Benefits and drawbacks of applied direct currents for soil improvement via carbonate mineralization

Dimitrios Terzis1*, Patrick Hicher1, and Lyesse Laloui1

1Swiss Federal Institute of Technology Lausanne (EPFL), Soil Mechanics Laboratory, EPFL ENAC IIC LMS Station 18 CH-1015, Lausanne, Switzerland

Abstract. The study presented herein adopts a new vision of the processes involved in carbonate mineralization induced by MICP from an electrochemical and crystal growth perspective. More precisely a specific line of focus refers to the species involved in the bio-chemical reactions and especially their net particle charge. By altering electro-chemical conditions via the application of direct electric currents, we observe distinctive trends related to: (i) overall reaction efficiency; (ii) carbonate mineralization/dissolution and (iii) spatial distribution of precipitates. The study introduces the concept of EA-MICP which stands for Electrically Assisted MICP as a means of improving the efficiency of soil bio-consolidation and overcoming various challenges which were previously reported in conventional MICP-based works. Results reveal both the detrimental and highly beneficial role that electric currents can hold in the complex, reactive and transport processes involved. An interesting finding is the “doped” morphology of calcite crystals, precipitated under electric fields, validated by microstructural observations.

1 Introduction

The present study aims to shed light on the critical role applied Electrokinetics (EKs) can hold in MICP. We postulate that the reactive solutions involved in MICP (rich in microbes, urea and calcium) represent electrolytes and therefore the electrically induced flow of solutes would impact significantly the fate and efficiency of the process to consolidate porous media via the formation of stable, 

\[ \text{CaCO}_3 \] minerals. Applying electric fields to soils have attracted much attention in the geotechnical field, mainly at laboratory scale, for soil remediation applications (Cardoso and Santos 2013, Citeau et al. 2015, Gingine and Cardoso 2019). Applied EKs enhance electrical motion forces that are effective in distributing either dissolved ions, charged particles or even organic ones. The most efficient electrically induced motion force is electromigration of dissolved ions where the electrical force \[ \mathbf{F} = q\mathbf{E} \], with \( q \) being the charge of the considered species and \( \mathbf{E} \) the electric field across the specimen, distributes the ions according to the electric potential distribution. Also, electrophoresis applies on particles or large molecules (typically bacteria) that possess a charged surface due to their polarization under the applied electric field. Previous studies have attempted to introduce electric fields for the application of MICP. More precisely, Keykha et al (2012, 2015) conducted experiments of applied electric currents to Pasteurii bacteria and reported the dependence of electrophoretic mobility of the microorganisms, related to their zeta potential, on the pH (which increases at the cathode and decreases at the anode) as well as the possibility to enhance the transport rate throughout sandy clay specimens by increasing the applied current (Keykha et al 2015). However, the overall effects of the electric field on the microbial communities remain to be clearly understood (Kim et al 2010). Other findings relate to the injection of calcium ions at the anode and carbonate ions originating from microbially hydrolysed urea at the cathode which results into a good rate of calcification throughout soft clay (kaolinite) by means of EKs transport (electromigration and electroosmosis) of the reactive species (Keykha et al 2014). In addition, EKs also seem to be a promising way of removing ammonium ions which can potentially become a source of environmental contamination (Keykha et al 2017). The present study focuses on the potentially detrimental and beneficial effects of applied EKs to suggest an efficient EA-MICP approach.

2 Materials and methods

The study investigates alternative Ca2+ sources and investigates their efficiency as reactive species under applied Direct Current (DC). The vast majority of MICP studies utilize CaCl2 due to its relatively lower price, availability from synthetic routes and high solubility compared to alternative sources (Terzis and Laloui, 2019a). CaCl2 is a purely inorganic salt which displays well known electrolysis reactions that are herein understood with respect to its influence over the microbial activity. With CaCl2 used as a calcium source, Cl- anions are the preferential species that undergoes oxidative reaction at the anode, as follows:

\[ \text{CaCl}_2 + 2e^- \rightarrow \text{Ca}^{2+} + 2\text{Cl}^- \]
(1) \[ 2Cl^- \rightarrow Cl_2(l) + 2e^- \]

due to local low pH conditions. Active chlorine reacts with water to produce hypochlorous acid according to the following reaction:

(2) \[ Cl_2(l) + H_2O \rightarrow HOCl + H^+ + Cl^- \]

which is in equilibrium with hypochlorite at pH around 7.5. Among all oxidants, hypochlorite is mainly known for its powerful disinfecting activity, being the main agent of bleach water. Amino and thiol groups of proteins are particularly sensitive to the electrophilic attack of the latter oxidant (S. Fukuzaki 2006). Even germs that are protected by their plasma membrane drastically suffer from the action of hypochlorite acid that is able to penetrate into the lipid bilayer of living cells and destroy them. We introduce Sporosarcina Pasteurii cells in the sand column experiments previously reported in Terzis and Laloui (2019b).

Electrical conductivity (EC) analysis coupled with Ion Chromatography measurements to detect ionic species (Ca\(^{2+}\) and NH\(^{4+}\)) and monitor the evolution of MICP under applied EKs by comparing it with conventional MICP, i.e. bio-cementation under no applied field. Three alternative Ca\(^{2+}\) sources (Table 1) are provided to the system under no and applied electric field. These are calcium chloride, calcium acetate and calcium lactate.

| Ca\(^{2+}\) source | Solubility (g/100ml at 20º C) |
|------------------|-----------------------------|
| CaCl\(_2\)       | 74.5                        |
| Ca(C\(_2\)H\(_3\)O\(_4\))\(_2\) | 34.7                       |
| CaL\(_2\)       | 4.08                        |

To monitor the EA-MICP process, electroconductivity (EC) measurements are used to provide insight into ionic species production/consumption (NH\(^{4+}\) and Ca\(^{2+}\)). Differential electro-conductivities between the injected, parent solution and the collected outflow (Fig. 2a) show that under applied electrical current, values of EC recorded in the field-free case diverge from those of applied field case after 3 days. Hereafter, \(\Delta EC\) settles to approximately +5 mS/cm in the field-free case, whereas in the case of both applied high 15V and low 5V electric potentials, no difference in EC is recorded. \(\Delta EC\) measurements are correlated to the ionic chromatography (IC) analyses which validate the depletion of NH\(^{4+}\) with relevant concentrations reaching almost zero after 6 days in the applied field cases. Contrary, in the field-free case, NH\(^{4+}\) production over time is captured (Fig. 2b). The Ca\(^{2+}\) concentration reaches that of the parent solution nominal concentration (0.5 M) in the applied field cases. This shows that no reaction occurred, whereas Ca\(^{2+}\) consumption is observed in the field-free experiment. For Cl\(^-\) ions, the same trends are observed in every case with fabric of mineralized calcite and compared to that under conventional MICP.

![Fig. 1. Schematic representation of the setup used to induce EA-MICP in sand columns of 65mm height to 45mm diameter; blue arrows indicate inflow and outflow of reactive solutes and red arrows the applied current via a cathode and anode.](image-url)
concentrations reaching the nominal parent solution concentration after 6 days.

Fig. 2 Monitoring the MICP and EA-MICP processes by (a) Electric conductivity measurements and (b) IC elementary analyses.

3.2 Sand column experiments on EA-MICP using Ca(C$_2$H$_3$O$_2$)$_2$ and CaL$_2$

Plots of the measured NH$_4^+$ concentration in the outflow solutions show the clear relationship between the increased sample’s resistivity with decreasing NH$_4^+$ content (Fig. 3a) with a linear correlation shown in Fig. 3b. Herein, the electrolysed chloride medium is responsible for the destruction of the bacteria cells, meaning no NH$_4^+$ production occurs. Looking at the recorded resistivity of the samples treated with CaL$_2$ and Ca(C$_2$H$_3$O$_2$)$_2$ under 5V applied voltage (Fig. 3c and 3d respectively), no relationship with the ammonium concentration can be extracted, meaning that electrochemical reactions take place, with NH$_4^+$ production and CaCO$_3$ mineralization resulting in quasi-null ΔR.

The mineralized CaCO$_3$ distribution is further illustrated in Fig. 4 for the samples produced using CaL$_2$ and Ca(C$_2$H$_3$O$_2$)$_2$ under two applied voltages and compared to the case of no applied field.

The F series (CaL$_2$ samples) in Fig. 4 shows that both F$_0$ (no field) and F$_1$ (5V field of fixed polarity) samples display a similar trend with a maximum content observed in the middle section, where the reactive solution enters and flows out of the specimen. However, we observe that under an applied 5 V electric field, the distribution of CaCO$_3$ shifts slightly towards higher contents both in the middle and towards the cathode end of the sample, where calcium is attracted. The same behaviour is observed for F$_2$ and F$_3$ samples which yield an extension of the cemented blocks towards the cathode with higher calcite contents than in the no field (F$_0$) case. The most homogenous distribution is yielded for F$_4$ sample, bio-cemented under alternating polarity of 15 V. This much improved homogeneity across the sample’s height is due to the successive electromigration of Ca$^{2+}$

Fig. 3 Plots of the recorded electrical resistance with NH$_4^+$ content measurements over time periods of applied EA-MICP under 5V applied voltage in: (a) calcium chloride with (b) the relationship between the sample’s resistance and NH$_4^+$ content, (c) calcium lactate and (d) calcium acetate media.

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upwards or downwards, depending on the polarity of the applied electric field (see Fig. 1).

Table 2. Variation of NH$_4^+$, Ca$^{2+}$ and pH for samples F and G during 17 batches of EA-MICP treatment.

| Sample | Infiltrations 1-9 | Infiltrations 10-17 | Total | pH |
|--------|------------------|--------------------|-------|----|
| F0     | NH$_4^+$ 0.28    | NH$_4^+$ 0.18      | 0.37  | 7.9 |
|        | Ca$^{2+}$ 0.19   | Ca$^{2+}$ 0.19     | 0.38  | 7.9 |
|        | NH$_4^+$ 0.19    | NH$_4^+$ 0.18      | 0.38  | 7.9 |
| F1     | NH$_4^+$ 0.36    | NH$_4^+$ 0.19      | 0.32  | 8.1 |
|        | Ca$^{2+}$ 0.17   | Ca$^{2+}$ 0.19     | 0.33  | 8.1 |
|        | NH$_4^+$ 0.14    | NH$_4^+$ 0.14      | 0.28  | 8.1 |
| G0     | NH$_4^+$ 0.14    | NH$_4^+$ 0.19      | 0.37  | 7.7 |
|        | Ca$^{2+}$ 0.17   | Ca$^{2+}$ 0.19     | 0.32  | 7.7 |
|        | NH$_4^+$ 0.18    | NH$_4^+$ 0.17      | 0.34  | 7.7 |
| F2     | NH$_4^+$ 0.30    | NH$_4^+$ 0.17      | 0.25  | 8.1 |
|        | Ca$^{2+}$ 0.17   | Ca$^{2+}$ 0.19     | 0.33  | 8.1 |
|        | NH$_4^+$ 0.14    | NH$_4^+$ 0.14      | 0.28  | 8.1 |
| G1     | NH$_4^+$ 0.14    | NH$_4^+$ 0.17      | 0.25  | 8.1 |
|        | Ca$^{2+}$ 0.17   | Ca$^{2+}$ 0.19     | 0.33  | 8.1 |
|        | NH$_4^+$ 0.18    | NH$_4^+$ 0.17      | 0.34  | 8.1 |
| F3     | NH$_4^+$ 0.28    | NH$_4^+$ 0.16      | 0.34  | 7.8 |
|        | Ca$^{2+}$ 0.16   | Ca$^{2+}$ 0.18     | 0.33  | 7.8 |
|        | NH$_4^+$ 0.17    | NH$_4^+$ 0.17      | 0.34  | 7.8 |
| G2     | NH$_4^+$ 0.16    | NH$_4^+$ 0.09      | 0.25  | 7.8 |
|        | Ca$^{2+}$ 0.17   | Ca$^{2+}$ 0.19     | 0.33  | 7.8 |
|        | NH$_4^+$ 0.18    | NH$_4^+$ 0.17      | 0.34  | 7.8 |
| G3     | NH$_4^+$ 0.11    | NH$_4^+$ 0.09      | 0.21  | 7.8 |
|        | Ca$^{2+}$ 0.15   | Ca$^{2+}$ 0.17     | 0.31  | 7.8 |
|        | NH$_4^+$ 0.17    | NH$_4^+$ 0.17      | 0.34  | 7.8 |

Fig. 4 Calcite content distributions across the bio-cemented columns' height under no-field (0) and three applied fields of 5, 10, 15 V for samples bio-cemented using CaL$_2$ (F series) and Ca(C$_2$H$_3$O$_2$)$_2$ (G-series).

The efficiencies of produced NH$_4^+$ and that of depleted Ca$^{2+}$ are shown in Table 2 and are expressed as the ratio of the measured NH$_4^+$ in the outflow over the theoretical NH$_4^+$ production, should urea had completely reacted. It should be noted that Ca(C$_2$H$_3$O$_2$)$_2$ treated samples were provided with 0.5 M concentration compared to 0.25M for CaL$_2$. Therefore, lower efficiency does not necessarily reflect lower calcite content. G samples reveal a different CaCO$_3$ distribution pattern with the exception of G0 (field-free) case where the same variations are observed with respect to F0 (field-free sample). G1 and G2 samples both display a local peak in calcite content in the middle section and a very strong depletion towards the bottom anode end. As far as the G1 sample is concerned, the drop observed in calcite content in the middle is compensated by the increase towards the cathode, due to electromigration of Ca$^{2+}$ ions which result in strong depletion in the anodic region. Thereby, supersaturation of the solution drops, preventing calcium carbonate precipitation. At 15 V applied voltage of fixed polarity, lower calcite content is yielded for the cathodic end while the rest of the distribution remains identical to that of 5 V fixed voltage. The sample G3 under alternating polarity did not yield sufficient bio-cementation and therefore its distribution is not shown in Fig. 4.

3.3 Microstructural observations on EA-MICP "doped crystals"

SEM observations on the texture and morphology of calcite binders which precipitated under applied electric field are carried-out and compared with the characteristics of field-free samples. Figs. 5 and 6 illustrate mineralized CaCO$_3$ yielded using CaL$_2$ and Ca(C$_2$H$_3$O$_2$)$_2$ as calcium sources respectively. Both figures compare the morphology of crystals which were produced via conventional MICP with those produced under EA-MICP. The selected observations highlighted the generally observed trend of minerals which grow larger planes, more stretched and elongated ones with less impurities or imperfections attributed to the presence of encapsulated cells.

Fig. 5 SEM observations revealing calcite binders precipitated using CaL$_2$ under no-field (top) and applied field of 5V (bottom) for the same magnification.

We postulate that this “doped” morphology of crystals is due to the formation of the crystalline lattice, composed of Ca$^{2+}$ and CO$_3^{2-}$, under electric fields which results into a more hierarchical, or less random, distribution of ions and therefore to different nucleation
forms. This finding implies that the effect of EA-MICP extends from influencing the calcite distribution and homogeneity across flow paths, as discussed in section 3.2, and impacts the microstructural and mechanical properties of the precipitated nuclei. Results on the mechanical behaviour of samples yielded under the various chemical and electrokinetic conditions will be the focus of a broader study, currently under development by the authors.

Fig. 6 SEM observations revealing calcite binders precipitated using Ca(C$_2$H$_3$O$_2$)$_2$ under no-field (top) and applied field of 5 V (bottom) for the same magnification

4 Conclusions

This work introduced the concept of EA-MICP and presented a series of benefits and limitations related to its application for soil bio-improvement. The complex reactive-transport phenomena involved in soil bio-cementation are herein understood as a bio-geo-chemical system which can be influenced significantly by the application of electric currents which induce electromigration of bacterial cells and dissolved solutes. Results present the positive effect of applied electric fields of varying force and polarity on the distribution of calcite binders when these precipitate using Ca(C$_2$H$_3$O$_2$)$_2$ and CaL$_2$ as Ca$^{2+}$ sources. Contrary, EA-MICP has a detrimental effect when using CaCl$_2$ for inducing bio-cementation due to the strong acidic front produced. An interesting finding is that application of electric fields results into enhanced morphologies for the mineralized binders, which grow and stretch their planes following more hierarchical patterns. This beneficial form of “doped” crystals is expected to influence the engineering properties of binders and that of the bio-cemented geomaterial. The work overall introduces a new concept for MICP which sets new ground for developments and optimization in the field of soil bio-improvement.

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