Research Article
Strength and Solidification Mechanism of Silt Solidified by Polyurethane

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Received 15 March 2020; Revised 14 August 2020; Accepted 23 August 2020; Published 18 September 2020

1. Introduction

Due to river transportation, the low-liquid-limit silt is widely distributed in the middle and lower reaches of the Yellow River, Huaihe River, and Haihe River in China. Such silt is often used as roadbed filler in construction in these regions. The soil is, however, considered poor subgrade soil in the construction industry due to its high silt content, its poor water stability, and the difficulty to compact it. Consequently, developing silt solidification methods appears particularly important [1]. However, there are only a few studies on the solidification of silt and most of them use the soil curing agent to stabilize the clay. For example, Alban et al. [2] proved the feasibility of stabilizing silt using cement and lime. Kolas et al. [3], Cheng et al. [4], and Rios et al. [5] stabilized clay and expansive soil with high-calcium fly ash, alkali-activated cement, and lime pozzolan, respectively, as the curing agent and explained the solidification mechanism from a microscopic perspective. However, the production of calcium-based inorganic substances like cement is very energy-consuming. Every ton of cement that is produced consumes 5000 MJ of energy; most of this is produced from primary energy sources, such as coal. In addition to pollution such as dust, the production of cement also emits massive amounts of CO₂ and SO₂; these separately aggravate the greenhouse effect, produce acid rain, and cause serious pollution to the environment [6, 7]. Many efforts have been made to develop green soil curing agents, and the most widely used ones currently include biological enzymatic curing agents, ionic curing agents, and organic polymer curing agents. Rafique et al. [8] introduced biological enzymes to reinforce silt roadbeds and evaluated the stabilization from the California Bearing Ratio (CBR) value. He et al. [9] and Junshuai Huo et al. [10] developed liquid ionic soil stabilizer (LISS) and EN-1 ionic soil curing agents to stabilize expansive soil and sandstone, respectively, and improved their UCS. Subsequently, Cai et al. [11] proposed a method to solidify soil with MgO and used it on the alluvial
silt of the Yellow River. The UCS of the silt was 5.1 MPa after 28 days of curing, which was a significant improvement over other similar soil curing agents.

However, the performance of soil curing agents is often improved by extending the curing time, which is not compatible with emergency construction projects. As a result, organic polymer curing agents were developed for such applications [12]. Initially, Masoumi et al. [13] and Rezaeimalek et al. [14, 15] used polypropylene fibers, polyvinyl acetate resins, styrene-acrylic liquid polymers, and methylene diphenyl diisocyanate (MDI) to cure sandy soil, but the strength was still dependent on the curing time. Therefore, Liu et al. [16] proposed to solidify sand–clay mixtures with polyurethane polymers. The strength and the erosion resistance of the solidified soil were significantly improved due to the high early strength of polyurethane. Subsequently, Wei et al. [17] added sisal fiber to polyurethane to stabilize the sandy soil and investigated the strengthening mechanism from a microscopic perspective. Zhang et al. used lignin, a byproduct of the paper industry, to strengthen soil and studied the physical and mechanical properties of the stabilized soil [18]; the results showed that the byproduct lignin exhibited a satisfactory performance of improving engineering properties of both cohesive soils and noncohesive soils with respect to strength, erosion resistance, and durability [19]. They established a model to study the relationship between the energy required during the plane shear process and the external workload and proved the effectiveness of their solidified soil through field tests [20, 21].

Most of the current research on soil stabilization focuses on clay, sandy soil, and expansive soil, whereas the studies on silt remained scarce. Therefore, a soil curing agent based on a permeable polyurethane polymer to improve the stability of silt roadbeds was developed. It has a low viscosity and a high early strength. UCS tests were conducted along with a microscopic analysis to evaluate the solidification properties and discuss the solidification mechanism of the polymer-treated silt. Our results provide a technical reference for the rapid and stable construction of roadbeds in silt-rich regions.

2. Materials and Methods

2.1. Test Materials. The polyurethane polymer used in the test was the 9802V polymer produced by the Henan Road Inspection Engineering Technology Research Center. Figure 1 illustrates that the polymer is a two-component curing agent, where the main component of material A is polyol with a density of 1.26 g/cm$^3$, and the main component of material B is polymer isocyanate with a density of 1.17 g/cm$^3$. The material is mainly composed of C, H, O, and N. Material A is mainly composed of hydrophilic low-viscosity polyether polyol, permeable diluent, and surfactant (nNCO-R-OCN); material B is mainly composed of isocyanate (nHO – OH).

Both materials were mixed with a mass ratio of 1:1 to form a polyurethane polymer. The curing time of the polymer is controlled by a catalyst, and its curing time at normal temperature ranges from 10 s to 24 h. In this study, the curing time of the polymer was set to 6 h. The mercury intrusion porosimetry (MIP) test was conducted by a mercury intrusion meter, namely, AutoPore IV 9500.

2.2. Preparation of the Specimens. The silt used in this test was taken from a depth of 5 m in a construction site in Zhengzhou. Table 1 shows the basic physical properties of the soil, and Figure 2 shows the distribution of the particle size of the soil. The soil used in this work was low-liquid-limit silt, as determined by accepted methods [22].

In addition, a mixing-compaction method was used to prepare the specimens. First, the plain soil was dried, broken, and filtered by a 2 mm sieve to obtain the test soil since the influence of particle gradation was ignored in this work. According to the preliminary test results, when the moisture content of the soil was less than 8% or the polymer content was more than 15%, it was difficult to form the specimens. When the polymer content was less than 5% or the moisture content of the soil was more than 15%, the distribution of the polymer in the soil was uneven, and the strength was only slightly higher than that of the specimens without the polymer. Soil samples were prepared with different moisture contents of 8%, 10%, 13%, and 15%; then different amounts of curing agent were added (5%, 10%, 13%, and 15%). Next, the curing agent containing materials A and B was mixed with a mass ratio of 1:1, and the product was added to the previously prepared soil samples with different moisture contents and mixed thoroughly. A triaxial compactor was used to prepare cylindrical specimens with a diameter of 39.1 mm and a height of 80 mm (Figure 3(a)). Three specimens were prepared for each set of parameters (moisture content, polymer content, and immersion time). Three specimens with the same parameters were prepared; a total
of 240 specimens were prepared and placed in a ventilated and dry place for 24 h. The reaction was the self-reaction of the two materials, A and B, and it occurred relatively quickly. Unlike cement-based materials, there was no need for complicated maintenance conditions; the product will have a certain strength after being left to stand for 24 h. And according to the results of the preliminary test, changing the curing conditions within 24 hours has little effect on its performance. This is also an advantage over other materials.

2.3. Test Method. To study the effect of the polymer on the solidification of the silt in a water environment, the specimens were immersed in water after standing for 24 h. The UCS test was carried out on specimens immersed in water for 1, 7, 14, 21, and 28 days, as suggested in the standard method [17, 22]. A TSZ-6 strain-controlled triaxial instrument (Figure 4) was used to perform the UCS test since the triaxial test is considered a UCS test when the confining pressure remains constant. At the same time, the strain rate was controlled at 0.5 mm/min according to a standard method [22]. The maximum strength and the strength at a strain of 15% determined the unconfined compressive strength. Figure 3(b) shows the specimens after the test. Representative samples were taken for XRD and EDS analysis in order to study the interaction mode of the polymer and the silt. Finally, representative samples were scanned by electron microscope and tested by mercury injection to determine the microscopic mechanism of solidification of silty soil by polyurethane (Figure 5).
3. Results and Discussion

3.1. Analysis of the UCS Test Results. To reduce measurement errors, three specimens for each set of parameters were tested in triplicate, and the specimen with a larger error was removed while the average value was reported. The UCS of the pure soil specimens with different moisture contents was also measured for direct comparison. Therefore, the following analysis uses a plain soil strength of 0.21 MPa as a reference.

3.1.1. Effect of Polymer Content on Strength of the Specimens. Figure 6 shows the effect of polymer content on strength of the solidified soils. The results for specimens with a moisture content of 8% and an immersion time of 28 days are similar with those obtained by Liu et al. [16] and Wei et al. [17] and show that the polymer content has a significant impact on the strength of the solidified soil. The polymer amount is directly positively correlated with the UCS of the cured soil specimen. For example, when the polymer content is 5%, the UCS is 0.47 MPa, but increases to 2.14 MPa when the polymer content is 10%. The strength increases by more than 3.5 times when the polymer doubles. When the polymer content increases from 13% to 15%, the strength increases from 2.74 MPa to 5.09 MPa. Compared with the specimen with a polymer content of 5%, the strength increases by 4.8 times and 9.8 times when the polymer content increases by 1.6 times and 2.0 times, respectively. The strength also increases between 2.24 and 24.24 times compared with the plain soil specimen. This clearly indicates that the permeable polymer solidifies the silt used in this work.

3.1.2. Effect of Immersion Time on Strength of the Specimens. Figure 7 shows the effect of immersion time on the strength of solidified soil specimens with a moisture content of 8%. The figure shows that UCS of solidified soil evolves linearly with the immersion time. Longer immersion times produced a higher UCS. Linear fit gives the linear relationship between the UCS and the immersion time for different polymer contents and indicates that the UCS is positively correlated with the immersion time. Compared with other cement-based soil curing agents, the permeable polymer developed in this study is more advantageous for the reinforcement of soil slopes in hydraulic engineering [23].

3.1.3. Effect of Soil Moisture Content on Strength of the Specimens. Figure 8 shows the effect of moisture content on the strength of solidified soil. For instance, the strength of specimen immersed in water for 28 days with a polymer content of 15% decrease when moisture content increases. When the moisture content increases from 8% to 10%, the strength decreases by about 25%. Similarly, when it increases from 10% to 13%, the strength decreases by about 56%. Finally, when it increases from 13% to 15%, the strength decreases by about 8%. However, for all specimens with a polymer content of 5%, the decrease in strength is much less significant (between 2.2% and 6.3%) when the moisture
content increases. It can be seen that soil with a high water content was not conducive to increasing the strength of the solidified soil [23].

3.2. Analysis of Mineral Composition of Solidified Soil. In order to explore the solidification mechanism of polymer and silt, XRD and EDS were used to analyze the chemical composition of solidified soil. It was investigated whether the strength of silt was increased due to the formation of new substances.

3.2.1. EDS Analysis. Comparing Figures 9(a) and 9(b), it can be seen that the C, N, and O content in the sample after the addition of the polymer had significantly increased, and the content of other elements was slightly reduced. This was because the polyurethane is a polymer compound with a large molecular weight. Both the isocyanate (R-N=C=O) and the polyol (nHO∼OH) have a higher C, N, and O content; these three elements account for the increase in the ratio, and this indicates that the polymer has effectively adhered to the surface of the silt particles [24].

3.2.2. XRD Analysis. As shown in Figure 10, before and after the addition of the polymer, the diffraction spectra of the samples were basically the same, but the diffraction intensity fluctuated slightly; however, no new diffraction peaks appeared. This shows that after the polymer was added to the silt, the mineral composition of the soil did not change after solidification, and no new phase substances were formed. From this, it could be judged that only a physical interaction between the silt and the polymer occurred [23].

Based on the results of EDS and XRD, the polymer did not react with the silt, and it was effectively attached to the surface of soil particles.

3.3. SEM Analysis. In order to further explore the solidification mechanism induced by polymer in silty soil, a scanning electron microscope (SEM) was used to image the soil samples.

3.3.1. Effect of Moisture Content on Strength of the Specimens. A series of samples with a polymer content of 15% were taken as representative examples to explore the effect of the water content of the silt on the strength of the solidified soil. Figure 11 shows the SEM images of the specimens with a polymer content of 15% and a moisture content of 8%, 10%, 13%, and 15%, at a 100 μm scale. The sheet-like film part appearing in the images is the polymer. The specimen with a moisture content of 8% has very smaller pores between the soil particles and the polymer fills almost all the voids between them. The polymer has a strong adhesion and forms a
dense film impermeable to water. The specimen with a moisture content of 15% has a loose soil particle distribution and the soil particles are not well bonded by the polymer. This is mainly because the soil specimens with a lower moisture content have less free water in the voids between the soil particles, and the polymer easily enters the pores to achieve a good filling. In the soil specimens with a high moisture content, only a small number of polymer molecules displace the free water to fill the pores and the excess of free water reduces the ability of the polymer to “wrap” soil particles, which reduces the adhesion to soil particles [25]. The presence of water affects the ability of polymer to fill voids within soil particles, hinders its adhesion performance, and prevents interconnections between soil particles. These negative effects are more pronounced when the moisture content increases.

3.3.2. Effect of Polymer Content on Strength of the Specimens. A series of samples with a water content of 8% were taken as representative to explore the effect of the polymer content on strength of solidified soil. Figure 12 shows SEM images of the specimens with a moisture content of 8% and a polymer content of 5%, 10%, 13%, and 15%, with a 100 μm scale. The angular objects are soil particles. The specimen with a polymer content of 5% has a more dispersed distribution of soil particles that are not connected into large sheets and are separated by large voids. The soil particles in the specimen with a polymer content of 15% are well connected and almost form large sheets. The polymer is able to fill the voids between them. In soils with the same moisture content, a higher polymer content results in a stronger connection between the soil particles, which improves the compressive strength of the soil. This result is consistent with the UCS tests.

3.3.3. Microscopic Mechanism of Silt Solidification Induced by the Polymer. Our results allow us to propose a mechanism for the silt solidification by the polymer. Figure 13(a) shows the SEM image of the specimen with a moisture content of 8% but without any polymer. The soil particles in the specimen are loosely distributed, the voids are relatively large, and the soil particles compress and deform under the action of external forces. Figures 13(b) and 13(c) are the SEM images of the specimens with polymer. Since the
polymer has a high "permeability," it fully penetrates into the voids between the soil particles, fills them, and "wraps" them. In short, the solidification by the polymer has two roles in the soil. Figure 13(b) shows that the polymer "covers" and "wraps" the soil particles to interconnect the particles and form sheets with a large area. The soil particles stick together to form a water-impermeable and air-impermeable "film." The presence of the film reduces the erosion of the strength of the specimen by water. The second role is illustrated in Figures 13(c) and 13(d): the polymer fills the voids between adjacent soil particles, forming a "bridge connection," making them stick with each other and thereby increase the density of the soil. Both actions by the polymer increased the ability of the soil particles to connect with each other; therefore, the permeability of the soil decreased; water affects the strength of the soil in a much reduced manner, and the compactness of the soil increases. As a result, the strength of the soil is significantly improved [17, 26].

3.4. Analysis by MIP. The SEM results show that the polymer has a filling effect between the soil particles. To study the void characteristics of the polymer-solidified soil, MIP was carried out on the specimens with a moisture content of 8% and different polymer contents. Figure 14 shows the corresponding pore distribution, and Table 2 lists the other physical characteristics of the specimens obtained by MIP. When the polymer content increases, the intermediate pore diameter, the pore diameter, and the total pore volume all gradually decrease. A higher polymer content densifies the soil. The decrease in the permeability also supports this conclusion [27]. The packing density in Table 2 shows that the specimen with a polymer content of 15% piles up more easily and becomes denser. The decrease of the porosity and the significant decrease of the permeability also show that the soil becomes denser when the polymer content increases. The results show that the solidification of the soil by the polymer effectively reduces the permeability of the soil and significantly increases its strength [28].

4. Conclusions

The purpose of this study was to illustrate the effect of the addition of polyurethane to soil. Based on the test results, polyurethane grouting material can be used as a soil curing agent, and it has been shown to have a significant curing effect on the silt.

The results showed that the addition of a higher amount of the polymer increased the strength of the solidified soil. Additionally, the strength of the solidified soil increased with the increase of the immersion time. However, a higher water content was not conducive to improving the strength of the solidified soil. Based on the result of the UCS and MIP, silt with an 8% water content mixed with 15% polymer can be regarded as the optimum concentration for yielding effective stabilization.
According to the results from EDS and XRD, the principle action of the polyurethane-cured silt was not that it reacts chemically with silt, but that it improved the soil through physical actions. It can be seen that the polyurethane mainly improved the strength of the silt through adhesion, wrapping, filling, and bridging.

Data Availability

The data used to support the findings of this study are available from the corresponding author upon request.

Conflicts of Interest

The authors declare that they have no conflicts of interest regarding the publication of this paper.

Acknowledgments

This work was supported by the National Key R&D Program of China (no. 2017YFC0405002) and the National Natural Science Foundation of China (NSFC) (no. 51679219).

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