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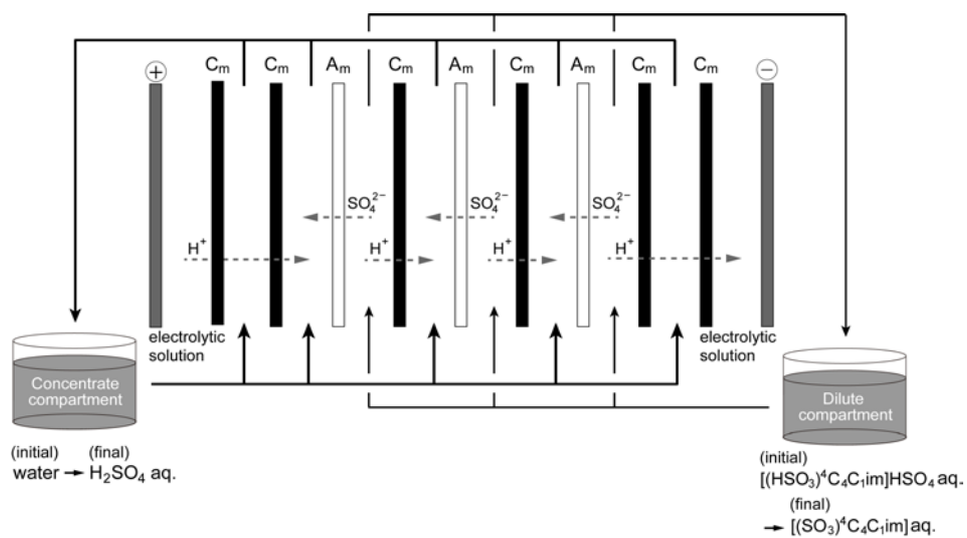
# Supporting Information

## Efficient hydrolysis of polysaccharides in bagasse by *in situ* synthesis of an acidic ionic liquid after pretreatment

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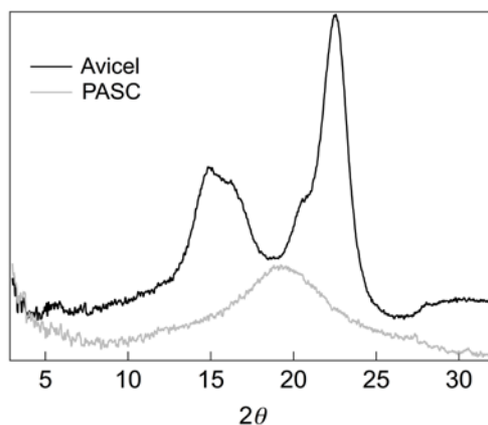
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<b>Figure S1.</b> Schematic diagram for the electrodialysis of $[(\text{HSO}_3)^4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$ . The actual apparatus comprises five pairs of anion exchange and cation exchange membranes. ....	<b>S2</b>
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**Figure S1.** Schematic diagram for the electrodedialysis of  $[(\text{HSO}_3)_4\text{C}_4\text{C}_{1\text{im}}]\text{HSO}_4$ . The actual apparatus comprises five pairs of anion exchange and cation exchange membranes ( $A_m$  and  $C_m$ , in the figure).

Figure S2 shows the X-ray scattering spectra of Avicel and PASC. Avicel has high crystallinity, and its crystallinity index<sup>[1]</sup> was 0.82. PASC shows the peak of amorphous cellulose, and its crystallinity was 0.00. It indicates that the efficient hydrolysis of PASC is due to the low crystallinity.



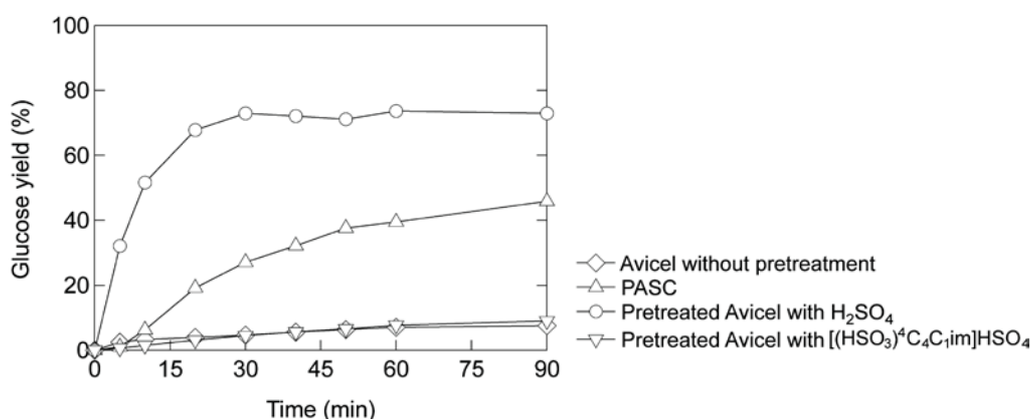
**Figure S2.** X-ray scattering spectra of Avicel and PASC.

<sup>[1]</sup> Park, S.; Baker, J. O.; Himmel, M. E.; Parilla, P. A.; Johnson, D. K. Cellulose crystallinity index: measurement techniques and their impact on interpreting cellulase performance. *Biothechnol. Biofuels* **2010**, *3*, 1-10.

Figure S3 shows time courses of glucose yield during hydrolysis of microcrystalline cellulose (Avicel) and phosphoric acid swollen cellulose (PASC) as partially amorphous cellulose in 1.0 M  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  solution under microwave at 100 °C. Avicel was hydrolyzed with the maximum yield of 8 % within 90min. PASC was hydrolyzed with the maximum yield of 46 % within 90min. Cellulose crystallinity was confirmed to prevent efficient hydrolysis with the  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  solution.

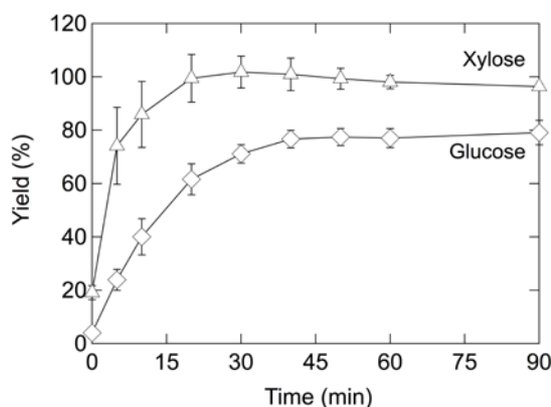
Figure S3 also shows time course of glucose yield when Avicel was pretreated with 72 wt%  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  solution followed by subsequent hydrolysis at 100 °C in the 1.0 M  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  solution solution. The glucose yield was 10% at 90min and similar to that without pretreatment. It shows that  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  does not have decrystallizing ability to cellulose.

When Avicel was pretreated with 72 wt%  $\text{H}_2\text{SO}_4$  solution and hydrolyzed with  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  at 100 °C, the glucose yield was 73% within 30min. These results clearly indicate that the hydrolytic method in this study is highly efficient from both viewpoints of yield and reaction time.



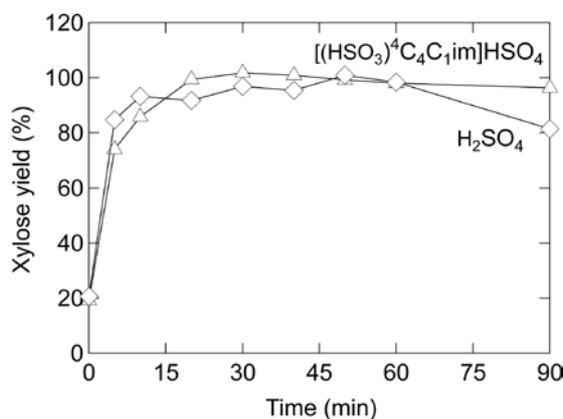
**Figure S3.** Time course of glucose yield during hydrolysis of Avicel and PASC with 1.0 M of  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  at 100 °C with or without pretreatments.

Figure S4 shows error bars in yield of glucose and xylose during hydrolysis in a 1.0 M  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  solution at 100 °C (basically the data is the same as shown in Figure 2). The error was not large in yield of both glucose and xylose.



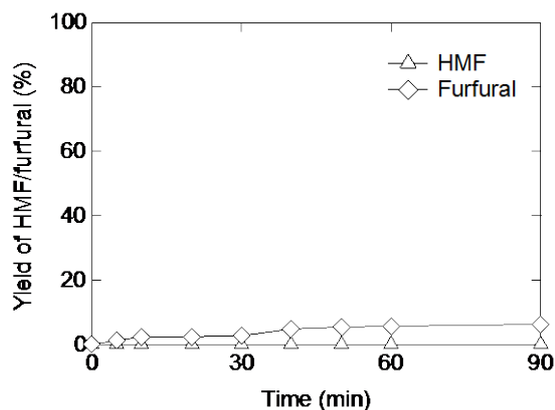
**Figure S4.** Time courses of yield of glucose and xylose during hydrolysis of bagasse pretreated with  $\text{H}_2\text{SO}_4$  in a 1.0 M  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  solution under microwave heating at 100 °C with error bars.

Figure S5 shows time course of xylose yield during hydrolysis of bagasse with 1.0 M of  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  or  $\text{H}_2\text{SO}_4$  at 100 °C under microwave after pretreatment with  $\text{H}_2\text{SO}_4$ . In the case of  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$ , maximum yield was 102 % at 30min while that with  $\text{H}_2\text{SO}_4$  was 100 % at 50min.



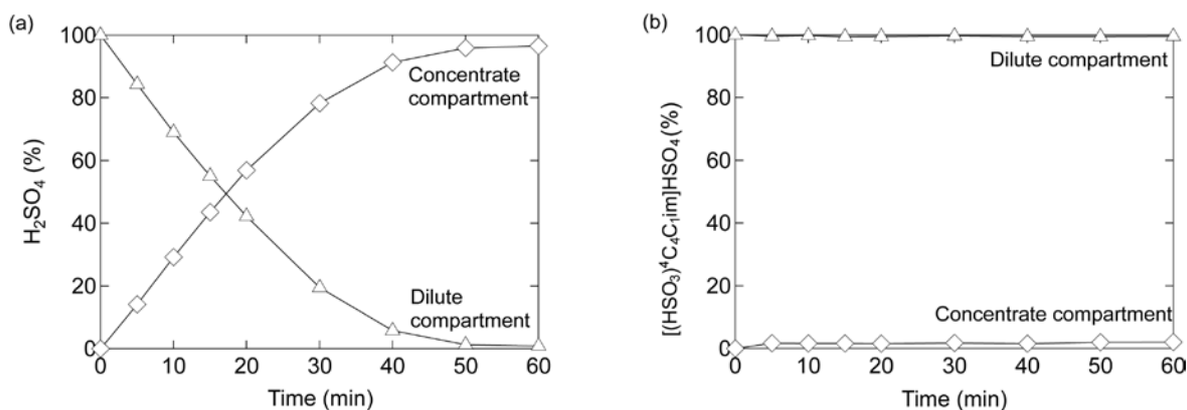
**Figure S5.** Time course of xylose yield during hydrolysis of bagasse with 1.0 M of  $[(\text{HSO}_3)_4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  or  $\text{H}_2\text{SO}_4$  at 100 °C under microwave after pretreatment with concentrated  $\text{H}_2\text{SO}_4$ .

Figure S6 shows time courses of yield of 5-(hydroxymethyl)furfural (HMF) and furfural during hydrolysis of the bagasse pretreated with H<sub>2</sub>SO<sub>4</sub> in 1.0 M [(HSO<sub>3</sub>)<sup>4</sup>C<sub>4</sub>C<sub>1</sub>im]HSO<sub>4</sub> solution under microwave at 100 °C. No HMF was detected during the hydrolysis. Little furfural generated during the hydrolysis and the yield was 6 % after 90min. These results clearly show that the present method does not lead to the significant decomposition of sugars unlike usual acid hydrolysis at high temperature.



**Figure S6.** Time courses of yield of HMF and furfural during hydrolysis of the bagasse pretreated with H<sub>2</sub>SO<sub>4</sub> in 1.0 M [(HSO<sub>3</sub>)<sup>4</sup>C<sub>4</sub>C<sub>1</sub>im]HSO<sub>4</sub> solution under microwave at 100 °C.

Figure S7 shows the time courses of concentration of  $\text{H}_2\text{SO}_4$  and  $[(\text{SO}_3)^4\text{C}_4\text{C}_1\text{im}]$  in the dilute and the concentrate compartments during electro dialysis of  $[(\text{HSO}_3)^4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$ . Almost  $\text{H}_2\text{SO}_4$  (97 %) was transferred from the dilute compartment to the concentration compartment after 60 min while almost  $[(\text{SO}_3)^4\text{C}_4\text{C}_1\text{im}]$  (99 %) remained in the dilute compartment. Electro dialysis is a suitable method to separate  $[(\text{HSO}_3)^4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$  into  $\text{H}_2\text{SO}_4$  and  $[(\text{SO}_3)^4\text{C}_4\text{C}_1\text{im}]$  for reuse.



**Figure S7.** Time courses of concentration of (a)  $\text{H}_2\text{SO}_4$  and (b)  $[(\text{SO}_3)^4\text{C}_4\text{C}_1\text{im}]$  in dilute and concentrate compartments during electro dialysis of  $[(\text{HSO}_3)^4\text{C}_4\text{C}_1\text{im}]\text{HSO}_4$ .