Research Article

Interface Bonding Properties between Nonwater Reaction Polyurethane Polymer Materials and Concrete

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The concentric pushing method was used to study the bonding properties between polymer and concrete. This paper studied the influence of polymer density, environmental temperature, and moisture content of concrete between polymer and concrete on the bond strength. The results indicated that the bond failure of specimens occurred mainly when the polymer was pushed out. Furthermore, increasing the polymer density increases the bond strength at the polymer-concrete interface but decreases as the moisture content of the concrete increases. The environmental temperature affects the curing time, and the bond strength increases with increasing temperature. Under the same condition, the bond strength was influenced by the roughness of the interface. This study provides references for the construction design and enhances polymer materials and matrix application for repairing cracks in concrete dams.

1. Introduction

Hydraulic structures are prone to cracks and concentrated leak channels during construction or operation. Many dam damage cases are caused by crack penetration. When repairing cracks in concrete dams, surface sealing, insertion, and grouting methods are generally used [1]. The grouting method is the best method to solve such problems. The grout used can be divided into cement grouting and chemical grouting [2–4]. The cement grouting method relies on the pumping pressure of the grouting equipment to fill the crack and act as the plugging slurry. However, the cement material has a long setting time, which is unsuitable for emergency or engineering rescue. After solidification, the cement slurry converts to a brittle material, resulting in repeated cracking [5–7]. The chemical grouting method relies on the pressure of the equipment to pump the slurry to the cracks and fill the cracks through expansion force [8]. Epoxy resin and polyurethane are generally used in polymer chemical grouting materials [9, 10]. Epoxy resin materials are similar to cement materials. After hardening, they are brittle and cannot withstand secondary cracking [11, 12]. In contrast, the elasticity of the polyurethane slurry in a specific density range is superior to the elasticity of the dam body and has a good sealing effect [13].

With the increasing number of large dams and reservoirs, the commonly used methods of emptying reservoirs are too expensive for deep diseases. Underwater repair is a more feasible method. However, the underwater environment requires a higher standard for repairing materials. Cracks in the dam body in a deep water environment can be repaired using submersibles. This means that the performance of repairing materials faces the adverse conditions of low temperature and wet interface [14–16]. Many
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2. Experiment

2.1. Materials and Model Design. The polymer was a two-component polymer that reacted and foamed. Component A consisted of isocyanates and polyisocyanates, and component B consisted of polyols and additives. Materials A and B were mixed and injected into steel models by a grouting gun. For polymer materials with densities of 0.1, 0.12, 0.2, 0.29, 0.4, 0.6, 0.8, 0.83, and 1.2 g/cm³, the polymer density was controlled by mass. The C60 concrete was used in the test. The test pieces of this experiment were based on a standard permeability test piece (upper bottom, 185 mm, lower bottom, 175 mm, and height, 150 mm) [29]. The center was preset with a hollow cylinder with a diameter of 90 mm filled with polymer (as shown in Figure 1(a)).

The polymer generates high temperatures and large expansion forces during the reaction [30]. Therefore, the specimens were placed in a special steel cylinder, and the top and bottom were connected using a flange plate and fixed with bolts. As shown in Figure 1(b), the position of the grouting hole was located at the center of the flange, and the venting hole was located at a short distance from the side of the grouting hole.

2.2. Specimen. In this test, 120 specimens were created to examine the bond strength between the polymer and concrete, mainly considering three parameters: concrete interface temperature, moisture content, and polymer density (as shown in Table 1). Hollow cylindrical specimens were used for the test. As shown in Figure 2, the concrete adopted a standard impermeability test model (upper, 185 mm, lower, 175 mm, and height, 150 mm). First, the center was cut, and a PVC pipe with a diameter of 90 mm was inserted to reserve a hole for the polymer. The prepared concrete was then poured and removed from the mold, and, 28 days after standard curing, the specimens were soaked in groups. The time control was divided into 0 h, 3 h, 10 h, and 24 h. The soaked specimens were wrapped in plastic wrap and placed in a constant temperature test room (0°C–15°C) for 24 h. After applying butter, they were placed in a steel cylinder. They were then covered with upper and lower flanges, bolts were tightened, and the grouting density was changed by controlling the grouting mass. As a result, the polymer expanded rapidly within the model, discharging air through the overflow hole. After grouting for 1 hour, all samples were unmolded and placed in the laboratory until testing.

After prolonged immersion, cracks in the concrete may be fully saturated. In this experiment, specimens were made using C60 grade high-strength concrete. The moisture content was measured as shown in Figure 3. From the figure, it can be seen that the moisture content of the specimens increased sharply in the early stage after soaking in water, but the growth rate was slow after 24 h. Therefore, the moisture content corresponding to the immersion time of 24 hours can be regarded as the saturated moisture content (about 2.66%).

2.3. Experiment Method. Due to the short hole in the center of specimens, it was difficult to anchor the low-density specimens. The main test was the bonding properties of the interface between polyurethane polymer and concrete. Therefore, the concentric pull-out test was not adopted, and the bond strength was determined by the concentric pushing test shown in Figure 4 [31]. The specimens were placed on a test bench with a special steel ring substrate. The top of the pusher was made of an 85 mm diameter disc, the center of which was aligned with the sample. The universal testing machine was equipped with a strain acquisition system that collected continuous displacement. The maximum load of the universal test machine was 100 kN, the load speed was 0.1 kN/s, and changes in load and displacement could be recorded automatically. Before official loading, we considered a preload of 1 kN [32]. Loading was stopped when the following happened: (1) the central hole of the concrete specimen was destroyed; (2) the polyurethane polymer core was cracked and damaged; and (3) the polyurethane polymer core did not slip along the concrete interface. Since the maximum load \( P \) was measured using a load cell with an accuracy of 0.01 kN, the average bond stress \( T \) can be taken as into account \( T = P/A \). The average bond strengths of the three groups of parallel specimens were taken as measurement values for those experimental specimens.
The specimens were classified into three groups to examine the effects of polymer density, prewetting of the concrete surface before grouting, and temperature after grouting on bond behavior. The impact of interfacial moisture on bond strength under dry and saturated conditions was evaluated in the first group. The temperature was 10°C, and the polymer density was 0.6 g/cm³. In the second group, specimens with temperatures of 0°C, 5°C, 10°C, and 15°C were tested to examine the effects of temperature. In the third group, samples with various polymer density were studied.
densities of 0.1, 0.12, 0.2, 0.29, 0.4, 0.6, 0.8, 0.83, and 1.2 g/cm³ were prepared to evaluate the effects of polymer densities.

3. Results and Discussion

Most of the 120 specimens in this test experienced slip damage at the polymer-concrete interface (as shown in Figure 5(a)). In addition, when a part of the annular concrete of the specimen was pushed out, it was partially damaged (as shown in Figure 5(b)). Some high-density specimens are damaged by the hardened surface of polyurethane, but this damage did not penetrate the polymer, and the polymer core was intact. It was found that the interface of the polymer was worn, and the uneven part of the surface was polished. The inner interface of the concrete was also worn, the concrete at the inner interface of the individual specimens was sheared, and the concrete skin was stuck off.

3.1. Average Compression. As shown in Figure 6, the compression value at the time of failure of the polymer bond in the center of specimens was recorded to determine the relationship between the compression value and the density. The results show that the greater the push-out forces for low-density specimens, the greater the compression deformation. The reason is that when the polymer density is low, many bubbles exist in the polymer, the compactness is low, and the compression deformation is large. The deformation of low-density specimens is greatly affected by the push-out force. The push-out force is proportional to the compression value for high-density specimens (1.2 g/cm³). As a result, the slope of the density-compression curve is less than the slope of low-density specimens. In high-density polymers, the microscopic morphology can be interpreted as the formation of isolated high-density bubbles within the polymer. By decreasing the size of the bubbles, the wall thickness of the bubbles increases. In addition, the delicate and uniform cell structure enhances the resistance of the material to deformation (as shown in Figure 7(b)).

3.2. Polymer-Concrete Bond-Slip Characteristics. As shown in Figure 8, the slip of the load end and that of the free end are basically the same, but the free end displacement is slightly delayed. The free end load-displacement curve intersects the coordinate axis. This indicates that the load end slightly slips in the early stages of loading, and no failure is transmitted from the bond interface to the free end. Thus, there is no slip on the free end, and external loads are resisted mainly by chemical cementation at this stage. According to bond-slip characteristics, the entire development can be simply divided into three stages.
3.2.1. Linear Slip Stage (OA). In the OA stage, the load-displacement curve is approximately linear. The thrust is mainly resisted by friction and mechanical bite force at this stage, and the chemical bonding force no longer works. As a result, the displacement increases faster, and the displacement of the free end is slightly smaller than that of the load end.

3.2.2. Ejection Stage. As the load continues to increase, the concrete on the bonding interface is worn out, the amount of slip is further accelerated, the load-displacement curve tends to be flat, and the load eventually reaches its peak.

3.2.3. Thrust Decay Stage. As the amount of slip increases, the concrete surface that comes into contact with the polymer wears severely, reducing the friction and mechanical bite force and the corresponding thrust.

3.3. Bond Strength. This test uses the bond strength between polymer and concrete under normal temperature (25°C) and dry conditions as a reference. Table 2 shows the trend of bond strength as a function of density.

3.3.1. Effect of Moisture Content. Cracks at certain engineering sites in dams are often moist because the water level constantly changes on rainy days and under the influence of open sluice gates [20]. Polyurethane materials are sensitive to the moisture level at the repair interface, which in turn affects the bond strength between the polyurethane polymer and the concrete on either side of the crack [33]. Specimens in Table 3 have the same density (0.6 g/cm³) and interface temperature (0°C). Four different wetting conditions were set, and soaking time was used as a screening condition for moisture levels. Using the moisture content as a quantitative index, the bond strength was measured as follows. It can be seen that the bond strength decreases with the increase of moisture levels (water content) under the same temperature condition, because the water reacted with isocyanate to form carbon dioxide to reduce the adhesion if the isocyanate in the two-component polyurethane material was less than 2% in the aqueous state. The reason may be that when the amount of isocyanate is small, the foam curing is not complete, and the structure of the cross-linking network is partly defective. At a temperature of 0°C and a polymer density of 0.6 g/cm³, the bond strength of the specimens at the soaking times of 3 h and 24 h decreased by 75.2% and 84.3%, respectively. Therefore, it is highly required to reduce the interfacial moisture between concrete and polymers in engineering applications. For example, a superplasticizer can be added to concrete to prevent a significant decrease in bond strength due to moisture [34].

3.3.2. Effect of Temperature. Figure 9 shows that, under the same conditions, the higher the temperature within a certain temperature range, the higher the bond strength. However,
under saturation conditions, the bond strength is reduced accordingly (as shown in Table 4). As can be seen from Table 5, the bond strength, corresponding to the temperature drop, decreased by 43.8%–57.1% in the dry state. FYhe lower the temperature, the more significant the decrease. In addition, as the temperature drops, the bond strength decreases from 65.2% to 83%. FYhis is because the higher the temperature of the specimen, the slower the temperature loss of the polymer during the reaction and the lower the temperature shrinkage stress. On the other hand, at the concrete interface, the higher the temperature of the specimens, the denser the compaction. FYherefore, differences between the reaction temperature of the low temperature specimens and the material in the reaction process are large, and temperature shrinkage stress is generated. After the reaction, the extrusion and cementation forces are offset by the temperature shrinkage stress, the concrete is disengaged from the concrete, and the bond force is reduced.

3.4. Effect of Polymer Density. To produce polymer water stop plugging, the effect of density on bond strength at low temperatures and under fully saturated interfacial conditions needs to be further investigated [35]. FYhe interface temperature of the specimens is 5°C, and it is fully saturated (24 h). The effect of polymer density on bond strength can be analyzed and compared to dry conditions. Figure 10 shows the relationship between bond strength and polymer density. Based on the linear fitting results, the bond strength between the polymer and the concrete interface is positively correlated with the polymer. The higher the density of the polymer, the higher the bond strength between the polymer and the concrete interface.

| Density ρ/(g/cm³) | 0.1 | 0.2 | 0.4 | 0.6 | 0.8 | 1.2 |
|-------------------|-----|-----|-----|-----|-----|-----|
| Bond strength (MPa) | 0.31 | 0.53 | 0.6 | 1.12 | 1.55 | 1.98 |

The absolute difference in the bond strength between the saturated and dry specimens between 0.1 g/cm³ and 0.4 g/cm³ is not significant, with a mean reduction of about 3.8%. However, the absolute difference in density between 0.4 g/cm³ and 1.5 g/cm³ is 1.0 and it is significantly larger for 1.2 g/cm³. The reduction rate is large, ranging from 67.2% to 72.7%. The difference between dense specimens (1.2 g/cm³) is much more remarkable. The bond strength between fully saturated low-density specimens and high-density specimens is relatively concentrated, indicating that an increase in polymer density in the water has a limited effect on the growth of bond strength. There are two reasons for this phenomenon. As shown in Figure 7, when the polymer density is relatively low (Figure 7(a)), there are many bubble cells in the polymer, the radius of the bubbles becomes large, and the bubbles are squeezed together. On the other hand, the polymer is combined with the concrete through the thin walls of the bubble chamber, reducing the bond strength. The higher the polymer density (Figure 7(b)), the smaller the radius of the bubble and the thicker the walls. This increases the distance between the bubble chamber and the concrete. Therefore, at the polymer-concrete interface, the thick walls of the bubbles cells bond with the concrete, increasing the

![Figure 7: SEM images of the polymer at densities of (a) 0.15 g/cm³ and (b) 0.57 g/cm³.](image)

![Figure 8: Bond-slip curves of the loading end and the free end.](image)

![Table 2: Relationship between bond strength and density.](image)
bond strength. In addition, the expansion force of the polymer increases with density. As a result, the intercalation and cementation of the polymer at the interface become stronger, resulting in higher bond strength [28]. The higher the density of the polymer, the higher the bond strength at the interface in a certain range. This means that the contact

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Table 3: Relationship between moisture content and bond strength.

| Number | Temperature (°C) | Moisture content (%) | Ave. moisture content (%) | Soaking time (h) | Bond strength (MPa) | Ave. bond strength (MPa) |
|--------|------------------|----------------------|---------------------------|------------------|---------------------|-------------------------|
| 1–3    | 0                | 0                    | 0                         | 0                | 0.42                | 0.48                    |
| 13–15  | 1.03             | 1.11                 | 0.95                      | 0.83             | 0.44                | 0.09                    |
| 25–27  | 0.47             | 1.08                 | 0.93                      | 1.3              | 0.64                | 1.12                    |
| 37–39  | 1.19             | 1.79                 |                           |                  | 0.2                 | 0.11                    |

Figure 9: Relationship between temperature and bond strength.

Table 4: Saturation conditions.

| Number | Density ρ/(g/cm³) | Soaking time (h) | Temperature (°C) | Ave. moisture content (%) | Bond strength (MPa) | Reduction rate (%) |
|--------|-------------------|------------------|------------------|---------------------------|---------------------|-------------------|
| 37–39  | 0.6               | 24               | 0                | 1.3                       | 0.19                | 83.0              |
| 40–42  | 0.6               | 24               | 5                | 1.22                      | 0.33                | 70.5              |
| 43–45  | 0.6               | 24               | 10               | 1.03                      | 0.33                | 70.5              |
| 46–48  | 0.6               | 24               | 15               | 0.76                      | 0.39                | 65.2              |

Table 5: Dry condition.

| Number | Density ρ/(g/cm³) | Soaking time (h) | Temperature (°C) | Ave. moisture content (%) | Bond strength (MPa) | Reduction rate (%) |
|--------|-------------------|------------------|------------------|---------------------------|---------------------|-------------------|
| 1–3    | 0.6               | 0                | 0                | —                         | 0.48                | 57.1              |
| 4–6    | 0.6               | 0                | 5                | —                         | 0.49                | 56.3              |
| 7–9    | 0.6               | 0                | 10               | —                         | 0.57                | 49.1              |
| 10–12  | 0.6               | 0                | 15               | —                         | 0.63                | 43.8              |
surface can withstand higher shear forces and pressures. However, the increase in polymer density has a limited effect on the growth of bond strength at the saturated interface. During the reaction, many bubbles are generated on the surface of the concrete, and the bond strength is significantly reduced due to intrusion. If the specimen polymer density exceeds a specific range, most of it will damage the concrete [36].

4. Conclusion

Through a concentric pushing test system for 120 adhesive specimens, the effects of temperature, density, and interfacial moisture on the bonding properties between polymers and concrete under special construction conditions were studied. From the results obtained, the following conclusions can be drawn:

1. When fully saturated, increasing the polymer density has little effect on bond strength. In contrast, increased polymer density has a significant effect on bond strength under dry conditions.

2. Interfacial moisture content is a significant factor affecting the bonding properties between the polymer and concrete. At the same interfacial temperature, the higher the interfacial moisture content, the
lower the bond strength between the polymer and concrete. When the interface is completely saturated, the interfacial bond strength follows this rule, but the overall bond strength is lower than that of a dry interface. For dam concrete structures, reducing interfacial water content and increasing the mass of the polymer are effective methods to improve the bonding force between the polymer and concrete.

(3) The bond strength between the polymer interface and the concrete interface is much less than the shear resistance of the polymer itself. As a result, most of the polymer cores are intact and are mainly damaged by ejecting, with less damage to the inner surface of the concrete.

**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.

**Authors’ Contributions**

Mingsheng Shi conceptualized the study. Han Tian performed data curation. Xijun Zhang performed formal analysis. Mingsheng Shi was responsible for funding acquisition. Xijun Zhang developed the methodology. Xijun Zhang wrote the original draft. Chaojie Wang reviewed and edited the article.

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