



Exfoliating nanomaterials in canola protein derived adhesive improves strength and water resistance†

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Wood adhesives are mainly produced using petrochemical-based resins. However, concerns over environmental and health effects have prompted a renewed interest on developing bio-based adhesives. The potential of using canola protein for adhesives was explored with limited success. Similar to other proteins, canola adhesives also have weak adhesion and water resistance. The objective of this research was to develop and characterize nanomaterial reinforced canola protein adhesive with improved water resistance and adhesion strength. Hydrophilic bentonite (bentonite), surface modified nanoclay (SM-MMT), graphite oxide (GO), and nanocrystalline cellulose (NCC) were exfoliated at different addition levels (0, 1, 3, 5, 10% w/w) using an improved solution intercalation method. A significant increase ($p < 0.05$) in both dry and wet adhesion was found at addition levels of 1–3% (w/w). NCC and GO showed superior effect over bentonite and SM-MMT. The improvement in adhesive strength at low nanomaterial addition was a result of adequate exfoliation of nanomaterials in protein matrix, and increased exposure of functional groups for enhanced interaction with wood surface. This study demonstrated for the first time the potential of nanomaterials for improving adhesion and water resistance of protein-derived adhesive. Canola protein adhesive developed in this study showed its potential for replacing synthetic adhesives as green alternative adhesive materials.

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Introduction

The wood adhesive industry is dominated by petrochemical-derived resins, such as urea formaldehyde (UF), phenol formaldehyde (PF) and melamine urea formaldehyde (MUF).^{1–3} However, increasing concerns on petrochemical based wood adhesives such as environmental impact, potential health hazards due to formaldehyde emission, and non-renewability have renewed the interest in developing green, renewable alternatives.^{3–6} Biobased adhesives, derived from protein, and polysaccharides were widely used before they were replaced by synthetic ones during the World War II.^{1,2,5,6} However, the challenge remains with regard to developing cost-effective and performance comparative adhesives from these biobased materials.² Proteins are preferred among other biobased polymers for preparing adhesives, due to their versatile functionalities as well as flexibility for modifications.^{2,7} Historically, casein, gelatin, blood and soy proteins were applied for various adhesive applications.^{2,4,8} More recently, the possibility of using other protein sources such as wheat gluten,⁹ cottonseed protein,¹⁰ triticale protein,^{5,11} and canola protein⁷ were studied.

Without exception, canola protein derived adhesives also showed poor water resistance and low adhesion strength.⁷

Canola is the second largest oil seed crop in the world and oil processing generates a great deal of meal, containing 35–40% w/w proteins. Irrespective to its high protein content, canola meal has limited uses mainly as a low value animal feed or as a fertilizer.^{7,12} Canola meal mainly consists of cruciferin (12S), napin (2S) and oleosin with approximate proportions of ~60%, ~20% and ~8% respectively.¹³ Cruciferin is a neutral protein (PI ~ 7.2) with molecular weight of ~300–310 kDa while napin is strongly basic (PI ~ 11.0) protein because of high proportion of amidated amino acids present in its structure with a molecular weight of ~12.5–14.5 kDa.¹⁴ Canola protein adhesive prepared by modifying with sodium bisulfite showed dry, wet and soaked adhesion strength values of 5.28 ± 0.47 MPa, 4.07 ± 0.16 MPa and 5.43 ± 0.28 MPa, respectively.¹⁵ Canola adhesive prepared with 0.5% sodium dodecyl sulphate (SDS) showed dry, wet, and soaked adhesion strength of 6.00 ± 0.69 , 3.52 ± 0.48 MPa, and 6.66 ± 0.07 MPa, respectively.¹⁶ By grafting poly(glycidyl methacrylate) into canola protein, our study showed improved dry, wet and soaked strength of 8.25 ± 0.12 MPa, 3.80 ± 0.15 MPa and 7.10 ± 0.10 MPa for respectively.⁷ However, the use of high amount of expensive poly(glycidyl methacrylate) polymer (2.7 g/2 g protein, w/w) prevents its future commercial exploration. Therefore, there is a need to look for new methods to improve water resistance and adhesion strength of canola protein.

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Nanomaterials are widely used in material science in order to change mechanical, electrical and chemical properties of the bulk material.⁶ For example, in composites research, adding nanomaterials were reported to improve flexural strength, elasticity, toughness and stability of the material.¹⁷ The potential of nanomaterials in improving adhesive performance was recently explored but with limited success. Kaboorani *et al.* (2011, 2012) observed slight improvement in adhesion and water resistance after adding montmorillonite,¹⁸ nano aluminum oxide,¹⁹ and nanocrystalline cellulose⁶ into polyvinyl acetate adhesives at low nanomaterial concentrations. Research on nanomaterial addition into protein based adhesives was extremely limited. Zhang *et al.* (2014) reported a decreased adhesion strength of polyisocyanate modified soy protein with the addition of montmorillonite, probably due to a nano scale blocking mechanism.²⁰ Li *et al.* (2016) recently studied the effect of modified sepiolite-based united crosslinked network in improving adhesion of soybean meal-based wood adhesive and reported an improvement of wet strength from 0.81 MPa to 1.18 MPa.³ Another recent study by Qi *et al.* (2016) reported an improvement of wet adhesion from 2.9 MPa to 4.3 MPa by exfoliating sodium montmorillonite at 8% w/w addition rate into soy protein isolate.²¹ Both studies suggest that “physical filling effect” of sepiolite,³ and nanomaterial induced cross-linking of protein network as the key factors contributing towards improving adhesion.^{3,21} It is well recognized that proper dispersion and exfoliation of nanomaterials is a critical factor in their applications.²² Therefore, it is necessary to develop new methods for exfoliating nanomaterials in protein-based adhesive matrix.

We hypothesized that a proper dispersion of nanomaterials into canola protein adhesive would improve water resistance and adhesion strength. The main objective of this study is to develop and characterize nanomaterial reinforced canola protein adhesive with improved water resistance and adhesion strength. Effects of addition levels and intercalation conditions of nanomaterials such as hydrophilic bentonite (Bento), surface modified montmorillonite (with 25–30%, w/w trimethyl octadecylammonium chloride–SM-MMT), nanocrystalline cellulose (NCC), and graphite oxide (GO) were studied in this research. These nanomaterials have been selected based on the strong evidence found in literature in improving functional properties of adhesives, composites and plastic research.^{6,18,23,24} Nanoclays are hydrated material made with either tetrahedral or octahedral stacks of silicate sheets.²⁵ Bentonite is typically considered as an impure form of clays that contain both montmorillonite and other crystalline structures, arranged in a octahedral sheet sandwich between two tetrahedral plates and an isolated additional octahedral plate.²⁶ Montmorillonite is a naturally hydrophilic and inorganic material that made with two stacked layers of tetrahedral sheets around a middle octahedral sheet, and usually contains hydrated Na⁺ or K⁺ ions.^{25,26} Montmorillonite used for this study is modified with 25–30% (w/w) trimethyl octadecylammonium chloride to improve interlayer spacing and hydrophobicity. GO consists of oxidized graphite sheets (or graphene oxide sheets) as their basal planes while surface of the sheet is decorated mostly with epoxide, hydroxyl, carbonyl and

carboxyl groups.²⁷ Nanocrystalline cellulose (NCC) is derived from acid hydrolysis of native cellulose and shows a rigid rod-like crystals with diameter in the range of 10–20 nm and lengths of a few hundred nanometers.²⁸

Experimental section

Materials and chemicals

Canola meal was a generous gift from Richardson Oilseed Ltd. (Lethbridge, AB, Canada). Hydrophilic bentonite (Bento), surface modified montmorillonite (SM-MMT), graphite and cellulose were purchased from Sigma-Aldrich (Sigma Chemical Co, St. Louise, MO, USA). All chemicals were from Fisher Scientific (Ottawa, ON, Canada) unless otherwise noted.

Method

Canola protein extraction. Proteins were extracted from canola meal as described by Manamperi *et al.*, (2010)¹² with slight modifications. Canola meal was ground to pass through 100-mesh size using a Hosokawa milling and classifying system (Hosokawa Micron Powder Systems, Summit, NJ, USA). Canola meal was mixed with mili-Q water in 1 : 10 (w/v) ratio; pH was adjusted to 12.0 using 3 M NaOH and stirred for 30 min followed by centrifugation (10 000g, 15 min, 4 °C). The supernatant was collected, readjust pH to 4.0 using 3 M HCl and centrifuged as above after 30 min stirring. The resulting precipitate was collected, freeze-dried and stored at –20 °C until further use.

Graphite oxide preparation. Graphite oxide nanoparticles (GO) were prepared according to Hummers and Offeman method²⁹ with slight modifications. In brief, 5 g of graphite and 2.5 g of NaNO₃ was mixed in a glass beaker where 120 mL of concentrated H₂SO₄ was slowly added while stirring for 30 min (200 RPM) in an ice bath to oxidize graphite. Then, 15 g of KMnO₄ was slowly added to the mixture while maintaining temperature at 35 ± 3 °C and stirred for 1 h. After the reaction, 92 mL of deionized water was added to the reaction mixture, and stirred for 15 min. Leftover KMnO₄ was neutralized by adding another 80 mL of hot (60 °C) deionized water containing 3% H₂O₂. After cooling to room temperature, the sample was centrifuged (10 000g, 15 min, 4 °C) to remove any remaining acid and chemicals. The precipitate was washed for three times as above to prepare oxidized graphite, followed by 5 min sonication (at 50% power output), before freeze drying.

Nanocrystalline cellulose preparation. Nanocrystalline cellulose (NCC) was prepared from cellulose samples as described by Cranston and Gray (2006)³⁰ with slight modifications. Cellulose hydrolysis was carried out by mixing 20 g of cellulose powder with 350 mL of 64% (w/w) sulfuric acid under continuous stirring for 45 min at 45 °C. The mixture was diluted 10 times with deionized water in order to suspend the reaction and centrifuged (10 000 g, 4 °C, 10 min) to remove excessive acid. The resulting precipitate was washed with deionized water, and centrifuged to remove any remaining chemicals. Extracted NCC was dialyzed for three days against deionized

water until neutral pH achieved, freeze-dried, and stored at -20°C until further use.

Exfoliation of nanomaterials and adhesive preparation. A solution intercalation method was developed to exfoliate nanomaterials into canola protein matrix. In brief, 3 g of canola protein was mixed with 20 mL of deionized water to make 15% w/v dispersion. Samples were stirred for 6 h (300 rpm, RT) to disperse canola protein, and pH was readjusted to 5.0 using 1 M HCl solution. Nanomaterials at various concentrations (to have final concentrations of 0%, 1%, 3%, 5%, and 10% w/w, nanomaterial/protein) were separately dispersed in 10 mL deionized water, stirred for 5 h at room temperature (300 rpm) and another 1 h at $45 \pm 3^{\circ}\text{C}$. Dispersed nanomaterials were sonicated for 3 min using a medium size tapered tip attached to a high intensity ultrasonic dismembrator (Model 500, Thermo Fisher Scientific INC, Pittsburg, PA, USA) providing intermittent pulse dispersion of 5 s at 3 s intervals and 60% amplitude. Resulting nanomaterial dispersions were homogenized for 2 min at 20 000 rpm using digital ULTRA TURRAX high shear homogenizer (Model T25 D S1, IKA® Works, Wilmington, NC, USA). Then, prepared nanomaterial dispersions were slowly added to protein dispersions dropwise while stirring for 15 min to have a final protein concentration of 10% w/v. Following the intercalation, protein-nanomaterial mixture was sonicated and homogenized as above and the pH of the adhesive mixture was readjust to 12.0 by adding 6 M NaOH solution. Negative controls were prepared by dispersing canola protein in deionized water at 10% w/v ratio and used as is. In the pH controls, canola protein samples were dispersed in deionized water at 10% w/v ratio, but adjust the pH to 12.0 similar to nanomaterial reinforced samples, without adding nanomaterials. Solution intercalation method developed by Zhang *et al.* (2014) was used to produce canola protein adhesives from SM-MMT and NCC for the purpose of comparing water resistance and adhesion of the method developed in our lab.²⁰

Adhesion strength measurement. Birch veneer samples with a thickness of 1.2 mm were cut into a dimension of 20 mm \times 120 mm (width and length) using a cutting device (Adhesive Evaluation Systems, Corvallis, OR, USA). They were conditioned according to the requirement of ASTM (American Society for Testing and Materials) standard method D2339-98 (2011)³¹ at 23°C and 50% relative humidity in a controlled environment chamber (ETS 5518, Glenside, PA, USA). The prepared adhesive samples were spread at an amount of 40 μL per veneer strand in a contact area of 20 mm \times 5 mm using a micropipette. After adhesive application, veneer samples were air dried for 5 min, followed by hot pressing at 120°C and 3.5 MPa for 10 min using Carver manual hot press (Model 3851-0, Carver Inc, In, USA). Dry adhesion strength (DAS) was measured according to ASTM standard method D2339-98 (2011) by measuring tensile loading required to pull bonded veneer using Instron machine (Model 5565, Instron, MA, USA) equipped with a 5 kN load cell and data was collected using Bluhill 3.0 software (Instron, MA, USA). Wet adhesion strength (WAS) and soaked adhesion strength (SAS) were measured according to ASTM standard D1151-00 (2013)³² using Instron tensile loading. WAS values were measured after submerging bonded veneer samples for 48 h in water (23°C)

while SAS was measured after reconditioning submerged wet samples for another seven days at 23°C and 50% relative humidity in a controlled environment chamber. In each strength testing (DAS, WAS and SAS) minimum four bonded veneer samples per replicate was used. All samples were clamped to Instron with a 35 mm gauge length and tested at 10 mm min^{-1} cross head speed.

Differential scanning calorimetry (DSC). Thermal properties in prepared nanomaterials and adhesive samples were analyzed using differential scanning calorimeter (Perkin-Elmer, Norwalk, CT, USA). Temperature and heat flow were calibrated using pure indium samples. Moisture in the samples was removed by freeze-drying followed by drying in a hermetic desiccator containing P_2O_5 for two weeks before analysis. Nanomaterial and adhesive samples were accurately weighed into T-Zero hermetic aluminum pans (~ 6 mg each), mixed with 60 μL of 0.01 M phosphate buffer, hermetically sealed with lids, and analyzed against an empty reference pan under continuous nitrogen purging. All samples were equilibrated at 0°C for 10 min and heated from 0 to 250°C at a ramping rate of $10^{\circ}\text{C min}^{-1}$. Thermodynamic data was collected and analyzed using Universal Analysis 2000 software for thermal transition changes in adhesives and nanomaterials (Perkin-Elmer, Norwalk, CT, USA).

Fourier transform infrared spectroscopy (FTIR). FTIR was used to characterize nanomaterial induced secondary structural changes of canola proteins using Nicolet 8700 Fourier transform infrared spectrometer (Thermo Eletron Co. WI, USA). Moisture in the samples was removed by freeze drying followed by drying in a hermetic desiccator containing P_2O_5 for two weeks before analysis. Graphite, GO, NCC, and nanomaterial reinforced adhesive samples were mixed with potassium bromide (KBr), and milled to make a fine powder before FTIR analysis. IR spectra in the range of $400\text{--}4000\text{ cm}^{-1}$ were collected using 128 scans at a resolution of 4 cm^{-1} . Collected data were processed and analyzed with Origin 2016 software (OriginLab Corporation, MA, USA), where the second derivative of FTIR spectra was used to identify the protein secondary structural changes.

X-ray diffraction. X-ray diffraction (XRD) of nanomaterial reinforced adhesive samples was performed using Rigaku Ultima IV powder diffractometer (Rigaku Co. Japan). Cu- $\text{K}\alpha$ radiation (0.154 nm) was used to collect the angle data (2θ) from 5 to 50 degrees. XRD data was processed using Origin 2016 software (OriginLab Corporation, MA, USA) for nanomaterial-reinforced adhesives to identify the dispersion pattern at different nanomaterial concentrations. Interlayer distance of nanomaterials were calculated using the Bragg's equation;³³ $\sin \theta = n\lambda/2d$, where λ is the wavelength of X-ray radiation used in the experiment, d is the spacing between diffraction lattice (interlayer spacing), and θ is the glancing angle (measured diffraction angle).^{18,34}

Transmission electron microscopy (TEM). Transmission electron microscopy analysis was performed using Philips/FEI transmission electron microscope (Model Morgagni, FEI Co, OR, USA) coupled with Gatan digital camera (Gatan Inc, CA, USA). For nanomaterials, samples were dispersed in ethanol at



a concentration of 0.5% w/w whereas adhesive samples were diluted to 100 fold with ethanol before TEM imaging. A drop of prepared solution was casted onto 200 mesh holey copper grid covered with carbon film and allowed for air drying before imaging. For NCC sample and adhesive containing NCC, 1% w/w uranyl acetate drop was added onto air dried drop in the copper grid in order to improve the contrast of NCC fibres.

Statistical analysis. Data were analyzed using analysis of variance (ANOVA) followed by Duncan's Multiple Range (DMR) test to identify the effects of each nanomaterial concentration on adhesion strength (dry, wet, soaked) using Statistical Analysis System Software (SAS version 9.4, SAS Institute, Cary, NC). Effect of nanomaterial concentrations on adhesion strength of each nanomaterial was evaluated at the 95% confidence level.

Results and discussion

Characterization of nanomaterials

Fig. 1 shows the diffraction angles and interlayer spacing of nanomaterials used in this study. Both bentonite and SM-MMT show similar crystalline peaks around diffraction angles of 19.7°, 34.9°, and 54.0° with interlayer spacing of 0.450 nm, 0.257 nm, and 0.167 nm, respectively. Kaboorani and Riedl (2011) also observed similar intense peaks in unmodified montmorillonite clay.¹⁸ The crystalline peaks at 6.6° and 28.7° in bentonite has shifted to 5.6° and 26.5° in SM-MMT whereas their corresponding interlayer spacing have shifted from 1.346 and 0.310 nm to 1.559 and 0.336 nm, respectively. Surface modification of montmorillonite with 25–30% (w/w) trimethylolctadecylammonium chloride polymer is known to cause changes of diffraction angle and the increase in interlayer spacing.^{35,36}

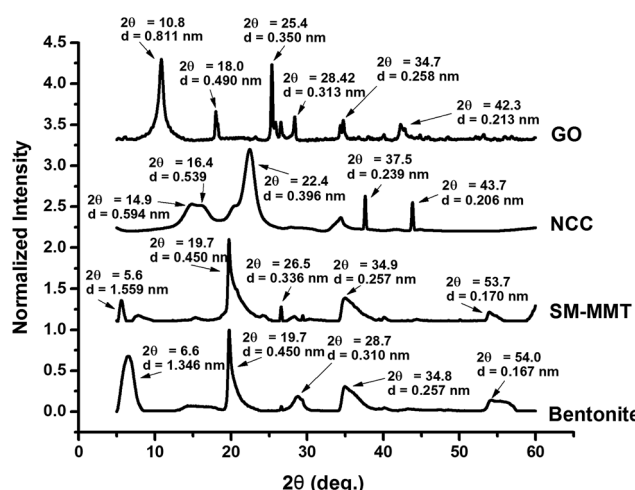


Fig. 1 X-ray diffraction patterns show glancing angle (θ) and interlayer spacing (d) of bentonite, SM-MMT (surface modified montmorillonite), NCC (nanocrystalline cellulose) and GO (graphite oxide) used in the adhesive preparation. Notes: X-ray diffraction data of the nanomaterials were collected at the glancing angle (2θ) range of 5–60° and interlayer spacing (d) was calculated according to the Bragg's equation: $\sin \theta = n\lambda/2d$.

Crystallinity and interlayer spacing of NCC depend on the method of preparation.^{37,38} NCC samples show three characteristic cellulose crystalline peaks at 14.9°, 16.4 and 22.4° with interlayer spacing of 0.594 nm, 0.539 nm and 0.396 nm, respectively. Cellulose crystalline peaks at similar diffraction angles were previously reported.^{37,38} In addition, two other minor peaks were shown at 37.5°, and 43.7° diffraction angles with interlayer spacing of 0.239 nm and 0.213 nm, respectively. GO samples show two major peaks at 10.8° and 25.4° with interlayer spacing of 0.811 nm and 0.350 nm. In addition, three minor peaks were observed in the prepared GO at 18.0°, 34.7° and 42.3° angles with d space of 0.490 nm, 0.258 nm and 0.213 nm, respectively. The interlayer spacing of GO mainly depends on their oxidation level and C : O ratio;³⁹ GO prepared for this study has a C : O ratio of 2.18 (Fig S1: ESI†). Krishnamoorthy *et al.* (2013) identified similar crystalline peaks for graphite oxide prepared under different oxidation conditions. They attributed the peak at 10.8° to an oxidation product whereas the peak at ~25.4° to crystallinity of graphite. In the same study they observed changes in diffraction angle and interlayer spacing with different oxidation conditions.³⁹ TEM images of nanomaterials are shown in Fig. 2. Both bentonite and SM-MMT showed the platelet like structure at ~80–150 nm whereas NCC samples appear to be in a long rod like fibers at ~60–90 nm diameters. GO appeared to be thin sheets stacked one another with average width of ~600–800 nm.

Dispersion of nanomaterials in canola protein

Previous studies on dispersing NCC, nanoclay, and Al_2O_3 nanoparticle in PVA adhesives suggested that homogeneous dispersion/exfoliation of nanomaterials is the key to improve adhesion strength.^{6,18,19} Fig. 3 shows the TEM images of nanomaterial dispersion in canola adhesive samples at different concentrations. At a 1% addition level, all four nanomaterials were exfoliated where they dispersed completely and randomly in the protein matrix.^{18,40} However, aggregation of clay platelets started to be visible at 3, 5 and 10% (w/w of protein) addition levels in bentonite and SM-MMT samples. Similar results were observed in previous studies on nanoclay and Al_2O_3 dispersed PVA adhesives where aggregation of nanoclay platelets were reported at concentrations greater than 4%.^{18,19} In terms of NCC and GO, exfoliation was observed up to 5% (w/w of protein) addition level whereas aggregation was visible at 10% (w/w of protein) addition level. The presence of surface hydrophilic groups such as $-\text{OH}$ and $-\text{COOH}$ in NCC and GO might be the reason for better exfoliation in canola protein matrix than those of bentonite and SM-MMT.

X-ray diffraction of nanomaterial dispersed canola protein adhesives are shown in Fig. 4. XRD patterns of dispersed nanomaterials supported the results observed in TEM. In a situation where nanomaterials are properly exfoliated in the matrix, crystalline peaks of original nanomaterial should not be visible since exfoliated nanomaterial could not generate identical peaks due to the absence of similar crystal lattice.^{18,40} XRD patterns of bentonite and SM-MMT exhibit a similar trend where the exfoliation of nanomaterials was observed only in 1%

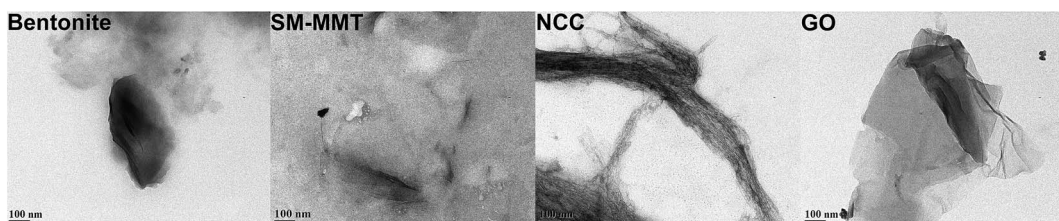


Fig. 2 Transmission electron microscopic images of bentonite, SM-MMT (surface modified montmorillonite), NCC (nanocrystalline cellulose) and GO (graphite oxide) used for adhesive preparation.

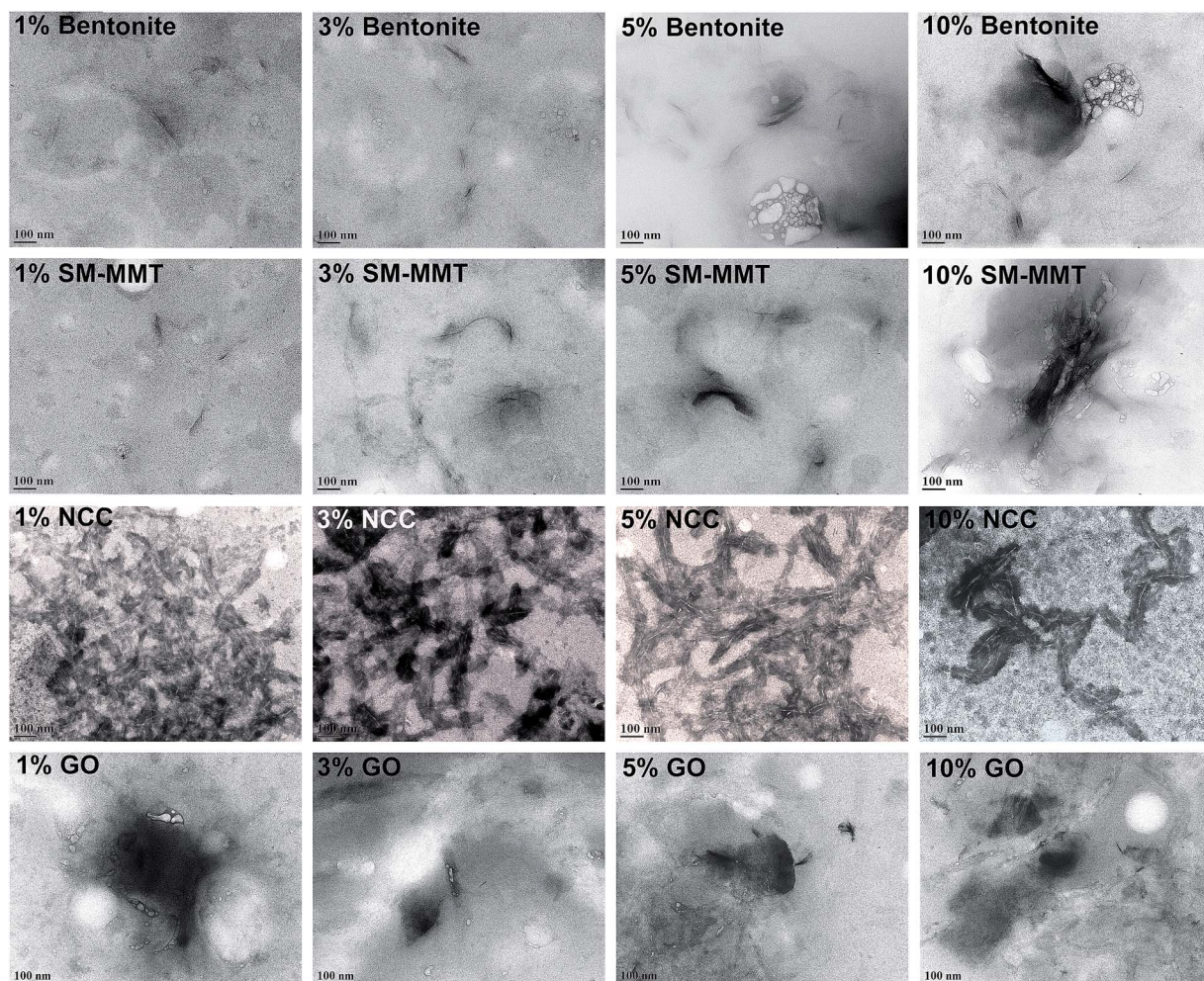


Fig. 3 Transmission electron microscopic images of canola protein adhesives after exfoliating 1%, 3%, 5%, and 10% (w/w nanomaterial/canola protein) levels of bentonite, SM-MMT (surface modified montmorillonite), NCC (nanocrystalline cellulose) and GO (graphite oxide).

(w/w of protein) addition level. Characteristic nanoclay peaks arising at 5.6° , 19.7° , and 34.8° start to appear in bentonite and SM-MMT incorporated adhesives after increasing the nanomaterial addition up to 3% or above. This can be due to partial exfoliation of nanomaterials or aggregation of nanoclay platelets at higher concentrations. Kaboorani and Riedl (2011, 2012) observed similar trend in XRD patterns where, nanomaterial loading above 4% exhibit crystalline peaks similar to their original nanomaterials in montmorillonite and nano Al_2O_3 dispersed in PVA adhesives.^{18,19}

In comparison, NCC and GO show better exfoliation in canola protein as original NCC and GO crystalline peaks were not visible in XRD patterns up to 5% addition level. However, at 10% (w/w of protein) addition level, both NCC and GO exhibit respective crystalline peaks in XRD patterns of prepared adhesives. In a previous study with NCC reinforced PVA adhesive, NCC showed exfoliation only up to 3% addition with improved wet adhesion than PVA adhesives reinforced with nanoclay.^{6,18} The improved solution intercalation method we used to disperse nanomaterials in canola protein might be the reason



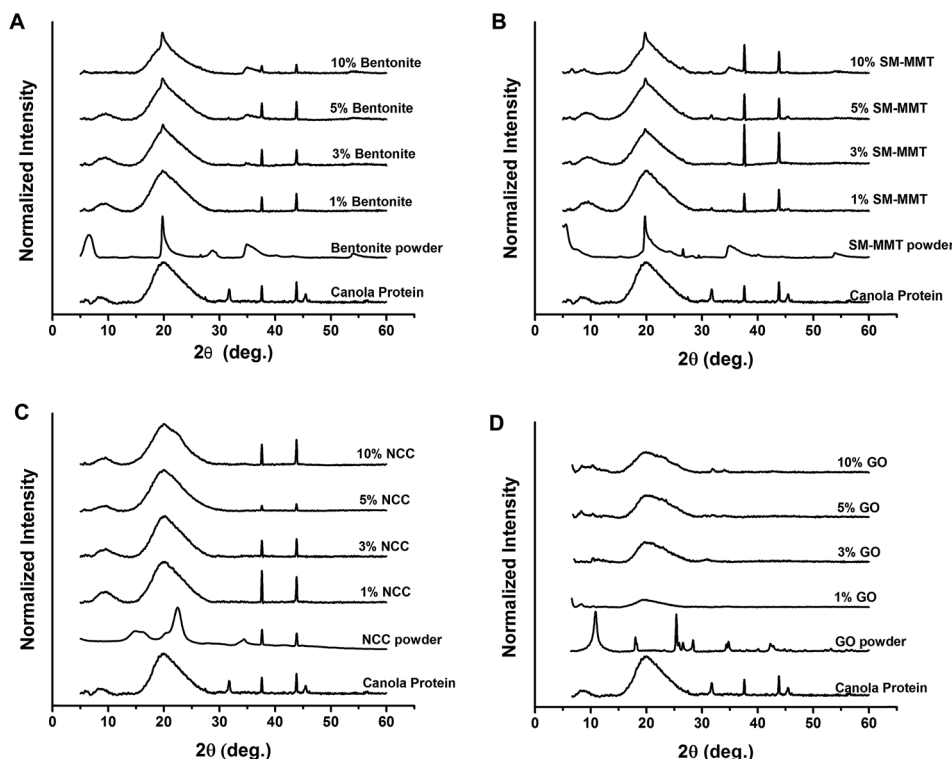


Fig. 4 X-ray diffraction patterns of canola protein adhesives after exfoliating 1%, 3%, 5%, and 10% (w/w nanomaterial/canola protein) levels of bentonite, SM-MMT (surface modified montmorillonite), NCC (nanocrystalline cellulose) and GO (graphite oxide).

for improved exfoliation observed in NCC and GO up to 5% addition level.

Effect of nanomaterial type and their concentration on adhesion strength

Effects of different nanomaterials and concentrations on adhesion strength are shown in Fig. 5. Adding nanomaterials at low concentrations significantly improved adhesion strength compared to both pH and negative controls. Bentonite significantly increased dry strength from 6.38 ± 0.84 MPa to 7.65 ± 1.33 MPa at 1% (w/w) addition and to 8.50 ± 1.27 MPa at 3% (w/w) addition (Fig. 5A). The wet strength was also increased from 1.98 ± 0.22 MPa (pH control) to 2.80 ± 0.50 MPa and 2.44 ± 0.29 MPa at 1% and 3% (w/w) addition respectively. However, the soaked strength was not affected by bentonite addition. A similar trend was observed with the addition of SM-MMT (Fig. 5B). At 1% SM-MMT addition, the dry, wet and soaked strengths were significantly increased up to 9.29 ± 1.53 MPa, 3.19 ± 0.57 MPa, and 6.87 ± 1.29 MPa respectively. However, the strength values were reduced at increasing bentonite/SM-MMT addition levels, which may be due to partial exfoliation or aggregation of nanomaterials at higher concentration as evidenced by TEM and XRD. Aggregation of nanomaterials at higher concentrations were previously reported in metal oxide nanomaterials and carbon nanotubes in different matrixes, mainly due to interaction with functional groups in polymer.^{41,42} Similarly, aggregation of bentonite and SM-MMT might reduce

the functional groups available for interacting with wood surface, thereby decreasing adhesion strength.

In comparison, both NCC and GO exhibited a better exfoliation even under higher addition levels up to 5% as evidenced by TEM and XRD, which was in good agreement with improved adhesion strength with NCC and GO addition (Fig. 5C and D). NCC significantly increased both dry and wet strength (10.37 ± 1.63 MPa and 3.57 ± 0.57 MPa respectively) at 1% (w/w) addition level while the highest soaked strength (8.98 ± 1.15 MPa) was observed at 3% (w/w) NCC addition. Unlike bentonite and SM-MMT, all tested NCC and GO addition levels significantly increased the adhesion and water resistance compared to negative and pH control samples. The highest dry and soaked strength for GO (9.27 ± 1.24 MPa, 7.78 ± 0.45 MPa respectively) was observed at 5% (w/w) concentration whereas the highest wet strength for GO (3.25 ± 0.36 MPa) was observed at 1% (w/w) concentration. Adhesive prepared by our exfoliation method exhibited significantly higher water resistance and adhesion than that prepared by the method reported by Zhang *et al.* (2014) (Table S1: ESI†).

Our study further supported the importance of uniform dispersion/exfoliation of nanomaterial in adhesive application, which was in good agreement with previous reports in improving mechanical properties of polymer matrix.^{6,18,19} The adhesive strength of soy protein adhesive was not improved when SiO_2 nano particles was not homogenously dispersed and exfoliated.⁴⁰ Adding high level of montmorillonite could even reduce the adhesive strength, due to a nano scale blocking



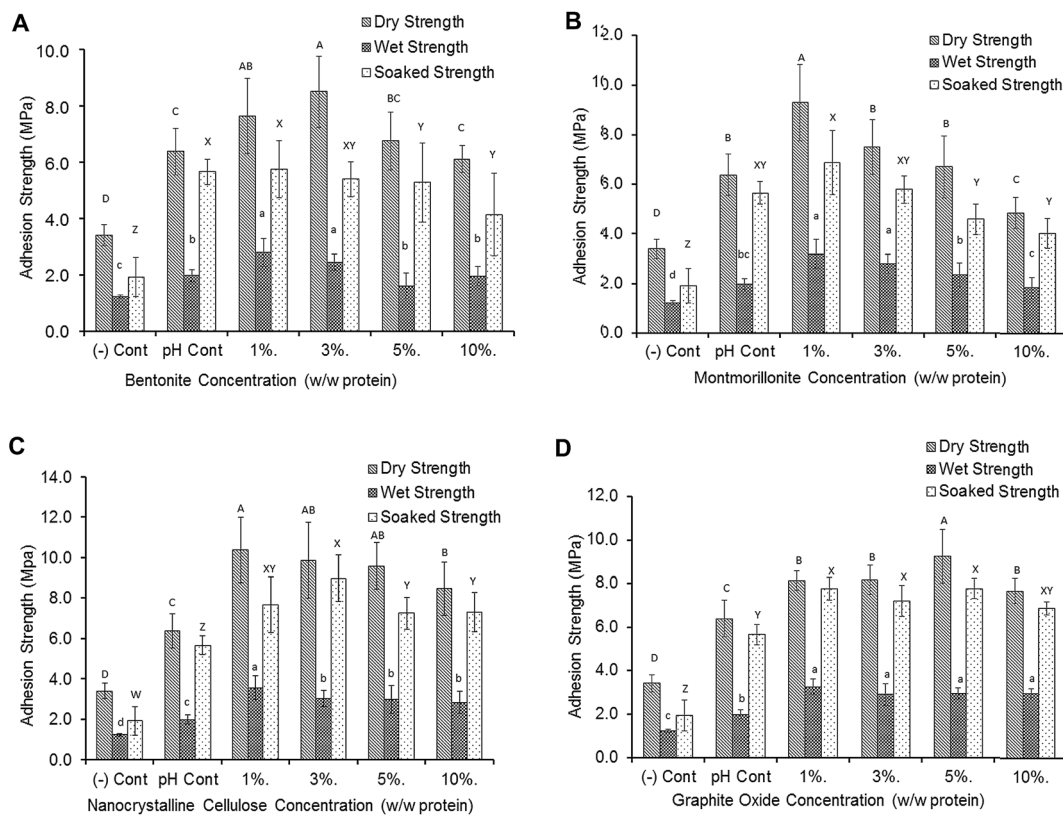


Fig. 5 Adhesion strength of nanomaterial exfoliated canola protein adhesives after exfoliating 1%, 3%, 5%, and 10% (w/w nanomaterial/canola protein) levels of bentonite, SM-MMT (surface modified montmorillonite), NCC (nanocrystalline cellulose) and GO (graphite oxide). All adhesive samples were prepared in triplicate ($n = 3$) and minimum 5 wood samples per replicate were used for each strength measurement. Adhesion data was analyzed using one-way ANOVA followed by Duncan test for mean separation for dry, wet and soaked adhesion separately. Different letters on the bar represent significantly different adhesion strength ($p < 0.05$).

mechanism.²⁰ Zhang *et al.* (2014) suggested that enhanced interactions between MMT and soy protein could make functional groups unavailable for reacting with wood surface, thus reducing adhesion strength. Adhesion between adhesive-wood interface results from various interactions including chemical bonding and mechanical interlocking.⁴³ The exfoliated nanomaterials have the potential to affect both chemical bonding and mechanical interlocking thereby increasing adhesion. The presence of nanomaterials in an exfoliated state in the adhesive matrix could act as a physical barrier for water penetration;³ it may also improve cohesion by increasing protein–protein interactions as a cross linker^{3,20,21} which ultimately improves wet and dry adhesion strength. In addition, adding nanomaterials would also induce protein structural changes by exposing hydrophobic and other buried functional groups⁴⁴ enabling reaction with functional groups present in surface and inner layers of the wood during adhesive penetration and subsequent curing.

Effect of nanomaterial on structural changes in canola protein based wood adhesives

Effects of nanomaterial addition on protein secondary structure are shown in Fig. 6. Protein secondary structural changes after modifications can be identified by processing amide I peak,

typically generated by C=O and C–N stretching vibrations at 1600–1700 cm^{-1} wavelength.^{45,46} The CPI pH control samples are predominated by β sheet structure^{45,46} with fitted peaks allocated at 1626 cm^{-1} , 1639 cm^{-1} , and 1676 cm^{-1} in the second derivative spectra, followed by α -helix and turns with peaks found at 1657 cm^{-1} and 1695 cm^{-1} respectively⁴⁵ (Fig. S2 – ESI†). In all nanomaterial added samples, in particular at concentrations over 3%, there is an increase in unordered structure as the peak at 1641 cm^{-1} increased.^{45,46}

At increasing bentonite addition, the relative proportion of β sheet structures (1628 cm^{-1} , 1676 cm^{-1}) was decreasing while that of unordered structures (1641 cm^{-1}) was increasing (Fig. 6a). Similar trend was also observed with SM-MMT addition (Fig. S2 and S3 – ESI†). In addition, at high nanomaterial concentrations, a peak shift from 1695 cm^{-1} towards 1691 cm^{-1} also represent turns in secondary structure of modified proteins.⁴⁵ In both type of nanoclay, increasing nanomaterial did reduce the relative proportion of α -helical structures as well (Fig. S2 and S3 – ESI†).

NCC addition reduced the relative proportion of β sheet structure (1628 cm^{-1} and 1675 cm^{-1} wavelengths) while the proportion of α -helical structure (1657 cm^{-1}) increased at high NCC concentrations (Fig. 6c). Unlike nanoclay, the unordered structure (1641 cm^{-1}) was not visible in second derivative



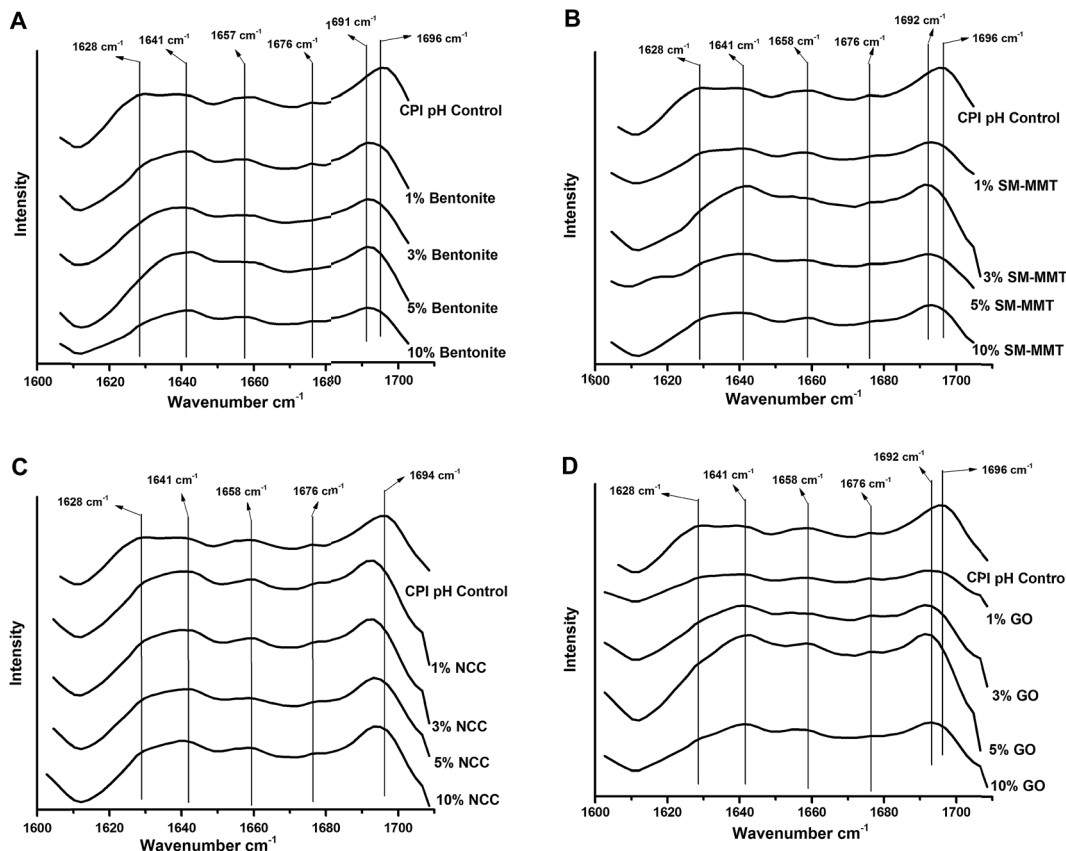


Fig. 6 Second derivative spectra of amide I peak in FTIR spectra showing protein secondary structural changes of adhesives prepared either with canola protein (CPI pH control), or by exfoliating bentonite, SM-MMT (surface modified montmorillonite), NCC (nanocrystalline cellulose), and GO (graphite oxide), at different nanomaterial addition levels (1%, 3%, 5%, and 10% w/w nanomaterial/canola protein).

spectra after peak fitting until concentrations up to 3% or above. At increasing GO concentrations, increase in unordered structure (1641 cm^{-1}) was more obvious compared to other nanomaterials, mainly at the expense of the relative proportions of α -helix and β sheet structure. Nanomaterial induced protein secondary structural changes were observed in previous studies as well^{44,47,48} where they reported decreased α -helix and β sheet structures⁴⁸ but increased β turns and unordered structures.^{44,48} These changes were attributed to the protein nanomaterial interactions such as nanomaterial induced the protein–protein interactions and exposed hydrophobic functional groups as a result of protein nanomaterial interactions, which was packed in core of protein structure.⁴⁴

Results obtained from FTIR analysis of modified adhesives support the trends observed in the adhesion strength of nanomaterial incorporated canola protein adhesives. At lower nanomaterial concentration, changes in secondary structure would expose more hydrophobic functional groups and enhance interactions with the wood surface thereby increasing adhesion strength and, specifically water resistance. Increasing nanomaterial concentrations, however, would lead to drastic change of secondary structure and promote strong nanomaterial–protein interactions, thereby reducing the potential functional groups available to react with wood surface,²⁰ which ultimately reduce adhesion strength and water resistance.

Therefore choosing an appropriate level of nanomaterial addition into canola protein matrix is important in improving adhesion strength and water resistance of canola protein adhesives.

Effect of nanomaterial on thermal properties of adhesives

Effects of nanomaterial additions at different concentrations on the thermal stability and enthalpy required for adhesive denaturation were shown in Table 1. Denaturation temperature (T_d) of a protein is an indication of thermal stability of protein.⁴⁹ Extracted canola protein used for this study exhibits an onset temperature of 72.20 ± 0.16 °C and T_d value of 90.91 ± 2.09 °C, which was comparable to previously reported onset temperature of 77.9 °C and denaturation temperature of 83.9 °C, by Wu and Muir (2008). The slight variations of denaturation temperatures are attributed to the method of protein extraction and compositional changes of extracted protein, which has an effect on thermal stability of the protein.⁴⁹ Thermal stability is one of the properties required in developing wood adhesives where hot pressing is required in adhesive curing. In general, adding nanomaterials increased the T_d of CPI adhesives compared to the controls which could be due to strong protein–nanomaterial and protein–protein interactions. Linse *et al.* (2007) attributed increased thermal stability and denaturation temperatures of



Table 1 Changes in thermal transitions of canola protein based adhesives after exfoliating bentonite, surface modified montmorillonite (SM-MMT), nanocrystalline cellulose (NCC) and graphite oxide (GO) at different nanomaterial concentrations (1%, 3%, 5%, and 10% w/w nanomaterial/canola protein)

Sample	Onset T° (°C)	Peak T° (°C)	Specific heat (J g ⁻¹ C ⁻¹)
CPI – control	72.20 ± 0.16	90.91 ± 2.09	1.44 ± 0.03
CPI pH control	88.55 ± 4.82	103.16 ± 4.74	2.53 ± 0.02
1% bentonite	86.25 ± 2.62	103.79 ± 2.28	2.69 ± 0.12
3% bentonite	91.75 ± 0.69	107.42 ± 0.18	2.60 ± 0.07
5% bentonite	90.32 ± 1.75	105.58 ± 2.51	2.20 ± 0.01
10% bentonite	87.69 ± 1.53	103.35 ± 2.14	2.14 ± 0.06
1% SM-MMT	85.98 ± 2.04	102.54 ± 2.41	2.32 ± 0.02
3% SM-MMT	85.80 ± 1.32	100.53 ± 1.61	1.92 ± 0.03
5% SM-MMT	78.22 ± 3.48	93.02 ± 4.82	2.12 ± 0.15
10% SM-MMT	84.34 ± 5.18	98.86 ± 3.44	1.84 ± 0.04
1% NCC	91.84 ± 1.73	105.15 ± 2.14	2.43 ± 0.04
3% NCC	85.40 ± 2.40	100.63 ± 2.82	2.12 ± 0.03
5% NCC	87.82 ± 1.28	101.69 ± 1.40	1.55 ± 0.06
10% NCC	84.34 ± 5.18	104.18 ± 0.08	1.56 ± 0.05
1% GO	79.17 ± 0.83	96.02 ± 0.88	2.88 ± 0.09
3% GO	80.66 ± 0.47	97.62 ± 1.32	2.52 ± 0.01
5% GO	82.39 ± 0.18	98.20 ± 1.64	2.11 ± 0.06
10% GO	80.50 ± 1.03	95.55 ± 2.94	1.48 ± 0.02

graphite oxide nano sheet incorporated soybean peroxidase to protein secondary structural changes.⁴⁷ However, further increasing nanomaterial concentrations resulted in decreasing T_d values of CPI adhesives. This can be a result of drastic changes in protein secondary structures as evidenced by FTIR data, where they created higher degree of unordered structures, resulting lower temperature requirement for denaturation.

Conclusions

A solution intercalation method was developed to exfoliate nanomaterials in canola protein matrix as evidenced by TEM and XRD analysis. Our study showed that nanomaterials at lower addition levels (at 1% w/w addition) could significantly improve the adhesion strength and water resistance of canola protein adhesives. However, decrease in adhesion strength at increasing nanomaterial addition levels were observed with the exception of NCC and GO, where adhesion was improved even at 3% and 5% w/w levels, respectively. Our study further supported the significance of uniform dispersion and exfoliation of nanomaterial in the protein matrix. Adding nanomaterials exposed more hydrophobic and other functional groups to react with wood surface, which would increase water resistance and adhesion strength. The improvement could also be attributed to the nanomaterial-induced cohesion. In addition, the properly exfoliated nanomaterials could act as physical barriers for water penetration, contributing to improved water resistance. Among four nanomaterials tested in this study, GO and NCC proved to be superior in terms of increasing functionality of canola protein adhesives compared to bentonite and SM-MMT. Results of this study provided evidence on the potential use of

nanomaterial to improve the adhesive properties of biobased wood adhesives, which may replace traditional synthetic adhesives as green alternative adhesive materials.

Conflict of interest

The authors declare no financial competing interest.

Abbreviations

Bento	Bentonite
SM-MMT	Surface modified montmorillonite
NCC	Nanocrystalline cellulose
GO	Graphite oxide
PVA	Polyvinyl acetate
XRD	X-ray diffraction
DSC	Differential scanning calorimetry
TEM	Transmission electron microscopy

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