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1 **Dry chemo-mechanical pretreatment of chickpea straw: effect and**
2 **optimization of experimental parameters to improve hydrolysis yields**

3

4 Mouna Aouine^{1,2}, Doha El Alami², Abdellatif Haggoud¹, Saad Ibsouda Koraichi¹,
5 Laurent Roumeas³, and Abdellatif Barakat^{2,3*}

6 ¹Laboratory of Microbial Biotechnology and Bioactive Molecules, Faculty of Sciences and Techniques,
7 Sidi Mohammed Ben Abdellah University, Fez, Morocco.

8 ²AgroBioSciences Department, Mohammed VI Polytechnic University, Benguerir, Morocco.

9 ³IATE, INRAE, University of Montpellier, Agro Institut, INRA, Montpellier, France.

10

11 **Abstract**

12

13 Dry chemo-mechanical (DCM) pretreatment is an interesting eco-friendly approach for
14 bioconversion of lignocellulosic biomass into sugars and other valuable molecules. In
15 this study, different dry DCM pretreatments were developed using a combination of
16 alkaline and vibro-milling “VBM”. High-resolution fractional factorial 2^{k-1} design
17 (FFD) was applied to evaluate statistically the effects of NaOH concentration (2-10 %),
18 impregnation ratio (20-50 %), milling time (10-60 min), milling frequency (15-30 Hz),
19 and ball diameter (1 or 2.5 cm) on reducing sugars release from chickpea straw (CS).
20 The optimal conditions ensuring the maximum concentration of reducing sugars (374.70
21 mg/g biomass) after 72 h of enzymatic hydrolysis were 10 % of NaOH, impregnation of
22 50 %, 60 min of VBM, frequency VBM of 30 Hz, and ball diameter of 2.5 cm.
23 Furthermore, an ethanol concentration of 17.81 g/L was obtained after simultaneous
24 saccharification and fermentation of the pretreated CS under the defined optimized
25 conditions.

26 **Keywords:** dry chemo-mechanical pretreatment, chickpea straw, reducing sugars,
27 fractional factorial design, ethanol.

28 **Abbreviations**

29 **CrI** Crystallinity index

30 **CS** chickpea straw

31 **DCM** dry chemo-mechanical

32 **FFD 2⁵⁻¹ design** fractional factorial design

33 **SEM** scanning electron microscopy

34 **SSF** simultaneous saccharification and fermentation

35 **VBM** Vibro-ball milling

36 **1. Introduction**

37 In Morocco, chickpea production tops the list of grain legumes cultivated. Its
38 agriculture occupies more than 65.9 hectares per 1000 cultivated hectares and its
39 production estimated at 440.6 quintals per 1000 quintals (Agricultural companion,
40 2015- 2016). Along with the production of chickpeas, there is also the generation of
41 straw co-product, which is widely used as an alternative forage in ruminant diet due to
42 its fibers and proteins content. In recent years, agricultural residues such as chickpeas
43 straw are increasingly considered in the production of biomolecules as a sustainable
44 way of managing these by-products. Moreover, when using land for sugarcane
45 production dedicated for bioethanol, accessibility to food will be limited. Hence, the
46 importance of valorizing agro-food residues. Production of carboxylates, biofuels

47 (biogas, bioethanol and biodiesel), energy (syngas via gasification) and valuable
48 chemicals can be carried out using agricultural residues (Barakat et al., 2014c). In
49 addition, the availability of chickpea straw (CS), and its low cost make this feedstock an
50 interesting lignocellulosic biomass for biofuel and chemical production. However, this
51 biomass is not much exploited in literature compared to wheat and rice straw and
52 bagasse.

53 Lignocellulosic biomass is a complex matrix mainly composed of cellulose,
54 hemicellulose, and lignin (Laurichesse and Avérous, 2014). The bioconversion of this
55 recalcitrant structure into fermentable sugars involves a pretreatment process in order to
56 eliminate lignin and make cellulose and hemicellulose more accessible to chemical or
57 enzymatic attack during saccharification and fermentation. In general, the pretreatment
58 of lignocellulosic biomass for bioethanol production at pilot and industrial scale is the
59 most important expensive compared to other steps and operations in biorefinery (Rocha-
60 Meneses et al., 2019; Rajendran et al., 2018). Therefore, it is essential to optimize this
61 process in function to biomass properties and valorization route. Several pretreatment
62 technologies have been developed so far, the most common are thermal, alkaline, dilute
63 acid, steam explosion and organosolv pretreatments (Memon and Memon, 2020; Liu
64 and Chen, 2017; Vargas et al., 2015). However, these technologies present some
65 disadvantages in terms of energy consumption, corrosion of equipments, and generation
66 of inhibiting molecules such as furans and involves some additional steps such as
67 separation and purification (Licari et al., 2016).

68 Dry chemo-mechanical (DCM) pretreatment is an eco-friendly approach with a
69 combination of alkaline and mechanical pretreatments (Barakat et al., 2014a).

70 Technically, lignocellulosic biomass first undergoes a chemical treatment by alkaline

71 impregnation at high solid loads, followed by a mechanical size reduction (Barakat et
72 al., 2014a). Studies has demonstrated that dry chemical pretreatment increases sugars
73 yield, reduces energy requirements, and decrease effluents generation (Chuetor et al.,
74 2019; Barakat et al., 2014). To take full advantage of this pretreatment procedure, it is
75 necessary to identify and optimize the parameters that affect the efficiency of the
76 pretreatment. Lazuka et al. (2017) investigated the effect of particle size and sodium
77 hydroxide (NaOH) impregnation on the microbial transformation of wheat straw. Their
78 results showed that the highest reducing sugar content was obtained after the treatment
79 with NaOH at 100 μm . These results are inconclusive, which focused only on two
80 factors, NaOH concentration and particle size. Thus, the integration of other factors
81 such as milling time, impregnation time, milling frequency, alkaline concentration...and
82 their interaction effects on biomass proprieties is very importance to develop an
83 efficiency mechanochemical pretreatment.

84 Up to day, there is no study in literature describing the use experiments design for
85 evaluating the influence of different DCM factors and their interaction on biomass
86 proprieties and accessibility by enzymes. Hence, in this study, a high-resolution
87 fractional factorial 2^{5-1} design (FFD 2^{5-1}) was implemented to evaluate the effects of
88 NaOH concentration, impregnation, milling time and frequency, and ball diameter on
89 biomass proprieties and sugars release from CS biomass. The effectiveness of
90 pretreatment was monitored by measuring reducing sugar concentrations after
91 enzymatic hydrolysis.

92 **2. Materials and methods**

93 **2.1. Biomass**

94 CS biomass was kindly harvested from a farm (Fez region, Morocco). CS samples were
95 air-dried and coarsely cut into a particle size of less than 1 mm using a hammer mill
96 (Retsch SM 100, Germany). Raw CS contained 27 % cellulose, 18 % hemicelluloses,
97 and 26 % lignin. The samples were dried to a moisture content of 8- 10 % and sealed in
98 plastic bags until use.

99 2.2. Pretreatment

100 Dry mechanochemical (DCM) pretreatment was carried out in two stages according to
101 (Barakat et al., 2014a). Briefly, CS samples were first impregnated with NaOH solution
102 for 5 hours at room temperature and then dried at 50 °C. The dried biomass was ground
103 using a vibrating ball mill (VBM).

104 2.3. Experimental design

105 A two-level FFD 2^{5-1} design was used for evaluating the experiments conditions that
106 affect reducing sugars release from CS biomass. As shown in **Table 1**, the selected
107 independent variables and their variation ranges were: NaOH concentration (g/100 g
108 biomass, 2 – 10 % wt), impregnation (water/ biomass, 20- 50 % wt), milling time (10-60
109 min), milling frequency (15-30 Hz) and ball diameter (1 and 2.5 cm). Those conditions
110 were based on (Barakat et al., 2014a) work. The concentration of reducing sugars
111 released (mg/g biomass) was retained as response variable. Data analysis was performed
112 via Minitab 18 software.

113 2.4. Enzymatic hydrolysis

114 The enzymatic hydrolysis was carried out in 50 mM sodium acetate buffer (pH 5) at a
115 solid loading of 10 % (w/v) of the biomass. The reaction mixture contained also a
116 cellulase Cellic CTec2 (Novozymes) with an activity loading of 20 FPU/g and an endo-
117 1,4-xylanase from *Trichoderma longibrachiatum* (Sigma-Aldrich) with an activity

118 loading of 20 U/g. Sodium azide (3 g/L) was added to prevent any microbial
119 contamination. Incubation was done at 50 ° C for 72 h with stirring at 100 rpm.
120 Reducing sugars contents in saccharification liquors were determined colorimetrically
121 using the dinitrosalicylic acid (DNS) method (Miller, 1959) and the absorbance was
122 measured at 540 nm. The experiments were performed in duplicate.

123 2.5. X-ray diffraction

124 The crystallinity of raw and pretreated CS samples was analyzed using an X-ray
125 diffractometer (XPERT-PRO) with a Cu tube at an accelerating voltage of 40 kV and a
126 current of 30 mA. Scans were conducted at a 2θ angle, between 8 and 28°, with a step
127 of 0.01°, and at a scan rate of 2°/min. The crystallinity index (CrI) of samples was
128 calculated according to the following equation (Segal et al., 1959):

$$129 \quad CrI (\%) = \frac{100*(I_{002} - I_{001})}{I_{002}}$$

130 where I₀₀₂ is the intensity of the diffraction from the 002 plane at 2θ= 22°, and I₀₀₁ is
131 the peak intensity of the amorphous zone at 2θ = 16°, in diffractogram.

132 2.6. Scanning electron microscopy

133 The morphological alterations occurred in the surface of CS biomass after the
134 pretreatment conducted at the optimal conditions were evaluated with scanning electron
135 microscopy (SEM) analysis. The analysis of SEM was performed using a scanning
136 electron microscope (JSM-IT500 InTouchScope™).

137

138 2.7. Simultaneous saccharification and fermentation (SSF) of raw and pretreated

139 CS

140 *2.7.1. Microorganism and growth conditions*

141 The fermentative yeast used in this study was *Pichia kudriavzevii*, a thermotolerant
142 strain selected and identified in our laboratory. Yeast cells were grown overnight at 35
143 °C in 100 mL of YPD liquid medium (10 g/L of yeast extract, 20 g/L of peptone, and 20
144 g/L of glucose) with an orbital agitation of 150 rpm. After incubation, cells were
145 collected by centrifugation for 10 min at 10,000 rpm, washed twice with sterile
146 deionized water and suspended in 0.9 % NaCl solution with an absorbance at 600 nm.

147 2.7.2. *Simultaneous saccharification and fermentation (SSF)*

148 Raw and pretreated CS was subjected to simultaneous saccharification and fermentation
149 (SSF) under the optimal conditions to produce bioethanol. For SSF experiments, 10% of
150 the biomass were suspended in 50 mM sodium acetate buffer (pH 5). Nutrients such as
151 yeast extract 5 g/L, peptone 5 g/L, K₂HPO₄ 1 g/L, and MgSO₄ 1 g/L were
152 supplemented. The fermentation medium was sterilized for 15 min at 120 °C. *P.*
153 *kudriavzevii* (10 %, v/v) and an enzymatic cocktail of Cellic CTec2 (Novozymes, 20
154 FPU/g) and endo- 1,4-xylanase from *Trichoderma longibrachiatum* (Sigma-Aldrich, 20
155 U/g) were subsequently added. SSF assays were carried out at 42 °C and 150 rpm. The
156 samples taken after 24, 48 and 72 h were centrifuged for 10 min at 10,000 rpm. The
157 concentration of bioethanol in the supernatant was estimated by the chromic acid
158 method (Caputi et al., 1968) and the absorbance was measured at 600 nm using a
159 spectrophotometer. SSF assays were performed in duplicate.

160

161 **3. Results and discussion**

162 3.1. Assessment of DCM pretreatment on reducing sugars release and cellulose
163 crystallinity

164 **Figure 1** displays the concentrations of reducing sugars released after 72 h of enzymatic
165 hydrolysis at 50 °C of untreated and DCM pretreated CS, as well as the crystallinity
166 index. The experiment conditions of each run are described in **Table 2**. It' can be seen
167 in **Table 2** and **Figure1** that, the concentration of reducing sugars ranged from 274.61
168 to 374.70 mg/g biomass. The highest concentration was observed in run 6, in which CS
169 sample was treated with 10 % of NaOH, 50 % of impregnation, 60 min of milling, 30
170 Hz of frequency, and 2.5 cm as ball diameter. However, the run 15 resulted in the
171 lowest concentration (276.50 mg / g biomass) (**Table 2**). Generally, all the trials
172 improved reducing sugars production over the untreated CS (274.61 mg / g biomass).
173 For the crystallinity measurements, the majority of the pretreated CS samples showed a
174 reduction in the crystallinity index, run 6 provided an important decrease reaching a
175 value of 23.50 % compared to 38.71% obtained with the raw CS.

176 Previously, sugar release was raised by 5-fold due to NaOH (2%v/v) addition under
177 60°C, and this increase was further enhanced when increasing temperature to 121°C
178 (McIntosh and Vancov, 2011). Overall, using alkali reagent at high doses (6-20%) can
179 lead to dissolution of nondegraded polysaccharides, while high temperature
180 pretreatment can enhance cellulose hydrolysis. However, under more severe
181 temperature conditions, enzymatic digestibility of lignocellulosic matrix can strongly
182 decrease due to phenolic and furans compounds (Carrere et al., 2016).

183 DCM pretreatment is based on the combined action of NaOH which solubilize lignin
184 and swells the structural components of lignocellulose and milling which reduces the
185 particle size and decrease cellulose crystallinity (Chuetor et al., 2021). The highest
186 concentration obtained in this study was 374.70 mg/g biomass. These results are in
187 agreement with previous studies reporting a remarkable enhancement in fermentable

188 sugars release from lignocellulosic biomass when DCM was applied (Lazuka et al.,
189 2017; Sambusiti et al., 2015). A significant negative correlation was observed between
190 reducing sugar yields and crystallinity index ($R = -0.51$). These findings join Barakat's
191 (2014) statement that DCM pretreatment reduces cellulose crystallinity and solubilize
192 lignin and acetyl groups, which increase saccharification rate. Similarly, Qu et al.
193 (2017) demonstrated that ball mill-assisted alkaline peroxide pretreatment of wheat
194 straw samples changed the crystalline cellulose into an amorphous form, resulting in
195 improved enzymatic hydrolysis due to the higher surface accessibility. However, the
196 only concern is that the solid/liquid ratio can drastically increase the energy
197 consumption during drying before mechanical pretreatment. An optimization of this
198 ratio is therefore necessary to improve the sustainability of this combination (Barakat et
199 al., 2014b).

200 3.2.DCM efficiency using fractional factorial design

201 To better understand the influence of each factor on the efficiency of DCM
202 pretreatment, a 2^{5-1} FFD was implemented. The factors evaluated were NaOH
203 concentration, impregnation, milling time, milling frequency, and ball diameter. The
204 concentration of reducing sugars released (mg/g biomass) was retained as response
205 variable (Table 2).

206 3.2.1. Analysis of the main and interaction effects

207 **Figure 2** represents the Normal plot and the Pareto chart of the standardized effects.
208 The graphical in **Figure 2** indicate that, all significant factors are positive,
209 demonstrating a direct correlation with the release of reducing sugars. In terms of main
210 effects, milling time (C) and ball diameter (E) were the most influencing variables.
211 Rezende et al. (2018) applied an FFD 2^{5-1} to optimize a sequential acid–alkali

212 pretreatment of elephant grass leaves. They found that extending ball milling time
213 before the chemical treatment has a positive significant effect on reducing sugar release.
214 Zhang et al. (2021) also proved that ball milling time factor evaluated in ball mill-
215 assisted alkaline peroxide pretreatment influenced the compositions, particle sizes,
216 morphology, and crystallinity of corn stalks biomass and hence, increased the yield of
217 xylo-oligosaccharides and fermentable sugars. For ball diameter, Khumalo et al. (2006)
218 claimed that larger sized grinding media would break complex matrix faster. In general,
219 when the diameter of the ball is large, the impact and compression of the biomass
220 particles trapped either between the grinding balls or between the grinding ball and the
221 mill casing is important.

222 In terms of binary interactions, NaOH concentration (A) impregnation (B) had the most
223 significant impact on reducing sugar production from CS biomass. Impregnation (B) is
224 individually insignificant variable, but it had a pronounced effect when is coupled with
225 NaOH concentration. Several researchers have approved that NaOH is very effective at
226 disrupting the lignocellulosic structure (Valles et al., 2021, Mukherjee et al., 2018, Kim
227 et al., 2016). Thus, in DCM pretreatment when the biomass particles are well sprayed
228 with high concentrations of NaOH, more lignin is solubilized, resulting in high
229 fermentable sugar yields. Huang et al. 2019 statically assessed during a wet-ball milling
230 the effect of various concentrations of NaOH on the enzymatic saccharification of
231 bagasse and pennisetum. Their results showed that glucose yields increased with
232 increasing NaOH concentration.

233 3.2.2. *Statistical modelling and ANOVA*

234 Analysis of variance (ANOVA) is reproduced in **Table 3**. Calculations were performed
235 at a confidence level of 95% (p-value < 0.05). The obtained F-value of 26.84 was

236 greater than the tabulated F-value of 3.35, while the p-value of 0.003 was less than 0.05.
237 These results indicate that the model selected in this study is statistically significant. As
238 predicted by the Pareto chart, the most significant influence on reducing sugars release
239 was exerted by milling time (p-value of 0,001), ball diameter (p-value of 0,002), and by
240 the coupled effect of NaOH concentration x impregnation (p-value of 0,002). The model
241 R-square was 0.9866 and the adjusted R-square was 0.9499, ensuring a good fitness of
242 the model. **Figure 3** depicts the analysis of the residuals. For the normal probability
243 plot (**Figure 3a**), the residuals were adjusted to a straight-line. As for the graph
244 comparing the residuals versus fits (**Figure 3b**), no pattern for the residual distribution
245 was recognized. Therefore, the constant variance assumption was satisfied.

246 3.2.3. *Response surface and validation of the optimum conditions for reducing* 247 *sugars release*

248 Contour and 3D surface graphs were used to visualize the interactions of the significant
249 independent factors (**Figure 4**). The categorical factor (ball diameter) was maintained
250 on 2.5 cm. As observed, a shift to higher values of factors provides higher
251 concentrations of reducing sugars, values exceeding 335 mg/g biomass.
252 According to the regression equation (1), the predicted maximum concentration of
253 reducing sugars of 391.58 mg/g biomass is attained by adopting the following optimal
254 conditions: NaOH concentration of 10 %, impregnation of 50 %, milling time of 60
255 min, frequency of 30 Hz, and ball diameter of 2.5 cm. To ensure the model validity, two
256 new experiments were conducted under the optimal conditions given above. The
257 average concentration of reducing sugars after 72 h of saccharification was 376. 68
258 mg/g biomass. This value is in reasonable agreement with the predicted concentration.
259 It was evident that applying FFD approach was advantageous since the importance of

260 each factor was established. Subsequently, the generation of fermentable sugars from
261 lignocellulosic biomass pretreated by the DCM method can be optimized by fixing the
262 settings of the factors on their maximum.

263

$$\begin{aligned} \text{Concentration of} &= 363.1 - 5.032 A - 1.676 B - 0.818 C - 1.453 D - 4.97 E & (1) \\ \text{reducing sugars (mg/g} &+ 0.1360 A*B + 0.0481 A *C + 0.00943 B*C + 0.0326 B*D \\ \text{biomass)} &+ 0.3716 B*E + 0.02622 C*D \end{aligned}$$

264

265 3.3. Effect of DCM pretreatment on surface morphology of CS biomass

266 **Figure 5** shows the scanning electron microscopy (SEM) images of untreated and
267 processed CS under the optimal conditions identified by the experimental design. As it
268 is clear, DCM pretreatment caused an obvious disruption of the structure morphology of
269 the biomass. In the raw CS (**Figure 5a**), the surface appeared dense, smooth without
270 apparent damage. Conversely, in **Figure 5b** the pretreatment gave an ultrafine powder
271 showing that the fibers were completely fragmented and the surface had cracks,
272 micropores and wrinkles. These observations are in agreement with those reported by
273 Zhang et al. (2021), Chen et al. (2019), and Qu et al. (2017), who indicate that
274 mechanochemical pretreatments generate fibrillated structures and cause a significant
275 breakdown of the plant cell walls, which facilitates the accessibility of enzymes at the
276 hydrolysis stage.

277 3.4. Simultaneous saccharification and fermentation (SSF) of the pretreated CS 278 for bioethanol production

279 **Figure 6** summarizes the SSF profile of *P. kudriavzevii* using untreated and pretreated
280 CS under optimal conditions. The totality of ethanol was produced after 48 h of
281 fermentation. For pretreated CS, the maximum ethanol concentration was 17.81 g/L, in
282 contrast, the raw CS provided only a concentration in ethanol of 9.54 g/L. In the study
283 of Sambusiti et al. (2015), Alkali-mechanically pretreated sugarcane bagasse yielded an
284 ethanol concentration of up to 7.45 g/L during the SSF process. Another study of
285 Monlau et al. (2019) demonstrated that mechanical fractionation by vibro-ball milling
286 (VBM) of solid separated digestate of agricultural biogas plants resulted in 4.9 g/L of
287 ethanol after the SSF test. In comparison with conventional alkaline pretreatment of
288 agricultural residues, Singh and Kumar (2020) obtained an ethanol concentration of
289 18.07 g/L from sodium carbonate pretreated rice straw. Likewise, a concentration of
290 17.26 g/L (0.48g of ethanol/g of glucose and xylose consumed) was generated from
291 alkaline pretreated sugarcane bagasse after SSF (Hilares et al., 2017). Overall, DCM
292 pretreatment followed by SSF technique has given promising results in terms of
293 bioethanol production from lignocellulosic biomass. It is worth noting that this
294 combination allows a reduction in processing time and steps, which is highly
295 recommended in transformation plants.

296 **4. Conclusion**

297 In this study, a high resolution 2^{5-1} FFD allowed simultaneous evaluation of five
298 variables and their coupled interactions in DCM pretreatment for reducing sugars
299 recovery from CS biomass. The results indicate that milling time, and ball diameter and
300 the interaction between NaOH concentration and impregnation had the most significant
301 effect on DCM efficiency. The highest concentration of reducing sugars was obtained
302 during pretreatment under 10 % of NaOH, impregnation of 50 %, milling time of 60

303 min, frequency of 30 Hz, and ball diameter of 2.5 cm. Moreover, an ethanol yield of
304 17.81 g/L was obtained through the SSF process using the thermotolerant yeast *P.*
305 *kudriavzevii* and CS pretreated under the optimized conditions. The statistical
306 optimization of DCM pretreatment enables intelligent eco-valorization of
307 lignocellulosic raw materials, particularly for integrated biorefineries.

308

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418 **Figure captions**

419 **Figure 1.** Reducing sugars (in mg/ g of biomass) released from DCM pretreated CS
420 samples after 72 h of enzymatic hydrolysis, and their crystallinity index according to
421 FFD 2^{5-1} design matrix. Untreated CS was included for comparison.

422 **Figure 2.** Normal plot of the standardized effects (a), Pareto chart of the standardized
423 effects (b). Panels were obtained after implementing backward selection using α -value
424 to enter of 0.05.

425 **Figure 3.** Plots of residuals (a) Normal probability (b) Residuals versus fits.

426 **Figure 4.** Contour and surface plots showing the interactions between the significant
427 independent factors.

428 **Figure 5.** SEM images of (a) untreated CS, (b) pretreated CS with: NaOH concentration
429 of 10 %, impregnation of 50 %, milling time of 60 min, milling frequency of 30 Hz, and
430 ball diameter of 2.5 cm. The numbers 1-3 in panel names denote images of the same
431 area at increasing magnifications

432 **Figure 6.** Ethanol production profile (in g/L) of *P. kudriavzevii* during SSF of untreated
433 and pretreated CS biomass.

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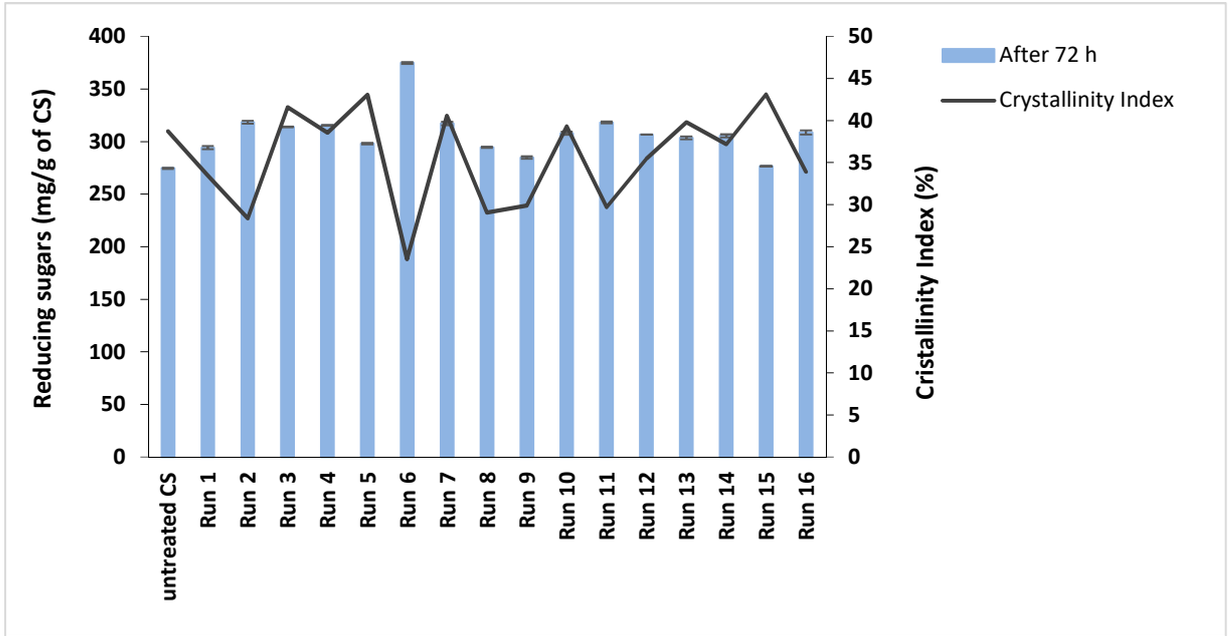
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442 **Figure 1**

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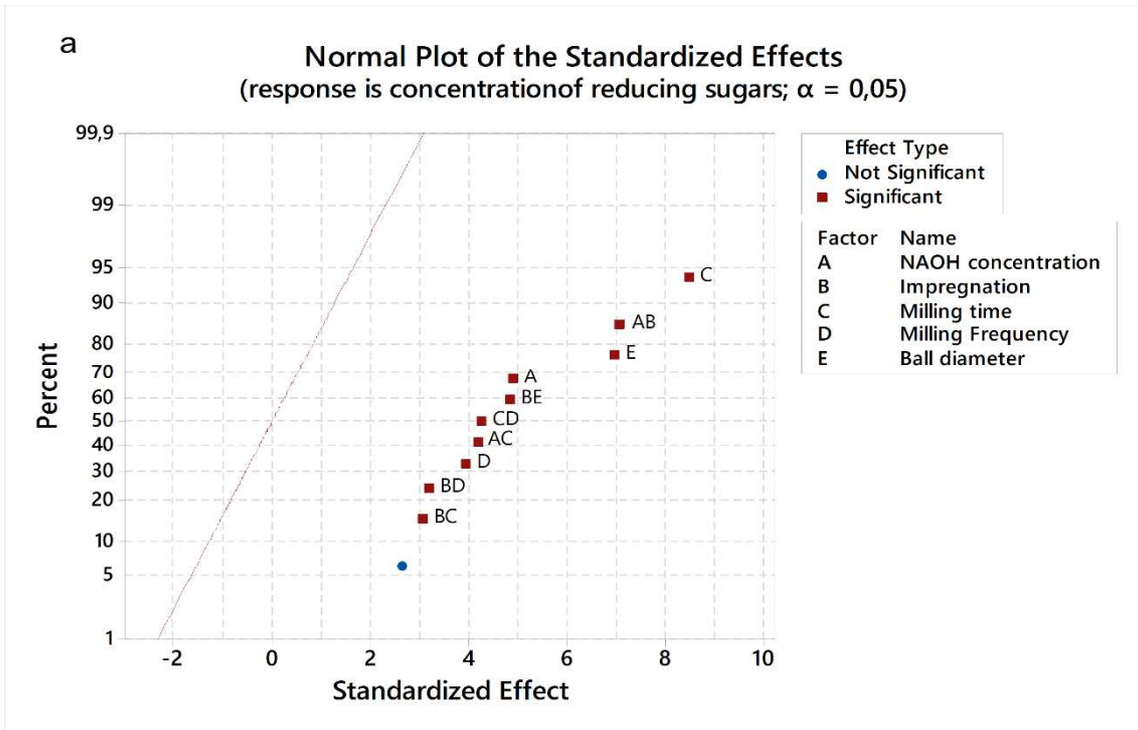
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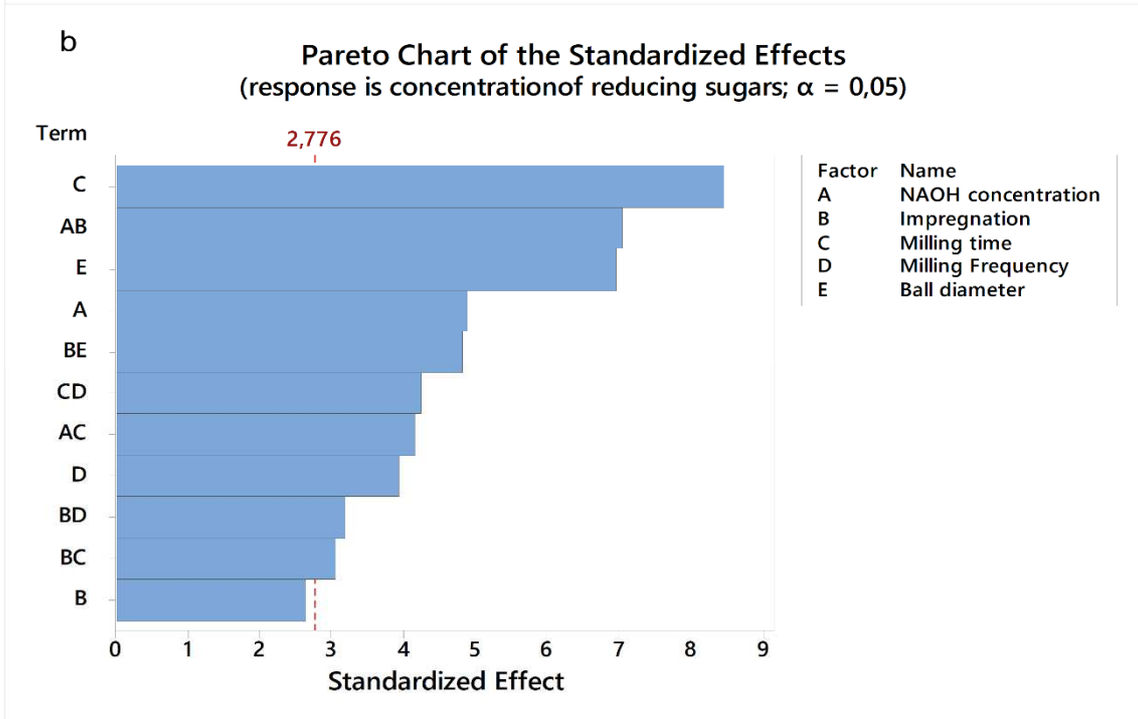
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460 **Figure 2**



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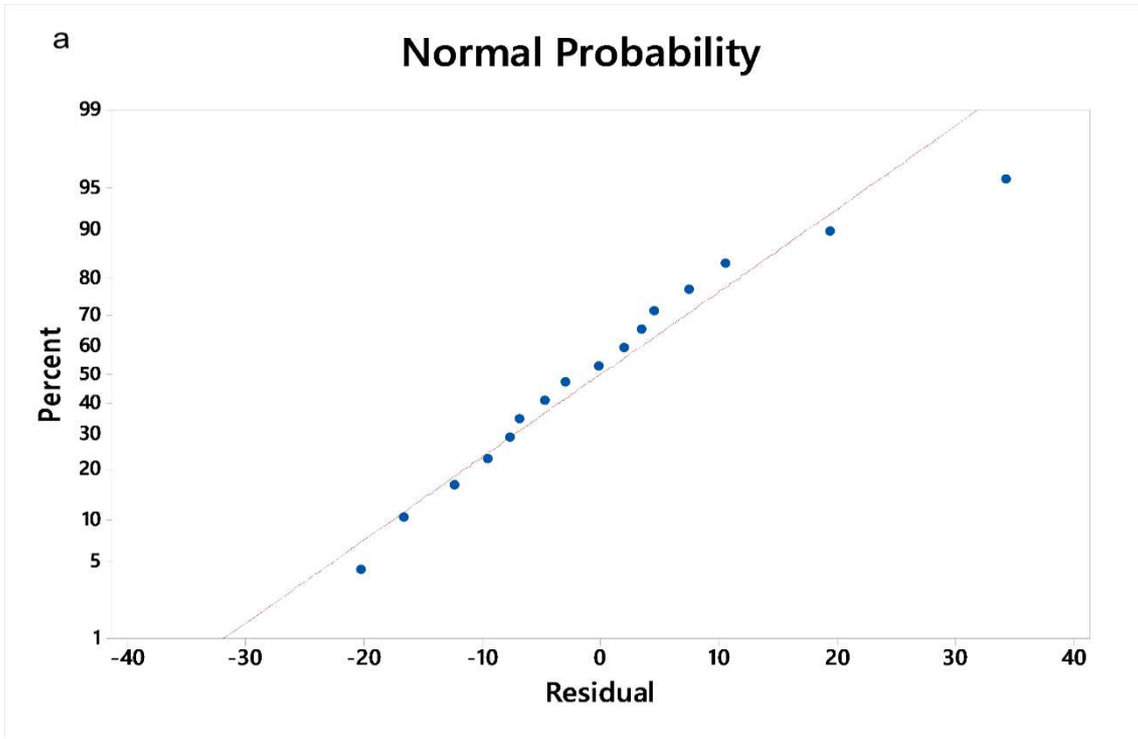
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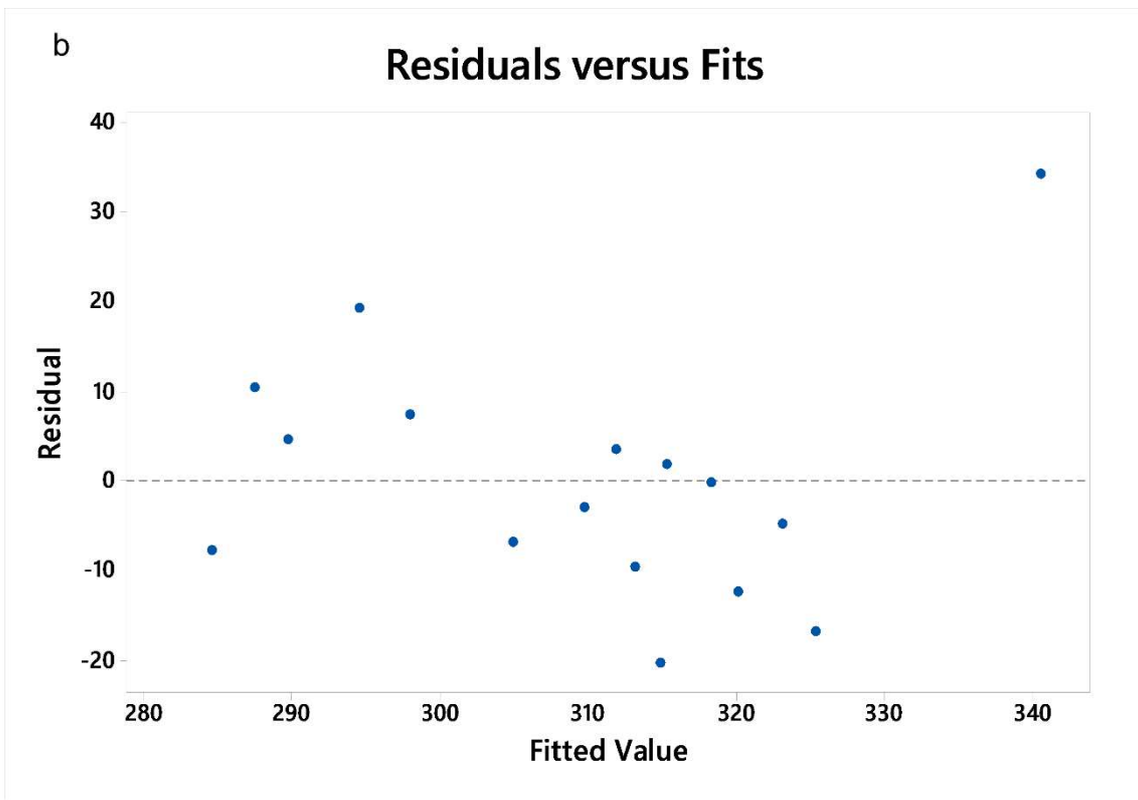
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466 **Figure 3**



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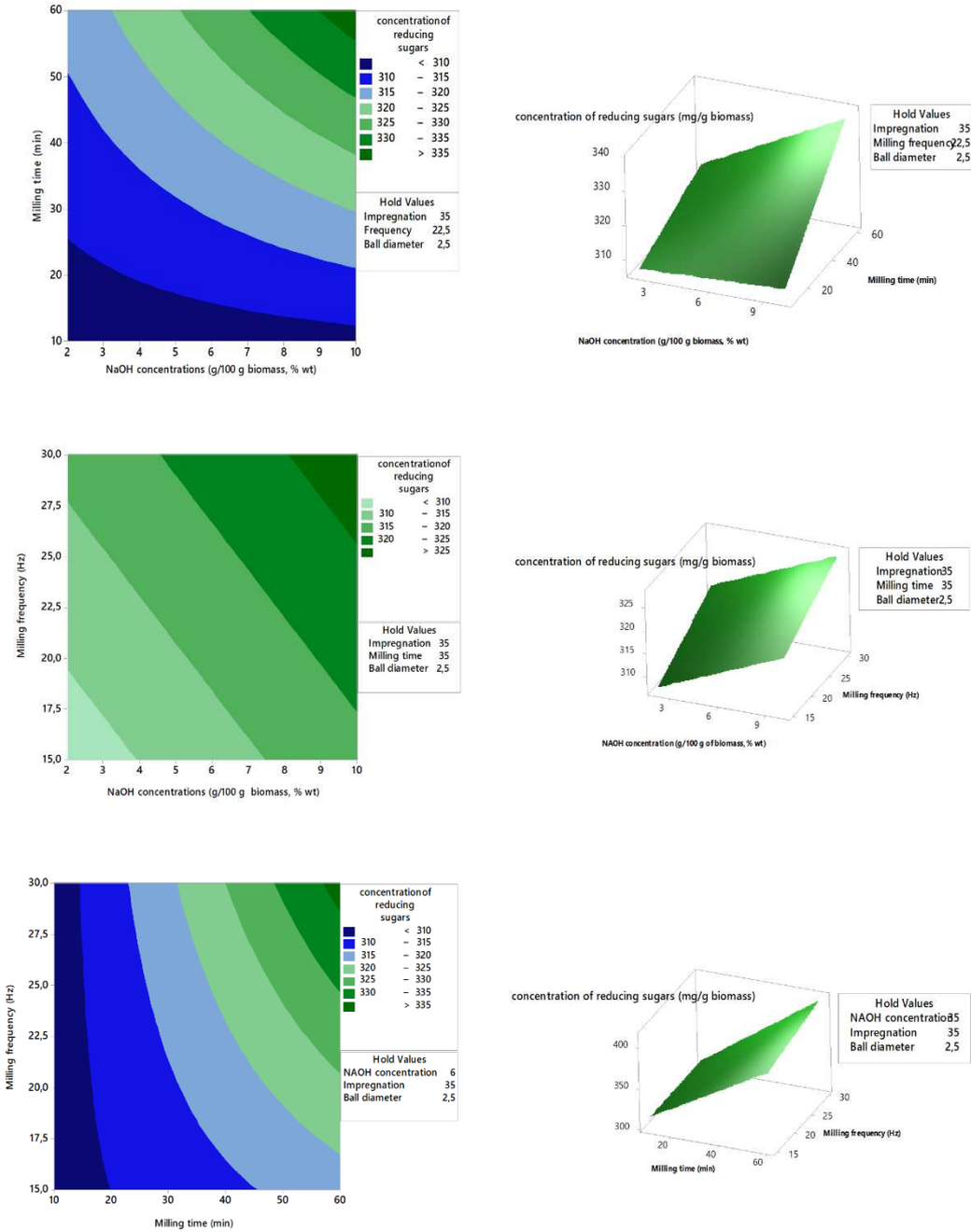


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471 **Figure 4**



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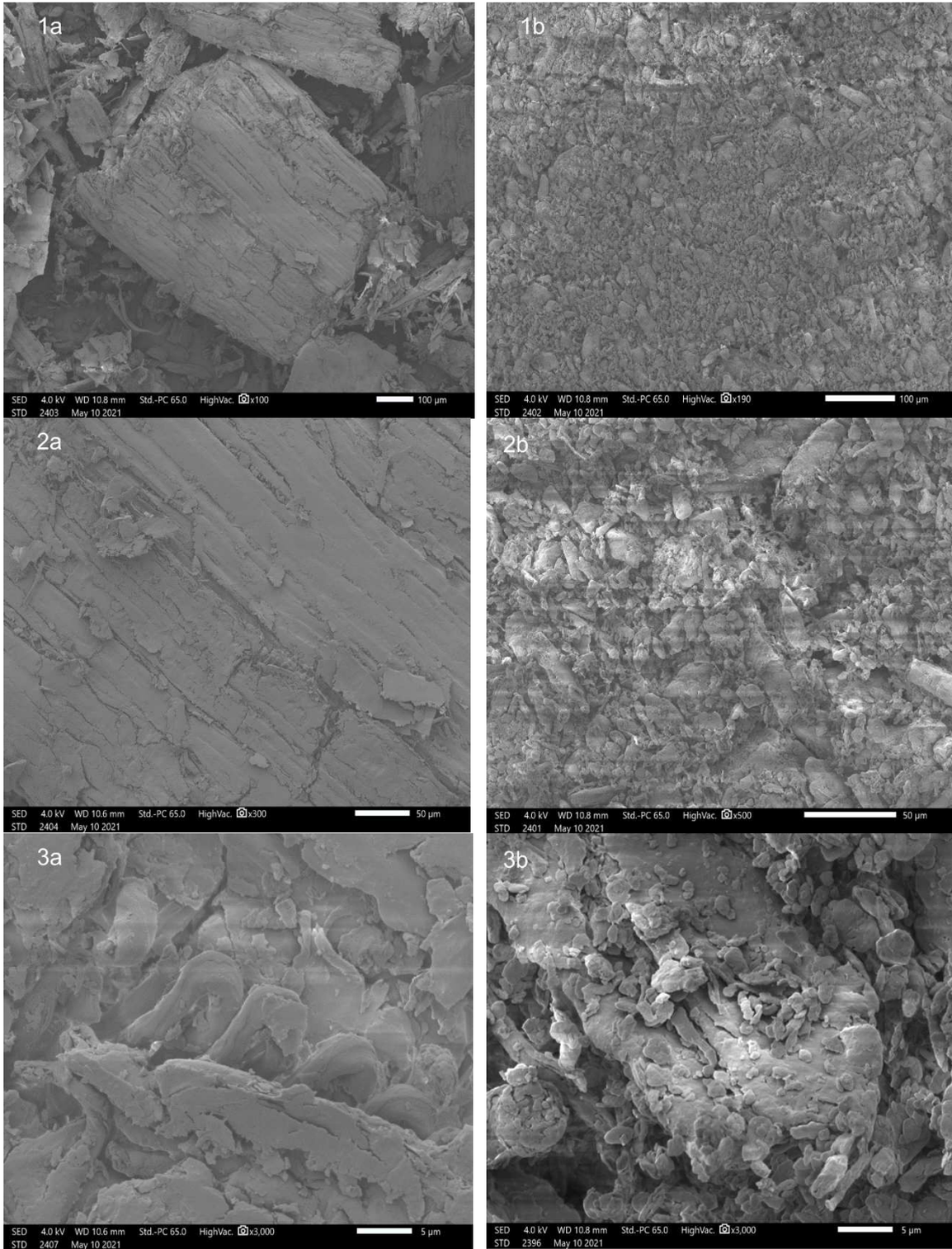
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477 **Figure 5**

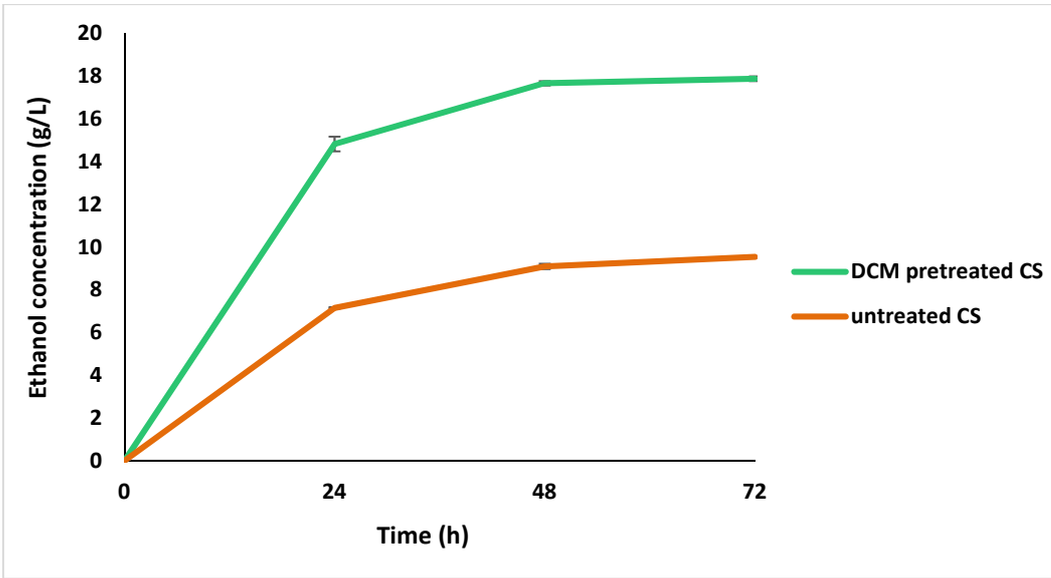
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480 **Figure 6**

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497 **Tables**

498 **Table 1.** Levels of the factors evaluated in the 2^{5-1} FFD

Factors	Low level (-)	High level (+)
A- NaOH concentration (g/100 g biomass, % wt)	2	10
B- Impregnation (water/biomass, % wt)	20	50
C- Milling time (min)	10	60
D- Milling frequency (Hz)	15	30
E- Ball diameter (cm)	1	2.5

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518 **Table 2.** Runs identification with the corresponding experimental conditions and the
 519 response of reducing sugars (mg/g biomass) released after 72 h of enzymatic hydrolysis.

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Runs	NaOH concentration (% wt)	Impregnation (% wt)	Milling time (min)	Milling frequency (Hz)	Ball diameter (cm)	Concentration of reducing sugars released (mg/g biomass)
1	10	20	10	15	1	294.19
2	2	20	60	30	2.5	318.39
3	2	20	10	15	2.5	313.88
4	10	50	10	15	2.5	315.29
5	2	20	10	30	1	298
6	10	50	60	30	2.5	374.70
7	10	50	60	15	1	317.25
8	10	20	10	30	2.5	294.54
9	10	50	10	30	1	284.78
10	2	50	60	15	2.5	307.80
11	10	20	60	30	1	318.15
12	2	50	10	30	2.5	306.70
13	2	50	60	30	1	303.52
14	2	20	60	15	1	305.37
15	2	50	10	15	1	276.50
16	10	20	60	15	2.5	308.62

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531 **Table 3.** ANOVA of the 2^{5-1} FFD describing the reducing sugars released after 72 h of
 532 enzymatic hydrolysis as a linear function of the selected variables.

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Source	Degrees of freedom	Sum of squares	Mean square	F-Value	P-Value
Model	11	6288.44	571.68	26.84	0.003
Linear	5	3553.49	710.70	33.37	0.002
A : NaOH concentration	1	509.51	509.51	23.92	0.008
B: Impregnation	1	149.60	149.60	7.02	0.057
C: milling time	1	1528.87	1528.87	71.78	0.001
D: Frequency	1	331.23	331.23	15.55	0.017
E: Ball diameter	1	1034.28	1034.28	48.56	0.002
2-Way Interactions	6	2734.95	455.83	21.40	0.005
A*B	1	1064.80	1064.80	49.99	0.002
A*C	1	370.59	370.59	17.40	0.014
B*C	1	199.96	199.96	9.39	0.038
B*D	1	215.79	215.79	10.13	0.033
B*E	1	497.17	497.17	23.34	0.008
C*D	1	386.64	386.64	18.15	0.013
Error	4	85.20	21.30		
Total	15	6373.64			

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**Raw chickpea straw
(CS)**

**Fractional factorial
design 2^{5-1}**

16 experiments

**Application of alkaline
vibro- ball milling
treatment**



**Pretreated chickpea
straw (CS)**

- ✓ **Reducing sugar yield of 374.70 mg/g biomass**
- ✓ **Reduction of cristallinity index to 23.50 %**

**Simultaneous
saccharification
and fermentation**

Bioethanol

17.81 g/L