Effect of some biotic factors on microbially-induced calcite precipitation in cement mortar

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Abstract Sporosarcina pasteurii, a common soil bacterium has been tested for microbial treatment of cement mortar. The present study also seeks to investigate the effects of growth medium, bacterial concentration and different buffers concerning the preparation of bacterial suspensions on the compressive strength of cement mortar. Two growth media, six different suspensions and two bacterial concentrations were used in the study. The influence of growth medium on calcification efficiency of S. pasteurii was insignificant. Significant improvement in the compressive as well as the tensile strength of cement mortar was observed. Microbial mineral precipitation visualized by Scanning Electron Microscopy (SEM) shows fibrous material that increased the strength of cement mortar. Formation of thin strands of fillers observed through SEM micrographs improves the pore structure, impermeability and thus the compressive as well as the tensile strengths of the cement mortar. The type of substrate and its molarity have a significant influence on the strength of cement mortar.

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1. Introduction

Concrete is the most extensively consumed construction material on the Earth. However, it is vulnerable to deterioration, corrosion, and cracks and hence loses its strength over a period of time. The consequent damage and loss of strength requires immensely expensive remediation and repair. Many organic and inorganic treatments have been adopted but as of now,
Materials and methods

S. pasteurii (ATCC 6453) was used throughout the study as the test organism, whereas E. coli DH5a (ATCC 53868) was used as a negative control for comparing the effect of the presence of microorganisms on cement mortar.

Growth media and bacterial concentration

S. pasteurii was grown in two different media, viz. (i) ammonium-sulfate and yeast extract (NH4-YE medium) – a specific broth for S. pasteurii, and (ii) Nutrient Broth (NB medium), a general medium that stimulates cellular metabolism. E. coli DH5a cells were grown in Lysogeny Broth (LB) media. Compositions of the three media (per liter) were as follows:

- **NH4–YE medium**: 20 g yeast extract; 10 g di-ammonium sulfate [(NH4)2SO4]; 0.13 M tris buffer (pH = 9.0); 20 g agar.
- **NB medium**: 3 g Nutrient Broth; 20 g urea [(NH4)2CO]; 10 g ammonium chloride [NH4Cl]; 25.2 mM sodium bicarbonate [NaHCO3].
- **LB medium**: 10 g tryptone; 5 g yeast extract; 10 g NaCl.

S. pasteurii cells were grown in the first two media separately, whereas E. coli DH5a cells were grown in LB medium. The cells were allowed to multiply overnight with 1% inoculum from freshly-prepared primary cultures. Cells were harvested by centrifugation at 3000g for 10 min and washed twice with phosphate buffer. Cell pellets were suspended in buffer to obtain a high-density stock of cells; and the cell concentrations were determined by recording absorbance at 600 nm. The final cell concentrations for different treatments were adjusted by diluting portions of these stocks on the basis of hemocytometer counts.

Measurement of turbidity or optical density (OD) is not a direct measurement of bacterial numbers, but an indirect measurement of cell biomass that includes both living and dead cells. Hence, to quantify viable cells, a plate-count method was used and cell counting by hemocytometer under microscope was also performed to have an accurate estimate.

Solutions for bacterial suspensions

Bacterial cells in appropriate number were suspended in water and phosphate buffered solutions containing varying concentrations of urea (CH2N2O) as a substrate for bacterial activity, and calcium chloride (CaCl2·2H2O) as a source of Ca2+ ion. A total of six solutions (S0 to S5) were used for assessing the influence of molarity of salts and buffer on the calcification efficiency of S. pasteurii. Compositions of these solutions are given in Table 1. The solutions S2 and S3, containing only the phosphate buffer in different proportions, were used as

Only little success has been achieved. As a result, there is an impending need for technology development and improvement to meet the demand of society. Interdisciplinary research at the confluence of microbiology and civil engineering has already proven its potential by utilizing the biological activity of living cells (Boquet et al., 1973; Stocks-Fischer et al., 1999; Newnham, 1997; Ramakrishnan et al., 2005; Bachmeier et al., 2002) for improving the quality of building materials.

One of the important characteristics of cells, which is not confined to human beings alone, is the ability of mineral precipitation for the formation of bones and teeth. Other living organisms, including some of the bacteria, also have the ability to form bones and teeth-like material such as natural pearls and shells. This phenomenon of biomineralization has inspired some investigators for developing innovative high-performance composites for construction applications (Ramakrishnan et al., 2005; Bachmeier et al., 2002).

The bacterial remediation technique has been proposed for repairing structures of historical importance to preserve their esthetic value, because the conventional techniques, such as epoxy injection, cannot be used to remediate cracks in these structures (Ramachandran et al., 2001; Bang and Ramakrishnan, 2001). The cracks in granite were effectively repaired structures of historical importance to preserve their esthetic value, because the conventional techniques, such as epoxy injection, cannot be used to remediate cracks in these structures (Ramachandran et al., 2001; Bang and Ramakrishnan, 2001). The cracks in granite were effectively repaired by immobilization of S. pasteurii (DeMuynck et al., 2010) has provided an in-depth comparison of different approaches with due attention to the background information.

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controls for highlighting the requirement of substrate and calcium source in bacterial calcification.

2.4. Preparation of bacterial mortar and specimens

Ordinary Portland cement Type I, conforming to ASTM (2008), was used for experiments. Locally available clean red sand conforming to ASTM (2008) was used as fine aggregate. Suspensions of bacterial cells in various solutions, as described above, were used for preparation of mortar specimens. Physical properties of the materials are given in Table 2.

A cement to sand ratio of 1:3 (w/w), and a solution to cement ratio of 0.6 (w/w) were maintained throughout the experiments. Cubes of 50 mm size were prepared for determining the compressive strength, whereas briquettes were used for determining the tensile strength of mortar. Bacterial cells at concentrations of 10⁸ and 10⁹ per ml in different solutions were added to the mortar. No additional food material, except that present in the diluted cultures, was supplemented in the mortar during mixing. Dry cement and sand were thoroughly mixed and then the solution containing bacteria was added. The solution without bacterial cells was used to prepare the control mortar blocks. Cubes were cast and compacted in a vibration machine, and all specimens were de-molded after 24 h of casting and then water-cured at room temperature (30 ± 2 °C) until compression and tension testing was done after 28 days. The compressive strength tests were performed on mortar cubes using 300 kN Toni Technik – Toni NORM compression testing machine according to ASTM (2007). The tensile strength tests were performed on mortar briquettes using 1000 lbs Zero-Max tensile testing machine in accordance with ASTM (2012). The specimens under test are shown in Fig. 1. SEM examinations were made on the broken sample pieces collected from the tested mortar cubes. Three cubes were tested for each of the treatments and the controls.

The five-character specimen IDs consisted of three letters followed by two digits. The first letter ‘M’ stands for mix; the second letter represents the bacterial genotype (D: E. coli DH5α; S: S. pasteurii; N: No bacterium – control); and the third letter represents the bacterial growth medium (N: No medium; L: LB medium; B: Nutrient Broth; Y: NH₄YE medium). Of the two digits, the first stands for the log value of the bacterial cell concentration; whereas the second represents the numeric in the solution ID.

2.5. Treatments

Experiments were designed to assess the effect of: bacterial growth medium, bacterial cell concentration and solutions on the compressive strength of the cement mortar. In all, eleven treatments were maintained, based on the following parameters:

(i) Bacterial strain: S. pasteurii
(ii) Bacterial growth media: NB and NH₄–YE, as described above
(iii) Bacterial cell concentrations: 10⁸ and 10⁹ cells/ml.
(iv) Solutions used for preparing bacterial suspensions: S0–S5, as described in Table 1.

2.6. Controls

Three types of control mortar specimens were prepared:

(i) Cement mortar in solutions but no bacteria (Specimens: MNN00, MNN01 and MNN02),
(ii) Cement mortar with E. coli DH5α (10⁸ and 10⁹ cells/ml), in solution S0 (Specimens: MDL80 and MDL90); in solution S1 (Specimens: MDL81 and MDL91); and in solution S2 (Specimens: MDL82 and MDL92).

Thus, nine different control specimens were prepared for studying the influence of different chemicals used for bacterial suspension and the presence of microorganisms in mortar on its strength. The effect of dead cells of E. coli DH5α and S. pasteurii was supposed to be the same.

2.7. Image analysis

Scanning Electron Microscope (FEL-Spectra) was used for detecting changes in microstructure of the formed phases. The broken specimens, collected after the compressive strength tests, were used for preparing thin sections by grinding the
solid and sound mortar chips with carborundum (silicon carbide), washed thoroughly in water and dried at a temperature of 90–100 °C (Winchell, 1949). The thin section slides were scanned by FEL-Spectra microscope. Three sections were prepared for each mix.

3. Results and discussion

The effect of varying bacterial concentrations and solutions on the compressive strength of cement-sand mortar was significant. The compressive strength of 28-days cured mortar cubes under different treatments is given in Table 3.

3.1. Control specimen

Calcification potential of a bacterium emanates from its ability to breakdown urea enzymatically (DeMuynck et al., 2010; Tittelboom et al., 2010). E. coli DH5α is known to have no urease activity; therefore, it was used as a control strain for exclusive assessment of the influence of experimental
parameters on the activity of *S. pasteurii* strain harboring active urease gene (Kim and Spizizen, 1985; Stocks-Fischer et al., 1999; Tiwari et al., 2014).

The influence of the bacterial concentration of *E. coli* DH5α and suspension on the compressive strength of the mortar is shown in Fig. 2, which allows for the following deductions:

(i) The increase in bacterial concentration from $10^8$ to $10^9$ cells/ml reduced the compressive strength of the mortar cubes slightly, the difference being nominal; an average of test results of the two concentrations was used for making comparisons. In solution S0, compressive strength of the mortar with bacteria (MDL80, MDL90) was reduced by 1.9%, whereas in the presence of solutions S1 and S2, the compressive strength of mortar containing bacteria (MDL81, MDL82, MDL91, MDL92) was reduced by 1.2% and 2.0% respectively. It indicates that the addition of blank strain *E. coli* DH5α had almost no effect on the compressive strength of the cement mortar.

(ii) The effect of solution S1 (urea: 0.2 M; CaCl$_2$: 0.2 M) on the compressive strength of cement-mortar could be assessed by comparing the compressive strength of (a) MNN00 (no bacteria; S0) with MN001 (no bacteria; S1), (b) MDL80 (bacteria: $10^8$ cells/ml; S0) with MDL81 (bacteria: $10^8$ cells/ml; S1), and (c) MDL90 (bacteria: $10^9$ cells/ml; S0) with MDL91 (bacteria: $10^9$ cells/ml; S1). Application of solution S1 in cement mortar resulted in an average reduction of 10.0% in the compressive strength of mortar.

(iii) Similarly, the effect of using solution S2 (No substrate solution; 0.1 M phosphate buffer, pH 7.8) on the compressive strength of cement-mortar could be assessed by comparing the compressive strength of (a) MNN00 (no bacteria; S0) with MNN002 (no bacteria; S2), (b) MDL80 (bacteria: $10^8$ cells/ml; S0) with MDL82 (bacteria: $10^8$ cells/ml; S2), and (c) MDL90 (bacteria: $10^9$ cells/ml; S0) with MDL92 (bacteria: $10^9$ cells/ml; S2). The addition of solution S2 in cement mortar resulted in an average reduction of 11.1% in the compressive strength of mortar.

The observed changes in compressive strength by incorporation of *E. coli* DH5α were less than 2% in almost all cases and there was no consistent improvement/degradation in strength due to the addition of this bacterium. The bacterium

| Table 3  | Compressive strength of cement mortar under different treatments. |
|----------|---------------------------------------------------------------|
| Mix set  | Bacteria | Medium | Concentration (cells/ml) | Suspension | Compressive strength (MPa) |
| Control specimens |  |  |  |  |  |
| MNN00 | – | – | – | – | S0 | 31.7 |
| MNN01 | – | – | – | – | S1 | 28.4 |
| MNN02 | – | – | – | – | S2 | 28.2 |
| MDL80 | DH5α | LB | $10^8$ | – | S0 | 31.2 |
| MDL90 | DH5α | LB | $10^9$ | – | S0 | 31.0 |
| MDL81 | DH5α | LB | $10^8$ | – | S1 | 27.9 |
| MDL91 | DH5α | LB | $10^9$ | – | S1 | 28.2 |
| MDL82 | DH5α | LB | $10^8$ | – | S2 | 27.8 |
| MDL92 | DH5α | LB | $10^9$ | – | S2 | 27.5 |
| Specimens containing *S. pasteurii* |  |  |  |  |  |
| MSB91 | SP | NB | $10^9$ | – | S1 | 38.8 |
| MSB92 | SP | NB | $10^9$ | – | S2 | 26.8 |
| MSY91 | SP | NH$_4$–YE | $10^9$ | – | S1 | 38.9 |
| MSY92 | SP | NH$_4$–YE | $10^9$ | – | S2 | 26.9 |
| MSB81 | SP | NB | $10^8$ | – | S1 | 37.0 |
| MSB82 | SP | NB | $10^8$ | – | S2 | 27.0 |
| MSY81 | SP | NH$_4$–YE | $10^8$ | – | S1 | 37.4 |
| MSY82 | SP | NH$_4$–YE | $10^8$ | – | S2 | 26.6 |
| MSY83 | SP | NH$_4$–YE | $10^9$ | – | S3 | 26.5 |
| MSY84 | SP | NH$_4$–YE | $10^8$ | – | S4 | 39.6 |
| MSY85 | SP | NH$_4$–YE | $10^9$ | – | S5 | 38.7 |

*SP = *S. pasteurii*; LB = Lysogeny Broth; NB = Nutrient Broth.

Figure 2 Influence of cell concentration of *E. coli* DH5α and solutions on compressive strength of mortar.
proved ineffective in the presence of the substrate solution also. In previous studies also, *E. coli* has shown no calcification potential (Ghosh et al., 2005). Bachmeier et al. (2002) assessed the calcification potential of some bacteria and used *E. coli* (pBR322) as a control, because it has no urease activity. This suggests that the selection of an active bacterial strain is important for improving the mortar strength.

### 3.2. Effect of growth medium

The influence of the growth medium for *S. pasteurii* at a cell count of $10^8$ cell/ml on the compressive strength of mortar is shown in Fig. 3. For solution S1 (urea: 0.2 M; CaCl$_2$: 0.2 M), improvement in the compressive strength of mortar with bacteria grown in both MSB81 and MSY81 media was closely similar (Average = 17.4%). As discussed above, addition of solutions results in 10–11% reduction in the compressive strength of mortar and therefore the gain in strength as a result of calcification was significant. When urea and calcium chloride are degraded and utilized due to bacterial activity, they no longer remain as impurities in the mortar and contribute toward calcification. Growth media were used in this study only for cultivation and maintenance of the bacterial cultures. Before adding to the mortar, the bacterial cells were pelleted out and washed with buffer, leaving very little amount of the medium to go into the mortar. Thus there was no significant carry over effect of the medium on the subsequent activity of *S. pasteurii* cells in the mortar. It is generally believed that the improvement in compressive strength of mortar is due to the calcite deposition on the microorganism cell surfaces and within the pores of cement–sand matrix, which plugs the pores within the mortar (Stocks-Fischer et al., 1999; Newnham, 1997; Ramakrishnan et al., 2005).

For solution S2 (no substrate solution; 0.1 M phosphate buffer, pH 7.8), there was an average decrease of 15.5% in the compressive strength of the mortar, using bacteria grown in either of the two media possibly because of the absence of substrate and the calcium source due to which the phosphate buffer acted as an impurity.

### 3.3. Effect of bacterial cell concentration

The effect of concentration of *S. pasteurii* cells in the solutions S1 and S2 used for making the mortar has been shown for cells grown on NB medium (Fig. 4) and for cells grown on NH$_4$–YE medium (Fig. 5).

In general, the compressive strength improved with increase in bacterial cell count for the solution S1 (Fig. 4). The enhancement in compressive strength for increase in concentration of bacteria from $10^8$ to $10^9$ cells/ml was 4.9% for bacteria grown in NB medium, and 4% for bacteria grown in NH$_4$–YE medium (Fig. 5), showing that the medium of growth for bacteria had no significant effect on its calcification efficiency in the mortars. However, the bacterial cell count had an impact on calcification level in mortars (Pacheco-Torgal and Labrincha, 2013). Ghosh et al. (2009), using *Shewanella* species, found that highest compressive strength of the mortar was attained with a cell count of $10^9$ cells/ml. However for healing of cracks, LeMetayer-Leval et al. (1999) have suggested a cell concentration of $10^6$ cells/ml to be optimal.

The increase in the concentration of bacteria from $10^8$ to $10^9$ cells/ml, suspended in solution S2, caused slightly more reduction in the compressive strength which might be due to increase in the biomass.
3.4. Effect of solutions in suspension

In addition to S1 and S2, three more suspensions (S3 to S5) were studied for bacterial growth in NH₄-YE medium. The effect of solutions on the compressive strength of mortar has been plotted in Fig. 6 for a bacterial concentration of 10⁸ cells/ml. All of the specimens (MSY81, MSY82, MSY83, MSY84, and MSY85) considered in figure are for S. pasteurii grown in NH₄-YE medium except the control (MNN00).

Figure shows that the maximum compressive strength of the mortar [24.9% more than the control (MNN00)] was attained for suspension S4 (specimen MSY84), containing phosphate buffer. The strength of mortar for suspensions S1 and S5 was 18.0% and 22.1% higher respectively. The reduction in the molarity of substrate solutions S4 and S5 could be one of the reasons for increase in the compressive strength of the mortar mixes using these suspensions (specimens MSY84 and MSY85). Moreover, phosphate buffers are known to enhance the activity of bacteria. With lower salt content, level of un-degraded residues acting as impurities would be low, causing little change in behavior of the cement mortar.

Microbial calcite formation is directly dependent on the availability of urea as a substrate for urease activity and Ca⁺² derived from an appropriate source (DeMuynck et al., 2010). As long as the whole or the major part of these salts is utilized by microbial activity, their increasing concentration would result in a greater calcification and a corresponding increase in the mortar strength. However, the unutilized amount of salts would obviously have a negative impact.

Figure 6  Influence of the solutions on compressive strength of mortar for S. pasteurii at a concentration of 10⁸ cells/ml.

Figure 7  SEM images of cement mortar specimen MNN00 (control).
3.5. SEM micrographs

The micrographs of the control plain cement mortar (MNN00) and of a better-performing bacteria-modified mortar (MSY84) are presented in Figs. 7 and 8 respectively. The selection of specimen MSY84 is based on its high compressive as well as the tensile strength. The direct tensile strength of specimens MNN00 and MSY84, determined using standard briquettes (ASTM, 1985), were found to be 2.7 and 3.7 MPa respectively. Thus the compressive strength of the mix MSY84 increased by 24.9% and its tensile strength by 37%, as compared to the plain cement mortar.

Figs. 7 and 8 indicate that in a mortar made with a $10^8$ cells/ml of \textit{S. pasteurii} in suspension S4, the pores are better filled with narrow strands of filler (Fig. 8); and a higher modification in pore size distribution is noticed, whereas no such filler material is observed in the micrographs of the control sample of plain mortar. This confirms the increase in compressive and direct tensile strengths of mix MSY84. The qualitative assessment of SEM images shows that the presence of narrow strands of filler is beneficial for modification of the porosity and pore-size distribution of cement mortar.

In congruence with our findings, several studies addressing the microbial calcification have shown deposition of calcite crystal in cement mortars in SEM images (DeMuynck et al., 2008; Ghosh et al., 2009), which demonstrate the real possibility of application of this technique in improving the cement mortars.

4. Conclusions

Addition of \textit{E. coli} DH5\textalpha, a control bacterium, causes almost no effect on the compressive strength of the mortar, which suggests that the choice of microorganism is important for improving the compressive strength of mortars.

This study suggests that the strength of the \textit{S. pasteurii} incorporated mortar increases due to mineral precipitation by the bacterial activity. Addition of \textit{S. pasteurii} has a positive effect on the compressive as well as the tensile strength of the mortar. The influence of growth medium on calcification efficiency of \textit{S. pasteurii} is insignificant. The type of substrate solution and its molarity have a significant influence on the strength of the mortar.

SEM examination reveals the growth of fibrous filler material within the pores due to the precipitation by \textit{S. pasteurii},
which confirms the increase in strength of the mortar observed. This improvement is attained by the modification of porosity and pore size distribution of the cement mortar it generates.

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