1	Enzyme kinetics approach to assess biocatalyst inhibition and deactivation caused
2	by [bmim][Cl] ionic liquid during cellulose hydrolysis
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### 20 Abstract

22	The aim of this work was to study the inhibition and deactivation of commercial					
23	enzyme cocktail (Cellic <sup>®</sup> Htec2) in the presence of [bmim][Cl] ionic liquid employing					
24	model cellulosic substrate, carboxymethyl cellulose (CMC). It turned out from the					
25	experiements – relying on enzyme kinetics appproach – that [bmim][Cl] could act as a					
26	competitive inhibitor. Furthermore, depending on the process conditions i.e. contact of					
27	enzyme solution with high concentration [bmim][Cl], severe biocatalyst inactivation					
28	should be also taken into account as a potential risk during the enzymatic cellulose					
29	hydrolysis even in as short process times as few minutes.					
30						
31	Keywords: lignocellulose, pretreatment, ionic liquid, enzymatic hydrolysis, inhibition,					
32	deactivation					
33						

### 1. Introduction

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Lignocellulose, as an inexpensive and abundant, renewable material is a 38 trending feedstock for new generation energy production technologies. Among its 39 constituents, cellulose is the most important one to be considered for biotechnological 40 applications (Liu et al., 2012). Cellulose is a well-known polymer molecule, built-up 41 by individual glucose monomers, which represent the primary source of fermentable 42 sugars. Unfortunately, the direct conversion of lignocelluloses to valuable products i.e. 43 44 biofuels is hindered by the limited access of biocatalysts to the cellulose regions (Kumar et al., 2015; Xu et al., 2016a). Therefore, to open up the complex, recalcitrant 45 structure and enhance the solubilization of such organic matter, a pretreatment is 46 47 recommended (Vancov et al., 2012). To accomplish this step, the use of ionic liquids 48 (IL) has been widely proposed because of their recognized potential for the efficient structural transformation of the crystalline cellulose, which increases the efficiency of 49 50 its consecutive enzymatic hydrolysis and thus, the glucose yield (Brandt et al., 2013; Maki-Arvela et al., 2010; Raj et al., 2016). Basically, two main routes to realize 51 52 enzyme-catalyzed hydrolyzis after dissolving cellulose in IL have been suggested. In the first one, the cellulose is separated from the IL solution and this, so-called 53 regenerated cellulose (having reduced degree of crystallinity and increased porosity) is 54 subjected for subsequent enzymatic hydrolysis (Tan et al., 2011; Zhao et al., 2009). In 55 56 the other one, referred as *one-pot or single-step* approach, the enzymatic hydrolysis of dissolved cellulose is conducted by adding cellulase enzymes directly to the IL phase 57 in a water-based buffer solution (Gunny et al., 2014; He et al., 2016; Shi et al., 2011; 58

Xu et al., 2014, 2016b), after which the glucose released can be converted to biofuels
or other alternative products such as glucose esters (Findrik et al., 2016).

To make sure that the best IL is chosen for the pretreatment of a particular 61 lignocellulose, a screening is advised (Zavrel et al., 2009) since its appropriateness is 62 dependent on factors such as the physicochemical properties, i.e. the anion and cation 63 constituents of the particular IL (Raj et al., 2016). It was found that among the various 64 ILs, those consisting of an imidazolium-ring, such as [bmim][Cl] are promising 65 candidates to dissolve considerable amount of cellulose (Engel et al., 2012; Tan et al., 66 2011; Zhao et al., 2009). However, besides that high capacity, ILs should meet 67 additional criteria e.g. biocompatibility with cellulase enzymes performing the 68 cellulose saccharification (Li et al., 2010). In this regard, issues with ILs (including 69 [bmim][Cl]) were observed attributed to their reportedly negative influence on 70 hydrolytic enzyme activity and stability (Engel et al., 2012; Li et al., 2013; Lozano et 71 al., 2011; Ouellet et al., 2011; Park et al., 2012; Salvador et al., 2010; Turner et al., 72 73 2003; Xiao et al., 2012; Zhao et al., 2009).

Although the enzyme inhibition and deactivation by ILs from these examples 74 are known to occur, to our knowledge, no thorough study has been dedicated so far to 75 evaluate these phenomena i.e. in terms of the mechanisms involved for the inhibition. 76 Therefore, in this work, comprehensive enzyme kinetics (well-known approach of the 77 biocatalysis area) was applied to get new insights to the enzymatic cellulose hydrolysis 78 process using a model cellulose substrate (carboxymethyl cellulose, CMC) and a 79 commercial enzyme solution (Cellic<sup>®</sup> Htec2) in the absence and presence of 80 [bmim][Cl], as one of the most widely employed ILs for lignocellulose pretreatment. 81

82	The novelty of this investigation is the findings presented for the first time using the						
83	enzyme kinetic approach, which can thus have the potential to contribute to the						
84	international knowledge and further expansion of the research area regarding the						
85	application of ionic liquids in the (ligno)cellulose-based biorefinery concept.						
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87	2. Materials and methods						
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90	2.1. Enzyme kinetics						
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02	2.1.1 Cellulose hydrolysis in the absence of [hmim][Cl] ionic liquid -						
92	2.1.1. Centrose nyurorysis in the absence of [binnin][Ci] fonce inquite						
93	Michaelis-Menten kinetics						
93 94	Michaelis-Menten kinetics						
93 94 95	Michaelis-Menten kinetics The enzymatic cellulose hydrolysis was considered as a multi-step process: (i)						
93 93 94 95 96	Michaelis-Menten kinetics The enzymatic cellulose hydrolysis was considered as a multi-step process: (i) the free (cellulase) enzyme (E) binds to the substrate (S) as reversible reaction (E+S),						
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92 93 94 95 96 97 98 99 99 100	<ul> <li>Michaelis-Menten kinetics</li> <li>The enzymatic cellulose hydrolysis was considered as a multi-step process: (i)</li> <li>the free (cellulase) enzyme (E) binds to the substrate (S) as reversible reaction (E+S),</li> <li>subsequently (ii) an enzyme-substrate complex (ES) is formed and afterwards, as a</li> <li>results of irreversible (ES) breakdown, (E) and product (P) are released (Zhang et al.,</li> <li>2010), in accordance with the classical Michaelis-Menten kinetics (Johnson and</li> <li>Goody, 2011). This model was thus applied to assess the enzymatic hydrolysis</li> </ul>						

$$103 \quad V = \frac{Vmax\left[S\right]}{Ks + \left[S\right]} \tag{1}$$

105	where V and $V_{max}$ are the actual (initial) and maximal (initial) product formation rates
106	(g product/g enzyme-min), respectively, and [S] is the actual (initial) substrate
107	concentration (g/L). $K_s$ is denoted as the half-saturation constant (g/L), equaling to an
108	[S] where $V=V_{max}/2$ . Experimental conditions i.e. in terms of range of [S] tested can be
109	found in Section 2.2.
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111	2.1.2. Cellulose hydrolysis in the presence of [bmim][Cl] ionic liquid –
112	inhibition kinetics
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114	Reversible enzyme inhibition may occur via four different reaction mechanisms
115	such as competitive (Eq. 2), uncompetitive (Eq. 3), linear-mixed (Eq. 4) and non-
116	competitive (special case of Eq.4, where $K_i = K'_i$ ) (Marangoni, 2003).
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118	$V = \frac{V_{max} [S]}{Ks\left(1 + \frac{[I]}{K_i}\right) + [S]} $ (2)
119	
	17 [c]

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$$V = \frac{V_{max}[S]}{Ks + \left(1 + \frac{[I]}{K_{i}}\right)[S]}$$
(3)

122 
$$V = \frac{V_{max} [S]}{Ks \left(1 + \frac{[l]}{K_i}\right) + \left(1 + \frac{[l]}{K'_i}\right) [S]}$$
(4)

where  $K_i$  and  $K'_i$  are the enzyme-inhibitor and (enzyme-substrate)-inhibitor dissociation constants (g/L), respectively, while [*I*] is the actual (initial) inhibitor concentration (g/L).

To determine which of the above is actually involved when an inhibition phenomenon is observed, a comprehensive kinetic study was carried out, requiring sets of experiments at various initial substrate and inhibitor (in this investigation, bmim][Cl] ionic liquid) concentrations (Marangoni, 2003). The related measurement details are specified in Section 2.2.

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### 133 **2.1.3.** Enzyme deactivation by undiluted [bmim][Cl] ionic liquid

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To study whether [bmim][Cl] ionic liquid – besides its potential inhibitory effect on the cellulose hydrolysis process – causes the deactivation of the enzyme used thoroughly in this work (see below in Section 2.2.), a first-order kinetics was taken (Eq. 5). It is a widely used model in the literature to investigate enzyme deactivation (Lencki et al., 1992; Sadana, 1988) and in particular, deactivation of cellulase in the course of cellulose hydrolysis (Zhang et al., 2010).

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142 
$$A = \frac{a_t}{a_0} = e^{-k_{de_1}t}$$
 (5)

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where the relative activity of the enzyme is *A*.  $a_t$  refers to the activity measured in the presence of the ionic liquid (with a particular concentration, herewith 50 mg/L, see Section 2.2.) after treating the enzyme with undiluted ionic liquid for various times (*t*). 147  $a_0$  is the activity measured in the presence of the ionic liquid (with a particular 148 concentration, herewith 50 mg/L) without previous incubation of enzyme with 149 undiluted ionic liquid (*t*=0, to be taken into account as baseline activity).  $k_{de1}$  is 150 denoted as the first-order rate constant of cellulase deactivation (min<sup>-1</sup>) and *e* is the 151 exponential term (2.718). By taking the natural logarithm of both sides in **Eq. 5** and 152 plotting ln *A* against *t*,  $k_{de1}$  is derived from the slope of the straight line (Lencki et al., 153 1992).

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### 155 **2.2.** Cellulose hydrolysis assays

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All cellulose hydrolysis measurements were performed in closed-top 250 mL 157 Erlenmeyer flasks (to prevent evaporation) with 100 mL total working volume, 158 ensuring vigorous stirring (400 rpm by magnetic bar) and constant 60 °C temperature. 159 Citrate buffer (100 mM, pH of 4.5) served as bulk phase thoroughly. The model 160 substrate used was low-viscosity carboxymethyl cellulose (CMC) (Sigma-Aldrich, 161 USA) with phisyco-chemical properties reported in the specification sheet available 162 for download at the manufacturer's official website, while Cellic<sup>®</sup> Htec2 163 (Novozymes<sup>®</sup>, Denmark) was utilized as enzyme source, which has an already 164 reported potential for the hydrolysis of cellulosic materials (Benjamin et al., 2014; Joe 165 et al., 2015; Song et al., 2014). Cellic<sup>®</sup> Htec2 is characterized with optimal working 166 pH and temperature of 4.5-5.5 and 60-75 °C, respectively (according to the application 167 sheet issued by the producer) and was provided in  $600(\pm 48)$  mg/L concentrations for 168

169 each tests. Data were always evaluated by considering the exact, actual mass of170 enzyme supplemented to the particular reaction vessel.

During the experiments without ionic liquid (Michaelis-Menten kinetics, Section 2.2.1.), substrate (CMC) concentrations investigated were 0.5, 2, 5 and 25 g/L. In case of enzyme inhibition tests (Section 2.2.2.), definite amounts of [bmim][Cl] (99 % purity, IoLiTec, Germany) were added (without further purification) to the citrate buffer to get the desired ionic liquid concentration (50, 100, 150, 200 and 250 mg/L) along with substrate concentrations of 1, 2.5 and 5 g/L.

In the course of enzyme deactivation study (Section 2.2.3),  $60(\pm 4.8)$  mg enzyme was first mixed with 5 mg (undiluted) [bmim][Cl] ionic liquid and held together for treatment (contact) times such as 1, 2, 3 and 10 minutes at 60 °C. Subsequently, the whole mixture was loaded to the above described citrate bufferbased reaction medium (resulting in an initial [bmim][Cl] and enzyme concentrations of 50 mg/L and  $600(\pm 48)$  mg/L, respectively) containing 2.5 g substrate/L.

183 Quantitative analysis of glucose (as final end-product of multi-step cellulose hydrolysis) released from cellulose hydrolysis was monitored based on the refractive 184 index (RI) change of the reaction mixture over time - relative to the RI of initial 185 reaction mixture (background) – pumped continuously (10 mL min<sup>-1</sup>) through Merck-186 Hitachi (RI-71) differential refractometer (at 37 °C), attached to the reaction vessel in a 187 closed-loop design. Preliminary calibration – to establish the relationship of RI change 188 and glucose concentrations - was accomplished by measuring the RI of reaction 189 mixture with well-defined glucose (reagent grade, Sigma-Aldrich, USA) contents. 190

191	Experiments were always commenced by the injection of the enzyme solution
192	to the reaction mixture. Kinetic parameters (i.e. $K_s$ , $V_{max}$ , $K_i$ , $kd_{el}$ ) were estimated by
193	linear regression using the least squares method in Matlab software.
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### 195 **3. Results and Discussion**

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## 197 3.1. Cellulose hydrolysis process kinetics without [bmim][Cl] ionic liquid 198

199 The progress curves obtained during the measurements can be seen in **Fig. 1**. It seems to infer that product (glucose) formation showed a directly proportional trend 200 with time, as reflected by the appreciably high  $R^2$  values (**Table 1**). Though to some 201 extent, various terms of experimental error e.g. data recording, observations, etc. 202 203 (Bakonyi et al., 2015) may make the determination of apparent, initial reaction rates vague, replicates undertaken (Fig. 1) indicated that the results were fairly reproducible 204 and consequently, basically reliable for further analysis. The confidential evaluation of 205 V is supposed to be performed using a good mass of data acquired within a limited 206 207 time, otherwise it can be misleading due to fact that initial cellulose hydrolysis rates decline remarkably as time passes (Carrillo et al., 2005). Hence, information recorded 208 in the first 10-15 minutes of the enzymatic hydrolysis tests was only taken into account 209 in this study. Similar period of time was considered by other researchers working in 210 this field, as well (Yeh et al., 2010). 211

It can be concluded from the time profiles of cellulose hydrolysis experiments (Fig. 1) that enhanced V values (slope of the trendlines fitted, Table 1) could be achieved along with increasing substrate concentrations. Applying the well-known Lineweaver-Burk double reciprocal plot ( $V^{-1}$  vs.  $[S]^{-1}$ ), the kinetic parameters of the Michaelis-Menten model could be computed and as a result,  $K_s$  and  $V_{max}$  were found as 2.57 g/L and 0.068 g glucose/g enzyme-min, respectively. The value of  $K_s$  is in the same order of magnitude reported by Yeh et al. (2010) for a range of microcrystalline cotton cellulose substrates and a cellulase enzyme produced by *T. reesei* ATCC 26921.

Generally speaking, the use of Michaelis-Menten kinetics for discussing the 220 experimental data requires a "mass of enzyme to mass of substrate ratio" <0.15, which 221 ensures that the mechanistic model is applicable (Bezerra and Dias, 2004, 2005) and 222 the substrate is not limiting. In our current investigation, apparent "mass of enzyme 223 mass to mass of substrate ratios" such as 0.024, 0.12, 0.3 and 1.2 were obtained, being 224 solely dependent on the initial substrate concentration because of the constant enzyme 225 loadings (600 mg/L). Although seemingly not all of them meet the criteria referred, a 226 decent correlation of the experimental outcomes with the fitted Michaelis-Menten 227 kinetics was established. This contradiction can be explained by considering the 228 properties of the enzyme source. Cellic® Htec2 is a commercial enzyme cocktail 229 preparation and not a highly purified cellulase. Such products can be seen as a mixture 230 of accessory enzymes such as cellulases and hemicellulases (Samayam and Schall, 231 2010), however, their exact composition and relative proportion of ingredients is 232 unknown (Barr et al., 2012). Thus, once a given mass of this enzyme solution is used 233 (e.g. 600 mg for every liter of reaction mixture in this study), only its limited portion 234 was actually the cellulase enzyme participating in the hydrolysis reaction, while the 235 rest consisted mainly of water, other enzymes i.e. endoxylanases (Benjamin et al., 236 2014; Joe et al., 2015; Song et al., 2014), etc. 237

Consequently, the real "cellulase enzyme mass to substrate mass ratio" was (probably orders of magnitudes) below the above defined threshold value (<0.15) even in the lower initial substrate concentration range (0.5-2 g/L), making the Michaelis-Menten theory valid within the respective experimental boundaries and leading to an adequate prediction of the hydrolysis process.

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# 3.2. On the inhibition of enzymatic cellulose hydrolysis by [bmim][Cl] ionic liquid

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It turned out in previous works examining cellulose hydrolysis in the mixture of 247 aqueous buffer and IL that the cellulose hydrolysis can be negatively affected by ionic 248 liquids (Kamiya et al. (2008; Turner et al., 2003). For instance, Kamiya et al. (2008) 249 summarized that biocatalyzed cellulose saccharification (applying commercialized 250 cellulase from *Trichoderma reesei*) was fully stopped in a citrate-buffer based reaction 251 252 mixture composing of >40 vol.% [emim][dep], meaning that the proportion of IL is a determining factor of the process. Moreover, another research carried out by Turner et 253 al. (2003) also confirmed the existence of IL-induced enzyme inhibition working with 254 [bmim][Cl], which was associated with the high, dissociated Cl<sup>-</sup> ion concentration in 255 256 the media, causing the interference of cellulose with ionic liquid.

Overall, it is clear by the literature examples that enzyme-mediated cellulose hydrolysis can be sensitive to the presence of ILs, including [bmim][Cl] employed thoroughly herewith. However, it is still of question which of the possible inhibition mechanisms (i.e. competitive, uncompetitive, mixed, non-competitive) stands behind once decreased reaction rates (as a side-effect of IL) are encountered. Therefore, a
throughout kinetic study was devoted to get new insights on this subject.

The analysis of the progress curves and consecutive use of Lineweaver-Burk 263 double reciprocal technique (Lineweaver and Burk, 1934) ( $V^1$  vs. [S]<sup>-1</sup>, along with 264 various inhibitor concentrations [1], 50-250 mg [Bmim][Cl]/L) yielded Fig. 2, which is 265 a clear identification of the competitive-type inhibition (Marangoni, 2003). Thus, in 266 the light of this diagnosis, the deviation of initial reaction velocities from those 267 projected by the Michaelis-Menten model (no inhibition occuring) took place by the 268 mechanism established in Eq. 2. When the data in Fig. 2 are subjected for an 269 estimation of the catalytic parameters, it can be observed that  $V_{max}$  remains unaffected 270 in accordance with the feature of competitive inhibition, meanwhile  $K_S$  is influenced 271 by the inhibitor concentration ([1]) and the enzyme-inhibitor dissociation constant  $(K_i)$ 272 (Eq. 2). Since the experimental results suggest competitive inhibition caused by the 273 [bmim][Cl] ionic liquid, it means that the inhibitor competes for the substrate-binding 274 275 site of the enzyme and when an enzyme-inhibitor complex is formed rather than an enzyme-substrate one, the enzyme cannot express its catalytic activity. Overall, it 276 seems to be the case that the cellulase enzymes present in Cellic<sup>®</sup> Htec2 suffer from an 277 inhibition in the presence of [bmim][Cl], even at low [1]. General strategies to 278 suppress this type of inhibition include the use of higher enzyme as well as substrate 279 loadings to increase the probability of enzyme and substrate interactions rather than 280 that of the enzyme and the inhibitor. In summary of the kinetic analysis on enzyme 281 inhibition tests (Fig. 2, Table 2), numerical estimate for  $K_i$  was obtained as 0.163 g/L. 282 In general, the increase of  $[I]/K_i$  will more notably reduce the affinity of the enzyme to 283

the substrate, causing the concomitant raise of  $K_s$ . For instance, once the [bmim][Cl] concentration in the reaction mixture reaches to  $K_i$ , it will double the original (uninhibited)  $K_s$  value, in accordance with the mechanistic model for competitive inhibition in **Eq. 2**.

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#### **3.3.** Cellulase enzyme deactivation by [bmim][Cl] ionic liquid

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The important work done by Turner et al. (2003) implied that not only an inhibition, but even deactivation of cellulase may occur when they are contacted with ionic liquids in the solution. As it was deduced, enzyme denaturation can be a possible threat and in such a case, overcoming strategies may be unable to refold, reactivate the cellulase. Deactivation (or in other words, irreversible inhibition) is one of the major phenomena known to slow the cellulose hydrolysis rate down (Bansal et al., 2009), leading essentially to a process of limited efficiency.

To judge whether the inactivation of cellulase enzymes in Cellic<sup>®</sup> Htec2 is to be 299 taken into account, tests (in accordance with Section 2.2.) were undertaken and the 300 enzyme solution was mixed and incubated with [bmim][Cl] for various durations, 301 followed by the determination of actual, initial reaction rates to reveal residual enzyme 302 activity. It can be drawn from Fig. 3 that the treatment time of enzyme by IL 303 considerably affected the achievable V values, providing a good indication that contact 304 of the cellulase with undiluted [bmim][Cl] – even when it happened for short times 305 only e.g. 1 min – was accompanied by the relative loss of enzyme activity. Taking the 306 advantage of the formula expressed in Eq. 5, kinetic analysis was conducted to 307 compute the rate constant of deactivation. Since in the bottom part of Fig. 3 a well-308

fitting linear relationship can be found, the first-order kinetics to predict biocatalyst 309 deactivation seems to be valid (Lencki et al., 1992) and  $k_{del}$  as 0.132 min<sup>-1</sup> could be 310 determined. A correlation of similar shape between relative cellulase activity and 311 various residence times in 10 % [bmim][Cl] was communicated by Salvador et al. 312 (2010), as well. However, in that investigation (Salvador et al., 2010), residual enzyme 313 activities were much closer to the original, since even after 40 minutes of contact with 314 IL, reduction was only 13-14 %, suggesting that the inhibition was mostly of 315 reversible nature and ascribed primarily to the change of thermodynamic water activity 316 under the different conditions. However, it was also shown that increased IL 317 concentrations are severely disadvantageous from an enzyme activity point of view 318 (Salvador et al., 2010; Xiao et al., 2012). Significant loss of Celluclast<sup>®</sup> cellulase 319 enzyme activity was observed by Engel et al. (2010) too in reaction mixtures 320 comprising of commercialized ionic liquids (10%), including [bmim][Cl]. 321

Keeping this in mind together with our results, it would appear that when 322 323 cellulase enzymes (even for short times) are exposed to concentrated ILs, deactivation effect may take over. This would explain why in our experiments the reaction rates 324 have notably dropped, causing that the enzyme lost approximately 25 % of its activity 325 relatively to its control  $(a_0)$  even after only 1 min of pre-incubation with undiluted 326 [bmim][Cl]. This phenomenon can be interesting when cellulase hydrolysis is 327 performed in situ, referred as one-pot process design. In this arrangement, the 328 cellulosic raw material is first pretreated with concentrated ionic liquid to help the 329 dissolution of the polysaccharide fractions and in that way, provide better accessibility 330 for the enzymes participating in their hydrolysis. Thereafter, the hydrolytic enzymes – 331

carried by an aqueous buffer solution – are loaded to the same vessel containing the 332 cellulose being dissolved in the IL (Shi et al., 2013). Many ionic liquids, depending on 333 their structural features, are hydrophilic (including [bmim][Cl]) and therefore soluble 334 in water to certain degrees (Huddleston et al., 2001). In these cases, ILs can be more or 335 less homogenously distributed in the whole reaction medium, which makes it possible 336 to decrease their concentrations to a sufficiently low threshold level. However, 337 appropriate dilution factor should be selected since even its low concentrations could 338 exhibit a hindering impact on the hydrolysis for the particular example of [bmim][Cl] 339 used in this work, as demonstrated by our results in Section 3.2. Besides dilution 340 method (Li et al., 2013), the engineering and application of cellulases with satisfactory 341 robustness to work under harsh conditions can be proposed (Nordwald et al., 2014; 342 Raddadi et al., 2013) i.e. via the development of halophilic, IL-tolerant cellulases 343 (Gunny et al., 2014; Xu et al., 2014; 2016b), their immobilization (Xu et al., 2016c) 344 which may better withstand the negative impact of ILs. In other cases, when the IL is 345 346 basically immiscible with the water-based solution, a biphasic (separated) system is formed (Kuroda et al., 2016), and saccharification can take place on the phase 347 boundary, where the probability of an enzyme-concentrated IL encounter is higher, 348 representing a threat on the time-stability of the process. For cellulose saccharification 349 and subsequent utilization technologies relying on regenerated cellulose, ionic liquid 350 residues should be removed as much as possible to avoid inhibition by a washing 351 process (Li et al., 2013; Ouellet et al., 2011) 352

354 Conclusions

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In this work, the effect of [bmim][Cl] ionic liquid on cellulose hydrolysis 356 catalyzed by Cellic® Htec2 enzyme solution was studied. It was found that the 357 enzymatic reaction could be fairly described by the Michaelis-Menten kinetics when 358 [bmim][Cl] ionic liquid was absent. However, the presence of this IL in low 359 concentrations significantly hindered the process via competitive inhibition, supported 360 by the kinetic evaluation. Furthermore, it was demonstrated that contacting the enzyme 361 with highly-concentrated IL even for short times could induce irreversible inhibition 362 (deactivation), which should be considered as an important aspect of technology 363 design. 364

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522							

526	Fig. 1 – Typical progress curves of cellulose hydrolysis (actual experimental					
527	conditions: absence of [bmim][Cl] ionic liquid, varied initial substrate					
528	concentration)					
529	blue diamond and red square: [S]=0.5 g/L; green triangle and purple cross: [S]=2 g/L;					
530	blue asterisk and orange dot: [S]=5 g/L; pink dash: [S]=25 g/L					
531						
532	Fig. 2 – Kinetic evaluation on enzyme inhibition test results obtained during					
533	cellulose hydrolysis with different initial [bmim][Cl] ionic liquid (inhibitor) and					
534	substrate concentrations					
535	blue diamond: no [bmim][Cl] added; red square: 50 mg/L [bmim][Cl]; green triangle:					
536	100 mg/L [bmim][Cl]; purple cross: 150 mg/L [bmim][Cl]; blue asterisk: 200 mg/L					
537	[bmim][Cl]; orange dot: 250 mg/L [bmim][Cl]					
538						
539	Fig. 3 –Kinetic evaluation on enzyme deactivation experiments					
540	blue dots: initial reaction rate; red square: natural logarithm of relative enzyme activity					

Table 1 – Example for progress curve analysis to deliver the initial reaction rates (V)
using the dataset of Fig. 1.

	Fitte				
	1. repetition		2. repetition		-
[S]	slope (g glucose/g		slope (g glucose/g		V (g glucose/g enzyme-
(g/L)	enzyme-s)	$\mathbf{R}^2$	enzyme-s)	$R^2$	min)
0.5	0.000180	0.98	0.000181	0.91	0.011
2	0.000430	0.19	0.000425	0.95	0.026
5	0.000837	0.99	0.000843	0.99	0.051
25	0.001013	0.99			0.060

Table 2 – Characteristics of fitted trendlines (Fig. 2) as the function of [bmim][Cl] concentration to estimate the maximal initial reaction rate (*Vmax*) and enzymeinhibitor dissociation constant ( $K_i$ ) for competitive inhibition model. The corresponding [S]<sup>-1</sup> range studied is presented in Fig. 2.

550

	[b	[bmim][Cl] concentration (mg/L)				
	50	100	150	200	250	
slope	45.71	54.71	63.99	81.04	96.27	
intercept	14.82	14.82	14.82	14.82	14.82	
$\mathbf{R}^2$	0.98	0.97	0.94	0.99	0.99	

551











