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Full text available at http://www.sciencedirect.com/science/article/pii/S0141391016300970

DOI: doi:10.1016/j.polymdegradstab.2016.03.038

Please, cite it as:

O. Gil-Castell, J. D. Badia, T. Kittikorn, E. Strömberg, M. Ek, S. Karlsson, A. Ribes-Greus. Impact of hydrothermal ageing on the thermal stability, morphology and viscoelastic performance of PLA/sisal biocomposites. Polymer Degradation and Stability 2016, 132; 87-96

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IMPACT OF HYDROTHERMAL AGEING ON THE THERMAL STABILITY, MORPHOLOGY AND VISCOELASTIC PERFORMANCE OF PLA/SISAL BIOCOMPOSITES

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Abstract

The influence of the combined exposure to water and temperature on the behaviour of polylactide/sisal biocomposites coupled with maleic acid anhydride was assessed through accelerated hydrothermal ageing. The biocomposites were immersed in water at temperatures from 65 to 85 °C, between the glass transition and cold crystallisation of the PLA matrix. The results showed that the most influent factor for water absorption was the percentage of fibres, followed by the presence of coupling agent, whereas the effect of the temperature was not significant. Deep assessment was devoted to biocomposites subjected to hydrothermal ageing at 85 °C, since it represents the extreme degrading condition. The morphology and crystallinity of the biocomposites were evaluated by means of X-Ray diffraction (XRD) and field emission scanning electron microscopy (FE-SEM). The viscoelastic and thermal performance were assessed by means of dynamic-mechanical-thermal analysis (DMTA) and thermogravimetry (TGA). The presence of sisal generally diminished the thermal stability of the biocomposites, which was mitigated by the addition of the coupling agent. After composite preparation, the effectiveness of the sisal fibre was improved by the crystallisation of PLA around sisal, which increased the storage modulus and reduced the dampening factor. The presence of the coupling agent strengthened this effect. After hydrothermal ageing, crystallisation was promoted in all biocomposites therefore showing more fragile behaviour evidencing pores and cracks. However, the addition of coupling agent in the formulation of biocomposites contributed in all cases to minimise the effects of hydrothermal ageing.

Keywords

Biocomposites, polylactide (PLA), natural fibres, sisal, hydrothermal ageing, degradation, performance, mechanical fibre effectiveness

1. Introduction

Poly(lactic acid) or polylactide (PLA) is a biodegradable, highly versatile polyester produced from renewable sources [1]. It is mainly based on agricultural bases such as corn, beet, wheat and other starchy products [2]. The development of fully biodegradable composite materials based on PLA and lignocellulosic fibres is a technological challenge [3] [4] [5] [6]. Natural fibres provide better mechanical properties including high strength and stiffness without compromising important factors such as biodegradability, density or cost [7] [8] [9] [10]. Composites made of polylactide and natural fibres such as kenaf [11], hemp [7], flax [3] and sisal fibre [12] have been investigated and used in several applications. However, adverse effects can occur due to the high affinity to water of these natural fibres [13] [14] [15]. The chemical nature or surface topology may influence the composite interfacial bond [16]. In a composite, a susceptible area of degradation is the fibre-matrix interface, where decoupling, cracks and delamination can take place as a result of water absorption [17] [18]. In order to improve this interface, some alternatives have been proposed, such as coating the whole composite with a thin layer of polypropylene [19], adding an anti-hydrolysis agent [20] or even looking for specific processing methods [21] [22]. Among these, the incorporation of a coupling agent to the system is a simple and effective option to improve the cohesiveness of the composite and thus, the hydrolytic resistance [12] [22] [23] [24] [25] [26]. Usually, the addition of a coupling agent promotes chemical reactions between functional groups of the polymer matrix in their chain ends (-OH, -COOH) and -OH groups characteristic of cellulosic natural fibres [25] [26]. Maleic anhydride (MAH) is one of the most widely used reactive couplers due to its good chemical reactivity, low toxicity and low potential to polymerize itself under free radical grafting conditions [27].

Several factors can catalyse the hydrolytic breakage of the ester bond, such as acidic conditions, high moisture or high temperatures. Previous studies have demonstrated the relative stability of PLA and natural fibre when immersed in water at temperatures below the glass transition [12] [20]. In these cases, a Fickian model seems to govern the water uptake process. Conversely, at temperatures above the glass transition high degradation rates take place and non-Fickian behaviour in water uptake have been reported [12]. A high degradation degree can be achieved when samples are subjected to high temperatures and, therefore, long-term properties can be studied after water saturation. Accelerated hydrothermal ageing tests are regarded as a suitable tool to bring the materials to an advanced state of degradation as they could be after a hypothetical hydrolytic degradation along their service life [12] [28] [29] [30].

This research was focused on the influence of combined water and temperature on the morphology, thermal and viscoelastic performance of polylactide/sisal biocomposites, with special interest on the effects of using coupling agent in the formulation of the biocomposites.

2. Experimental procedure

2.1. Materials and reagents

Polylactide (PLA) 3251D was purchased from Natureworks (Minnetonka, USA) as pellets, with a glass transition in the 65-70 °C range. Sisal fibres coming from a farming crop, were supplied by the Thai Royal Project [31]. Their length was about 0.5-1 mm, tensile strength of 550 MPa, tensile modulus of 30 GPa and density about 1.5 g·cm⁻³. Dicumyl peroxide (DCP) 98% (Sigma-Aldrich, Sweden AB) and maleic anhydride (MAH) M188-99% (Sigma-Aldrich, Sweden AB) were used as free radical initiator and coupling agent, respectively.

2.2. Sample preparation

Prior to processing, virgin PLA (VPLA) and sisal fibre were dried in an oven at 80 °C during at least 12 hours and kept in zip bags. The fibre contents in the biocomposite were formulated as 10%, 20% and 30% by weight with and without coupling agent. In case of not using MAH, composites were prepared in an internal mixer (Brabender, Germany) during 5 minutes at 180 °C at the speed of 50 rpm. The resulting biocomposites were labelled as PLA10, PLA20 and PLA30. Biocomposites containing the coupling agent were prepared by incorporating MAH 2.5% and DCP 0.3% by weight in molten polymer and mixing during 5 minutes at 180 °C at the speed of 50 rpm. These samples were labelled as PLA10C, PLA20C and PLA30C. The compounded PLA/sisal biocomposite was then ground and the granules dried at 80 °C in the oven during at least 12 h. The final samples were obtained by means of a compression moulding equipment (Fontijne Presses, Netherlands). The biocomposites preparation procedure consisted of preheating the press to 200 °C for 2 min and applying a compression force of 150 kN during 2 min. An automatic cooling procedure into the press was performed. All the compression moulding operations were performed under vacuum conditions. The thickness of the sheet-shaped samples was 0.5±0.1 mm. Finally, all compounded biocomposites were dried at 50 °C in a vacuum oven (Heraeus Vacutherm 6025, Germany), then kept in zip bags and placed in a desiccator for further analyses at normalized lab conditions according to ISO 291 [32].

2.3. Hydrothermal ageing procedure

A normalized water absorption test reported in the ISO 62 - method 1 [33] was adopted as hydrothermal ageing environment. Temperature specifications were adapted to the desired ageing conditions. Samples of VPLA and composites were submerged into distilled water at six different temperatures (60, 65, 70, 75, 80 and 85 °C), below and above the polymer glass transition (~70 °C) and below the cold-crystallisation (~105 °C) [12], until water saturation was reached. In order to establish a common criterion, for a given temperature, the test extent was considered as the triple of the saturation time for the material with slower saturation rate, in order to assure that the mass uptake profiles followed an asymptotic horizontal behaviour, meaning the completion of the saturation. Concretely 8.0, 6.9, 3.3, 3.1, 2.5 and 2.3 hours were considered for the assays performed at 60, 65, 70, 75, 80 and 85 °C, respectively.

2.4. Characterisation

2.4.1. Water uptake

After saturation of biocomposites by water, the specimens were gently wiped to get rid of surface moieties. The amount of absorbed water (M_s) was calculated according to the **Equation 1**, where m_s and m_0 are the saturation mass and initial mass of the specimens, respectively. A weight difference procedure by means of a scale (Mettler Toledo AB135-S) with a precision of 0.1 mg was implemented. The average of quintuplicates was considered as a representative value.

$$M_S = \frac{m_S - m_0}{m_0} \times 100 \qquad (Equation 1)$$

2.4.2. X-Ray diffraction (XRD)

An X-Ray diffraction instrument (X'Pert Pro diffractometer model PW3040/60) was used to determine the crystal size of the biocomposites. Dried specimens of 100 mm² were considered for the analysis. The Cu K α radiation ($\lambda = 1.5418$ Å) was generated with a tension of 45 kV and

current of 35 mA and then monochromatized by using a Ni filter of 20 μ m. The experiment was measured and recorded in a reflection mode at the angular range of 5–60° (2 θ) and a rate of scanning of 0.05° per 10 s. An average spectrum of three different specimens was considered as representative for each sample.

In order to determine the mean size of the crystalline domains, the Scherrer's equation was exploited (**Equation 2**), where *K* is a dimensionless shape factor (0.9), λ is the X-Ray wavelength (0.154 nm), β is the line broadening at half the maximum intensity after subtracting the instrumental line broadening in radians and θ is the Bragg angle. The peak of 16.8° (2 θ) was observed to calculate the crystal size, which represented the diffraction of the (200)/(110) plane.

$$L_{hkl} = \frac{\kappa\lambda}{\beta\cos\theta}$$
 (Equation 2)

2.4.3. Thermogravimetric analysis (TGA)

Thermal stability data were obtained by thermogravimetry by means of a Mettler-Toledo TGA 851 series (Columbus, OH). The samples, with a mass of about 4 mg were introduced in TGA Mettler-Toledo perforated alumina crucibles, with capacity of 70 μ l. The samples were analysed in the temperature range of 25 to 800 °C with a heating rate of 10 °C·min⁻¹, under atmosphere of oxygen at a flow rate of 50 ml·min⁻¹. Experiments were repeated at least three times and the averages were considered as representative values.

2.4.4. Dynamic-mechanical-thermal analysis (DMTA)

Dynamic mechanical thermal analyses were conducted by the two-point bending mode in single cantilever clamping with 6 mm of effective length between clamps, by means of Mettler-Toledo DMA861e (Columbus, OH). The displacement was checked before all the experiments. The deformation force was set at 0.1 N. Experiments were carried out by heating from 35 °C to 130 °C with isothermal steps of 2 °C at 1 Hz of frequency and the displacement amplitude set at 10 μ m. Analyses were performed at least three times to ensure reproducibility.

2.4.5. Field emission scanning electron microscopy (FE-SEM)

The matrix-fibre interface was analysed by means of a Zeiss Ultra 55 Field emission scanning electron microscope (Oberkochen, Germany) to determine the morphology of the cryofracture obtained by breaking the samples after their immersion into liquid N_2 during 1 min. The aged and non-aged samples were cut into small pieces and dried at 50 °C in a vacuum oven for 24 h and then kept in desiccator during 48 h before FE-SEM sample preparation. The pieces were mounted on metal studs and sputter-coated with platinum during 10 s using a Leica EM MED020 high resolution sputter coater (Wetzlar, Germany). The analyses were performed at room temperature at a voltage of 3 kV.

3. Results and discussion

3.1. Influence of fibre, coupling agent and temperature on water absorption capability of *PLA/sisal biocomposites*

Table 1 shows the percentage of absorbed water at equilibrium (M_S) for VPLA and biocomposites at the hydrothermal ageing temperatures (T_{HA}) chosen for the study. VPLA attained a ~1% of M_S , regardless the T_{HA} . Due to presence of abundant hydroxyl groups on the cellulose molecules of

the sisal fibre, liable to interact with water molecules [34], the biocomposites showed M_S values from 3% to 10%.

$M_{S}(\%)$ -								
	60	65	70	75	80	85		
VPLA	1.1	1.2	0.8	0.9	1.1	1.1		
PLA10	3.6	3.6	3.0	3.7	3.2	3.2		
PLA20	6.3	6.3	6.1	7.1	6.1	6.2		
PLA30	9.4	8.7	10.4	9.3	6.3	10.3		
PLA10C	4.9	5.1	4.2	3.9	2.3	3.0		
PLA20C	7.4	6.9	5.9	6.1	5.1	5.2		
PLA30C	8.3	8.0	7.6	8.1	7.7	9.0		

Table 1. Water saturation mass (M_S) for VPLA and composites at all T_{HA} .

In order to assess the influence of fibre, coupling agent and ageing temperature, a statistical factorial analysis (SFA) was applied [35] [36]. **Figure 1** shows that the amount of fibre was the most influent factor, followed by the presence of the coupling agent in the composite. However, the variation of temperature slightly modified the percentage of absorbed water at equilibrium (M_s) by biocomposites and thus, was not considered for further analyses. Samples subjected to an accelerated hydrothermal ageing at 85 °C were therefore chosen to focus the study on the influence of fibre content and coupling agent.



Figure 1. Main-effects plot of relations between fibre, coupling agent and hydrothermal ageing temperature for water saturation mass (*Ms*).

3.2. Impact of hydrothermal ageing on thermal stability of PLA/sisal biocomposites

Thermal stability was assessed by means of thermogravimetric analysis (TGA) in order to ascertain the state of chemical degradation in the biocomposites. **Figure 2** shows the thermogravimetric (TG) and derivative thermogravimetric (DTG) curves for PLA/sisal biocomposites before and after the hydrothermal ageing. All curves showed a general weight-loss of 98-99%, regardless the formulation of the biocomposites. They exhibited four differentiated weight-loss stages, in accordance with reported results for sisal fibre [37]. The first slight initial loss can be related with remnant water present in both the matrix and fibres [38]. Then, the second decomposition stage, from 222 °C to 315 °C, is assigned to the decomposition of hemicellulose. The third stage, from 315 °C to 400 °C, was caused by decomposition of both the PLA matrix and

cellulose, which peaks at 356 °C and 325 °C respectively, are often overlapped. Finally, the fourth weight-loss from 380 °C to 500 °C is related to the degradation of char formed during previous decomposition.



Figure 2. Decomposition TG (a) and DTG (b) curves prior to hydrothermal ageing and TG (c) and DTG (d) curves after hydrothermal ageing for VPLA and biocomposites.

The peak decomposition temperature (T_{peak}) remained almost constant prior to and after hydrothermal ageing for all materials. The evolution of T_0 , calculated as the temperature in which weight-loss was 5% is plotted in **Figure 3**. Before hydrothermal ageing, the presence of sisal induced lower T_0 , due to its lower thermal stability, which resulted in a difference of 30 °C between VPLA (315 °C) and PLA30 (285 °C). The degradation of the hemicellulose seems to be responsible for this decrease. A decrease in the molecular weight of the PLA matrix during processing can also contribute to reduction of the onset degradation temperature [12]. Even though, T_0 was higher for biocomposites than for pure sisal fibre, attributed to protection of fibre by polymer matrix, according to Albano et al. [39]. The presence of coupling agent considerably reduced T_0 in all cases (~10 to 20 °C). This effect may be assigned to the reduction in molecular weight due to incorporation of dicumyl peroxide (DCP) followed by a maleic anhydride (MAH) branching reaction [12] [40].



Figure 3. Evolution of the onset temperature (*T*₀) prior to and after hydrothermal ageing at 85 °C for VPLA and biocomposites.

After hydrothermal ageing, two different behaviours were observed. For a given non-coupled biocomposite, hydrolytic reactions may have degraded the fibre-matrix interface, promoting peeling and delamination, thus exposing sisal fibre to ambient environment, showing the subsequent general decrease in T_0 . However, when coupling agent was added, an increment of T_0 was determined. Thus, though the presence of the coupling agent induced a reduction of thermal stability during processing, the biocomposites were more resilient to thermal decomposition after hydrothermal ageing. This might be explained by the presumable formation of crystals as a shell around sisal fibres [16], enhancing the fibre-matrix interface, as discussed in the next sections.

3.3. Fibre-matrix interface: morphology of cryofractured biocomposites

Fracture morphology before and after hydrothermal ageing was studied by field-emission scanning electron microscopy (FE-SEM). **Figure 4** shows the micrographs of the cryofractured surface for VPLA and biocomposites before and after the hydrothermal ageing at 85 °C. For non-aged materials, VPLA showed an apparently smooth surface while for biocomposites, the presence of fibres provoked a completely different fracture surface characterised by pores, holes and fibre tails.

During processing, the suggested crystallisation might have induced internal rearrangements that confer more fragile performance, resulting in fibre detaching, holes and pores. Concerning coupled biocomposites, better adhesion between fibre and matrix seemed to take place, but holes and traces still appeared after fracture.



Figure 4. FE-SEM micrographs for VPLA and PLA/sisal biocomposites prior to and after hydrothermal ageing at 85 $^{\circ}C$ (3 kV, 100×, 100 μ m).

After hydrothermal ageing, PLA matrix showed a rougher texture in VPLA and biocomposites. This change in the polymeric texture are in agreement with the hypothesis of the crystallisation occurred during hydrothermal ageing. The increase in fibre dimensions caused by swelling may deteriorate the interface, influencing the mechanical properties as shown afterwards, and resulted in cracks and rifts around fibres. However, the presence of coupling agent contributed to reach better polymer/fibre interface. **Figure 5** shows the detail of PLA30 and PLA30C for the sake of comparison. Prior to ageing, a small gap between fibre and matrix could be seen for PLA30, while a kind of polymeric covering of fibres by PLA matrix was found for PLA30C. After ageing, the separation of fibres and polymeric phase increased for PLA30. However, the presence of coupling agent in PLA30C seemed to reduce delamination.



Figure 5. FE-SEM detail micrographs for PLA30 and PLA30C prior to and after hydrothermal ageing at 85 °C (3kV, $1000\times$, 10 μ m).

3.4. Development of crystallisation during hydrothermal ageing

It has been suggested in previous sections that the influence of water in the polymer matrices in combination with the temperature effect and the presence of sisal fibres might be a consequence of microstructure change, and thus X-Ray diffraction was considered to counter-check results.

Figure 6 shows the diffraction patterns of VPLA and biocomposites prior to and after hydrothermal ageing at 85 °C. The crystallinity of PLA matrix can be found by the reflection peaks positioned at 16.8°, 19.1° and 22.6°, which represented the diffractions of (200)/(110), (203), (210) planes respectively [41]. Prior to the hydrothermal ageing, VPLA showed reflections at 16.8° and 19.1° related to typical PLA crystalline domains. Biocomposites also showed characteristic reflections of a semicrystalline microstructure, more intense when sisal fibre augmented. In addition, a wide reflection around 22.5° was observed, which corresponded to the diffraction of the (002) plane of the sisal fibre, as described by M. Benítez-Guerrero et al. [42]. This reflection associated to sisal presumably overlapped the peak at 22.6°, characteristic of the PLA crystalline domains. After hydrothermal ageing, a general reduction of the wide flat signal

between 10° and 30° corresponding to the amorphous morphology of PLA, was shown. This reduction was compensated by the increase of the sharp peaks at 16.8° and 19.1°, suggesting an increment in crystallinity in aged materials.



Figure 6. Diffraction patterns of VPLA and biocomposites prior to and after hydrothermal ageing at 85 °C.

Several studies demonstrated that a diffraction at 24.1° corresponding to (016) plane is related to the presence of crystalline domains in PLA in the α mode [43] [44] [45]. Conversely, an absence of the peak at 24.1° indicates that the structure of the crystalline domains of PLA are ordinated according to the α' mode, which occasionally is corroborated by a reflection at 24.6° [46]. The differences between the α' and the α crystalline modes are mainly based on a slight rearrangement of the chain conformation and packing in the unit cell, especially related to the side groups, to achieve a more energy-favourable state, i.e. the reduction of unit cell dimension [44]. The transition from α' to α form usually takes place by an increase of the crystallisation temperature $(T_c \ge 105 \text{ °C})$ [47]. The phase composition determination in polymorphic polymers is typically evaluated from the ratio of the intensity of the reflections of one crystal mode to the sum of the heights of intensities of the reflections of all polymorphs. However, a precise estimation cannot be performed for PLA samples, as the two polymorphs have very similar profiles with slight differences, e.g. a small shift in peak position and/or appearance of very weak reflections [47]. Taking into account the complex differentiation of both polymorphs for PLA based materials a particular structure cannot be defined. Even so, the peak absence at 24.1° suggested the presence of the α' mode for all materials.

In order to evaluate the crystalline structure from X-Ray diffraction studies, the mean size of the crystalline domains (L_{hkl}) was calculated from the peak at 16.8°, which represented the diffraction of (200)/(110) plane, according to **Equation 2**. Figure 7 shows the obtained results along with the crystallinity degree values obtained in a previous study by means of calorimetric analyses [12]. Accordingly, the crystallinity of these materials can be qualitative and quantitatively ascertained. Prior to hydrothermal ageing, greater crystallinity degree involving bigger crystal size was found for biocomposites. During processing, the presence of sisal fibres promoted the formation of bigger crystalline domains than those found for VPLA. After hydrothermal ageing at 85 °C, the crystallinity degree grew in all cases. However, the generation of the new crystalline populations was not directly related with a mean size increment. This fact can be understood in terms of the competitive behaviours of nucleation and growth of crystalline domains. VPLA and PLA/sisal biocomposites with low fibre content showed an increase of crystal size, assignable to

a more relevant growth of crystals. In contrast, the increment of $L_{(200/110)}$ for biocomposites was lower as fibre content increased. Concretely, the crystalline size for PLA30 and PLA30C almost remained constant after hydrothermal ageing, which may be related to a more prominent nucleation process. This fact was in line with the observation of transcrystalline regions throughout the fibres which implies the generation of small crystalline domains around fibres, perpendicular to the fibre surface [48]. Actually, natural fibres as jute or sisal into a PLA matrix are known to generate a particular oriented microstructure, as demonstrated by Le Duigou et al. [16]. Summing up, hydrothermal ageing promoted in composites the transcrystallisation phenomenon, more perceptible as fibre content increased.



Figure 7. Mean size of the crystalline domains ($L_{(200/110)}$) for VPLA and biocomposites prior to and after hydrothermal ageing at 85 °C. Percentage values gathered in bars correspond to the crystallinity degree (X_c), as obtained from calorimetric analyses in a previous study [12].

3.5. Impact of hydrothermal ageing on dampening, storage modulus and fibre effectiveness

Dynamic mechanic thermal analysis (DMTA) is a widely used technique for investigating the viscoelastic behaviour of polymeric materials, giving information about the stiffness and damping characteristics of the material by means of the storage modulus (E'), the loss modulus (E') and the tangent of the out-of-phase angle delta ($tan \delta$) [49]. The determination of those properties over a range of temperatures is important in polymer and composite characterisation prior to and after ageing and degradation tests [50] [51] [52] [53] [54].

Table 2. Glass-rubber relaxation temperature (T_{GR}) for VPLA and biocomposites prior to and after hydrothermal
ageing at 85 °C.

<i>T_{GR}</i> (°C)	VPLA	PLA10	PLA20	PLA30	PLA10C	PLA20C	PLA30C
Non-aged	62.61	63.37	63.54	63.70	63.63	63.58	63.88
Hydrothermal ageing at 85 °C	74.15	72.27	72.12	71.95	72.56	71.87	73.87

The dampening gives information about the internal friction of the biocomposites and consequently would be influenced by the molecular motion in the interface, which can be studied in terms of tangent of delta (*tan* δ , i.e. E''/E') [55]. In the **Figure 8a**, the *tan* δ is represented as a function of temperature for VPLA, PLA30 and PLA30C before and after hydrothermal ageing. The peak in the dampening is known to take place within the glass transition and its maximum is often recorded as the glass-rubber relaxation temperature (T_{GR}) [55]. Temperature values of *tan* δ peak are gathered in **Table 2**. Non-aged materials showed similar T_{GR} . However, after hydrothermal ageing, the value shifted to temperatures around ~10 °C higher for all cases. This

increment was related to the crystalline phase formation during ageing, which hindered the movement of amorphous polymer chains. Moreover, some peaks could be observed after glass transition. Internal chain rearrangements previous to the polymer cold-crystallisation, connected with the interactions between fibre and PLA matrix might be the cause of the appearance of this atypical behaviour. **Figure 8b** shows the peak heights of *tan* δ as a function of fibre content and coupling agent prior to and after hydrothermal ageing at 85 °C. This parameter is usually considered as a useful indicator of the dampening behaviour. Due to processing, a reduction of both the height and the broadening of the relaxation was observed in the presence of sisal. This effect was greater with increasing the fibre content. As suggested by Pothan et al. [56], for a composite with a low fibre ratio, the polymer chains are relatively free to move. However, when the fibre content is increased, the mobility of the polymer chains is reduced, hindered by the reinforcement and crystallinity formed during the preparation of composites. After hydrothermal ageing, the profiles of *tan* δ showed a peak about 60% lower and considerably wider for all studied compositions. All materials, including VPLA, showed equivalent constraints of chain motion of the amorphous phase due to the increased crystallinity during ageing.



Figure 8. (a) Tangent delta (*tan* δ) as a function of temperature prior to and after hydrothermal ageing at 85 °C for VPLA, PLA30 and PLA30C; (b) *tan* δ peak height as a function of fibre content for VPLA and composites prior to and after hydrothermal ageing at 85 °C.

Figure 9 shows the evolution of the storage modulus (E') from the glassy to the rubbery state for all the biocomposites before and after the hydrothermal ageing. VPLA exhibited a similar behaviour as in literature, showing an E' around ~2200 MPa [4] [57] [58]. Regarding to biocomposites, prior to hydrothermal ageing, the crystallisation around sisal induced a reduction of mobility and deformation of the matrix. The presence of coupling agent improved the fibre-matrix interface. In agreement with literature, it caused an effective stress transfer from matrix to fibres through the crystalline domains [55] [59] [60] [61] [62]. Along the DMTA experiment, a drastic drop of E' was observed within the glass-rubber relaxation. Afterwards, E' started to increase again around 80 °C which confirmed the typical effect of cold-crystallisation [4] [12]. This change in trend occurred at lower temperatures as the fibre content increased, reaching higher E' once the cold-crystallisation was completed.

After hydrothermal ageing, the glass-rubber relaxation turned into a smoother transition and the cold-crystallisation phenomenon disappeared, in agreement with the complete crystalline development suggested in previous sections. Two differentiated behaviours were found above and below the glass-rubber relaxation temperature T_{GR} . On the one hand, below the T_{GR} , the deterioration of the fibre-matrix interface caused by water resulted in lower storage modulus for the biocomposites. However, above the T_{GR} , the crystalline structure formed during the

hydrothermal degradation promoted a higher storage modulus for composites in the rubbery state. This effect was higher for coupled biocomposites.



Figure 9. Storage modulus (*E'*) as a function of temperature for VPLA and composites prior to (left) and after hydrothermal ageing (right).

Figure 10 shows the *E'* values taken at 40 °C in the glassy state with the purpose of quantifying the effects of fibre and coupling agent as well as the effect of hydrothermal ageing. For non-aged samples, *E'* increased by ~20% (2200 to 3000 MPa) from VPLA to PLA30. For a given fibre percentage, coupling agent generally promoted an increase in *E'*, greater when fibre content increased, reaching the highest value for PLA30C (3300 MPa). After hydrothermal ageing at 85 °C, *E'* slightly increased for VPLA (2200 to 2800 MPa) while it decreased for biocomposites. In general, the presence of sisal fibre reduced *E'*. This reduction was greater the higher the fibre content was. A reduction from 2800 to 600 MPa between VPLA and PLA30 was found. The coupling agent always contributed to minimise the effect of hydrothermal ageing around ~10%.



Figure 10. Storage modulus (*E'*) in the glassy state as a function of fibre content for VPLA and composites prior to and after hydrothermal ageing at 85 °C.

Finally, the mechanical fibre effectiveness, i.e. MFE, was calculated according to **Equation 3** [56] [63], where E'_g and E'_r are the storage modulus values in the glassy and rubbery region respectively. In general terms, the higher the value of the constant MFE is, the higher the effectiveness of the filler is considered.

$$MFE = 1 - \frac{(E'_g/E'_r)_{comp}}{(E'_g/E'_r)_{resin}}$$
(Equation 3)

Figure 11 shows the values of the *MFE*, taking the measured E' values at 40 °C and 100 °C as E'_g and E'_r respectively. Prior to hydrothermal ageing, the effectiveness of the filler was significantly higher when the sisal content increased, in agreement with other reports [59] [60] [61]. The most effective fibre percentage was always 30% by weight, which revealed active stress transfer between fibre and matrix through the crystalline phase formed both during processing and the DMTA experiment. After hydrothermal ageing, an increase of *MFE* was found for all cases, as a result of the induced crystalline formation around fibres in the so-called transcrystalline region. The influence of coupling agent could be perceived at low fibre contents, while it was negligible for biocomposites with fibre content higher than 20% by weight.



Figure 11. Values of the *MFE* as obtained by Equation 3, for VPLA and composites prior to and after hydrothermal ageing at 85 °C.

4. Conclusions

An accelerated hydrothermal ageing test was a suitable procedure to evaluate the combined influence of water and temperature on the morphology of PLA/sisal biocomposites, and the subsequent effects on the thermal and viscoelastic stability of the materials.

After the preparation of biocomposites, the sisal fibre promoted crystallisation which altered the viscoelastic properties, increasing the storage modulus. However, lower thermal stability was established for biocomposites. The presence of coupling agent slightly affected the crystalline formation, increased the storage modulus and displaced the thermo-decomposition process towards higher temperatures.

Hydrothermal ageing boosted crystallisation for all studied biocomposites, both in terms of crystalline degree and mean size of crystals. It also provoked a general reduction of the storage moduli of biocomposites, along with a decrease of the dampening performance, due to chemical degradation, therefore showing more fragile behaviour evidencing pores and cracks. The mechanical effectiveness of the fibres in aged biocomposites was improved by the addition of fibres. Results showed that PLA/sisal biocomposites with coupling agent were resilient to hydrothermal ageing.

Acknowledgements

The authors would like to acknowledge the Spanish Ministry of Economy and Competitiveness, through the Research Project ENE2014-53734-C2-1-R and UPOV13-3E-1947. The Spanish Ministry of Education, Culture and Sports is also thanked for the predoctoral FPU grant of O. Gil-Castell (FPU13/01916). Generalitat Valenciana is thanked for the APOSTD14/041 program for J.D. Badia. The Chilean Government is thanked for the 13CEI2-21839 project. The financial support given by the Prince of Songkla University and KTH Royal Institute of Technology is gratefully acknowledged.

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