Strength, Stiffness, and Microstructure of Wood Ash Stabilized Marine Clay

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Abstract: The world’s population is growing at a rapid pace, thus increasing the need for shelter, which, because of increased carbon emissions, is making our planet less inhabitable. Thus, supplementary cementitious materials (SCMs) are used to reduce the embodied carbon emissions in the building sector. Wood ash, as a replacement for cement in soil treatment, seems to be a promising material. In this study, we considered the strength, stiffness, and microstructural behavior of marine clay treated with cement and wood ash as a cement replacement. Since clay is abundant in nature, it could help stabilize waste to improve the mechanical behavior of produced composites. Portland cement (7%, 10%, and 13%) was replaced with various amount of wood ash (5% and 10%) with two different dry densities (1400 and 1600 kg/m3) and three distinct curing periods (7, 28, and 60 days). Unconfined compressive strength, direct shear, porosity, pulse velocity, x-ray diffraction, and scanning electron microscopy tests were performed on selected specimens to evaluate the structural and microstructural effect of clay–wood ash–cement interaction. The results revealed that the replacement of cement with 5% of wood ash yielded superior performance. The microstructure investigation of wood ash–cement–clay blends further showed the formation of a densified matrix with stable bonds. Furthermore, the porosity and strength properties of blends developed unique relationships, which were further confirmed by other supplementary materials and soils.

Keywords: strength; stiffness; clay; stabilization; microstructure; wood ash; waste

1. Introduction

Not long ago, our ancestors were hunters and gatherers. They settled on fertile lands filled with animals to hunt. As soon as the soil became infertile and food sources grew extinct, these hunters and gatherers sought new homes. This process continued until they reached the fertile lands of Mesopotamia between the Tigris and Euphrates rivers, which extend from Eastern Anatolia to the Persian Gulf. These lands were so fertile that people did not need to seek new homes; thus, they settled there and built homes. Upon examining early settlements, settling next to fertile lands required the construction of homes on the strong ground using soft alluvium soils rich with minerals needed for farming. Since that time, the increase in population has created demands for speedy construction. According to the Cyprus Chamber of Civil Engineers [1](Site Investigation Database), most of this increase in construction activities has taken place in areas close to the sea where marine clays are present. It is known that marine clays are problematic for construction because they lack strength and compressibility, which leads to excessive settlements that have low bearing capacities [2–8].
Soil stabilization techniques have been developed to improve the properties of the poor-quality soil so that inadequate soil would be able to serve the current needs. One of the most widely used stabilization techniques is mixing soil with several binding agents to enhance the mechanical properties of inadequate soils. Soil-cement and soil-lime stabilization have been extensively studied. However, in addition to the soil stabilization approach, it became necessary to consider sustainability to decrease the carbon footprint and environmental pollution. In that sense, the utilization of waste products, such as wood-ash within soil stabilization as a partial replacement for cement, could be an effective approach. It is known that cement production is one of the main sources of CO₂ emission worldwide [9].

Consequently, using wood-ash as partial replacement to cement will decrease the cement used in soil stabilization and decrease the carbon emission caused by cement production. Additionally, it will be beneficial in terms of waste management since a waste product can be used in soil stabilization [9]. In today’s world, climate change and global warming are forcing world leaders to find more effective sustainable solutions [9,10]. Concrete is used in the construction of various structures necessary to accommodate the rapidly increasing world population. The statistics show that approximately 1 cubic meter of concrete is used per person [11]. Approximately 1 ton of carbon dioxide gas is released into the atmosphere during the production of 1 ton of Ordinary Portland cement, which affects the strength of the concrete components without SCMs [12]. Additionally, the global average of carbon release is established as 0.86, according to Miller et al. study [13].

In line with these developments, experts are seeking alternative ways to reduce carbon gas to address the issue of sustainability worldwide [14–19]. Cement companies around the world have been widely using SCMs, especially in the last 25 years. Scrivener et al. [20] offers more detailed projection of the construction sector for better sustainability and reviews ongoing studies on SCMs. Additionally, given the unavailability of the traditional industrial by-product of SCMs in certain regions, the limited global supply and its effects on sustainability strategies are highlighted in Scrivener et al.’s study [20]. To reduce the amount of clinker used in cement production, many universities and research organizations are using SCMs, e.g., fly ash, bottom ash, rice husk ash, marble powder, and wood ash (WA), and evaluating different alternative wastes [11,21–26]. Fly ash, bottom ash, rice husk ash, marble powder, and wood ash (WA) are commonly used SCM. The positive effect of these wastes on both fresh and hardened concrete properties benefits the industry. In addition to improving the concrete properties, SCM should make a positive contribution to the soil properties. Therefore, simply improving concrete properties is insufficient; rather, the soil properties must be improved [14,22]. SCMs promote gel formation, provide denser structure, and reduce the amount of ettringite formation in the paste. A homogeneous interstitial transition zone can be achieved with the use of SCMs [27–29]. Wood ash incorporation, as a partial replacement, generally reduces the porosity of the concrete by having a filler effect. However, a high amount of carbon makes it unsuitable for structural grade concrete applications [23,30]. The sum of alumina, silica, and iron oxides falls below the limit values specified in ASTM C618 [18,31,32]. In addition, the loss in ignition value is far above the limit value set in the standards. However, previous research has demonstrated that these values do not have a negative effect on concrete properties [9,18,30]. WA is a light and porous material. Its quality largely depends on the species of trees, burning temperature, and moisture content available during cutting operations [18]. The factor governing the composite’s behavior is related to the calcium-to-silica (Ca/Si) ratio. As this ratio increases, the expansion also increases. The growth of cracks is minimized as the Ca/Si ratio is reduced [28,33]. Based on previous findings, a high amount of silica reduces the pH of the system by reducing the Ca/Si ratio while providing a higher density when the ratio is high. Furthermore, the addition of WA to the concrete increases the water demand to ensure internal curing and helps the formation of better bonds in later stages [33,34]. The filler effect and poor pozzolanic activity of WA improve the strength beyond 90 days. The silica/alumina (Si/Al) ratio also plays an important role in the development of a strong bond by increasing the stability of the structure [10,28,33].

Microscopic investigations showed that WA contains irregular and angular particles. Some research has demonstrated that WA incorporated composites comprising flaky and elongated shape particles.
particles with porous carbon spheres. Carbon spheres contain micro and macropores and are distributed evenly[10,18,34]. Many studies have shown that portlandite and ettringite needles are also present. Crystalline phases are mostly dominated by calcium particles and show pozzolanic activity, which helps the densification and stabilizes a less crack-free microstructure. Tobermorite gel in WA can be considered a semicrystalline, while calcium silicate hydrate gel can be considered amorphous. Mullite crystals affect the durability properties of the composites [23,27,28,35]. The incorporation of WA increases water demand and creates internal pressure inside the matrix. If this pressure continues to increase, it deteriorates the composites’ structure. Naik and Kraus [19] investigated the physical and chemical properties together with the microstructure of WA, incorporating cement-based materials. In their study, various WA samples were collected from the United States and Canada. The authors found that the source of WA significantly affected the behavior of the composites. They suggested using WA to control low strength and structural grade applications. Siddique et al. [34] evaluated the WA as a replacement for fine aggregate using X-ray diffraction. They reported that the WA contained crystalline and amorphous phases. Furthermore, they found that 5% of WA is optimal in terms of better matrix properties. Due to WA’s higher absorption capacity, the final composites are adversely affected above this substitution level. Shearer and Kurtis [17] evaluated the possibility of using coal ash and biomass ash as a source of SCM. They reported that the increase in aluminum and silica concentration in pore solution most probably occurred after 7 days of curing, thereby gaining further strength in later ages. They also concluded that ashes have a positive effect on durability. Stolz et al. [23] characterized WA in foam-based composites. They reported that up to a 20% strength loss was recorded when WA was incorporated into a system as a 10% cement substitution. The elastic modulus reduction was approximately 35% at the 20% substitution level. However, they mentioned advantages of using WA in projects where acoustical and sound permeability is important. Elinwa [29] examined the effects of calcined clay on mortar and concrete properties. Samples composed of clay varied from 0-40% by mass of cement. As the clay content increased, the strength continuously decreased. The results of the study indicated that 10% calcined clay was found to be optimal in terms of better strength and durability.

Consoli et al. [36] obtained correlations between water-cement and porosity-cement ratios to evaluate the unconfined compressive strength of soils. The authors reported a significant increase in the strength of soil by the addition of cement only. The effect of cement content increased linearly with strength, whereas it showed an exponential relationship with the reduction in porosity. In other words, as the soil-cement products fill the voids of the soil, the strength increases consequently. Consoli et al. [36] proposed a correlation to evaluate the unconfined compressive strength of cement stabilized soils. According to the authors, the ratio between voids and cement, in other words, $\eta/C_v^{0.28}$ where $\eta$ is porosity and $C_v$ is volumetric cement content, best represents the UCS by some empirical factors. Another research by Consoli et al. [37] claimed that the unconfined compressive strength increases when the lime content and density are constant, and the amount of fly ash increases. In line with the previous study, the reduction in porosity revealed an exponential relationship with strength, whereas the amount of fly ash showed a linear trend. Ekinci et al. [22] concluded that mixing marine clay with Portland cement and copper slag had negative effects on strength properties; however, the inclusion of lime activated the pozzolanic effects, and an improvement in strength properties was observed. Some researchers studied the shear strength parameters of wood ash that included fine-grained soil by examining Mohr-Coulomb shear envelope and concluded that improvement occurs both in cohesion and friction angle until an optimum value of 10% addition of wood ash. However, above that value, i.e., at 12.5% addition, a decrease in shear parameters was observed [38]. Consoli et al. [39] established a reliable relationship between the unconfined compressive strength, splitting tensile strength and cohesion (C), and friction angle ($\phi$) in case of lime addition to soils. More recently, Soltani et al. [51] established a universal predictive model for the UCS of artificially cemented fine-grained soils.

WA alone does not possess pozzolanic activity, but due to the presence of some silica and alumina, it shows better engineering properties. Generally, clay particles contain enough silica and some alumina. The interaction of clayey soils with WA seems promising for future trends. To date,
no research has investigated the effects of WA–clay–cement blends in terms of strength, stiffness, and microstructure. This study investigated the effect of the partial replacement of cement with wood ash in marine clay stabilization by considering unconfined compressive strength, initial shear modulus, and shear strength parameters. Scanning electron microscopy imaging was used to analyze the microstructure and highlight the observed behavior of the blends. In addition, a parametric model was created to determine the amount of cement needed to establish the value of the studied parameters (UCS, shear strength, and initial shear modulus) for the optimal amount of wood ash. Finally, an equation was proposed to evaluate the necessary amount of cement to reach a target value for the studied parameters (i.e., unconfined compressive strength, shear strength, and initial shear modulus) to attain the optimal amount of wood ash.

2. Materials and Methods

The physical and chemical properties of marine clay and wood ash were evaluated. Unconfined compression tests were performed along with direct shear and ultrasonic pulse velocity tests. Portland cement has been replaced with various amounts of wood ash of two different dry densities and three distinct curing periods. Additionally, scanning electron microscopy was performed on selected specimens to further analyze the microstructure.

2.1. Materials

2.1.1. Clay

The soil was collected from the northern coast of Cyprus. Characterization tests, such as Atterberg limits, sieve analysis, and specific gravity, were conducted according to ASTM to determine the physical properties as well as the consistency of this soil. The grain size distribution of clay is shown in Figure 1 (note that it is well-graded with a \( D_{50} \) particle diameter of 0.0035 mm). Table 1 shows the physical characteristics of the clay. In addition, this clay is categorized according to USCS (ASTM, 2017b [40]) as an inorganic clay with low to medium plasticity (CL). The specific gravity of wood ash was determined in accordance with ASTM D854 – 14 and found to be 2,61. X-ray diffraction analysis was done with X’Pert Pro MPD diffractometer using monochromatic Cu-Ka radiation. The XRF analysis of this clay showed that it composes high amounts of silica, calcium, and alumina. Based on the XRF analysis of clay samples (Table 2), the calcium was the main compound by mass. The XRD test results of clay identified Illite as the peak at 9 and possibly pyrite as peak 35. The other peaks are illustrated in Figure 2, and EDX of pure clay is shown in Figure 3.
Figure 1. The grain size distribution of studied Clay and Wood Ash.
### Table 1. Physical properties of marine-deposited clay, wood ash, London clay and Portugal silty sand.

| Properties                          | Marine Clay | Wood Ash | London Clay | Portugal Silty Sand |
|-------------------------------------|-------------|----------|-------------|---------------------|
| Liquid limit (%)                    | 40          | -        | 78          | 39                  |
| Plastic limit (%)                   | 21          | -        | 30          | 34                  |
| Plasticity index (%)                | 19          | -        | 48          | 5                   |
| Specific gravity                    | 2.61        | 1.7      | 2.75        | 2.64                |
| Fine gravel (4.75 mm < diameter < 20 mm) (%) | -          | -        | -           | -                   |
| Coarse sand (2.00 mm < diameter < 4.75mm) (%) | 2          | -        | -           | -                   |
| Medium sand (0.425 < diameter < 2.00 mm) (%) | 3          | -        | -           | -                   |
| Fine sand (0.075 mm < diameter < 0.425 mm) (%) | 27         | 100      | 2           | 1.5                 |
| Silt (0.002 mm < diameter < 0.075 mm) (%) | 19         | -        | 48          | 65,5                |
| Clay (diameter < 0.002 mm) (%)      | 49          | -        | 50          | 33                  |
| D₅₀ particle diameter (mm)           | 0.0035      | 0.18     | 0.002       | 0.006               |
| USCS class                          | CL          | SP       | CH          | ML                  |

### Figure 2. X-ray diffraction of studied clay specimen.
2.1.2 Wood Ash (WA)

The WA used in this study was obtained from the remnants of pruned and burned olive trees. It contained particles with a mean size of $D_{50} = 0.18$ mm (Table 1). The particle size distribution is shown in Figure 1, along with that of the clay. The chemical analysis (standard) results are shown in Table 2. This wood ash consists mainly of calcite and silica. Similar findings were confirmed on the EDX analysis shown in Figure 4. The specific gravity of wood ash was determined in accordance with ASTM C188 – 17 in a standard flask used to determine the specific gravity of cement. The specific gravity can vary depending on the source and the heat of combustion, as explained by Naik et al. [41]. Therefore, specific gravity tests were performed on every lot of provided wood ash, and an average value of 1.7 was taken into consideration (variation of $G_s < 0.02$).

**Table 2.** Chemical analysis of the Portland cement and wood ash.

| Compound | Clay (%) | Portland Cement (%) | Wood Ash (%) |
|----------|----------|---------------------|--------------|
| SiO$_2$  | 22.1     | 21.2                | 18.1         |
| Al$_2$O$_3$ | 7.6      | 5.1                 | 3.3          |
| Fe$_2$O$_3$ | 6.7      | 2.5                 | 2.8          |
| CaO      | 55.5     | 64.7                | 44.4         |
| MgO      | 2.3      | 0.9                 | 2.8          |
| K$_2$O   | 2.1      | 0.2                 | 5.2          |
| SO$_3$   | 1.1      | 1.5                 | 0.9          |
| Loss in ignition | 2.6 | 2.5 | 22.1 |

Figure 3. EDX of pure clay sample
2.1.3. Cement

The cement used in the study was Portland cement (C) Type I, as specified in ASTM (2018). The Blaine fineness of cement was 289 m²/kg. The chemical analysis of this cement is shown in Table 2. Like wood ash, the main components of the cement are calcium oxide and silica.

Distilled water was used to mix and cure the composites.

2.2. Methods

To assess the mechanical effects of wood ash, i.e., cement mixture on clay, the preparation methods and testing regimes outlined below were adopted. Figure 5 shows setups for the tests carried out.

![Figure 5](image-url)  
**Figure 5** Tests setups; a) Load Frame for UCS, b) Pundit for Ultrasonic Pulse Velocity (UPV), c) Direct shear box for shear strength parameters.
2.2.1. Molding and Curing of Specimens

For the unconfined compression-strength tests, cylindrical specimens were prepared with a 100 mm height and 50 mm diameter to keep a 2:1 ratio specified in the standard ASTM C39 [42]. First, the targeted dry density was specified. In this research, the targeted densities were 1400 and 1600 kg/m^3 based on the maximum dry density of the clay at the optimum moisture content (1600 kg/m^3). As stated by Basma et al. [43], to obtain representative soil properties, the soil should be air-dried rather than oven-dried. Therefore, clay samples obtained from the site were air-dried at room temperature and pulverized with mortar and pestle. Next, the amounts of clay, cement, and wood ash were calculated based on the aforementioned density. These proportions of dry materials were then mixed under dry conditions until uniform distribution was achieved (at least 5 min). After that, the pre-calculated amount of water was slowly added and mixed with the binder until a homogenous mixture was reached. The amount of water to be used was determined from the compaction curve of the clay (see supplementary file). Two densities were chosen, where the first one corresponded to the maximum density (1600 kg/m^3), and the second one corresponded to a lesser density (1400 kg/m^3). This mixture was then separated into three equal layers and compacted statically in a split mold to the desired density using the compaction method proposed by Selig and Ladd [44]. After the sample was formed, it was removed from the mold, and the dimensions measured. The specimens were cured according to ASTM C 511 [45].

The mix of the specimens was based on the relative contents of Portland cement (C) and wood ash (WA). C was defined as the mass of Portland cement divided by the mass of the dry soil, whereas WA was defined as the mass of the wood ash divided by the mass of Portland cement, as it was a partial replacement for cement.

Porosity was calculated using a modified version of Equation 1 proposed by Consoli et al. [46], that considers dry density (ρ_d), and the mass of the marine clay (M_S), Portland cement (M_C), and wood ash (M_WA). The corresponding specific gravities are marine clay (G_S), Portland cement (G_C), and wood ash (G_WA), respectively.

\[ \frac{\rho_d}{\text{total mass of solid}} \left( \frac{M_S}{G_S} + \frac{M_C}{G_C} + \frac{M_WA}{G_WA} \right) = 100 - 100 \]  

(1)

Depending on the porosity cement index (η/C_W), a unique relationship was developed to predict the behavior of cement-treated soils [47], which only accounted for cement. More recently, Ekinci et al. [22] proposed a more general index X_{iv}, which accounts for all binder contents. In this study, Ekinci et al.’s [22] parameter was modified in an attempt to predict the strength of each mixture, where X_{iv} was calculated from the modified Equation 2, where V = M/ρ is true for all materials used:

\[ X_{iv} = \frac{\nu_s + \nu_c + \nu_{WA}}{\nu} \]  

(2)

Table 3 provides all the necessary molding data, including material contents, curing periods, dry density, and the type of tests conducted.

| Soil type              | Cement contents (%) | Wood Ash Content (%) | Molding dry density (kg/m³) | Curing periods (days) | Test Type       | Normalization index (%) | q* for normalization (kPa) |
|------------------------|---------------------|---------------------|-----------------------------|-----------------------|----------------------|-------------------------|---------------------------|
| Marine Deposited Clay  | 7, 10 and 13        | 1400 and 1600       | 7                           | UCS, Go, Direct Shear*, SEM* | \( \eta/(X_{iv})^{0.32} = 25 \) | 1650                     |
|                        |                     |                     | 28                          |                        | \( \eta/(X_{iv})^{0.32} = 25 \) | 1800                    |
|                        |                     |                     | 60                          |                        | \( \eta/(X_{iv})^{0.32} = 25 \) | 2050                    |
| Marine Deposited Clay  | 7, 10 and 13        | 5%                  | 1400 and 1600               | 7 UCS, Go, Direct Shear*, SEM* | \( \eta/(X_{iv})^{0.32} = 25 \) | 1700                     |
|                        |                     |                     | 28                          |                        | \( \eta/(X_{iv})^{0.32} = 25 \) | 2100                    |
|                        |                     |                     | 60                          |                        | \( \eta/(X_{iv})^{0.32} = 25 \) | 2400                    |
For direct shear tests, another set of samples with an average height of 30 mm and a diameter of 60 mm were prepared. The preparation procedure was similar to that of the previous specimens. Specimens were prepared in a specifically built (in the dimensions of test specimen) split mold and statically compacted into three equal layers while maintaining the same densities for the unconfined compressive strength specimens. Formed specimens were then cured for seven days in the direct shear device under the selected normal pressure.

2.2.2. Unconfined Compression Strength

A UCS test was conducted to investigate the effects of the addition of WA on the compressive behavior of cement-treated clay, according to ASTM C39 [42]. The fully automatic testing system (max. capacity: 20 kN, accuracy of 0.005 kN) was used. A constant strain rate was applied and controlled throughout the testing program.

2.2.3. Ultrasonic Pulse Velocity Test

In this study, the ASTM C 597-02 [48] was used to determine the ultrasonic pulse velocity. Samples tested during this stage were prepared, cured, inundated, and dried, as stated in the UCS test. The device used was MATEST Ultrasonic Tester Model C368. Two transducers were connected to the samples on both sides using grease. The distance between the transducers (the length of the specimen) was accurately determined. After that, the device sent a velocity wave, and subsequently, the time for the wave to travel through the sample was measured. The distorted form of the wave was viewed on the device’s screen.

The shear modulus was obtained from the product of the density of the specimen and the square of the shear wave velocity measured by the pundit. This is illustrated in the following formula:

\[
G_0 = \rho \cdot v^2
\]

where \(G_0\) is the shear modulus, \(\rho\) is the density of the sample, and \(v\) is the wave velocity measured from the device.

2.2.4. Direct Shear Tests

A direct shear test was performed on the aforementioned cylindrical samples of the 30 mm x 60 mm dimensions in accordance with ASTM D3080 - 11 [49] under consolidated drained conditions. Following the preparation of specimens, a computer-controlled apparatus was used for this test purpose. This apparatus was equipped with two load cells of 2-ton capacities for horizontal and vertical strength and two displacement transducers. The test started with the consolidation stage, where the sample was taken gradually to the required confining pressure and kept under consolidation for seven days to cure. Finally, the shearing stage started (strain rate: 0.1 mm/min) under the normal pressures (\(\sigma_1\)) of 100, 200, and 300 kPa up to a maximum displacement of 8 mm.

2.2.5. Scanning Electron Microscopy (SEM)

Once the UCS was finished, scanning electron microscopy (SEM) was applied to the samples to study the microstructure of the blends. For this purpose, a Jeol JSM-6480LV scanning electron was used.

3. Results and Discussion

3.1. Influence of Porosity/Binder Index on Unconfined Compressive Strength
Figures 6a, b, and c show the relationship between UCS and the adjusted porosity/binder index \((\eta/X iv^{0.32})\) of specimens prepared with 1400 and 1600 kg/m³ dry densities; 7%, 10%, and 13% cement content; and 7, 28, and 60 days of curing. Furthermore, the Figures 6a, b, and c present 5% wood ash, 10% wood ash, and no wood ash replacement, respectively.

A power relationship between \(q_u\) and \(\eta/X iv^{0.32}\) revealed good agreement for all blends. Previous empirical studies on various types of soils have shown that the exponent might vary slightly between 0.21 and 0.35. Therefore, in this study, 0.32 was chosen to be the best-fit exponent. Figure 6 shows that a unique correlation was developed for each blend.

Figure 6 and the available literature indicate that a reduction in compressive strength is due to an increase in porosity and reduction of blend content. Furthermore, in Figures 6a and 6b, where 10% and 5% of wood ash were used, respectively, the 7 and 28 days of curing did not affect the \(q_u\) at high porosity. On the other hand, curing for 60 days yielded an improvement. The contribution in all curing periods was more pronounced at low porosity.

The findings for the double binder of cement and both 5% and 10% wood ash in Figures 6(a) and 6(b) show a reduction in porosity due to wood ash replacement. In response to such replacement, compressive strengths at all curing periods increased. However, such behavior was not observed in Figure 6(c), where no wood ash enhancement was used.

Furthermore, aligning all the regression curves obtained from Figures 6(a), 6(b), and 6(c) into 6(d) revealed that the strength of the cement blend with 10% wood ash at all curing periods was nearly the same as that with cement alone; however, it reduced the porosity, and 5% wood ash incorporation accelerated the process of strength gain, yielding much more strength gain compared to cement alone at 60 days. The reason for this result is that the cement stabilized the coarse particles of portion with the hydration products from cement and wood ash to stabilize the clay fraction present in the mix.

These features are also evident in Figure 7, where \(q_u\) is plotted against \(\eta/X iv^{0.32}\) for all the tested blends. It is clear that incorporating 10% of wood ash as a replacement for cement reduces \(q_u\) and increases \(\eta/X iv^{0.32}\) compared to the reference samples. Therefore, to further encourage the rate of strength gain by activating the pozzolanic reactions, 5% of wood ash replacement was found to be
the most appropriate dosage, and the increase in $q_u$ was observed at all curing times. Wood ash, as 10% of cement replacement, reduced the compressive strength of the specimens relative to 5% wood ash+cement treated specimen for all curing times. Based on the SEM monograph that was further investigated, wood ash replacement enhanced the microstructure, reducing the available pores that were filled with hydration products. The clay–wood ash interaction shows better matrix properties and alumina in wood ash, acting as an accelerator for the chemical reactions. Overall, 5% of wood ash seems to optimize engineering properties. When compared with the porosity-strength value, an additional increase in the replacement value decreased performance compared to a 5% replacement level. Increasing the replacement level reproduces the amount of fine grains in the mixture, leading to an increase in the water demand. Thus, the weakening of the bonds increases the porosity and decreases the strength of the composites. Additionally, recent studies proved the usefulness of microstructural analysis of cement-based materials [34,50,51].

![Figure 7. Variation of unconfined compressive strength ($q_u$) with adjusted porosity/binder index for all blends considering cement only, cement+5% wood ash and cement+ 10% wood ash.](image)

Consoli et al. [52], influenced by Diambra et al. [53], established a general model via normalizing their results by dividing the established power equations to a particular value of strength. Using such an approach helped the authors predict the mechanical performances of cement-treated granular soils. In current study, 252 specimens with specific $q_u$ were chosen at $\eta/X_{0.32}$ = $V = 25$ for all the tested blends, as all of them had $V = 25$ (Table 3). As shown in Figure 8, Equations 4 (10 % wood ash), 5 (Cement only), and 6 (5% wood ash) yielded high coefficients of determination ($R^2$).
Figure 8. Normalization of unconfined compressive strength \( q_u \) (for the whole range of \( \eta/X_{iv0.32} \)) with adjusted porosity/binder index for all blends considering cement only, cement+5% wood ash and cement+10% wood ash.

\[
q_u = q_u(\eta/X_{iv0.32}=25) \times 2.6 \times 10^4 (\eta/X_{iv0.32})^{-3.16} \quad R^2 = 0.91,
\]
\[
q_u = q_u(\eta/X_{iv0.32}=25) \times 2.4 \times 10^4 (\eta/X_{iv0.32})^{-3.15} \quad R^2 = 0.96
\]
\[
q_u = q_u(\eta/X_{iv0.32}=25) \times 1.5 \times 10^5 (\eta/X_{iv0.32})^{-3.69} \quad R^2 = 0.91
\]

Furthermore, Figure 9 combines the wood ash replaced and cement-only unconfined compressive results normalized with the adjusted porosity/binder index. It is revealed that, by using Equation 7, a higher coefficient of determination \( R^2 = 0.93 \) was obtained.
Figure 9. Normalization of unconfined compressive strength \( (q_u) \), (for the whole range of \( \eta/X_{iv}^{0.32} \)) by dividing for \( q_u \) at \( \eta/X_{iv}^{0.32}=25 \). Considering all studied blends.

\[
q_u = 4.7 \times 10^4 (\eta/X_{iv}^{0.32})^{-3.34} \quad R^2 = 0.93
\]  

Equation 7 provides an important contribution, as it enables the \( q_u \) for specific blends of clays with 5% and 10% wood ash and cement cured for specific periods to be obtained using only one test. If possible, this test should be performed using three identical specimens to obtain a representative value of the strength at the chosen value of \( \eta/X_{iv}^{0.32} \). Based on the available literature and data, it is suggested to use \( \nabla \) values near 25. Based on the available literature and data, it is suggested to use \( \nabla \) values near 25.

To validate the applicability of Equation 7, the results of other authors on different soil types and blends were used in accordance with Equation 7 to predict the \( q_u \). Figure 10 shows the unconfined compression test results obtained by Consoli et al. [54] for cement-stabilized Portugal silty sand (Table 1) together with the line predicted using Equation 7. As suggested above and in the literature, \( \nabla = 25 \) was chosen, and the corresponding \( q_u \) was found to be 6300 KPa from the curve of Consoli et al. [54]. Substituting the value of \( q_u \) with \( X_{iv}^{0.32} = 25 \) in Equation 7 and varying \( X_{iv}^{0.32} \), a curve was plotted together with the laboratory test data. Figure 8 demonstrates that the predicted curve represents the experimental data. From the same paper by Consoli et al. [54], compression test results were obtained for London clay stabilized with cement. The data were taken from the specimens with \( X_{iv}^{0.32} = 25 \) and \( q_u \) (for \( \nabla = 25 \)) = 1100 KPa. By substituting these values into Equation 7 and varying \( X_{iv}^{0.32} \), a curve was plotted together with the laboratory test data. Figure 10 shows the curve obtained using Equation 7 that represents the experimental data, and the fit of the plotted curve shows a high degree of similarity with the curve plotted by experimental results.
Figure 10. Curve obtained using Equation 7 and laboratory testing data for Portugal Silty sand cement blend tested for unconfined compression and LC plus Portland cement blend tested for unconfined compression.

To summarize, creating a sample at 25\% porosity $X_{iv}$ content will enable any strength to be predicted. One can argue that to obtain the predicted curve, it is necessary to have the initial idea of the variation of $X_{iv}$. However, 25 is the value found in the previous literature. The relationship developed in this study to predict the $q_u$ was confirmed to yield successful predictions under all investigated conditions for fine- and coarse-grained soils treated with up to three different binders and cured up to 60 days.

3.2. Effect of Porosity/ Binder Index on Initial Shear Modulus

The curves plotted in Figure 11, $G_0$ vs. $X_{iv}$, and Figure 7, $q_u$ vs. $X_{iv}$ of the wood ash blends, are quite similar. Figure 11 shows a similar analysis for $G_0$. The results show a good correlation between $G_0$ and $X_{iv}$ for 5\% and 10\% wood ash mixes, where $R^2$ is found to be 0.8 for both mixes tested at 7, 28, and 60 days of curing. In this study, wood ash replacement improved the strength and stiffness of the clay matrix. Moreover, at high porosity, the effect of wood ash was not monitored; however, as the porosity decreased due to an extended curing period, the 5\% wood ash replacement showed improved strength. Data were more scattered than the compressive strength, although the correlation was good. The filler effect of the wood ash at 10\% did not allow us to highlight the 5\% wood ash contribution due to the pozzolanic reaction. This was also due to a denser matrix with the 10\% wood ash replacement rather than the 5\%.
A relationship of initial shear modulus as a function of porosity/binder index for compacted wood ash can be established. Like compressive strength analysis, the results can be normalized by dividing the established power equations by a specific value of stiffness. In this study, 106 specimens with specific initial shear modulus were chosen at $X_{0.32} = 25$ for all of the tested blends, as all of them had $\nabla = 25$ [$G_0$ (for $\nabla = 25$) = 3600 MPa (10% wood ash), 3800 MPa (5% wood ash)]. As shown in Figure 12, the form in Equation 8 yields good coefficients of determination ($R^2$) of about 0.80.
Normalization of initial shear modulus \( G_0 \) (for the whole range of \( \eta/X_{iv}^{0.32} \)) by dividing for \( G_0 \) at \( \eta/X_{iv}^{0.32} = 25 \) for considering all studied blends.

\[
G_0 = G_0(\eta/X_{iv}^{0.32}=25), 4.4 \times 10^3 (\eta/X_{iv}^{0.32})^{-2.61}, R^2 = 0.81,
\]

Equation 8 provides an important contribution, as it enables the initial shear modulus for blends of clays with 5% and 10% wood ash and cement-only cured for specific periods to be obtained using only one test for the first time.

3.3. Shear Strength Parameters

Typical shear strength and normal stress data with failure envelopes of marine clay, 5% wood ash, and 10% wood ash cement-replaced specimens with \( \eta/X_{iv}^{0.32} = 25 \) under effective normal stress of 100, 200, and 300 kPa and seven days as the curing period are shown in Figure 13. Table 4 shows that cohesion increased with the increase in cement content. Furthermore, the replacement of cement with 5% wood ash resulted in a further increase in cohesion, where a further increase in wood ash replacement to 10% resulted in reduced cohesion. An increase in cohesion at 5% wood ash replacement indicates the introduction of bonding strength along with clay particles. Stabilizing the marine clay with cement alone resulted in an increased friction angle from 27 to 42, where a further increase of cement content to 10% resulted in a slight decrease in friction angle to 39–40. Wood ash replacement did not seem to influence the friction angle of the specimens. Additionally, the brittleness index recommended by Bishop [55] was utilized to explore the softening response of blends with wood ash inclusion at varying cement contents. The brittleness index is defined as:

\[
I_B(\%) = \frac{\tau_{peak} - \tau_{post-rupture}}{\tau_{peak}} \times 100,
\]

Figure 12. Normalization of initial shear modulus \( G_0 \) (for the whole range of \( \eta/X_{iv}^{0.32} \)) by dividing for \( G_0 \) at \( \eta/X_{iv}^{0.32} = 25 \) for considering all studied blends.
Figure 13. Variation in direct shear failure envelopes for blends considering marine clay, 7% cement, 10% cement, cement + 5% wood ash, and cement + 10% wood ash.

Table 4. Shear strength parameters and average brittleness ratio of all blends.

|                  | Friction Angle (ϕ) | Cohesion (kPa) | Brittleness Index (%) |
|------------------|--------------------|----------------|-----------------------|
| Marine Clay      | 27                 | 10             | 0.00                  |
| 7% Cement        | 42                 | 57             | 42.60                 |
| 7% Cement 10% Ash| 42                 | 91             | 25.87                 |
| 7% Cement 5% Ash | 41                 | 123            | 25.48                 |
| 10% Cement       | 39                 | 164            | 54.06                 |
| 10% Cement 10% Ash| 40              | 216            | 32.87                 |
| 10% Cement 5% Ash| 39                 | 233            | 21.40                 |

where $\tau_{\text{post-rupture}}$ is the shear stress obtained at the maximum horizontal displacement. The average brittleness index of 100, 200, and 300 KPa consolidated specimens for each tested blend is shown in Table 4, which also shows that an increase in cement content increased the average brittleness index. Further, wood ash replacement was monitored to reduce the brittleness, with optimal reduction achieved at 5% wood ash replacement. The results are similar to unconfined compressive strength and initial shear modulus.

Table 4 shows that clays alone have a negligible brittleness index. Cement-clay blends have higher brittleness index value compared to clay-wood ash groups. Increasing the amount of wood ash causes an increase in the brittleness index. Additionally, cohesion value causes more stable matrix properties; hence, as it increases, the brittleness index decreases. Considering the number of clay-cement blends, as cement content increases, the brittleness index increases. However, 5% of wood ash replacement has better matrix properties and more cohesion at lower brittleness index value. It seems that cement and clay alone do not provide enough bonds, and less calcium silicate hydrate gel is formed during their interaction (i.e., less dense structure). In this study, this reaction was offset by wood ash.

3.4. Micro-Structure

Based on the XRF data in Table 2, XRD data in Figure 2, and EDX data in Figure 3, clay particles used in this study were of an Illite clay origin and contained mainly calcite ($\text{CaCO}_3$) and quartz ($\text{SiO}_2$).
Additionally, some amounts of alumina and iron were also available. The availability of aluminous elements in clay accelerated the chemical interaction with cement and wood ash particles to form a densified composite structure. Figure 14 shows the SEM of pure wood ash sample, and Figure 15 shows the SEM of pure clay sample. Additionally, Figure 17 shows the SEM of the clay-ash-cement sample.

Figure 14. SEM of pure wood ash sample

More reactive particles were observed on the SEM monograph (Figure 16). The portlandite phase is believed to be effective in producing secondary CSH gel when mixed with clay. This depends strongly on the mineralogy and treatment of the clay. Clays generally only have sufficient pozzolanic reactivity for the use as SCMs once calcined, and 2:1 clays (such as montmorillonite) are typically less reactive compared to kaolinite. As stated in Li et al. [56] study, it would have been beneficial to conduct a bound water or heat release test to measure the reactivity of the clay to understand the exact behavior of the composites. More gel formation due to secondary CSH reactions can help reduce the brittleness index and improve the strength of the composites. This is evident in Figure 15. Compared to clay particles, wood ash particles are irregular in shape as seen in Figure 14 and Figure 15.
Figure 15. SEM of pure clay sample

However, flaky and elongated particles need more water to become saturated and contain more voids. These voids can be considered as a water reservoir for later hydration process. The water in pores is being used by the chemical reactions to produce more gel, as seen in Figures 16b-d, to improve the strength and reduce the brittleness index. From the data presented in an Figure 1, the d50 of the wood ash is higher than that of the clay (and also the cement), so its addition would reduce the overall available surface area. Additionally, agglomerates of wood ash particles with no inter-particle bonding effectively act as large pores in the microstructure. These large pores might be acting as water reservoirs for later curing ages. Thus, lesser availability of cement decreases the strength and hence increases the brittleness of the matrix.
The scanning electron microscopy images, conducted on (a) wood ash at (b) 7 days, (c) 28 days and (d) 60 days, cured cement + wood ash blend marine deposited clays.

Based on the SEM micrographs shown in Figs. 14 and 16-a, and chemical analysis, the wood ash composed mainly of calcite and silica. The brighter colored structures indicate a glassy surface. These products have little pozzolanic due to their inert character. However, the elongated irregular shaped particles formed mainly from calcium, quartz, portlandite, and a small amount of merwinite indicated crystalline phase. They are cementitious minor oxide compounds, such as magnesium, manganese, and phosphorous. The angular, flaky, and elongated particles with sharp edges can be seen in Figure 14 and Figure 16a. This heterogeneity in the mixture increases porosity, followed by a decrease in strength. Naik et al. [23] and Siddique et al. [34] reported the same key findings for wood ash. The availability of a large amount of calcium led to the formation of more CSH gel. The marine clay is not calcined and is dominated by 2:1 clays, so it will have much lower pozzolanic reactivity compared to calcined clay used in SCMs [56]. Limestone may react, but its effects can be seen at later ages as seen in Figure 16-c. Li et al. [56] showed that each blend by the newly proposed method called “R³ technique” can be easily quantified. This method can help for better understanding the reactivity mechanisms.

Additional reaction with clay resulted in the denser matrix, as shown in Figures 16b-d. Needle-like structure, laminar phases, and angular flakes containing the porous region were also observed, as shown in Figures 16b-d. Figure 16b displays the unreacted carbon particles. These particles loosely bonded and contained more water. The carbon particles were inert, and they did not react easily, as seen in Figure 16b. This finding is in line with previous researches.

Additionally, hexagonally shaped calcium hydroxide plates were observed, as seen in Figure 16b. The porous nature of carbon particles seems to have an advantage with clay mixes. Some fibrous phases were also observed, as in Figure 16b. Adding clay into the system increased the rate of reactions. The particles connected better with each other, and pores filled up with hydrated products.
or transformed into a stable CSH gel. However, some pores and unreacted particles still appeared in SEM micrograph 16b. The unreacted carbon particles and aluminate hydrates decreased, as seen in Figure 16c. This can be attributed to the amount of water remaining in pores during mixing operation, leading to the additional water reservoir for reactions. This water can then be used to continue chemical reactions, as illustrated in Figures 16c and 16d.

The EDX data also confirmed that wood ash contains larger amounts of silicon, oxygen, and calcium and less aluminum. At the beginning of chemical reactions, the silicon and oxygen particles formed weaker bonds compared to aluminum and oxygen atom. Aluminum acted as a catalyst for the chemical reactions, and it was effective in wood ash blends. The wood ash used in this study was obtained from the local houses. Generally, the heat produced during the burning of wood did not exceed 100°C. This is consistent with the high number of unreacted particles observed in SEM and high LOI in chemical analysis of wood ash. LOI value was very high compared to the recommended levels in cement (higher than 6%). This makes the wood ash not suitable for load-bearing applications. However, for controlled low strength or moderate strength applications, it can be used. Most of the research conducted on wood ash was collected from the biomass operation where heat reached more than 500°C. At this stage, fewer unreacted carbon particles were available. However, it seems that more gel was formed when unreacted particles came in contact with the clay mix. Additionally, water in pores is readily available for continuous hydration; thus, the denser matrix will be formed, as seen in Figures 16c and 16d.

Based on the EDX data shown in Figure 4, the wood ash sample contained mainly calcite and silica; therefore, both crystalline and amorphous structures were available. This is consistent with the SEM analysis of the pure wood ash sample in Figure 14. The availability of the amorphous phase indicated pozzolanic. However, the crystalline phase indicated the availability of inert particles. Accordingly, the hydration was slower, and it needed more curing for pores to be filled with hydration products. Especially, SEM monograph (Figure 16 and Figure 17) clearly shows this situation. Additionally, a high amount of CaO in wood ash helped form strong bonds when interacting with clay and cement, as shown in Figure 17.

Figure 17. SEM of Clay-cement ash sample
The number of pores decreased, and the unreacted particles transferred to a stable denser gel structure. CSH gel and some calcium aluminate hydrate flakes formed as seen in Figure 16b-d and Figure 17. However, incorporating wood ash increased the alkalinity of the system, destabilizing the expansive aluminate gel. An increase in the consumption of the portlandite and refinement of pores by lowering the aluminate phase improved the composite’s microstructure, as observed in Figures 16c and 16d and cement-clay-wood ash interaction in Figure 17. Similar findings were reported by Mehta and Monteiro [57] and Siddique et al. [34]. The ettringite formation declined due to the addition of illite origin clay and cement as shown in SEM graphs. In current study, the total oxide amount (24.2) was lower than 70%.

The wood ash used in this study might have been inert and acted as a filler material. This can be seen in Figures 16c and 16d, showing a continuous decrease in porosity. It is also evident from the SEM results that wood ash-clay blended mixtures (Figure 17), especially wood ash that had high LOI, needed more curing to improve their matrix properties. Better bonding was attained beyond 60-days. However, the available pores, as shown in Figure 16d, did not generate problems, since the number of pores was lower. The pores were mainly from the clay particle. In the beginning, the distance between the particles was high. The particles were then separated from each other. However, the distance diminished in later ages due to formation of densified matrix, as seen in Figure 16d.

Additionally, a few bundles of calcium aluminate silica hydrate needles (strâlingite) were formed. Poorly reacted matrix transformed into a densely packed, more stable structure. Additionally, most clay particles seemed to react at this stage. A negligible amount of expansive gel formation was observed in Figure 16d. Formed gels combined with a strong bond. This improved the strength of the composites.

Additionally, the amorphous phase transformed into a defect-free crystalline phase. Arrangement of particles here seems to be more ordered with lesser pores, as observed in Figure 16d. It is believed that beyond the 60-days of curing, the performance of the composites will further improve. Many researchers [10,28,58] have reported similar findings.

5. Conclusions

From the results presented in this study, the following conclusions can be drawn:

- The mechanical response of the wood ash blend is most beneficial at 5% wood ash replacement. The 5% of wood ash replacement seems to be the optimum replacement percentage to improve matrix properties.
- By testing one sample, the UCS and initial stiffness can be predicted for all other wood ash content and cement content at any curing time and density.
- The proposed adjusted porosity/binder index formula can predict the strength and initial stiffness of other binder products, which was also confirmed in other studies.
- The replacement of cement with 5% wood ash improves cohesion, and above this replacement level to 10% results in a reduction of cohesion.
- The long-term effective cohesion and friction angle parameters are aligned with the results of the unconfined compressive strength and initial stiffness. However, more tests should be carried out to develop a correlation of those shear parameters with the binder index.
- Based on the microscopic investigation, marine clay makes an excellent contribution to gel formation. Heterogeneity of the interstitial transition zone improves with the incorporation of the wood ash into a system. The high LOI value of wood ash diminishes the negativity at later ages, acting as a water reservoir to enhance the hydration process to produce better-densified matrix properties. The porosity slightly reduces with the help of cement-clay-wood ash interaction. The duration of laminar phases, which were observed in an SEM monograph, decrease after 28 days; further, particles are interconnected with a strong bond. The authors believe that this beneficial effect would be more visible after 60 days and recommend the microscopic investigation beyond 60 days, especially for clay blends.
• Reusing unsuitable soil and hazardous wastes will reduce environmental and financial impacts. Improving soil with additives will facilitate the use of the available soil on site. In addition to the environmental contribution of cement usage reduction, using waste material, such as WA, will enable safe disposal of those harmful materials.
• The incorporation of such material on site does not require any specific tool; field application is conventional and straightforward.
• Conducting bound water or heat release test to measure reactivity would be beneficial for mixtures composed of clay with cementitious constituents.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Figure S1: Compaction curve of the clay used with the two chosen densities.

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