Study on the Softening Mechanism and Control of Red-Bed Soft Rock under Seawater Conditions

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Abstract: Red-bed soft rock easily softens and disintegrates when it comes into contact with water, which is the main factor restricting the application of soft rock as an engineering filler. Therefore, research on the influence of seawater on soft rock softening has great significance for the application of soft rock in marine engineering. To examine the softening mechanism of soft rock under seawater conditions, two kinds of soft rock softening experiments, as well as ion inhibition tests of soft rock softening, were performed under seawater and pure water conditions, and the results were compared. The variation in the soft rock composition, the deformation and failure characteristics of soft rock under the influence of sea water, and the variation in main cations in the softening process of soft rock are examined successively; the influences of different ions on soft rock softening are further analysed. Based on the analyses, the softening mechanism and control method of soft rock under sea water conditions are expounded. The study showed that soft rock softening was inhibited by seawater, which decreased the softening degree of soft rock. The main cations in seawater had an inhibitory effect on soft rock softening, and the order of inhibition was Ca^{2+} > Mg^{2+} > Na^+. According to the inhibitory effect of ions on soft rock softening, we propose that seawater or calcium salt should be added to reduce the softening of soft rock in soft rock engineering and improve the mechanical strength of soft rock; in addition, soft rock can be considered a raw material in marine engineering. The experimental results have great significance for studies of the disintegration mechanism and inhibitory rules of soft rock under the influence of seawater and provide a theoretical basis for the application of soft rock in marine engineering, such as in artificial reef engineering and coastal dike engineering.

Keywords: red-bed soft rock; seawater; ion; softening mechanism; control

1. Introduction

In recent years, with the construction of man-made islands and reefs, maritime airports, undersea tunnels, undersea military bases and other marine projects, marine engineering has entered an era of great advancements. In engineering construction, a large amount of geotechnical material is often needed as engineering filler. Although red rock is widely distributed in coastal areas, red rock layers easily soften and collapse in water, and so, in red rock distribution areas, especially coastal areas, the widely distributed soft rock. Alternatively, inland granite, basalt and other geotechnical materials are chosen as engineering raw materials, resulting in high transportation and time costs. Large amounts of manpower and material and considerable financial resources are consumed, and the protection of the natural ecological environment and sustainable development will also be greatly impacted. However,
if soft rock and local materials can be used according to local conditions, project costs will be greatly reduced, and the impact on the natural environment will be minimized to the greatest extent. Therefore, for the widely distributed red-bed soft rock, an application objective in marine engineering is to solve the problems of softening and disintegrating under seawater conditions. In the past, research on soft rock softening control has mainly focused on soft rock modification experiments [1–13] and soft rock deformation control methods [14–24]. In general, durability characteristics are tested prior to design in seawater conditions. Through theoretical research, e.g., Li et al. [9,10], the feasibility of modifying soft rock by electrochemistry has been assessed according to the physical, chemical and mechanical properties of the soft rock itself. Two mechanisms of the electrochemical modification of clay minerals are reviewed: the stability mechanism of electroosmotic dehydration and cation substitution and the mechanism of structural and property changes. Liu et al. [5] selected a salt solution as the hydrolysate to conduct collapse resistance tests of red sandstone and proposed that the improvement by mixing with 7% lime is the best engineering measure to restrain the collapse of class I red sandstone. Liu [11] studied the effect of the solution concentration on the expansion and deformation of mudstone. A KCl solution effectively inhibited the expansion and deformation of reconstructed mudstone samples. Yang et al. [12] used chemical modification, cement modification and composite modification to change the structure of soft rock from a microscopic point of view and found that composite modification can form a hydrophobic film on the surface of rock particles. Furthermore, the cementation force between particles was improved. In addition to indoor modification tests of soft rock, scholars have performed road use research on soft rock [25–29]. These studies provide a solid theoretical basis for the control of soft rock softening. If soft rock softening control methods can be used in marine engineering construction, it will greatly aid marine engineering involving soft rock.

Research on the deformation control of soft rock mainly focuses on the failure mechanism and deformation control of soft rock roadways. Numerical experiments such as those by Wang et al. [14] are carried out on the basis of considering the arch strength, ground stress, mechanical parameters of the surrounding rock, deformation of the surrounding rock, range of the plastic zone and the mechanical properties of the supporting unit by comparing and analyzing different factors. Thus, the failure and control mechanisms of deep soft rock roadways are determined. Yuan et al. [15] revealed the compound failure mechanism of a Mesozoic soft rock roadway in the Shajihai mining area, which involved the molecular expansion, shear slip and structural disturbance of the weak structural surface, and proposed coupling support technology with large-deformation constant bolt resistance as the core. Li et al. [21] proposed that the instability mechanism of soft rock roadways caused by a high content of confined water consists of fracturing, seepage, silting and closing processes based on two experimental cycles of the surrounding rock; these processes decrease rock strength and lead to roadway instability. However, at present, experimental studies on soft rock softening control under seawater conditions are still limited to the damage characteristics and stability analysis stage of coastal rock bank slopes [30–35].

Therefore, this paper takes sandy mudstone as the research object, and saturated tests of soft rock under seawater conditions are performed to study the evolution of the soft rock strength under the influence of seawater and the variation in the main cations in solution during the softening process of soft rock; in addition, the characteristics of composition change, deformation and failure of the soft rock are examined. The main cations are studied, and the effects of the different ions on soft rock softening are examined. Based on the results, the softening mechanism and the softening control method of soft rock under seawater conditions are expounded. The study provides experimental and theoretical bases for the study of the effective utilization of soft rock in marine engineering.
2. Soft Rock Softening Test

2.1. Softening Test of Soft Rock under Seawater Conditions

2.1.1. Purpose of the Test

The purpose of this experiment is to study the macroscopic and microcosmic variations in red-bed soft rock under seawater conditions and to examine the inhibitory mechanism of seawater on soft rock softening.

2.1.2. Test Content and Process

The sandy mudstone widely distributed in South China is selected as the research object. The geologic age of the red sandy mudstone we used is the cretaceous period. The main minerals of the sandy mudstone are quartz, calcite, anorthite, tremolite and clay minerals. In this experiment, the sandy mudstone was made into standard 50 $\times$ 100 mm cylinders. The element composition of three selected rock samples is determined via scanning electron microscopy, the energy spectrum and the electron backscattering diffraction system in a thermal field emission environment. According to the energy spectrum, the contents of C, O, K, Al, Fe, Si and other elements are measured for the three samples of natural soft rock. Table 1 provides the percentage of elements in each rock sample. The sum of the Si and Ca contents in the three rock samples is approximately 90%.

The test process mainly includes the following:

1. The rock samples were immersed in two kinds of soaking solutions for 30 days. The pH value and main cationic concentration of the solution were measured in the first 7 days, and the pH value was measured every 12 h, while the ion concentration was measured every 24 h.

2. During the initial 24 h of soaking, macroscopic changes in the soft rock were observed at intervals of 1 h, and the fracture development and disintegration were recorded.

3. A uniaxial compressive strength test was carried out on the natural rock samples after 30 days of immersion.

2.2. Inhibition Testing of the Ions on Soft Rock Softening

2.2.1. Testing Programme

The concentrations of the four main cations in seawater are in the order of Na$^+$ > Mg$^{2+}$ > Ca$^{2+}$ > K$^+$, and so MgCl$_2$, CaCl$_2$ and NaCl are selected as the dissolved salts in the experiments, and K$^+$ is not considered in the experiments.

The experiments consisted of observing the effects of MgCl$_2$, CaCl$_2$ and NaCl solutions with two different concentrations of 0.1 mol/L and 1 mol/L on soft rock softening and to examine the inhibitory effect of the different ions and ion concentrations on soft rock softening.

Because it is difficult to maintain complete consistency in sample concentration and solute solubility in the experiments, the following assumptions are made:

---

Table 1. Relative content percentage of elements in the rock samples.

<table>
<thead>
<tr>
<th>Sample Label</th>
<th>C</th>
<th>O</th>
<th>Mg</th>
<th>Al</th>
<th>Si</th>
<th>K</th>
<th>Ca</th>
<th>Fe</th>
</tr>
</thead>
<tbody>
<tr>
<td>01</td>
<td>11.79</td>
<td>48.46</td>
<td>0.52</td>
<td>5.01</td>
<td>21.21</td>
<td>2.51</td>
<td>7.85</td>
<td>2.66</td>
</tr>
<tr>
<td>02</td>
<td>13.73</td>
<td>49.33</td>
<td>0.37</td>
<td>3.43</td>
<td>16.24</td>
<td>1.49</td>
<td>14.44</td>
<td>0.96</td>
</tr>
<tr>
<td>03</td>
<td>12.89</td>
<td>49.54</td>
<td>—</td>
<td>3.43</td>
<td>16.24</td>
<td>1.49</td>
<td>13.91</td>
<td>0.64</td>
</tr>
</tbody>
</table>
After preparation of the solution was completed and maintaining the solution for 30 min, the initial concentration of the solution remained constant, and it was assumed that there was no solute precipitation.

After each sampling, it is assumed that the concentration of the solution remains consistent before and after sampling and does not change with the change in volume.

2.2.2. Test Process

The ion variations in each ionic solution were measured every 24 h for 7 days, and the softening coefficient of the rock samples after 30 days of immersion was calculated. Then, the inhibitory influences of the ions on soft rock softening were compared and analysed.

3. Softening Law of Soft Rock under Seawater Conditions

3.1. Effect of Seawater on Soft Rock Softening

3.1.1. Macroscopic Variation Law of the Rock

Figure 1 shows the macroscopic changes in the soft rock after soaking for 2 h, 6 h and 18 h.

![Macroscopic changes in soft rock softening](image)

After considering and referring to some standard test methods [36–38], the macroscopic changes in the soft rock softened in seawater and pure water are compared, and when the rock sample is immersed for 2 h, the rock sample clearly collapses, and soft rock-caving occurs at the bottom of beaker. Fine debris particles are observed in the seawater, but in pure water, powder is observed. After soaking for 6 h, there are distinct cracks on the surface of the soft rock immersed in pure water, while in seawater, debris flakes can easily be seen on the surface of the soft rock, but there are no cracks on the surface of the soft rock.

![Macroscopic changes in soft rock softening](image)
With the progress of softening and disintegration, the cracks on the surface of the soft rock in pure water gradually expand. After 18 h, cracks appeared on the upper surface of the soft rock in seawater, while the cracks in pure water were more developed. From these observations, it can be deduced that the soft rock softening rate is higher, the water-rock interactions are stronger, and the degree of disintegration is higher in pure water.

3.1.2. Change in the Ion Concentration

The largest concentration changes in Ca$^{2+}$, Mg$^{2+}$, Na$^+$ and K$^+$ are mainly determined. The concentration curves of the ions with time are shown in Figures 2 and 3.

![Figure 2](image-url) Concentration change curves of the main cations after pure water immersion.

![Figure 3](image-url) Concentration change curves of the main cations after seawater immersion.

The curves show that the change in cationic concentration of the aqueous solution during sample saturated with water has the following characteristics:

1) In pure water immersion, the dissolution of soft rock is very strong, large amounts of Ca$^{2+}$, Mg$^{2+}$, Na$^+$ and K$^+$ are ionized, and the concentration of each ion increases. These ions are the main components of soluble minerals, which are often present in the ionic state in solutions. When sample immersion lasts approximately 120 h, the dissolution of minerals gradually weakens, the upward trend of the ion concentration slows down, and ion exchange increases.

2) In seawater immersion, there are high concentrations of Ca$^{2+}$, Mg$^{2+}$, Na$^+$ and K$^+$ in the initial solution. With the process of softening, the cations are continuously consumed under the actions of water and rock, and the ion concentration decreases gradually. When the initial solution...
reaches a certain saturated state, the softened soft rock tends to be stable, and the concentration of metal cations in the solution remained stable.

(3) Through the variation curve of the cationic concentration of the solution during the soft rock saturated with water, it is found that the concentrations of Ca\(^{2+}\), Mg\(^{2+}\), Na\(^{+}\) and K\(^{+}\) change greatly, which indicates that the softening and disintegration of soft rock in seawater may be related to the change in the main ion content in seawater.

3.1.3. Uniaxial Compressive Strength Test

With reference to the relevant experimental standards [17, 39], it is difficult to sample and conduct testing, but good results are obtained with respect to sampling success rate and disturbance resistance with self-made sampling equipment. The uniaxial compressive strength of the soft rock samples and the natural state of the soft rock samples soaked for 30 days under the two water conditions are tested. The results are provided below (see Table 2).

<table>
<thead>
<tr>
<th>Group</th>
<th>Compressive Strength (MPa)</th>
<th>Average Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Natural</td>
<td>25.8</td>
<td>26.7</td>
</tr>
<tr>
<td>Pure Water</td>
<td>10.02</td>
<td>10.21</td>
</tr>
<tr>
<td>Seawater</td>
<td>13.36</td>
<td>13.97</td>
</tr>
</tbody>
</table>

The experimental results (see Figure 4) show that the uniaxial compressive strength of the rock samples immersed in pure water and seawater decreases notably, which indicates that the overall strength of soft rock decreases after water-saturated. At the same time, according to the experimental results, the uniaxial compressive strength of the soft rock in seawater is 13.67 MPa, which is higher than that of the rock sample in pure water (10.12 MPa). From the shape variable, it can be concluded that, in the natural state, the strain of the rock sample at the time of failure is greater than that after water-saturated.

Because the whole test process is conducted under still indoor water conditions, the influences of other physical factors are excluded, and the results of the collapse and strength tests of the soft rock under pure water and seawater conditions are compared and analysed. It is considered that a series of physical and chemical reactions have taken place between the main cations and the soft rock in seawater under the actions of water and rock, which has suppressed soft rock softening.
3.2. Inhibitory Influences of the Ions on the Softening of Soft Rock

3.2.1. Variations in Ion Concentration

Due to the large concentration difference between the different ions, the data are incrementally processed based on the initial values.

That is, the increment is

\[ \Delta = N_{t=1} - N_{t=0} \]  

(1)

where \( N_{t=1} \) denotes the solute ion concentration and \( N_{t=0} \) represents the initial ion concentration.

The ion increment curves of the three solutions with concentrations of 0.1 mol/L and 1 mol/L, are shown in Figures 5 and 6, respectively.

As shown in Figures 5 and 6, the growth rates of the \( \text{Na}^+ \) and \( \text{Ca}^{2+} \) concentrations are basically consistent, but the growth rate of \( \text{Mg}^{2+} \) is lower than those of the first two ions. The author proposes the following two reasons:

(1) As shown in Table 1, Ca is the main component of soft rock, which occurs in the calcareous cement that connects the clay particles. When the soft rock encounters water, the soluble calcareous cement dissolves. The concentration of \( \text{Ca}^{2+} \) increases at a high rate in the early stage of softening.
The rapid increase in the Na⁺ concentration is related to the dissolution of soluble minerals, and it is possible that albite dissolves and Na⁺ precipitates.

\[2\text{NaAlSi}_3\text{O}_8 + 9\text{H}_2\text{O} + 2\text{H}^+ \leftrightarrow \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4 + 4\text{H}_2\text{SiO}_4 + 2\text{Na}^+\]  \hspace{1cm} (2)

(Dissolution of albite)

(2) In the MgCl₂ solution, the lattice substitution of Mg²⁺ is limited to the montmorillonite structure, and the negative charge formed after substitution is compensated by Na⁺ and Ca²⁺. In the NaCl and CaCl₂ solutions, the amounts of Na⁺ and Ca²⁺ consumed to compensate the negative charges are smaller than that of Mg²⁺ consumed by substitution. Moreover, the Mg content in soft rock is low, and the amount of Mg²⁺ formed by the dissolution reaction is also low.

For the above two reasons, the concentration of Mg²⁺ is lower than those of Na⁺ and Ca²⁺.

In addition, the ion concentration variations for different concentrations of the same ionic solution are plotted in Figures 7–9.

![Figure 7](image1.png)

**Figure 7.** The Na⁺ concentration increase curve for the NaCl solution.

![Figure 8](image2.png)

**Figure 8.** The Mg²⁺ concentration increase curve for the MgCl₂ solution.
According to the ion concentration increment curve for each solution, the increase in ion concentration gradually slows with time and tends to stabilize or even decrease, as shown in Figure 8 for example. In the 1 mol/L MgCl$_2$ solution, when the softening time reached 120 h, the concentration of Mg$^{2+}$ stopped increasing; on the contrary, there was a small decrease. It can be considered that in the beginning, the soft rock is softened mainly by the dissolution of soluble minerals, and the Mg minerals in the soft rock dissolve by the solution to form Mg$^{2+}$. When rock softening has proceeded for approximately 120 h, the soluble minerals on the surface of the soft rock are almost completely dissolved. As a result, dissolution decreases, but the exchange of adsorbed ions is enhanced, and lattice replacement consumes the free Mg$^{2+}$, which reduces the concentration of Mg$^{2+}$ in the solution; hence, there is a small decrease in the concentration. Because of the continuous actions of water and rock, mineral dissolution and ion exchange-adsorption always occur at the same time, so it can be predicted that the concentration of Mg$^{2+}$ will be in a dynamic equilibrium state in the continued softening process, but will eventually stabilize.

3.2.2. Analysis of the Softening Coefficient

The evaluation of the mechanical properties of soft rock is mainly characterized by the softening coefficient. The rock-softening coefficient refers to the ratio of the uniaxial compressive strength when the rock is saturated to the compressive strength in the dry state, which is represented by $\eta$. Because the compressive strength of rock after saturated with water is lower than that in the dry state, the softening coefficient is generally less than 1, and the expression is as follows:

$$\eta = \frac{R_s}{R_d}$$

where $R_s$ denotes the average uniaxial compressive strength in the saturated state and $R_d$ denotes the average uniaxial compressive strength in the dry state.

After 30 days of immersion, uniaxial compression tests were carried out on the three groups of soft rocks, and the results are shown in Table 3 below.
Table 3. Uniaxial compressive strength comparison table.

<table>
<thead>
<tr>
<th>Solution</th>
<th>Compressive Strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.1 mol/L</td>
</tr>
<tr>
<td>NaCl</td>
<td>11.68</td>
</tr>
<tr>
<td>CaCl₂</td>
<td>13.67</td>
</tr>
<tr>
<td>MgCl₂</td>
<td>12.86</td>
</tr>
</tbody>
</table>

The experimental results (see Table 4) show that the saturated uniaxial compressive strength of soft rock in the 1 mol/L solution is clearly higher than that in the 0.1 mol/L solution. For the different solutions with the same concentration, the saturated uniaxial compressive strength of the soft rock in the CaCl₂ solution is higher than that in the MgCl₂ solution, and that in the MgCl₂ solution is higher than that in the NaCl solution, but the uniaxial compressive strength of these groups of soft rock is higher than that of the soft rock in pure water to a certain extent. This phenomenon shows that the CaCl₂ solution maintains the strength of soft rock to a certain extent, followed by the MgCl₂ and NaCl solutions.

Table 4. Softening coefficient comparison table.

<table>
<thead>
<tr>
<th>Solution</th>
<th>Softening Coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.1 mol/L</td>
</tr>
<tr>
<td>NaCl</td>
<td>0.44</td>
</tr>
<tr>
<td>CaCl₂</td>
<td>0.52</td>
</tr>
<tr>
<td>MgCl₂</td>
<td>0.49</td>
</tr>
</tbody>
</table>

According to the above analysis and the softening coefficients, it can be concluded that the order of the ionic inhibitory effect on soft rock is \( \text{Ca}^{2+} > \text{Mg}^{2+} > \text{Na}^+ \), and the higher the ion solubility is, the more notable the softening effect on soft rock is.

Through regular analysis, the ionic inhibition on soft rock softening may be related to the following influencing factors of ion exchange-adsorption:

1. \( \text{Ca}^{2+} \) and \( \text{Mg}^{2+} \) are bivalent cations. In the lattice substitution of montmorillonite and kaolinite, the exchange capacity of divalent ions is greater than that of the monovalent \( \text{Na}^+ \).
2. According to the influence of the ion radius on the exchange capacity, because the order of the ion radius is \( \text{Ca}^{2+} > \text{Mg}^{2+} > \text{Na}^+ \), the order of the ion exchange capacity is \( \text{Ca}^{2+} > \text{Mg}^{2+} > \text{Na}^+ \).
3. In the three solutions, the higher the concentration of the ionic solution is, the more soft rock softening is inhibited.

4. Softening Mechanism of Soft Rock under Seawater Conditions

According to the experimental analysis and the characteristics of clay minerals, the softening mechanism of soft rock under seawater conditions includes the following processes: water absorption, expansion and disintegration of clay minerals, ion exchange-adsorption, and dissolution of soluble minerals.

4.1. Mechanism of Water Absorption, Expansion and Disintegration of the Clay Minerals

1. Because kaolinite, Illite and other clay minerals have small particles and are strongly hydrophilic, when interacting with seawater, the water molecules enter the layered clay mineral particles and form polarized water molecular layers between the particles. These water molecular layers can continuously absorb water and expand, resulting in an increasing gap between the particles. At the same time, water molecules enter the unit cell interlayers of the clay minerals, forming an interlayer water layer in the minerals. Relatively speaking, it is easier for water molecules
to enter the interparticle gap than to enter the interlayer of each particle. The former causes the 
external expansion of clay minerals, called intragranular expansion, and the latter causes internal expansion, called intra-particle expansion. Existing studies [40] show that the physical and chemical reactions between Illite and water can cause the expansion of soft rock, which can increase the original volume by 50% to 60%. The chemical reaction process is as follows:

\[
K_{0.9}Al_{2.9}Si_{3.1}O_{10}(OH)_2 + nH_2O \rightarrow K_{0.9}Al_{2.9}Si_{3.1}O_{10}(OH)_2 \cdot nH_2O
\]

(4)

(2) According to the two kinds of expansion of clay minerals, the clay particles of soft rock can absorb much water during water immersion, which increases the unit cell distance or thickens the diffusion layer, and the minerals expand. When the expansion stress exceeds the bonding effect, the clay cements are destroyed and disintegrated, while the clastic particles gravitationally disintegrate due to the lost connections. In addition, due to the uneven water absorption and expansion of clay minerals, an uneven stress distribution gradually develops in the soft rock, resulting in a large number of micropores, which destroy the original internal structure of the rock sample. Finally, the phenomenon leads to the fragmentation and disintegration of rock particles.

4.2. Ion Exchange-Adsorption

In the early stage of the interaction between soft rock and seawater, the water-rock interactions are mainly the dissolution of soft rock and cation exchange-adsorption, and the dissolution of soluble minerals is gradually weakened and tends to be stable due to the high ion concentration in the solution.

Second, the process of ion exchange-adsorption between water and rock is affected by many factors, such as the mineral composition, structure compactness, pH value and chemical composition or concentration. In an environment of no energy exchange or less energy exchange with the outside world, the total charge of cations adsorbed by clay minerals remains unchanged, and most of the exchange and adsorption of ions is carried out under these conditions, and the ion exchange between water and clay minerals will reach an equilibrium. When the hydrochemical environment changes, this balance will be disturbed, so that the ions will spread rapidly in a short period of time and ion exchange and adsorption will occur again, and the structure of soft rock will also change. Therefore, when the structure of soft rock changes, clay minerals disintegrate, or physicochemical changes result in imbalanced charges, the adsorption and exchange of ions will be dynamically balanced.

Compared with pure water, seawater is equivalent to changing the external hydrochemical environment of soft rock, resulting in a residual negative charge of the clay minerals due to homomorphism, and the non-saturated negative charge between the crystal layers makes the clay particles have strong surface properties. The residual negative charge is balanced by the adsorption of cations such as K\(^+\), Na\(^+\), and Ca\(^{2+}\) from the seawater, leading to a change in the interlayer spacing, which affects the suction and repulsive force between particles and affects the physical and mechanical properties of the clay minerals.

The factors affecting ion exchange-adsorption are as follows:

1. Ion valence number: the larger the ion valence number is, the more prone to ion exchange;
2. Ion radius: the larger the ion radius is, the smaller the ion hydration radius is, the stronger the adsorption and exchange capacities of ions are. Usually, the order of the ion exchange capacity from weak to strong is as follows:

\[
\text{Li}^+ < \text{Na}^+ < \text{K}^+ (\text{NH}_4^+) < \text{Mg}^{2+} < \text{Ca}^{2+} < \text{Ba}^{2+} < \text{Al}^{3+} < \text{Fe}^{3+} < \text{H}^+
\]

3. Ion concentration: the higher the ion concentration is, the stronger the exchange capacity.
4.3. Dissolution and Corrosion of the Soluble Minerals

In addition to macroscopic expansion and disintegration and microscopic ion exchange and adsorption, the water-rock effects can cause the dissolution or corrosion of soluble minerals, resulting in a change in the structure of the soft rock. According to the variation in the ion concentration of the soft rock before and after immersion in seawater, the variation range of active ions in the aqueous solution, such as Na$^+$, Ca$^{2+}$, K$^+$, and Mg$^{2+}$, is large. Therefore, it can be inferred that the dissolution and corrosion of minerals occur in the interaction process between soft rock and water.

The possible dissolution reactions \[41,42\] are as follows:

\[
2\text{KAl}_3\text{Si}_3\text{O}_8 + 9\text{H}_2\text{O} + 2\text{H}^+ \leftrightarrow \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4 + 4\text{H}_4\text{SiO}_4 + 2\text{K}^+ \tag{5}
\]

(Dissolution of potash feldspar)

\[
2\text{NaAlSi}_3\text{O}_8 + 9\text{H}_2\text{O} + 2\text{H}^+ \leftrightarrow \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4 + 4\text{H}_4\text{SiO}_4 + 2\text{Na}^+ \tag{6}
\]

(Dissolution of albite)

According to the above possible reactions, the pH value during soft rock softening in seawater changes as follows (Figure 3): the pH value increases gradually, then decreases gradually, and then remains stable. The author believes that in the early stage of soft rock saturated with water, clay minerals absorb water and expand when they are immersed in water, and soluble minerals such as potassium feldspar and albite dissolve, thereby consuming the H$^+$ in water and precipitating Na$^+$ and K$^+$; thus, the pH value of the solution increases gradually in the beginning stage. When the concentrations of the Na$^+$ and K$^+$ produced by the chemical reactions are too high, the ions are adsorbed onto the interlayer surfaces of the crystal layers of clay minerals, which balances the excess negative charges, and enhances the bonding force between the layers. The dissolution or corrosion of minerals is weakened, and the precipitation of metal cations is reduced. Thus, the pH value of the solution decreases. When the two processes reach an equilibrium, the pH value in the solution will remain at a stable value, and the pH value of this experiment is finally maintained at approximately 7.5–7.6. The analytical results are in good agreement with the experimental results.

5. Soft Rock Softening Control under Seawater Conditions

5.1. Control Principle Based on the Solubility Product Rule and Same Ion Effect

According to the previous tests, Ca$^{2+}$, Mg$^{2+}$, K$^+$, Na$^+$ and other main mineral ions dissolve and precipitate in the process of soft rock softening. The presence of large amounts of Ca$^{2+}$, Mg$^{2+}$, Na$^+$, and K$^+$ in seawater is considered based on the solubility product rule in basic chemistry \[43\]. The precipitation and dissolution of any electrolyte in solution is subject to a dynamic equilibrium. The index of the dissolving capacity of a material is the solubility product constant $K_{sp}$, and $K_{sp}$ is the product of the coefficient power of the ionic concentration of the saturated solution of the insoluble electrolyte at a certain temperature. For example, the ionization equilibrium of insoluble electrolyte $X_aY_b(s)$ is:

\[
X_aY_b(s) \rightleftharpoons aX^{m+} + bY^{n-} \tag{7}
\]

When the system is in equilibrium, the system obeys the law of chemical equilibrium:

\[
[X^{m+}]^a [Y^{n-}]^b = K_{sp} \tag{8}
\]

The product of the power of ion concentration coefficients of the insoluble electrolytes in solution is called the ion product $Q_i$:

\[
Q_i = c_{X_{m+}}^a c_{Y_{n-}}^b \tag{9}
\]
(1) \( Q_i = K_{sp} \): the solution is a saturated solution with no precipitation occurring and has obtained
dynamic equilibrium;

(2) \( Q_i < K_{sp} \): the solution is not saturated and there is no precipitation. If there are solids in the
system, the solids will dissolve until the solution becomes saturated.

(3) \( Q_i > K_{sp} \): the solution is oversaturated, and the equilibrium moves to the left with increasing
precipitation, while the ion concentration in the solution is reduced depending on the saturation
of the solution.

According to the solubility product rule, to inhibit the dissolution of the minerals of the sandy
mudstone during immersion, it is necessary to increase the concentration of dissolved ions in the
solution in such a way that the ion product \( Q_i > \) solubility product \( K_{sp} \). Soluble electrolytes containing
the same ions should be added to the solution to force the equilibrium to move to the left, which
achieves the same ionic effect [44].

Therefore, based on the softening inhibition test results, because there is a large amount of free
\( \text{Ca}^{2+}, \text{Na}^+ \) and \( \text{Mg}^{2+} \) in the three solutions, according to the solubility product rule, the authors think
that during soft rock saturated with water, the presence of metal cations in the solution suppresses the
dissolution of the soluble clay minerals with the same ions. Due to the aforementioned ionic effect, the
equilibrium moves to the left, the dissolution of clay minerals decreases, the structure of the soft rock is
more stable, and the bonding force between the particles is more intense. The collapse resistance of soft
rock is improved, and the strength remains higher. According to the same ionic effect, the inhibitory
effect of each ion solution on soft rock softening in the tests can be well explained.

Previous experiments have proven that at different concentrations of the same ions,
high-concentration ionic solutions have a greater inhibitory effect on soft rock softening. The same
ionic effect is explained as follows. Taking the 1 mol/L and 0.1 mol/L \( \text{CaCl}_2 \) solutions as examples, it
can be understood that solid \( \text{CaCl}_2 \) can be added to the 0.1 mol/L \( \text{CaCl}_2 \) solution under the condition
that the soft rock and other external factors are consistent to increase the \( \text{CaCl}_2 \) concentration of the
solution to 1 mol/L. In the process of continuously increasing the \( \text{Ca}^{2+} \) concentration, when \( Q_i > K_{sp} \) is
satisfied, the equilibrium moves to the left and the concentration of \( \text{Ca}^{2+} \) decreases through calcareous
precipitation. This process promotes precipitation, suppresses the dissolution of minerals and ensures
the stability of the soft rock structure. It is verified that a high-concentration ionic solution has a greater
inhibitory effect on soft rock.

5.2. Soft Rock Softening Control Method under Seawater Conditions

According to the inhibition test of the ionic solution on the softening of red rock, the results
show that high contents of \( \text{Na}^+ \), \( \text{Ca}^{2+} \) and \( \text{Mg}^{2+} \) have a certain inhibitory effect on the softening
and disintegration of soft rock in seawater, and the higher the concentration is, the more notable the
inhibitory effect. In this experiment, the order of the inhibitory effect of the ions is \( \text{Ca}^{2+} > \text{Mg}^{2+} > \text{Na}^+ \),
so the control method of soft rock softening under seawater conditions can be studied from the point
of view of the ions.

(1) When a project is executed in a coastal area, a certain proportion or volume of seawater can be
considered a soft rock softening inhibitor, which improves the mechanical properties of the soft
rock packing structure to a certain extent. For marine engineering or coastal engineering, when
the engineering strength requirement is not high, soft rock can be considered as foundation filling
for the project, which will greatly reduce the construction cost.

(2) In addition, when the project is executed in an inland area, a certain amount of calcium salt can
be added to the soft rock packing structure, and the overall stability and mechanical properties of
the soft rock can be improved by relying on the inhibitory effect of calcium ions on the softening
of soft rock.
6. Conclusions

In this paper, sandy mudstone is the research object, and soft rock softening tests under pure water and seawater conditions and soft rock softening inhibition tests of ionic solutions are designed and carried out, and the following conclusions are drawn:

(1) In the same external environment, seawater has a certain inhibitory effect on the softening of soft rock compared with pure water, and the inhibitory effect is related to the main cations. The order of the inhibitory effect was \( \text{Ca}^{2+} > \text{Mg}^{2+} > \text{Na}^+ \).

(2) According to the inhibitory effect of the main cations on soft rock, calcium salt can be added to soft rock, supplemented by a cement slurry, silicone and other traditional chemical modifiers to modify the soft rock to reduce its softness and the disintegration of soft rock.

This research can provide guiding references and establishes the theoretical basis for the application of soft rock filling and softening control in marine artificial islands, reef engineering and coastal embankment engineering projects and other types of marine engineering construction projects.

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**References**


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