Regenerated Silk and Carbon Nanotubes Dough as Masterbatch for High Content Filled Nanocomposites

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Regenerated silk (RS) is a natural polymer that results from the aggregation of liquid silk fibroin proteins. In this work, we observed that RS dispersed in aqueous solution undergoes a reversible solid/liquid transition by programmed heating/cooling cycles. Fourier transform infrared, atomic force microscopy imaging and Raman measurements of the RS reveal that the transition from random coil to β-sheet structures is involved in this liquid–solid transition. The reversible solid-liquid transition of silk fibroin was then found to be helpful to prepare polymer-like carbon nanotube (CNT) dispersions. We demonstrate that the gelation of RS makes the CNTs with the consistency of a dough with polymeric behavior. Such RS can disperse carbon nanotubes at high concentrations of tens of weight percent. Finally, such carbon nanotube dough has been used for the realization of rubber composites. With this method, we pave the way for handling nanopowders (e.g. CNTs or graphene related materials) with safety and reducing the filler volatility that is critical in polymer-processing.

Keywords: carbon nanotubes, regenerated silk, phase transitions, electrical conductivity, polymer composites

INTRODUCTION

Regenerated silk (RS) is a natural polymer made by the coagulation of silk fibroin that is a an aggregation of proteins with short and long chains; the combination of intermolecular interactions via hydrogen bonds between the chains leads to the formation of β-sheet structures that have a high crystalline local order. Increasing the fibroin concentration in a solution enhances the probability to generate chain interactions where the β-sheets form a stable gel transition. The dehydration of such crystalline structures results in an irreversible liquid-solid transition due to thermodynamic cross-linking of the β-structures that undergo gelation with time (Ayub et al., 1993; Hanawa et al., 1995; Kang et al., 2000; Wang et al., 2008). Some physical and/or chemical methods have been proposed to enhance the gelation kinetics of silk fibroin, including pH change, mechanical sonication and addition of salts (Matsumoto et al., 2006; Gong et al., 2012; Wu et al., 2012; Kapoor and Kundu, 2016). A reversible sol-gel transition of silk fibroin exposed to acidic or basic vapors and, more recently, in hydrogel-based silk fibroin have also been reported (Terry et al., 2004; Yin et al., 2017).

Liu et al. (2014) demonstrated the thixotropy (i.e., a time dependent shear thinning property) of silk fibroin via dissolution nanofibrils gel in alcohol and sodium chloride while Bai et al. (2014) also showed a reversible sol—gel transition by self-assembly of nanofibers into supramolecular
aggregations without altering the β-sheet contents. Numata et al. (2011) reported the gelation of the regenerated silk induced with ethanol and with a heterogeneous network structure made of a β-sheet and fibrillar structures.

The shape, dimension and the aspect ratios over 1,000 of carbon nanotubes (CNTs) with their ability to undergo large deformations without damage suggests that interesting parallels may be drawn with silk fibroin. However, attention turns to their behavior and potential processability in liquid suspension. There are a lot of experiments done with most common solvents to get stable dispersions of CNTs (Fukushima et al., 2003; Bergin et al., 2009; Davis et al., 2009); however, generally the most common solvents used, such as N-methyl-2-pyrrolidone, dimethylformamide, and 1,2-dichlorobenzene, can disperse only CNTs with specific size and dimension at low concentrations. Recently, Chiou et al. (2018) dispersed carbon nanotube powders in m-cresol at very high concentrations giving them a dough-like consistency. However, m-cresol is highly volatile, and the processing requires a controlled environment.

In this study we exploit the utilization of salts to dramatically reduce the gelation time of silk fibroin. The results demonstrate that regenerated silk drop solution can undergo liquid-solid transitions within a few min by thermal annealing. The proposed method induced a rapid gelation of CNTs that, when added to RS solution at high concentrations up to tens of weight percent, show a transition dough state that exhibit polymer-like and viscoelastic properties. Finally, as proofs of the concept, such polymer-like CNTs were used as masterbatch to realize a rubber composite.

MATERIALS AND METHODS

For the preparation of RS film, commercial B. mori silk cocoons were boiled for 1 h in distilled water solution of 0.025 wt% NaHCO₃, rinsing with distilled water every 30 min to remove the sericin. The degummed silk (i.e., 0.23 g) was then added to a CaCl₂ (i.e., 24 g)—water (i.e., 68 ml)—ethanol (i.e., 55 ml) solution and stirred 12 h at 40°C yielding an 1 wt% solution. NANOXYL® NC7000™ multi-walled carbon nanotube powder (average diameter 9.5 nm, average length 1.5 μm, volume resistivity 10⁻⁴ Ohm·cm) was used. In 1 ml of solution, 10 mg of CNTs up to 50 and 2 mg of regenerated silk were dispersed to yield, at maximum, a RS concentration of about 4 wt%. The liquid-solid transition was monitored by the optical density (OD) method, measuring the absorbance at wavelength 550 nm and temperature of 80°C at different times. Fourier transform infrared (FTIR) analysis was performed in a Jasco FTIR FT/IR-615 spectrometer, equipped with an ATR mode in the wave number range from 400 to 4,000 cm⁻¹. The spectra were deconvoluted by firstly smoothing the signal with a polynomial function with a 15-point Savitski—Golay smoothing function, subtracting a linear baseline and applying Gaussian deconvoluting curves by Origin 9 software. Rheological data for the RS/CNT dough were recorded on circular specimens (ca. 12 mm wide, and 0.13 mm thick) on an ARES rheometer (Rheometric Scientific). Frequency and strain sweeps, as well as constant strain amplitude, were performed at constant temperature to find the storage modulus (G′) and the loss modulus (G″) of RS and RS/CNT samples at different CNT concentrations. The electrical resistance of the RS/CNT solid dough was obtained with a Keithley 6517B electrometer/high resistance meter equipped with an 8009 test fixture, according to the ASTM D257 that is the worldwide laboratory standard for sensitive measurements. Briefly, the basic method used to determine the resistance is a two-step method: first a test voltage is applied to the sample and the subsequent current is measured; then, the test voltage value and measured current value are applied and the electrical resistance is calculated.

For the atomic force microscopy (AFM) and Raman characterization, dispersion of RS and RS/CNT was drop casted on silicon wafers and glass slides. The sample was heated at 80°C for 1 h in air conditions. The solid samples were placed in AFM chamber which is an insulated box to minimize environmental noise and building vibration at room temperature. AFM instrument (Model: Bruker Dimension Icon) operated at intermittent contact mode using silicon cantilever (Model: ScanAsist). All the AFM images are produced as a function of time by keeping the other parameters (i.e., scan speed, scan size) constant.

Raman spectroscopy was carried out at room temperature by using 50X objective lens using LASER source of 532nm at 5mW and grating 2400 l/mm. The spectral range was analyzed between 200cm⁻¹ to 3000cm⁻¹. The Raman spectra were recorded as a function of displacement (i.e. line map) and function of time (i.e. collection of spectra at single region at different time intervals).

For the realization of the polymer composite, an acrylonitrile butadiene rubber (NBR; KNB35), LEVAPREN 700 and vulcanizing agent LUPEROX® F40MF were used. Rubber compounds were prepared by dissolving separately 1.6 g of NBR, 3.1 g of LEVAPREN and 0.25 g of LUPEROX in chloroform and then mixing together with RS/CNTs dough with a DISPERMAT® and left to evaporate onto a Teflon mold until the complete evaporation of the solvent was achieved. Carbon nanotubes powder, for a comparison purpose, was sequentially added to the rubber by melt mixing (Banbury mixer) at about 100°C for 10 min. The vulcanizing agent was added just before the extraction of the composite to avoid any cross-linking effect. The composites prepared with both procedures were then vulcanized at 180°C for 5 min in a thermofluid-heated press.

Tenstile properties were measured according to ASTM D 412 specifications, on an Instron dynamometer (Model 4301), at 25°C at a crosshead speed of 500 mm min⁻¹. At least five specimens of each sample type were tested. The samples were then cut into dumbbell specimens Die A with dimension of ~140 × 25 × 2 mm.

RESULTS AND DISCUSSION

The liquid RS extracted from solution and a drop cast on silicon substrate at 80°C shows a liquid–solid transition (Figure 1a) that recovers the liquid state under cooling at 25°C within few minutes, which is lower than that of conventional silk-fibroin
FIGURE 1 | (a) Photos showing the reversible liquid-solid transition observed along the heating/cooling time of RS. (b) ATR-FTIR spectra in the amide I region for RS liquid solution annealed at 80°C at different times. (c) ATR-FTIR spectra in the amide I region for RS solid phase cooled at room temperature at different times. (d) AFM topography of solid silk fibroin imaged at different time intervals at room temperature are showing dendritic structure of silk fibroin with several branches. The inset represents a peculiar region analyzed at different time intervals. AFM topography of a zoomed region analyzed up to 25 min; a transformation in the silk-fibroin structure is marked by dashed circle. The scale bar represents the degradation of the structure as a function of time. (e) AFM image of the dendritic fibroin network; the dashed line represent the topographic profile of different altitude of the fibroin structure.
FIGURE 2 | (a) Photos showing the reversible liquid-solid transition observed along the heating/cooling time of RS/CNTs. (b) ATR-FTIR spectra in the amide I region for RS/CNTs liquid solution annealed at 80°C at different times. (c) Annealing time effect on β-sheet content at 80°C assessed by FTIR amide I band analysis for RS

Continued)
The liquid-solid transition of fibroin results from the formation of physical cross-links made of a β-sheet that makes the optical aspect of the film more opaque. In its liquid state, fibroin is a hydrophobic molecule surrounded by ordered water. In this state, the ordered structure of the water hinders the fibroin aggregation but when the temperature increases, the positive entropic term overpowers that of heat or enthalpy and local dehydration takes place, enhancing the hydrogen bonding between the chains (Connelly et al., 1964). Increasing the annealing time provides the increased chain interactions through dehydration of the system (Urry et al., 1997), and thus accelerates the solidification.

The RS also exhibits a transition from the solid to the liquid state by setting the substrate at room temperature (Figure 1a). Although silk fibroin contains a high content of β-sheet in the solid state, the return to the liquid phase may also imply the existence of metastable fibroin structures within the β-sheet fraction, likely those composed of silk fibroin and Ca²⁺ ions (Ling et al., 2016). Calcium ions can capture water molecules from the atmospheric and hence, the more calcium ion is in silk fibroin solution, the more water molecules, as a solvent, can be captured, resulting in liquid transition.

Changes in the structure of the RS were detected by FTIR analysis. ATR-FTIR spectra of the RS in the amide I region were measured at different times of annealing, as reported in Figure 1b. The β-sheet (crystalline) content was determined by the deconvolution of the amide I region and by estimating the ratio between the peak area in the wavenumber region of 1622–1637 cm⁻¹, which is the main absorbance region of β-sheet crystal in amide I (Hu et al., 2006), and the whole area of the amide I region comprising the peaks of the structural components, including turns and random coils. The dashed lines mark the adsorption bands at 1650 and 1625 cm⁻¹, which are characteristic of the random-coil and β-sheet structures of silk proteins, respectively, according to previous studies (Dong et al., 1990; Goormaghtigh et al., 1990; Mourou et al., 1997; Jung, 2000; Teramoto and Miyazawa, 2005; Tretinnikov and Tamada, 2001). The structure of RS shifted predominantly to a β-sheet structure by increasing the annealing time (Figure 1b), whereas random-coil structures are prevalent in the liquid state and vice-versa by cooling at 25°C (Figure 1c). The β-sheet content (calculated from the amide I bands over the amide I region comprising the peaks of the structural components) in the liquid state was unexpectedly high (≈15%). Moreover, the β-sheet fraction increases with increasing the annealing time and was as high as 45% when the film was completely dry.

Figure 1d shows the images of single fibroin molecules obtained by AFM. The concentration of the RS sample was 1 wt % . According to Inoue et al. (2000), the silk fibroin molecule consists of a rodlike part with stringlike parts on each end of the rod. The observed size is 100 nm in width and 14 nm in height (Figure 1e). According to the ATR-FTIR analysis, the silk fibroin shows the disappearance of the dendritic structure at different time intervals at room temperature (Figure 1d).

The RS/CNT solution, which demonstrated a similar behavior (Figure 2a), showed a higher β-sheet content of about 60% once dried (Figures 2b,c). More indicative of such transition is the rheological behavior. In Figure 2d, the plots of storage (G’ and loss (G’’) modulus vs. annealing/cooling cycles (increasing/decreasing values) are shown. According to the interpretation of the FTIR spectra both G’ and G’’ increase with annealing time; interestingly both G’ and G’’ are independent from the strain for the RS sample, instead they decrease with increasing the strain amplitude for the RS/CNTs sample (Figure 2e). For viscoelastic systems filled with rigid solids, this is known as the Payne effect (Payne, 1962). According to this theory, solid fillers may exist in chainlike formation or aggregates within the solid polymer. If there are attractive forces, the particles are held together in single agglomerates that behave like a rigid group under deformation. However, as the magnitude of the stress increases with increasing deformation, these groups will be broken down into smaller units and the elasticity of the material will change. Then, the number of connections N rescales as:

\[ N = N_0 [1 + (\gamma / \gamma_c)^2m]^{-1} \]  

where, \( N_0 \) is the initial connection density, \( m \) is the network structure factor, \( \gamma \) and \( \gamma_c \) are the strain and the yield strain, respectively. This leads to the equation:

\[ G’ = G’_\infty + (G’_0 - G’_\infty) / [1 + (\gamma / \gamma_c)^2m] \]  

where \( G’_0 \) or \( G’_\infty \) are the storage moduli at low or high strain, respectively. The best fit of the data (e.g., 0.53) reported in Figure 2e with Equation 2 results in a structure factor close to \( m = 0.5 \), which is typical for filled elastomers. From the rheological study, we can assume that the RS/CNTs sample behaves as a viscoelastic material.

RS can process CNTs up to tens of weight percent; the photos in Figures 2a, 3a show the liquid CNT solution and its dough state as the annealing time in RS was increased. After drying at 80°C, a disk-shaped sample shrinks isotropically maintaining its shape, resulting in a stiff solid object that could be further handled (Figure 3a). Upon cooling, the dough-like state remains stable for at least 20 min (Figures 2a, 3a). As reported in the FESEM images of Figure 3b, the CNTs are well-dispersed by the RS. From this analysis, we suggest that the interaction between silk fibroin and CNTs allows their dispersion.
FIGURE 3 | (a) The CNTs in RS solution can be processed to obtain a dough. Once annealed, the RS/CNT dough shows a contraction and can be taken by hands as a freestanding solid sample. (b) Morphology of RS/CNTs dried sample in FESEM showing well separated CNTs after being processed in RS. (c) Optical image of...
In order to confirm this hypothesis, we reported the Raman characterization of RS and RS/CNT samples in Figures 3c–h. Raman spectra of the regions reported in Figure 3c show the typical Raman features of CNTs i.e., the so-called G-line that is a characteristic feature of the graphitic layers, the defective graphitic structure (D-line), the radial breathing mode (RBM) and the resonant 2D mode (Dresselhaus et al., 2005) (Figure 3d).

Figures 3e,g show the Raman spectra of solid RS, analyzed as a function of time at room temperature. The spectra of Figure 3e are similar to those already published (Rousseau et al., 2004) and the assignment of the major bands observed around 600, 800, and 1100 cm$^{-1}$ are based on published Raman results on silk, proteins, and polypeptides (Rousseau et al., 2004). The sub-peaks (Figures S1–S3 in the Supplementary Material), due to the C-C
stretching vibration at around 1,103 cm$^{-1}$ in the spectrum of the RS, show a blue-shift as a function of time that is associated with the β-sheet reduction, as suggested by Zheng et al. (1989). On the contrary, after addition of CNT in the RS (Figures 3f,h), the peak positions at 1,087, 1099.85, and 1113.65 cm$^{-1}$, show a red-shift as a function of time, illustrating a significant contribution from CNT.

The transition to solid state has also been confirmed by rheological, and viscoelastic properties. The transition to polymer-like state suggests that the nanotubes form a cohesive network in RS (Payne, 1962). After annealing, above 10 mg/mL, the CNT network is not free to flow, leading to a freestanding dough (Figure 3a). It becomes solid-like with increased storage modulus (Figure 4A). The loss modulus decreased more slowly than the storage modulus, giving the dough a sufficient level of viscous character for extrusion. In view of the similarity of our RS/CNTs dough with a viscoelastic medium, as reported in Figure 2e, the elastic properties of the samples were modeled by applying the scaling law:

$$\gamma_c \approx f^{-(1+d_f)/(3-d_b)}$$

which have been successfully demonstrated for polymer gels (Shih et al., 1990; Shaffer and Windle, 1992; Boland et al., 2016). These studies showed that both the storage modulus $G'$ and the limit of linearity of the strain $\gamma_c$ exhibit a power-law with respect to particle concentration $(f)$ that is $G' \approx f^{4.1}$ and $\gamma_c \approx f^{-2.1}$ (Sonntag and Russel, 1987; Buscall et al., 1988). Taking the $\gamma_c$ values from Figures 2e, 4A and plotting versus the CNT wt% in Figure 4A, we obtain the best fit with a power law with exponent $-2.22$. Such a value is consistent with the rheological model developed by Shaffer and Windle (1992), who considered the structure of a gel network as a collection of fractal flocs, which are closely packed throughout the sample with $d_f$ and $d_b$, indicating the fractal dimensions of the network and its backbone, respectively. They observed that in the strong-link regime, where there is an interfloc interaction, the elastic constant (i.e., $G'$) increases but the limit of linearity (i.e., $\gamma_c$) decreases with increasing particle concentration.

The higher CNT concentrations of this dough-like state are also accompanied by a threshold in the electrical properties. For example, the increase of the CNT concentration from $\approx 1$wt% to $\approx 10$wt% was accompanied by the onset of electrical conductivity (Figure 4B), which can be attributed to the start of a percolation threshold, through an interconnected pathway of nanotubes.

The dough state is of potential interest for polymer processing (De Gennes, 1979; Huang et al., 2012; De Volder et al., 2013). The dough can be added to the polymer matrices as masterbatch without handling powders, moreover, the dough state eliminates the use of solvents. Figure 5 shows the mechanical results of the experiment, where rubber composite has been prepared by blending NBR with RS/CNTs dough. The nanocomposite was then vulcanized under a hot plate press at 180°C. At 10 wt% loading of CNTs in NBR, the toughness modulus of the RS/CNT composite increased by $\approx 16\%$, in comparison with a similar composite prepared with CNT powder (Table 1). Such preliminary findings suggest that RS/CNTs dough could be used for the processing of polymer nanocomposites.

### CONCLUSIONS

In this study, the transition from solid to liquid state of RS was obtained by the addition of calcium chloride salts to water based silk fibroin solution. It was observed that the presence of salts stimulated the water absorption and this reaction led to a reversible transition. Moreover, the processing parameters including annealing time, allows the control of the molecular organization of silk fibroin in β-sheets structures. Such an inexpensive method to assemble silk in solid state was used for optimizing the dispersion of carbon nanotubes at high concentrations of tens of weight percent. The results show the advantage to use silk fibroin as a natural dispersant to obtain CNTs with the consistency of polymers. As a proof of concept, RS/CNT dough was used to prepare a polymer composite with improved deformability and toughness modulus, with respect to the composite prepared by the addition of CNT powder. Such findings pave the way the utilization of nanopowders dough as master batch in the processing of polymer composites and to overcome the problems of handling nanopowders in industrial implants including graphene and related materials.

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**TABLE 1** | Polymer nanocomposites (e.g. 10 wt%) mechanical properties (i.e., elongation at break, tensile strength, stress at 50 and 100% of strain and toughness modulus).

| Sample        | Elongation at break (%) | Tensile strength (MPa) | Stress at 50% (MPa) | Stress at 100% (MPa) | Toughness modulus (MPa) |
|---------------|--------------------------|------------------------|---------------------|----------------------|-------------------------|
| NBR           | 246 ± 10                 | 8.35 ± 0.39            | 1.5 ± 0.1           | 3.8 ± 0.2            | 10.27 ± 0.24            |
| NBR/CNTs      | 283 ± 18                 | 13.30 ± 0.44           | 2.0 ± 0.1           | 7.8 ± 0.4            | 18.81 ± 0.31            |
| NBR/RS/CNTs   | 344 ± 9                  | 12.38 ± 0.49           | 1.5 ± 0.1           | 3.8 ± 0.2            | 21.29 ± 0.30            |
AUTHOR CONTRIBUTIONS

All authors listed have made a substantial, direct and intellectual contribution to the work, and approved it for publication. In particular SB contributed to the RS and composites realization. MT AFM imaging and its corresponding discussion. AD Raman data and its associated discussion and critical revision.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmats.2019.00060/full#supplementary-material

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary Information

Figure S1: (a) Optical image of RS/CNT marked with different regions R1-R10 over glass substrate. (b) Raman spectrum correspond to the regions marked in panel (a). (c) Region 5 (green colour) is showing the presence of CNT peaks along with Silk fibroin.

Figure S2: Peak positions range between 1080 cm\(^{-1}\) to 1110 cm\(^{-1}\) of silk fibroin varies at different time interval from 0 minute to 21 minutes. The spectra obtained at t = 0 minute is used as reference for other panels distinctly indicating transformation in the silk fibroin with time.
**Figure S3:** Peak positions range between 1080 cm$^{-1}$ to 1115 cm$^{-1}$ of silk fibroin+CNT varies at different time interval from 0 minute to 28 minutes. The spectra obtained at $t = 0$ minute is used as reference for other panels.