, mutagenic and carcinogenic nature of the constituent
hydrocarbons (Souza et al. 2014; Varjani, 2017). Another concern is that hydrocarbon pollutants can accumulate
in the high trophic levels of the food chain (Semple et al. 2001, Truskewycz et al. 2019).

69 Soil is a key natural resource that is essential to sustain life on Earth. Many key functions of soil are carried out 70 by microorganisms, which also play important roles in various biogeochemical cycles (Gao et al. 2014; Jansson 71 and Hofmockel, 2018). However, soil also acts as an ultimate repository for contaminants, accumulating 72 contamination through precipitation and sedimentation from air and water. This contamination affects the soil 73 microbiota, leading to a disruption of its functions (Gao et al. 2014; Wyszkowski et al. 2020). While 74 physiochemical remediation methods such as landfilling, and incineration are available, more environmentally 75 safe, efficient and cost-effective remediation methods are sought to mitigate the impacts of soil diesel

76 contamination (Shahsavari et al. 2013).

77 Biodegradation is the natural attenuation of toxic contaminants by autochthonous microorganisms. However, the 78 process is slow, and enhancement is needed to deal with high contaminant concentrations, the associated 79 nutritional imbalances and toxicity to intrinsic microbial populations (Sarkar et al. 2016). Thus, bioremediation 80 techniques are employed under controlled conditions to enhance biodegradation. Bioremediation is an 81 exploitation of the metabolic capabilities of microorganisms to transform contaminants into innocuous, 82 mineralised products (Guarino et al. 2017). The efficiency of the biodegradation process is associated with the 83 enzymatic potential of the resident microorganisms, enabling them to digest the various substrates in the relevant 84 degradation pathways (Das and Chandran, 2011; Eze et al. 2022; Bekele et al. 2022). 85

Hence, microbial populations are the agents of biodegradation and play a vital role in the decontamination of 86 petrogenic hydrocarbons in soil (Sheppard et al. 2011, Shahsavari et al. 2013). As such, analysis of the 87 microbial community dynamics during bioremediation is essential to understand the response and adaptation of microbes to pollution and monitor the bioremediation process (Fuentes et al. 2014). More specifically, analysis 88 89 targeting the 16S rRNA gene as a molecular marker enables profiling of microbial communities and has been 90 usefully applied to characterise microbial communities involved in bioremediation of various environments 91 (Yergeau et al. 2012; Sarkar et al. 2016; Wu et al. 2020). Monitoring the presence and abundance of 92 hydrocarbon degradation genes in the degrading microbiome is also required to fully appreciate its catabolic 93 potential. This can also provide useful biomarkers for estimating the bioremediation potential of contaminated 94 sites (Yang et al. 2015; Lima-Morales et al. 2016). For example, the *alkB* gene is very important in the aerobic 95 transformation of aliphatic hydrocarbons as it encodes an alkane monooxygenase enzyme that hydrolyses 96 alkanes to their corresponding primary or secondary alcohols and has been evaluated in a number of studies 97 (Gielnik et al. 2019; Garrido-Sanz et al. 2019; Kawagoe et al. 2019).

98 Bacteria are instrumental in degrading hydrocarbon pollutants in soil and utilising the resulting metabolites for 99 energy and growth, via the tricarboxylic acid (TCA) cycle (Kulshreshtha, 2012). Metabolites associated with 100 hydrocarbon degradation have also been successfully identified in studies (Yanto and Tachibana 2013; Umar et 101 al. 2017). Following soil contamination, biodegradative bacterial strains that are resistant to the toxicity of soil 102 pollutants, can detoxify soil and make nutrients available for other quiescent populations to grow, leading to 103 changes in microbial community structure and succession (Zucchi et al. 2003; Wu et al. 2016). As a result, 104 community composition varies over time, with those species best able to exploit the metabolic breakdown 105 products dominating as bioremediation progresses (Reddy et al. 2011). Bacterial genera associated with the 106 biodegradation of petroleum hydrocarbons have been shown to include Acinetobacter, Arthrobacter, Bacillus, 107 Flavobacterium, Nocardia, Pseudomonas and Vibrio (Chandra et al. 2013; Varjani, 2017). Among the different 108 species associated with bioremediation, Pseudomonas spp. in particular are known to be versatile in the 109 biodegradation of hydrocarbons, especially those found in diesel fuel (Dussán and Numpaque, 2012). However, 110 knowledge of the microbial species involved in diesel remediation using food wastes in temperate countries is 111 limited.

Shahsavari et al. (2013) observed high degradation rates in diesel and gasoline contaminated soil amended with crop residues. As soil is the most expensive medium to decontaminate, the use of crop residues and food waste is cost efficient and advantageous (Agamuthu et al. 2013). Brewery spent grain is a readily available food byproduct with high nutritional content (Thomas and Rahman, 2006; Lynch et al. 2016). It has been successfully

used to stimulate bioremediation and its effluents have been used for diesel bioremediation, indicating strong

potential (Agarry and Latinwo, 2015). However, these uses have been mostly under tropical conditions

118 (Agamuthu et al. 2010; Abioye et al. 2012). BSG was chosen as a biostimulant in this study because, as well as

releasing nutrients for autochthonous bacteria during biodegradation, it is known to have its own resident
 microflora and so may provide both biostimulation and bioaugmentation benefits (Robertson et al. 2010). It

would also allow reuse of an industrial by-product that might otherwise be put to landfill as waste (Bianco et al.

121 would also allow reuse of all industrial by-product that hight otherwise be put to failurin as waste (122 2020).

123 Previous work on diesel biodegradation using BSG has tended to focus on the reduction of TPH and culturable 124 bacterial count. Changes in the associated catabolic genes, metabolites, and community succession over time 125 during diesel bioremediation using BSG in a temperate soil have not been investigated. This time-course study 126 was designed to illuminate any correlations between breakdown metabolites, gene abundance and community 127 shifts in response to biodegradation, with the aims of evaluating the impact of BSG supplementation on diesel 128 bioremediation in soil and answering the following questions. How does BSG influence biodegradation? What 129 are the metabolites associated with this biodegradation process and how do they fit into known hydrocarbon 130 breakdown pathways? Which bacterial species are associated with biodegradation, how does their abundance 131 change over time and does BSG supplementation favour known hydrocarbon-degrading bacteria and catabolic 132 genes?

133

134 **2.** Methods 135

136 2.1 Soil preparation and experimental design

137 Soil was collected (10-30 cm depth) from an uncontaminated pristine agricultural farmland using sterile 138 implements, air dried and sieved using a 4 mm mesh (Shahsavari et al. 2013). The pH of the soil was determined 139 to be 6.8 after a 1:2.5 soil: distilled water dilution (Emmett et al. 2010). Soil characteristics were determined (by 140 Anglian Soil Analysis Ltd, UK) as listed in Table 1. The three treatments used in this study were soil + diesel 141 (S+D) unamended natural attenuation, soil + diesel + BSG (S+D+G) amendment, and soil alone (S) control. In 142 triplicate, 20 ml of diesel was spiked into 2000 g soil to achieve a 1% (v/w) contamination. To ensure 143 homogeneity the diesel was added to 25 % of the soil and mixed thoroughly with a stainless-steel implement 144 before adding the remaining soil (Fernández et al. 2011). BSG (10%) was then added to treatments as required 145 and mixed in thoroughly. BSG had a moisture content of 75%, a pH of 5.3 and was stored at 4 °C for four days 146 prior to use.

148	Soil property	Value
149	Soil texture	Sandy/Loam
150	рН	6.8
151	Moisture (%)	26.04
152	Organic matter (%)	12.0
153	Sand (%)	50.44
154	Silt (%)	41.84
155	Clay (%)	7.72
156	Total organic carbon (%)	6.96
157	Potassium (mg kg ⁻¹)	15.4
158	Magnesium (mg kg⁻¹)	21.1
159	Phosphate (mg kg ⁻¹)	17.5
160	Nitrate (mg kg ^{−1})	3.5

147 Table 1 Physical and chemical characteristic of the soil used in this study

161

Sterile counterparts of treatments (sterile soil + diesel, and sterile soil + diesel + sterile BSG) were also analysed 162 163 to confirm biotic loss. Sterilisation was achieved prior to diesel spiking by autoclaving for one hour at 121 °C on three alternate days (Molina-Barahona et al. 2004; Emam et al. 2014). For each treatment, 2000 g soil was 164 165 incubated in triplicate 5 litre pots. Treatment pots were covered with Gore-Tex cloth and incubated at 15 °C ± 3 166 °C. Treatments were oxygenated by mixing twice a week using a sterile spatula and the moisture content was maintained by the weekly addition of 5 % (v/w) sterile distilled water. Composite samples for analysis were 167 168 obtained from each pot on days 0, 2, 5, 7, 12, 14, 21, and 28 by collecting 5 g samples from the four corners and 169 the centre and mixing together (Chagas-Spinelli et al. 2012).

170 2.2 Determination of TPH removal and detection of hydrocarbon metabolites

171 2.2.1 GC Analysis

172 TPH was determined using a modified US EPA 8015 technique (Bento et al. 2005; Suja et al. 2014). Hexane was 173 used for diesel extraction with mechanical shaking (Geilnik et al. 2019). Aliquots of the extract (1 ml) were 174 transferred in triplicate to 1.5 ml gas chromatography vials and GC analysis was carried out using an Agilent 175 Technologies 7890A system equipped with a flame ionization detector (FID) and autosampler (7693). A 30 m x 176 0.32 mm x 0.25 µm capillary column (19091J-413E HP-5, Agilent Technologies, UK) was used with helium as 177 the carrier gas at a flow rate of 2 ml min⁻¹, hydrogen gas at a flow rate of 30 ml min⁻¹ and air at a flow rate of 300 178 ml min^{-1.} The temperature program used was a modified version of that given by Bento et al. (2005). The initial 179 temperature was 50 °C with isothermal operation for 5 min, followed by heating to 270 °C at a constant rate of 10 °C min⁻¹ and a final 5 min isothermal operation. Samples from each time point and triplicate standards were 180 181 analysed on the same run.

- 182 2.2.2 Percentage TPH reduction, extent of aliphatic TPH (C10-C28) fractions removal and biodegradation
 183 rate
- 184 Percentage TPH reduction was calculated using the formula:
- 185 % TPH reduction = [(TPH of control TPH treatment) / TPH control] x 100, with day 0 TPH being used as
- a control for each treatment (Bento et al. (2005). The TPH Standard Mix 1 (Sigma Aldrich, UK), with known
- concentration for each of the (C10 C28) analytes, was used to obtain the calibration curve for each fraction.
 Concentrations of each fraction in the soil hydrocarbon mixture were then determined based on the calibration
- 189 curve of each corresponding standard fraction. Retention times of each fraction (analyte) in the standard were
- 190 compared to the sample chromatogram to determine target compounds and the total peak areas of both standard
- 191 and analyte fractions were determined. The biodegradation rate was determined using the formula $\frac{c}{c_0} = e^{-k^t}$
- (Baek, et al. 2004), which is same as: $C = C_0 e^{-k^t}$, (Abioye et al. 2012); where C is the concentration of the TPH fractions (mg kg⁻¹) at time t, C₀ is the initial concentration of the TPH fractions (mg kg⁻¹), t is time (day ⁻¹) and k is the biodegradation rate constant (day⁻¹).
- 196 2.2.3 GS-MS Analysis
- 197 GC-MS analysis was carried out using an Agilent GC-MS 7890A/5975C series instrument with a 30 m x 0.32 198 mm x 0.25 µm capillary column (19091S-433E HP-5MS, Agilent Technologies, UK). Helium was the carrier 199 gas with a flow velocity of 1 ml min⁻¹ and I μ l of sample was injected into the column in a splitless mode. The 200 analytical conditions were an initial temperature of 50 °C, with isothermal operation for 1 min followed by heating to 120 °C at a constant rate of 20 °C min⁻¹ and a final heating to 310 °C at a constant rate of 4 °C min⁻¹ 201 202 (Xu and Lu, 2010) with a 5 min isothermal operation. The column was directly connected to an electron 203 ionisation mass spectrometer with an electron energy of 70 eV, producing ions that are characterised according 204 to mass-to-charge ratio and relative abundance. 205
- 206 2.2.4 Detection of hydrocarbon degradation metabolites and BSG metabolic potential

Following the GC-MS analysis to identify compounds present in the treatments at each time point, the G3835AA
Mass Hunter Mass Profiler Professional Software (Agilent Technologies, UK) was used to analyse the MS data
(identified compounds) and the identified metabolites were used to determine the oxidative pathways utilised in
the breakdown of diesel with and without BSG. The abundance of each compound in the treatments, and
compounds differing significantly between the treatments overall, were determined.

- 2.3 Determination of colony forming units (CFUs) for the enumeration of heterotrophic and hydrocarbon
 degrading bacteria
- Composite samples of 10 g from each treatment pot were transferred to sterile bottles containing 100 ml of 0.2 %
 (v/v) sterilised sodium pyrophosphate and mixed on a shaker at 150 rpm for 30 minutes at 20 °C. Thereafter, 1
 ml of the soil suspension from each bottle was 10-fold serially diluted in sterile saline to give dilutions 10⁻¹ to 10⁻⁶. Following dilution, 0.1 ml of each suspension was plated onto R2A agar for enumeration of heterotrophic
 bacteria, and onto oil agar for enumeration of hydrocarbon-degrading bacteria. These media were incubated at 30
 °C for 24 hours and 25 °C for 7 days, respectively.
- 222 223
- 224 2.4 Quantification of catabolic genes and bacterial community composition profiling

225 2.4.1 DNA extraction from treatments

Microbial community DNA was extracted from bioremediation treatment samples (1 g), with and without BSG
at each time point (days 0, 5, 12 and 21), using the EZNZA soil DNA kit (Omega Bio-Tek, Inc., USA) (Dineen
et al. 2010). Purity of extracted DNA was estimated by measuring absorbance at 260nm and 280nm, using a
micro volume spectrophotometer (Nanodrop technologies, USA) and calculating the 260/280 ratio, which was
required to be between 1.8 to 2.0.

231 2.4.2 qPCR quantification of diesel catabolic genes

232 Three hydrocarbon catabolic genes, alkB (Powel et al. 2006), catA and xylE (Shahsavari et al. 2016), were 233 assayed quantitatively using PCR primers as listed in Table 2. Quantification was performed by real time PCR 234 (qPCR) using 1:10 dilutions of the extracted community DNA in sterile nuclease-free water (Shahsavari et al. 235 2016). The assays were performed in a Rotor Gene Q thermocycler (Qiagen, UK) using the 2x Kapa Sybr® Fast 236 qPCR Master Mix Universal kit (Sigma Aldrich, UK). Each reaction (20 µl) contained 2x Kapa Sybr Fast qPCR Master Mix (10 µl), forward primer (0.2 µM), reverse primer (0.2 µM), PCR-grade water (8.2 µl) and DNA 237 238 template (1 µl). The amplification programme for the *alkB* gene included initial denaturation at 95°C (5 mins), followed by 40 cycles of denaturation at 95°C (10 s), annealing at 50°C (30 s), extension at 72°C for 30 seconds 239 240 and primer-dimer removal and signal acquisition at 80°C for 10 seconds. Thermal cycling programmes for both 241 *catA* and *xylE* genes comprised an initial denaturation step at 95° C (5 mins), followed by 40 cycles of 242 denaturation at 95°C (10 s), annealing at 58°C (30 s), extension at 72°C (30 s), with primer-dimer removal and 243 signal acquisition at 80°C for 10 seconds. Reactions were run in triplicate and negative controls (PCR-grade

244 water) were included in all amplifications.

245 In an initial experiment to evaluate the methodology (Nnadi, 2019), PCR amplicons from genes of interest in the 246 amended treatment were verified to be of correct band size by electrophoresis on a 2% (w/v) agarose gel. Bands 247 were visualised using a Chemi Doc™ MP Imaging System (Bio-Rad Laboratories Ltd. Hertfordshire, UK), 248 excised under UV radiation and extracted using the QIAquick Gel Extraction Kit (QIAGEN, USA). Following 249 confirmation of their DNA sequences, the cleaned PCR products were used as positive controls for the standard 250 curve. CT values of the treatments were related to the standard curve. Gene copy numbers were calculated using the formula: number of copies = $(ng/ul DNA \times 6.022 \times 10^{23}) / (PCR product length in base pairs x 1 x 10⁹ x 650) /$ 251 (Staroscik, 2004) where Avogadro's number of 6.022×10^{23} is the number of molecules/mole DNA and 650 Da is 252 253 the average weight of a base pair. Results were expressed as \log_{10} of gene copy numbers per g dry soil (log 10^g) 254 ¹).

255 Table 2 Details of the primers used for the detection and quantification of target hydrocarbon catabolic genes

256

Target gene	Primer name	Annealing temperature °C	Sequence (5' – 3')	Amplicon size	References
Alkane monoxygenase	alkB	50	F: AAC TAC ATC GAG CAC TAC GG R: TGA AGA TGT GGT TGC TGT TCC	100 bp	Powell et al. 2006
Catechol-1,2- dioxygenase	catA (C12O)	58	F: ACVCCVCGHACCATYGAAGG R: CGSGTNGCAWANGCAAAGT	470 bp	Sei et al. 1999 Shahsavari et al. 2016
Catechol-2,3- dioxygenase	<i>xylE</i> (C23O)	58	F: AAGAGGCATGGGGGGCGCACCGGTTCGATCA R: CCAGCAAACACCTCGTTGCGGTTGCC	380 bp	Azhari et al. 2010 Shahsavari et al. 2016

²⁵⁷

258 2.4.3 High-throughput sequencing of 16SrRNA gene amplicons for bacterial community profiling

259 Samples of microbial community DNA extracted from bioremediation treatments with and without BSG at each

time point (days 0, 5, 12 and 21), as used for catabolic gene quantification, were also used for community

profiling via analysis of the 16S rRNA gene. V4 variable region PCR primers 515 – 806 (Caporaso et al. 2011)

were used in a single-step 30 cycle PCR using the HotStarTaq Plus Master Mix Kit (Qiagen, USA) under the

following conditions: 94°C (3 mins), followed by 28 cycles (5 cycle used on PCR products) of 94°C (30s), 53°C

for (40 s) and 72°C (1 min), after which a final elongation step at 72°C (5 mins) was performed.

Sequencing was performed at MR DNA, Shallowater, TX, USA using an Ion Torrent PGM system. Sequence
 data were processed using a proprietary analysis pipeline (MR DNA, Shallowater, TX, USA). In summary,

- sequences were depleted of barcodes and primers, then sequences < 150bp, with ambiguous base calls and with
 homopolymer runs exceeding 6bp were removed. Sequences were denoised, OTUs generated, and chimeras
- removed. Operational taxonomic units (OTUs) were defined by clustering at 3% divergence (97%
- similarity). Final OTUs were taxonomically classified using BLAST against a curated database derived from
- 271 Green Genes, RDPII and NCBI (<u>www.ncbi.nlm.nih.gov</u>, DeSantis et al. (2006), <u>http://rdp.cme.msu.edu</u>). Data
- was submitted to the Sequence Read Archive (SRA) at NCBI with the BioProject accession number
- 273 PRJNA861128. https://www.ncbi.nlm.nih.gov/sra/PRJNA861128
- 274 2.5 Statistical Analysis
- 275

IBM SPSS 24 statistics software was used to determine significant differences among treatments in respect to
TPH reduction, microbial count, and catabolic gene copy numbers. All data were tested for normality using the
Shapiro-Wilk test. Analysis of variance (ANOVA) and the least significant difference (LSD) *post hoc* test were
used to determine significant differences among treatment means of more than two independent variables, while
the independent T test was used to determine significant differences between two independent variables.

- 281 Significance was determined as p < 0.05.
- 282

283 3 Results

- 284 3.1. Determination of TPH removal and detection of hydrocarbon metabolites
- 285 3.1.1 Percentage TPH reduction

The natural attenuation treatment (S+D) and the treatment amended with BSG (S+D+G) produced a rapid reduction in TPH of 78% and 84% respectively by day two (Table 3). Further reductions of 92% in the natural attenuation, and 96% in the amended treatments were observed after one week of incubation. After 28 days of incubation, a final 99% reduction was observed in the amended treatment and a 93% reduction in the natural attenuation treatment. Statistical analysis revealed a significant difference in TPH total peak heights between the two treatments throughout the experiment except at day 0. The treatment amended with BSG showed a significantly higher percentage TPH reduction compared to the natural attenuation, unamended treatment.

- 293
- 294

Table 3 Percentage reduction of TPH during bioremediation of diesel contaminated soil treatments with
 (S+D+G) and without (S+D) BSG

			297				
	% TPH Reduction in treatments						
Days	S+D	S+D+G	299				
2	78	84					
5	89	92					
7	92	96					
12	92	97					
14	92	97					
21	93	98					
28	93	99					

300 301 302

3.1.2 Extent of aliphatic TPH (C10-C28) fractions removal and biodegradation rate

303 The extent of aliphatic TPH fractions (C10-C28) removal in both natural attenuation (S+D) and BSG amended 304 treatments (S+D+G) was evidenced by the reduced concentrations of the TPH fractions over time, during the 305 bioremediation (Fig 1). The results show a 100% decline of the C10 fraction by day 12 in both treatments. This 306 was the only hydrocarbon fraction to be fully removed in the natural attenuation treatment. However, a complete 307 decline of the C10, C12 and C28 fractions was observed in the amended treatment by day 28. These three fractions had initial concentrations of 700 mg kg⁻¹, 2400 mg kg⁻¹ and 500 mg kg⁻¹ respectively at the start of the 308 309 incubation. The concentrations of the C14 to C26 fractions reduced over time but persisted until day 21 in both 310 treatments. However, the reduction of these fractions was significantly greater in the amended treatment than in 311 the natural attenuation treatment. The biodegradation reaction rate constant (k) was significantly higher in the 312 amended treatment (0.1021 day⁻¹) compared to the natural attenuation treatment (0.0590 day⁻¹). The first order 313 linear model r values were 0.8699 and 0.9585 respectively for these two treatments.

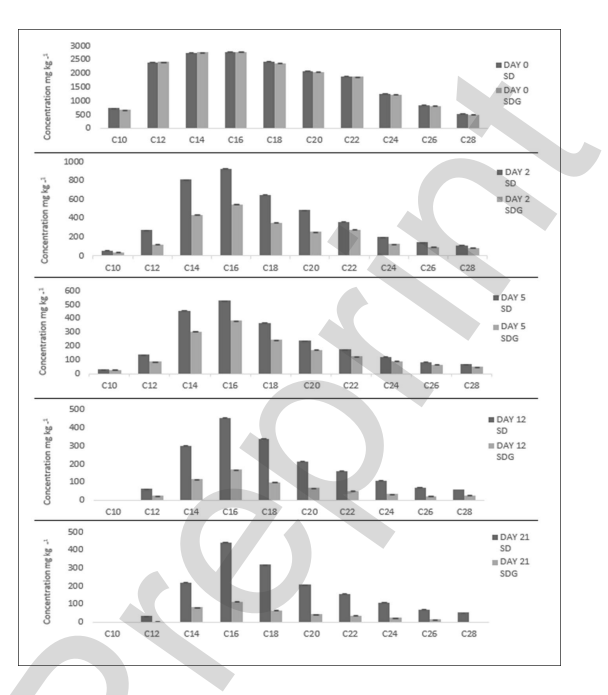


Fig 1 Extent of aliphatic TPH (C10 - C28) fractions removal, with and without BSG, over time, during diesel biodegradation. Results represent the means of three replicates. Error bars show standard error. SD = soil + diesel, SDG = soil + diesel + BSG amendment.

323 3.1.3 Detection of hydrocarbon degradation metabolites and BSG metabolic potential

Compounds detected by GC-MS from the treatments (S+D and S+D+G) at different stages during the

bioremediation, were assessed to determine the presence of known metabolites in the degradation pathways of
 both aliphatic and aromatic hydrocarbons, based on their functional groups such as alcohols, ketones, esters,
 aldehydes, carboxylic acids, and esters.

328 Results for aliphatic hydrocarbon degradation (Table 4) revealed that ketones, being metabolites of the

subterminal oxidation pathway, were present in both unamended (S+D) and amended (S+D+G) treatments,

- throughout the bioremediation process, while aldehydes, being metabolites of the terminal oxidation pathway,
- were absent. However, carboxylic acids, which are further metabolites of the terminal oxidation pathway,
- resulting from aldehyde oxidation (van Elsas 2007; Chikere et al. 2011; Varjani, 2017), were present at the startand up until day 5 in the unamended treatment but were only present at the start in the amended treatment. For
- aromatic hydrocarbons however, aromatic ketones, which are oxidation products of the *ortho*-cleavage pathway,
- were present in both treatments throughout the bioremediation. Aromatic aldehydes, which are oxidation
- products of the *meta*-cleavage pathway, were only observed from the start of the experiment on days 0 and 5 in
- unamended treatments but were present on days 5, 12 and 21 in the amended treatments.

338 Table 4 Hydrocarbon degradation metabolic compounds present in treatments, over time during bioremediation, including diesel control

 \bigcirc

339

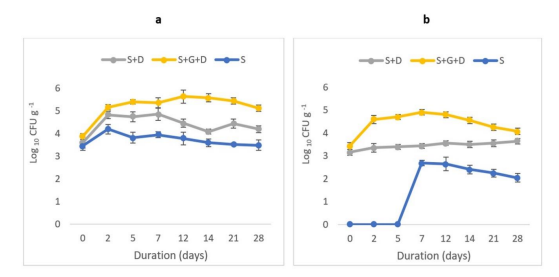
	TREATMENTS										
				D	ay O	D	ay 5	Da	ay 12	Da	ay 21
				SOIL	SOIL	SOIL	SOIL	SOIL	SOIL	SOIL	SOIL
HYDROCARBON	FUNCTIONAL	TYPE OF	DIESEL	AND	DIESEL	AND	DIESEL	AND	DIESEL	AND	DIESEL
ТҮРЕ	GROUP	COMPOUND	CONTROL	DIESEL	AND BSG	DIESEL	AND BSG	DIESEL	AND BSG	DIESEL	AND BS
ALIPHATIC	C-H	STRAIGHT CHAIN	+	+	+	+	+	+	+	+	+
HYDROCARBONS	–OH	ALCOHOLS		+	+	+	+	+	+	+	+
	-H-C=O	ALDEHYDES		-	-	-	-	-	-	-	-
	-R-C=O	KETONES		+	+	+	+	+	+	+	+
	OHC=O	CARBOXYLIC ACIDS		+	+	+	-	-	-	-	-
	-0-C=0	FATTY ACID ESTER		+	+	+	+	+	+	+	+
	\bigcirc										
AROMATIC	\mathbf{i}	BENZENE RINGED	+	+	+	+	+	+	+	+	+
HYDROCARBONS	–OH	CATECHOL (Alcohol)	+	+	+	+	+	+	+	+	+
	-H-C=O	ALDEHYDES		+		+	+	-	+	-	+
	-R-C=O	KETONES		+	+	+	+	+	+	+	+
	OHC=O	CARBOXYLIC ACIDS		+	+	+	+	+	+	-	-
	-O-C=O	FATTY ACID ESTER		+	+	+	+	+	+	+	+

340 3.2 Determination of colony forming units (CFUs) for the enumeration of heterotrophic and hydrocarbon
 341 degrading bacteria

Changes in heterotrophic bacterial CFUs are shown in Fig 2a. CFUs in the unamended diesel contaminated soil
samples (S+D) increased at the start of the experiment and peaked at 5.0 log 10 g⁻¹ on day 7 after which they
decreased continually. The CFUs in the diesel contaminated samples amended with BSG (S+D+G), increased
continually from day 0 and peaked at 5.6 log 10 g⁻¹ on days 12 and 14 before decreasing gradually. However,
the heterotrophic bacterial CFUs remained significantly higher in the amended treatment (S+D+G). The soil
control treatment (S) also showed increased CFUs after day 0 but peaked at 4.2 log 10 g⁻¹ on day 2. This

treatment had the lowest CFUs.

349 The soil control treatment (S) had no hydrocarbon utilising bacterial growth until day 7 after which the 350 population slowly declined. Hydrocarbonoclastic CFUs in the unamended treatment (S+D) remained similar 351 throughout the experiment (Fig 2b), with a 6% increase on day 12 when it peaked at 3.6 log 10 g⁻¹. The CFUs in 352 the amended treatment (S+D+G) had the highest CFUs and showed a rapid CFU increase of 34% on day 2 as compared to the level at day 0. It then peaked at 4.9 log 10 g⁻¹ on day 7 with a 43% increase compared to the 353 354 unamended treatment (S+D). After this time, a gradual decrease in CFU's occurred until day 28 at which point a 355 12% increase in CFUs was seen as compared to the unamended treatment. A negative correlation was observed 356 between TPH concentration and hydrocarbon degrading CFUs in both unamended (r = -0.858) and amended (r = 357 -0.926) treatments.



358

Fig 2 Mean colony forming units (CFU's) for the enumeration of heterotrophic bacteria (a) and hydrocarbon degrading bacteria (b) in treatments with and without BSG. Results represent the means of three replicates.
Error bars show standard error. S+D = soil + diesel, S+D+G = soil + diesel + gain amendment, S = control soil.

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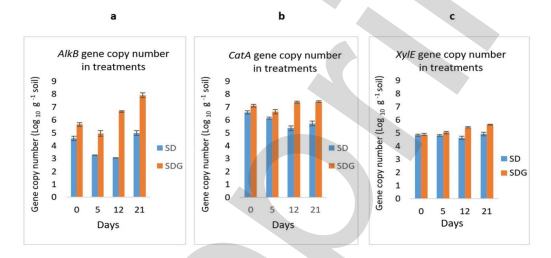
363 3.3 Quantification of diesel catabolic genes and bacterial community composition.364

365 3.3.1 Quantification of diesel catabolic genes by qPCR

366 Gene copy number quantification in this study was limited to the contaminated soils, with and without BSG, to 367 determine the bioremediation potential of BSG. The gene copy numbers of all three catabolic genes in this 368 study, increased with the addition of BSG. From day 5 until the end of the experiment, the alkB gene copy 369 numbers (Fig 3a) in the amended treatment (S+D+G) were significantly higher than that of the unamended 370 treatment (S+D). Also, a very distinct difference in gene copy numbers was observed in the amended treatment 371 on day 12 while that in the unamended treatment remained constant. Nevertheless, there was an increase in gene 372 copy numbers in both treatments after this time. From the start to the end of the experiment, gene copy numbers in the unamended treatment (S+D) increased by 22.8% from 4.57 $\log_{10} g^{-1}$ to 5.61 $\log_{10} g^{-1}$ while that in the amended treatment (S+D+G) increased by 40.4% from 5.65 $\log_{10} g^{-1}$ to 7.93 $\log_{10} g^{-1}$. 373 374 375

376 A significant difference in the catA gene copy numbers was observed (Fig 3b), between the amended (S+D+G) 377 and unamended (S+D) treatments from the start and throughout the experiment. The rapid reduction in 378 percentage TPH and concentration of aliphatic hydrocarbon (C10-C28) fractions also occurred at the start of 379 the experiment and was associated with a negative correlation in hydrocarbon degrading bacterial CFUs. 380 However, like the alkB gene, a distinct difference in gene copy numbers was evident between the amended and 381 unamended treatments from day 12, after which it plateaued until the end of the experiment. Nevertheless, a 13.2% decrease in the *catA* gene copy numbers, from 6.59 $\log_{10} g^{-1}$ to 5.72 $\log_{10} g^{-1}$, was observed in the natural 382 attenuation treatment S+D, from day 0 to the end of the experiment. In the amended treatment, S+D+G, catA 383 384 gene copy numbers increased by 4.7% from 7.10 $\log_{10} g^{-1}$ to 7.43 $\log_{10} g^{-1}$.

Gene copy numbers for the *xylE* gene (Fig 3c), like the *alkB* gene, were significantly higher in the amended treatment from day 5 and throughout the experiment. However, as with both the *alkB* and *catA* genes, a distint difference in gene copy numbers was evident from day 12. A 1.9% increase in copy number of the *xylE* genes was observed in the naturally amended treatment, S+D, from day 0 to the end of the experiment increasing from 4.84 log₁₀ g⁻¹ to 4.93 log₁₀ g⁻¹ while a 15.4% increase was observed in the amended treatment, S+D+G, from 4.89 log₁₀ g⁻¹ to 5.64% log₁₀ g⁻¹.



³⁹¹

Fig 3 Gene copy numbers of the *alkB* gene (**a**), *catA* gene (**b**) and *xylE* gene (**c**) in treatments with and without BSG over time, during the bioremediation of diesel contaminated soil. Results represent the means of three replicates. Error bars show standard error. S+D = soil + diesel, S+D+G = soil + diesel + gain amendment.

395

396 3.3.2 High-throughput Sequencing of 16S rRNA gene amplicons for community profiling

397 Results monitoring the bacterial community changes and dynamics during the transformation of aliphatic and 398 aromatic hydrocarbons in the diesel contaminated soil during bioremediation, were profiled in terms of the 399 relative abundance (Fig 4) and percentage abundance (Fig 5) of the twelve most dominant bacterial populations 400 in the treatments. Aliquots from the same community genomic DNA samples, with and without BSG, used in 401 the qPCR assay for the three catabolic genes in this study, were used for this analysis for comparability with 402 each other and with the control soil alone and BSG alone samples. DNA yield from contaminated samples (ug) 403 were all high and the purity based on absorbance at 260nm and 280nm using a Nanodrop (ND-2000, Thermo 404 Fisher Scientific, UK) were between 1.8 to 2.0 as expected. Diesel contamination did not appear to impact 405 genomic DNA recovery from the soil.

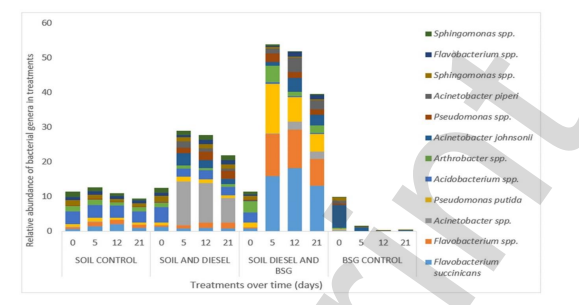
406 The results in Fig 4, show that the treatment amended with BSG had the highest relative abundance of bacterial populations. These results are consistent with those from culture-based methods in which treatments amended
408 with BSG supported the highest heterotrophic and hydrocarbonoclastic bacterial populations. However, the
409 bacterial population on day 0 were similar for all treatments except that of the grain control. The grain control, however, did not continually sustain microbial growth on its own.

411 As shown in Fig 5, we observed in the soil alone control treatment (S), that the percentage abundance of

412 Flavobacterium succinicans increased from around 7% on day 0 to 20% on day 12. Thereafter, it reduced to

413 11% on day 21. Culturable hydrocarbon degrading bacteria were first observed on day 7 in this treatment. Apart

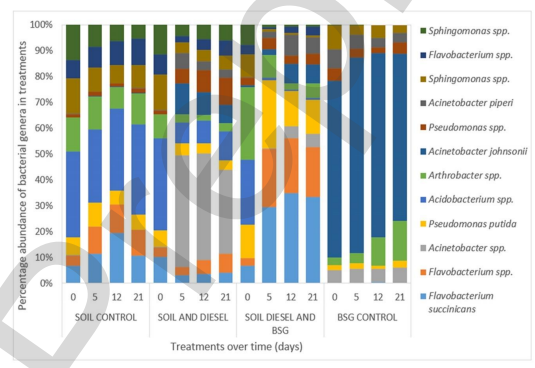
414 from these changes, no significant shifts in the percentage composition of other bacterial genera in this treatment 415 were observed.





417 Fig 4 Relative abundance of bacterial genera present in samples showing the bacterial population profile of
 418 treatments during diesel bioremediation over time.

419 420



421 422

422 Fig 5 Bacterial community profile showing percentage abundance of bacterial populations during diesel423 bioremediation.

424 In the BSG alone control treatment, the percentage abundance of *Arthrobacter* spp. increased from 2% to 15%

from the start of the experiment up till day 21 (Fig 5). Slight decreases in the abundance of *Pseudomonas* spp.,

426 *Acinetobacter piperi* and *Sphingomonas* spp. were also observed. As with the control soil alone treatment, there 427 were no significant changes in the percentage composition of other bacterial populations in this treatment with

428 *Acinetobacter johnsoni* being the most abundant species in this treatment throughout the experiment.

429 In contrast, the percentage composition of bacterial populations in the diesel contaminated soil treatments

430 changed extensively. A rapid shift in bacterial community composition was observed in the unamended soil and

431 diesel treatment after day 0. The population profile revealed three *Acinetobacter* species in the treatment by day
432 5 namely: *Acinetobacter* species 45%, *Acinetobacter johnsonii* 12%, and *Acinetobacter piperi* 5% while

433 Acidobacterium spp. reduced. Changes in species abundance were observed hereafter though no more shifts in

434 community composition were seen. After day 5 the three *Acinetobacter* species gradually reduced while two

435 Flavobacterium species increased. Acinetobacter spp. were the most abundant species in this treatment though it

reduced from 45% to 41% and 32% on days 5, 12 and 21, respectively. *Acinetobacter johnsonii*, reduced from

437 12% to 8% and 5% on days 5, 12 and 21 respectively and *Acinetobacter piperi* reduced from 5% to 3% and
438 2.5% on days 5, 12 and 21. *Flavobacterium* spp., however, increased from 3% to 5% and finally to 9% on day

438 2.5% on days 5, 12 and 21. *Flavobacterium* spp., however, increased from 3% to 5% and finally to 9% on days
439 5, 12 and 21 while *Flavobacterium succinicans* increased from 2% to 3% and 5% on the same days.

440 *Pseudomonas* spp. also increased from 5% to 8% and finally to 10%.

441 Addition of BSG to the contaminated soil on day 0 may have contributed to the increased percentage abundance 442 of the Arthrobacter spp. and Pseudomonas putida observed in the amended treatment (S+D+G) compared to the 443 unamended soil and diesel (S+D), and soil control (S) treatments (Fig 5). A rapid shift in community 444 composition was observed in this treatment by day 5, resulting in the increased abundance of Flavobacterium 445 succinicans, Flavobacterium spp. and Pseudomonas putida. Thereafter, this treatment sustained these three 446 species as the most abundant throughout the experiment. At the same time, Arthrobacter spp. declined and 447 Acidobacterium spp. were almost eliminated. As with the unamended treatment, a reduction in the abundance of 448 Acidobacterium spp. following contamination was also observed in the amended treatment. Acinetobacter piperi 449 was also observed in this treatment after day 0. After day 5, a second shift in bacterial community composition 450 was observed in the amended treatment during which Acinetobacter spp. was seen and the abundance of two 451 Acinetobacter species increased. Acinetobacter johnsonii increased from 2% to 8% and 9% and Acinetobacter 452 piperi increased from 4% to 9% and 8% on days 5, 12 and 21 respectively. At the same time, however, the 453 abundance of Pseudomonas putida, reduced from 28% to 14%.

454

455 Discussion

456 The enhancement of the bioremediation process by stimulating with BSG was evident by the significant 457 reduction in percentage TPH and high biodegradation rates observed. Similarly, Agarry and Latinwo (2015) 458 reported a high rate of 79% TPH reduction after 28 days when using brewery spent effluent for the 459 bioremediation of a 10% (w/w) diesel contaminated soil. The potential of BSG to enhance the bioremediation 460 process was also made obvious by the significant increase in both heterotrophic and hydrocarbonoclastic 461 bacterial populations, along with a negative correlation between petroleum hydrocarbon degradation and 462 bacterial CFUs (Margesin et al. 2003; Shahsavari et al. 2013). The CFUs observed in the soil only control reveal 463 that the soil has got indigenous hydrocarbon degraders which began utilising diesel as their only carbon source 464 after a lag phase of about one week. Similarly in their study, Alisi et al (2009) reported that in soils without prior 465 hydrocarbon contamination, a 2-day lag phase was observed before CO2 evolution and 6 days were required for 466 the soil microbial community to become active.

467

468 The initial rapid TPH reduction observed in this study was contributed to by all the aliphatic TPH (C10 - C28) 469 hydrocarbon fractions. However, although the C10, C12 and C28 aliphatic fractions in the amended treatment 470 were completely degraded, the C12 fraction, having a lower molecular weight despite its higher initial 471 concentration, degraded faster than the C28 fraction. Thus, it is likely that the most labile and low molecular 472 weight fractions were metabolised in the initial rapid phase while the more resistant and higher molecular 473 weight fractions were degraded in a later, second phase (Ros et al. 2010; Karamalidis et al. 2010; Varjani and 474 Upasani, 2017). The initial rapid degradation in both the natural attenuation and amended treatments may also 475 have resulted from autochthonous hydrocarbon degraders being present in the soil (Shahsavari et al. 2013). 476 Although the soil was pristine, hydrocarbonoclastic microbes are known to be ubiquitous and following 477 contamination, it has been observed that microbial communities in pristine soils adapt well to contaminants 478 resulting in rapid degradation (Kingston, 2002; Margesin et al. 2013; Schwarz et al. 2018).

The degradation pathways of aliphatic hydrocarbons show that, depending on the position of the methyl group
initially attacked by the oxygenase enzyme, alkanes may be oxidised to either primary or secondary alcohols
(van Elsas et al. 2007). Further oxidation of primary alcohols produce aldehydes and fatty acids while secondary

(van Elsas et al. 2007). Further oxidation of primary alcohols produce aldehydes and fatty acids while secondary
 alcohols produce ketones and esters (Varjani, 2017). The degradation pathways of aromatic hydrocarbons,

483 however, indicate that they are initially oxidised to catechol (Benzene-1, 2-diol) (Das and Chandran 2011; Eze

484 et al. 2021). Thereafter, cleavage of the benzene ring occurs in either of two routes: The *ortho*-cleavage

485 pathway, which involves cleavage between carbons 1 and 2 catalysed by catechol-1, 2-dioxygenase to produce

ketones and esters, and the *meta*-cleavage pathway, which involves cleavage between carbons 2 and 3 by

487 catechol-2, 3-dioxygenase to produce aldehydes and carboxylic acids (Chikere et al. 2011; Olajire and Essien,
488 2014). From the elucidated pathways, and as observed during this study, ketones and aldehydes are key

489 distinguishing metabolites in determining the catabolic pathways utilised by the bacterial community.

490 Aldehydes being the distinguishing metabolite for the terminal oxidation pathway of aliphatics and *meta*-

491 oxidation pathway of aromatics (Okoh 2006; van Elsas 2007; Olajire and Essien 2014). Ketones on the other

hand, distinguish the activity of the subterminal oxidation pathway of aliphatics and the *ortho*-oxidation
pathway of aromatics (van Elsas 2007; Varjani et al. 2017).

494 According to Tsugawa et al. (2011), to deduce the metabolic activity of microbes associated with hydrocarbon 495 biodegradation, an analysis of metabolites in the biological samples is required. Thus, to deduce the metabolic 496 pathways in this study, we identified ketones and aldehydes from the literature (Varjani 2017) as the signature 497 metabolites that differentiate between the terminal and subterminal pathways in aliphatic hydrocarbon 498 degradation catalysed by alkane monoxygenase and the ortho and meta degradation pathways of aromatic 499 hydrocarbon degradation catalysed by catechol-1, 2-dioxygenase and catechol-2, 3-dioxygenase respectively 500 (Varjani and Upasani, 2017; Fuchs et al. 2011). Aliphatic hydrocarbons may have been degraded mainly via the 501 subterminal oxidative pathway as suggested by the presence of aliphatic ketones and esters, in both the natural 502 attenuation and amended treatments, throughout the biodegradation process (Table 4). However, the presence of 503 aliphatic fatty acids (carboxylic acids), though aliphatic aldehydes were not detected, suggests that a rapid 504 oxidation of aldehydes to carboxylic acids may have occurred, particularly in the BSG amended treatments. The 505 carboxylic acids may then have been oxidised via the β -oxidation pathway to the tricarboxylic acid (TCA) 506 cvcle.

507 For aromatic hydrocarbon degradation however, aromatic ketones were present throughout the degradation 508 indicating that degradation was mostly via the ortho-cleavage pathway catalysed by catechol 1, 2-dioxygenase 509 (Fuchs et al. 2011). Aromatic aldehydes, which are the key differentiating metabolites of the meta-cleavage 510 pathway catalysed by catechol-2, 3-dioxygenase (Varjani and Upasani, 2017), were only present at the start of 511 the degradation in the natural attenuation treatment. However, the amended treatment with BSG supported its 512 presence as from day 5 of the degradation process and throughout the experiment (Table 4). Thus, the 513 metabolites present at different times during the bioremediation process in this study, have shown that the 514 breakdown of diesel was mostly via the subterminal oxidation pathway for the aliphatic hydrocarbon content of 515 diesel and via the ortho-oxidation pathway for the aromatic hydrocarbon content of diesel.

516 The amendment of diesel contaminated soil with BSG appeared to enhance metabolism via these two pathways, 517 while also sustaining metabolism via the *meta*-cleavage pathway for aromatic hydrocarbon degradation and 518 speeding up the metabolism of carboxylic acids via the β -oxidation pathway for aliphatic hydrocarbons in terms 519 of synthesis or appearance of metabolic substrates and catabolism of metabolic products. Thus, diesel 520 degradation in this study fits the elucidated pathways, is consistent with the literature (Yanto and Tachibana, 521 2013; Ghosal et al. 2016; Varjani and Upasani, 2017) and provides evidence that the amendment of

522 contaminated soil with BSG enhances the oxidative breakdown of hydrocarbons.

523 Knowledge of the degradation pathway, as determined by the metabolites present, provided information on the 524 degradation genes to be assayed and related the degradation pathways to their associated degradation genes. 525 DNA sequencing targeting the 16S rRNA gene amplicon, on the other hand, provided information regarding the 526 community bacterial populations associated with the biodegradation; thus, correlating these degradation genes 527 with the bacterial population harbouring them and further confirming the bioremediation potential of BSG. 528 Among the three associated genes of the degradation pathways in this study, the *alkB* gene, which is key in the 529 aerobic biodegradation of aliphatics (Powell et al. 2006; Rojo, 2009; Wang et al. 2010; Garrido-Sanz et al. 530 2019), had the highest percentage increase from the start up until the end of the bioremediation especially in the 531 amended treatment. This, interestingly, tallies with the presence of the aliphatic ketone metabolites all through 532 the biodegradation experiment. Similar increase in *alkB* gene copy numbers have been recorded during the 533 bioremediation of a 1% diesel and engine oil contaminated soil using plant residues (Shahsavari et al. 2013).

534 The combination of microbiome analysis with GC-MS evaluation of metabolites adopted in this study, is useful 535 in elucidating the bacterial community dynamics during diesel bioremediation. The results tally with that of the 536 elucidated degradation pathways such that the abundance of aromatic ketone metabolites from the start of the 537 experiment (Table 4), was reflected in the abundance of the catA gene, which was the only gene having a 538 significantly higher copy number from the start of the degradation. This coincided with the rapid hydrocarbon 539 biodegradation observed at the onset of the degradation process. Since the catA gene encodes the catechol-1, 2-540 dioxygenase enzyme, responsible for metabolising aromatic hydrocarbons through the ortho-cleavage pathway 541 and having ketone metabolites, its activity may have contributed to the statistically significant reduction of 542 hydrocarbons in the treatments amended with BSG. A similar rapid initial degradation phase was also observed 543 in the study of Ros et al. (2010).

544 Pseudomonas putida, which was present in abundance in the treatment amended with BSG from the start of the 545 experiment and sustained all through the experiment in this treatment (Fig 5), declined after day 5. Similarly, 546 the catA gene copy numbers (Fig 3b) also declined by day 5. Pseudomonas putida is known to metabolise both 547 aliphatic and aromatic hydrocarbons (Marcus, 2003) and have the catA gene that metabolises aromatic 548 hydrocarbons via the ortho-cleavage pathway catalysed by catechol-1, 2-dioxygenase (Harwood and Parales, 549 1996; Nelson et al. 2002). Pseudomonas putida was also present in the treatment without BSG, in which 550 hydrocarbons were also reduced during the initial rapid biodegradation. Nevertheless, the abundance of P. 551 putida was greater in the amended treatment than in the unamended treatment. This suggests that P. putida may 552 have been actively involved in the metabolism of aromatics catalysed by catechol-1, 2-dioxygenase to produce 553 ketones at the start of the experiment and contributed to the rapid initial degradation observed. Further 554 experiments showing the breakdown pattern of aromatic hydrocarbons, revealed that aromatic hydrocarbons 555 were mostly degraded at the initial degradation phase (Nnadi, 2019).

556

557 A major shift in microbial community in favour of Acinetobacter species was observed following the 558 introduction of diesel to the natural attenuation treatment. Acinetobacter is a known hydrocarbon degrading 559 genus, identified as having the xylE gene encoding catechol-2, 3- dioxygenase that catalyses the degradation of 560 aromatic hydrocarbons via aldehyde metabolites (Méndez et al. 2010; Fuentes et al, 2014). This gene has also 561 been detected in Pseudomonas spp. (Méndez et al. 2010; Fuentes et al. 2014). Acinetobacter was identified in 562 the unamended soil and diesel treatment immediately after the start of the degradation process. It is not 563 surprising, then, that aromatic aldehydes, were identified in this treatment at the start of the degradation. 564 However, as the abundance of Acinetobacter johsonii and Acinetobacter piperi decreased in the unamended 565 treatment after day 5, both species increased in abundance until the end of the experiment in the amended 566 treatment. This was also reflected by the presence of aromatic aldehydes in the amended treatment as from day 5 567 up till the end of the experiment and although Acinetobacter spp. persisted in the unamended treatment, 568 aldehydes were no longer seen in the unamended treatment.

569

570 Community profiling has shown that the genera Acinetobacter and Pseudomonas, among others, may be 571 responsible for the metabolism of hydrocarbons in this study. These genera were also observed to be present in 572 BSG. Flavobacterium, however, was seen to be the most abundant genera in the amended treatment. 573 Nevertheless, it is likely that a synergy of microbes is necessary for the complete degradation of hydrocarbon 574 contaminants rather than a single species. This tallies with previous studies showing that species of 575 Acinetobacter, Arthrobacter, Flavobacterium, Nocardia, Pseudomonas, and Vibrio are associated with 576 petroleum hydrocarbon degradation (Varjani, 2017; Chandra et al. 2013). Since Flavobacterium have been 577 identified as degraders of hydrocarbons (Zhang et al. 2006; Shahsavari et al. 2013; Varjani, 2017), their increase 578 in the soil alone control treatment may be the reason behind the presence of hydrocarbon utilising bacterial 579 CFUs in this treatment as from day 7. Acidobacterium spp. however, reduced in abundance following 580 hydrocarbon contamination. This is interesting to note as they are ubiquitous and mostly found in soils but not 581 known hydrocarbon degraders (Naether et al. 2012). 582

583 The adoption of molecular techniques in this study provided a culture independent approach in the elucidation 584 the bacterial population dynamics during the biodegradation of diesel contaminated soil amended with BSG. 585 Changes in the abundance and an enrichment of autochthonous aerobic hydrocarbon degraders resulting in shifts 586 in the bacterial population in their favour, was made evident. Also, the community profiling of the BSG alone 587 control treatment showed that the grain supports its own microflora; most of which are known hydrocarbon 588 degraders. This supports the potential of supplementation with BSG as not just a biostimulation treatment but 589 also bioaugmentation. 590

Conclusion

594 The findings of this study, including microbiological, metabolite and genetic analysis to assess bioremediation 595 provides a more informed understanding of the process. It also elucidates the correlation between these 596 parameters in monitoring bioremediation to allow for the design of more effective interventions. It has 597 demonstrated that the amendment of diesel contaminated soil with BSG enhanced biodegradation under 598 controlled conditions reflecting temperate environments. BSG contains intrinsic hydrocarbon degrading species 599 including Pseudomonas putida and Acinetobacter piperi and promoted increased gene copy numbers of the 600 alkB, catA and xylE genes. Organic by-products such as BSG can thus provide a valuable contribution to 601 bioremediation as well as reducing potential landfill disposal making it an environmentally viable option.

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628	
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Dear Editor and Reviewers,

Thank you for your suggestions and feedback. We have studied the comments carefully and made corrections to the manuscript which we hope will meet your approval.

Manuscript number: ACMI-D-22-00154 Title: Bacterial community dynamics and associated genes in hydrocarbon contaminated soil during bioremediation using brewery spent grain Authors: Mabel Owupele Nnadi; Lewis Bingle; Keith Thomas

Reviewer 1 Comments to Author: General comments

This paper presents an interesting study dealing with the enhanced degradation of total petroleum hydrocarbons through the addition of brewery spent grain. This is of interest to researchers in the field of environmental bioremediation. The results are clear and convincing. However, in some cases the results were also discussed in the results section, which is not appropriate. In addition, a number of claims lack supporting references. I recommend moving all discussions to the "Discussion" section unless the authors want to combine both sections. Specific comments that address these and other changes are provided below.

Response: Thank you for the informative comments and helpful suggestions. The discussion previously in Section 3.1.3 of the Results section, have been moved to the Discussion section in lines 480 - 494 of the manuscript and more references have been included as suggested.

Introduction

Line 82-84: "The efficiency...degradation pathways." Add one or two more references after this sentence. One relevant reference is the recent study: <u>https://www.nature.com/articles/s43247-022-00526-2</u>

Response: Thank you for this suggestion for improvement. The two references below have been cited in line 84 of the manuscript and added to the reference list in lines 691 and 656.

Eze MO, Thiel V, Hose GC, George SC, Daniel, R (2022) Bacteria-plant interactions synergistically enhance biodegradation of diesel fuel hydrocarbons. Commun Earth Environ 3, 192. <u>https://doi.org/10.1038/s43247-022-00526-2</u>

Bekele GK, Gebrie SA, Mekonen E, Fida TL, Woldesemayat AA, Abda EM, Tafesse M, Assefa F (2022) Isolation and characterization of diesel degrading bacteria from hydrocarbon contaminated sites, flower farms, and soda lakes. Int J Microbiol, 2022:5655767. <u>https://doi.org/10.1155/2022/5655767</u>

Materials and Methods

Line 140: Add the word "in" between the words "used" and "this".

Response: The word 'in' has been added to the manuscript on line 140. Thank you for identifying this clarification.

Line 171-175: How many times was extraction performed per sample? Granted, the effectiveness of n-hexane over other solvents such as DCM for the extraction of diesel fuel hydrocarbons was clearly demonstrated in the study http://dx.doi.org/10.1039/C9RA10919F. However, using non-standardized instruments like shakers (without sonication) once may not lead to a complete extraction of residual diesel fuel. The standard methods of extraction include Soxhlet extraction (USEPA Method 3540C), ultrasonic extraction (USEPA Method 3550C), accelerated solvent extraction (ASE) and microwave-based extraction method.

Response: Thank you for identifying this limitation and suggestions for improvement. The extractions were performed for triplicate samples of each treatment per time point. Thereafter, three aliquots were analysed from each triplicate with a total of nine samples per treatment at each time point. The method is comparable to reported literature (Gielink et al. 2019) in terms of mechanically shaking in hexane and was also evaluated for comparability during initial experiments. And although all the diesel fuel may not have been completely extracted, the same method was used all through the experiment to give comparable results of percentage degradation. We have edited the manuscript in lines 171 - 173, to reflect the wording in the literature referred to, during the experiment.

Line 186-187: Use the formula function in MS Word to express %TPH reduction in order to avoid any ambiguity associated with subtraction, division and multiplication here.

Response: Thank you for this clarification. The formular has been updated in line 185 of the manuscript.

Line 188-190: "The concentration...in the samples." How?

Did you make use of an internal standard that have the same chemical characteristics as your target analyte? Or did you prepare calibration curves for all target compounds and then determined the concentration of each analyte using the parameters of its calibration curve? Please provide more details of the approach you used.

Response: Thank you for suggesting this improvement. The TPH Standard Mix 1 (Sigma Aldrich, UK), with known concentration for each of the (C10 - C28) analytes, was used to obtain the calibration curve for each fraction. Concentrations of each fraction in the soil hydrocarbon mixture were then determined based on the calibration curve of each corresponding standard fraction. Retention times of the analytes were compared to the sample chromatogram to determine target compounds and total peak areas of both standard and analyte fractions were determined. The manuscript has been updated in lines 186 - 191 to include this information.

Results

The results are clearly presented. In some cases, some of the results were also discussed in this section, which is not appropriate, unless the authors decide to combine "results and discussion". I recommend moving discussions to the "Discussion" section. See specific comments below.

Line 325-333: Move this part to "Discussion" section and provide sufficient references as mentioned below.

Response: Thank you for this suggestion. Lines 325 - 337 have been moved to lines 480 - 494 in the Discussion section of the manuscript. The references have also been included in the reference section of the manuscript.

Line 328-329: "The degradation pathways...to catechol". Provide supporting references from published articles. For example, a detailed analysis of these pathways including the mechanisms leading to catechol as well as meta/ortho-cleavages of catechol can be found in doi.org/10.3390/genes12010098 and doi.org/10.1007/BF00186968

Response: Thank you for suggesting this improvement. The sentence has been updated in line 483 - 485 and the references added to the reference section of the manuscript.

Line 335-337: "Aldehydes being...pathways of aromatics." Provide supporting references.

Response: Thank you for identifying and suggesting this improvement. References have been added to this statement in lines 491 - 492.

Line 341-342: "However, carboxylic acids...resulting from aldehyde oxidation," Provide supporting references just after the word "oxidation" and before the continuation of the sentence.

Response: Thank you for identifying and suggesting this improvement. References have been added to this statement in line 332 of the Results section.

Reviewer 2 Comments to Author: In this study, the authors examined the bioremediation of diesel-contaminated soil amended with BSG. They monitored for degradation of petroleum hydrocarbons and for recovery of CFUs. They then determined the bacterial pathways involved in degradation of petroleum hydrocarbons and determined that the bacterial community shifts towards Acinetobacter and Pseudomonas. Overall, the authors found that amendment with BSG resulted in an improved reduction in TPH, and that the bacterial community shifts to break down the hydrocarbons.

The introduction was well-written and made the topic accessible to microbiologists outside of the bioremediation field.

Some minor points to address/areas to improve:

Figure 2: The scale of the y-axes should be the same in Fig 2A and 2B

Response: Thank you for the informative comments and suggestions improvement. Figures 2A and 2B have been updated in line 359 of the manuscript.

What is the cause of the sudden increase in CFUs of hydrocarbon degrading bacteria at Day 5 in the control soil?

Response: Thank you for noting this. The limited growth of hydrocarbon degrading bacteria CFUs up till day 5 is not very clear but may be the result of a long lag phase due to the very small initial numbers of culturable hydrocarbon degrading bacteria in the soil prior to exponential growth by day 5. Similarly, Alisi et al (2009) reported that in soils without prior hydrocarbon contamination, 6 days were required for the soil microbial community to become active. They also observed a 2-day lag phase before CO_2 evolution. This has been updated in the manuscript in lines 463 - 467.

Figure 3: The scale of the y-axes should be the same in Fig 3A-C.

Response: Thank you for identifying this clarification. Figures 3A – C have now been updated in line 392 of the manuscript.

In Fig. 3, are gene copy numbers in the soil+diesel and soil+diesel+BSG shown relative to gene copy numbers in control soil? If so, this should be made clear. If not, how do they compare to the gene copy numbers of the control soil?

Response: Thank you for identifying this clarification. The scope of this study was limited to the quantification of gene copy numbers in the contaminated soil, with and without BSG, to determine the biodegradation potential of BSG. Nevertheless, as this knowledge is useful, it would be considered in future evaluations. The manuscript has been updated to clarify this in lines 367 - 368.

Does diesel impact the recovery of genomic DNA from the soil?

Response: Thank you for identifying this clarification. From the results obtained during this study, the extracted DNA yield from contaminated samples in terms of concentration (ug), were all high and the purity based on absorbance at 260nm and 280nm using a Nanodrop (ND-2000, Thermo Fisher Scientific, UK) were between 1.8 to 2.0 as expected. Thus, it is unlikely that diesel impacted genomic DNA recovery. Lines 403 – 406 of the manuscript has been updated to include this.