Critical Review of the Parameters Affecting the Effectiveness of Moisture Absorption Treatments Used for Natural Composites

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Abstract: Natural composites can be fabricated through reinforcing either synthetic or bio-based polymers with hydrophilic natural fibers. Ultimate moisture absorption resistance at the fiber–matrix interface can be achieved when hydrophilic natural fibers are used to reinforce biopolymers due to the high degree of compatibility between them. However, the cost of biopolymers is several times higher than that of their synthetic counterparts, which hinders their dissemination in various industries. In order to produce economically feasible natural composites, synthetic resins are frequently reinforced with hydrophilic fibers, which increases the incompatibility issues such as the creation of voids and delamination at fiber–matrix interfaces. Therefore, applying chemical and/or physical treatments to eliminate the aforementioned drawbacks is of primary importance. However, it is demonstrated through this review study that these treatments do not guarantee a sufficient improvement of the moisture absorption properties of natural composites, and the moisture treatments should be applied under the consideration of the following parameters: (i) type of hosting matrix; (ii) type of natural fiber; (iii) loading of natural fiber; (iv) the hybridization of natural fibers with mineral/synthetic counterparts; (v) implantation of nanofillers. Complete discussion about each of these parameters is developed through this study.

Keywords: moisture treatments; interfacial adhesion; diffusivity; hydrophilicity; hydrophobicity

1. Introduction

The natural fiber-based composites have numerous implementations in different industries since they are good acoustic absorbers and thermal insulators. Furthermore, they can be used as indoor humidity regulators for building through using coatings and layers of natural composites in interior walls’ structure [1,2]. Natural fiber-based composites have the potential to be implemented in civil engineering applications such as manufacturing materials for the construction of buildings when the durability and structural performance are considered. Appropriate treatments to improve the durability performance of natural composites will make them in general more competitive to synthetic fiber-based composites [3–5]. The natural fiber-based composites are the most widely used as poroelastic materials in the automotive industry sector. In addition to their higher specific mechanical properties, natural composites can effectively reduce the level of acoustic noise inside the vehicle cabin, similar to synthetic polyurethane foam and commercial rock wool fibers [6–8]. The broad field of application of natural composites can be attributed to the properties of natural fibers, which are characterized by low weight, affordable cost, easy processing, flexibility, biodegradability, and environmentally friendly fibers. The properties of natural composites depend mainly on the characteristics of the reinforcing fiber and the adhesion between the matrix and the fiber [9–11].
are a number of drawbacks that should be resolved in order to disseminate the implementation of natural composites in various industries, such as inadequate adhesion between the hydrophobic hosting matrix and raw fibers, low wettability by non-polar plastics, and high level of water uptake [12–14]. The natural fiber is composed of cellulose, hemicellulose, lignin, pectin, wax, water-soluble substances, and fat. Cellulose is regarded as the main framework element of the fiber structure. It imparts stiffness, strength, and structural stability to the fiber. However, cellulose and hemicellulose are the major constituents of natural fiber that absorb the highest amount of moisture. Lignin provides effective shielding for hemicellulose/cellulose against harsh environmental circumstances such as humidity and temperature [15–17].

The absorption behavior of moisture can be described according to two main models: (i) pseudo-Fickian behavior, where the weight gains of water didn’t reach equilibrium after take-off; and (ii) linear Fickian behavior, where the weight gains of water gradually achieved equilibrium after quick initial take-off. The higher moisture absorption facilitates the microbial attack, leading to a process called biodegradation. The transport of moisture inside the natural composites can be facilitated through different mechanisms such as the diffusion of water inside the microgaps among the polymer chains. The water absorbed in the natural composites is classified into bound and free water. The bound water are dispersed molecules of water that are bounded to the polar groups of the polymers, while the free molecules of water have the capability to transport independently through cracks and voids. When the natural composites are exposed to moisture, the molecules of water diffuse into the composite and attach onto hydrophilic groups of natural fibers, forming an intermolecular bonding of hydrogen with the fibers that mitigates the interfacial adhesion at the fiber/matrix interface. The degradation of natural composite’s properties take place when the swelling of cellulosic fibers create stress at the interfacial regions, leading to the mechanism of micro-cracking in the matrix close to the swollen fibers, which emphasizes transport and capillarity through micro-cracks. With the excessive absorption of water, the bound water increases while the free water decreases. Water-soluble materials begin to leach from fiber, and finally lead to ultimate debonding between the matrix and fiber. The aforementioned diffusion mechanism of water molecules into the natural composite structure is illustrated in Figure 1 [18–25].

![Figure 1](image_url)

**Figure 1.** The degradation of interfacial adhesion due to moisture absorption: (a) development of micro-cracks due to expansion of swollen fiber, (b) water molecules diffused in the bulk matrix flow along the fiber–matrix interface, (c) water-soluble components leach from fiber, (d) ultimate matrix–fiber debonding takes place. Reproduced with permission from [18].

The moisture absorption causes all three dimensions of the laminated composite to increase. Nevertheless, the increase of the composite thickness is very large relative to the increase in width and length. The natural fibers are influenced by water uptake in two directions: (i) swelling of the fiber itself; and (ii) changes in the fiber’s density through the weight of absorbed moisture. The water uptake of natural composite is attributed to the polar and hydroxyl groups found at natural fibers. The hydrolysis (chain segmentation of polymer) and swelling emerging from moisture absorption
induce debonding between matrix and fibers, delamination of the laminated composite, and physical
damage to the polymeric matrix, which increase the mechanical and thermal degradation of natural
composites’ properties [26–30]. Nevertheless, the minor swelling of natural fibers has a positive effect
on the mechanical properties of natural composites, since this emphasizes the mechanical interlocking
between the matrix and fiber [31]. The water uptake is responsible for increasing the mobility of
the side groups and molecular chains, which leads to the reversible plasticization of the polymer
matrix [32–35]. The matrix plasticization increases the fracture toughness, while it also reduces the
strength, stiffness, fatigue durability, and natural frequencies of the natural composite [36–45]. When
the sisal fiber-reinforced polypropylene (PP) composite is immersed in hot water at 90 °C, the tensile
strength and modulus are reduced with the increasing immersion time in the water, while the impact
strength increases initially with the immersion time. The variety in behaviors between impact and
tensile properties can be explained through the swelling effect of the reinforcing fiber and plasticization
of the interface between the fiber and the matrix [46]. The diffusion of water molecules into some
types of polymer films could substantially change the intermolecular interaction, which significantly
contributes to increasing the thermal conductivity, and as a result reduces the thermal insulation
properties of natural composites [30].

The water uptake absorption behavior of natural composites is largely affected by many factors
such as the chemical and physical treatments of natural fibers, fiber loading, fiber distribution, fiber size
and shape, linear mass density of natural yarns, permeability of the fiber, crystallinity of hosting resin,
hydrophilicity of each composite’s constitute, void content especially at the fiber–matrix interface,
duration of exposure to humid environment, fabrication method, conditioning medium circumstances,
including temperature, and RH. The surface treatments of natural fibers reduce the hydrophilic nature
of natural fibers and introduce significant changes to the surface morphology. The architecture of
the fabric has a significant impact on the amount of moisture absorbed by the natural yarns. For
example, the twill weave fabrics have optimum mechanical performance even after absorption of
moisture. For certain yarn architecture, a lower linear mass density of reinforcing natural yarns
provides the composite with greater resistance toward moisture, while retaining at the same time
optimum mechanical properties during the moisture exposure period [47–54]. The moisture absorption
of natural composites is influenced by the fiber arrangement within the structure of the composite.
For example, the non-woven form of the hemp fiber-reinforced polylactic acid (PLA) composite
absorbs a higher amount of moisture relative to the aligned fiber composite structure. This can
be attributed to the increased porosity resulting from the increased complexity of the matrix flow
path during the fabrication process [55]. The surface hydrophobicity of the natural fibers can be
enhanced through the application of appropriate chemical treatments, additives, nanocoatings, and
bio-based coatings [56–59]. If the coating is appropriately applied and the material characteristics of
the coating are carefully chosen, the diffusion of water molecules can be reduced by 50%. Generally,
coatings are only effective in offering short to medium-term protection against moisture [60,61].
Natural composites with a higher content of voids absorb a higher amount of moisture relative to the ones without voids [62]. Managing the manufacturing environment of natural composites such
as maintaining it under vacuum and low humidity is fundamental to reduce the void content that
could be created during the manufacturing process. Drying of natural fibers is usually needed to
eliminate excessive moisture content. The moisture absorption is directly correlated with the type of
natural fiber. For example, flax and hemp fibers have a lower capability to absorb moisture during
the manufacturing process of composite relative to ramie and cotton fibers [63]. The hybridization of
natural fibers with other synthetic or natural fibers characterized by low moisture absorption tendency
is considered an effective technique to reduce the moisture absorption of natural composites [64].
The ambient RH has a direct effect on the moisture content of the composite as well as the average at
which it accumulates [65,66]. Additionally, it is demonstrated that the increasing temperatures increase
the equilibrium moisture absorption and rate of water diffusion due to the activation of the diffusion
process [67–69]. However, it was reported in the literature that in some cases, the diffusion coefficient
increases with decreasing temperatures, since the water molecules bind to the fiber surface through the amorphous waxy materials, which facilitates the water sorption [70].

There is an urgent need to consider both the material durability under different weathering conditions (temperature, rain, air, solar radiation, and oxygen) and life cycle effect during the composite design stage [71–73]. The water uptake and swelling behavior of natural fiber-based composites should be studied thoroughly in order to evaluate the consequent impacts on the performance of composite parts. The swelling coefficient can be calculated through the following equation:

\[ T\% = \left( \frac{T_2 - T_1}{T_2} \right) \times 100\% , \]  

where \( T_1 \) and \( T_2 \) are the initial and final thicknesses of the composite, respectively, while the water uptake can be calculated through the following relation:

\[ W\% = \left( \frac{W_2 - W_1}{W_1} \right) \times 100\% , \]  

where \( W_1 \) and \( W_2 \) are the initial and final weights after immersion [74,75]. The moisture absorption behavior can be described through: (i) diffusivity or the coefficient of diffusion, which characterizes the transport rate of water molecules inside the composite, and controls the duration required to reach the equilibrium moisture state; (ii) the moisture content at equilibrium condition, which describes the composite affinity for water; and (iii) the duration of transient, which is directly influenced by the thickness of the composite [76,77]. To the best of our knowledge, none of the reviewed studies in this review investigate the major parameters that control the efficiency of chemical and physical treatments on improving the moisture absorption properties of natural composites. The combining effect of these parameters along with applying chemical and/or physical treatments could increase or decrease the efficiency of the treatment used. Therefore, this study provides a significant contribution to the existing literature through summarizing the factors that affect the moisture absorption treatments used for the most common natural fiber-based composites which are illustrated in Figure 2 and discussed subsequently in the forthcoming sections of this study.

![Figure 2. Main parameters affecting the efficiency of chemical/physical treatments.](image)

2. The Treatment Types of Natural Composites

The surfaces of natural fibers are composed of non-cellulosic substances (lignin, hemicellulose, and pectin) and waxes, which prevent fibers from forming adequate adhesion with the hosting matrix. It is recommended to increase the hydrophobicity of the natural fibers in order to produce natural composites with improved mechanical, tribological, and moisture absorption properties. This can be achieved through introducing the surfaces of natural fibers to physical and chemical treatments [78–81]. Treating the natural fibers with chemical and physical methods eliminates the hydrophilic elements from the surface of the fiber. Hence, the content of lignin is increased, leading to the considerable reduction of the fiber’s hygroscopic behavior [82,83]. The effect of various types of treatments on enhancing the moisture absorption behavior of natural composites can be realized through exploring the improvement attained on the moisture absorption behavior for different types of natural composites, as shown in Figure 3. It can be noted that some types of treatments have negative consequences on the
water uptake behavior of some composites, such as jute fiber/UP and sisal fiber/starch, which are treated with hot water and alkalization methods, respectively. This reduction can be attributed to the removal of hydrophobic components such as lignin from the treated fiber’s structure, which as a result induces the composite to absorb a higher amount of moisture.

![Figure 3](image-url)

**Figure 3.** The effect of different treatment methods on the improvements attained on the moisture absorption behavior of natural composites. 30 wt. % Abaca/PP (1920 h) [84], Bagasse/UP (720 h) [85], Bamboo/PES (1440 h), immersion in boiling water (2.0 h) [86], 17 wt. % Coir/PES (1440 h) [87], 40 vol. % Jute/UP (960 h) [88], 15 wt. % Kenaf/epoxy (816 h) [81], Oak wood (168 h) [89], Spruce wood (168 h) [89], 15 wt. % Sisal/starch [90].

The moisture absorption treatments are classified into chemical and physical treatments according to their capabilities of changing the chemical composition and structure of natural composites. The chemical and physical treatments that are commonly used to improve the moisture absorption properties of various types of natural composites are discussed in detail in Sections 2.1 and 2.2, respectively.

### 2.1. Chemical Treatments

The chemical treatments can eliminate the OH coating from the natural fibers and increase the roughness of fiber surface, leading to a noticeable increase in the mechanical interlocking of the natural fiber with the hosting resin [91–93]. With increasing chemical treatment time, the treated fibers exhibited a significant increase on the surface roughness, and the impurities are eliminated to the highest extent, as shown in Figure 4. It can be inferred from the scanning electron microscope (SEM) images of sisal fibers treated with sodium bicarbonate that a noticeable reduction occurred at the fiber diameter for all of the treated fibers due to the elimination of a specific portion of the natural oil covering the surface the fiber’s cell wall in addition to lignin, waxy, and hemicellulose substances. Upon the removal of lignin and hemicellulose from the fiber’s structure, the internal constraints located within the interfibrillar region are released. Hence, the cellulose fibrils reorganized themselves into a highly compacted structure, leading to a higher dense packing of cellulose fibrils’ chains [94].

The natural fibers can be treated through using one of the following chemical treatments:
2.1.1. Acetylation

Applying acetic anhydride treatment to various types of natural fibers can enhance the moisture resistance properties of natural composites. This improvement is attributed to the elimination of lignin and hemicellulose elements from the treated fibers [16,95]. Through the acetylation process, the acetyl groups (CH$_3$CO) react with the hydroxyl groups (OH) of the fiber, which makes the surface of the fiber more hydrophobic [96,97].

2.1.2. Benzoylation

In this treatment, fibers are treated with alkali solution with a specific NaOH concentration first, followed by benzoyl chloride treatment. Then, the OH groups of the fibers are further substituted by the benzoyl group, which then attaches to the backbone of the cellulose [16].

2.1.3. Treatment by Peroxide

The peroxide treatment includes the grafting of PE adheres onto the surface of the natural fibers. The peroxide treatment mechanisms include an alkali pretreatment on the fibers, which are then immersed with dicumyl or benzoyl peroxide (about 6% concentration) in acetone aqueous for a duration of 30 min [16,98].

2.1.4. Treatment by Isocyanate

The chemical bond formed by isocyanate offers a strong covalent linkage between the matrix (especially thermoplastic one) and the fiber. Furthermore, the moisture that exists on the fiber surface reacts with isocyanate and forms urea, which can react more with the hydroxyl groups of the celluloses. This secondary reactions lead to better linkage between the fiber and matrix, and better moisture barrier properties of the fiber [16,99].

2.1.5. Acidic Treatments

In this treatment, different types of acids can be used in order to improve the hydrophobic nature of natural fibers such as stearic, hydrochloric, and acrylic acids. The hydrophilic hydroxyl groups of the fibers react with the carboxyl group of the treating acid and consequently enhance the moisture resistance properties. Moreover, the treating acid enhances the mechanical interlocking of the fiber with the hosting matrix through the fibrillation of the fiber bundle [16,100]. However, removing non-cellulosic elements in a wide range through treatments such as hydrochloric acid induces the degradation of the fibers and considerably impacts the transverse mechanical properties of the composites [101,102].

2.1.6. Alkalization

This treatment is the most feasible method that can effectively emphasize the interfacial adhesion between the fiber and matrix, leading to significant improvement on the moisture absorption properties.
of the treated composite [15,103–106]. The hydrophilic hydroxyl groups located at the surface of natural fibers can be reduced through treating them with NaOH solution. Furthermore, this treatment can eliminate a specific portion of the lignin, wax, pectin, hemicellulose, and oil that is covering the materials [107]. After applying NaOH treatment, the surface of the fiber becomes rougher and more organized because of the removal of microvoids, which enhances the capacity of stress transfer among the ultimate cells [17,108]. Furthermore, it mitigates the diameter of the fiber, increasing the aspect ratio of the fiber. This participated in increasing the surface area effectively for better adhesion with the reinforced matrix. Treating natural fibers with a lower concentration of alkaline solution is found to be better than treating them with higher concentration solutions [109,110]. Treating hemp, coir, kenaf, and alfa fibers with 5–6% of NaOH cleans the fiber surface through eliminating the amorphous compounds, and increases both the crystallinity index of the fiber bundle and the hydrogen groups at the fiber surface [111–116]. The treatment time and concentration of alkali can adversely influence the properties of the fibers. If the fibers are not washed adequately, the alkali will progress to degrade the treated fibers long after exposure, leading to the considerable breakdown and swelling of the fiber. Furthermore, there are some undesirable consequences associated with the alkalization process, such as high surfactant content, high pH values, and contaminated wastewater remaining after treatment [60,117]. Consequently, the concentration of alkaline solution should be carefully selected in order to avoid the deterioration of the structural properties of natural fibers such as the considerable fibrillation of the natural fiber bundle, which occurs when the fiber is treated with a higher concentration of NaOH, as demonstrated in Figure 5 [118].

![Figure 5](image_url)

Figure 5. The alkalization effect on fibrillation of hemp fiber: (a) raw, (b) 5 wt. % NaOH and (c) 10 wt. % NaOH. Reproduced with permission from [119].

The improvement that can be achieved from alkalization is subjected to the type of natural fiber. For example, treating the water hyacinth fiber-reinforced UP composite with alkali solution does not change the moisture absorption resistance considerably. This can be attributed to the physical effect that take place in the cell wall of the fiber, creating many more micro and nano hollow cavities, crevices, and porosities that enable the fiber to absorb more moisture [120]. The alkalization has an adverse effect on the moisture absorption properties of specific types of natural fibers [81]. For instance, treating sisal fibers with 5% NaOH makes the sisal/starch composite absorb a higher amount of moisture relative to the untreated one, while there is a slight reduction noticed in the diffusivity of the treated composite [121]. The alkalization of banana fibers swells the structure of the treated fiber because of considerable alteration in the morphology, structure, dimensions, and mechanical properties [122]. The combination of silane and alkali treatments exhibits a superior enhancing effect on the moisture absorption properties relative to single treatment [123–125]. More details about different types of silane coupling agents are discussed in the next section.
2.1.7. Coupling Agents and Grafting

The coupling agent promotes the interfacial interaction between the polymer matrix and natural fibers, enhancing the compatibility at the matrix–fiber interface. Strengthening the interfacial adhesion reduces the water uptake behavior, since the gaps and voids content at the fiber–matrix interface is minimized to the highest extent. Silane coupling agents introduce improvement to the moisture absorption behavior and diffusivity of different natural composites up to 89% and 96%, respectively, as demonstrated by the values listed in Table 1 [126–128]. The coupling agent can serve two main functions: it reacts with the functional groups of the matrix, and then it reacts with the –OH groups found in cellulose. Specifically speaking, the silane molecule links between cellulose and resin by creating a chemical bond with the cellulose surface through a siloxane bridge, while its organofunctional group bonds to the polymer matrix [129]. This reaction process reduces the number of hydroxyl groups found on the fiber surface. The chains of hydrocarbon emerging from silane application prevent the fiber swelling through the cross-linked network that is formed due to covalent bonding between the fiber and matrix. The interaction between silane-treated fibers and the hosting matrix is much stronger than that of alkali-treated ones [16,130].

Table 1. The improvements on moisture absorption properties attained from using coupling agents.

<table>
<thead>
<tr>
<th>References</th>
<th>Composite</th>
<th>Immersion Time (Hours)</th>
<th>Coupling Agent</th>
<th>Moisture Improvement %</th>
<th>Diffusivity Improvement %</th>
</tr>
</thead>
<tbody>
<tr>
<td>[131]</td>
<td>50 wt. % Alfa pulps/LDPE</td>
<td>480</td>
<td>HDS</td>
<td>37.14</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>MPS</td>
<td>25.71</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>MRPS</td>
<td>14.29</td>
<td>-</td>
</tr>
<tr>
<td>[86]</td>
<td>Bamboo matting/polyester</td>
<td>1440/2 (^1)</td>
<td>APTES</td>
<td>62.75/89</td>
<td>95.38</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>TMSPMA</td>
<td>37.25/55.5</td>
<td>87.05</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>VTMOE</td>
<td>52.94/84.2</td>
<td>95.81</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>TESPTS</td>
<td>13.73/24.5</td>
<td>16.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>APTMOS</td>
<td>31.37/66.67</td>
<td>55.53</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>OTMOS</td>
<td>50.98/36.67</td>
<td>90.48</td>
</tr>
<tr>
<td>[132]</td>
<td>30 wt. % Ijuk/PP</td>
<td>240</td>
<td>VTMS</td>
<td>5.22</td>
<td>-</td>
</tr>
<tr>
<td>[25]</td>
<td>Jute/epoxy</td>
<td>336 (^2)</td>
<td>GLYMO</td>
<td>8.0</td>
<td>-</td>
</tr>
<tr>
<td>[133]</td>
<td>10 wt. % Luffa/PP</td>
<td>960</td>
<td>MS</td>
<td>34.3</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>AS</td>
<td>39</td>
<td>-</td>
</tr>
</tbody>
</table>

\(^1\) Immersion in boiling water. \(^2\) Immersion time for one cycle of absorption and desorption of moisture.

Treating natural fibers with different types of silane coupling agents considerably reduces the water uptake behavior of the treated fibers. Treating sisal fibers with 1% of an MPS coupling agent reduces the water absorption from 123.8% to 64.4% and the equilibrium moisture content from 12.8% to 1.7%. Vinyltrimethoxysilane, \(\gamma\)-aminopropyl triethoxy silane, and GPS are silane coupling agents that can be used to effectively reduce the water uptake tendency of oil palm, sisal, and jute fibers, respectively [128]. The MA grafted polymer is one of the most common methods used to reduce the moisture absorption capability of natural composites [133]. Moreover, graft copolymerization reduces the hydrophilic properties of biopolymers without altering their inherent properties [134]. The MA can reduce the hygroscopic behavior through the reaction with –OH groups at the surface of the natural fiber to create ester linkages, which slightly decreases the moisture absorption values [135]. The grafting of MA is frequently used with PP resin, which has an outstanding improving effect on the moisture absorption properties, as can be inferred from the values listed in Table 2.

It can be concluded from the values listed in the table above that the alkalization of fibers prior to grafting induces the composite to absorb a higher amount of moisture relative to silane-treated fibers. Removing the non-cellulosic elements of natural fibers after alkalization influences the capability of MA to link the fibers with the hosting resin.
Table 2. The effect of MA grafting on moisture absorption properties.

<table>
<thead>
<tr>
<th>References</th>
<th>Composite</th>
<th>Aging Conditions</th>
<th>Coupling Agent</th>
<th>Moisture Content % (Untreated)</th>
<th>Diffusivity, m²/s (Untreated)</th>
</tr>
</thead>
<tbody>
<tr>
<td>[84]</td>
<td>30 wt. % Abaca/PP</td>
<td>95% RH at 50 °C for 1920 h</td>
<td>MAPP</td>
<td>9.62, (15.09)</td>
<td>-</td>
</tr>
<tr>
<td>[136]</td>
<td>50 wt. % Kenaf/PP</td>
<td>24 h</td>
<td>2 wt. % MAPP</td>
<td>1.05</td>
<td>-</td>
</tr>
<tr>
<td>[52]</td>
<td>Jute/PP</td>
<td>95% RH at 23 °C for 18 h</td>
<td>2 wt. % MAPP</td>
<td>21.5</td>
<td>-</td>
</tr>
<tr>
<td>[133]</td>
<td>10 wt. % Luffa/PP</td>
<td>960 h</td>
<td>2 wt. % MAPP</td>
<td>28.4</td>
<td>-</td>
</tr>
<tr>
<td>[77]</td>
<td>50 wt. % Rice husks/PP</td>
<td>Untreated fiber + 2% MAPP</td>
<td>9.91, (11.39)</td>
<td>$1 \times 10^{-13}$</td>
<td>$(2.37 \times 10^{-13})$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Untreated + 2% MAPP + 1% SEBS-g-MA</td>
<td>9.38</td>
<td>$8.37 \times 10^{-14}$</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>NaOH + 2% MAPP</td>
<td>15.31</td>
<td>$1.50 \times 10^{-13}$</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>NaOH + 2% MAPP + 1% SEBS-g-MA</td>
<td>14.35</td>
<td>$1.33 \times 10^{-13}$</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Silane + 2% MAPP</td>
<td>9.45</td>
<td>$4.65 \times 10^{-14}$</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Silane + 2% MAPP + 1% SEBS-g-MA</td>
<td>9.71</td>
<td>$5.65 \times 10^{-14}$</td>
<td></td>
</tr>
<tr>
<td>[126]</td>
<td>60 wt. % WF/PP</td>
<td>23 °C and 50% of RH for 48 h</td>
<td>5% MAPP</td>
<td>11.57, (14.57)</td>
<td>-</td>
</tr>
<tr>
<td>[137]</td>
<td>15 wt. % Wood flour/PP</td>
<td>96 h</td>
<td>1 wt. % A-1100</td>
<td>1.29, (1.39)</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1 wt. % A-174</td>
<td>1.14</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 wt. % MAPP</td>
<td>1.20</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>1% A-1100 + 5 wt. % MAPP</td>
<td>1.09</td>
<td>-</td>
<td></td>
</tr>
</tbody>
</table>

2.1.8. Polymeric Coatings

In order to prevent the fast diffusion of water molecules into the composite structure specifically in the interfacial region between the fiber and matrix, it is recommended to apply a suitable thickness of resin-rich surface at the composite surface and to keep this resin layer intact and free of cracks throughout the service life of the composite [138]. Natural composites can be coated through films fabricated from either synthetic polymers or natural materials, which proved their capability on reducing the moisture absorption [139]. Applying coatings to the natural fibers introduces a higher degree of moisture resistance relative to chemically treated ones [56]. Plastic adhesive tape of PP with a thickness of 0.1 mm is used to coat the short jute fiber/PLA composite. It is demonstrated that the water uptake behavior can be efficiently retarded after applying this coating [140]. Acrylated epoxidized soybean oil polymer is used as a coating for the flax fiber-based composite. It is found out that the moisture absorption property is enhanced through the applied coating [141]. A film composed of PE is implemented to coat the hemp fiber-based composite. After applying the coating, the water uptake behavior is enhanced considerably. The PE film inhibits the water molecules from penetration into the composite, since the polymer film has no tendency to absorb moisture. The water uptake test showed that the moisture absorption is reduced by 88.5%, 84.8%, and 68.5% when composite samples are exposed to ageing conditions for 2 h, 24 h, and 120 h, respectively [142]. A jute fiber-reinforced epoxy composite is coated with acrylic paint and epoxy resin. Relative to the raw composite, the applied coatings increase the moisture resistance of the treated composite. The alkalization of natural fibers prior to the application of coatings increases the composite’s capability to resist moisture [143]. It is important to highlight that there are specific types of coatings that have an adverse effect on the moisture absorption resistance and consequent fiber swelling. For example, treating the surface of oil palm ash, which is used to reinforce the natural rubber composite, with liquid epoxidized natural rubber retards the crosslink density of the filler with the hosting matrix, and mitigates both the moisture absorption and swelling resistances of reinforcing fillers [144].

2.2. Physical Treatments

The physical methods such as the fabrication of hybrid yarns, calendaring, thermal treatment, and stretching do not alter the chemical composition of natural fiber. Physical treatments change
the surface and structural properties of the fiber, and hence affect the mechanical bonding with the hosting matrix [145]. Thermal treatment can be used to treat wood composite panels in order to impede water molecules from penetrating inside the composite structure. This treatment increases the compactness of the wood structure both in the panel cross-section as well as on the surface due to the macromolecular rearrangement and water loss. This compactness provides the effective stabilization of the wood assembly under water aging circumstances [89]. Fewer hydrophilic fibers can be produced by a retting process where a high amount of waxes can be deposited on the fiber’s surface. The steam explosion technique (STEX) competes effectively with the retting process in terms of producing fibers with less hydrophilicity. During the STEX process, additives and steam are diffused into the fiber interspaces of green fiber bundles due to the applied process conditions of increased temperature and adequate pressure. Consequently, the center lamella can be improved at ideal conditions of reaction. The subsequent abrupt relaxation of the steam leads to an efficient breaking up of the green bast fiber, which results in substantial decomposition into individual fibers (DDA). Relative to raw fibers, Duralin and DDA-treated fibers (both of them retrieved from the STEX process) clearly exhibit the most hydrophobic surfaces [146]. Manipulating the natural fillers with plasma treatment increases their hydrophilic behavior, providing them with better sorption aspects. This treatment eliminates part of the lignin content, transforming it to a surface with a higher degree of hydrophilicity [147,148]. Through oxygen plasma treatment, different functional groups can be used to introduce different functional groups on the surface of natural fibers, and these functional groups can establish strong covalent bonds with the matrix, leading to strong interfacial adhesion between the fiber and matrix. Furthermore, plasma treatment increases the surface roughness and leads to better adhesion with the hosting matrix through mechanical interlocking [149,150]. The treatment by fungi can be employed to eliminate the non-cellulosic constituents such as wax from the surface of the fiber by the effect of specific enzymes. The fungi generate hyphane, which creates tiny holes on the surface of the fiber and generates a coarse interface for efficient interlocking with the matrix [16,84]. Ideally, this treatment method leads to a noticeable increase in the surface hydrophilicity due to the elimination of hemicellulose and lignin, leaving the cellulosic backbone exposed to the environment [151]. It is important to realize that the physical treatment methods such as hot water, enzyme, and electron beam radiation compromise the flexural properties of composites due to the over extraction of the fiber’s extractives. For example, enzymes severely dissolve the hemicellulose as an interfibrillar matrix, which leads to a great reduction of the fiber’s properties. However, chemical treatment techniques provide the treated fibers with more dimensional stability, higher resistivity toward fungal decay, and moisture absorption [124].

3. Type of Hosting Matrix

The efficiency of the physical and chemical treatments applied on natural fibers is largely affected by the structure and type of hosting matrix. Polymeric and cementitious matrices are the most frequently hosting mediums used in the fabrication of natural composites. In the subsequent sections, complete discussion about the moisture absorption behavior of these matrices and their composites is developed in Sections 3.1 and 3.2.

3.1. Polymeric Matrix

Thermoplastic resins have various advantages relative to thermoset polymers in the fabrication of biocomposites such as the ease of molding complex parts, low processing, and design flexibility. PP is the most familiar thermoplastic used in the fabrication of natural composites due to its good mechanical aspects, relative high temperature resistance, reasonable impact resistance, outstanding processibility, and low density [152]. Polyvinyl chloride, PE, PP, and polystyrene are frequently used as thermoplastic hosting matrices for natural fibers. Epoxy, VE, PES, and phenol formaldehyde are thermoset matrices, which can be embedded with natural fibers [153]. According to Figure 6, the type of polymeric resins and the duration of immersion of these resins in water play a key role
in controlling the equilibrium moisture content penetrated in the polymer matrix. It can be inferred from the same figure that epoxy and PP exhibit an optimum stable performance at both short and long immersion durations compared to other thermostet and thermoplastic resins, respectively. Additionally, the HDPE experiences a considerable change in moisture absorption behavior when the duration of water immersion is increased from 24 h to 48 h.

Figure 6. The equilibrium moisture content for most widely used polymers in natural composites. Epoxy (24 h) [78], epoxy (792 h) [76], epoxy (5616 h) [154], HDPE (24 h) [78], HDPE (48 h) [155], LDPE (24 h) [78], nylon 6 (24 h) [78], nylon 6,6 (24 h) [78], polybenzoxazine (800 h) [156], PBS (250 h) [157], PES (24 h) [78], PP (24 h) [78], PP (240 h) [132], polystyrene (24 h) [78], TPU (168 h) [158], UP (720 h) [85], UP (960 h) [88], and VE (24 h) [78].

The biopolymer matrix has a higher tendency to absorb moisture relative to its synthetic counterpart. The water molecules can penetrate into the most of organic polymers, leading to significant alteration in their mechanical, chemical, and thermophysical properties. The major consequence of the moisture absorption is on the polymer itself through saponification, plasticization, hydrolysis, and other degradation mechanisms, leading to both irreversible and reversible modifications in the structure of the polymer [138]. Jute fibers are used to reinforce bio and synthetic epoxy resins in order to investigate the moisture absorption effect on the properties of fully biodegradable composite. It is demonstrated that the type of the epoxy has a significant effect on the water uptake content of the composite. The investigations show that the fiber swelling and moisture absorption are higher in the bio-epoxy-based composite relative to the synthetic one. The higher swelling rate noticed for the fully green composite is justified, since the bio-epoxy contains a specific percentage of cellulose [159]. Generally, soy protein resins absorb a higher quantity of water, which mitigates their mechanical aspects. The moisture absorption characteristics of soy protein can be significantly improved through immersing the protein film in benzilic acid [60]. The compacted structure of the VE polymer justifies the lower coefficient of water diffusion, higher strength, chemical stability, and stiffness of its composites when they are immersed in water [160]. For example, the sisal fibers/VE composite displays 24% lower moisture absorption at equilibrium relative to the sisal/epoxy composite [161]. Thus, it can be concluded that the variences in moisture absorption behaviors among polymers are attributed to the degrees of crystallinility, hydrophobicity, and structure compactness of the hosting resins [162].

The moisture absorption properties of polymeric resins can be improved through increasing their stiffness, reinforcing them with synthetic fillers, and finally through blending them with proper resins characterized with low hydrophilicity [163]. For instance, the resistance of PLA toward weathering conditions, including moisture absorption, can be enhanced through reinforcing it with specific loadings of synthetic fibers [54,164,165]. However, the incorporation of natural fillers such as natural
rubber into a PLA/rice starch composite introduces moderate enhancements in tensile properties, but it improves the biodegradability of the PLA/rice starch composite when it is exposed to a humid environment [166]. Combining PP with polyamide-6 is applied commonly to reduce both the weight of the resulting composite and its moisture absorption tendency. PP is characterized by low cost, density, and high water uptake resistance, while the polyamide-6 is known for its high affinity for moisture and dimensional stability [167]. The micro-cracking of the matrix originating from freezing and fiber swelling rarely takes place when the hosting matrix is characterized with high ductility such as PP. However, the thermosetting resins such as UP have a high degree of brittleness, and hence cracks propagate quickly inside their structure [168].

3.2. Cementitious Matrix

The cumulative effect of hydroxyl groups, cell wall pores, and open lumen makes the natural fibers susceptible to water sorption, which induces the dimensional instability. The moisture absorption has a considerable effect on the mechanical strength of the natural fiber-reinforced cement composite [169,170]. The cyclic wetting and drying of natural fibers at temperatures exceeding 70 °C accelerates their degradation in a cement matrix more efficiently compared to static aggressive aging conditions. Compared to natural fibers immersed in distilled water, the moisture absorption content of fibers soaked in Ca(OH)₂ liquid is higher. The durability of cement-based composites in a humid operating environment has fundamental importance for applications in decoration, repair, and construction. The improvement of moisture resistance is attributed to the coupled changes of the matrix surface property and the microstructure of the cementitious composite. The degradation of natural fibers in a cement matrix passes through the following steps: (i) the exposure of holocellulose resulting from the degradation of lignin and the portion of hemicellulose; (ii) the reduction of the stability and integrity of cell walls originating from the degradation of hemicellulose; (iii) the dispersion of cellulose microfibrils emerging from the degradation of intramolecular hydrogen bonding; (iv) the entire degradation of cellulose microfibrils resulting from the alkaline hydrolysis of amorphous components. There are different techniques that can be used to reduce the degradation of natural fibers in a cement matrix, such as:

1. Managing the value of pH related to the pore solution at relatively low levels [171,172].
2. Combining pozzolanic additives, namely metakaolin and silica fume, with natural fiber-based cement composites [173].
3. Incorporating nano reservoir silts within the binders of the composite [174].
4. Applying a polymeric coating on the surface of the natural fiber. The raw natural fiber is incapable of developing a compact interface of the fiber–cement composite due to the hydrophilic nature of the fiber. The polymer molecules diffuse into the fibers, particularly between the filling lumens and fiber cells, reducing the moisture absorption. Consequently, the application of a polymer coating decreases the concentration of OH and enhances the strength of the natural fiber, which improve its interfacial integrity and bonding strength [175,176].
5. Reinforcing cement with mineral fillers such as basalt fibers, which can be classified as non-hazardous and environmentally friendly substances [177].

However, the effect of the hosting matrix type should be addressed with the type of natural fiber, which is used to improve its structural performance, in order to determine the degree of compatibility at the interfacial region between the fiber and matrix. More information about the effect of reinforcing fiber is discussed in detail through Section 4.

4. Type of Natural Fibers

The natural fibers have varying moisture absorption behavior, since they have different interfacial bond strengths in addition to various morphologies and chemical compositions in terms of lignin and hemicellulose loadings [75]. The chemical composition and structure of jute fibers make them
different from other natural fibers, since they absorb moisture, and its moisture regain characteristic is very high. For example, jute fibers absorb a lower amount of moisture relative to banana fibers at the same immersion period [178]. Additionally, due to the porous configuration of the ground tissues, bamboo fibers absorb a higher amount of moisture relative to more common bast fibers such as hemp, kenaf, and flax fibers [179,180]. The OSF-reinforced PP composite shows lower moisture absorption relative to the wood-based PP composite. This behavior can be attributed to the aspect ratio and morphology of OSF, which are quite different from wood fillers, even though both fillers have the same chemical compositions [126]. The type of natural fiber has a more significant effect on water uptake behavior compared to the type of chemical and physical treatment conducted on the natural fiber [181]. Biofibers such as cotton burr and stem, guayule bagasse, and oak fibers are used to reinforce HDPE resin modified with two types of coupling agents in order to prepare biofiber-based composites. The water uptake test shows that the coupling agent has a minimal effect on the moisture absorption of the composite. The guayule bagasse fibers have a great influence on decreasing the moisture absorption tendency of the composite [182]. The effect of the natural fiber types on the efficiency of the moisture resistance treatments can be fully realized through exploring the percentage of improvements on moisture absorption behavior, as shown in Figure 7.

\[ \text{Moisture Improvement (\%)} \]

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Moisture Improvement (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coconut (plasma)</td>
<td>-156.41%</td>
</tr>
<tr>
<td>Coir (plasma)</td>
<td>-1.06%</td>
</tr>
<tr>
<td>Hemp (17.5 wt. % NaOH)</td>
<td>18.71%</td>
</tr>
<tr>
<td>Hemp (0.7 wt. % NaClO₂)</td>
<td>50.00%</td>
</tr>
<tr>
<td>Jute (acetlylation)</td>
<td>65.00%</td>
</tr>
<tr>
<td>Pine (acetlylation)</td>
<td>45.64%</td>
</tr>
<tr>
<td>OPEFB (5% NaOH)</td>
<td>42.74%</td>
</tr>
<tr>
<td>OPEFB (10% Latex)</td>
<td>42.74%</td>
</tr>
<tr>
<td>OPEFB (TEVS)</td>
<td>43.98%</td>
</tr>
<tr>
<td>OPEFB (acetlylation)</td>
<td>44.40%</td>
</tr>
<tr>
<td>OPEFB (Peroxide)</td>
<td>44.40%</td>
</tr>
</tbody>
</table>

Figure 7. The improvement percentage of different treatments on moisture absorption behavior of natural fibers. Coconut plasma [147], coir (plasma) [183], hemp (17.5 wt. % NaOH) [184], hemp (0.7 wt. % NaClO₂) [184], jute (acetlylation) [96], pine (acetlylation) [96], OPEFB (5% NaOH) [70], OPEFB (10% Latex) [70], OPEFB (TEVS) [70], OPEFB (acetlylation) [70], and OPEFB (Peroxide) [70].

It can be inferred from the values clarified in the figure above that the samples of OPEFB fiber, which are treated with five different types of treatments, exhibit approximately the same percentage of improvement on moisture absorption behavior. However, treating coconut and coir fibers with the same moisture treatment technique (plasma method) leads to great variation between these two fibers in terms of moisture absorption resistance’s properties. This behavior implies that applying plasma treatment on different types of natural fibers having various chemical compositions could introduce adverse impact on the moisture absorption behavior of the fiber under treatment if the moisture treatment technique excessively eliminates non-cellulosic components from the fiber structure such as lignin, which consequently induces the natural fiber to absorb higher quantities of moisture from the surrounding environment. In next sections, the most commonly used natural fibers are discussed in terms of their effects on mitigating the moisture quantities absorbed by chemically/physically treated composites.
4.1. Cellulose Fiber Composites

The properties of pure PLA resin can be easily deteriorated at high temperature in the existence of residual moisture or water. Therefore, cellulose fibers are commonly used to reinforce PLA resin composites in order to increase the resistance of PLA resin under hygrothermal conditions. The cellulose fiber/PLA composites show a poor adhesion at the cellulose fiber–PLA interface, but their structures exhibit well-dispersed morphology. To improve the water uptake behavior and mechanical properties of cellulose fiber composites, the surfaces of the fibers are coated with suitable coatings, chemically/physically treated, or hybridized with secondary nanofillers [185]. For example, the silane coupling agent is used to improve the mechanical and physical properties of the cellulose nanofibers reinforced bio-epoxy composite. The treated composite exhibits the lowest content of voids relative to the raw composite due to the effect of silane treatment on improving the wettability of cellulose nanofiber in the bio-epoxy resin [186]. Cellulose fibers are used to reinforce the water-based resin composite. The implementation of water-based resin in fabricating the composite is favorable due to its similarity with cellulose. The moisture absorption of the resulting composite is 100 folds lower than that of the neat cellulose sheet [187].

4.2. Wood Plastic Composites

Biopolymers are considered suitable materials to be implemented in manufacturing WPCs, which have the potential to substitute engineered wood and petroleum-based plastics in building construction applications. Nevertheless, the dissemination of WPCs in construction applications is still limited due to the concerns related to its long-term durability, especially in high wet and humidity environments. It was highlighted in the literature that the type of hosting polymer matrix has a considerable effect on the mechanical performance of WPCs. For example, PP-based WPCs have lower resilience properties relative to PHBV-based ones. Furthermore, the performance of chemically treated PP-based WPCs are lower than that of raw PHBV-based WPC reinforced with 20% of wood powder [188]. The moisture absorption and swelling of WPCs are largely affected by the source of the wood fiber, size of the wood particle, surface treatment of the wood, orientation of the wood fiber, conditioning temperature, exposed surface area of wood fillers, and increased proportionally with the increasing content of these fillers. Composites reinforced with large wood particles exhibit high thickness swelling and water diffusivity. The molecules of water can diffuse in WPCs through the interface between the fiber and the matrix, fiber lumen, and fiber cell [189,190]. The penetration of water molecules into the structure of WPCs decreases the mechanical properties of the composite considerably. Wood powder composed of hard particles can effectively increase the stiffness of WPCs. However, when these particles absorb moisture, they become ductile, and hence the stiffness of the composite is reduced. When the WPC is redried, the wood particles are able to regain their hardness, and can contribute efficiently in composite stiffness. However, exposing WPCs to a humid environment leads to the swelling of the wood powder in the hosting resin, leading to the initiation of micro-cracks in the composite structure, which prevents the composite from restoring its original stiffness [191–194]. The fillers of wood powder are qualified candidates to reinforce thermoplastic polymers such as PP, which improves the biodegradability of these polymers. For example, the sawdust–PP composite has a higher degree of biodegradability over neat PP resin due to its accelerated rate of degradation, resulting from hydrolysis and microbial attack [195].

Various techniques have been investigated in the literature in order to improve the resistance of wood fillers toward moisture absorption, including the application of superhydrophobic coatings and using hot water extraction. It is found out that the application of coatings to wood powder has a trivial effect on the moisture absorption behavior [196]. On the other hand, liquid hot water can be used to extract hemicellulose from wood, which improves the moisture absorption resistance of WPC. This improvement can be attributed to the increase in compatibility between the thermoplastic matrix and reinforcing wood fillers [197]. Using coupling agents to treat WPC reduces the moisture absorption, thickness swelling, and moisture diffusion in the composite’s structure. The optimum loading of the
coupling agent can be determined based on the content and size of the wood [190,198,199]. Following better processing methods during the manufacturing of WPCs is another route that can be followed to improve the moisture absorption behavior. For example, the water uptake behavior of injection molded PP-based WPC can be effectively improved through increasing the screw speed and residence time during the manufacturing process, which produces small-sized wood particles, leading to a noticeable reduction in the rate of moisture absorption in the range of 10% up to 40% without any change in the mechanical properties [200,201]. Moreover, increasing the content of the polymer in WPC improves the weatherability performance of the composite, such as the suppression of surface cracks resulting from various weathering conditions [202].

4.3. Flax Fiber Composites

The anisotropy and morphology of flax fibers have a considerable effect on the diffusivity of water. The diffusivity of water across and along the direction of the fibers is considerably higher relative to the diffusivity across the thickness direction [203]. The influence of the moisture absorption during the manufacturing process of flax fiber/UP composite on the interfacial adhesion strength could be limited [204]. The interaction between flax fibers and the PP hosting matrix depends on van der Waals interactions that are sensitive to water. The adhesive pressure originating from residual stress could be released through the effect of moisture plasticizing [205]. Structural damages can be easily initiated in flax fiber-based composites exposed to moisture aging conditions. These damages are motivated by fiber swelling, fiber degradation, the washing out of soluble substances—specifically the cement of the fiber bundle, the poor interfacial strength between the fiber and matrix, and the plasticization of the hosting matrix [5,206–209]. In order to develop economically feasible flax fibers composites characterized with better mechanical and environmental performance, it is recommended to improve the hydrophobic behavior through treating them with appropriate chemical and/or physical treatments [152]. Duralin, peroxide, benzoylation, and the grafting of MA are the most effective treatment methods that can be used to improve the water uptake behavior of flax fibers composites, as demonstrated by the improvement values shown in Figure 8.

The moisture absorption and diffusivity of flax fiber-based composites can be considerably reduced through hybridizing flax fibers with synthetic fibers such as glass fibers. Nevertheless, the hybridization of flax fibers has an adverse effect on tensile strength when the hybrid composite is exposed to a hygrothermal environment characterized with high temperatures and humidity [210]. The thermal treatment of flax fibers can effectively mitigate the swelling of flax fibers and their sensitivity to moisture absorption through changing the chemistry of the fiber without considerably altering the morphology of elementary fibers [211]. The hosting resin contributes effectively to reducing the moisture absorption potential of flax fibers. For example, the PLA matrix decreases the water uptake behavior of flax/PLA composites by around 40% relative to a neat flax matrix [212]. Twill flax fiber-reinforced thermoplastic resin (acrylic) composites absorb a lower amount of moisture relative to flax fiber-reinforced thermoset (epoxy) composites. This behavior can be attributed to the higher porosity loading of the flax fiber-reinforced epoxy composite relative to the flax fiber-reinforced acrylic [213]. Two flax fiber-based composites of flax/polyurethane and flax/bio-epoxy have been compared in terms of their moisture absorption characteristics. It is found out that polyurethane absorbs a lower quantity of water and at a slower rate than bio-epoxy resin.
During the manufacturing process of flax fiber/UP composite on the interfacial adhesion strength, the thermal treatment of flax fibers characterized with high temperatures and humidity is found to improve the hydrophobic behavior through treating them with appropriate substances. Interfacial adhesion in the hemp/epoxy composite is increased with increasing fiber loading and when the diameter of the hemp fiber is reduced. This behavior can be attributed to the improved fiber surface area at lower diameters, leading to a great reduction of the tensile and flexural properties of the composite. The packing structure mitigates the porosity inside the composite structure, which participates in reducing the moisture absorption.

The chemical bonding between urethane pre-polymer and hydroxyl groups of cellulosic fibers reduces the number of available hydroxyl groups. Therefore, it is recommended to use polyurethane resin as the hosting matrix for most cellulosic fibers in order to attain good durability, moisture resistance, and environmental stability properties [217].

### 4.4. Hemp Fiber Composites

The packing arrangement and chemical treatment such as the weave patterns of hemp fibers have a considerable effect on the moisture absorption and porosity of the hemp fiber-based composite. The most effective treatment methods for hemp-based composites are demonstrated by the improvement values shown in Figure 8. Improving the moisture absorption behavior of flax fiber-based composites. 38 vol.% flax fiber/PP (1440 h) [214], 10 wt. % flax fibers/HDPE/LDPE (24 h) [215], dew-retted (RFB) flax/PP and Duralin upgraded flax/PP (RH = 100% for 4500 h) [146], flax fiber/PLA (408 h) [216], 60 wt. % short flax fiber bundle/5 wt. % MAPP (E43) (5040 h) [135].

The chemical bonding between urethane pre-polymer and hydroxyl groups of cellulosic fibers reduces the number of available hydroxyl groups. Therefore, it is recommended to use polyurethane resin as the hosting matrix for most cellulosic fibers in order to attain good durability, moisture resistance, and environmental stability properties [217].
bond strength with fully synthetic epoxy resin, while they have a good interfacial bond strength with green epoxy resin [228].

5. The Effect of Fiber Loading

The water uptake, diffusivity, and the resulting thickness swelling are increased with the increasing content of fibers in natural composites, which leads to an acceleration of the biodegradation rate of the composite [229–231]. Composite structures with higher thicknesses have a higher capability to absorb moisture relative to thinner ones. Consequently, an optimum fiber loading should be selected to attain balance in all aspects of composites, including mechanical properties [232]. Referring to Figure 9, the fiber content along with water immersion time play a key role in controlling the quantity of moisture absorbed by natural composites. For instance, epoxy composites reinforced with around 20 wt. % and 46 wt. % of recycled cellulose fiber that is immersed in distilled water for 5616 h exhibit the highest amount of moisture absorption. This behavior is attributed to the two main factors of the high cellulosic content in natural composite, and the long immersing duration of the composite in distilled water.

![Figure 9](image-url)

Figure 9. The effect of fiber loading and immersion time (shown in the above data points) on the percentage of moisture absorbed by natural composites. Banana fiber/polyester [74], CF/geopolymer [28], Lantana camara fiber/epoxy [68], Padauk sawdust wood/biodegradable PBS [157], Pineapple fiber/HDPE [155], RCF/epoxy [154], Sisal/PES [232], SPF/epoxy [76], SPF/TPU [158], Unshredded waste newspaper/PES [233].

This conclusion can be further verified through investigating the moisture absorption behavior of cotton fabric-reinforced geopolymer composites. The long immersing duration of the composite in distilled water (3192 h) induces the composite to absorb around 18% of the moisture from the conditioning medium, even though the composite is reinforced with less than 8 wt. % of cotton fabric. Reinforcing natural composites with higher loadings of natural fibers is a desirable optimization objective, since the natural fibers improve the overall mechanical properties of natural composites. A trade-off can be achieved between the loading of natural fibers and the moisture absorption behavior of natural composites through hybridizing the composite structure with other natural and/or synthetic fibers characterized by a lower tendency to absorb moisture. More details about the hybridization of natural composites will be provided in Section 6.

6. Hybridization of Natural Composites

Based on the non-renewable/renewable source of resin/fiber, the polymeric composites can be divided into three main types: (i) synthetic polymeric composites; (ii) entire renewable (green) polymeric composites; (iii) partially renewable composites [15]. The hybrid composite can be fabricated
through the combination of two types of different fibers, which are used to reinforce the common hosting matrix. The hybridization of the natural composite is considered to be an effective method to reduce the moisture absorption [234]. The effect of hybridization on the moisture absorption behavior for both entire and partially natural hybrid composites can be better realized through exploring the moisture improvement values listed in Table 3, which are calculated relative to unhybridized composites. It can be noted that the hybridization of natural composites has an adverse impact on the coefficient of water diffusion due to the increasing load of voids in the composite structure and poor interfacial adhesion between the laminates of hybrid composites.

### Table 3. The effect of hybridizing natural fibers on moisture absorption properties.

<table>
<thead>
<tr>
<th>References</th>
<th>Hybrid Composite</th>
<th>Moisture Improvement %</th>
<th>Diffusivity Improvement %</th>
<th>Immersion Time (Hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>[235]</td>
<td>Sisal fibers hybridized banana at (50:50) reinforced epoxy.</td>
<td>11.48</td>
<td>-12.13</td>
<td>50</td>
</tr>
<tr>
<td>[87]</td>
<td>7 wt. % Glass hybridized 13 wt. % coir-reinforced polyester.</td>
<td>39.16</td>
<td>-</td>
<td>1440</td>
</tr>
<tr>
<td>[236]</td>
<td>15% Glass fibers hybridized 15% jute-reinforced 70% UP.</td>
<td>58.36</td>
<td>-309.79</td>
<td>504</td>
</tr>
<tr>
<td>[237]</td>
<td>5 wt. % Glass hybridized 35 wt. % hemp-reinforced 55 wt. % PP.</td>
<td>6.76</td>
<td>-</td>
<td>3600</td>
</tr>
<tr>
<td></td>
<td>10 wt. % Glass hybridized 30 wt. % hemp-reinforced 55 wt. % PP.</td>
<td>21.31</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td></td>
<td>15 wt. % Glass hybridized 25 wt. % hemp-reinforced 55 wt. % PP.</td>
<td>37.11</td>
<td>-</td>
<td></td>
</tr>
</tbody>
</table>

In this section, the entire and partially natural hybrid composites are discussed with further details through Sections 6.1 and 6.2, respectively.

### 6.1. Entire Natural Hybrid Composites

The hybridization improves the dynamic mechanical properties, including the loss and storage moduli, and reduces the moisture absorption capabilities of natural composite [238–240]. The fully natural hybrid composites find different applications in the construction and industrial sectors. Green biocomposites are developed through gluing hemp shiv with wheat straw. The developed composite exhibits hygric and thermal performances which contribute to mitigate the energy demands of building and guarantee the hygrothermal comfort of users [241,242]. Using natural fibers characterized with relatively high impedance against moisture absorption is one of the options to improve the water uptake behavior of natural fiber composites. Mineral fibers such as basalt fibers are frequently used to hybridize natural composites in order to improve their moisture resistance and durability, in addition to their mechanical properties. Usually, basalt fiber-based composites are used to fabricate the skin laminates of the hybrid sandwich composites in order to protect the natural fibers with lower moisture resistance, such as flax and hemp fibers, from absorbing the moisture found in the surrounding environment [243–246]. The basalt fibers enhance the interfacial adhesion, mitigating the moisture absorption, and consequently postponing the structural degradation of basalt/flax hybrid laminated composites. The moisture absorption changes the failure mechanism of flax–basalt hybrid composites, as demonstrated in Figure 10, due to the plasticization phenomena. Furthermore, the moisture absorption induces the delamination between basalt layers and interior flax-based layers [247]. Structural flaws such as delamination, matrix cracking, and voids increase the capability of hybrid composites to absorb moisture and mitigate the strength of the moisture-saturated composite due to the degradation of the interface between the matrix and fiber [248]. The most critical factor that promotes the structural failure in hybrid composites is the content of porosity and air voids. It is recommended to dry the natural fibers through the fabrication process before resin injection in order to attain fibers characterized with good wetting properties as well as prevent any generation of gas bubbles during curing [249,250].
A superior moisture absorption improving effect can be attained through using chemical/physical treatments associated with hybridization. Poplar wood flour and recycled newspaper fiber are combined in the recycled PP matrix modified with MAPP. This composite includes a higher fraction of recycled newspaper fiber, which shows maximum moisture absorption due to the presence of a high quantity of holocellulose. The incorporation of MAPP has an advantageous influence on both the thickness swelling and moisture absorption through enhancing the quality of adhesion between the fibers and polymer [251].

6.2. Partially Natural Hybrid Composites

Hybridizing natural fibers with synthetic ones such as glass or carbon fibers is an efficient technique that is used for controlling the vibration damping and enhancing the mechanical and moisture absorption properties of partially natural hybrid composites [252]. The incorporation of natural fibers such as jute and flax fibers to carbon and glass fiber-based composites improves the flexural properties, ductility, and impact strength of the hybrid composites, while the carbon and glass fibers participate in enhancing the moisture absorption behavior, stiffness, strength, and thermal stability of the hybrid composites [253–257]. Glass fiber is considered an economically feasible fiber characterized with high tensile strength, low moisture absorption capability, and high dimensional stability. Glass fiber-based composites have a good capability to resist various weathering conditions, including exposure to moisture and heat aging conditions [209,258–262]. For instance, the amounts of moisture absorption for the kenaf/VE composite are 97.95% and 98.35% higher than that of the glass/VE composite under tensile and flexural loading conditions, respectively. Likewise, the saturated weight gain of flax fiber composites is 12-fold higher than that of glass fiber composites. Additionally, the incorporation of two external glass fiber layers considerably improves the moisture absorption of the glass/flax hybrid composite by more than 180% compared to the flax/epoxy composite. Therefore, the increasing content of synthetic hydrophobic fiber in hybrid natural composite mitigates the content and absorption speed of moisture [258,263–271]. Glass fibers can be used to improve the moisture absorption properties of natural sandwich composite structures [272]. Skin layers of woven jute fabric are used to cover the particleboards’ core composite, which is composed of a mixture of short glass fibers and wood particles. The hybrid composite shows good dimensional stability and outstanding moisture resistance relative to commercially available wood-based composites [273].

Using a coupling agent such as MAPP adds further improvement to the tensile strength and moisture absorption behavior of hybrid composites [274]. The hybrid composite of glass/bamboo
fibers reinforced with PP and modified with MAPP is immersed in water for 1200 h at 25 °C. Due to the hybridization with glass fibers, a reduction in moisture absorption of 4% relative to unhybridized composite has been recorded [275]. The synthetic fibers can be organized in a randomly close-packed structure in order to prevent the water molecules from diffusing inside the composite structure, and hence hinder them from contacting and binding with the surfaces of hydrophilic natural fibers [87]. When the water molecules diffuse into a hybrid composite, composite failure takes place due to the jute fiber swelling, which creates swelling stresses leading to the failure of the composite. The moisture absorption weakens the interfacial bonding strength between the fiber and matrix, reduces the stiffness of the natural fiber, leads to the stress corrosion of synthetic fibers, and plasticizes the polymeric matrix [276]. For example, the water penetration into woven glass/long kenaf degrades the fracture toughness due to the weakening of the interfacial bond strength between the fiber and polymer [277]. A glass and twill flax fiber-reinforced epoxy hybrid composite is immersed in water at 55 °C. Delamination between the glass and flax fiber-based layers is initiated due the moisture absorption. This leads to a reduction in the maximum tensile strength of the moisture saturated composite [278]. On the contrary, it is demonstrated that in some cases, the molecules of water that penetrate into the cracks and cavities within the composite structure increase the flexibility of the hybrid composite [279].

7. Incorporation of Nanofillers

Nanotechnology can be implemented to improve the moisture absorption properties of natural fibers composites through using nanofillers and nanotechnology-based coatings. The particle forms of some types of natural fibers can add considerable improvement to the water uptake behavior of natural composites relative to other forms of mats and short fibers [280–282]. Polyethyleneimine-reduced graphene oxide fillers can be used to improve the tortuous diffusion pathway for water molecules and the surface hydrophobicity of methyl cellulose composite films, which can be used in water vapor barrier application [283]. The nanofillers such as fly ash particles function as additive and reinforcing fillers, filling the air gaps and voids that exist in the structure of sustainable composites [284]. The incorporation of metal oxide particles into the structure of natural composites improves their moisture absorption and swelling properties. This behavior is attributed to the good capability of hydrophobic metal oxides to bond with hydrophobic polymers [285]. Reinforcing kenaf fibers composites with magnetite particles offers dual-functional properties such as paramagnetic and electromagnetic shielding functions in addition to moisture absorption resistance [286]. TiO$_2$ fillers can be used to enhance the water uptake behavior for natural and synthetic fiber-based composites. Incorporating 0.1 wt. % of TiO$_2$ filler into glass fiber-reinforced polymer composite decreases the diffusivity by 9% [287]. Furthermore, 18% of water sorption reduction can be achieved upon the grafting of TiO$_2$ particles on the surface of flax fibers that are used to embed PLA composites. An oxidization treatment is carried out on the surface of flax fiber in order to enhance the interfacial adhesion of TiO$_2$ film to flax fibers, as shown in Figure 11. This enhanced adhesion prevents the penetration of water molecules and propagation of cracks in the interphase region [288, 289]. Likewise, the incorporation of SiC nanoparticles into cellulose fiber-reinforced epoxy strengthens the interfacial adhesion between the fiber and matrix [48]. Carbon nanotubes (CNTs) are used to lower the moisture absorption tendency of wood filler-reinforced HDPE. With the increasing content of CNTs, the equilibrium time (the time needed to attain the equilibrium thickness swelling) and thickness swelling are reduced. This reduction is attributed to the barrier properties of CNTs, which hinder the water diffusion into the composite structure. CNTs can inhibit the moisture penetration into the composite through two main mechanisms: (i) CNTs hinder the diffusion of water into deeper regions of the composite through filling the air gaps and voids; (ii) the surface of CNTs tends to immobilize the amount of the moisture due to its hydrophobic nature [290, 291]. The effect of natural nanofillers such as cellulose and clay reinforcements on the moisture absorption behavior of natural composites is further investigated through Sections 7.1 and 7.2, respectively.
Nanotechnology can be implemented to improve the moisture absorption properties of synthetic fiber-based composites. Small amounts of silane coupling agents are usually needed to treat nanoclay in order to produce nanocomposites characterized with higher resistance to hygrothermal aging conditions, along with the better retention of flexural and tensile strength properties. The enhancing effect of silane coupling agents can be attributed to the influence of salinization on exfoliation, as well as creating covalent bonds with the hosting matrix [296]. A glass fiber-reinforced VE composite has been modified with 1.0 wt. % of raw and (3-Aminopropyl)triethoxysilane (APTES)-treated montmorillonite (MMT) nanoclay. When the nanoclay comes into contact with moisture, swelling among layers takes place, which as a result changes the mechanical properties of the resulting composite. A composite reinforced with 7.1. Cellulosic Reinforcements

CNCs and cellulose nanofibers are natural particles characterized by a high surface area and high strength. The cellulose nanofillers can be extracted from natural fibers such as sisal fibers without considerable quantities of lignin and hemicellulose. The implementation of CNCs is being investigated for different uses, since it is stiffer than aluminum and stronger than steel. CNCs may be used to manufacture multi-scale composites with advanced mechanical performance, long service life, and good durability at different weathering conditions. Hence, crystalline and viscous CNCs can be implemented to manufacture high performance cement nanocomposites that have an increased tortuosity, which hinders the diffusion of water into the composite structure [153,169,292]. CNCs can be used to mitigate the water sensitivity of starch/poly vinyl alcohol composite films. This moisture stability originates from the creation of strong hydrogen bonds among the hydroxyl groups from constituents of the film through a three-dimensional network. With an increasing loading of starch, the water sorption of the film is reduced further [293]. The moisture behavior and tensile properties of starch-based composites can be improved through incorporating bacterial cellulose (BC) nanofibers in the composite structure. Both BC and starch are hydrophilic in nature, which could render the high moisture absorption of the composite. The higher resistance toward moisture absorption for BC/starch composites relative to a pure starch matrix is attributed to the strong hydrogen bonding created at the matrix–fiber interface and to the higher resistance of BC nanofibers to weathering conditions [294].

7.2. Clay Nanoparticles

Nanofillers such as nano clay and nano silica carbide can be incorporated into natural fiber-based composites in order to improve their water uptake behavior through improving their mechanical properties in wet conditions. The clay nanolayer forms an impermeable medium to hinder the flow of water and force the water to flow through an indirect path, and consequently, a longer time is needed for water diffusion [295]. In addition to natural composites, clay nanoparticles can be used to improve the moisture absorption properties of synthetic fiber-based composites. Small amounts of silane coupling agents are usually needed to treat nanoclay in order to produce nanocomposites characterized with higher resistance to hygrothermal aging conditions, along with the better retention of flexural and tensile strength properties. The enhancing effect of silane coupling agents can be attributed to the influence of salinization on exfoliation, as well as creating covalent bonds with the hosting matrix [296]. A glass fiber-reinforced VE composite has been modified with 1.0 wt. % of raw and (3-Aminopropyl)triethoxysilane (APTES)-treated montmorillonite (MMT) nanoclay. When the nanoclay comes into contact with moisture, swelling among layers takes place, which as a result changes the mechanical properties of the resulting composite. A composite reinforced with
silane-treated MMT exhibits better mechanical properties after immersion in sea water due to the improved interfacial adhesion at the MMT–VE interface [297]. Nanoclays alleviate the reduction that occurs in the mechanical properties of carbon fiber-reinforced polymers after exposure to sea water, and improve the impedance of the composites toward moisture absorption. A composite in which 2.0% nanoclays are dispersed in its matrix absorbed 0.39% of sea water, while the control composite absorbed 0.67% of water. This improvement can be justified to the outstanding barrier capacity along with the high aspect ratio and surface area of clay nanofillers [298]. Organoclay is another type of clay nanofillers that can significantly improve the water uptake behavior of natural composites. The effectiveness of organoclays on moisture absorption reduction depends on many factors, such as the polarity of organic moieties and the morphology of nanoclays in the hosting resin [296]. Organoclays (ammonia salts) are used to fabricate poly vinyl alcohol/functional clay aerogel composites. Depending on the RH, a five-fold reduction can be achieved for composites reinforced with organoclay relative to those reinforced with traditional clays. The main outcome of the aforementioned study shows that the moisture sorption capacity is mitigated by up to 40% after 30 days of aging under a RH of 85% due to the increase in the interlayer space, which forms a more tortuous path for heat to pass through [299].

8. Conclusions

This review demonstrates through previously published experimental data that treating natural composites with chemical and/or physical treatments alone does not provide the treated composites with adequate resistance toward moisture absorption. Actually, there are a number of parameters that govern the efficiency of the chemical or physical treatment method on improving the moisture absorption behavior of natural composites, including: the proper selection of the hosting matrix and natural fiber, adjusting the fiber loading and duration of the moisture exposure with applied moisture treatment, the hybridization with hydrophobic and minerals fibers, and lastly, the implementation of natural and synthetic nanofillers. A complete comparison between the aforementioned parameters in terms of their capability on improving the moisture absorption behavior of natural composites is discussed in Table 4. Choosing an appropriate type of natural fiber and hosting matrix considerably affects the efficiency of the treatment applied on natural composites. The compatibility between hosting resin and reinforcing fiber is a very critical parameter that determines the degree of bond strength at the fiber–matrix interface. The loading of the natural fiber and the aging time are directly correlated to each other. Increasing the fiber content and aging duration induces the natural composites to absorb higher quantities of moisture from the conditioning medium, regardless of the applied chemical and/or physical treatments. The hybridization of natural fibers with synthetic fibers reduces the amount of moisture treatments that are needed, which can be specified through the loading ratio of synthetic to natural fibers. For example, if the hybridization is done at a 50:50 ratio, the amount of treatment that is needed is decreased by 50%. This can be attributed to the capability of hybridizing fibers to form excellent interfacial adhesion with the synthetic hosting resin, which prevents the water molecules from passing to the interior hydrophilic natural fibers. It is noteworthy to highlight the adverse impact of hybridizing natural composites on reducing the efficiency of moisture absorption treatments, since the hybridization increases the coefficient of water diffusion inside the composite structure. This can be attributed to the voids and gaps that exist in the structure of the hybrid composite in addition to the poor interfacial adhesion between the laminates of different reinforcing fibers. The role of secondary nanofillers is magical when these fillers are used along with chemical/physical treatments, since these fillers perform multiple tasks at the interface, including: (a) strengthening the mechanical interlocking through increasing the surface roughness of the natural fibers, (b) alleviating the damaging effect of delamination emerged from moisture absorption through the pinning mechanism, (c) improving the overall barrier properties of natural composites through offering additional tortuous paths to composite structure that decelerate the penetration of water molecules, and (d) eliminating the air gaps and voids generated due to the poor compatibility between the hydrophilic natural fiber and hydrophobic hosting resin through filling them.
Table 4. Comparison between moisture treatment parameters of natural composites.

<table>
<thead>
<tr>
<th>No.</th>
<th>Treatment Parameter</th>
<th>Moisture Resistance</th>
<th>Moisture Diffusivity</th>
<th>Interfacial Adhesion</th>
<th>Efficiency of Treatment</th>
<th>Void Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Type of hosting matrix</td>
<td>Hydrophobic synthetic resins provide protection for the</td>
<td>Heavy chemical/physical treatments are not needed,</td>
<td>Interfacial adhesion is weak due to poor</td>
<td>The incorporation of MA in the</td>
<td>The thermoset resins such as epoxy contain a specific loading of voids,</td>
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<tr>
<td></td>
<td></td>
<td>hydrophilic natural fiber regardless of fiber type.</td>
<td>since the hydrophobic polymers deaccelerates the</td>
<td>covalent bonding between the hydrophobic</td>
<td>hydrophobic resin increases the</td>
<td>which facilitates the penetration of water molecules. Thermoplastic</td>
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<td></td>
<td></td>
<td>Consequently, choosing an appropriate type of resin is</td>
<td>diffusion of water molecules.</td>
<td>and hydrophilic components.</td>
<td>capability of most polymers to</td>
<td>polymers such as PP are highly recommended for natural composite applications.</td>
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<td></td>
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<td>more important relative to the type of natural fiber.</td>
<td></td>
<td></td>
<td>resist moisture.</td>
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</tr>
<tr>
<td>2.</td>
<td>Type of natural fiber</td>
<td>It is recommended to use natural fibers with a higher</td>
<td>The packing arrangement of natural fibers play a key role</td>
<td>The type of natural fiber has the highest</td>
<td>The type of natural fiber</td>
<td>The moisture treatments increase the surface roughness of natural fibers,</td>
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<tr>
<td></td>
<td></td>
<td>degree of compatibility with hydrophobic resins in order</td>
<td>in reducing the water diffusivity.</td>
<td>impact on the interfacial adhesion’s bond</td>
<td>controls the success of the</td>
<td>which improves the fiber wetting in the hosting resin, leading to a</td>
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<td></td>
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<td>to improve the moisture absorption properties. The</td>
<td>The woven arrangement of natural fibers is recommended.</td>
<td>strength. Natural fibers with higher</td>
<td>chemical/physical treatment</td>
<td>considerable reduction in the void content at the fiber-matrix</td>
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<td></td>
<td></td>
<td>overtreatment of natural fibers should be avoided.</td>
<td></td>
<td>surface areas and aspect ratios induce</td>
<td>process. treating specific</td>
<td>interphase region.</td>
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<tr>
<td>3.</td>
<td>Fiber loading</td>
<td>The fiber loading reduces the moisture absorption</td>
<td>Applying a polymeric coating on natural composites that</td>
<td>The efficiency of moisture treatments is</td>
<td>With increasing fiber content,</td>
<td></td>
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<tr>
<td></td>
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<td>behavior of the natural composite. With the increasing</td>
<td>is reinforced with higher loadings of natural fiber is an</td>
<td>limited at higher contents of natural fibers,</td>
<td></td>
<td>The void content is greatly reduced with the increased loading of</td>
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<td></td>
<td></td>
<td>content of cellulose in the natural composite, the</td>
<td>essential requirement for reducing the moisture</td>
<td>and using another parameter for moisture</td>
<td></td>
<td>synthetic/mineral fibers in the composite structure due to the good</td>
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<td></td>
<td>application of efficient chemical/physical treatments</td>
<td>diffusion due to the increasing content of air voids at</td>
<td>behavior improvement such as hybridizing</td>
<td></td>
<td>wetting capability of these fibers with the hosting matrix.</td>
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<td></td>
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<td>is recommended.</td>
<td>the fiber-matrix interface.</td>
<td>composites with synthetic/mineral fibers is</td>
<td></td>
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<tr>
<td>4.</td>
<td>Hybridization of natural</td>
<td>The hybridization alleviates the need for chemical/</td>
<td>The diffusivity can be improved with minimal chemical</td>
<td>The hybridization enhances the efficiency</td>
<td>The void content is greatly</td>
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<td></td>
<td>fiber</td>
<td>physical treatments if proper synthetic or mineral</td>
<td>physical treating efforts if the external layers of</td>
<td>of moisture treatments applied on natural</td>
<td>reduced with the increased</td>
<td>reduced with the increased</td>
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<td>fibers are used to cover the external layers of</td>
<td>synthetic/mineral fibers prevent the water molecules</td>
<td>composites, since the layers of synthetic/</td>
<td>loading of synthetic/mineral</td>
<td>loading of synthetic/mineral</td>
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<td></td>
<td></td>
<td>natural composites. The moisture absorption resistance</td>
<td>from penetrating into the interior composite’s core.</td>
<td>mineral fibers do not need additional</td>
<td>fibers in the composite</td>
<td>fibers in the composite structure due to the good wetting capability of these</td>
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<td>is improved with the increasing content of synthetic/</td>
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<td>treatments.</td>
<td>matrix.</td>
<td>fibers with the hosting matrix.</td>
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<td></td>
<td></td>
<td>mineral fibers.</td>
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<tr>
<td>5.</td>
<td>Implementation of nanofillers</td>
<td>Nanofillers improve the moisture barrier properties of</td>
<td>Nanoreinforcements can effectively reduce the water</td>
<td>The improvement attained on interfacial</td>
<td>The hybridization enhances the</td>
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<td>composites. It should be considered as a second option</td>
<td>diffusivity through the additional tortuous paths that</td>
<td>adhesion is proportional to the loading of</td>
<td>efficiency of moisture</td>
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<td>in hand after choosing a suitable type of natural fiber.</td>
<td>are created in the composite structure.</td>
<td>synthetic/mineral fibers used due to the</td>
<td>treatments applied on natural</td>
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<td>capability of these fibers to form strong</td>
<td>composites, since the layers of</td>
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<td>covalent bonding with the hosting matrix.</td>
<td>synthetic/mineral fibers do</td>
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<td>not need additional treatments.</td>
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Nomenclature

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>AA</td>
<td>Acrylic acid</td>
</tr>
<tr>
<td>AN</td>
<td>Acrylonitrile</td>
</tr>
<tr>
<td>APTES</td>
<td>3-aminopropyltriethoxy silane</td>
</tr>
<tr>
<td>APTMOS</td>
<td>Aminopropyltrimethoxy silane</td>
</tr>
<tr>
<td>AS</td>
<td>(3-aminopropyl)-triethoxysilane</td>
</tr>
<tr>
<td>BC</td>
<td>Bacterial cellulose</td>
</tr>
<tr>
<td>CNCs</td>
<td>Cellulose nanocrystals</td>
</tr>
<tr>
<td>CNTs</td>
<td>Carbon nanotubes</td>
</tr>
<tr>
<td>GLYMO</td>
<td>y-glycidoxypropyltrimethoxysilane</td>
</tr>
<tr>
<td>GPS</td>
<td>3-glycidoxypropyltrimethoxysilane</td>
</tr>
<tr>
<td>HDPE</td>
<td>High-density polyethylene</td>
</tr>
<tr>
<td>HDS</td>
<td>Hexadecyltrimethoxy-silanes</td>
</tr>
<tr>
<td>LDPE</td>
<td>Low density polyethylene</td>
</tr>
<tr>
<td>MA</td>
<td>Maleic anhydride</td>
</tr>
<tr>
<td>MAPP</td>
<td>Maleic anhydride grafted polypropylene</td>
</tr>
<tr>
<td>MMT</td>
<td>Montmorillonite</td>
</tr>
<tr>
<td>MPS</td>
<td>c-methacryloxypropyltrimethoxy silane</td>
</tr>
<tr>
<td>MRPS</td>
<td>c-mercaptopropyltrimethoxy silane</td>
</tr>
<tr>
<td>MS</td>
<td>3-(trimethoxysilyl)-1-propanethiol</td>
</tr>
<tr>
<td>NDS</td>
<td>Natural digestion system</td>
</tr>
<tr>
<td>OPEFB</td>
<td>Oil palm empty fruit bunch</td>
</tr>
<tr>
<td>OTMOS</td>
<td>n-Octyltrimethoxy silane</td>
</tr>
<tr>
<td>OSF</td>
<td>Olive stone flour</td>
</tr>
<tr>
<td>PBS</td>
<td>Poly (butylene succinate)</td>
</tr>
<tr>
<td>PE</td>
<td>Polyethylene</td>
</tr>
<tr>
<td>PES</td>
<td>Polyester</td>
</tr>
<tr>
<td>PLA</td>
<td>Poly lactic acid</td>
</tr>
<tr>
<td>PP</td>
<td>Polypropylene</td>
</tr>
<tr>
<td>RH</td>
<td>Relative humidity</td>
</tr>
<tr>
<td>SEBS</td>
<td>Styrene ethylene butadiene styrene-grafted maleic anhydride</td>
</tr>
<tr>
<td>TESPTS</td>
<td>Bis (3-(trimethoxysilyl) Propyl tetrasulphide</td>
</tr>
<tr>
<td>TEVS</td>
<td>triethoxy vinyl silane</td>
</tr>
<tr>
<td>TMSPMA</td>
<td>3-trimethoxysilyl propyl methacrylate</td>
</tr>
<tr>
<td>TPU</td>
<td>Thermoplastic polyurethane</td>
</tr>
<tr>
<td>UP</td>
<td>Unsaturated polyester</td>
</tr>
<tr>
<td>VE</td>
<td>Vinyl ester</td>
</tr>
<tr>
<td>VG</td>
<td>Vinyl grafting</td>
</tr>
<tr>
<td>VTMEO</td>
<td>Vinyltris (2-methoxethoxy) silane</td>
</tr>
<tr>
<td>VTMS</td>
<td>Vinyltrimethoxy silane</td>
</tr>
<tr>
<td>WF</td>
<td>Wood flour</td>
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<tr>
<td>WPCs</td>
<td>Wood plastic composites</td>
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</table>

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