

Performance of Self-Healing Bacterial Concrete to Repair for Micro Cracking

Shaik Shahul ¹, G. Nagalakshmi ²

¹ PG Scholar, Department of Civil Engineering, Chalapathi Institute of Technology, Mothadaka, Andhra Pradesh, India, 522016

² Assistant Professor Department of Civil Engineering, Chalapathi Institute of Technology, Mothadaka, Andhra Pradesh, India, 522016

Abstract: The employment of carbonate-producing bacteria as a new approach to enhance the qualities of concrete has attracted a lot of interest since it is thought to be innocuous to the environment, natural, and maybe advantageous. This is a result of the favorable implications attached to these traits. The use of microbially induced carbonate precipitation as a remedy for several problems affecting concrete, such as fracture healing, reduction and change of porosity and permeability, and more, has been the subject of much investigation. Concrete crack healing is one of the topics that have been researched. Additionally, it has been shown that the procedure of bacterial carbonate precipitation, also known as bio deposition, contributes to the development of concrete's compressive strength. There has not yet been a thorough investigation of the research relating to the appropriate bacterial solution dose and its effect on concrete durability. This might occur as a result of the project not receiving enough time to complete it. As a result, it has been decided that an investigation will be conducted in order to determine the proper quantities of bacterial solution required for concrete. To do this, several concrete cube samples will be made using varying amounts of bacterial solution, such as 15 ml, 30 ml, 45 ml, 60 ml, and 75 ml, respectively. These quantities will be added to the appropriate moulds. This will allow you to determine the proper dosage of the bacterial solution to employ. In order to determine the optimal dosage that should be used, these various samples are also put through a battery of tests using a variety of laboratory techniques, such as the properties of materials, slump cone test, a compressive strength testing machine, an ultrasonic pulse

velocity test, plate count cells, and scanning electron microscopes, Rapid Chloride Penetration Test (RCPT), Acid attack test.

Keywords: Bacterial Concrete, Crack Repair, Bacterial Carbonate, Compressive Strength, Rapid Chloride Penetration Test (RCPT), Ultrasonic Pulse Velocity, Plate Count Cells, Scanning Electron Microscopes.

1. INTRODUCTION

Carbonate-producing bacteria have attracted lots of interest as a promising, natural, environmentally friendly novel technique to improvement of concrete characteristics. Considerable research has been conducted on utilizing microbial-induced (Dilja Rose Joseph, LifeJohn 2017)[10] carbonate precipitation to mitigate several concrete problems such as crack repair, reduction and modification of porosity, and permeability. Furthermore, bacterial carbonate precipitation (bio deposition) has shown positive influences on compressive strength improvement of concrete and also, it also reduces water absorption and carbonation of concrete as an alternative surface treatment (Pipat Termkhajornkit et.al. 2009) [1]. As part of metabolism, some bacteria (S. Sunil Pratap Reddy et.al. 2010) [2] (C.C. Gavimath et.al. 2012) [4] produces enzyme urea which catalyzes the hydrolysis of urea to generate carbonate ions without an associated production of protons which leads to CaCO_3 precipitation in presence of calcium ions. Therefore, bacteria cells not only provide a nucleation site for CaCO_3 precipitation due to their negatively charged cell walls, but also create an alkaline environment inducing further growth of CaCO_3 crystals. One ml of urea is hydrolyzed intra-cellularly to 1 ml of ammonia and 1 ml of carbonate, which is presented in Eq. (1). According to Eq. (2), carbonate hydrolyzes to ammonia and carbonic acid. Eqs. (3) and (4) demonstrate former

products subsequently equilibrate in water to form bicarbonate, ammonium, and hydroxide ions. The latter causes pH increase resulting in the formation of carbonate ions [Eq. (5)], which in the presence of soluble calcium ions precipitate as CaCO_3 [Eq. (6)]. Eq. (7) is the overall reaction, which demonstrates that ammonium and calcium carbonate are the products of added urea and calcium to the system

1.1. Aim of the Work

- Bacteria belonging to the bacillus family should be mixed in with the bacteria that are already present in order to produce bacterial concrete (*Bacillus subtilis*).
- to establish the ideal number of bacteria that should be employed in the manufacturing of bacterial concrete. The goal of this research is to achieve this.
- The method of serial dilution was used in order to determine the total number of viable bacterial cells.
- An ultrasonic pulse velocity test will be used for the purpose of determining whether or not there are openings.
- We use SEM to ascertain whether or not there are voids inside the internal structure of the concrete being examined.
- Investigate bacterial activity from the point of view of chemistry.
- Investigate how the change affects the characteristics of the concrete, such as how its compressive strength and permeability are affected by the transformation.

2. MATERIALS

Cement: For the sake of this specific experiment, Portland cement of the standard 53 grade kind, which can be procured with relative ease, was used. The cement that was utilised was subjected to a battery of tests in accordance with IS: 4031-1988 to assess its different properties. The findings indicated that the cement was compliant with the various criteria of IS: 12269-1987 and had a specific gravity of 3.15. The examinations were carried out in accordance with the recommendations that were included in IS: 4031-1988.

Fine Aggregate: In the course of our investigation, we made use of GODAVARI sand, which provided conclusive evidence that zone III is relevant in light of IS-383's requirements. It was established that the specific gravity of sand is 2.386, giving it a value.

Coarse Aggregate: The component of concrete known as the coarse aggregate is the component that is both the most permeable and the most robust of all of the components that make up concrete. It is possible that drying shrinkage and other dimensional changes brought about by the passage of moisture may be decreased to a level that is more controllable if coarse aggregate is used. During the course of our experiment, we made use of an aggregate that was able to get through a 20mm IS-Sieve but was still able to be captured on a 12.5mm sieve. This allowed us to have the best of both worlds. Because of this, we were able to enjoy the benefits of both settings. After doing more research, it was discovered that the aggregate had a specific gravity of 2.994.

Bacteria: *Bacillus Subtilis* as shown in figure.1 is selected because it produces Calcium Carbonate which is main component for cement.

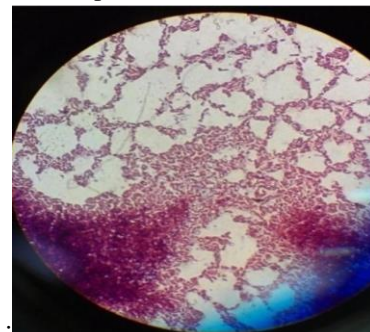


Figure 1: *Bacillus Subtilis*

Table 1: Physical Properties of Cement

Sr. No.	Particulars	cement
1	Grade	53
2	Specific gravity	3.15
3	Standard consistency %	32
4	Fineness	3
5	Initial setting time	30 min
6	Final setting time	600 min

Table 2: Properties of Fine Aggregate.

Sr. No.	Particulars	cement
1	Specific gravity	2.386
2	Fineness modulus	3.06
3	Bulk density	1451kg/m ³
4	Voids ratio	0.644

Table 3: Properties of Coarse Aggregate

Sr. No.	Particulars	cement
1	Specific gravity	2.994
2	Fineness modulus	7.17
3	Bulk density	1594 kg/m ³
4	Voids ratio	0.878

3. METHODOLOGY

The mix ratio of M30 is 1:1.97:2.96 having the following different mixes confirming with IS: 10262-2009 [12].

3.1 Cultivation of Bacteria

Bacillus Subtilis, a pure bacteria culture, is kept on nutrient agar slants. On nutrient agar slants, it develops irregular dry white colonies. Two colonies of bacteria (Harshali J et.al 2016) are seeded into 350 mL of nutrient in a 500 mL conical flask as shown in figure 2 and incubated at 37 degrees Celsius using a 150 rpm orbital shaker incubator. Peptone, NaCl, and yeast extract are the main ingredients in the bacterial culture medium.



Figure 2: Bacterial Solution

3.2 Slump Cone test

The container is filled with bacterial concrete in three layers, whose workability is to be examined, and the metal plate, which serves as the

base, is placed on a smooth surface. A standard 16 mm (5/8 in) diameter steel rod, rounded at the end, is used to tamper each layer 25 times. The top surface of the mould is struck off (levelled with mould top aperture) by screening and rolling motion of the tamping rod once the microbiological concrete is entirely filled. The Moulds remained securely against their base throughout the procedure, preventing them from moving owing to the pouring of concrete through handles or foot-rests. The cone is gently and carefully removed vertically when the filling has been done and the concrete has been levelled, unsupported bacterial concrete will now slump as shown in figure 3.2. The slump is measured by inserting the cone next to the slumped concrete and tamping the area with the tamping rod. With scale, the height of concrete is determined to be 110mm for conventional concrete and 50mm for bacterial concrete, indicating a decrease in height to that of Moulds.

3.3 Compressive Strength Test

After withdrawing the specimen from the water for the required curing time, the surplus water was scraped from the surface. The bearing surface of the testing machine needs to be cleaned. (M. Manjunath et.al 2014) [7] The numerous sample specimens were placed in the machine one after the other, with the load applied to the opposing sides of the cube cast. The specimen is perfectly placed as shown in figure 3.3 on the machine's base plate. The force is delivered gradually, without shock, and at a rate of 5.2 KN/sec until the specimen fails. The maximum load was recorded, as well as any unexpected characteristics in the type of failure. Concrete cubes in the CTM machine, both before and after crushing. Each bacterial concrete sample's readings 15ml, 30ml, 45ml, 60ml and 75ml were taken each time after curing interval of 7 days, 14 days and 28 days.

3.4 Plate count test

An experiment using the plate count technique was carried out in order to ascertain the overall number of viable cells that are present in a culture of bacteria. This was accomplished by counting the number of plates used in the experiment. This approach is used to determine the number of cells that are capable of reproducing under certain circumstances in order to offer an answer to the problem that the research was attempting to investigate.

3.5 Scanning Electron Microscope (SEM)

An investigation of the calcium carbonate crystals that had been deposited was carried out with the use of a scanning electron microscope. The shape of the crystals as well as their mineralogical make-up were the primary focuses

of this investigation (SEM). In order to create the SEM micrographs, a Jeol JSM 5600 LV type Philips XL 30 that was connected to an EDX unit was used. The voltage that was used to accelerate was changed to 30 kilovolts. The resolution was changed to W, and the magnification was increased as high as 400,000 times its initial size (3.5 nm). After having a layer of carbon applied to their surface, the samples were subsequently given a covering of gold to cover their exposed areas.

3.5 Rapid Chloride Penetration Test (RCPT)

The RCPT is carried out by keeping track of the quantity of electrical current that flows through a sample that is 6 hours long and has aspects of 50 millimetres in thickness and 100 millimetres in circumference. The sample also has dimensions of 50 millimetres in thickness and 100 millimetres in circumference. In most laboratories, the sample is prepared as a slice that is taken from the centre of a cylinder. The voltage between the two ends of the sample is kept at a constant direct current of sixty volts during the whole of the test. The first lead is immersed in a solution containing 3.0 percent sodium chloride (NaCl), and the second lead is immersed in a solution containing 0.3 M sodium hydroxide (NaOH). The charge that is permitted to travel through the sample is used to offer a qualitative evaluation of the permeability of the concrete, as stated in the Table. This is done using the charge that is allowed to go through the sample.

4. RESULTS AND DISCUSSIONS

4.1 Compressive Strength Test results

Compressive strength of concrete cube was carried out confirming IS: 516-1959 after curing period of 7,14,28 days. The results so obtained are tabulated below with their respective graph.

Table 4: Compressive Strength results for 7 days in Mpa

Type of concrete	Compressive strength of concrete after 7days			Mean
	Sample 1	Sample 2	Sample 3	
M1	20.86	20.70	21.16	20.90
M2	25.89	30.20	29.17	28.42
M3	29.76	33.65	32.48	31.96
M4	33.84	32.74	33.27	33.28
M5	32.58	37.27	34.85	34.89
M6	35.82	37.67	35.70	36.39

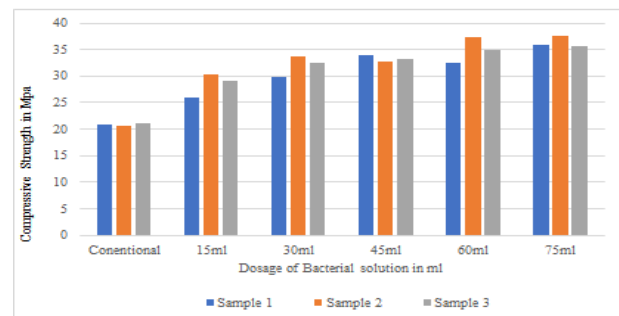


Figure 3: Compressive Strength Graph for 7 Days

Table 4 and graph 3 explains about the strength for 7 days of curing the specimens. The five samples of every solution i.e., 15ml, 30ml, 45ml, 60ml, and 75ml are tested for 7 days and 28 days. As dilution of bacteria increases the strength increases. By adding the bacillus subtilis the strength has gained nearly to target mean strength even in 7 days

Table 5: Compressive Strength for 14 days in Mpa

Type of concrete	Compressive strength of concrete after 14 days			Mean
	Sample 1	Sample 2	Sample 3	
M1	22.82	26.72	25.83	25.12
M2	43.04	37.14	40.05	40.07
M3	44.24	42.93	43.55	43.57
M4	45.46	44.85	46.15	45.48
M5	45.64	46.28	45.75	45.89
M6	46.96	46.74	46.55	46.75

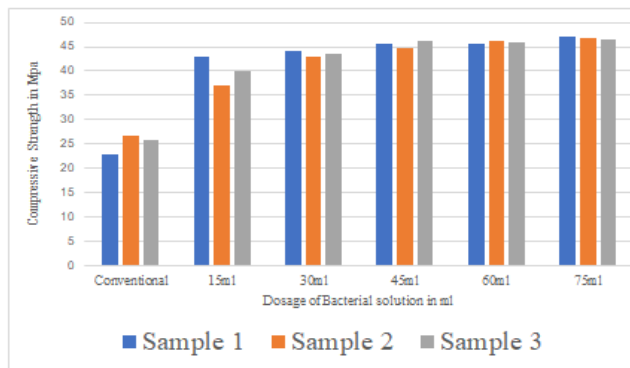


Figure 4: Compressive Strength graph for 14 days

Table 5 and graph 4 describes about the bacterial concrete specimens compressive strength after 28 days of curing at room temperature. The increment of the solution increases the strength of concrete. But after 45ml solution the strength decreases gradually because of the bacterial content which are effecting the strength characteristics are increased

Table 6: Compressive Strength for 28 days in Mpa

Type of concrete	Compressive strength of concrete after 28 days			Mean
	Sample 1	Sample 2	Sample 3	
M1	31.40	33.99	36.81	34.06
M2	42.61	48.39	46.96	45.98
M3	54.56	51.89	53.22	53.22
M4	51.69	55.46	53.74	53.63
M5	53.80	54.15	53.96	53.97
M6	49.50	52.95	51.08	51.17

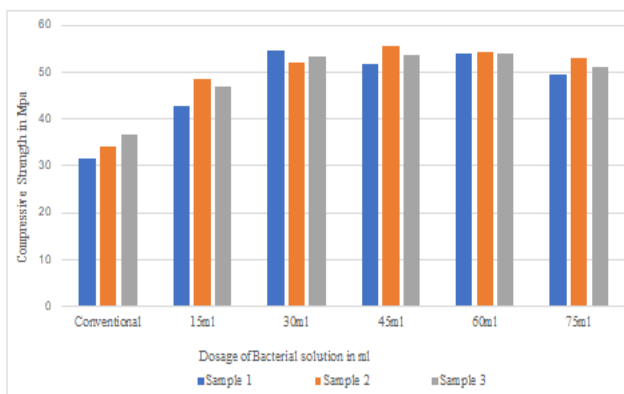


Figure 5: Compressive Strength graph for 28 days

graph 5 explains about the strength for 7 days of curing the specimens. The five samples of every solution i.e., 15ml, 30ml, 45ml, 60ml, and 75ml are tested for 7 days and 28 days. As dilution of bacteria increases the strength increases. By adding the bacillus subtilis the strength has gained nearly to target mean strength even in 7 days

4.2 Ultrasonic Pulse Velocity

An ultrasonic pulse velocity test was carried out in order to determine whether or not the interior structure of the concrete cubes included any cavities. After the exercise was completed, the findings were analysed and collated into table no. 4, which can be seen further down in this table. These findings are based on the data. This was the situation with each and every specimen that went through the testing procedure.

Table No 7: Ultrasonic Pulse Velocity Reading

Property of Concrete	RCC Member	Prob. Distance mm	Time Micro sec	Velocity Km/sec	Probing Method
Conventional concrete	Cube	150	29.3	5.120	Direct
Bacterial concrete					
15 ml	Cube	150	29.70	5.13	Direct
30 ml	Cube	150	28.60	5.30	Direct
45 ml	Cube	150	29.30	5.17	Direct
60 ml	Cube	150	30.40	4.98	Direct
75 ml	Cube	150	29.90	5.12	Direct

Table 8: Ultrasonic Pulse velocity after healing

Type of concrete	UPV TEST			
	Cube Before CTM Test	Cube after CTM Test (50% Load applied)	Cube after 7 days of curing	Cube after 28 days of curing
M1 (Conventional)	5.12	2.80	2.90	3.10
M2 (15 ml)	5.13	2.77	2.92	3.17
M3 (30ml)	5.30	3.10	3.27	3.45
M4 (45ml)	5.17	3.47	3.56	3.78
M5 (60ml)	4.98	2.97	3.20	3.34
M6 (75ml)	5.12	2.66	2.97	3.13

Table No 9: Plate Count Test Result.

bacterial suspension in ml	bacteria Number
15 ml	68 X 10 ³
30 ml	77 X 10 ³
45 ml	89 X 10 ³
60 ml	48 X 10 ³
75 ml	32 X 10 ³

Table 10: Rapid Chloride Penetration Test M30 grade concrete

Mix	Specimens (Coulombs)		Mean (Coulombs)
Control Concrete	Specimens- 1	1681	1665
	Specimens- 2	1642	
	Specimens- 3	1672	
15 ml	Specimens- 1	1674	1650
	Specimens- 2	1623	
	Specimens- 3	1652	
30 ml	Specimens- 1	1660	1636
	Specimens- 2	1665	
	Specimens- 3	1645	
45 ml	Specimens- 1	1672	1664
	Specimens- 2	1668	
	Specimens- 3	1652	
60 ml	Specimens- 1	1672	1678
	Specimens- 2	1686	
	Specimens- 3	1675	
75 ml	Specimens- 1	1660	1660
	Specimens- 2	1655	
	Specimens- 3	1665	

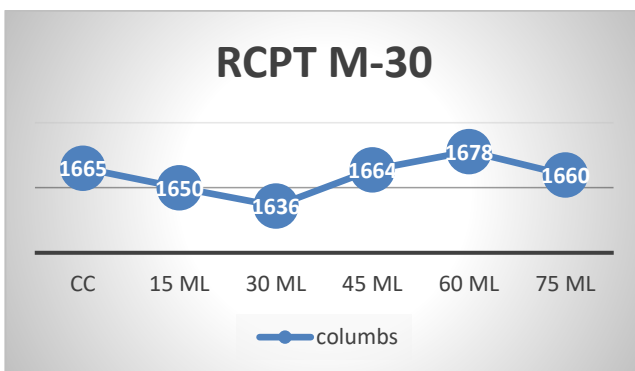


Figure 6: RCPT coulombs

4.3 Scanning Electron Microscope (SEM)

This helps to ensure accurate results. In addition to this, it offers an exceptional level of protection to the spores that are contained within the samples by offering resistance against the pressure that is generated inside the samples as a direct consequence of the production of microstructures. The resistance to the pressure that is formed in the samples is what allows this to be achieved. This validates the creation of calcium carbonate with a comparable crystalline structure and shows that the crystal growth recorded in SEM pictures is similarly. The patterns found in the outcomes of the study carried out by Wiktor and Jonkers are compatible with the variations in CaCO_3 formation that take place depending on whether or not a carrier chemical is present. In addition, this reveals that the crystal

formations that were captured by SEM photos

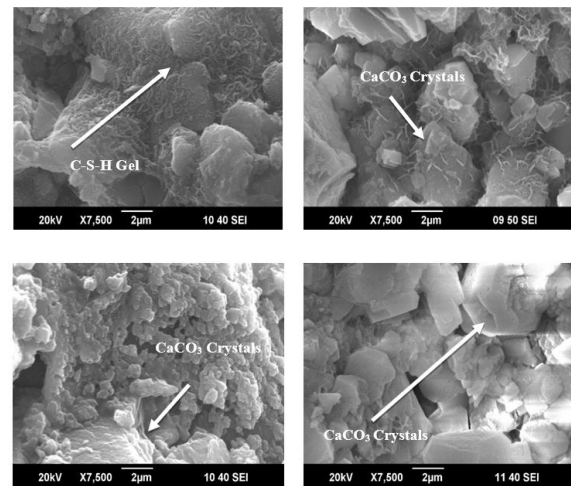


Figure 7: 7 days bacterial concrete effects on SEM

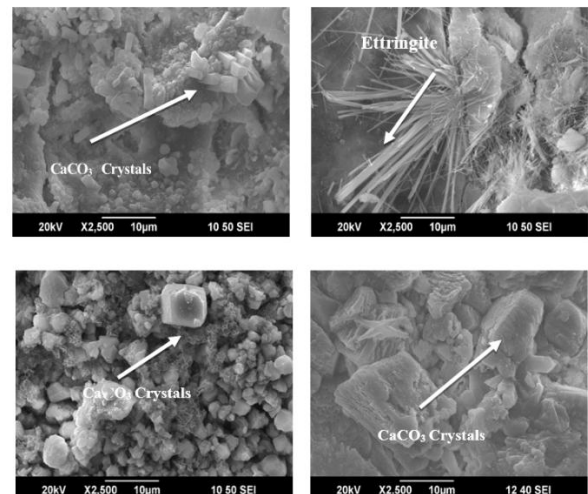


Figure 8: 28 days self-healing bacteria crack reductions

5. CONCLUSION

The highlight the production processes that have been investigated and the performance that have been measure on bacterial self-healing concrete. In conclusion, any type of bacteria with ability to metabolically convert calcium source into calcium carbonate can be used in producing autogenous healing concrete. It is important to provide protection to bacteria in concrete to sustain the self-healing ability throughout the life span of concrete. Bacterial concrete has lower strength compared to conventional concrete about the same composition. However, bacterial concrete able to fully repair visible crack autogenously compared to conventional concrete.

- Incorporating "Bacillus Subtilis" at the correct concentration results in concrete more strength.
- The increasing of bacterial solution, the strength increases up to 60 ml and then strength decreases.
- As compared with conventional concrete, the concrete specimen containing 45ml solution of bacteria increases 25% strength considering the average of three samples after 28 days of curing.
- The concrete containing the 45ml bacterial solution is good to use for crack repairing purposes.
- Ultrasonic pulse velocity probing in direct method maximum in 30 ml of bacteria used velocity is increased.
- The maximum bacterial plate count in 45 ml
- Maximum Rapid Chloride Penetration values are 60 ml dosage of bacterial concrete.
- The resistance of concrete against the assault of acid is noticeably higher than that of regular, typical concrete. It has been shown that bacterial concrete with 60 ml as a substitute offers significantly improved resistance to acid assault.
- A crack in concrete that is between one and forty-five micrometres wide may be healed in a period of thirty days, demonstrating that there is a viable cure for microcracks.
- Oxygen is the agent that may cause corrosion. However, since bacteria consume oxygen, the rate of corrosion can be lowered while the bacteria continue to thrive.
- The formation of cracks will be healed at an earlier stage than was originally envisaged, which will result in an increase in the service life of the structure beyond its projected life.

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