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Preparation of Activated Carbon and Silica Particles from Rice Straw

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Supporting Information

ABSTRACT: An efficient three-step process using toluene/ 5 ethanol, NaClO₂, and KOH has been successfully devised to 6 isolate pure cellulose from rice straw while generating two 7 filtrates as activated carbon and silica precursors. The NaClO₂ 8 dissolution filtrate contains oxidized lignin and hemicellulose 9 10 as carbon precursors as well as sodium carbonates as activating agents for direct carbonization (800 °C) into highly porous 11 $(0.90 \text{ cm}^3/\text{g})$, high specific surface area (997 m²/g), activated 12 carbon particles (100-500 nm). The KOH dissolution filtrate 13 contains mainly potassium silicate that could be precipitated 14



- by dilute acidified poly(ethylene oxide) and calcinated (500 °C) to pure, uniformly sized (100-120 nm), nonporous silica 15 nanospheres. Deriving these additional activated carbon and silica particles along with nanocellulose creates advance materials 16
- while fully utilizing all major components in rice straw, the highest quantity agricultural crop byproduct in the world. 17
- KEYWORDS: Activated carbon, Porous, Monodispersed silica, Green chemistry, Rice straw utilization, 18

INTRODUCTION 19

20 Lignocellulosic biomass, such as agricultural crop residues, 21 forestry byproducts, and municipal waste, is a rich source of 22 renewable energy and materials. Lignocellulosic materials have $_{\rm 23}$ been converted to biofuel such as ethanol, $^{1-3}$ hydrogen, and 24 other combustible gases,^{4,5} as well as nanocellulose,³⁻⁵ 25 carbon,^{2,5} and silicon-based^{6,7} materials. Cellulose nanocrystals 26 and nanofibrils have been isolated from various major 27 agricultural residuals including rice straw,⁶ wheat straw, and 28 soy hulls.⁷ Direct thermal processing of various biomass has 29 also generated charcoals,⁴ activated carbon,⁸ and silicon-based 30 advanced materials including mesoporous silica⁹ as well as 31 silicon carbide and nitride.¹⁰ Simultaneous derivation of parallel 32 products from specific biomass for full utilization has not been 33 given much attention.

Rice straw represents the largest agricultural byproduct 34 35 around the world. Rice is the largest cereal crop and the highest 36 valued agricultural commodity (\$187 billion in 2011) in the 37 world.¹¹ Although third in quantity (723 million MT in 2011) 38 behind sugar cane and maize, rice production generates the 39 highest quantity of byproduct as rice straw accounts from 1 to 40 1.5 kg per kg of rice grain harvested.¹² Direct use has presented 41 some shortcomings as rice straw is a marginal feed compared 42 with other cereal straw and causes severe furnace fouling and 43 ash production when burned to produce steam.

Other utilization of rice straw has mainly involved single 44 45 product development by converting certain components while 46 removing others employing biochemical, chemical, and/or 47 thermal processes. Ethanol production was improved by 48 enzymatic saccharification of cellulose and hemicellulose,¹³ 49 while hydrogen-rich gas production was made efficient via 50 microwave-assisted pyrolysis of rice straw.¹⁴ Long, highly 51 crystalline, cellulose fibers have been extracted from rice

straw via alkali and enzymatic methods.¹⁵ Activated carbons 52 were prepared from rice straw by KOH activation and s3 carbonization at 800–900 °C^{16,17} or preoxidation at 200 °C s4 and $(NH_4)_2HPO_4$ activation at 700 °C¹⁸ to reach 1900 and s5 1150 m^2/g specific surface areas, respectively, comparable to 56 commercially available activated carbon products. Silica nano- 57 spheres were prepared from alkali dissolution of burned straw 58 ashes and then acid precipitation¹⁹ to better control nano s9 particle sizes from 16 to 100 nm.²⁰ Uniform nanodisks were 60 also prepared by sulfuric acid precipitation of dissolved silica 61 from rice straw ashes, followed by freeze-drying.²¹ Silicon 62 carbide and nitride nanorods were fabricated by direct pyrolysis 63 of rice straw under nitrogen from 1400 to 1800 °C.²

The significant lignocellulosics, that is, 32-47% cellulose, 65 19-27% hemicellulose, and 5-24% lignin, as well as 7-20% 66 silica and trivial amount of waxes and minerals in rice 67 straw^{20,23–27} present opportunities for simultaneous develop- 68 ment of multiple products. We have developed a simple 69 approach to isolate cellulose, lignin/hemicelluloses, and silica 70 components in rice straw with minimal chemical and energy 71 input.⁶ Pure cellulose was extracted from rice straw via an 72 efficient NaClO₂-KOH process with at least 36% yield, where 73 dewaxing in 2:1 v/v toluene/ethanol extraction was followed by 74 lignin removal via acidified NaClO₂ and then hemicellulose and 75 silica isolation by KOH. Cellulose nanocrystals and nanofibrils 76 have been derived from pure rice celluloses via sulfuric acid 77 hydrolysis^{6,28} and 2,2,6,6-tetramethylpyperidine-1-oxyl 78 (TEMPO)-mediated oxidation,²⁸ respectively, and then 79 assembled into fibrous and porous matrixes. 80

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Figure 1. Scheme for (a) isolation of celluloses, lignin-rich filtrate 1, and silica-rich filtrate 2 from rice straw; (b) activated carbon from filtrate 1; and (c) silica from filtrate 2.

The goal of this work was to develop additional advanced 81 82 materials from the as-derived extracts of two dissolution 83 streams in isolating pure cellulose toward utilization of all 84 major rice straw components, that is, lignin, hemicelluloses, and 85 silica. NaClO₂ and KOH were used as the former is a strong 86 oxidant that could break the strong bonding among the 87 lignocellulosics, and the latter could dissolve silica in rice straw. 88 The NaClO₂ dissolution stream from the dewaxed rice straw 89 contains oxidation products of lignin and hemicellulose as well 90 as sodium byproducts. As lignin and hemicelluloses were 91 intrinsically bound to each other and thus very difficult to be 92 completely separated without sacrificing one or another to 93 soluble fragments, it is advantageous and economic to use this 94 lignin/hemicelluloses mixture together as a carbon source, 95 while sodium salts already present could serve as the activating 96 agents to produce porous structures without needing further 97 addition of chemicals. The KOH dissolution stream from 98 NaClO2-treated rice straw should contain mainly potassium 99 silicate, which may serve as a precursor to silica. These 100 precursors in both streams would be recovered by precipitation 101 using ethanol for lignin/hemicelluloses carbon precursors and 102 acidified poly(ethylene oxide) for the silica precursors. These 103 two precursors could be thermally converted to carbon and 104 silica products via a single-step pyrolysis and calcinations at 800 105 and 500 °C, respectively.

106 **EXPERIMENTAL DETAILS**

107 **Chemicals.** Rice straw (Calrose variety) used in this study was 108 from the 2009 harvest in the Sacramento valley in northern California. 109 Toluene ($C_6H_5CH_3$, ACS grade, Fisher Scientific), ethanol (CH₃CH₂OH, anhydrous, histological grade, Fisher Scientific), 110 sodium chlorite (NaClO₂, 80%, Fluka), glacial acetic acid 111 (CH₃COOH, 99.7%, ACS GR, EMD), poly(ethylene oxide) (PEO) 112 (M_w = 600 kDa, Sigma-Aldrich), and potassium hydroxide (KOH, 113 85%, EM Science) for fractionating rice straw components were used 114 as received. Water used was purified by a Milli-Qplus water 115 purification system (Millipore Corporate, Billerica, MA). All 116 concentration and yield percentage values were based on weight 117 unless otherwise specified. 118

Preparation of Activated Carbon and Silica Particles from 119 Rice Straw. Fractional extraction of lignin/hemicelluloses and silica 120 from rice straw was first performed by the NaClO2 treatment as shown 121 in Figure 1. Rice straw was washed thoroughly with water to rid of dirt, 122 fl dried, and then milled (Thomas-Wiley Laboratory Mill model 4, 123 Thomas Scientific, U.S.A.) to pass through mesh 60. Rice straw 124 powders were Soxhlet extracted with 2:1 toluene/ethanol (v/v) at 55 125 °C for 24 h to remove all lipophilic and hydrophilic nonstructural 126 components (Figure 1). The dewaxed rice straw powder was placed in 127 1.4 wt % NaClO₂ under acidic condition (pH ~5 adjusted by 10% 128 acetic acid) at a 15 mL/g liquid-to-solid ratio at 70 °C for 6 h and 129 filtered to yield holocellulose/silica and filtrate 1. α -Cellulose was 130 isolated from holocellulose by extracting with 5 wt % KOH at a 20 131 mL/g liquid-to-solid ratio at 70 °C for 24 h to give pure cellulose and 132 dissolved silica and hemicelluloses in filtrate 2. Both filtrate 1 and 2 133 were then concentrated by heating at 90 °C to half their volumes. The 134 remaining filtrates 4 and 5 contained mostly water with very little 135 solutes. Ethanol in filtrate 4 may be recovered by distillation. 136

Preparation of Activated Carbon. Ethanol was added to the $_{137}$ concentrated filtrate 1 at 3 times its volume to precipitate alkali-soluble $_{138}$ lignin together with some hemicelluloses (LH). The LH powders were $_{139}$ dried in oven at 60 °C for 12 h, placed in a quartz tube (2 cm inner 140 diameter), and then <u>pyrolyzed</u> in a furnace (Mini-Mite, Lindberg/ 141 Blue) at 10 °C/min to 105 °C and held for 0.5 h. Then the LH 142

¹⁴³ powders were heated to 800 $^\circ$ C and held for up to 0.5 h, all under ¹⁴⁴ flowing N₂ at 100 mL/min. Finally, they were cooled to ambient ¹⁴⁵ temperature in 12 h. The particles were washed with 5% HCl and ¹⁴⁶ water to remove residual salts and other small hydrocarbon impurities ¹⁴⁷ and then dried at 105 $^\circ$ C for 0.5 h to yield activated carbon (AC).

Preparation of Silica Particles. Acidified PEO solutions at 0.2%, 148 149 0.35%, 0.5%, and 1% were prepared by dissolving 0.04, 0.07, 0.1, and 150 0.2 g of PEO, respectively, in 20 g of 2% HCl and stirred for 6 h. Concentrated filtrate 2 was added slowly to each acidified PEO 151 152 aqueous solution to form a milky white mixture and centrifuged 153 (Centrifuge 5804R, Eppendorf) at 5000 rpm for 15 min. The 154 precipitate was then washed with water thoroughly and dried in a 155 desiccator as silica and lignin/hemicelluloses/PEO (SHLP). The dried 156 SHLP powder was placed in a quartz tube (2 cm inner diameter) and 157 calcinated under flowing air (100 mL/min) in a furnace (Mini-Mite, 158 Lindberg/Blue). The sample was heated at 10 °C/min to 105 °C and 159 then 500 °C; the sample was held for 0.5 h at each temperature. The 160 obtained silica after calcination was cooled under flowing air (100 mL/ min) to room temperature in 12 h. Filtrate 4, SHLP, and silica were 161 referred to those obtained at 0.35% PEO unless stated otherwise. 162

Analytical Methods. The yields were calculated by the percentage 163 164 of extract or product mass over the original rice straw mass, both 0.1 165 mg accuracy (BP 300S, Satorius). The aqueous filtrates from the 166 extraction process were examined by ultraviolet-visible spectroscopy 167 (UV-vis) (Evolution 600, Thermo Scientific). UV samples were placed in polystyrene cuvettes with a 1 cm path length. The chemical 168 composition and structure of LH, AC, SHLP, and silica were analyzed 169 170 by Fourier transform infrared spectroscopy (FTIR) (Nicolet 6700, 171 Thermo Scientific). All FTIR spectra were collected from samples 172 dried at 60 °C for 12 h and pressed with anhydrous KBr powders into 173 pellets. The elemental analysis of LH, AC, SHLP, and silica were 174 carried out by energy-dispersive X-ray spectroscopy (EDX) adjunct to 175 a scanning electron microscope (SEM) (FEI-XL 30, FEI). All samples 176 are stored in oven at 60 °C for 24 h before EDX measurements.

The morphology and structure of LH and SHLP as well as their 177 pyrolyzed products AC and silica were observed by SEM (FEI-XL 30, 178 FEI). The SEM samples were sputter coated with gold for 2 min and 179 then observed under a working voltage of 5 kV. The structures of AC 180 and silica particles were also examined by transmission electron 181 182 microscopy (TEM) (JEOL 3000, JEOL). TEM samples were prepared by dispersing a small amount of AC and silica particles in water (~0.01 183 184 g/L) and sonicated (2510, Branson) for 60 min first, and then a drop 185 of the sonicated suspension was placed onto a carbon grid and dried in 186 air

187 The thermal properties of LH and SHLP were analyzed by 188 differential scanning calorimetry (DSC) (DSC-60, Shimadzu) and 189 thermogravimetric analysis (TGA) (TGA-50, Shimadzu). DSC 190 samples were tightly packed in aluminum cells with press-sealed lids, 191 while TGA samples were placed in a platinum pan. LH samples were 192 heated to 60 °C and held for 2 h, then heated to 105 °C and held for 193 0.5 h, and finally heated to 550 and 800 °C for DSC and TGA, 194 respectively, and held for 0.5 h under 50 mL/min nitrogen flow. SHLP 195 samples were heated to 105 °C and held for 0.5 h and then heated to 196 500 °C and held for 0.5 h under a 50 mL/min air flow.

For surface area and pore characteristics, LH, AC, SHLP, and silica 197 were dried in oven at 50 °C for 48 h and then measured at 77 K by a 198 199 nitrogen adsorption-desorption analyzer (ASAP 2020, Micrometics). The surface area was calculated from the isotherm using the 200 Brunauer-Emmett-Teller (BET) equation in the linear region 201 202 where relative pressure P/P_0 ranged from 0.05 to 0.1. The mesopore surface areas of LH, AC, SHLP, and silica were derived from the 203 adsorption branch, whereas pore- and neck-size distributions were 2.04 205 derived from both adsorption and desorption branches of the isotherm 206 using the Barret-Joyner-Halenda (BJH) method. The micropore surface area and pore hydraulic diameter distribution (0.7-1.6 nm) of 207 208 AC were derived from the t-plot using the Mikhail, Brunauer, and 209 Bodor MP method²⁹ and the Harkins and Jura equation.³⁰ Micropore 210 volume $(V_{\rm mp})$ was derived from the tangent line of a contiguous range 211 of *t*-plots using the surface area of the filled pores via eq 1,

217

$$V_{\rm mp} = \frac{(S_n - S_{n+1}) \times (t_n + t_{n-1})}{2} \times 15.47 \tag{1}_{212}$$

where S_n and t_n are the surface area derived from the slope of tangent 213 and the thickness of absorbed layer at the *n* point in the *t*-plot, 214 respectively, and 15.47 was the constant of converting the gas volume 215 to liquid volume at STP. 216

RESULTS AND DISCUSSION

Yields and Efficiency of Sodium Chlorite Isolation. The 218 NaClO₂-KOH process for isolating rice straw components 219 involved three steps, yielding pure cellulose as well as filtrates 1 220 and 2 (Figure 1a). The washed and dried rice straw was 221 refluxed in a 2:1 toluene/ethanol (v/v) ratio at 55 °C for 24 h 222 to yield 94.7% dewaxed rice straw and a green-yellow distillate 223 containing surface oils, pigments, and possibly organic soluble 224 lignin. The dewaxed rice straw was then treated by 1.4% 225 NaClO₂ at pH 5 adjusted by acetic acid at 70 °C for 6 h and 226 then 5% KOH at 70 °C for 24 h to obtain pure cellulose as well 227 as the NaClO₂ and alkaline dissolution streams, that is, filtrates 228 1 and 2, respectively. Adding ethanol to filtrate 1 under the 229 basic condition precipitated the dissolved lignin/hemicelluloses 230 (LH) at 5.7% of the original rice straw mass (Figure 1b). 231 Adding the basic filtrate 2 to the dilute acidified PEO 232 precipitated silica along with remaining lignin/hemicellulose 233 (SHLP) at similar yields of 22.1-23.7%, increasing to 0.2-1% 234 PEO concentrations (Figure 1c). At 0.35% PEO, the silica/ 235 hemicelluloses/lignin (SHL) yield was 20.7%, assuming all 236 PEO precipitated. 237

Chemical Structures of Extracts and Final Products. $_{238}$ NaClO₂ and KOH treatments yielded respective filtrates 1 and $_{239}$ 2, which were precipitated as precursors to be subsequently $_{240}$ converted to AC and silica particles, respectively. The $_{241}$ compositions and structures of both filtrates, their precipitates, $_{242}$ and final products were examined by UV–vis, FTIR, and $_{243}$ elemental analysis to determine their compositions and to $_{244}$ assess the isolation process. $_{245}$

UV–Vis Spectra of the Filtrates. The UV–vis spectrum 246 of filtrate 1 exhibited strong broad absorbance below 300 nm 247 (Figure 2), ascribing to partially oxidized and/or decomposed 248 f2 lignin from NaClO₂, possibly a polylignol dehydrogenative 249 copolymer of sinapyl alcohol, coniferyl alcohol, p-coumaryl 250 alcohol, and hydroxycinnamic acid esters.^{31,32} Filtrate 2 showed 251 reduced yet clear UV absorbance below 300 nm, indicating the 252



Figure 2. UV–vis spectra of filtrates from rice straw isolation shown in Figure 1.

253 presence of some alkaline-soluble lignin and incomplete lignin 254 removal in the previous NaClO₂ step. The broad absorption 255 peak in the 360 nm region of the filtrate 2 spectra is consistent 256 with the expected alkaline-soluble lignin, possibly various lignin 257 derivatives of o,p-dihydroxystilbenes, α -carbonyl phenolics, 258 esters of p-coumaric, and ferulic acids,³³ as well as charge-259 transfer complexes from the carbonyl and carboxyl-conjugated 260 heteroaromatic compounds.³⁴ Filtrate 3 showed very little 261 absorbance below 300 nm, indicating effective precipitation of 262 lignin in LH by ethanol. Filtrate 4 also exhibited reduced peak 263 intensity below 300 nm, showing the presence of a small 264 amount of alkaline-soluble lignin in this final filtrate. The UV-265 vis spectrum suggested that most lignin in rice straw was 266 extracted by NaClO₂ dissolution in filtrate 1 and recovered by 267 ethanol precipitation in LH, while some lignin present in the alkaline filtrate 2 was mostly precipitated by PEO in SHLP, 268 269 leaving yet a small residual lignin in filtrate 4. Therefore, lignin 270 in rice straw was recovered by precipitation mainly in LH from 271 filtrate 1 and partially in SHLP from filtrate 2, with a residual 272 amount in the final filtrate 4.

FTIR Spectra of Precipitates and Final Products. The Transform for the LH precipitate from filtrate 1 showed major lignin characteristic peaks, including those between 1600 and 1000 cm⁻¹ from partially oxidized phenylpropanoids and multiple hydroxyl peaks (Figure 3). Lignin was evident by the

f3



Figure 3. FTIR spectrum of LH, AC, SHLP, and silica.

278 strong 1450 and 1040 cm⁻¹ peaks associated with aromatic 279 C=C stretching and aromatic =C-H in-plane bending, 280 respectively. The prominent peak at 3459 cm⁻¹ and two minor 281 peaks at 2930 and 2835 cm⁻¹ were attributed to OH stretching 282 in the respective hydrogen-bonded and free hydroxyls, whereas 283 the ones at 1090 cm⁻¹ were from C-O deformation in the 284 aliphatic hydroxyl and ether groups. The broad band at 1630 285 cm⁻¹ could be from moisture peaks overlapped with various 286 C=O stretching, C=C stretching in phenylpropanoid side 287 chains, and aromatic skeleton vibration. Therefore, LH 288 precipitated from filtrate 1 consisted mainly of oxidized lignin 289 from reaction with the strongly oxidative NaClO₂.

SHLP precipitated from filtrate 2 with diluted PEO showed characteristic peaks of silica, lignin, hemicelluloses, and PEO (Figure 3). Silica was clearly evident by the prominent Si-O-Si asymmetric and symmetric stretching peaks at 1090 and 460 294 cm⁻¹, respectively, as well as a Si-O deformation peak at 795 295 cm⁻¹. The presence of lignin was evident by the benzene C==C 296 stretching peak at 1460 cm⁻¹. However, the peak intensity decreased greatly compared to that of LH, suggesting the ²⁹⁷ reduced lignin composition in SHLP. The broad shoulder at ²⁹⁸ 1205 cm⁻¹ was ascribed to the various C–O–C stretching from ²⁹⁹ hemicelluloses and PEO. The broad shoulder at 1640 cm⁻¹ was ³⁰⁰ from moisture overlapping with C=C stretching and various ³⁰¹ esters and ketone structures, such as β ketone ester, $\alpha - \beta$ ³⁰² unsaturated ketone, and β diketone (enol form) in hemi- ³⁰³ celluloses. The broad 3430 cm⁻¹ peak as well as minor peaks at ³⁰⁴ 2930 and 2835 cm⁻¹ were ascribed to hydroxyls from moisture, ³⁰⁵ lignin/hemicelluloses, and silica surfaces. ³⁰⁶

The pyrolytic product of LH from heating at 800 °C showed 307 little characteristic FTIR peaks of either lignin or hemicelluloses 308 as expected. The broad OH stretching peak around 3440 cm⁻¹ 309 greatly decreased in intensity compared to LH, clearly showing 310 the loss of hydroxyls and is consistent with transformation of 311 hydrophilic lignin/hemicelluloses to hydrophobic carbon. The 312 only two small peaks at 1454 and 1380 cm⁻¹ were ascribed to 313 aliphatic C–H bending. Thus, the FTIR of pyrolyzed LH is 314 consistent with conversion to carbon with few lignin and 315 hemicelluloses features. 316

The FTIR spectrum of SHLP calcinated at 500 °C showed 317 only silica characteristic peaks, that is, Si-O-Si asymmetric 318 and symmetric stretching at 1080 and 460 cm⁻¹, respectively, 319 and Si-O deformation at 780 cm⁻¹, as well as the broad 320 hydroxyl stretching peaks at 3440 and 1630 cm⁻¹, indicating 321 absorbed moisture or the hydrophilic nature of silica. None of 322 the lignin, hemicelluloses, and PEO characteristic peaks was 323 observed, confirming their removal upon calcination. 324

UV-vis results clearly showed that lignin-rich LH and silica/ 325 hemicelluloses-rich SHLP were successfully isolated in filtrates 326 1 and 2, respectively, while FTIR confirmed that the isolated 327 lignin in LH and hemicelluloses in SHLP were oxidized and 328 degraded by the oxidative $NaClO_2$ and then thermally 329 converted to carbon and silica, respectively. 330

Elemental Analysis of Extracts. The elemental compo- 331 sitions by EDX show LH to consist of 38.0% C, 42.8% O, 332 18.6% Na, and 0.6% Si (Table 1). The high Na and O contents 333 ti

Table 1. Elemental Compositions (wt%) of Crude LH and SHLP and Their Respective Pyrolyzed Products: AC (800 $^{\circ}$ C, 30 min) and Silica (500 $^{\circ}$ C, 0.5 h)

	С	0	Si	Na
LH	38.0	42.8	0.6	18.6
AC	81.7	15.8	2.5	0
SHLP	16.6	51.7	29.7	0
silica	0	53.0	47.0	0

suggested LH contained oxy-salts of sodium, possibly sodium ³³⁴ carbonates, oxidative products of lignin/hemicelluloses, and ³³⁵ acetic acid in the NaClO₂ step. The continuous evolution of ³³⁶ odorless bubbles when adding crude LH powder into dilute ³³⁷ aqueous HCl indicated gaseous products and was consistent ³³⁸ with the presence of carbonate salts in crude LH. To further ³³⁹ investigate the compositions of inorganic salts, LH was ³⁴⁰ calcinated at 500 °C for 6 h. The yellow color of LH turned ³⁴¹ to a white color after calcination (Figure S1, Supporting ³⁴² Information) and was shown by EDX to have the elemental ³⁴³ composition of Na₂CO₃ (Figure S1, Supporting Information), ³⁴⁴ confirming the presence of sodium carbonates in crude LH. As ³⁴⁵ LH was precipitated from aqueous solution, sodium carbonate ³⁴⁶ should exist in the form of monohydrate after drying at 60 °C. ³⁴⁷ Under this assumption, monohydrate sodium carbonate ³⁴⁸



Figure 4. Thermal characteristics of crude LH (a,b) and SHLP (c,d): (a,c) DSC and (b,d) TGA thermographs.

349 content in the crude LH precipitate would be 50.1%, leaving 350 49.9% to be lignin and hemicelluloses (LH). Such a unique 351 composition allowed LH to be directly pyrolyzed into AC, without the need to add additional activating chemicals, making 352 353 this pathway superior than previously reported ap-354 proaches.¹⁶⁻¹⁸ SHLP consisted of 16.6% C, 51.7% O, and 29.7% Si, showing its elemental compositions to be mainly 355 356 silica (64%) and lignin/hemicelluloses/PEO (36%). This three-357 step toluene/ethanol, NaClO2, and KOH isolation process isolated 5.3% waxes and organic extracts, 36% highly pure 358 cellulose,³ 2.8% LH, and 20.7% SHL, or 65% of precursors, 359 leaving about 35% rice straw components as soluble from 360 361 NaClO₂ oxidation and KOH dissolution in the remaining 362 filtrates.

The chars pyrolyzed from LH after dilute HCl washing 363 364 consisted of 81.7% C, 15.8% O, and 2.5% Si, confirming 365 substantial carbon with little silica impurities. After calcinating 366 SHLP at 500 °C for 0.5 h, only 47% Si and 53% O were 367 detected, while no carbon was present. This mass percentage is equivalent to a 1:2 Si/O atomic ratio or that of pure silica. 368

371 372 373 374 375 377

f4

f4

Thermal Properties of Extracts. The thermograms of LH 369 370 showed a very sharp exothermic peak at 60 °C and a small exotherm around 105 °C (Figure 4a) with respective 24.4% and 5.3% mass losses (Figure 4b), reflecting the transition from sodium carbonate decahydrate to monohydrate as well as moisture loss from lignin and hemicelluloses. The endotherm at 300 °C is attributed to the decomposition of hemicelluloses, 376 while two exotherms at 430 and 475 °C were likely from decomposition and further charring of lignin, corresponding to 378 the major 19.2% mass loss, that is, from 70.7% at 105 °C to 379 51.5% at 550 °C. The solid mass was lowered to 47.6% when 380 heated to 800 °C initially and slightly lowered to 45.6% when 381 held for additional 30 min.

The thermal analysis further supported the compositions 382 derived from EDX that the crude LH at ambient temperature 383 was a mixture of sodium carbonate monohydrate and 384 decahydrate as well as lignin/hemicelluloses. Assuming the 385 mass loss at 105 °C is solely caused by the transformation of 386 sodium carbonate monohydrate to the anhydrous form, the 387 weight percentages of sodium carbonate monohydrate and Na 388 in crude LH were estimated to be 48.3% and 17.9%, 389 respectively (eqs 1 and 2), very close to the 50.1% estimated 390 from EDX elemental analysis. These values are slightly higher 391 than the atomic compositions due to the missing hydrogen in 392 the EDX. The composition of crude LH precipitate at 105 °C 393 would thus be estimated to be 44.4% anhydrous sodium 394 carbonate and 55.6% lignin/hemicelluloses (eq 3). Therefore, 395 the final pyrolyzed product could be a mixture of lignin char 396 and various forms of sodium, sodium carbonate, sodium oxide, 397 and elementary sodium, all of the latters removed in the wash 398 to obtain 1.3% activated carbon (AC) from the starting rice 399 straw. 400

SHLP showed an endothermic shoulder below 105 °C 401 (Figure 4c) accompanied by a slight 4% mass loss from 402 moisture (Figure 4d). A broad exothermic peak centered at 350 403 $^{\circ}$ C was observed along with a significant 33% mass loss, that is, $_{404}$ from 96% at 105 °C to 63% at 500 °C, due to thermal 405 decomposition of hemicelluloses/lignin/PEO. This quantity 406 was consistent with the 64% silica from the EDX analysis. 407 When held at 500 °C for 0.5 h, the DSC thermograph baseline 408 shifted endothermically, while the solid mass slightly decreased 409 from 63% to 61.5%, indicating further combustion of the 410 residual carbon char, obtaining 14% pure silica from the starting 411 rice straw. 412



Figure 5. SEM (a,b,d,e) and TEM (c,f) of (a) LH, (b,c) AC, (d) SHLP, and (e,f) silica. Insets in SEM: scale bar = 20 µm.

$$Na_{2}CO_{3}\bullet H_{2}O_{(at\,60\ ^{\circ}C)}(wt\%) = \frac{Mass loss from Na_{2}CO_{3}\bullet H_{2}O to Na_{2}CO_{3}(wt\%) \times Molar mass of Na_{2}CO_{3}\bullet H_{2}O}{Molar mass of H_{2}O \times Mass percentage of solid residue_{(at\,60\ ^{\circ}C)}} = \frac{5.3 \text{ wt\%} \times 124}{18 \times 0.756} = 48.3 (wt\%) \quad (1)$$

$$Na_{(at\,60\ ^{\circ}C)}(wt\%) = Na_{2}CO_{3}\bullet H_{2}O_{(at\,60\ ^{\circ}C)}(wt\%) \times \frac{Molar mass of Na \times 2}{Molar mass of Na_{2}CO_{3}\bullet H_{2}O} = 48.3 \times \frac{46}{124} = 17.9 (wt\%) \quad (2)$$

$$Na_{(at\,60\ ^{\circ}C)}(wt\%) = Na_{2}CO_{3}\bullet H_{2}O_{(at\,60\ ^{\circ}C)}(wt\%) \times \frac{Molar mass of Na_{2}CO_{3}\bullet H_{2}O}{Molar mass of Na_{2}CO_{3}\bullet H_{2}O} = 48.3 \times \frac{46}{124} = 17.9 (wt\%) \quad (2)$$

$$Na_{2}CO_{3(at\,60\,^{\circ}C)}(wt\%) = \frac{Na_{2}CO_{3(at\,60\,^{\circ}C)}(wt\%)}{Na_{2}CO_{3(at\,60\,^{\circ}C)}(wt\%) + (100 - Na_{2}CO_{3}\bullet H_{2}O_{(at\,60\,^{\circ}C)}(wt\%))} \times 100 = \frac{41.3}{41.3 + 100 - 48.3} \times 100 = 44.4 (wt\%)$$
(4)

424 On the basis of the TGA analysis, the LH component 425 contained 51% sodium salt or about 2.8% yield of lignin/ 426 hemicelluloses from NaClO₂ dissolution of rice straw (e.g., 427 filtrate 1). SHLP consisted of 61.5% silica, 29.3% lignin/ 428 hemicelluloses, and 9.2% PEO or 14% silica and 6.7% lignin/ 429 hemicelluloses yields from the KOH dissolution stream (e.g., 430 filtrate 2). From these and our prior report on cellulose,³ this 431 three-step toluene/ethanol acidic NaClO₂–KOH process has 432 shown to isolate 5.3% waxes and organic extracts, 36% celluloses, 9.5% lignin/hemicelluloses, and 14% silica from 433 rice straw, that is, 65% of rice straw. The lignin/hemicelluloses 434 and silica were eventually converted to 1.3% AC and 14% pure 435 silica.

Surface and Morphology of Activated Carbon and $_{437}$ Silica. The LH powders were gray in color and appeared as $_{438}$ packed particulates in over 20 μ m sizes (Figure 5a). After $_{439}$ fs pyrolysis at 800 °C for 30 min, the obtained AC appeared as $_{440}$ irregularly shaped porous clusters of 100–500 nm-sized 441



Figure 6. Nitrogen adsorption-desorption isotherms of (a,b,c,d) of LH and AC (800 °C, 0.5 h, N_2) and (e,f) SHLP and silica (500 °C, 0.5 h, air): (a,e) isotherm, (b) BJH neck-size distribution, (c,f) BJH pore/cavity width distribution, and (d) micropore hydraulic diameter distribution.

442 particles (Figure 5b), which are much smaller than the ACs in ⁴⁴³ the previous studies.^{16–18} These much shorter diffusion lengths contribute to faster adsorption rate, which is a desirable 444 property of AC absorbent. The TEM images showed the semi-445 crescent AC particles had about a 450 nm Feret diameter. The 446 randomly distributed slit-shaped micropores and small 447 mesopores were observed within the particle (Figure 5c). 448 Such porosity was generated by sodium carbonate activation as 449 450 well as pyrolytic decomposition of hemicelluloses and removal 451 of sodium byproducts from washing.

The SEM showed smaller and more narrowly distributed SHLP particles with increasing PEO concentrations (Figure S2, Supporting Information). Semi-spherical particles from 100 to 454 200 nm sizes were obtained at 0.2% PEO. With increasing PEO 455 concentrations to 0.35%, more uniformly sized nanospheres 456 between 100 and 150 nm were formed (Figure 5d-f), possibly 457 due to better capping of particles preventing them from 458 aggregating and/or growing into larger particles. At 0.5 wt % 459 PEO and above, however, the particles became irregularly 460 shaped and polydispersed in sizes less than 50 nm, possibly 461 from easy sintering during calcination. Only the SLHP particles 462 precipitated from 0.35% acidified PEO remained similar in size 463 after calcinations. Such a PEO precipitation approach has 464 proven to be fast and effective, without the need for the time- 465

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	BET surface area (m²/g)	BJH adsorption cumulative surface area between 1.7 and 300 nm width (m^2/g)	single-point desorption pore volume (cm³/g)	BJH adsorption cumulative volume of pore between 1.7 and 300 nm (cm ³ /g)	<i>t</i> -plot-derived micropore surface area between 0.7 ando 2 nm width (m ² /g)	<i>t</i> -plot-derived micropore volume (cm ³ /g)
LH	3	3	0.006	0.006	0	0
AC	997	249	0.90	0.58	592	0.27
SHLP	7.9	6.6	0.015	0.020	0	0
silica	7.2	6.1	0.014	0.019	0	0

466 consuming aging process. The obtained silica spheres were 467 much more uniform in both size and shape than those 468 previously reported.^{17,20} The high size uniformity is crucial to 469 fine-tune the properties of silica particles to be used as additive/ 470 fillers for composites or further functionalized for other 471 applications.

Porous Structure of Activated Carbon and Silica. The 472 473 as-prepared LH powders appeared flaky and showed a H2 474 hysteresis in a BET nitrogen adsorption isotherm (Figure 6a) 475 with a very low surface area of 3 m²/g and a pore volume of 476 0.006 cm³/g, indicative of their nonporous nature (Table 2). 477 The extremely low surface area was attributed to secondary 478 porosity, for example, interparticulate mesopores distributed 479 between 3 and 30 nm, and peaked at 10 nm as shown in the 480 pore width distribution (Figure 6b). Carbonization at 800 °C 481 for 30 min significantly increased the BET surface area of the 482 submicrometer-sized AC particles to 997 m²/g and total pore $_{483}$ volume to 0.9 cm³/g, that is, a specific surface area over 330 484 times of that of the LH precursor. The BJH surface area and 485 pore volume of the meso/macropores between 1.7 and 300 nm 486 were estimated to be 249 m^2/g and 0.58 cm^3/g , respectively, 487 and those of the *t*-plot-derived micropores below 2 nm were 488 approximated to be 592 m²/g and 0.27 cm³/g, respectively 489 (Table 2) Therefore, the surface area was mostly attributed to 490 the micropores, while the pore volume was attributed to the 491 larger mesopores and macropores. The type IV isotherm with 492 noticeable H4 hysteresis observed on AC (Figure 6a) suggests a ⁴⁹³ slit-like micro/mesoporous structure.³⁵ The micropores ranged 494 from 0.8 to 1.6 nm in diameter, peaking at 0.95 nm (Figure 6d), 495 whereas the mesopores were bimodally distributed between 2 496 and 6 nm as well as between 10 nm to submicrometer (Figure 497 6c), consistent with TEM observations (Figure 5c). The 498 adsorption-desorption isotherms exhibited characteristic step-499 down at 0.45 P/P_0 , while the pore width distribution derived 500 from the desorption branch showed a distinct peak around 4 501 nm from cavitation of nitrogen during desorption (Figure 6b), ⁵⁰² indicating mesopores connected by slit-like micropores ⁵⁰³ consistent to similar observations by others.^{36,37} The internal 504 micropores and mesopores contributed mostly to the high surface area and pore volume of AC, while the large 505 506 macropores likely from interparticle spacing (secondary porosity) as shown in the SEM (Figure 5b) contributed little. 507 Both SHLP and silica exhibited type II BET nitrogen 508 509 adsorption isotherms, typical of nonporous and macroporous 510 materials with weak affinities to nitrogen (Figure 6e). Both had very similarly low BET surface area and pore volume of about 7 511 $512 \text{ m}^2/\text{g}$ and 0.015 cm³/g, respectively, as well as BJH mesoporous surface area and pore volume of about 6 m^2/g and 0.002 cm³/g, 513 514 respectively (Table 2). Only macropores larger than 100 nm 515 were observed in both (Figure 6f). There was no evidence of 516 mesopores. These observations were consistent with their low 517 surface areas and pore volume that were mainly attributed to 518 the external particle surfaces and interparticulate spacing. In 519 calcination of SHLP, the organic hemicelluloses, lignin, and

PEO components were removed, leaving only silica with no 520 porosity as indicated by the nitrogen adsorption—desorption 521 analysis. This nonporous structure suggests that the organic 522 components are either mainly capped on the silica particle 523 surface and/or well mixed with silica to create micropores that 524 collapse during calcinations or are below the 0.3 nm nitrogen 525 adsorption—detection level of the instrument used.

CONCLUSION

This study has demonstrated the feasibility to optimally isolate 528 the lignin, hemicellulose, and silica components from rice straw, 529 along cellulose previously reported,³ and efficiently convert 530 them into advanced materials. An efficient three-step toluene/ 531 ethanol, NaClO₂, and KOH isolation process has been 532 successfully devised for isolating two-thirds of crude rice 533 straw into at least 36% pure cellulose from rice straw while 534 generating two filtrates, 2.8% lignin-rich LH and 20.7% silica- 535 hemicellulose-rich SHL as precursors for activated carbon and 536 silica particles, respectively. The NaClO₂ dissolution stream 537 from the dewaxed rice straw contains oxidized lignin and 538 hemicelluloses as carbon precursors as well as sodium 539 carbonates as activating chemicals that can be precipitated for 540 direct carbonization at 800 °C to yield 1.3% activated carbon 541 particles with submicrometer sizes between 100 and 500 nm as 542 well as high specific surface area (997 m^2/g) and pore volume 543 $(0.90 \text{ cm}^3/\text{g})$. The KOH dissolution stream from NaClO₂- 544 treated rice straw contains mainly potassium silicate and a small 545 amount of lignin/hemicelluloses and could be precipitated by 546 dilute (0.35%) acidified poly(ethylene oxide) and calcinated to 547 yield 14% pure, nonporous, silica nanospheres in 100 to 120 548 nm diameters. This is the first report of highly efficient 549 processes that optimally isolated and fully utilized all major rice 550 straw components, the highest quantity agricultural crop 551 byproduct in the world. Together with nanocellulose products 552 reported,³ 80% of the isolated components or 51.5% of crude 553 rice straw was converted into high quality particulates of highly 554 porous activated carbon particles and uniformly sized non- 555 porous silica nanoparticles. These high quality nanomaterials 556 can provide broadly available feedstock for advanced materials. 557

ASSOCIATED CONTENT	558
Supporting Information	559
EDX of LH after calcination and SEMs of silica particles	560
precipitated by acidified PEO of different concentrations before	561
and after calcination. This material is available free of charge via	562
the Internet at http://pubs.acs.org.	563
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Notes	567
The authors declare no competing financial interest.	568

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