Structural changes of corn starch during fuel ethanol production from corn flour

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Abstract

The key factor in production of fuel ethanol by simultaneous saccharification and fermentation is the efficient conversion of granular starch into ethanol. Most difficult stage in the process is the enzymatic hydrolysis of starch granules. Their supramolecular structure as well as crystallinity and presence of complexing agents are key factors for the hydrolysis process. The aim of the study was to examine structural changes in starch granules during the simultaneous processes of saccharification and fermentation of corn flour in the long-term repeated Simultaneous Saccharification and Fermentation (SSF) process with complete recycling of the stillage liquid fraction. The SSF experiments were performed using corn flour as a raw material, the STARGEN 001 preparation as a hydrolytic enzyme, and Red Star Ethanol Red (Saccharomyces cerevisiae) fermentation yeasts. Residual starch structure after the 4th, and the 7th operation cycle was examined using scanning electron microscopy, X-ray diffractometry, IR spectroscopy as well as gel permeation chromatography. In spite of accumulating glycerol, organic acids and inorganic ions in the fermentation broth, the repeated batch SSF process, with stillage recycling into the fermentation phase conducted on corn flour with the use of the STARGEN 001 enzyme preparation, was found to run efficiently. The amount of unhydrolyzed residual starch was independent of the number of operation cycles. Hydrolysis of starch resulted in the formation of porous granules and a small amount of undigested granules and pyramid-shaped residuals. Crystalline and amorphous regions were evenly digested. The molecular mass distribution of residual starch after the SSF process significantly differed from that of native starch both in the region corresponding to amylopectin and to amylose, while the most distinctive changes with respect to the amylopectin/amylose ratio, i.e. in the 4th cycle the amylopectin content decreased by up to 19%.

Key words: ethanol, simultaneous saccharification and fermentation, native corn starch

Introduction

Biological technologies of ethanol production have been known for a long time (Lim et al., 2003; Kosaric et al., 1996). They involve process of initial starch processing, i.e. starch gelatinization at high temperature followed by enzymatic hydrolysis. These two processes are characterized by high energy consumption, usually accounting for approx. 30 to 40% of entire energy required for ethanol production (Lim et al., 2003). There is a constant pursuit for energy-efficient technologies that would ensure high productivity and cost effectiveness. One of the possible ways to achieve this is using a direct one-step method for ethanol production by simultaneous hydrolysis and fermentation in one SSF (Simultaneous Saccharification and Fermentation) reactor and elimination of the gelatinization step using enzymes that can hydrolyze native starch (Kosaric, 1996; Kroumov et al., 2005). As the immediate utilization of hydrolysis products reduces enzyme inhibition, it is possible to obtain ethanol productivity that is 25-40% higher when compared with the traditional two-step method (Öhgren et al., 2007). Moreover, using native corn starch in the SSF process makes it possible to considerably reduce the costs of starch mash heating. Additionally, SSF decreases the costs and duration of the process, since it is no longer necessary to use two separate fermenters, owing to the elimination of preliminary fermentation stage which takes place already during hydrolysis (Kobayashi et al., 1998). Since hydrolysis usually requires higher temperatures than fermentation, finding optimum
The key factor in the above-described process is connected with efficient conversion of granular starch into conditions, i.e. ambient temperature and acidity suitable for both processes, poses a serious problem. For most strains of distiller’s yeast, the optimal growth temperature is 30-32°C, whereas in the SSF process, the temperature of the medium is 35-37°C, which is higher than the tolerance limit of distiller’s yeast (Mensonides et al., 2002; Laluce et al., 2002).

Starch can be hydrolyzed in its native, granular form and this process occurs efficiently in plants. However, commercial enzyme preparations commonly used in the starch-processing industry such as Thermamyl (Novozymes, Denmark), an outstandingly heat-stable α-amylase produced by genetically modified Bacillus strains, or BAN 480L (Novozymes, Denmark), a bacterial α-amylase produced by selected strains of Bacillus amyloliquefaciens, are effective only towards gelatinized starch (Sarikaya et al. 2000; Uhling, 1998). These enzymes were designed for processes in which starch is hydrolyzed at elevated temperatures in order to improve the reaction kinetics. It is known that gelatinization considerably enhances the reactivity of starch towards amylolytic enzymes. Hydrolysis of granular starch is significantly slower than that of dissolved starch, as the former process requires the step of unwinding of double helix in starch macromolecules. Thus, chains with restricted mobility, i.e. complexed or crystallized, are hydrolyzed with difficulty (Oates, 1997). The efficiency of enzymatic hydrolysis of granular starch depends on its origin, because different starches vary considerably in terms of their supramolecular structure (Oates, 1997; Dona et al., 2010). A comparison of the four cheapest commercial native starches in terms of their susceptibility to amylolysis allows us to set them in the following diminishing order: corn > wheat > cassava > potato (Sarikaya et al., 2000; Piyachomkwan et al., 2007).

The susceptibility to enzymatic hydrolysis also largely depends on the enzymes used (Sarikaya et al., 2000), which suggests a potential to genetically engineer better and more efficient enzyme(s) isoforms. A commercial enzymatic preparation exhibiting amylolytic activity to native starch, STARGEN 001, has been recently introduced to the market. STARGEN 001 preparation was developed by Genencor International (Palo Alto, CA) and it is recommended for simultaneous hydrolysis of native starch and ethanol fermentation. This enzyme preparation makes it possible to eliminate double pH regulation of the medium (liquefaction step – optimal pH 5.0-6.0, the saccharification step – optimal pH 4.8-5.0). Furthermore, when using STARGEN 001, there are no limitations related to mash viscosity, so mediums of high density can be used in the process. Employing STARGEN 001 ensures energy and water conservation and results in higher ethanol productivity by avoiding the loss of fermenting sugars, which may occur during heating (the Maillard reaction). Additionally, it does not require the use of an activator, such as calcium ions for most amylases.

There are also two important factors related to water management that determine economic effectiveness of bioethanol production. Firstly, the increasing global water shortage results in an elevation of its costs. Secondly, distillery stillage is ranked among especially burdensome industrial effluents, particularly in large bio refineries. One of the methods to reduce water costs incurred by distilleries is to reutilize stillage in the production process, which not only facilitates reasonable utilization of the by-product, but also significantly reduces the demand for industrial water (Larsson et al., 1997; Montesinos and Navarro, 2000). In order to improve the economical effectiveness of the SSF process, the zero-discharge fermentation system is applied. This means that the solid-containing whole stillage is separated using a decanter centrifuge. Thereafter, the solid phase is dried in a rotary dryer to produce DDGS (Dried Distillers Grain with Solubles), a valuable co-product used for livestock feed, whereas the liquid phase is evaporated in a double-effect evaporator (Bialas et al., 2010). Another way of improving this zero-discharge fermentation system is applying the repeated SSF process with complete recycling of the stillage liquid fraction. In such a system, the liquid phase is recycled into the SSF process (Bialas et al., 2010). This methodology meets the requirement of lowering the cost of process water. Scientific literature provides a range of suggestions for the reutilization of distillery stillage for technological purposes (Kim et al., 1997; Lu et al., 2003; Navarro et al., 2000). However, there are few described processes, where the whole stillage or its liquid fraction is repeatedly turned back and reutilized to prepare a new batch of medium. So far the possibility of returning stillage to the SSF process with native starch hydrolysis has not been investigated, therefore a research project on the subject has been designed.
ethanol. Most difficult stage is the enzymatic hydrolysis of starch granules, in which the key role is played by their supramolecular structure, crystallinity and presence of complexing agents. Aim of the present study was to examine structural changes in starch granules during the simultaneous processes of saccharification and fermentation with full stillage recycling.

Materials and methods

Microorganism
Freeze-dried distiller’s yeast, Red Star Ethanol Red (Saccharomyces cerevisiae), obtained from Lesaffre Company (Marcq en Baroeul, France) was used in this study for the production of ethanol from corn mashes. The number of the living cells for packing was > 2.0 × 10¹⁰/g, as specified by the manufacturer.

Starch material
Commercially available corn flour (BIO CORN, Ziebice, Poland) was used as a raw material for fermentation. It had a median diameter of 250 μm and contained 84% starch and 0.1% ash.

Enzymes
A mixture of granular starch hydrolyzing enzymes, containing Aspergillus kawachi α-amylase expressed in Trichoderma reesei, and glucoamylase from Aspergillus niger, was used in this study (STARGEN 001TM, Genecor International, Palo Alto, CA). The enzymatic activity of this set of enzymes was ≥ 456 GSHU/g (Granular Starch Hydrolyzing Units), as specified by Genencore International. In addition, fungal acid protease GC 106 (Aspergillus niger), also obtained from Genencore International, was added to mashes. The enzymatic activity of GC 106 was ≥ 1000 SAPU/g (Spectrophotometric Acid Protease Units), as specified by the manufacturer.

Simultaneous saccharification and fermentation
The SSF experiments were performed in a 5 l batch bioreactor (BIOFLO III, New Brunswick Scientific, New Brunswick, NJ) with a 4.0 l working volume under non-aerated conditions, at 35°C, 200 rpm. A slurry of raw corn flour in water (25% w/v) was prepared and saccharification was carried out by adding 2.05 ml (per kg corn flour dry matter) of Granular Starch Hydrolyzing (GSH) enzyme preparation (STARGEN 001). pH of the fermentation broth was measured at each sampling and adjusted to 5.0 by an addition of either 10% H₂SO₄ or 20% NaOH. In all cases, the medium was supplemented with acid protease GC 106 (40 μl/kg corn flour dry matter) and chloramphenicol (50 mg/l of the fermentation medium). Fermentation was started by the addition of freeze-dried distiller’s yeast Red Star Ethanol Red (0.5 g/l of fermentation medium). The conditions of SSF process were optimized using the multifactorial experimental approach (Bialas et al., 2009). Samples were obtained and analyzed for starch, glucose and ethanol concentrations after fermentation.

Recycling of stillage
After the fermentation period was completed, mash containing ethanol was pumped to a continuous distillation column (UOP3CC, Armfield, UK). At the top of this column, operating at 78.5°C and reflux ratio of 4:1, carbon dioxide was stripped from the ethanol solution which led to (an) ethanol concentration of 93-95% (volume based) in the side stream. The liquid fraction collected at the bottom of the distillation column was first cooled down to 30°C and then centrifuged at 4000 × g for 20 min. The supernatant was used instead of water to dilute native corn starch for the next SSF run. The recirculation of stillage was performed at a constant recirculation degree, with 75% of fresh water being replaced.

Analysis of starch structure
Samples from the 4th and 7th fermentation cycles were taken for analyses. The structure of the starch fraction in corn flour was also examined.

The morphology of starch granules was examined by scanning electron microscopy (SEM). Starch samples for analyses were isolated and prepared as follows: corn flour or the sediments after the 4th and the 7th fermentation cycles were suspended in a 20% NaOH solution and shaken for 10 minutes at room temperature. Then, the samples were centrifuged at 4000 × g for 15 minutes. Starch fractions were collected and purified by repeatedly suspending in water, shaking and centrifugation. Finally, the purified granules were dried and then deposited on a copper disc, coated with gold using a Jeol JEE 400 vacuum evaporator and analyzed under an SEM (Jeol JSM 5200) at 10 kV accelerating voltage.

Gel permeation chromatography (GPC) was carried out using a Waters Company apparatus (Alliance HPLC System 2695) equipped with a refractometric detector (RI) Waters 2414. Three UltraHydrogelTM columns arranged in a series were used. The obtained data were
Table 1. Time course of SSF processes with stillage recycling after hydrolysis in the 4th and the 7th cycle of the SSF process

<table>
<thead>
<tr>
<th>Fermentation time [h]</th>
<th>Operational cycle</th>
<th>Ethanol content [g/l]</th>
<th>Starch content [g/l]</th>
<th>Disaccharides content [g/l]</th>
<th>Glucose content [g/l]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>4th</td>
<td>0 ± 0.00*</td>
<td>210 ± 0.12</td>
<td>0.2 ± 0.03</td>
<td>15.1 ± 0.55</td>
</tr>
<tr>
<td>0</td>
<td>7th</td>
<td>0 ± 0.00</td>
<td>210 ± 0.17</td>
<td>0.5 ± 0.05</td>
<td>12.9 ± 0.22</td>
</tr>
<tr>
<td>72</td>
<td>4th</td>
<td>87 ± 0.54</td>
<td>43 ± 0.61</td>
<td>0.4 ± 0.03</td>
<td>2.1 ± 0.09</td>
</tr>
<tr>
<td>72</td>
<td>7th</td>
<td>91 ± 0.32</td>
<td>40 ± 0.52</td>
<td>0.1 ± 0.01</td>
<td>9.2 ± 0.08</td>
</tr>
</tbody>
</table>

* Three repetitions of measurement were conducted for each sample, the given standard deviation concerns measurement variability.

The X-ray diffractometry was performed with a TUR 62 Carl Zeiss X-ray diffractometer under the following conditions: X-ray tube CuKa (Ni filter), voltage 30 kV, current 15 mA, scanning from $Q = 2^\circ$ to $18^\circ$. To avoid the influence of relative humidity on relative crystal-
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Fig. 2. X-ray diffraction patterns of corn starch: native, hydrolyzed in the 4th and the 7th cycle of the SSF process

Fig. 3. IR spectra of corn starch: native, hydrolyzed in the 4th and the 7th cycle of the SSF process

linity, starch samples were placed in a desiccator and conditioned for 48 h in the atmosphere of 92% relative humidity.

The Fourier Transform Infrared Spectroscopy (FTIR) measurements were performed in the solid state with a FTIR Bruker IFS 113v spectrometer under the following conditions: KBr/pellet (200 mg/1.5 mg), resolution at 2 cm⁻¹.

Analytical methods

Samples for chemical analyses were first centrifuged at 4000 × g for 10 min. at 4°C (Multifuge 3SR, Germany), filtered through a 0.22 μm membrane filter (Millipore, USA) and then analyzed on an HPLC system (Merck Hitachi, Germany). Glucose, disaccharides and ethanol were separated on an Aminex HPX-87P apparatus (Bio-Rad, USA) at 30°C using 5 mM H₂SO₄ solution as the mobile phase at a flow rate of 0.6 ml/min, and then detected with a refractive index detector (Model L-7490, Merck Hitachi, Germany). The starch content was analyzed according to the enzymatic method developed by Holm et al. (1986). Block diagram of the experiment is demonstrated in Fig. 1.

Results and discussion

The repeated batch SSF process with stillage recycling was conducted on native starch, with the use of novel GSH enzymes, potentially being an alternative for the traditional technology of fuel ethanol production. Detailed results of the analysis of combined hydrolysis and fermentation process were presented by Bialas et al. (2009, 2010). Within this study, only 4th and 9th fermentation cycles were investigated (Table 1).

As a result of fermentation of native corn starch after 72-h fermentation, the concentration of ethanol was 87 g/l for the 4th and 91 g/l for the 7th cycle. Starch utilization rate in both cases was approximately 80%, while process efficiency accounted for 78% (4th) and 84% (9th) theoretical yield. The probable cause of incomplete substrate utilization in SSF process was decrease in the activity of amylolytic enzymes contained in STARGEN. It could have also been caused by mash acidification below optimal pH. Actions aimed at intensifying starch hydrolysis (like increasing enzyme addition, pH regulation, increasing stirring rate) would not be economically justified taking into account the high yield of fermentation process.

According to the data supplied by Genecor International Inc., theoretically, when using STARGEN 001 and corn mash of 33% s.s. density, final ethanol concentration of 16-18% may be obtained. However, these values have not been confirmed by any publications or descriptions of currently run production processes. Deviations of results recorded in this study from the values reported by the enzyme manufacturer may be caused by several reasons. A considerable role is played by the raw material – its origin, species and strain, which influences the proportions of individual starch fractions, and also by the milling rate of flour itself, which determines starch availability for enzymes. The SSF process using STAR-
Fig. 4. Scanning electron micrographs (SEM) of corn starch: native (A), hydrolyzed in the 4th (B) and the 7th (C) cycle of the SSF process

Table 2. Molecular weight distribution of native corn starch and corn starch hydrolyzed in the 4th and the 7th cycle of the SSF process

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Native starch</th>
<th>4th cycle</th>
<th>7th cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>M_w (g/mol)</td>
<td>area (%)</td>
<td>M_w (g/mol)</td>
</tr>
<tr>
<td>Amylopectin</td>
<td>1.53 × 10^6</td>
<td>64.2</td>
<td>1.49 × 10^6</td>
</tr>
<tr>
<td>Amylose</td>
<td>1.07 × 10^9</td>
<td>35.8</td>
<td>1.01 × 10^6</td>
</tr>
</tbody>
</table>

α molecular weights (g/mol) were calculated on the basis of GPC profiles presented in Fig. 4
b coefficient of variation did not exceed 0.3% for any calculated molecular weight

GEN 001 has also been described previously (Wang et al., 2005 and 2006; Sharma et al., 2007).

There are a number of structural factors that determine starch resistance to enzymatic attack. Its crystal structure is considered to be one of the most important factors in this regard (Planchot et al., 1997). A-type starches, such as those from normal-genotype cereals, are much more susceptible to hydrolysis by α-amylase than B-type starches, such as high-amylase cereals or potato starches. Moreover, it was postulated that the hydrolysis rate of starch granules largely depends on the distribution of semi-crystalline and crystalline layers (Gallant et al., 1997). This hypothesis was confirmed by SEM analysis of residual starch granules after partial α-amylase hydrolysis and was the basis of “blocklet” concept of the supramolecular organization of starch (Gallant et al., 1997). Following this hypothesis one could expect that residual starch would exhibit a higher degree of crystallinity than native starch. As it is shown in Fig. 2, residual starch after the SSF process showed no changes in crystallinity in comparison with native starch both in terms of the type of X-ray diffraction pattern and relative crystallinity.

This observation was similar to that made by Zhang et al. (2006) that both crystalline and amorphous regions in native cereal starches are evenly digested by the mixture of α-amylase and amyloglucosidase. The above-mentioned discrepancies can be related to different mechanisms of enzymatic hydrolysis catalyzed by enzymes of different biological origin (Robertson et al., 2006). The supramolecular organization of starch may also be examined using IR spectroscopy. According to the data reported by van Soest et al. (1995), the IR spectrum is sensitive to changes in the short-range structure in the C-C and C-O stretching regions at 1300-800 cm⁻¹. The absorbance band at 1047 cm⁻¹ is sensitive to the amount of ordered or crystalline starch, while the band at 1022 cm⁻¹ is characteristic of amorphous starch (van Soest et al., 1995). However, the FTIR technique does not differ between the A- and B-type of crystallinity, and the differences observed between different starches are not related to this factor (Sevenou et al., 2002; Lewandowicz and Soral-Smietana, 2004). The IR examination demonstrated no changes caused by the SSF process either in the functional groups or in the finger-print range (Fig. 3), and even the 1047 cm⁻¹
and 1022 cm\(^{-1}\) bands did not change as a result of the hydrolysis process.

According to Planchot et al. (1995), in the first stage of hydrolysis by \(\alpha\)-amylases normal corn starch granules are randomly furrowed, then the pores are penetrated through the outer shells of granules, and finally the enlarged, individual pores form external grooves and internal corrosion channels. However, it was also reported that native corn starch granules have small pores (more precisely – radial, tube-like channels that connect the central cavity to the external environment) (Sarikaya et al., 2010; Huber and Miller, 1997; Huber and Miller, 2000). Zhang et al. (2006) demonstrated that hydrolysis of native corn starch starts with an increase in pore size until an almost complete fragmentation of the granules. However, in the last stage of hydrolysis, a few undigested small granules exist alongside pyramid-shaped residuals (Gallant et al., 1997). In the SEM images of starch remaining after individual cycles of the SSF process (Fig. 4), starch granules with pores were predominant. Undigested starch granules and pyramid-shaped residuals were less prevalent. The above observation suggests that there is a margin in the SSF process, and the degree of hydrolysis can be enhanced.

The experimental data presented above showed that both amorphous and crystalline fractions were uniformly digested by the STARGEN 001 amyloytic enzyme preparation. Since crystalline regions are formed only by amylopectin (Robertson et al., 2006), both the amylose and the amylopectin fractions should be simultaneously hydrolyzed. This hypothesis was, to some extent, confirmed by the GPC analysis (Fig. 5).

This method is conventionally applied for the determination of molecular weight distribution in polymers. However, in the case of starch, the determination of the actual values of number \((M_n)\) or weight \((M_w)\) average molar masses is especially difficult due to the extremely high molecular mass of amylopectin as well as problems with dissolving starch. As a consequence, the data regarding molecular mass of different native starches reported by different researchers vary considerably (Swinkles, 1985; Aberle et al., 1994; Han and Lim, 2004).

The main problem in the determination of the molecular weight distribution in starches is associated with dissolving the entire starch sample without cleaving any of the macromolecules. The most frequently recommended starch solvents are aqueous solutions of sodium hydroxide or aqueous di-methyl sulfoxide. However, both the solvent and the solubilization method strongly affect the occurrence of a polysaccharide breakdown. Moreover, the most popular pullulan and dextran molecular weight standards are characterized by slightly different molecular structures and far lower molecular weights when compared to starch, so they can be recommended only for the assessment of the molecular weight of amylose (Lin et al., 2003; Chen et al., 2010; Braun 1993; Jackson, 1991). However, to answer the question regarding the susceptibility of the amylose and amylopectin fractions to amyolysis, knowledge of the absolute values of molecular weight is not essential, in contrast with information on the changes in both molecular weight and percentage of individual fractions. As it is shown in Figure 5 and Table 2, the molecular mass distribution of residual starch after the SSF process actually differed from that of native starch both in the region corresponding to amylopectin and amylose. These changes were mainly in regard to the amylopectin/amylose ratio. As a result of the SSF process, proportion of the low-molecular fraction (amylose) increased. This phenomenon strongly confirmed the statement of the Zhang group on an even digestion of the amylose and amylopectin fractions during amyolysis of starch granules (Zhang, 2006).

**Conclusions**

In conclusion, it needs to be stated that the enzymatic preparation produced by Genencor International...
using genetically modified microorganisms provides effective hydrolysis of native corn starch, owing to which efficiency of ethanol production may be increased with a simultaneous cost reduction as a result of the elimination of the energy-intensive starch liquefaction process run at high temperatures. The amount of unhydrolyzed residual starch is independent of the number of operation cycles. Hydrolysis of starch occurs with the predominant formation of granules with pores as well as an additional small amount of undigested granules and pyramidal-shaped residuals. Crystalline and amorphous regions are evenly digested. This observation is also confirmed by the determination of changes in molecular mass distribution caused by the SSF process.

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References


