

Experimental Optimisation of Various Cultural Conditions on Urease Activity for Isolated Sporosarcina Pasteurii Strains and Evaluation of Their Biocement Potentials

Author

Omoriege, Armstrong Ighodalo, Khoshdelnezamiha, Ghazaleh, Senian, Nurnajwani, Ong, Dominic Ek Leong, Nissom, Peter Morin

Published

2017

Journal Title

Ecological Engineering

Version

Accepted Manuscript (AM)

DOI

[10.1016/j.ecoleng.2017.09.012](https://doi.org/10.1016/j.ecoleng.2017.09.012)

Rights statement

© 2017 Elsevier. Licensed under the Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International Licence (<http://creativecommons.org/licenses/by-nc-nd/4.0/>) which permits unrestricted, non-commercial use, distribution and reproduction in any medium, providing that the work is properly cited.

Downloaded from

<http://hdl.handle.net/10072/375624>

Griffith Research Online

<https://research-repository.griffith.edu.au>

- 1 Experimental optimisation of various cultural conditions on urease activity for isolated
2 *Sporosarcina pasteurii* strains and evaluation of their biocement potentials
- 3 Armstrong Ighodalo Omoregie, Ghazaleh Khoshdelnezamiha, Nurnajwani Senian,
4 Dominic Ek Leong Ong and Peter Morin Nissom
- 5 Armstrong Ighodalo Omoregie: *PhD Candidate*, Faculty of Engineering, Computing and
6 Science, Swinburne University of Technology, Sarawak Campus, Jalan Simpang Tiga, 93350
7 Kuching, Sarawak, Malaysia. Tel: +60146823127,
8 Email: aomoregie@swinburne.edu.my.
- 9 Ghazaleh Khoshdelnezamiha: *PhD Candidate*, Faculty of Engineering, Computing and Science,
10 Swinburne University of Technology, Sarawak Campus, Jalan Simpang Tiga 93350 Kuching,
11 Sarawak, Malaysia. Tel: +60168941252,
12 Email: gKhoshdelnezamiha@swinburne.edu.my.
- 13 Nurnajwani Senian: *PhD Candidate*, Faculty of Engineering, Computing and Science,
14 Swinburne University of Technology, Sarawak Campus, Jalan Simpang Tiga 93350 Kuching,
15 Sarawak, Malaysia. Tel: +60128839239, Email: nSenian@swinburne.edu.my.
- 16 Dominic Ek Leong Ong: *Associate Professor and Director*, Swinburne Sarawak Research
17 Centre for Sustainable Technologies, Swinburne University of Technology, Sarawak Campus,
18 Jalan Simpang Tiga 93350 Kuching, Sarawak, Malaysia. Tel: +6082260631,
19 Email: elong@swinburne.edu.my.
- 20 Peter Morin Nissom: *Associate Professor and Associate Dean (Science)*, Faculty of
21 Engineering, Computing and Science, Swinburne University of Technology, Sarawak Campus,
22 93350 Kuching, Sarawak, Malaysia. Tel: +6082260939,
23 Email: pmorin@swinburne.edu.my.
- 24

25 Corresponding author:

26 Armstrong Ighodalo Omoregie, *PhD Candidate*, Faculty of Engineering, Computing and

27 Science, Swinburne University of Technology, Sarawak Campus, Jalan Simpang Tiga, 93350

28 Kuching, Sarawak, Malaysia. Tel: +60146823127,

29 Email: aomoregie@swinburne.edu.my.

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50 **Abstract**

51 Microbially induced carbonate precipitation (MICP) is a process that has emerged as an
52 attractive alternative ground improvement technique in Geotechnical and Civil
53 Engineering using ureolytic bacteria for soil strengthening and stabilisation. Locally
54 isolated *Sporosarcina pasteurii* from limestone cave samples of Sarawak were found to
55 have high urease-producing abilities. Optimisation of various cultural conditions
56 (incubation temperature, initial pH medium, incubation period and urea concentration)
57 were performed using conductivity and optical density measurements to determine the
58 maximum specific urease activity. In addition, an *in vitro* biocement test was done to
59 define the prospect of using these bacterial isolates in civil engineering work for the
60 improvement of soils with inferior properties. The experimental results showed that
61 urease activities were optimum at 25 to 30°C, pH 6.5 to 8.0, 24 hr incubation and 6 to
62 8% (w/v) urea concentration. It was also demonstrated that biocementation using the
63 local ureolytic bacteria can improve the strength of poorly graded soils. However, the
64 efficiency of the MICP process in improving the soil's strength varied among samples
65 treated with different bacterial cultures.

66 **Keywords:** Microbial induced carbonate precipitation; *Sporosarcina pasteurii*; Urease
67 activity; Bacterial optimisation; Biocementation

68

69

70

71

72

73 **1. Introduction**

74 Microbial induced carbonate precipitation (MICP) is a new ground improvement
75 technique for soils which incorporates ureolytic bacteria such as *Sporosarcina pasteurii*
76 (*S. pasteurii*) to induce carbonate precipitates via urea hydrolysis, thus filling the pores
77 of loose soils and binding soil particles together (Rahim et al. 2015, Jiang et al. 2016).
78 MICP is an effective and environmentally-friendly technology often used to solve
79 various environmental problems such as soil instability and concrete crack (Anbu et al.
80 2016). During MICP process, ureolytic bacteria generate metabolic products (CO_3^{2-})
81 which react with calcium ions (Ca^{2+}) in the microenvironment (Anbu et al. 2016),
82 urease turns the urea molecule (non-conductive) into two charged ions: ammonium
83 (NH_4^+ , positively charged) and carbonates (CO_3^{2-} , negatively charged) (Cuzman et al.
84 2015). These biochemical reactions result in the precipitation of calcium carbonate
85 (CaCO_3) minerals (Jiang et al. 2016).

86 MICP is highly desirable because of its natural availability, effectiveness and
87 sustainability (Al-Thawadi 2008). Biocementation is one of the most reported methods
88 of MICP in the literature. It uses a biochemical binding agent to improve the strength
89 and stiffness properties of soils via microbial activity and its products (Ivanov and Chu
90 2008). Achal et al. (2009a) used a simplified MICP treatment technique for biocement
91 production. The sand columns were injected with *S. pasteurii* (NCIM 2477) and
92 cementation solution under gravimetric free flow direction for the duration of 120 hr.
93 Their finding showed that the CaCO_3 contents precipitated in the sand columns were
94 mostly deposited at the upper layer (40%) which led the reduction of soil porosity and
95 permeability. A large-scale (100 m^3) biocement study by van Paassen et al. (2010) on
96 poorly graded sands showed that when *S. pasteurii* (DSM 33) was injected into a sand

97 box container. They reported having unconfined compressive strength (UCS) results
98 between 0.7 and 12.4 MP, with a nonhomogeneous CaCO₃ contents (12.6 to 27.3%) for
99 their treated sample.

100 Another technique which is also used to treat loose sand is the surface percolation.
101 Cheng and Cord-Ruwisch (2012) reported using this technique to treat Poly Vinyl
102 Chloride (PVC) sand columns (length of 1 m). *Bacillus sphaericus* (DSM 23526) and
103 cementation solution were poured alternatively from the top of the unsaturated sand
104 columns in order to increase the chances of bacterial immobilisation. Their findings
105 showed the UCS strength of the sand column was 390 kPa and there was no clogging at
106 the injection point. They suggested the percolation technique produced higher strength
107 than the submerged flow method previously reported by Whiffin (2004). The use of
108 surface percolation technique was later tested on a larger container (2 m) by Cheng and
109 Cord-Ruwisch (2014) as an attempt to minimise the clogging formation often found in
110 small-scale columns around the injection points which often leads to un-even CaCO₃
111 content distribution. They reported having a relatively homogenous distribution of
112 cement in the sand column, but the UCS results they obtained varied between 850 kPa
113 and 2067 kPa for the coarse samples.

114 There are extensive reports on microorganisms capable of inducing CaCO₃ precipitates
115 for various MICP applications. In a review by Sarayu, Iyer, and Murthy (2014), only a
116 few of the tabulated microorganisms with abilities to induce CaCO₃ precipitate can
117 produce urease enzyme and are non-pathogenic. Ureolytic microbes are preferred for
118 MICP process because via urea hydrolysis, the mechanism is easily controlled with the
119 capacity to induce a substantial amount of CaCO₃ within a reasonable time period
120 (Dhami, Reddy, and Mukherjee 2014). *S. pasteurii* has been reported to be an ideal

121 ureolytic bacteria for biocement applications because of its high urease activity and
122 inability to cause harmful diseases (Wei et al. 2015). In most reported studies, several
123 type strains of *S. pasteurii* such as NCIM 2477, DSMZ 33, MTCC 1761, NCIMB 8841
124 and ATCC 11859 are purchased from different microorganism culture collection centres
125 (Achal et al. 2009a, Wei et al. 2015, Abo-El-Enein et al. 2013, Achal et al. 2009b, Li,
126 Cheng, and Guo 2013). Most locally isolated ureolytic bacteria are from water, soils and
127 sludge samples (Mohammed 2013, Helmi et al. 2016, Arias, Cisternas, and Rivas 2017).
128 However, exploitation on ureolytic bacteria isolated from limestone cave regions for
129 effective biocement application is still very limited. This is because of the notion that
130 caves are generally a region considered as extreme environments, unfavourable
131 microbial growth (Tomczyk-Żak and Zielenkiewicz 2015).

132 Study on optimisation is essential in biotechnological processes because it provides
133 optimum conditions for preferred microbial growth and proliferation (Karamba et al.
134 2016). Each microorganism has its own desired conditions for the maximum enzyme
135 production (Kumar and Takagi 1999). Physical parameters such as temperature and pH
136 play essential roles in enhancing microbial biomass production (Hamzah et al. 2012).
137 Hence, it is necessary to optimise environmental conditions and bacterial growth which
138 will aid large scale microbial production for field biocement application. It is evident
139 that most studies utilise purchased ureolytic bacterial strains for biocement applications
140 while limited studies have reported the use of locally isolated ureolytic bacteria. This
141 present study investigated the effect of various parameters on urease activity for four
142 locally isolated ureolytic bacteria (*S. pasteurii*). These parameters include temperature,
143 initial pH medium, incubation period and urea concentration. A laboratory-scale test
144 was also conducted, as well as a comparative analysis in terms of surface strength to

145 determine the possibility of using these locally isolated ureolytic bacteria for biocement
146 applications.

147 **2. Materials and Methods**

148 2.1. Bacterial strain and growth condition

149 Four ureolytic bacteria were isolated from limestone cave samples of Sarawak,
150 Malaysia using enrichment culture technique and Christensen's medium to isolate and
151 screen for ureolytic bacteria, previously described by Omoregie et al. (2016). These
152 ureolytic bacteria (NB33, LPB21, NB28 and NB30) were selected due to their high
153 specific urease activity (19.98, 23.97, 19.28, and 20.09 urea hydrolysed.min⁻¹.OD⁻¹
154 respectively) when compared to other isolated bacteria and the representative strain (*S.*
155 *pasteurii* DSM 33) (17.75 mM urea hydrolysed.min⁻¹.OD⁻¹) which served as a control
156 strain throughout this study. The bacterial cultures were grown in standard 250 mL
157 conical flasks containing sterile 125 mL of production medium. The growth medium
158 consisted of Nutrient broth (13.0 g.L⁻¹, HiMedia *Laboratories Pvt. Ltd*), supplemented
159 with urea (4% w/v, Bendosen *Laboratory Chemicals*). The initial pH of the media was
160 adjusted to 7.5 using 0.1 M NaOH or 0.1 M HCl before sterilisation (Reyes et al. 2009).
161 Sterile Urea substrate (by 0.45 µm filter sterilisation) was added post-autoclaving to
162 prevent chemical decomposition under autoclave condition. The cultures were
163 temporarily stored in the fridge at +4°C.

164 2.2. Monitoring methods

165 2.2.1. Urease Measurement

166 Conductivity (mS.cm⁻¹) method was used to determine the urease activity (mM urea
167 hydrolysed.min⁻¹) of the ureolytic bacteria. **Conductivity measurement is an analytic**

168 process used to determine the enzymatic rate of reaction because it is easy to operate
169 and an inexpensive assay system (Al-Thawadi 2008). For urease assay, 1.0 mL of 24 hr
170 cultivated bacterial cultures were inoculated into universal bottles containing 9.0 mL of
171 1.11 M urea solution and monitored for 5 min at $25\pm 2^{\circ}\text{C}$ (Whiffin 2004) and the
172 respective conductivity values were measured by immersing the probe of the
173 conductivity meter (Walk LAB conductivity pro meter, Trans Instruments COMPRO)
174 into the bacterial-urea solution. At the end of the assay, a graph was plotted using values
175 from conductivity ($\text{mS}\cdot\text{cm}^{-1}$) measurement against time (min). The conductivity
176 variation rate ($\text{mS}\cdot\text{cm}^{-1}\cdot\text{min}^{-1}$) was acquired from the slope of the plotted graph, which
177 was then multiplied by a dilution factor. This was taken as the ratio of the stock bacteria
178 culture to the sampling bacteria culture before inoculating into the urea solution (Zhao
179 et al. 2014). The specific urease activity ($\text{mM urea hydrolysed}\cdot\text{min}^{-1}\cdot\text{OD}^{-1}$) was derived
180 by dividing the urease activity ($\text{mM urea hydrolysed}\cdot\text{min}^{-1}$) by the bacterial biomass
181 (OD_{600}) (Whiffin 2004).

182 2.2.2. Optical density

183 Optical density (OD) was used as a biomass concentration indicator for the ureolytic
184 bacteria based on the turbidity of the culture. Prior to measuring the OD of the bacterial
185 cultures, a blank (un-inoculated medium) was used to calibrate the spectrophotometer
186 readings. Subsequently, three millilitres (3 mL) of the aliquot was sampled from the
187 bacterial culture and transferred into a cuvette (10 mm). The OD was measured by using
188 a spectrophotometer (GENESYSTM 20, Thermo Fisher Scientific) at a wavelength of
189 600 nm.

190 2.3. Effect of cultural conditions on urease activity

191 The protocol adopted for optimisation of cultural conditions aimed at evaluating the
192 effect of individual condition using one-factor-at-a-time technique and integrate it as the
193 standard level before optimising the parameter. Various cultural conditions affecting
194 urease activity viz., incubation temperature (20-45°C), initial medium pH (6.0-8.5),
195 incubation period (studied up to 96 hr with 24 hr sampling) and urea concentration (2-
196 10%) were studied. The bacterial cultures were grown in nutrient broth medium (13.0
197 g.L⁻¹, HiMedia Laboratories Pvt. Ltd) and supplemented with urea (4% w/v, Bendosen
198 Laboratory Chemicals) by 0.45 µm sterile syringe filter. The overnight grown bacteria
199 were inoculated (2% v/v) into separate sterile conical flasks (containing 125 mL
200 nutrient broth) under aerobic batch conditions with agitation at 130 rpm. The
201 conductivity and OD₆₀₀ were measured and used to determine the specific urease
202 activity at the end of the cultivation period.

203 2.3.1. *Effect of incubation temperature (°C)*

204 The influence of different temperatures ranging from 20±2 to 45±2°C with an interval
205 of 5°C was carried out by incubating the ureolytic bacteria cultures for 24 hr. The initial
206 pH of the growth medium used was attuned to pH 7.5 with the use of 1 N NaOH and
207 1 N HCl. The temperature that promoted the highest enzyme activity was used for
208 subsequent steps of the investigation.

209 2.3.2. *Effect of initial medium pH*

210 The effect of distinctive pH on the ureolytic activity from the selected isolates was
211 determined by examining urease activity at different pH ranging from 6.0 to 8.5 with an

212 interval of 0.5. The initial medium pH that promoted the highest enzyme activity was
213 used for subsequent steps of the investigation.

214 2.3.3. *Effect of incubation period (hr)*

215 The optimal incubation period was determined by incubating the ureolytic bacteria
216 culture at different selected incubation periods ranging from 24 to 96 hr with an interval
217 of 24 hr. The incubation period that promoted the highest enzyme activity was used for
218 subsequent steps of the investigation.

219 2.3.4. *Effect of urea concentration (%)*

220 The influence of urea substrates with varied concentration for enzyme production was
221 studied. Different urea concentration ranging from 2 to 10% (w/v) with an interval of
222 2% (w/v) was selected. The urea concentration that promoted the highest enzyme
223 activity was used during bacterial cultivation for biocement experiment.

224 2.4. Small scale biocementation test

225 2.4.1. *Bacterial culture and cementation solution*

226 Each of the ureolytic bacterial cultures was grown in separate sterile shaking flasks (250
227 mL capacity) containing nutrient broth medium (13.0 g.L⁻¹, HiMedia Laboratories Pvt.
228 Ltd). These bacterial cultures were then incubated using the previously optimised
229 cultural conditions. The bacterial consortia used for biocement treatment was
230 formulated by using a modified method of Hamzah, Phan, Abu Bakar, et al. (2013). The
231 consortia consisted of an equal portion (1:1:1:1, v/v) of the isolates (NB33, LPB21,
232 NB28 and NB30) by mixing 25 % (v/v) of each bacterial culture in shake flask. At the
233 end of each incubation, the OD and enzyme activity of the bacterial cultures were
234 measured immediately before each biocement treatments were performed. The

235 cementation solutions used to treat the sand columns consist of nutrient broth (13 g.L⁻¹,
236 HiMedia Laboratories Pvt. Ltd), urea (1 M, Bendosen Laboratory Chemicals), calcium
237 chloride (1 M, Sigma-Aldrich Co. LLC), sodium acetate (0.17 M, HiMedia
238 Laboratories Pvt. Ltd) and ammonium chloride (0.0125 M, HiMedia Laboratories Pvt.
239 Ltd) were adapted from Cheng et al. (2014) and Weaver et al. (2011). All the
240 cementation solution components were autoclaved except urea and CaCl₂, which were
241 added after the solution was autoclaved.

242 2.4.2. Preparation of sand columns

243 The sand used for the biocement test was classified as poorly graded medium sand in
244 accordance with British Standards, BS5930. The particle sizes selected ranged from fine
245 sand (0.075 mm) to fine gravel (4.75 mm). Re-informed paper tubes served as the
246 moulds used in this experiment having an internal diameter of 75 mm and length of 49
247 mm. Each column (mould) was packed with 294.73 g of autoclaved sand. All columns
248 were placed on flat surfaced polypropylene sheet; five holes were drilled on the surfaces
249 of the polypropylene sheets to allow the effluents of the cementation solution to pass
250 through. The polypropylene sheets containing drilled holes were later covered with
251 Whatman filter papers. A plastic container was placed below the polypropylene sheet to
252 accumulate the effluents. The top of each column was covered with a layer of scouring
253 pads (Scotch-Brite™) as filters to prevent disturbance on the surfaces of the sands
254 during biocement treatments. Seven sets of sand columns were prepared and each set
255 contained three replicates. The biocement test was done in triplicates per bacterial
256 treatment. As shown in Figure 6, Set 1 was treated with positive control (*S. pasteurii*
257 DSM 33), set 2 was treated with consortia, set 3 was treated with LPB21, set 4 was

258 treated with NB33, set 5 was treated with NB28 and set 6 was treated with NB33. Set 7
259 was the negative control (no bacteria) and was only treated with cementation solution.

260 2.4.3. *Biocementation method*

261 Prior to the beginning of the treatment, the sand samples were mixed with bacteria
262 culture, calcium chloride and urea solution before being compacted into their respective
263 columns. The sand columns were treated with the bacteria and cementation solutions
264 using surface percolation technique. The sands were treated with separate isolates,
265 bacterial consortia, positive control (*S. pasteurii* DSM 33) and negative control (sand
266 specimen treated with cementation solution only). The bacterial cultures were harvested
267 at their respective late exponential phases before being mixed with the air-dried sand
268 specimens. The columns were treated twice daily with the 80 mL ureolytic bacteria
269 culture and 80 mL cementation solution. However, the treatment was split into two
270 series of treatment and added twice daily. The MICP treatment was performed by
271 introducing 80 mL of bacterial culture and 80 mL of cementation solution into the sand
272 specimens at an interval of 12 hr for a duration of 96 hr. The treatments of the sand
273 columns were performed inside a fume hood (LabCraft, BASIX 52). Upon completion
274 of the treatments, all the sand columns were cured at room temperature for a duration of
275 14 days before the treated sand were being removed from their respective mould.

276 2.4.4. *Surface strength measurement*

277 The surface strength measurements of the treated sand were obtained by using a pocket
278 penetrometer (ELE International, 38-2695) as suggested by Al-Thawadi (2008). The
279 penetrometer used had a reading scales from 0 to 700 psi (0 to 4.83 MPa). The pocket
280 penetrometer was used to measure the surface strength by pushing the tip of the
281 penetrometer into the soil to a depth of approximately 0.25 inches and three selected

282 surface regions were tested on each of the cemented sand. The readings of the loaded
283 weight were recorded when the samples were completely penetrated (breakage).

284 2.5. Statistical analysis

285 A standard deviation for each experimental result obtained was calculated using the
286 Excel spreadsheets available in the Microsoft Excel (version 2016). One-way analysis
287 of variance (ANOVA) and post hoc Turkey's test was analysed using StatPlus program
288 to indicate any significant difference between groups. The level of significance was set
289 at 0.05.

290 3. Results and Discussion

291 The four ureolytic bacteria isolated from limestone cave samples of Sarawak showed
292 high degrees of similarity (96-98%) to their respective closest bacterial species as
293 shown in Table 1. The BLAST results suggested that the isolates were closely related to
294 bacteria from the *S. pasteurii* group. Achal and Pan (2011) have previously reported
295 isolating *S. pasteurii* group from an extreme environment. The microbial cell wall such
296 as teichuronic-peptide play a role in the pH homeostasis at alkaline pH, hence allowing
297 for a survival in extreme environments such as caves (Aono, Ito, and Machida 1999).

298 3.1. The effect of incubation temperature (°C)

299 The specific urease activity varied in the bacterial cultures grown at different
300 temperatures (Figure 1). Isolate NB33 (25.32 mM urea hydrolysed.min⁻¹.OD⁻¹), isolate
301 NB30 (41.98 mM urea hydrolysed.min⁻¹.OD⁻¹) and control strain (23.03 mM urea
302 hydrolysed.min⁻¹.OD⁻¹) produced their respective maximum specific urease activity
303 when incubated at 30°C, while 25°C was observed to be the maximum specific urease
304 activity for isolate LPB21 (29.81 mM urea hydrolysed.min⁻¹.OD⁻¹) and isolate NB28

305 (26.26 mM urea hydrolysed.min⁻¹.OD⁻¹) when compared to other incubation
306 temperature. Among the parameters that could affect urease activity, the temperature is
307 a critical factor that needs to be controlled because it usually varies from one organism
308 to another and can have a substantial effect on urease production (Coulon et al. 2004,
309 Kumar and Takagi 1999, Soon et al. 2014). Hence, in this study, the temperature was
310 first chosen as a parameter to optimised. The result presented in Figure 1 suggests that
311 at 25°C, the maximum specific urease activity for isolate LPB21 and NB30 were
312 observed. On the other hand, at 30°C, the maximum specific urease activity for isolate
313 NB33, NB28 and the control strain were observed. It was obvious as shown in Figure 1
314 that these isolates were able to grow at the studied temperature range variation from
315 20°C to 45°C and could still display urease activities, which advocates the possibility of
316 these isolates to mesophilic microorganisms (Willey, Sherwood, and Woolverton 2008).
317 Statistical analysis using ANOVA with Tukey's procedure showed there were
318 significant differences for the mean at 25°C for isolate LPB21 (P-value = 1.74E-04, M=
319 29.81), as well as 30°C for isolate NB33 (P-value = 4.06E-05, M= 25.32) and isolate
320 NB30 (P-value = 8.91E-07, M= 41.98). Two separate temperatures (25°C and 30°C)
321 were selected to cultivate the bacterial isolates, for the rest of the experiments in this
322 study, and were performed in different incubation shakers. These two temperatures
323 showed maximum specific urease activity for the different isolates. LPB21 and NB30,
324 was cultivated at 25°C, while 30°C was used to cultivate isolate NB33 and NB28. The
325 recommended temperature for cultivating the control strain is 30°C.

326 3.2. The effect of initial medium pH

327 In Figure 2, the maximum specific urease activity was observed in the medium of pH
328 6.5 for isolate NB33 (23.71 mM urea hydrolysed.min⁻¹.OD⁻¹), whereas isolate LPB21

329 and control strain showed their respective maximum specific activity at pH 7.5 with
330 33.74 and 21.43 mM urea hydrolysed.min⁻¹.OD⁻¹, respectively. Isolate NB30 and isolate
331 NB 28 showed their individual maximum specific urease activities at pH 8.0 with 30.92
332 and 34.51 mM urea hydrolysed.min⁻¹.OD⁻¹. The pH of a microorganism's growth
333 medium plays an important role capable of inducing the morphological changes of the
334 microbe, instigate its enzyme secretion and affect its stability (Sethi and Gupta 2014). *S.*
335 *pasteurii* is known to grow well in medium ranging from 6.5 to 8.0 (Sakai and Imachi
336 2015), but the effect on urease productivity for isolated bacterial strains are limited.
337 Hence, the effect of the initial medium pH (6.0 to 8.5) on urease activity for the bacteria
338 cultures were studied. Analysis of variance with Tukey's procedure showed there were
339 significant differences for the mean at pH 6.5 for isolate NB33 (P-value = 1.97E-05,
340 M= 23.71), at pH 7.5 for control strain (P-value = 0.004, M= 21.43) and isolate LPB21
341 (P-value = 4.56E-04, M= 33.74), while at pH 8.0 for isolate NB28 (P-value = 1.84E-04,
342 M= 34.51) and isolate NB30 (P-value = 2.24E-07, M= 30.92). The results illustrated in
343 Figure 2 indicate that all the bacterial cultures had their respective maximum specific
344 urease activity when incubated in growth medium with initial pH at 6.5, 7.5 and 8.0. A
345 study by Gat et al. (2014) suggested that the pH values of *S. pasteurii* (DSM 33) for
346 urea hydrolysis during incubation is at pH 7.39 after being incubation at 28 hr. During
347 fermentation, urea hydrolysis is expected to lead to an increase in the pH value due to
348 the production of ammonium (Gat et al. 2014). The enzyme activities at pH 7.5 and 8.0
349 in Figure 2 were the most noticeable values when compared to other pH values. On the
350 other hand, when the bacterial cultures were inoculated into growth medium at pH 6.0
351 and 8.5, their respective specific urease activities were found to be lower than the
352 aforementioned pH values. It has been reported in the literature that the optimum pH of

353 most microbial ureases is near neutral, while some alkali-tolerant bacteria such as *S.*
354 *pasteurii*, *Bacillus sphaericus* and *Bacillus megaterium* have their optimum pH ranging
355 between 7 to 9.5 (Soon et al. 2014).

356 3.3. The effect of incubation period (hr)

357 In Figure 3, the maximum specific urease activity was observed at 24 hr incubation
358 period for all the bacteria; isolate LPB21 (25.98 mM urea hydrolysed.min⁻¹.OD⁻¹),
359 isolate NB33 (27.93 mM urea hydrolysed.min⁻¹.OD⁻¹), isolate NB28 (25.54 mM urea
360 hydrolysed.min⁻¹.OD⁻¹), isolate NB30 (29.70 mM urea hydrolysed.min⁻¹.OD⁻¹) and the
361 control strain (22.08 mM urea hydrolysed.min⁻¹.OD⁻¹). Based on observations and
362 reports from other studies (Qabany, Soga, and Santamarina 2012, Stocks-Fischer,
363 Galinat, and Bang 1999), it was suggested that for higher urease activity, ureolytic
364 bacterial cells have optical density (OD₆₀₀) values in the range of 0.8 to 1.2
365 (10⁷ cells/mL) which was achieved over a 24-hr incubation period. Hence, in this study,
366 24 hr was chosen as the starting incubation time, and the experiment was then
367 monitored over a 24-hourly incubation interval, to assess the effect of incubation time
368 on the specific urease activities of the isolates. Analysis of variance with Tukey's
369 procedure suggested there were significant differences for the mean at 24 hr for the
370 control (P-value = 8.70E-07, M= 22.08), isolate LPB21 (P-value = 3.88E-04, M=
371 25.98), isolate NB33 (P-value = 8.79E-07, M= 27.93), isolate NB30 (P-value= 3.15E-
372 04, M= 29.70) and isolate NB28 (P-value = 1.10E-02, M= 25.54). The incubation
373 period is an essential parameter for enzyme production (Gautam et al. 2011). To
374 elucidate the effect of different incubation period on urease production and microbial
375 growth, experiments were conducted at different selected incubation period from 24 to
376 96 hr. The data analysed showed that during fermentation period, the specific urease

377 activity for all the bacterial cultures reached their maximum values at 24 hr incubation
378 period as presented in Figure 3. Further incubation (48 to 96 hr) of the cultures revealed
379 that their specific urease activity reduced drastically. **This finding is true only among**
380 **the tested incubation period.** A few possible reasons for this reduction of enzyme
381 production might have occurred during the fermentation might be due to the reduction
382 of essential nutrients, insufficient sugar contents in the medium, due to inhibitory
383 metabolites (Ali et al. 2016) or absence of sufficient hydrolyzed urea as nitrogen for
384 energy (Achal et al. 2009b, Burne and Chen 2000). In addition, the drastic reduction of
385 specific urease activity observed in Figure 3 might be due to saturated active sites of the
386 microbial enzyme by the substrate molecules (Robinson 2015) which occurs when
387 ureolysis has been completely used up.

388 3.4. The effect of urea concentration (%)

389 In Figure 4, the maximum specific urease activity was observed at 6% of urea
390 concentration for isolate LPB21 (32.36 mM urea hydrolysed.min⁻¹.OD⁻¹) and isolate
391 NB28 (25.98 mM urea hydrolysed.min⁻¹.OD⁻¹), while 8% urea concentration was
392 observed to show the maximum specific activity for isolate NB33 (33.95 mM urea
393 hydrolysed.min⁻¹.OD⁻¹), isolate NB30 (39.21 mM urea hydrolysed.min⁻¹.OD⁻¹) and
394 control strain (24.66 mM urea hydrolysed.min⁻¹.OD⁻¹). **Bacteria need a source of**
395 **nitrogen to support their maximal growth because nitrogen is a key building block of**
396 **protein, enzymes and nucleic acids (Hamzah, Phan, Abu-Bakar, et al. 2013).** ANOVA
397 with Tukey's procedure showed there were significant differences for the mean at 8%
398 for control strain (P-value = 0.00, M= 24.66), and isolate NB30 (P-value= 2.62E-04,
399 M= 39.21), while at 6% for isolate LPB21 (P-value = 6.43E-04, M= 32.36) and isolate
400 NB28 (P-value = 3.00E-02, M= 25.98). The result illustrated in Figure 4 suggests that

401 urea concentration at 6% (w/v) and 8% (w/v) were most favourable for the specific
402 urease activities of the bacteria isolates and control. Ammonia, a nitrogen source for
403 most bacteria can be detrimental or toxic to their environment when present in high
404 concentration due to cytotoxic effect (Hess et al. 2006). However, a higher
405 concentration of ammonia can be advantageous to ureolytic bacteria by assisting their
406 Adenosine triphosphate (ATP) generation which can supply more metabolic and
407 enzymatic activities (Mempin et al. 2013). On the other hand, increasing the urea
408 concentration can result in a decrease of the bacteria's biomass production (Cheng and
409 Cord-Ruwisch 2013). This is because the energy utilised by ATP for microbial
410 replication and growth would be reduced and then be used to generate and supply
411 energy enzymatic reactions (Mempin et al. 2013). The finding also showed that these
412 isolates can grow and produce substantial urease when 4% (w/v) and 10% (w/v) of urea
413 were supplemented into their respective growth medium, but when 2% (w/v) of urea
414 was used to grow the bacterial cultures, their specific urease activities were significantly
415 low. Hence, it is recommended that higher concentration of urea be used to cultivate
416 these ureolytic bacteria for MICP process.

417 3.5. Biocementation treatment

418 In Figure 5 (a), there was no visual observation of calcium carbonate precipitates at the
419 top layer of the columns during the initial period of immersion of the bacterial culture
420 and cementation, however on the third day of inoculation, white precipitates were seen
421 on all triplicate samples of the columns containing bacterial cultures (Figure 5 (b)). On
422 the other hand, none of the columns containing the negative control displayed any
423 visible precipitation on their respective top layers. The white precipitates on the top
424 layers of the biocemented sand shown in Figure 5 and Figure 6 were also reported by

425 Zhao et al. (2014) and Chu, Stabnikov, and Ivanov (2012), indicating the presence of
426 nucleation sites for MICP due to addition of more bacterial solution in order to promote
427 more urease enzyme. The influence of microbial cementation on granular behaviour is
428 dependent on the ability of the bacteria to move freely throughout the pore spaces of the
429 sand and on sufficient particle-particle contact per unit volumes at which cementation
430 will occur. Hence, this quicker formation of calcium carbonate precipitates at injection
431 points of the bacteria and cementation solution prevents more precipitates from flowing
432 freely down the columns and causing a non-uniformity of calcium carbonate
433 precipitates (Achal et al. 2009a, b). The columns holding the biocemented sands were
434 carefully removed at the end of the curing period as shown in Figure 6. All the
435 biocemented sand samples appeared to remain intact after removal from the columns. It
436 was also observed that the scouring pads (Scotch-Brite™) which were used to prevent
437 any disturbance of the column's top surfaces was not very productive during injection
438 of the cementation solution. However, the hardness of the biocemented sands was not
439 affected. The negative control contained only the cementation solution was done to rule
440 out the possibility that precipitates found in the sand columns were only as results
441 microbial urea hydrolysis and not any other process. Surface percolation method was
442 used for biocement treatment on the sand samples because with the help of gravity and
443 capillarity, the bacterial culture and cementation reagents were easily transported into
444 the sand particles (Cheng and Shahin 2016). The main merit of utilising this biocement
445 method is that it does not require heavy machinery (Mujah, Shahin, and Cheng 2016),
446 making it cost effective, simple and a practical approach due to the free movement of
447 the liquid used. Typically, during biocementation process using percolation method, the
448 solutions are alternatively oozed onto the surface of the sands while the sand column

449 was kept in a vertical position (Cheng and Cord-Ruwisch 2014, Dhami, Reddy, and
450 Mukherjee 2016) with the top and bottom of the column fully unsealed as shown in
451 Figure 5. Since the objective of this study was to exploit the capability of using locally
452 isolated ureolytic bacteria to improve the strength of loose soils under simple biocement
453 treatment method, the downward vertical flushing using percolation method was
454 selected rather than using other soil treatment methods via injection (Rowshanbakht et
455 al. 2016, Liang, Mohamed, and Donovan 2016) or pressurized flow (Soon et al. 2014,
456 Martinez et al. 2013). These methods often require hydraulic injection of the
457 cementation reagents and physical removal of the effluents (Cheng and Cord-Ruwisch
458 2014). It is vital to take note that even though percolation method feasible, it must be
459 performed cautiously so as not distort the surface of the sand column. This distortion
460 can be seen in **Figure 6** for one of the sand columns treated with isolate NB28 which
461 occurred during the treatment process Cheng and Cord-Ruwisch (2012) reported that
462 based on MICP using ureolytic bacteria, percolation treatment method does well in
463 saturated and unsaturated sands and produces higher strengths than submerged flow
464 method used by van Paassen et al. (2010) and Whiffin, van Paassen, and Harkes (2007)
465 in waterlogged soils.

466 3.6. Soil surface strength

467 In Figure 7, the strength measured for the biocemented sand treated with different
468 ureolytic bacteria are 582.33 psi for isolate LPB21, 626.67 psi for isolate NB33, 573.33
469 psi for isolate NB30, 700 psi for isolate NB28, 533.33 psi for bacterial consortia and
470 563.33 for the positive control strain. However, the negative control was too soft to
471 measure and could not yield any result. The highest strength measured was 700 psi for
472 biocemented sand treated with isolate NB28 while the lowest strength measured was

473 533.33 psi for consortia. One of the factors which may have affected the biomass
474 synergy of the bacterial consortia could be attributed to insufficient oxygen in the
475 microenvironment (Hamzah, Phan, Abu-Bakar, et al. 2013). It is also suggested
476 bacterial consortia construction with varied biomass concentration (v/v %) be optimised
477 to achieve a highly-desired biomass production and urease activity (Hamzah et al.
478 2016). Among all the biocemented sand, the sample treated with isolate NB28 reached
479 the maximum reading of the penetrometer and none of its samples cracked during this
480 surface strength test, unlike other samples. It was also observed that the sand treated
481 with bacterial cultures and cementation solutions were slightly more cemented in areas
482 closest to the point of injection regions. Visual observation after the strength test also
483 indicated that there were much more precipitates on the surface of the biocemented
484 sands than other areas. The bacterial consortia might not have performed better than
485 the individual bacterial isolates because of a low number of the bacteria cells. This led
486 to the sand that was treated with the bacterial consortia to have the lowest strength result
487 because of low biomass and urease production. Statistical analysis suggested there were
488 no significant differences among the surface strengths treated with local isolates and the
489 control. These local bacteria have potential applications for soil surface treatment using
490 MICP technology, especially in tropical rainforest climate such as Malaysia which has
491 abundant loose sands, posing challenges to engineers during the early stage of
492 construction. Some of these Malaysians soils often experience further soil softening due
493 to an extreme and prolonged downpours, which can be problematic for engineers hence
494 MICP treatment is strongly needed (Soon 2013).

495 **4. Conclusions**

496 In this study, a series of experimental tests were conducted to determine the optimised
497 specific urease activities for four locally isolated ureolytic bacteria (*S. pasteurii*) using
498 different parameters: temperature, pH, incubation period and urea concentration. In
499 addition, an *in vitro* laboratory biocement test was performed using MICP process to
500 investigate the feasibility of using these four isolates for strengthening and stabilisation
501 of poorly graded soil. The following conclusions can be drawn from this report:

502 (1). Based on the one-factor-at-a-time method used to investigate the effect of different
503 parameters on specific urease activities, the optimum incubation temperatures were
504 25°C (for isolates LPB21 and NB28) and 30°C (for isolates NB30 and NB33). The
505 optimum initial pH medium was pH 6.5 (for NB33), pH 7.5 (for LPB21) and pH
506 8.0 for (NB30 and NB28). Isolates LPB21 and NB28 showed their optimum urea
507 concentration at 6% (w/v), while isolates NB30 and NB33 showed their optimum
508 urea concentration at 8% (w/v). **Moreover**, all four locally isolated ureolytic
509 bacteria showed their respective optimum incubation period at 24 hr. These
510 findings suggest that urease activity for the isolates were favoured by mesophilic
511 temperature, alkaline pH, short incubation period and high urea concentration.

512 (2). The use of percolation method as an inexpensive and simple MICP process to treat
513 low rigidity soils was effective. The results showed that local ureolytic bacteria
514 were able to induce calcium carbonate precipitates comparable to the performance
515 of the control strain (*S. pasteurii* DSM 33). However, the efficiency of the MICP
516 process in improving the soil's strength varied among the samples which were
517 treated with different isolates, bacterial consortia, and the control strain.

518 (3). Local strength of the treated sand columns obtained from penetrometer test showed
519 that among all the bacteria used during the treatment via MICP process, isolate

520 NB28 (700 psi) had the highest surface strength, while the bacterial consortia
521 (533.33 psi) had the lowest surface strength. This could be due to different bacterial
522 cell concentration and urease produced by the microbes.

523 The test results attained from this present study suggest these isolates can serve as
524 alternative ureolytic bacteria for improvement of soil strength via MICP process as a
525 sustainable technology. Although, this study only investigated the use of percolation
526 method for these four isolates, further works are currently being conducted using these
527 ureolytic bacteria employing other alternative treatment methods in attempts to improve
528 the distribution of calcium carbonate contents in soil particles, reduce formation of
529 clogging at regions close to treatment points and enhance the soil's strength. Large-
530 scale bacterial production using these isolates are ongoing for biocement field
531 application.

532 **Acknowledgements**

533 This study was supported by Research, Consultancy & Future Projects under Swinburne
534 Sarawak Research Grant (SSRG 2-5502). Special thanks go to the science laboratory
535 technicians for provision of some experimental materials which contributed to the
536 completion of this study.

537 **References**

538 Abo-El-Enein, S. A., A. H. Ali, Fatma N. Talkhan, and H. A. Abdel-Gawwad. 2013.
539 "Application of microbial biocementation to improve the physico-mechanical
540 properties of cement mortar." *Housing and Building National Research Center* 9
541 (1):36-40. doi: 10.1016/j.hbrcj.2012.10.004.

542 Achal, V., A. Mukherjee, P. C. Basu, and M. S. Reddy. 2009a. "Lactose mother liquor
543 as an alternative nutrient source for microbial concrete production by
544 *Sporosarcina pasteurii*." *Journal of Industrial Microbiology & Biotechnology*
545 36 (3):433-438. doi: 10.1007/s10295-008-0514-7.

546 Achal, V., A. Mukherjee, P. C. Basu, and M. S. Reddy. 2009b. "Strain improvement of
547 *Sporosarcina pasteurii* for enhanced urease and calcite production." *Journal of*
548 *Industrial Microbiology & Biotechnology* 36 (7):981-988. doi: 10.1007/s10295-
549 009-0578-z.

550 Achal, V., and X. Pan. 2011. "Characterization of urease and carbonic anhydrase
551 producing bacteria and their role in calcite precipitation." *Current*
552 *Microbiology* 62 (3):894-902. doi: 10.1007/s00284-010-9801-4.

553 Al-Thawadi, Salwa. M. 2008. "High Strength In-Situ Biocementation of Soil by Calcite
554 Precipitating Locally Isolated Ureolytic Bacteria." PhD, School of Biological
555 Sciences and Biotechnology, Murdoch University.

556 Ali, S. R., Z. Anwar, M. Irshad, S. Mukhtar, and N. Tariq. Warraich. 2016. "Bio-
557 synthesis of citric acid from single and co-culture-based fermentation
558 technology using agro-wastes." *Journal of Radiation Research and Applied*
559 *Sciences* 9 (1):57-62. doi: 10.1016/j.jrras.2015.09.003.

560 Anbu, P., C. H. Kang, Y. J. Shin, and J. S. So. 2016. "Formations of calcium carbonate
561 minerals by bacteria and its multiple applications." *Springerplus* 5:250-262. doi:
562 10.1186/s40064-016-1869-2.

563 Aono, R., M. Ito, and T. Machida. 1999. "Contribution of the cell wall component
564 teichuronopeptide to pH homeostasis and alkaliphily in the alkaliphile *Bacillus*
565 *lentus* C-125." *Journal of Bacteriology* 181 (21):6600-6606.

- 566 Arias, D., Luis. A. Cisternas, and M. Rivas. 2017. "Biom mineralization of calcium and
567 magnesium crystals from seawater by halotolerant bacteria isolated from
568 Atacama Salar (Chile)." *Desalination* 405:1-9.
- 569 Burne, R. A., and Y. M. Chen. 2000. "Bacterial ureases in infectious diseases."
570 *Microbes and Infection* 2 (5):533-542.
- 571 Cheng, L., and R. Cord-Ruwisch. 2013. "Selective enrichment and production of highly
572 urease active bacteria by non-sterile (open) chemostat culture." *Journal of*
573 *Industrial Microbiology and Biotechnology* 40 (10):1095-1104. doi:
574 10.1007/s10295-013-1310-6.
- 575 Cheng, L., and R. Cord-Ruwisch. 2014. "Upscaling Effects of Soil Improvement by
576 Microbially Induced Calcite Precipitation by Surface Percolation."
577 *Geomicrobiology Journal* 31 (5):396-406. doi: 10.1080/01490451.2013.836579.
- 578 Cheng, L., M. A. Shahin, R. Cord-Ruwisch, M. Addis, T. Hartanto, and C. Elms. 2014.
579 "Soil Stabilisation by Microbial-Induced Calcite Precipitation (MICP):
580 Investigation into Some Physical and Environmental Aspects." 7th International
581 Congress on Environmental Geotechnics Melbourne, Australia.
- 582 Cheng, L., and R. Cord-Ruwisch. 2012. "In situ soil cementation with ureolytic bacteria
583 by surface percolation." *Ecological Engineering* 42:64-72. doi:
584 10.1016/j.ecoleng.2012.01.013.
- 585 Cheng, L., and M. A. Shahin. 2016. "Urease active bioslurry: a novel soil improvement
586 approach based on microbially induced carbonate precipitation." *Canadian*
587 *Geotechnical Journal* 53 (9):1376-1385. doi: 10.1139/cgj-2015-0635.

588 Chu, J., V. Stabnikov, and V. Ivanov. 2012. "Microbially Induced Calcium Carbonate
589 Precipitation on Surface or in the Bulk of Soil." *Geomicrobiology Journal* 29
590 (6):544-549. doi: 10.1080/01490451.2011.592929.

591 Coulon, F. , E. Pelletier, L. Gourhant, and D. Delille. 2004. "Effects of temperature
592 warning during a bioremediation study of natural and nutrient-amended
593 hydrocarbon-contaminated sub-Antartic soils." *Cold Regions Science and
594 Technology* 40:61-70. doi: 10.1016/j.chemosphere.2004.10.007.

595 Cuzman, O. A., K. Richter, L. Wittig, and P. Tiano. 2015. "Alternative nutrient sources
596 for biotechnological use of *Sporosarcina pasteurii*." *World Journal of
597 Microbiology and Biotechnology* 31 (6):897-906. doi: 10.1007/s11274-015-
598 1844-z.

599 Dhami, N. K., M. S. Reddy, and A. Mukherjee. 2014. "Application of calcifying
600 bacteria for remediation of stones and cultural heritages." *Frontiers in
601 microbiology* 5:304. doi: 10.3389/fmicb.2014.00304.

602 Dhami, N. K., M. S. Reddy, and A. Mukherjee. 2016. "Significant indicators for
603 biomineralisation in sand of varying grain sizes." *Construction and Building
604 Materials* 104:198-207. doi: 10.1016/j.conbuildmat.2015.12.023.

605 Gat, D., M. Tsesarsky, D. Shamir, and Z. Ronen. 2014. "Accelerated microbial-induced
606 CaCO₃ precipitation in a defined coculture of ureolytic and non-ureolytic
607 bacteria." *Biogeosciences* 11 (10):2561-2569. doi: 10.5194/bg-11-2561-2014.

608 Gautam, S. P., P. S. Bundela, A. K. Pandey, J. Khan, M. K. Awasthi, and S. Sarsaiya.
609 2011. "Optimization for the production of cellulase enzyme from municipal
610 solid waste residue by two novel cellulolytic fungi." *Biotechnology Research
611 International* 2011:810425-810433. doi: 10.4061/2011/810425.

- 612 Hamzah, A., C. W. Phan, N. F. Abu-Bakar, and K. K. Wong. 2013. "Biodegradation of
613 Crude Oil by Constructed Bacterial Consortia and the Constituent Single
614 Bacteria Isolated From Malaysia." *Bioremediation Journal* 17 (1):1-10. doi:
615 10.1080/10889868.2012.731447.
- 616 Hamzah, A., M.A. Zarin, A.A. Hamid, O. Omar, and S. Senafi. 2012. "Optical Physical
617 and Nutrient Parameters for Growth of *Trichoderma virens* UKMP-1M for
618 Heavy Crude Oil Degradation." *Sains Malaysiana* 41 (1):71-79.
- 619 Hamzah, A., C. Phan, N. Abu Bakar, and K. K. Wong. 2013. "Biodegradation of crude
620 oil by constructed bacterial consortia and the constituent single bacteria isolated
621 from Malaysia." *Bioremediation Journal* 17 (1):1-10.
- 622 Hamzah, A., S. N. M. Salleh, K. K. Wong, and S. Sarmani. 2016. "Nutrient
623 Amendments of Inorganic Fertiliser and Oil Palm Empty Fruit Bunch and Their
624 Influence on Bacterial Species Dominance and Degradation of the Associated
625 Crude Oil Constituents." *Soil and Sediment Contamination: An International
626 Journal* 25 (3):256-265. doi: 10.1080/15320383.2016.1124834.
- 627 Helmi, F. M., H. R. Elmitwalli, S. M. Elnagdy, and A. F. El-Hagrassy. 2016. "Calcium
628 carbonate precipitation induced by ureolytic bacteria *Bacillus licheniformis*."
629 *Ecological Engineering* 90:367-371. doi: 10.1016/j.ecoleng.2016.01.044.
- 630 Hess, D. C., W. Lu, J. D. Rabinowitz, and D. Botstein. 2006. "Ammonium toxicity and
631 potassium limitation in yeast." *PLOS Biology* 4 (11):351. doi:
632 10.1371/journal.pbio.0040351.
- 633 Ivanov, V., and J. Chu. 2008. "Applications of microorganisms to geotechnical
634 engineering for bioclogging and biocementation of soil in situ." *Reviews in*

635 *Environmental Science and Biotechnology* 7 (2):139-153. doi: 10.1007/s11157-
636 007-9126-3.

637 Jiang, N., Hideyoshi. Y, Koji. Y, and K. Soga. 2016. "Ureolytic activities of a urease-
638 producing bacterium and purified urease enzyme in the anoxic condition:
639 Implication for subseafloor sand production control by microbially induced
640 carbonate precipitation (MICP)." *Ecological Engineering* 90:96-104. doi:
641 10.1016/j.ecoleng.2016.01.073.

642 Karamba, K. I., S A. Ahmad, A. Zulkharnain, M. A. Syed, K. A. Khalil, N. A.
643 Shamaan, F. A. Dahalan, and M. Y. Shukor. 2016. "Optimisation of
644 biodegradation conditions for cyanide removal by *Serratia marcescens* strain
645 AQ07 using one-factor-at-a-time technique and response surface methodology."
646 *Rendiconti Lincei* 27 (3):533-545. doi: 10.1007/s12210-016-0516-8.

647 Kumar, C. G., and H. Takagi. 1999. "Microbial alkaline proteases from a bio-industrial
648 view point." *Biotechnology Advances* 17:561-594. doi: 10.1016/S0734-
649 9750(99)00027-0.

650 Li, M., X. Cheng, and H. Guo. 2013. "Heavy metal removal by biomineralization of
651 urease producing bacteria isolated from soil." *International Biodeterioration &*
652 *Biodegradation* 76:81-85. doi: 10.1016/j.ibiod.2012.06.016.

653 Liang, C., A. Shahin. M, and M. Donovan. 2016. "Influence of Key Environmental
654 Conditions on Microbially Induced Cementation for Soil Stabilization." *Journal*
655 *of Geotechnical and Geoenvironmental Engineering* 143 (1):04016083-
656 04016093. doi: 10.1061/(ASCE)GT.1943-5606.0001586.

657 Martinez, B. C., J. T. DeJong, T. R. Ginn, B. M. Montoya, T. H. Barkouki, C. Hunt,
658 B. Tanyu, and D. Major. 2013. "Experimental Optimization of Microbial-

659 Induced Carbonate Precipitation for Soil Improvement." *Journal of*
660 *Geotechnical and Geoenvironmental Engineering* 139 (4):587-598. doi:
661 doi:10.1061/(ASCE)GT.1943-5606.0000787.

662 Mempin, R., H. Tran, C. Chen, H. Gong, K. K. Ho, and S. Lu. 2013. "Release of
663 extracellular ATP by bacteria during growth." *BMC Microbiology* 13:301-301.
664 doi: 10.1186/1471-2180-13-301.

665 Mohammed, T. A. 2013. "Isolation, Characterization and Application of Calcite
666 Producing Bacteria from Urea Rich Soils." MSc Islamic University of Gaza.

667 Mujah, D., M. A. Shahin, and L. Cheng. 2016. "State-of-the-Art Review of
668 Biocementation by Microbially Induced Calcite Precipitation (MICP) for Soil
669 Stabilization." *Geomicrobiology Journal* 34 (6):524-537. doi:
670 10.1080/01490451.2016.1225866.

671 Omoregie, A. I., N. Senian, Y. L. Phua, N. L. Hei, D. O. E. Leong, I. R. H. Ginjom, and
672 P. M. Nissom. 2016. "Ureolytic Bacteria isolated from Sarawak Limestone
673 Caves show High Urease Enzyme Activity comparable to that of *Sporosarcina*
674 *pasteurii* (DSM 33)." *Malaysian Journal of Microbiology* 12 (6):463-470.

675 Qabany, A. A. , K. Soga, and C. Santamarina. 2012. "Factors affecting efficiency of
676 microbially induced calcite precipitation." *Journal of Geotechnical and*
677 *Geoenvironmental Engineering* 138:992–1001.

678 Rahim, S., M. A. Z. Seyed, N. Ali, and C. O. Brendan. 2015. "Improving sand with
679 microbial-induced carbonate precipitation." *Proceedings of the Institution of*
680 *Civil Engineers - Ground Improvement* 168 (3):217-230. doi:
681 10.1680/grim.14.00001.

682 Reyes, R. G., L. L. M. A. Lopez, K. Kumakura, S. P. Kalaw, T. Kikukawa, and F.
683 Eguchi. 2009. "Coprinus comatus, a newly domesticated wild nutraceutical
684 mushroom in the Philippines." *Journal of Agricultural Technology* 5 (2):299-
685 316.

686 Robinson, P. K. 2015. "Enzymes: principles and biotechnological applications." *Essays*
687 *in Biochemistry* 59:1-41. doi: 10.1042/bse0590001.

688 Rowshanbakht, K., M. Khomehchiyan, R. H. Sajedi, and M. R. Nikudel. 2016. "Effect
689 of injected bacterial suspension volume and relative density on carbonate
690 precipitation resulting from microbial treatment." *Ecological Engineering* 89:49-
691 55. doi: 10.1016/j.ecoleng.2016.01.010.

692 Sakai, S., and H. Imachi. 2015. "Methanocellales." In *Bergey's Manual of Systematics*
693 *of Archaea and Bacteria*, edited by M. Garrity. George, R. Boone. David and W.
694 Castenholtz. Richard, 722. New York, United States: John Wiley & Sons, Ltd.

695 Sarayu, K., N. R. Iyer, and A. R. Murthy. 2014. "Exploration on the biotechnological
696 aspect of the ureolytic bacteria for the production of the cementitious materials--
697 a review." *Applied Biochemistry and Biotechnology* 172 (5):2308-2323. doi:
698 10.1007/s12010-013-0686-0.

699 Sethi, S., and S. Gupta. 2014. "Optimization Of Cultural Parameters For Cellulase
700 Enzyme Production From Fungi." *BIolife* 2 (3):989-996.

701 Soon, N. W. 2013. "Improvements In Engineering Properties of Tropical Residual Soil
702 By Microbially-Induced Calcite Precipitation." Master of Engineering Science,
703 Department of Civil Engineering, Universiti Tunku Abdul Rahman.

704 Soon, N. W., L. M. Lee, T. C. Khun, and H. S. Ling. 2014. "Factors Affecting
705 Improvement in Engineering Properties of Residual Soil through Microbial-

706 Induced Calcite Precipitation." *Journal of Geotechnical and Geoenvironmental*
707 *Engineering* 140 (5):1-11. doi: 10.1061/(asce)gt.1943-5606.0001089.

708 Stocks-Fischer, S., J. K. Galinat, and S. S. Bang. 1999. "Microbiological precipitation
709 of CaCO₃." *Soil Biology and Biochemistry* 31:1563-1571.

710 Tomczyk-Żak, K., and U. Zielenkiewicz. 2015. "Microbial Diversity in Caves."
711 *Geomicrobiology Journal* 33 (1):20-38. doi: 10.1080/01490451.2014.1003341.

712 van Paassen, L. A., R. Ghose, T. J. M. van der Linden, W. R. L. van der Star, and M. C.
713 M. van Loosdrecht. 2010. "Quantifying Biomediated Ground Improvement by
714 Ureolysis: Large-Scale BiogROUT Experiment." *Journal of Geotechnical and*
715 *Geoenvironmental Engineering* 136 (12):1721-1728. doi:
716 10.1061/%28ASCE%29GT.1943-5606.0000382.

717 Weaver, T., M. Burbank, A. Lewis, R. Lewis, R. Crawford, and B. Williams. 2011.
718 "Bio-Induced Calcite, Iron, and Manganese Precipitation for Geotechnical
719 Engineering Applications." *Geo-Frontiers 2011: Advances in Geotechnical*
720 *Engineering*, Dallas, Texas, USA.

721 Wei, S., H. Cui, Z. Jiang, H. Liu, H. He, and N. Fang. 2015. "Biom mineralization
722 processes of calcite induced by bacteria isolated from marine sediments."
723 *Brazilian Journal of Microbiology* 46 (2):455-464. doi: 10.1590/S1517-
724 838246220140533.

725 Whiffin, V. S. 2004. "Microbial CaCO₃ Precipitation for the production of Biocement."
726 PhD, School of Biological Sciences & Biotechnology, Murdoch University.

727 Whiffin, V. S., L. A. van Paassen, and M. P. Harkes. 2007. "Microbial Carbonate
728 Precipitation as a Soil Improvement Technique." *Geomicrobiology Journal* 24
729 (5):417-423. doi: 10.1080/01490450701436505.

730 Willey, J. M., L. M. Sherwood, and C. J. Woolverton. 2008. *Prescott, Harley, and*
731 *Klein's Microbiology*. 7th ed. New York, United States of America: McGraw-
732 Hill Higher Education.

733 Zhao, Q., L. Li, C. Li, M. Li, F. Amini, and H. Zhang. 2014. "Factors Affecting
734 Improvement of Engineering Properties of MICP-Treated Soil Catalyzed by
735 Bacteria and Urease." *Journal of Materials in Civil Engineering* 26
736 (12):04014094-04014104. doi: 10.1061/(ASCE)MT.1943-5533.0001013.

737

738 **List of Table**

739 **Table 1:** Bacteria identification based on 16S rRNA sequencing

740 **List of Figures**

741 **Figure 1:** The effect of different temperature on urease activity. Cultivation of
742 ureolytic bacteria in NB-medium in 250 mL conical flasks incubated at 20 to 45°C
743 for 24 hr. Vertical error bars indicate standard deviation. The analysis of variance
744 (ANOVA) with Tukey's procedure was used to compare the variance between
745 different groups with the variability within each of the groups. The level of
746 significance was set at 0.05 (*).

747 **Figure 2:** The effect of different pH on urease activity. Cultivation of ureolytic
748 bacteria in NB-medium in 250 mL conical flasks incubated for 24 hr. Vertical error
749 bars indicate standard deviation. The analysis of variance (ANOVA) with Tukey's
750 procedure was used to compare the variance between different groups with the
751 variability within each of the groups. The level of significance was set at 0.05 (*).

752 **Figure 3:** The effect of different incubation period on urease activity. Cultivation of
753 ureolytic bacteria in NB-medium in 250 mL conical flasks incubated for 24 hr.
754 Vertical error bars indicate standard deviation. The analysis of variance (ANOVA)
755 with Tukey's procedure was used to compare the variance between different groups
756 with the variability within each of the groups. The level of significance was set at
757 0.05 (*).

758 **Figure 4:** The effect of different urea concentration on urease activity. Cultivation of
759 ureolytic bacteria in NB-medium in 250 mL conical flasks incubated for 24 hr.
760 Vertical error bars indicate standard deviation. The analysis of variance (ANOVA)

761 with Tukey's procedure was used to compare the variance between different groups
762 with the variability within each of the groups. The level of significance was set at
763 0.05 (*).

764 **Figure 5:** Treatment of sand column using locally isolated bacteria, consortia,
765 positive and negative controls. (a) setup of sand columns before treated with
766 ureolytic bacteria and cementation solution (Left). (b) sand columns during treatment
767 with bacteria and cementation solution (right).

768 **Figure 6:** Treated sand removed from their respective columns. The biocement
769 specimens were left to cure for 14 days before being removed from their respective
770 moulds.

771 **Figure 7:** Surface strength of the biocemented sand samples. A pocket penetrometer
772 (ELE International, 38-2695) was used to test the surface strength. Vertical error bars
773 indicate standard deviation.

774

775

776

777

778

779

780

781

782

783

784 **Table 1:** Bacteria identification based on 16S rRNA sequencing

Isolate ID	Accession number	Closest match	Base pair	Similarity
NB33	KX212190	<i>Sporosarcina pasteurii</i> WJ-4 [KC211296]	1198	97%
LPB21	KX212191	<i>Sporosarcina pasteurii</i> fwzy14 [KF208477]	1385	97%
NB28	KX212192	<i>Sporosarcina pasteurii</i> WJ-5[KC211297]	1280	96%
NB30	KX212196	<i>Sporosarcina pasteurii</i> fwzy14 [KF208477]	1279	98%

785

786

787

788

789

790

791

792

793

794

795

796

797

798

799

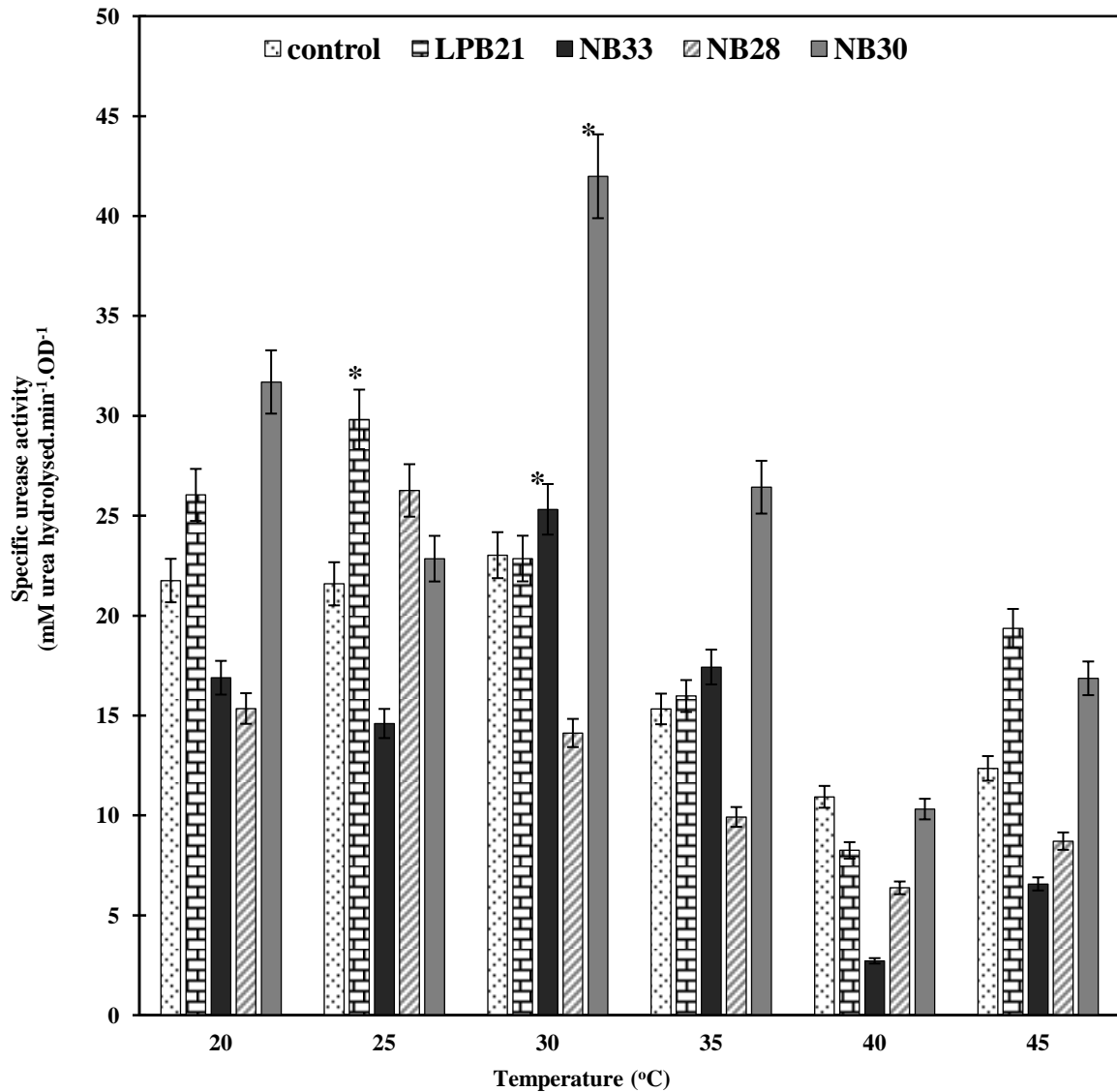
800

801

802

803

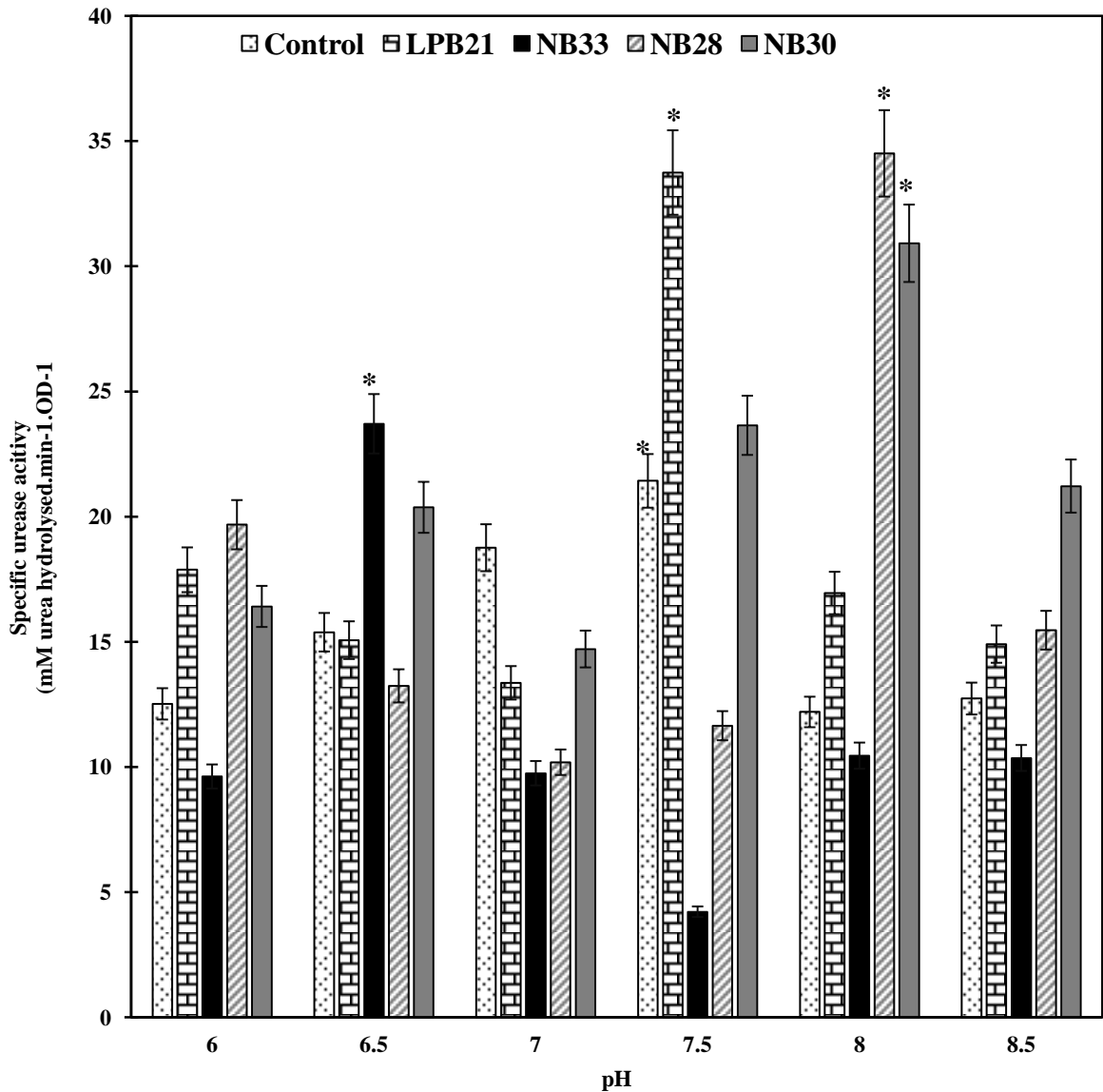
804



805

806 **Figure 1:** The effect of different temperature on urease activity. Cultivation of ureolytic
 807 bacteria in NB-medium in 250 mL conical flasks incubated at 20 to 45°C for 24 hr.
 808 Vertical error bars indicate standard deviation. The analysis of variance (ANOVA) with
 809 Tukey's procedure was used to compare the variance between different groups with the
 810 variability within each of the groups. The level of significance was set at 0.05 (*).

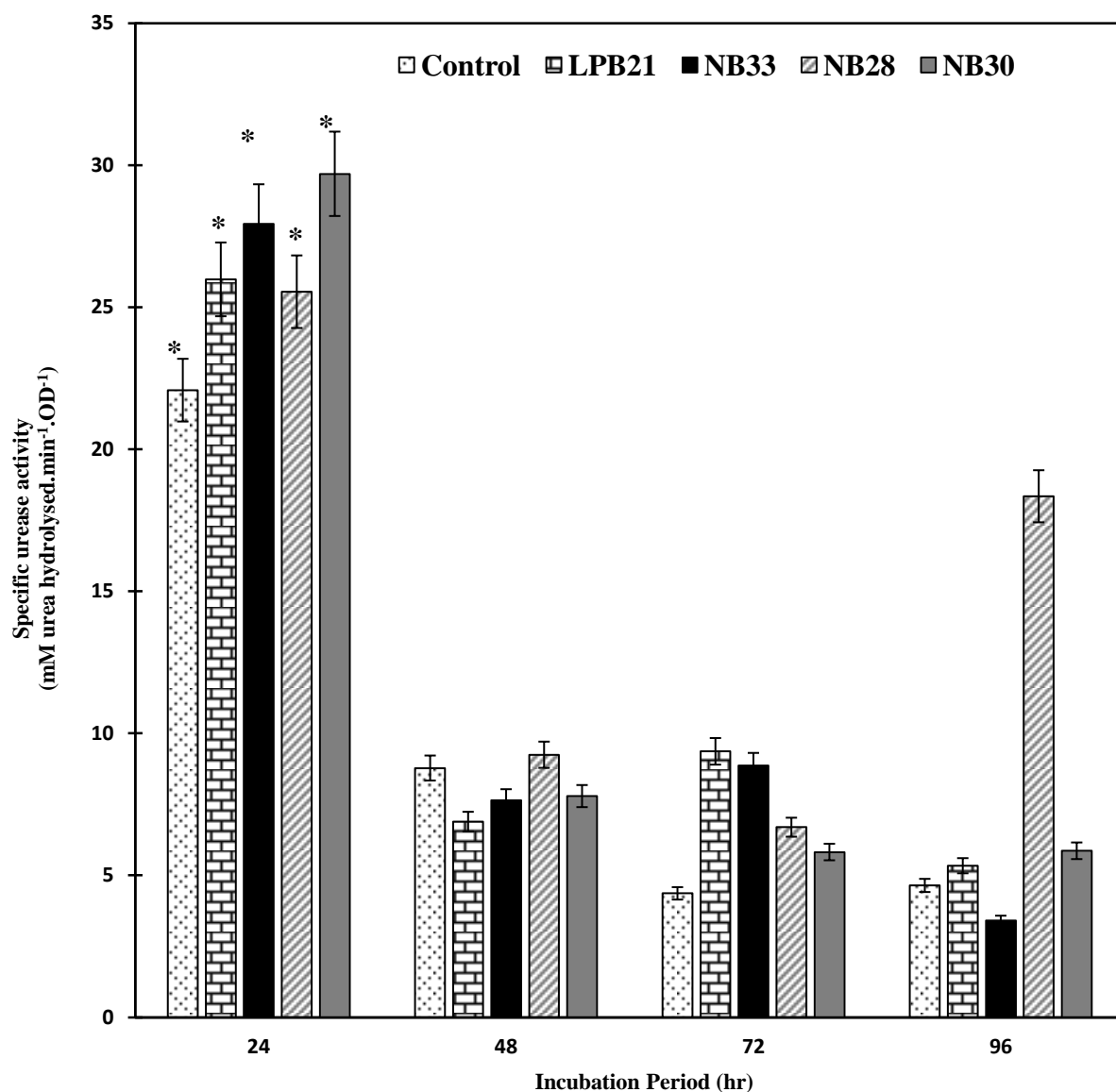
811



812

813 **Figure 2:** The effect of different pH on urease activity. Cultivation of ureolytic bacteria in
 814 NB-medium in 250 mL conical flasks incubated for 24 hr. Vertical error bars indicate
 815 standard deviation. The analysis of variance (ANOVA) with Tukey's procedure was used
 816 to compare the variance between different groups with the variability within each of the
 817 groups. The level of significance was set at 0.05 (*).

818

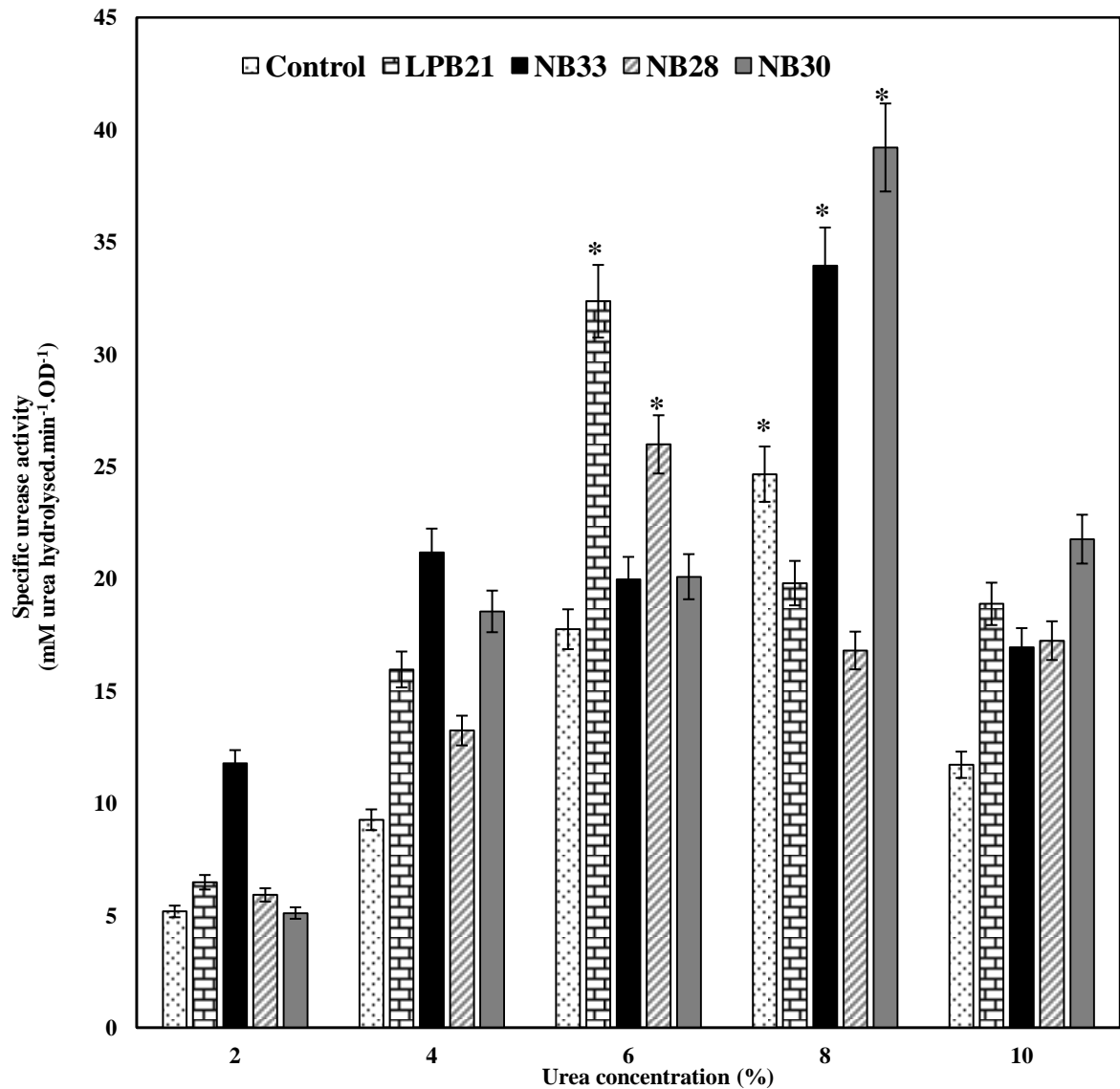


819

820 **Figure 3:** The effect of different incubation period on urease activity. Cultivation of
 821 ureolytic bacteria in NB-medium in 250 mL conical flasks incubated for 24 hr. Vertical error
 822 bars indicate standard deviation. The analysis of variance (ANOVA) with Tukey's
 823 procedure was used to compare the variance between different groups with the variability
 824 within each of the groups. The level of significance was set at 0.05 (*).

825

826



827
828

829 **Figure 4:** The effect of different urea concentration on urease activity. Cultivation of
830 ureolytic bacteria in NB-medium in 250 mL conical flasks incubated for 24 hr. Vertical
831 error bars indicate standard deviation. The analysis of variance (ANOVA) with Tukey's
832 procedure was used to compare the variance between different groups with the variability
833 within each of the groups. The level of significance was set at 0.05 (*).

834

835

836

(a)

(b)



837

838 **Figure 5:** Treatment of sand column using locally isolated bacteria, consortia,
839 positive and negative controls. (a) setup of sand columns before treatment with
840 ureolytic bacteria and cementation solution (Left). (b) sand columns during
841 treatment with bacteria and cementation solution (right).

842

843

844

845

846

847

848

849

850

851

852

853

854



Set 1: +ve control Set 2: Consortia Set 3: LPB21 Set 4: NB33 Set 5: NB28 Set 6: NB33

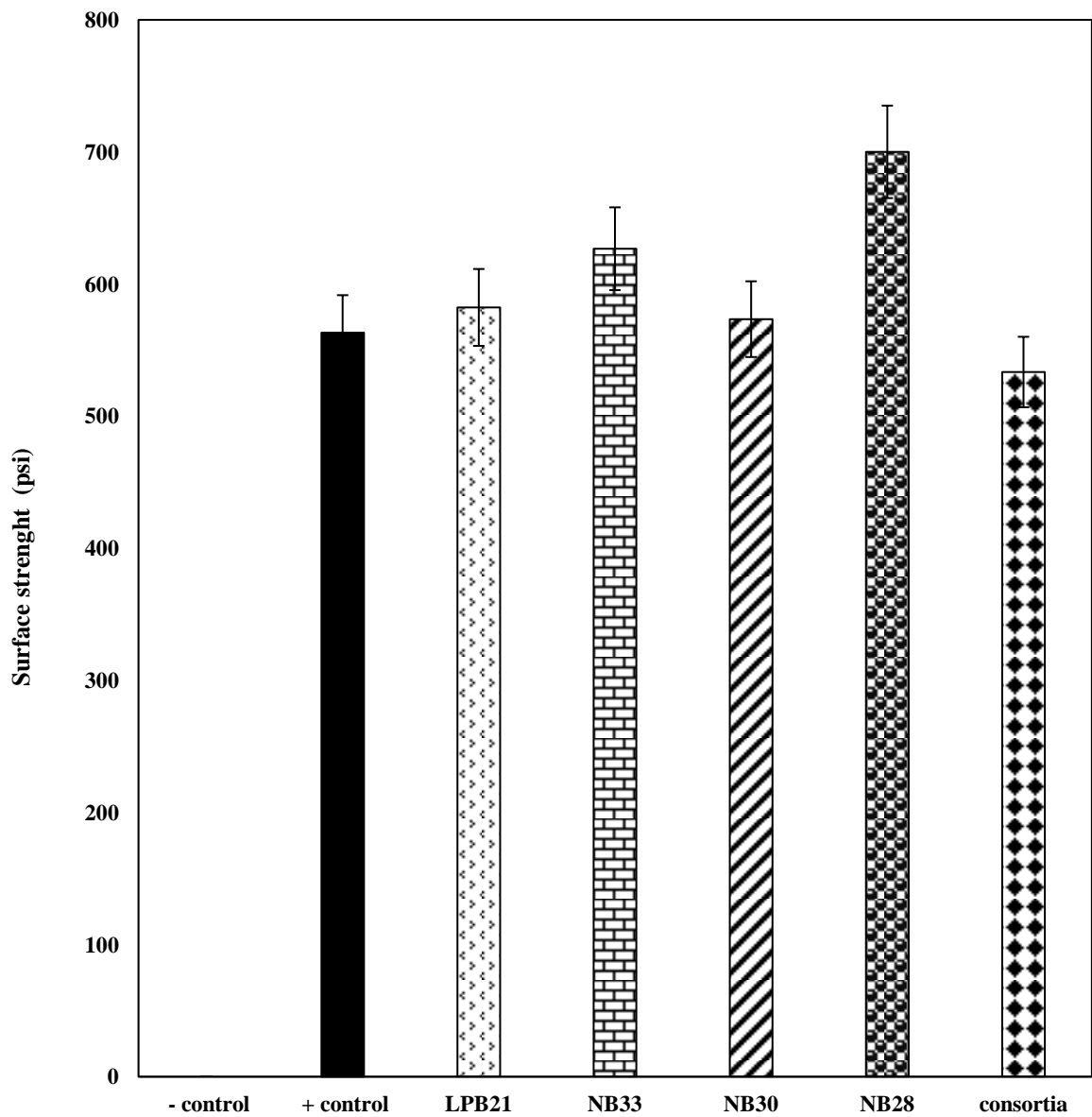
855

Figure 6: Treated sand removed from their respective columns. The biocement specimens were left to cure for 14 days before being removed from their respective moulds.

858

859

860



861 **Figure 7:** Surface strength of the biocemented sand samples. A pocket penetrometer (ELE
 862 International, 38-2695) was used to test the surface strength. Vertical error bars indicate
 863 standard deviation.

864

865

866