ELECTROKENITIC CORROSION TREATMENT OF CONCRETE

Inventor: Henry E. Cardenas, Ruston, LA (US)

Assignee: Louisiana Tech University Foundation, Inc., Ruston, LA (US)

Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 198 days. This patent is subject to a terminal disclaimer.

Appl. No.: 11/734,155
Filed: Apr. 11, 2007

Related U.S. Application Data
Continuation-in-part of application No. 11/341,074, filed on Jan. 27, 2006, now Pat. No. 8,377,278.
Provisional application No. 60/790,867, filed on Apr. 11, 2006.

Int. Cl.
G01N 27/447 (2006.01)
C04B 41/45 (2006.01)

U.S. Cl.
C04B 41/4564 (2013.01)

Field of Classification Search
CPC .......................... C04B 41/4564 (2013.01)

References Cited
U.S. PATENT DOCUMENTS
5,871,872 A * 2/1999 Matijevic et al. .................. 430/7
6,387,244 B1 * 5/2002 Lyublinski ..................... 205/734

FOREIGN PATENT DOCUMENTS

OTHER PUBLICATIONS

Primary Examiner — Gurpreet Kaur

A method and apparatus for strengthening cementitious concrete by placing a nanoparticle carrier liquid in contact with a first surface of a concrete section and inducing a current across the concrete section at sufficient magnitude and for sufficient time that nanoparticles in the nanoparticle carrier liquid migrate through a significant depth of the concrete section.

20 Claims, 5 Drawing Sheets
References Cited

OTHER PUBLICATIONS


Dumas, English Translation by Schreiber Translation Inc.*


Cited by examiner

* cited by examiner
Figure 1

Figure 2

Figure 3

Figure 4
Estimated Depth Range of Probable Cracking

Estimated Stress Level of Probable Cracking

Outside Surface at 25.4 mm

Figure 7 Von Mises Stress vs. Radial Distance Plot for Untreated Case

Figure 8 Von Mises Stress vs. Radial Distance Plot for Treated Case
Figure 9 Crack Penetration for Untreated and Treated Specimens
Field of Invention

The present invention relates to methods and apparatuses for improving the characteristics of concrete. One embodiment of the invention includes a method and apparatus for increasing the strength of concrete by using an electrical field to move particles into pores within the concrete.

Background of Invention

The movement of particles into porous materials such as concrete involve several processes, including adsorption, liquid diffusion, capillary absorption, bulk laminar flow, and electrokinetic transport. Electrokinetic transport includes ionic conduction, electrophoresis, and electroosmosis. Ionic solution conductivity accounts for the overwhelming majority of conductivity measured in cement-based materials. In an aqueous system (cement concrete structures generally retain a certain moisture content in most conditions), ions can be induced to drift in response to an applied electric field. Electrophoresis is characterized by the movement of a solid particle dispersed in an electrolyte under the influence of an electric field. Electroosmosis is the induced flow of water through a porous medium such as sand, clay or concrete when an electric potential is applied across the medium.

Depending on the degree of saturation of a concrete sample, any or all of the above transport processes may occur and a number of structural factors may influence the transport processes. Concrete is a mixture of sand, stone (or other aggregate) glued together with a hardened cement paste that is porous. This pore structure is the dominant microstructural feature governing transport. Pore structure originates from the microstructure when water, anhydrous cement grains, and aggregate are mixed. Capillary pore structure initially assumes the shape of the space occupied by mix water. However, hydration of the cement yields calcium silicate hydrate (C-S-H) the primary binder in hardened cement paste. The capillary pore structure is developed as these hydration products form. Capillary pores tend to dominate transport processes and specific structural characteristics of capillary pores which influence transport include pore volume of the sample, size distribution, tortuosity, and connectivity. The aggregate present in the concrete may influence transport in different ways. Low porosity aggregate tends to impede mass transport by blocking more direct paths through the hardened cement paste pores. Conversely, there can be high porosity at the paste-aggregate interfacial zones. Microcracks and bleed paths also influence particle transport. Microcracks form during drying of the calcium silicate hydrate layers which shrink and lead to tensile stress and cracking. Tensile stress do to plastic shrinkage, stresses from applied loads, thermal expansion or freezing pore water may also inducing microcracking. Bleed paths occur when prior to setting, water accumulates around aggregate and moves toward the surface of the cement paste. Discrete flows can join together to form bleed paths which remain after setting of the cement paste.

Changes in water content of hardened cement pastes have significant impacts on transport mechanisms and rates. At relative humidities above 45%, evaporable pore water is said to exist. Above this threshold, while the permeability of gases is increasingly blocked by liquid water barriers, the transport of aqueous ions or particles progresses more rapidly as the presence of evaporable capillary water increases. Thus, water content is an important factor affecting electrokinetic transport in concrete.

Brief Description of Invention

Preferred embodiments of this invention include a method and apparatus for strengthening cementitious concrete by placing a nanoparticle carrier liquid in contact with at least a first surface of a concrete section and inducing a current across the concrete section at sufficient magnitude and for sufficient time that nanoparticles in the nanoparticle carrier liquid migrate through a significant depth of the concrete section. These particles react with calcium ions liberated from resident calcium hydroxide to form strong phases that increase the strength of the concrete.
concrete section 3 to be treated. Positioned within each container 9 will be an electrode 6 and a conductive liquid. The electrodes 6 will be connected to an electrical power source 11.

In the embodiment seen in FIG. 1, the left electrode 6 will be anode 7 and the right electrode 6 will be cathode 8. Likewise, the conductive liquid in the left container 9 will be a nanoparticle carrier liquid 4 and the conductive liquid in the right container 9 will be an electrolyte solution 5. In this embodiment, the nanoparticle carrier liquid 4 is a colloidal alumina suspension such as product no. 8676 provided by Nalco Chemical Corp. of Chicago, III., which comprises 10% by weight of 2 nm alumina particles in a chloride stabilized carrier liquid. The electrolyte solution 5 in this embodiment is a saturated calcium hydroxide (Ca(OH)2) solution that is less than 1 molar. However, the nanoparticle carrier liquid 4 may contain any liquid containing particles which tend to strengthen concrete when moved into the pore spaces or microcracks existing in concrete. As used herein, nanoparticle generally means a particle that is less than 1 micron in size. A nanoparticle carrier liquid is generally a liquid containing a substantial percentage of nanoparticles, but such liquid may also include some percentage of particles larger than 1 micron. In many embodiments, the nanoparticle carrier liquid will be a colloidal suspension carrying nanoparticles of compounds such as alumina, silica, sodium silicate, or other silicates and aluminates. Other embodiments may include nanoparticle liquids of latex, or polymer particles. However, the carrier liquid is not necessarily limited to suspensions and solutions since vapors and air could be as carrier fluids. Typical weight percent concentrations of nanoparticles could be about 1 to about 60 weight percent, about 10 to about 50 weight percent, about 30 to about 50 weight percent, or any range between 1 and 60 weight percent. Likewise, electrolyte solution 5 could be any number of conductive suspensions or solutions such as cement, hydroxide, tap water, rain water, and pure water that has been allowed to absorb conductive ions from its surroundings (e.g., water in contact with the concrete). However, electrolyte solutions containing aggressive species such as chlorides or sulfates are not typically preferred.

Electrodes 6 may be formed of any material which is sufficiently conductive to carry out the function of the present invention. Exemplary electrode materials include zinc, cadmium, steel, aluminum, copper, monel, or other conductive metals or conductive-oxide-coated versions of these metals. Post-tensioned steel and other applications that may be susceptible to hydrogen embrittlement hazards are less preferred for Electrodes 6. In many embodiments, it is advantageous to form the anode 7 from a relatively corrosion resistant but conductive material such as graphite, a conductive polymer, or a mixed-metal-oxide titanium alloy.

In the embodiment of FIG. 1, the conductive liquids in containers 9 act to provide a uniform electric field across the portion of surfaces 20A and 20B which are in contact with the liquids. The electric field will induce movement of charged particles in the carrier liquid toward the electrode having the opposite charge of the particle and thus into the concrete section 3. Many factors affect the velocity at which particles move into and within the concrete, including the particle size, particle charge, pore fluid concentration, pore wall and particle zeta potential, pore fluid viscosity, electroosmotic coefficient of permeability, fluid pressure acting in opposition to particle transport, thickness of concrete, the size and number of micro-cracks, the porosity (both pore volume and tortuosity) of the concrete, the moisture content of the concrete and the strength of the electric field. If the electric field strength is stated in terms of current density, it may vary from as little as 10 mA/cm² (or less) to more than 1000 mA/cm². The time necessary for nanoparticles to disperse though the entire thickness of concrete section 6 is of course based upon the thickness of the section and the nanoparticle velocity.

As mentioned above, one nanoparticle carrier is colloidal alumina. Alumina may be considered a "pozzolan," which may be defined as a siliceous, aluminous or siliceous and aluminous material which in itself possesses little or no cementing property, but will in a finely divided form and in the presence of moisture chemically react with calcium hydroxide at ordinary temperatures to form compounds possessing cementitious properties. In regards to alumina, a pozzolanic reaction, may be described as the aluminous pozzolans reacting with the (non-cementitious) calcium hydroxide in the hydrated cement paste to produce (highly cementitious) calcium aluminate hydrates which yield higher strength and dramatically reduce the permeability of the concrete. This reaction may be represented by the formula:

\[ \text{CH} + \text{A} + \text{H} \rightarrow \text{C}_2\text{H}_2\text{A}_5\text{H}_2\text{O}_7 \]

where CH is Ca(OH)2, A is Al2O3, H is H2O, and C-A-H is calcium aluminate hydrate. Where the pozzolan is silica, a similar pozzolanic reaction may be described as:

\[ \text{CH} + \text{S} + \text{H} \rightarrow \text{C}_2\text{S}_4\text{H}_6 \]

where S is SiO2, and C-S-H is calcium silicate hydrate.

Sodium silicate may be another nanoparticle suspension utilized in certain embodiments of the present invention. Sodium silicate is any one of several compounds containing sodium oxide, Na2O, and silica, SiO2, or a mixture of sodium silicates with varying ratios of SiO2 to Na2O solids contents, and viscosity. Traditionally, sodium silicates are classified according to the acid from which they are derived as Orthosilicate Na2SiO3; Metasilicate Na2Si2O5; Disilicate Na2Si2O5; Tetrasilicate Na2Si4O10. The sodium silicate species can change from one type to another depending upon conditions and the relative concentrations of each one that is present. This process may provide more of the actual species undergoing the reaction. Sodium silicate (in the case of Meta-silicate) enters pores and combines with calcium hydroxide to form C-S-H gel as follows:

\[ \text{Na}_2\text{SiO}_3 + \text{H}_2\text{O} + \text{Ca(OH)}_2 \rightarrow \text{C}_2\text{Ca}_3\text{Si}_5\text{O}_{14}\text{H}_2\text{O} + 2\text{NaOH} \]

As used herein, "nanoparticle carrier liquid" includes (but is not limited to) any liquid containing nanoparticles (including ions) in suspension, solution, or dissolved, which react with other components to form strong phases to increase the strength of concrete. In certain embodiments, the nanoparticles will be particles or ions which react with calcium hydroxide (either resident in or driven into the concrete) in order to form this stronger phase.

In addition to pozzolans combining with calcium hydroxide originating in the cement section itself, certain embodiments of the present invention employ Ca(OH)2 in the electrolyte solution 5 to increase the availability of Ca2+ ions. For example, if colloidal silica is the nanoparticle carrier 4 in FIG. 1, it will be understood that locations of anode 7 and cathode 8 would be reversed in order to move the negatively charged silica particles into concrete section 3. Likewise, the positive Ca2+ ions in the electrolyte solution 5 would be moved into concrete section 3, where such Ca2+ would be available for the pozzolanic reaction described above. Although this polarity arrangement may tend to remove OH- ions from the concrete, the abundance of OH- in the concrete means any resulting removal of OFF is inconsequential to the overall treatment process.
Although FIG. 1 suggests containers 9 are enclosed (i.e., additional liquid is not shown being added), this need not always be the case. In some embodiments, the volume of nanoparticle liquid in container 9 will have sufficient nanoparticles to completely treat the concrete section 3. However, in other embodiments, additional nanoparticle carrier liquid may be fed into container 9 (i.e., exchanged with the depleted carrier liquid) if a greater mass of nanoparticles is needed for complete treatment.

Alternate embodiments of present invention applying similar electrokinetic transport principles as described above may be seen in FIGS. 2-5. FIG. 2 represents a concrete section 3 having reinforcing steel or “rebar” 13. A container 9 will be formed around one end of concrete section 3 such that the nanoparticle carrier liquid 4 is in contact with the surface 21. A wire, mesh, or flat plate electrode 12 will be positioned in the nanoparticle carrier liquid 4 and will extend substantially the length of concrete section 3 (i.e., the length running perpendicular to the cross-sectional plane shown in FIG. 2). In this embodiment, the rebar 13 will be connected to current source and serve as an electrode (e.g., cathode 8). When plate electrode 12 forms the anode and rebar 13 forms the cathode, positively charged nanoparticles (e.g., alumina) in carrier liquid 4 will be driven toward rebar 13 by the electric field established between the electrodes. Clearly, the distance which must be traveled by the nanoparticles is greater in FIG. 2 than FIG. 1, thereby suggesting the use of a higher current density or a longer treatment duration than might be necessary for the arrangement of FIG. 1.

Although the embodiments of FIGS. 1 and 2 contemplate the treatment of a rigid cured concrete section 3, FIG. 3 illustrates an alternate embodiment that could be employed to treat an uncured concrete section 15. A form 16 will be provided which, as is well known in conventional concrete production, will contain the uncured flowable concrete mix in the intended shape of the ultimate concrete section. The sides of form 16 will be higher than the depth of the concrete section in order to accommodate a fluid (explained below) overlying the placed concrete mix. A wire mesh electrode 14A will be positioned in the bottom of form 16 prior to the pouring of any concrete mix. Wire mesh electrode 14A can be of various types of mesh, preferably with a large enough opening to allow water to pass readily through and a small enough opening to ensure the distribution of a uniform electric field across the substrate. In one embodiment, a mesh with a 1/16 inch opening size may be placed in direct contact with the surface. In another embodiment, a large mesh with an opening size of 12 inches may be located 6 inches from the concrete surface. Such meshes will generally conform to the bottom dimensions (width and length) of the form 16. However, alternate embodiments of wire mesh electrode 14A could be greater or less than the bottom dimensions of form 16. Likewise, in other embodiments not specifically illustrated, rather than a mesh electrode, a conductive plate could be positioned in the bottom of form 16. Alternatively, any reinforcing steel intended for use in the concrete section 15 could serve as the electrode. After placement of electrode 14A, a conventional cement paste mix (e.g., water, cement, course and fine aggregate) will be placed in form 16 in an amount sufficient to produce the desired thickness of the concrete section. Optionally, the mix design may be adjusted with more water and less cement powder to optimize cost. A second mesh electrode 14B is spaced over the top of the uncured cement mix. In the embodiment shown, the mesh electrode 14B will be spaced slightly above (e.g. at least 0.5 cm) the top of the placed concrete mix. However, in other embodiments, the mesh can also be touching the concrete or be positioned just beneath the concrete surface. A nanoparticle carrier liquid 4 is then carefully introduced onto of the concrete mix in a manner that does not disturb the concrete mix and in an amount sufficient to immerse the mesh electrode 14B, assuming it is not cast in. Thereafter, an electric current is applied between the electrodes 14A and 14B in order to induce the migration of nanoparticles through the concrete section. In the embodiment shown, a current density of about 10 mA/cm² to about 1000 mA/cm² could be employed (although current densities outside this range are possible) and in one particular example, the current density would be about 500 mA/cm². The duration of current application will depend on factors such as the desired depth to which nanoparticles are being directed, the magnitude of the current density, and the water content of the cement mix. In many embodiments, the duration should be sufficient to move particles into all of the locations that are subject to the electric field, since application of an electric field to the concrete without nanoparticle strengthening may result in a weakening of the porosity of the concrete. If the treatment occurs over several days, the concrete may be more than sufficiently cured to remove form 16 and pull wire mesh 14A from the bottom of the concrete section. Although the method of FIG. 3 has been described as being initiated with uncured concrete mix, the method could be applied to any hardened cement paste (i.e., the concrete having reached the binder phase that holds all the stone and sand in a fixed matrix).

A further embodiment seen in FIG. 4 utilizes a different method for placing the nanoparticle carrier liquid into contact with the concrete section 3 (a cured concrete section in the example of FIG. 4). Placed against at least one surface of concrete section 3 will be a flexible porous material 17 which is capable of at least partially absorbing and retaining a nanoparticle carrier liquid brought into contact with the flexible porous material 17. In one example, flexible porous material 17 is a sponge material. Such sponge materials could include natural sponges, e.g., a porous rubber or cellulose product having properties similar to a natural sponge. Flexible porous material 17 will typically cover the general area of concrete section 3 into which nanoparticles are to be introduced. The thickness of flexible porous material 17 may vary in different embodiments. In one example, flexible porous material 17 may be approximately 2.5-7.6 cm thick. However, various embodiments, the porous material need only be thick and flexible enough to accommodate most of the surface topography of the substrate being treated so that wet electrical contact is maintained during treatment. FIG. 4 also illustrates a reservoir 18 of nanoparticle carrier liquid 4 which communicates with flexible porous material 17 via supply line 19. The transfer of fluid could be accomplished through a gravity feed system as suggested in FIG. 4 or though some type of pumping arrangement. Reservoir 18 will replenish the nanoparticle carrier liquid 4 in flexible porous material 17 as the liquid evaporates and as nanoparticles are driven from flexible porous material 17 into concrete section 3. A mesh electrode 14B will be positioned over flexible porous material 17. On the surface of concrete section 3 opposite mesh electrode 14B, another flexible porous material will be positioned to bring an electrolyte solution into wet electrical contact with that surface of concrete section 3. The mesh electrode 14A will be positioned on the flexible porous material. Although not explicitly shown in FIG. 4, certain embodiments could include a reservoir of electrolyte solution to supply the flexible porous material 17. Likewise, the porous material could...
be positioned under a mesh electrode of conductive fabric, e.g., a fabric containing a weave of flexible graphic wire. As in the embodiments described in FIGS. 1-3, application of a current between mesh electrodes 14A and 14B at a sufficient magnitude and for sufficient duration will induce nanoparticles to move into and through at least a significant portion and preferably the entire depth of the concrete section 3.

Many variations of the method described in FIG. 4 are within the scope of the present invention. The material 17 need not be flexible and there may be applications where a comparatively rigid porous material may be employed. Additionally, it may not always be necessary to have the material 17 re-supplied with liquids from some external source such as reservoir 18. Rather, in certain applications it may suffice to simply expose material 17 to nanoparticle and electrolyte liquids at the initial stage of the treatment process and this will provide sufficient nanoparticles for the complete treatment of the concrete section. Alternatively, material 17 could be refreshed with liquids at one or two points in the treatment process (as opposed to continuous supply from a reservoir). Although FIG. 4 illustrates separate mesh electrodes 14A and 14B positioned over porous material 17, other embodiments might include metal fibers (or other conductive materials) incorporated into porous material 17, thereby combining the liquid retaining function and the electrode function into a single section of material. Likewise, if rebar is present in the concrete section 3 (for example near the side on which mesh electrode 14A rests), then the rebar may substitute for electrode 14A (and eliminate the need for porous material 17 under electrode 14A). Also, porous material 17 could be applied to an uncured concrete mix to replace the pool of carrier liquid described in the embodiment of FIG. 3.

FIG. 5 illustrates another embodiment using a porous material to retain a nanoparticle carrier liquid in contact with a concrete section. In the cross-sectional view of FIG. 5, the concrete section is a concrete column section 25. As is typical with concrete columns, column section 25 will include a series of rebar members 15. In FIG. 5, a continuous section of porous material 26 is wrapped around column section 25 and a mesh electrode 14 is either intermeshed or positioned atop porous material 26. In this embodiment, one or more of rebar members 13 will act as the electrode 8. Typically, the center most rebar member(s) 13 will act as electrode(s) 8 in order to move nanoparticles as far as possible toward the center of column section 25 (i.e., nanoparticles are not expected to migrate any further inward than the most central rebar position of electrode 8). Although not shown, a nanoparticle carrier liquid reservoir could be connected to porous material 26. Additionally, porous material 26 need not be continuous around the circumference of the column, but could be placed in discrete sections to cover a substantial portion of the column’s circumference (with the same being true for mesh electrode 14).

In a further alternate embodiment of the present invention, electrokinetic transport arrangements similarly to those described above may be used for corrosive species extraction from a concrete structure. It is well known that the migration of aggressive chemical species such as chloride to the reinforcing metal (e.g., “rebar”) in concrete structures can corrode the rebar and significantly weaken the concrete structure. In a similar manner, carbon dioxide, sulfates, or nitrates absorbed into pore fluid in proximity to the rebar can lower the pH of the surrounding area and induce corrosion. The use of an electric field to move chlorine and other corrosive species away from the rebar and/or raise the pH in the vicinity of the rebar can act to extend the useful life of the concrete structure.

The structural elements required to carry out this chlorine extraction process may be similar to that seen in the preceding figures, particularly FIGS. 2 and 5. In one embodiment, the negative pole of the power supply (cathode) is connected to the rebar. This insures that negatively charged chlorine species are repelled from the rebar. In this embodiment, the positive pole of the power supply (anode) will typically be external to the concrete section. For example, the anode could be a bar electrode in the electrolyte liquid such as FIG. 2, or a mesh electrode placed on the concrete structure as in FIG. 6, or any other method of positioning an electrode in order to generate an electric potential which will urge negatively charged species away from the rebar. There may also be situations where the anode is not external to the concrete section. A still further alternate embodiment of the corrosive species extraction method could be carried out using a “cross flow” configuration, i.e., without an electrode connected to the rebar, but distal from the rebar such that the electric current flows across the concrete section and any rebar therein (e.g., FIG. 1). Although the figures show the treatment being applied to one or more entire sides of the concrete section, the invention naturally also includes applying the treatment to a surface portion which is less than the entire side.

The magnitude of the electrical field applied to the concrete section may vary significantly from embodiment to embodiment depending on various conditions of the particular concrete structure being treated. In general, application of a higher magnitude electrical field (e.g. current densities over about 100 mA/ft²) for any significant length of time may cause the concrete to soften and/or become more porous, thereby suffering a loss in strength (both compressive and tensile). On the other hand, increasing the magnitude of the electrical field decreases the treatment time required to obtain sufficient extraction and thereby significantly reduces the cost of treating concrete structures. In many embodiments, it would be desirable to employ electrical fields having current densities of between about 100 mA/ft² and about 2000 mA/ft² (or any range of current densities therebetween), although higher current densities are within the scope of the present invention. In one embodiment, a range of 500 mA/ft² and about 1500 mA/ft² may be employed whereas in another embodiment, a range of 300 mA/ft² to about 700 mA/ft² may be employed. There could also be embodiments where current densities less than 100 or greater than 2000 mA/ft² are employed.

In the embodiments shown in the figures, the electric field is enhanced by maintaining both the electrode(s) and the surface of the concrete section in contact with a fluid. In preferred embodiments, this fluid is an electrolyte solution. The fluid may be contained over one or more surfaces of the concrete section and an electrode(s) placed in the fluid (FIG. 1); the fluid may be pooled on one surface of the concrete with one electrode positioned in the pool and the other electrode attached to the rebar (FIG. 2); or an absorbent material containing the fluid may be positioned between the electrode(s) and the concrete section (FIGS. 4 and 5). When the rebar is used as an electrode 6, the best results may be obtained by placing a fluid (and opposing electrode 6) on opposite sides of the concrete section as seen in FIG. 10A, creating electric fields in both directions away from rebar 13. However, the working environment will often not allow more than one surface of the concrete section to be placed in contact with the electrolyte/carrying fluid 4. In particular, it may only be a lower surface of a structure which is accessible. In such cases, the containment 9 for electrolyte/carrying fluid 4 will be positioned on the bottom of concrete section 3 as seen in FIG. 10B.
over corrosive species above rebar 13, significantly reducing
the concentration of corrosive species below rebar 13 through
electrokinetic treatment will tend to reduce the concentra-
tion of corrosive species above rebar 13 by way of inherent con-
centration/diffusion forces. Nevertheless, the illustrated fig-
ures only represent selected examples of techniques for
bringing a fluid into contact with the surface(s) of the concrete
section and any alternative technique accomplishing this goal
is included within the scope of the present invention.

A concrete section will have an initial compressive and
tensile strength prior to beginning the treatment process. In
order to compensate for any weakening of the concrete struc-
ture resulting from application of the electrical field, in cer-
tain embodiments of the present invention it will be desir-
able to subject the concrete structure to a repair or re-strengthen-
ning process similar to that either described above or one of the
processes described below. In general, the strengthening pro-
cess will comprise driving certain microparticles or nanopar-
ticles into the concrete pores utilizing the electrical field. In
certain embodiments, the concrete structure will in a first step
be subject to an electric field for the removal (or at least
migration away from the rebar) of chloride or other corrosive
species. Then a second step will consist of driving micropar-
ticles into the concrete structure using the electric field. In
other embodiments, the step of chloride extraction and micro-
particle migration will partially overlap or alternatively, both
steps may be carried out simultaneously for the entire treat-
ment process.

In the most typical embodiments, the microparticles will be
less than about 10 microns in average diameter and this defi-
nition includes nanoparticles which are less than 1 micron.
Certain embodiments include microparticles in size ranges
such as about 500 nm to about 1 nm (or any range therebe-
tween), about 100 nm to about 10 nm, or about 50 nm to about
20 nm. The microparticles could be of a generally uniform
size (e.g., all particles about 20 nm) or a mixture of particle
sizes within a given size range, including a graduation of
particles sizes in one of the above ranges. In certain embed-
iments, particles of one size may be used in one portion of the
treatment and then particles of a different size used in another
portion of the treatment. For example, during the initial stages
of treatment, positive particles of 20 nm could be migrated
toward the cathode rebar while not significantly impeding the
chlorine species moving away from the rebar because the 20
nm particles are too large to enter and obstruct all pores in the
concrete. Then at a later stage of treatment, 2 nm particles
could be migrated toward the rebar and cause the concrete
pores to become impassable to even the smaller chlorine or
other harmful compounds (e.g., CO₂) which tend to lower the
pH of the concrete pore water.

Many different microparticles or nanoparticles could be
employed in the present invention. In certain embodiments, the
microparticles described above (e.g., silicates or alumi-
nates) could be employed. In embodiments where the rebar
acts as the cathode, the microparticles will generally need to
have a net positive charge in order to be driven into the con-
crete structure. In one embodiment, the microparticle will be
a silicon species combined with a positive species such as
alumina coated silica or lithium, alumina coated silica. Like-
wise, a large number of substances could form the positive
species component of the microparticle. In some embed-
ments, the positive species component could be a metal oxide
such as alumina or zirconia. The positive species component
could be polymers such as poly allyl dialkyl methyl amno-
nium chloride (PDDMA). Other species of either net charge
could be vinyl acetates, vinyl acetate-ethylenes, styrene-buta-
dienes, vinylidene chlorides, or epoxies. It could also be an
crylic compound with a functional group having a positive
net charge such as acrylic esters (acrylates) or acrylic copol-
mers. The above is merely an exemplary list of possible
positive species which could be combined with a silica spe-
cies and all positive species which physically or chemically
bond with a silicon species should be considered within the
scope of the present invention.

The microparticles utilized in the current invention are not
limited to the above examples combined with a silica species.
In alternate embodiments, one or more positive species alone
(i.e., not combined with silica) could be driven into the con-
tcrete to render it less permeable. The positive species could be
one described above, but other positive species could be
employed. In many instances, it will be desirable for the
positive species to have some tendency to strengthen the
concrete. For example, alumina, zirconia, lithium silicates
(lithium coated silica or lithium stabilized silica), calcium
compounds, iron or iron oxides covered with a positive spe-
cies. Many inert species capable of filling the concrete pores
will provide a strengthening effect due to the reduction of
porosity. These can include most metal oxides.

Not all process embodiments of the present invention
require strengthening of the concrete section. Certain
embodiments will lower the pH in the vicinity of the rebar and
then migrate microparticles into the concrete surface to form
a substantially impermeable outer surface on the concrete
section. Depending on the microparticle size and the size of
the pores in the concrete section, it may only be necessary
to migrate microparticles a few millimeters into the concrete
surface to render the surface substantially impermeable.
Thus, the concrete section is sealed, but microparticles have
not penetrated to a sufficient depth to measurably strengthen
the concrete section as a whole.

In many instances, it will assist the microparticle to remain
suspended in the carrier solution if a stabilizing agent is
employed. In one embodiment, an acetate-based stabilizing
agent such as acetic acid may be used. In other embodiments,
sodium, chloride, potassium, or ammonium may be used.
Other stabilizing agents could include almost any salt. De-
pending on the microparticle employed, simply maintain-
ing an appropriate pH in the solution (with no added stabiliz-
ing compound) may cause the microparticle suspension to
remain stable for long periods. Stabilizing ion concentra-
tions are often dosed at a fraction of a percent. As a nonlimiting
example, about a 50 percent concentration of colloidal silica
would require stabilizing dose of sodium of about 0.40 per-
cent (as Na₂O). In these cases, pH values preferably should be
well above 7 for good stability. For an alumina suspension,
the pH would preferably be in the about 3.5 to about 4 range.

The length of time the concrete section is subject to the
electric field may vary greatly from embodiment to embed-
diment depending on factors such as depth of the concrete
section being treated, the strength of the electric field, the
corrosive species and its concentration being removed, the
age, porosity, pore size, temperature and tortuosity of the
b. placing a microparticle carrier fluid in contact with at least one part of said concrete section, thereby returning the strength of said concrete section to at least said initial compressive and tensile strength.

c. applying a current at sufficient magnitude and for sufficient time to:
   i. remove at least a portion of a chlorine species present in or on said concrete section; and
   ii. move microparticles in said carrier liquid migrate through at least one quarter of said depth of said concrete section, thereby returning the strength of said concrete section to at least said initial compressive and tensile strength.

2. The method according to embodiment 1, said first surface being exposed to said liquid containing microparticles for about 15 to about 50 percent.

3. The method according to embodiment 1, wherein said microparticle carrier fluid includes microparticles of said first surface and said absorbent material brings microparticles into contact with said concrete section.

4. The method according to embodiment 1, wherein said absorbent material is used in structures at other locations.

5. The method according to embodiment 1, wherein said absorbent material is used in structures at other locations.

6. The method according to embodiment 1, wherein said absorbent material includes absorbent material for said concrete section.

7. The method according to embodiment 1, wherein said absorbent material includes absorbent material for said concrete section.

8. The method according to embodiment 1, wherein said absorbent material includes absorbent material for said concrete section.

9. The method according to embodiment 1, wherein said absorbent material includes absorbent material for said concrete section.

10. The method according to embodiment 1, wherein said absorbent material includes absorbent material for said concrete section.

11. The method according to embodiment 1, wherein said absorbent material includes absorbent material for said concrete section.

12. The method according to embodiment 1, wherein said absorbent material includes absorbent material for said concrete section.

13. A method of treating concrete comprising:
   a. providing a section of concrete including metal reinforcing extending into said concrete section for purposes of re-strengthening.
   b. placing a liquid in contact with at least one part of said concrete section; and
   c. applying a current at sufficient magnitude and for sufficient time to:
      i. remove at least a portion of a chlorine species present in or on said concrete section; and
      ii. move microparticles in said carrier liquid migrate through at least one quarter of said depth of said concrete section, thereby returning the strength of said concrete section to at least said initial compressive and tensile strength.

14. The method according to embodiment 13, wherein said absorbent material includes absorbent material for said concrete section.

15. The method according to embodiment 13, wherein said absorbent material includes absorbent material for said concrete section.

16. The method according to embodiment 13, wherein said absorbent material includes absorbent material for said concrete section.

17. The method according to embodiment 13, wherein said absorbent material includes absorbent material for said concrete section.

18. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

19. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

20. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

21. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

22. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

23. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

24. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

25. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

26. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

27. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

28. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

29. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

30. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

31. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.

32. The method according to embodiment 13, wherein a second, positive electrode comprises a wire mesh, or conductive fabric, or a fabric-reinforced conductive matrix wetted with a conductive medium.
reduce treatment access to the pore structure and thus reduce blocks were treated and tested 16 months following produc-
ction that would otherwise be available. Electrokinetic treat-
ment period. Saturation was established to minimize absorp-
tion to applying the nanoparticles in the methods described in FIGS. 1-5. For example, it is envisioned that the nanoparticle carrier liquid could be a thick viscous liquid with a consist-
cy to paint. The carrier liquid would then be “painted” onto the surface of the concrete section with an electrode (e.g., a mesh electrode) placed directly on the painted section of the cement. Alternatively, particles could be used that are transported through the air using a powder coating wand. The substrate would preferably be wet so that particles that absorb onto the wall could continue to migrate in the electric field. Another alternative could involve the use of a conductive gel. The gel would be loaded with particles, applied to the concrete substrate and driven into the concrete using and an electrode immersed in the gel, or the substrate. The gel may be vacuumed and recycled for a future application. All such variations and modifications should be consid-
ered within the scope of the claims.

EXAMPLES

Example 1

Tests were conducted to determine the impact of the electrokinetic treatment on common heavy and light weight masonry blocks. A heavy weight block is a hollow load-bearing concrete block 8-by-8-by-16-inches nominal size, having two hollow sections, and weighing from 40 to 50 pounds when made with heavyweight aggregate, such as sand, gravel, crushed stone, or air-cooled slag. The same size block is also shown in Table 1.

The heavy weight block specimens were labeled as fol-

Set 1 contains Experimental Block 1 (LEB1) and Control Block 1 (LCB1),

Set 2 contains Experimental Block 2 (LEB2) and Control Block 2 (LCB2), and

Set 3 contains Experimental Block 3 (LEB3) and Control Block 3 (LCB3).

The heavy weight block specimens were labeled as fol-

Set 1 contains Experimental Block 1 (HEB1) and Control Block 1 (HCB1),

Set 2 contains Experimental Block 2 (HEB2) and Control Block 2 (HCB2), and

Set 3 contains Experimental Block 3 (HEB3) and Control Block 3 (HCB3).

The treatment circuit was set for constant voltage. For the light weight specimens a current of 38 mA was applied for Set 1 and a current of 76 mA was applied for Set 2 and Set 3. The heavy weight blocks were all treated with a current of 76 mA. Some current and voltage drift was observed and was mainly due to electrode polarization. In addition, the loss of ions due to the C-S-H reaction caused resistance to go up and thus an increase in voltage. Current and voltage readings were taken every 24 hrs before and after adjusting the current reading back to the original value.

The five-day electrokinetic treatment period was con-

The maximum compressive load test was conducted to determine the change in the load resistance of the block after the treatment. The impact on load resistance in each Set of blocks is also shown in Table 1.
The percentage of increase in load resistance in light weight blocks was greatest for specimen LEB3 with a value of 138% as compared to 134% for specimen LEB2 and 119% for specimen LEB1. The average increase in load resistance among the light weight blocks was 130%. The maximum increase in load resistance in heavy weight blocks was observed in specimen HEB3 with an increase of 110%. The average increase in load resistance for the heavy weight blocks was 100%.

Current was supplied at the density of 9 mA/m² in order to minimize damage to the blocks. For an average surface area of 0.24 m² the resultant current was 38 mA. This was the applied current for specimen LEB1. It was later taken into account that the masonry blocks could be treated with double the current density since the particles tended to heal the damage from the current. This observation enabled the use of a current of 76 mA for blocks from Set 2 and Set 3. This healing assumption was demonstrated by the compressive load resistance observed. The load resistance increase in light weight blocks was higher for specimens LEB2 and LEB3 with an average increase of 136% as compared to a 119% increase for specimen LEB1. This demonstrated a difference of only 19%. This behavior also suggests that the different current densities did not produce a significant difference in result and that five days may not be required to achieve a given level of block performance.

There was a significant enhancement in the load resistance due to treatment especially in view of the largeness and the structure of the masonry block. For the average enhancement in load resistance was increased by 130% for light weight blocks and by 100% for heavy weight blocks. The average enhancement in load resistance for all the treated light weight blocks was 2200%. It is clear from these results that the mixture of ions and suspended silica particles in sodium silicate reacted sufficiently with calcium hydroxide to block the pores of the masonry block thereby bringing about a significant reduction in permeability.

**Example 2**

Thermal shock is a severe condition that military aircraft pavements experience. Normal strength concrete loses 10-20% of its original compressive strength when the temperature is increased to 300° C. It can lose up to 60% of its strength when heated to 600° C. To address this problem, specimens were tested that consisted of mixtures of Type I Portland cement. Cylindrically shaped specimens were made from Portland cement paste and cast into polyethylene vials. The specimens were 50.8 mm in diameter and 50.8 mm in height. Nanoparticle treatment commenced immediately following batching. Two batches of specimens were made with water/cement (w/c) ratios of 0.4, and 0.5. The Type I Portland cement was manufacture by Lonestar Industries Inc. An electrokinetic drive circuit was connected similar to that shown in FIG. 3. Three specimens from each batch of nine were treated electrokinetically with 2 nm colloid alumina nanoparticles for 14 days. The colloidal alumina was poured gently at the top of each Portland cement specimen. This pond was replenished daily. The current was set to provide a current density of 1.1 A/m². The power supply was set in current control mode. During this period, control specimens were stored in limewater. After 14 days, both the treated and control specimens were placed in a furnace and heated for 36 hrs at 550° F. (288° C.). The specimens were removed from the furnace and water quenched. After quenching, the specimens were capped with a sulfur capping compound. These specimens were tested in compression.

Cracks appeared on both the treated and untreated specimens after quenching following a thermal exposure at 550° F. (288° C.) for 36 hours. Compressive tests were conducted on specimens in accordance with ASTM standard C150 in the following categories.

1) Non treated and untreated,
2) Non treated and oven tested, and
3) Treated and oven tested specimens.

The strength values are presented in Table 1. These values provide a comparison of the above mentioned cases for the 0.4 and 0.5 w/c ratios. The load application time in each case was 70 s. Each value represents an average of three specimens. It was observed that the heated controls were stronger than the specimens that were heated and quenched. The specimens that were nanoparticle treated prior to heating and quenching were stronger than the untreated cases as well as the controls.

**TABLE 1**

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Failure Load (lbs)</th>
<th>Increase in Load Resistance (%)</th>
<th>Specimen</th>
<th>Failure Load (lbs)</th>
<th>Increase in Load Resistance (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LCBI</td>
<td>7200</td>
<td>119</td>
<td>HCB1</td>
<td>13600</td>
<td>101</td>
</tr>
<tr>
<td>LEB1</td>
<td>15800</td>
<td></td>
<td>HEB1</td>
<td>27400</td>
<td></td>
</tr>
<tr>
<td>LEB2</td>
<td>9900</td>
<td>134</td>
<td>HEB2</td>
<td>14800</td>
<td>105</td>
</tr>
<tr>
<td>LEB3</td>
<td>22300</td>
<td>130</td>
<td>HEB3</td>
<td>30300</td>
<td></td>
</tr>
<tr>
<td>LEB3</td>
<td>9600</td>
<td>138</td>
<td>HEB3</td>
<td>16100</td>
<td>110</td>
</tr>
<tr>
<td>LEB3</td>
<td>22800</td>
<td></td>
<td>HEB3</td>
<td>33800</td>
<td></td>
</tr>
</tbody>
</table>
Quenching produced cracks on all the specimens. For this reason compression testing was needed to discern the level of thermal damage. Table 2 contains a summary of the compression test results. Specimens from the 0.5 w/c batch that were heated to 550° F (288° C.) for 36 hrs (and quenched) exhibited a 26% reduction in compression strength as compared to the control specimens. Those specimens that were treated with nanoparticles exhibited a 5% increase in strength compared to the control specimens. When compared to the untreated and heated specimens, the nanoparticle treated specimens exhibited a strength increase of 40%. This work indicated that the nanoparticle treatment, applied to the 0.5 w/c ratio case, provided significant resistance to compressive strength degradation that would otherwise result from this thermal exposure.

A second batch of specimens having 0.4 w/c ratio was also prepared and tested. In this batch, the untreated and heated specimens lost 22% of the compressive strength compared to the control specimens. If the treated and heated specimens were compared with the control specimens, the treated and heated specimens exhibited 46% more strength. Treated and heated specimens exhibited an 116% increase in strength compared to the untreated and heated specimens.

Normally the lower the w/c ratio the higher the strength of the specimens. Results from Table 3 show that the strength of the control specimens of 0.4 w/c increased by an average value of 4% compared to the 0.5 w/c specimens. In the untreated and heated case the strength of the 0.4 w/c specimens was increased by 7%. For the treated and heated specimens this w/c-related strength increase was 64%. This indicates that by using this nanoparticle treatment the compressive strength of the specimens increases even after severe thermal exposure.

All the specimens cracked after quenching from 550° F. (288° C.). From the outside, both the treated and untreated looked similar. Compressive tests indicated that the thermal cracking may have occurred adjacent to the surface of the specimen but not deep within the core. The core strength may have played a role in restricting the thermal cracking. Quenching caused high thermal gradients on the surface of the both the treated and untreated specimens. Due to these gradients, high thermal stresses were expected on the surface of the specimens. When the thermal stresses exceeded the tensile strength of the specimens then cracking would be expected to occur.

A thermal analysis model was developed using ANSYS (Release 7.0) software to assess the possible extent of thermal cracking induced by quenching. A sequential analysis was done on both treated and untreated specimens for calculating thermal stresses. The thermal analysis results were compared with the experimental results. The properties of both the untreated and treated specimens are presented in Table 3. This table consists of handbook values adapted for constructing a thermal computer model analysis of how the concrete responded to being quenched. The model indicates how far the thermal cracking reaches and this information was used to determine the strength of the uncracked interior referred to as the core region or core strength.

A plot was made of thermal stresses vs. radial distance from the center towards the outer surface in the untreated case as shown in FIG. 7. A similar plot was developed for the treated case as shown in FIG. 8. These stresses were calculated using ANSYS (finite element software). Thermal stresses were calculated to predict the extent of tensile stresses that could cause cracking on the surface of a specimen. The stress curve plotted assumed no surface cracking but may possibly indicate the depth of surface cracking. The depth of possible surface cracking was indicated by the level of stress which exceeds the typical tensile strength range of 3.6 to 7.2 MPa.

A calculated modulus of elasticity (64 GPa) provided thermal stresses nearer to the tensile strength of the material than was provided using the literature value of 413 GPa. The typical level of tensile strength is in the range of 3.6-7.2 MPa. The stresses obtained were approximately 5 times this stress level. Contrarily when the reported modulus of elasticity (413 GPa) was used for the analysis, the stresses it induced were up to 100 times the tensile strength of the material. Based on these calculations, it appears that the modulus of elasticity had a high positive impact on the stresses calculated in the specimens. In future work, a better estimate of this modulus should be obtained.

The finite element analysis results showed that the temperature gradient was higher for the treated specimens as compared to the untreated specimens. The higher thermal gradient is expected as a result of a low heat transfer rate, but the important thing here is that a higher thermal gradient can provide higher thermal stresses. High displacements in treated specimens were observed as compared to the untreated specimens especially at the outer surface. The elevated coefficient of thermal expansion for the treated cases (~2x higher) is causing high displacements.

The thermal stress profiles due to quenching indicate a high stress state from the surface inward reaching 4 mm for the untreated case and 2 mm for the treated case. Since tensile strengths for these materials typically range from 3.6 MPa to 7.2 MPa, it is not likely that the actual stresses attained these values. Thermal cracking clearly provided a great deal of stress relief adjacent to the surface. Deeper within the speci-
men cores the model predicted high tensile stresses at or above the material tensile strength range. Since the remaining compressive strengths observed for these heated specimens were fairly high (see Table 2), it is clear that the tensile stresses predicted in the model were not attained. Revisiting the cracked region adjacent to the surface, one may take the 4 mm and 2 mm ranges of predicted high stresses, as demarcations of the beginning of the uncracked core in these cases. FIG. 9 contains an illustration of these cores. These core sizes were used to estimate the remaining core strength.

Thermal analysis results indicated that the crack penetration in the untreated case is up to 4 mm into the surface, whereas in the treated case it is only up to 2 mm. The stress in the core was calculated by taking the product of the apparent diameter stress and apparent diameter area and dividing by the core area. The apparent and core stress values for the untreated and treated cases are presented in Table 4. The estimated core strength of the treated specimen was increased by 17% compared to the case of the 0.5 w/c (the mass ratio of water to cement powder used to create the paste). A high value yields a very porous or weak hardened cement paste and thus a week concrete. 0.5 is considered a moderate value for the w/c ratio.

### TABLE 4

<table>
<thead>
<tr>
<th>Case</th>
<th>$S_{app}$ (MPa)</th>
<th>$S_{core}$ (MPa)</th>
<th>$S_{app}$ (MPa)</th>
<th>$S_{core}$ (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Untreated</td>
<td>11.7</td>
<td>16.5</td>
<td>12.5</td>
<td>17.6</td>
</tr>
<tr>
<td>Treated</td>
<td>16.4</td>
<td>19.3</td>
<td>27</td>
<td>31.8</td>
</tr>
</tbody>
</table>

The impact of the w/c ratio on the response to the nanoparticle treatment was significant. The 0.4 w/c ratio specimens responded with 65% more strength than 0.5 w/c ratio cases. From Table 2, experimental results showed that the compressive strength of the treated specimens after thermal exposure increased an overall average of 70% but the increase in resistance to thermal cracking was not clearly demonstrated.

Furnace test results showed both the treated and untreated specimens cracked at 550°F (288°C) following quenching. This work indicated that the nanoparticle treatment, applied to the 0.4 w/c and 0.5 w/c ratio cases, provided significant resistance to compressive strength degradation after thermal exposure. The results showed that there was an increase in compressive strengths at 550°F (288°C) for the treated cases. Finite element analysis indicated a crack-inducing tensile stress in untreated specimens up to a 4 mm depth into the surface whereas in the treated case it is only up to 2 mm. The estimated, uncracked core strength of the treated specimens exhibited an average value that was 50% higher than that of the untreated specimens. The Impact of w/c ratio on the response to the nanoparticle treatment was significant with the 0.4-w/c ratio specimens responding with 65% more strength than the 0.5-w/c ratio cases.

I claim:
1. A method of treating concrete comprising:
   a. providing a section of concrete having a depth, an initial compressive strength, and an initial pH;
   b. placing a first nanoparticle carrier fluid in contact with at least a first surface of said concrete section, said first nanoparticle carrier fluid comprising solid nanoparticles in a colloidal suspension;
   c. applying a first current at sufficient magnitude and for sufficient time to:
      i. remove at least a portion of a chlorine species present and/or raise the pH in said concrete section; and
      ii. move said solid nanoparticles in said first nanoparticle carrier fluid through at least one quarter of said depth of said concrete section, thereby returning the strength of said concrete section to at least said initial compressive strength;
   d. placing a second nanoparticle carrier fluid in contact with at least the first surface of said concrete section, said second nanoparticle carrier fluid comprising solid nanoparticles in a colloidal suspension, wherein said solid nanoparticles of the second nanoparticle carrier fluid are smaller in diameter than said solid nanoparticles of the first nanoparticle carrier fluid; and
   e. applying a second current at sufficient magnitude and for sufficient time to:
      i. remove at least a portion of a chlorine species present and/or raise the pH in said concrete section; and
      ii. move said solid nanoparticles in said second carrier fluid through at least one quarter of said depth of said concrete section.
2. The method according to claim 1, wherein said nanoparticles comprise a silicon species combined with a positive species such that said nanoparticles have a net positive charge.
3. The method according to claim 2, wherein said silicon species is silica.
4. The method according to claim 2, wherein said positive species is an oxide of aluminum and/or zirconium.
5. The method according to claim 3, wherein said positive species is a polymer.
6. The method according to claim 2, wherein said nanoparticles are alumina coated silica.
7. The method according to claim 1, wherein said first and second nanoparticle carrier fluids each include a stabilizing agent.
8. The method according to claim 7, wherein said stabilizing agent is acetate based.
9. The method according to claim 1, wherein said section of concrete includes reinforcing metal and said reinforcing metal acts as an electrode.
10. The method according to claim 9, said reinforcing metal is a cathode.
11. The method according to claim 1, wherein said first current has a density of at least about 100 mA per square foot.
12. The method according to claim 1, wherein said first current has a density of at least about 300 mA per square foot.
13. The method according to claim 1, wherein said strength of said concrete section is increased by at least 25 percent.
14. The method according to claim 1, wherein prior to placing said first nanoparticle carrier fluid in contact with said first surface, sufficient current is induced across said first surface for sufficient time to remove at least a portion of said chlorine species.
15. The method according to claim 1, wherein said nanoparticles are present in said first nanoparticle carrier fluid at an initiation of treatment.
16. The method according to claim 2, wherein said nanoparticles having a net positive charge are present in said first nanoparticle carrier fluid at an initiation of treatment.
17. The method according to claim 9, wherein the corrosion potential of said reinforcing metal is raised to more positive than about -0.2 volts.
18. A process for treating concrete comprising:
a. providing a section of concrete having a depth, an initial pH, and an initial strength;
b. applying a first stage of treatment comprising:
   i. inducing a first current across the concrete section;
   ii. applying the first current at sufficient magnitude and for sufficient time to remove at least a portion of a chlorine species present and/or raise the pH in the concrete section and electrokinetically transport a plurality of first nanoparticles through at least one quarter of the depth of the concrete section to increase the concrete section’s strength; and
c. then applying a second stage of treatment comprising:
   i. inducing a second current across the concrete section; and
   ii. applying the second current at sufficient magnitude and for sufficient time to electrokinetically transport a plurality of second nanoparticles into the concrete section, wherein the plurality of second nanoparticles are smaller in average diameter than the plurality of first nanoparticles.

19. A strength and anti-corrosion treatment process for concrete, the process comprising:
a. providing a section of concrete having a depth, an initial pH, and an initial strength; and
b. applying a current at sufficient magnitude and for sufficient time to:
   i) remove at least a portion of a chlorine species present and/or raise the pH in the concrete section; and
   ii) electrokinetically transport colloidal suspended solid nanoparticles through at least one quarter of the depth of the concrete section to increase the concrete section’s strength, wherein the colloidal suspended solid nanoparticles have diameters ranging between 1 nm to 500 nm and have solid, non-deformable cores.

20. The process according to claim 19, wherein the solid nanoparticles are selected from a group consisting of: alumina, silica, silicates, aluminates, latexes and polymers.

* * * *