## **Facile synthesis of chitin nanocrystal decorated on 3D cellulose aerogel as a new multi-functional material for waste water treatment with enhanced antibacterial and anti-oxidant properties**

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We report fabrication and application of multi-functional hybrid bio-aerogels based on cellulose nanofibers (CNF)and chitin nano crystals (CNC)developed through environmental friendly freeze-drying process. Herein, organic non-solvent approach such as acid hydrolysis was used to extract CNF and CNCfrom corn husk and shrimp shell respectivelyfollowed bycharacterisationusing IR spectroscopy and electron microscopy. Usual wired like and infrequent semi-square type of morphology were detected for CNF and CNCrespectivelyduring electron microscopic analysis. The aerogels with two different quantities of CNC on CNFwere

made (AR1 and AR2) and compared them with reference to the neat CNF aerogel (neat AR). Interestingly, for AR2, under the electron microscopy, a maple seed like morphology was observed compared to neat AR and AR1.Their ability in water treatmentwere determinedby conducting adsorption experimentusing methylene blueand Rhodamine 6G, found that AR2 has enhanced removal percentage and adsorption capacity. To addressing their economic viability, we conducted several rounds of studies and evaluated the reusability of AR2. Anti-bacterial and anti-oxidant activities of aerogels were studied using disc diffusion method against four bacterial species such as St. aureus, E. coli, S. typhimurium and B. Cereus and carefully discussed the best condition for the determination of minimal inhibitory concentration. The prepared material having higher CNC content (AR2) shows multi-functional behaviour and in the future further studies are anticipated to produce them in industrial scale for commercial application **19. Download** (AR1 and AR2) and compared them with reference to the near CNF accepted vect AR1, 37 Interestingly, for AR2, under the electron microscopy, a maple seed late morphology was<br>
38 observed compared to read A

#### **1. Introduction**

Government, industries, and customers are gradually demanding products developed from sustainable and renewable resources that are biodegradable, carbon neutral, non-petroleum, and have low environmental, human or animal health and safety risks. For the past thousand years natural, cellulose based materials have been used by our society as engineering materials and their use continues in paper production and in textile field world widely. Cellulose is a linear polysaccharidewhich contains glucose molecules in a ringed and flat ribbon like conformation. 54 The repeating unit of cellulose comprised of two anhydroglucose rings  $(C_6H_{10}O_5)_n$ ; n= 10000 to 15000, where n is depended on the cellulose source material linked together through an oxygen covalently to C1 of one glucose ring and C4 of adjoining ring (1→4 linkage) and so called the β  $1 \rightarrow 4$  glucosidic bond<sup>1</sup>. Cellulose mainly occurs as microfibrilated form having both amorphous and crystalline regions. The isolation of crystalline region and dissolution of amorphous region

makes cellulose into nanocrystals usually known as cellulose nanofibrilshaving similar properties of raw one. The low mechanical stability and charge less surface is main disadvantage of cellulose and cellulose nanofiber which can be overcome using reinforcement of other substances during the material processing time.Another polysaccharide known as chitin is a natural, renewable and biodegradable polymer, the second most abundant biopolymer after 64 cellulose<sup>2</sup> which can be found in exoskeleton shells of arthropods such as crabs, shrimps, beetles 65 etc and cell walls of fungi and yeasts $3-5$ . Chitin also a linear polysaccharide containing repeated units of β-(1→4)-2-acetamido-2-deoxy-β-D-glucose and β-(1→4)-2-amino-2-deoxy-β-D-67 glucose<sup>6</sup>. Chitin has two hydroxyl groups and an acetamide group makes chitin very crystalline 68 with surface charge and strong hydrogen bonding<sup>7</sup>. Similar to cellulose, chitin also occurs in nature as microfibrilated form having both amorphous and crystalline regions and these fibrils 70 are typically embedded in a protein matrix depending on the origin of sources<sup>8</sup>. The crystalline regions of chitin can be extracted by excluding amorphous region and it is known as chitin nanocrystals or nanowhiskerswhich is widely used as reinforcing nanoparticles. Several methods 73 can be employed for the extraction of chitin nanocrystals such as acid hydrolysis $8-17$ , mechanical 74 treatment<sup>7,18</sup>, electrospinning<sup>19</sup>, gelation<sup>20</sup>, and TEMPO-mediated oxidation<sup>21,22</sup>. Due to the diverse properties of chitin nanocrystals such as biodegradability, functionality, hydrophilicity, eco-friendly, and ease of processing, it is used in many field of application which includes  $\mu$  biomedical, water purification, packing and protein immobilization<sup>23</sup>. Recently Lin et. al reported fungus – derived chitin nanocrystals and their dispersion stability evaluation in aqueous media. In their study, chitin nanocrystals which avoided possible safety risks were extracted from mushrooms using protein/mineral purification and HCl hydrolysis. Such nanocrystals of **Solution** makes exillated into nanonystatis usually known as cellulese nanofficitioning similar properties<br> **Solution** on the ownloaded subsidity and charge less surface is main dissolvanting of<br> **Excelutions and cellule**  81 chitin's were in α-crystalline structure with a length and width of  $143 \pm 24$  and  $10 \pm 2$  nm 82 respectively<sup>24</sup>.

Water pollution from industrial as well as public effluents (dyes, heavy metals, oils etc) results in acute shortage of fresh water in many parts of the world in which dyes and heavy 85 metals can severely damage people and living things<sup>25</sup>. Consequently there is an increasing demand for advancedadsorptive materials having high selectivity, high efficiency and high 87 adsorption capacity<sup>25,26</sup>. The dye effluents are one of major contaminants in water. Different 88 types of adsorptive materials including activated carbon<sup>27</sup>, carbon nanotubes<sup>28</sup>, clays<sup>29</sup>, magnetic 89 oxides<sup>30</sup> etc are being used for the adsorption of dyes from waste water. Among them, 3D porous materials seem to be the most auspicious class of high performance adsorbent due to their high porosity, light weight and large surface to volume ratio. Because of these inherent physio-chemical characteristics, they can adsorb a large quantity of dye molecules from dye polluted waste water according to their surface charges. However, complex fabrication procedures and/or high raw materialcost have diminished their wide spread applicability. Therefore, there is every need to develop innovative, renewable/sustainable, eco-friendly and robust high performance adsorbent materials. **S2.** chitin's were in a-crystalline attneutie with a length and width of  $143 + 24$  and  $10 + 2$  am  $92$  respectively<sup>24</sup>.<br> **ACCEPT ACCEPT ACCEP** 

Xiong et.al demonstrated a facile and environmentally friendly approach to prepare  $F_{e3}O_4/Ag$  nanofibrillated cellulose nanocomposite aerogel having excellent catalytic properties for the reduction of 4-nitophenol. This prepared aerogels also showed high antibacterial activity 100 against model microbe *S. aureus*<sup>31</sup>. Zhou et. al prepared polyaniline (PANI) – decorated cellulose aerogel with strong interfacial adhesion and enhanced photocatalytic activity via 102 dissolve/regeneration route using ionic liquid<sup>32</sup>. In this study, cellulose nanofibril-chitin nanocrystal hybrid aerogel have been developed for the first time using environmentally friendly

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104 freeze drying process<sup>33</sup>. Theirapplication in dye waste water treatment, anti-bacterial and anti-105 oxidant properties were evaluated very carefully. Cellulose is the richest natural, renewable, 106 biodegradable and biocompatible polymer in the world<sup>34</sup>. Cellulose nanofibrils have high aspect 107 ratios and high surface areas, hence they certainly form an entangled web like structure<sup>35,36</sup>. 108 Cellulose nanofibers can be prepared using different methods including 2, 2, 6, 6 tetramethylpiperidine-1-oxyl (TEMPO)-mediated oxidation<sup>36,37</sup>, mechanical shearing<sup>38</sup>, and 110 high-pressure homogenization<sup>39,40</sup>. The aerogels made from cellulose nanofiber usually show high surface areas, reasonable mechanical, thermal properties and also low densities<sup>41–43</sup>. They have been used in a variety of applications, such as thermal insulation<sup>34</sup>, anti-bacterial agent<sup>44</sup> 113 and oil absorbents<sup>45,46</sup>. Chitin nanocrystals is an economical polymer which contains desired 114 properties such as water solubility, compatibility and functionality. New hybrid aerogel 115 structures based on chitin nanocrystal and cellulose nanofiber having an innovative maple seed 116 type morphology have been made use of for the first time for the macromolecular engineering of 117 new class of aerogels which is an excellent advanced bio-based green materialfor dye adsorption 118 from aqueous solution. Interestingly, the acetamide groups in chitin nanocrystal control anti-119 bacterial properties of aerogels. 104 **Treex drying process<sup>22</sup>**. The<br> **The magnetize were evaluated very carefully. Cellulous to the rachiest notation and and<br>
105 biologizable and biocompatible pelymer in the world<sup>44</sup> Cellulous numelibrished inclusive** 

120 **2. Experimental Section** 

#### 121 **2.1.Materials**

The cellulose extracted from local dried corn husk has been used in the present study. The chitin used for producing the chitin nanocrystal, was a commercially supplied fully bleached shrimp shell flakes. Sulfuric acid,sodium hypochlorite solution, methylene blue(MB), rhodamine 6G (Rh) and other chemicals were obtained from sigma Aldrich.They were used without further

- 126 purification. Millipore-MilliQ distilled water was employed during the complete experiments.
- 127 Table 1 shows the specification of dyes used in this study.

128 **Table 1.** Specifications of MB and Rh

Generic name	<b>Methylene blue</b>
Color Index Number	52015
Molecular formula	$C_{16}H_{18}CIN_3S$
Molecular weight	319.85
$I_{\text{max}}$ (nm)	609
Generic name	<b>Rhodamine 6G</b>
Color Index Number	45160
Molecular formula	$C_{28}H_{31}N_2O_3Cl$
Molecular weight	479.01
$I_{\text{max}}$ (nm)	530

129

#### 130 **2.2.Preparation of Cellulose Nanofibers**

Cellulose nanofiber (CNF) prepared from corn husk were collected from the local area, Slovenia and thoroughly washed with running tap water followed by soaked in Millipore-MilliQ water later dried under sunlight. The washed corn husk was bleached (treated) with aqueous 134 sodium chlorite (1.7 wt % NaClO<sub>2</sub>in water). The bleaching treatment was performed at 90 °C for 2h followed by acetic acid was added to reach pH 4 to enhance the bleaching reaction. The resultant fibers were repeatedly washed in distilled water until the pH became neutral and later 137 dried at 60  $\degree$ C for 12 h in an air circulating oven. The nanofibers were prepared from treated corn 138 fibers by following already reported method of Khawas et.al<sup>47</sup> with suitable modifications. In brief, the methodology is as follows: the dried corn fibers were crushed and treated with sodium hydroxide solution (4 %) at boiling condition for 2 h under thorough stirring to solubilize lignin, pectin and hemicelluloses. This step was done at least 3 times for complete purification of the fibers. The resulting fibers were hydrolyzed with 65 % (w/w) sulfuric acid for about 60 min **The purification.** Millipote-MilliO distilled water was employed during the complete experiments.<br>
127 Table 1 shows the specificantist of MB and Rh<br> **Contention and Rh**<br>
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#### **Page 7 of 33 New Journal of Chemistry**

using mechanical stirrer to dissolute the amorphous region and to isolate the required nanofibers. Subsequently, the suspension was diluted with ice cubes to quench the reaction followed by washing and centrifuging at 10000 rpm for 20 min. The CNF suspension was homogenized by using PRIMIX-Homomixer homogenizer for 15 min and stored in refrigerator.

**2.3.Preparation of Chitin Nanocrystals** 

Chitin nanocrystal (CNC) were prepared by following the study reported by Gopi et 149 al.<sup>48</sup> with suitable modifications. The powdered shrimp shells were suspended in 5 % aqueous potassium hydroxide solution for 6 h under boiling condition in order to remove the proteins and minerals from it. The obtained residue was rinsed with distilled water, filtered and kept for agitation at room temperature followed by bleaching with 17 g of sodium hypochlorite solution 153 for 2 h at 80°C. The resulting dispersion was hydrolyzed with sulfuric acid, filtered and washed with three times and CNC solution transferred into dialysis bag. The dialysis was continued until the pH of CNC solution reached 6 followed by ultrasonicated for 5 min. The CNC solution was stored in refrigerator to prevent from bacterial growth. using mechanical stime to distolute the amorphous region and to isolate the required numefibers.<br>
144 Subsequently, the suspension was dideed with ice codes to quench the reaction followed by<br>
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#### **2.4.Preparation of Cellulose Nanofiber/Chitin nanocrystal hybrid aerogels**

The Cellulose Nanofiber/Chitin nanocrystal hybrid aerogels (CNC/CNF) were fabricated 159 by a cost effective environmental friendly freeze-drying method. 6 and 12.5 mL of  $1 \text{mg } \text{mL}^{-1}$ CNC aqueous dispersion were mixed with 50 mL of 1.2 wt% CNF to get the final concentrations of 1 % and 2 % and named as AR1 and AR2 respectively.The resulting mixture was sonicated using a probe sonicator for 10 min to overcome the agglomerates and finally casted in petridishes. Neat CNF aerogel (neat AR) without the addition of CNC was made as a reference. 164 The whole solutions were precooled in a  $3^{\circ}$ C refrigerator to avoid macroscopic fracture during

165 the freezing step. Later the precooled samples were frozen at  $-73$  °C in a dry ice- iso-propanol 166 solution and freeze dried in a lyophilizer at a condenser temperature of -88.0 °C under vacuum for 4 days to fabricate aerogels. Finally, the samples of aerogel were stored in vacuum oven for further characterization.

**2.5.Fourier transform infrared spectroscopy** 

Fourier transform infrared spectroscopy (FTIR) analysis was used to detect the chemical moieties present in CNF, CNC and CNF/CNC hybrid aerogel recorded using Shimadzu IR-470 IR spectrophotometer. The FTIR spectra for every samples were attained over the wavelength 173 ranging from 400 to 4000 cm<sup>-1</sup> using potassium bromide disk and the experiments were carried 174 out with a resolution of  $2 \text{ cm}^{-1}$ . **one** *Recenting* **step. Later the precediod samples were frozen at -73 <sup>c</sup>C in a dry ises iso-proposal<br>
166 solution and freeze thred in a lyophilizer id a considerer temperature of -88.0 <sup>c</sup>C under vacuum<br>
167 ion 4** 

**2.6. X-ray diffraction analysis** 

X-ray diffraction analysis was employed to determine the crystalline nature of CNF and CNC. The powdered materials of CNF and CNC were placed in the sample holder and leveled to get uniform and total X-ray exposure. The X-ray diffraction patterns of CNF and CNC achieved with as X-ray diffractometer (Bruker Diffractometer D8 Advanced Model) at room temperature 180 with 2 $\Theta$  from 5 to 60<sup>o</sup> and with scan rate of 2<sup>o</sup> min<sup>-1</sup>

#### **2.7. Transmission electron microscopy**

Transmission electron microscopy ((TEM) JEOL-2100 model) was used to determine the morphology and dimension of CNF and CNC. Prior to TEM analysis diluted suspension of CNF and CNC pasted on a copper grid and dried at room temperature. The analysis was carried out with accelerating voltage of 25 KV.

#### **2.8. Field emission scanning electron microscopy**

The morphology of CNF/CNC hybrid aerogel was observed using field emission scanning electron microscopy ((FESEM (JEOL JSM-820 Model))with operation voltage of 20 189 kV. All samples were dried at 40  $^{\circ}$ C and coated with vanadium to avoid following charging before SEM observation.

#### **2.9. BET analysis**

Nitrogen adsorption-desorption isotherms were measured using a Quantachrome® ASiQwin™, and Brunauer – Emmet – Teller (BET) and Barret – Joyner – Halenda (BJH) analysis were obtained using the Autosorb software (Quantachrome®). The BET analysis was performed at relative vapor pressures of 0.08 – 0.3 (*P/P0*) and the BJH analysis was done from desorption isotherm branch. The pore-size distribution and surface area were calculated using BJH and BET methods. **136 2.8. Field emission scanning electron microscopy**<br>
187 The morphology of CNECNC hybrid seroged was observed tosing field emission<br>
188 scanning electron microscopy ((FESEM OEOL JSM-S20 Model))with operation values of

#### **2.10. Swelling studies**

199 Circularly shaped samples with diameter  $\approx$  2 cm of ASTM standard D5890 were weighed and immersed in toluene contained in test tubes with airtight stoppers kept at room temperature for 72 h. Afterwards the initial weight, swollen weight and deswollen weight were taken using a high sensitive electronic balance device. Cross link densities of the aerogel samples were calculated from swelling experiments using Flory-Rehner equation.

#### **2.11.Thermal gravimetric analysis**

Thermal gravimetric analysis (TGA) was conducted for CNF/CNC hybrid aerogels by using Perkin Elmer Pyris 1 thermogravimetric system under nitrogen purge, with a flow rate 100 mL

#### 209 **2.12. Differential scanning calorimetry**

210 Thermal properties of CNF/CNC aerogels were studied by differential scanning 211 calorimetry (DSC) with DSC instrument Mettler Toledo  $822<sup>e</sup>$  using nitrogen atmosphere. The 212 DSC stands as a thermal technique by which I and II order of polymer materials phase transitions 213 can be easily determined. About  $9-11$  mg of samples were heated to  $200^{\circ}$ C from room 214 temperature and stands isothermally for 15 min to exclude the effect of the thermal history 215 during sample processing. Subsequently, the samples were cooled and reheated to  $200^{\circ}$ C. 216 Experiments were carried out with heating/cooling rate of 10  $^{\circ}$ C min<sup>-1</sup>. The phase transition 217 temperatures were determined from the second DSC cycle. **202 online 1. The saming rate was 10 °C** min<sup>2</sup> and the temperature ranging from 40 to 200 °C. All<br> **202 actions were letted 40 °C** for *J* underline try<br> **102 Decrease is an electrometric of CNUCNC serugels** wer

## 218 **2.13. Performance of Cellulose nanofiber/Chitin nanocrystal hybrid aerogel on waste**  219 **water treatment**

220 The dye removal studies of aerogels were carried out using methylene blue (MB) and 221 Rhodamine 6G (Rh  $6G$ ). 1000 mgL<sup>-1</sup> dye concentration in water was used as stock solution. From the stock solution, dilutions of  $10 \text{ mgL}^{-1}$  concentrations of dyes were made and kept at 223 neutral pH. All the analysis was demonstrated at room temperature with the help of UV-VIS 224 spectrophotometer (Agilent technologies, Cary 60 UV-VIS). The samples  $(1 \text{ cm}^2)$  hanged into a 225 glass chamber with a magnetic stirrer and 250 mL of dye solution transferred into it. The 226 adsorption of MB and Rh 6G were measured at the absorbance maxima  $\lambda_{\text{max}} = 609$  nm and  $\lambda_{\text{max}} =$ 227 530 nm respectively. The percentage of removal of MB and Rh 6G with different adsorbents 228 were calculated by following eq.  $(1)^{49}$ .

$$
R = \frac{c_i - c_0}{c_i} \cdot 100\% \text{ (1)}
$$

230 where  $C_i$  (mgL<sup>-1</sup>) is the initial concentration of solution and  $C_0$  (mgL<sup>-1</sup>) is the final concentration of solution.

The following eq. (2) was used for calculating the dye adsorption capacity of the adsorbents: $^{49}$ 

233 
$$
q_e = \frac{(C_i - C_0)V}{m}
$$
 (2)

234 where  $q_e$  is expressed in mg/g, V is the volume of solution and m is the mass of adsorbent.

## **2.14. Antimicrobial and antioxidant activity studies of Cellulose nanofiber/Chitin**

#### **nanocrystal hybrid aerogel**

Four bacterial species were employed as test organisms which include *Staphylococcus aureus, Escherichia coli, Salmonella typhimurium, Bacillus cereus*. The bacteria were maintained in Mueller – Hinton (MH). Inoculate were prepared by adding an overnight culture of the organism in MH broth to obtain an OD600 0.1. The cells were allowed to grow until they obtain the McFarland standared 0.5 (approximately 108 CFU/mL). The suspension was then diluted to 1: 1000 in MH broth to obtain 106 CFU/mL. The antibacterial and antioxidant activities were carried out using neat AR, AR1 and AR2.  $R = \frac{C_1 - C_0}{C_1}$  • 100 % (1)<br>
239 where  $C_1$  (mgL<sup>-1</sup>) is the mind concentration of solidion and  $C_0$  transfer  $\frac{C_0}{C_1}$  ( $\frac{C_1}{C_2}$  ( $\frac{C_2}{C_1}$  The Following eq. (2) wis used for calculating the dye ad

#### **3. Results and discussions**

#### **3.1.Fourier transform infrared spectroscopy**

The fourier transform infrared (FTIR) spectroscopy was used to analysis the chemical moieties present in CNF, CNC, neat AR and AR2 shown in the figure 1. The FTIR spectra of CNF and 248 neat AR show a broad peak at a frequency of  $3300 \text{ cm}^{-1}$  which corresponds to polysaccharide 249 hydroxyl group. The peaks at  $2900-2800$  cm<sup>-1</sup> represent the C-H symmetrical stretching

250 vibrations. The FTIR spectrum of CNC in which, peaks at  $1663 \text{ cm}^{-1}$ ,  $1630 \text{ cm}^{-1}$  correspond to 251 amide I and peak at  $1556 \text{ cm}^{-1}$  corresponds to amide II bands. These are typical well characterized peaks for chitin alredy reported in the scientific literature<sup>15</sup>. Peak at 3451 cm<sup>-1</sup> 253 represents -OH group and peak at  $3259 \text{ cm}^{-1}$  attributed to -NH group. The FTIR spectrum of 254 AR2 having 2 wt.% of CNC shows the same peaks as of CNF or neat AR and also along with 255 the peaks of CNC such as  $1663 \text{ cm}^{-1}$ ,  $1630 \text{ cm}^{-1}$  and  $1556 \text{ cm}^{-1}$  which correspond to amide I and 256 amide II bands respectively. AR2 also shows peak at 3259 cm<sup>-1</sup> attributed to -NH group. To the 257 better understanding about the interaction between CNC over neat AR, FTIR spectra of the same 258 were compared.One thing can be observed that neat AR have a well-defined single peak around  $3300 \text{ cm}^{-1}$  corresponds to -OH functionality. But in AR2, the peak multiplies and a combined 260 multiple peak of both -OH (3300 cm<sup>-1</sup>) and amide group (3259 cm<sup>-1</sup>)observed which possibly 261 due to the hydrogen bonding interaction between the CNF and CNC. Compared to neat AR, AR2 262 have new peaks at  $1663 \text{ cm}^{-1}$ ,  $1630 \text{ cm}^{-1}$  correspond to amide I and peak at  $1556 \text{ cm}^{-1}$ 263 corresponds to amide II bands of CNC which were absent in AR. This confirms better interaction 264 and nanoscale reinforcement of CNC over AR2. The similar observation has been reported by 265 Sun et.al in chitosan (chitin)/cellulose composite biosorbents prepared using ionic liquid for 266 heavy metal ions adsorption. They reported that hydroxyl stretching vibrations for chitosan 267 (chitin) and cellulose located at  $3436 \text{ cm}^{-1}$  and  $3343 \text{ cm}^{-1}$  were broadened and shifted to  $3413$ 268 cm<sup>-1</sup> indicated that stronger intermolecular hydrogen bonds interaction<sup>50</sup>. From the demonstrated 269 FTIR spectra of neat and nanocomposites hybrid aerogel it can be confirmed the presence of 270 CNC in AR2 and the reinforcement doesn't make any changes in chemical moieties present in 271 CNF based aerogel. **259** bydroxy] geong. The posits at 2000-2600 cm<sup>+</sup> represent the C-H symmetrical stretching<br> **250** vibrations. Let F1R spectrum of CNC in winch, posits at 1665 cm<sup>-</sup>, 1690 cm<sup>-</sup> corresponds<br>
251 amide 1 and posit at 1556



273 **Figure 1.** FTIR spectra of a) CNF, b) CNC, c) neat AR and d) AR2.

## 274 **3.2. X-ray diffraction (XRD) analysis**

275 X-ray diffraction (XRD) analysis were employed to reveal crystallinity of CNF and CNC shown in figure 2. The XRD profile of CNF have two major peaks corresponds to  $2\Theta = 16.1^{\circ}$ 276 277 and  $22.8^{\circ}$  which is characteristically structure of cellulose I and sharp peak at  $22.8^{\circ}$  is due to 278 higher crystallinity of cellulose. In the case of CNC, two major diffraction peaks at  $2\Theta = 9.8^\circ$  and 279 19.5<sup>o</sup> along with three minor peaks at  $12.6^\circ$ ,  $23.9^\circ$ , and  $26.8^\circ$  were observed. In CNC, the high 280 crystalline nature and structure is due to the van der Waals and hydrogen bonding interaction in 281 it.



Figure 2. XRD spectra of CNF and CNC

#### **3.2. Transmission electron microscopy**

The transmission electron microscopy (TEM) images and size distribution graphs of both CNF and CNC in different resolutions given in figure 3. Let us examine the TEM images of CNF which is shown in figure 3 a. Uniform suspension of CNF was achieved after mild acid hydrolysis of purified corn husk which is shown in the insight of figure 3 a. A wire like morphology can be observed for CNF with slight agglomeration which may due to the insufficient ultra-sonic treatment. Agglomeration also could be possible due to the water evaporation from the CNF suspension during the sample preparation.The average diameter from the size distribution graph was found to be in the range of 60-120 nm.Similar CNF morphology 293 were reported by Asha et.  $al<sup>51</sup>$  in their studies. **New York of Chemistry Chemistry (CNC)**<br> **New York (CNC)**<br> **OF THE CHEMIST PUBLIC CONTRACT CON** 

Let us now examine the TEM images of CNC. The CNC isolated from shrimp shell using sulfuric acid hydrolysis shows a different morphology compared to other reported studies. The 296 TEM image of CNC show a needle type or rod like structure in most of the reported studies<sup>15,52</sup>. In our present study, CNC shows semi-square crystal morphology which is shown in the figure 3. The crystalline nature and morphology of the product depends on the raw materials and chemical treatment process. An inconsiderable agglomeration also can be observed from the TEM images of CNC which can influence dispersion behavior on CNF suspension. We also observed bright crystalline nature of powdered CNC (achieved by lyophilizing) by visually, which is shown the inset of figure 3 b. The average size from the size distribution graph was found to be in the range of 20-100 nm. Let us now cannise the TEM images of CNC. The CNC isolated from shrimp shell using<br> **295** Fallmans ed CNC show a medie type or red like stuckure in most of the reported studies<sup>192</sup>.<br>
29 The total manuscript SUC sh



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- **Figure 3.** TEM images of a) CNF, b) CNC and their size distribution diagrams.
- **3.3. Field emission scanning electron microscopy**

The morphologies under different resolutions of neat aerogel, AR1 and AR2 were observed and analysed using field emission scanning electron microscopy (FESEM) as shown figure 4. The images of neat AR show random, non-oriented porous structure having distribution in size and these pores were large in size. The fibers were not well interconnected and aggregation of fibers also can be observed. Nanofibers have a tendency to attract each other and this could be a reason for the above-mentioned aggregations. While the images of AR1 show, smaller pores compared to neat AR due to the small addition of CNC which may influence the morphological behaviour of CNF aerogel. In the other case, the images of AR2 containing higher amount of CNC show a notable morphological behaviour compared to neat AR and AR1. The morphological character of AR2 can be compared with layer-by-layer of maple seed. From a clear observation, it can be understood thata single maple seed contains a well oriented vein, these veins interlocked between each other and also a rough texture on the surface. The FESEM images of AR2 is very similar to multi-layer maple seed type morphology demonstrated here. The fiber orientation in AR2 is also mimicking the maple seed which clearly observed from the FESEM image of the same. Upon higher amount of CNC reinforcement, ithas a tendency to locate in between CNF that reduces the intermolecular interaction between the fibers. Consequently, it hinders the chances of aggregation resulting to obtain fiber networking leading to branch like maple seed morphology. This type of morphology is expected to have potential applications in various fields such as water treatment (adsorption, filtration), drug delivery, tissue engineering etc. **19.2 The morphologies under different resolutions of neat acroged, AR1 and AR2 were 900** observed and analysed unit first emission someonic electron microscopy (FESEM) is aboved and unitspectively direct AR intervene



**Figure 4.** FESEM images of aerogels

### **3.4. BET analysis**

BET surface area, pore diameter and pore volume of neat AR, AR1 and AR2 given in 331 table 2. It can be observed that the pore volume is decreasing gradually from neat AR  $(0.24 \text{ cc/g})$ with an increasing in levels of CNC content. AR2 has lesser pore volume (0.029 cc/g) compared to AR1 (0.039 cc/g) since higher level of CNC content in AR2 significantly changes the pore structure. A decrease in pore diameter can also be observed from neat AR to AR1 and then AR2,

- 335 this arises by filling the pores of AR2 with the higher CNC compared to AR1. BET surface area
- 336 value of AR2 was 17.508 nm which is higher than AR1 and neat AR, this increase in surface
- 337 area can improve the efficiency of adsorption performance.

338 **Table 2.** Specifications of MB and Rh

Aerogels	Pore volume $(cc/g)$	Pore diameter (nm)	<b>BET</b> surface area $(m^2/g)$
Neat AR	0.24	.595	11.473
AR1	0.036	.564	13.608
AR <sub>2</sub>	0.029	.555	17.508

339

#### 340 **3.5. Cross-link density studies**

According to the theory of swelling behavior of cross-linked polymers, strong bonding such as chemical cross link between the cellulose nanofiber and chitin nanocrystal chains prevent the molecules from completely surrounded by the fluids, but they can cause swelling. The percentage of increase in swollen mass was calculated by the following equation. **Chiemists by filling the ports of AR2** with the higher CNC compared to AR1. BET surface area<br> **339** value of AR2 was 17.508 much the higher than AR1 and non AR, this increase in surface<br> **339** area can improve the

345 *Swollen mass increase* (
$$
\%
$$
) =  $[(m_1 - m_0)/m_0] \times 100$  (3)

346 where  $m_0$  initial weight of the sample and  $m_1$  is weight of sample after immersing into the 347 solvent in which the solvent specimens cleared by using filter paper.

348 Cross-link density is usually calculated from equilibrium swelling data by means of the Flory-349 Rehner equation<sup>53–55</sup>.

350 Cross-link density  $V = 1/2 M_c$  (4)

351 
$$
M_c = \frac{[-\rho_r V_s V_{rf}^{1/3}]}{[\ln(1 - V_{rf}) + V_{rf} + \chi V_{rf}^2} \tag{5}
$$

352 
$$
V_{rf} = [(d - fw)\rho_r]/[((d - fw)\rho_r]) + (A_s/\rho_s)]
$$
 (6)

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353 where  $M_c$  is the molecular weight of polymer between cross-links,  $\rho_r$  is the density of polymer, 354  $V_s$  and  $V_{rf}$  are molar volume of the solvent and volume fraction of the polymer in the swollen 355 sample respectively. In which d represent deswollen weight f is the volume fraction of filler,  $w$ 356 is the initial weight of the material,  $A_s$  is the adsorbed amount of solvent by the material. The 357 interaction parameter  $\chi$ , shown in eqn (4) and is given by Hildebrand equation

$$
358 \qquad \chi = \beta + \left[ V_s \left( \rho_s - \rho_\rho \right)^2 / RT \right] \tag{7}
$$

359 here *β* is the lattice constant, *R* represent the universal gas constant, *T* is absolute temperature,  $\rho_s$ 360 and  $\rho_{\rho}$  are the solubility parameter of solvent and solubility parameter of the polymer 361 respectively. Table 1 shows the cross-link density values obtained by swelling experiments. 362 Aerogel containing higher amount of CNC (AR2) exhibit higher cross-link density value except 363 AR1 such as  $V = 0.420 \times 10^4$  (mol cm<sup>-3</sup>) and  $V = 3.4124 \times 10^4$  (mol cm<sup>-3</sup>) for AR1 and 364 AR2 respectively. The functional groups (hydroxyl and acetamide groups) of CNC on cellulose 365 nanofiber can act as a network forming agents able to react with the CNF chains leading to the 366 formation of strong networking structure in the aerogel. Solen where  $M_n$  is the molecular weight of polymer between cross-links,  $\rho_n$  is the density of pubymer.<br> **S54**  $V_n$  and  $V_n$  are molecular volume of the subvent and volume faction of the pubymer in the svolume of<br> **S54** 

#### 367 **3.6. Thermal gravimetric analysis**

368 The thermal gravimetric analysis (TGA) curves for AR1 and AR2 in comparison with 369 reference neat AR are shown in figure 5. All materials show decrease in mass at  $100^{\circ}$ C which 370 obviously due to the evaporation of adhering moisture from the materials. By careful evaluation 371 of obtained data, we can observe that AR2 losses only 5.2 % of mass at this temperature which is 372 low compared to other reported studies<sup>56,57</sup>. This may due to the freeze-drying process that 373 sublimate the whole water moieties from the sample. Also, the mass loss of AR2 at 100  $^{\circ}$ C lower 374 than neat AR and AR1 (8.4 % and 7.8 % respectively) probably due to fine dispersion of CNC

on CNF thus leaving no vacant space for entrapment of water molecules. The AR2 with 376 relatively higher CNC content unveiled an onset weight loss at around 290  $\degree$ C, which was higher than neat AR and AR1 having lower CNC content. The TGA results showed that thermal stability of AR2 aerogel was the best among all the samples (neat AR and AR1). This suggested that there existstrong interaction between CNF and CNC in AR2.



**Figure 5.** TGA curves of a) neat AR, b) AR1 and c) AR2.

**3.7. Differential scanning calorimetry** 

The differential scanning calorimetry (DSC) curves of neat AR, AR1 and AR2 are shown 384 in figure 6 a. It can be seen that the melting temperatures  $(T_m)$  of neat AR and AR1 (104.9<sup>o</sup>C)are 385 much lower than AR2 (126.6<sup>o</sup>C). The high melting temperature of AR2 is due to the uniform dispersion of CNC in CNF as mentioned in morphological part, which prevents the motion of CNF chains. As a result, more energy is needed for the molecular motion of CNF chains. The

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more is the crystalline region, the higher will be the melting temperature. A graphical representation given in figure 6 b showing better organization and alignment of nanofibers in AR2 having higher amount of CNC that can be leads to higher crystallinity and melting point.



**Figure 6**a) DSC curve of aerogels b) Graphical representation of aligned fibers in AR2

## **3.8. Performance of Cellulose nanofiber/Chitin nanocrystal hybrid aerogel on waste water treatment**

The effect of contact time on the dye water treatment using MB and Rh are shown in figure 7. It is clearly observed that the percentage of removal (removal (%)) and adsorption 397 capacity  $(q_e)$  of neat AR and AR1 are low compared to AR2. The percentage of removal and adsorption capacity of AR2 were found to be 80.07 % and 8 mg/g for MB, 92.89 % and 9.2 mg/g for Rh respectively. A schematic representation of mechanism of MB and Rh adsorption onto AR2 is shown figure 8. The AR2 material contains higher amount well dispersed CNC on CNF and the CNCs are enriched with acetamide groups. During the course of adsorption, the deprotonated carbonyl group present in acetamide (reactive group in CNC) creates a negative charge species as shown in the scheme (inside the green one). These negatively charged CNChas **New Your State Constitution region, the higher will be the melting temperature. A graphical <b>one of** proposition and interval and algument of constrained published and algument of particle DOI: 10.1039/2017. AND the memb an electrostatic interaction between positively charged dye molecule which is shown as doted lines. These electrostatic interactions favour efficient dye adsorption over the CNC decorated AR2. It is also important to consider the well-designed morphological texture in AR2 as already discussed under morphological part which could be anotherpossibility of the enhanced dye removal from water.



410 **Figure 7.** Removal(*R*) and adsorption capacity ( $q_e$ ) of neat AR, AR1 and AR2 on a) MB and b)

Rh.



**Figure 8.** Schematic representation of mechanism of MB and Rh adsorption onto AR2.

### **3.9. Round studies of material**

Round studies were carried out using AR2 on MB and Rh to understand the reusability of material. After the first-round of adsorption experiment the sample AR2 was dried at room temperature and washed several times using distilled water. Totally 5round experiments were conducted with the same drying and washing protocols and the results are shown in figure 9. During the first round, 80.07 % MB and 92.87 % Rhcould be removedand the removal efficiency 420 was gradually found decreased and in the final round i.e. in  $5<sup>th</sup>$ round 30.2 % MB and 52.9 % Rh could be removed using AR2. The decreasing efficiency up reuse could be explained as follows: During the several adsorptions process the active sites of material were occupied with dye 423 molecule and few of them only could be removed during thoroughwashing procedure<sup>58</sup>. As a result, subsequent adsorption process, active sites required to adsorb dye molecules reduces. Thereby a reduction in adsorption takes place in the round studies. Even though AR2 can be reused by simple washing and drying process in several times, one has to sacrifice the efficiency. **Figure 8. Schematic representation of mechanism of MB and Rh adsorption onto AR2.**<br> **As As Counter Accepted Manuscript Counterpart Constant Constant Constant Constant Affect Counterpart Constant Affect Counterpart 2017** 





**Figure 9.** Reusability of AR2 on removal of Rh and MB

## **3.10. Antimicrobial and antioxidant activity studies of Cellulose nanofiber/Chitin nanocrystal hybrid aerogel**

**Determination of antibacterial activity.**The discs such as neat AR, AR1 and AR2 were placed on Mueller-Hinton agar plates, previously swabbed with the target bacterial isolate at a concentration of 106 CFU/mL. In one disc, the respective organic solvent was added as negative control to determine possible inhibitory activity of the solvent. This preparation was incubated 435 for a period of 24 h at 30  $^{\circ}$ C. Antibacterial activity was defined as the diameter of the clear inhibitory zone formed around the discs. The MIC of the extract was determined by tube dilution technique in Mueller-Hinton broth (supplied by Merck) according to NCCLS. The range of concentration used was 156.25 to 5000 µg/mL. The four last vials of each bacterium with no growth from the MIC procedure were streaked onto nutrient agar (NA) plates. The antibacterial activities of aerogels neat AR, AR1 and AR2 against both gram-positive and gram-negative bacteria strains were examined by detecting of the minimum inhibitory concentrations (MICs). Each sample were assayed three times and the results are shown in the figure 10 (a) and table 3. From these results,it can be observed that the antibacterial activity of AR2 is the best among all other samples (neat AR and AR2) that produced zones of inhibition against E. coli, S. typhimurium, St. aureus and B. cereus. **3.40.** And<br>**nanocrystal hybrid acrogel**<br>**A22 Determination of antibacterial activity. The disc wash as near AR, AR1 and AR2 was<br><b>Determination of antibacterial activity**. The disc was a stand a RA AR1 and AR2 was<br>**313** 

**Determination of antioxidant activity.** The antioxidant activity of the aerogels neat AR, AR1 and AR2 was evaluated using DPPH (2, 2-diphenyl-1-picrylhydrazyl from Merck, Damstadt, Germany) radical scavenging assay. Briefly, 3 mL of aerogel dispersions were mixed with 1 mL of 1mM methanolic solution of DPPH. The mixture was vortexed and incubated in the dark at ambient temperature for 30 min. When the DPPH solution was mixed with the sample mixture which is acting as a hydrogen atom donor, a stable non-radical form of DPPH is

452 obtained with simultaneous change of the violet color to pale yellow. The absorbance was then 453 measured at 517 nm. The percentage of DPPH free radical quenching activity was determined 454 using the following equation

$$
DPPH\,scavending\,activity\% = \frac{A_{DPPH} - A_{Extract}}{A_{DPPH}}\,X\,100 --- --- --- (8)
$$

455 where A<sub>DPPH</sub> is the absorbance value at 517 nm of the methanolic solution of DPPH and  $A<sub>Extract</sub>$  is the absorbance value at 517 nm for the sample extract. Each sample was assayed two 457 times and results shown in the figure 10 (b) and table 4. From this results, it can be observed that 458 the antioxidant activity of AR2 was high compared to neat AR and AR1 as evaluated by free 459 radical scavenging assay. **Assumed with simultaneous change of the violat color to pala yellow. The absorbance was then<br>
<b>ASS** measured at 517 nm. The percentage of DPPH free radical quenching activity was determined<br> **OPPH screw angle accepted**<br>



461 **Figure 10.** neat AR, AR1 and AR2 for a) antimicrobial b) antioxidant studies



460

Table 3. Antibacterial properties of aerogels

			MIC % $(w/V)$		Mean MIC $\%$ (w/V)
<b>Sample</b>	<b>Bacterial strains</b>	<b>Assay</b>	<b>Assay</b>	<b>Assay</b>	
neat AR	E. Coli	6.2	7.5	7.8	7.17
	S. typhimurium	6.8	7.2	7.8	7.27
	<i>St.</i> aureus	7.1	7.6	8	7.57
	B. cereus	6.9	7.9	7.8	7.53
AR1	E. Coli	5	3.8	4.6	4.47
	S. typhimurium	4.8	5	5.2	5.00



463

### 464 **Table 4.** Antioxidant properties of aerogels



The CNC embedded bio-aerogels of cellulose (mainly AR2 which contains more CNC) in which the cells stick to each other and often these cells adhere to a surface. These adherent cells are frequently embedded within a self-produced matrix of extracellular polymeric substances (EPS). Furthermore, from all the analysis it is revealed that AR2 is showing a promising result of antibacterial and antioxidant properties compared to neat AR and AR1 which may be due to the presence of acetamide group in chitin. Hence it would be interesting to investigate the potentiality of this aerogel for possible applications in different fields. 3.  $\frac{32.00 \text{ cm/s}}{R \cdot \text{Coul}}$ <br>
AR2  $\frac{R \cdot \text{Coul}}{R \cdot \text{Coul}}$ <br> **ACCEPTE 2018**<br> **SUPPOSE ACCEPTED Accepted**<br> **OF CHEMISTRON CONSULS 12.2** 13.3<br> **OF CHEMISTRON CONSULS 12.2** 13.3<br> **ACCEPTED ACCEPTED**<br> **SUPPOSE ACCEPTED ACCE** 

472 **4. Conclusion** 

A facile green method was used to fabriacte multi-functional bio-hybrid aerogel based on CNF decorated with CNC. The morphology and propoiertis of the new nanostrucutred hybrids were charcterised using TEM, FESEM, DSC and FTIR. The chemical moeties of CNF, CNC and AR2 were analysed using FTIR spectroscopy and confirmed the presence of CNC on CNF aerogel. The CNF fibers showed wire like morphology having 60-120 nm in diameter while the CNC showed semi-square morphology having 20-100 nm in size. This semi-square morphology

for CNC has been reported for the first time as compared to other studies in the literature. Subsequently bio-hybrid aerogels are made with different percentage of CNC (1 % and 2 %) decorted on to CNF using environmentaly friendly freeze drying method. Their morphologies were investigated with respective the reference (neat AR) and found maple like morphology for AR2. The multi functional efficiency of AR2 was evaluated using various studies such as dye water treatemnt, anti-bacterial and anti oxidant studies and realized that AR2 is a promising aerogel material for the above mentioned applications. Further studies to be anticipated in the future for industrial scale applications. **APS** For CNC has been reported for the first time as compared to other studies in the literature.<br> **AUG** Moseopotally bis-hybrid are<br> **Manuscriptor** and university of CNC (1.9<sub>2</sub> and 2.94)<br> **AS2** decorted on to CN1 using

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