Bioproduction of Ethanol in Separate Hydrolysis and Fermentation and Simultaneous Saccharification and Fermentation from Cassava Stalks

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Abstract: Cellulose biomass is being investigated as a potential substrate for bioethanol production. Cassava stalks were successfully converted to ethanol by fermentation using Saccharomyces cerevisiae TISTR5048, S. cerevisiae KM1195, S. cerevisiae KM7253 and co-culture of S. cerevisiae TISTR5048 and Candida tropicalis TISTR5045. The objective of this study was to assess the ethanol production from cassava stalks by dilute-acid pretreatment and enzymatic hydrolysis that were convertible into ethanol by mono-culture and co-culture of yeast strain. Cassava stalks 1.5% (w/v) in 0.1 M sulfuric acid was pretreated for 30 min at 135 °C under the pressure of 15 lb/inch². The pretreated cassava stalk suspensions were neutralized to pH 5.5 for saccharification process. The enzyme solution (α-amylase, amyloglucosidase, cellulase, xylanase and pectinase solubilized in buffer pH 5.0) was used for hydrolysis of pretreated cassava stalk at 50 °C for 24 h. The hydrolyaste was supplemented with additional nutrients. The culture was incubated at 30 °C. The pretreatment of the stalk with dilute-acid resulted sugar yield of 0.57 g/g dry matter from enzymatic hydrolysis, which was higher than dilute-alkaline-pretreated and distilled water-pretreated stalk. The sugar hydrolysate was bioconverted to ethanol with separate hydrolysis and fermentation (SHF) and simultaneous saccharification and fermentation (SSF). The highest ethanol yields of 98.43% and 95.29% were obtained in SHF and SSF, respectively by S. cerevisiae KM1195. The fermentation time of SSF process was 24-32 h shorter than that of the SHF (∼ 56 h), but not significantly leading to difference in ethanol production (5.42 g/L-6.22 g/L for SSF; 5.9 g/L-6.23 g/L for SHF).

Key words: Ethanol, cassava stalk, fermentation.

1. Introduction

Cassava (Manihot esculenta Cranz) is considered an important source of food and dietary calories for a large population in tropical countries in Asia, Africa and Latin America. Cassava ranks as the world’s sixth most important food crop and is the basic food for more than 700 million people in several countries [1-2]. It has the remarkable capacity to adapt to various agro-ecological conditions. It is also considered as a low-risk crop. In view of its drought-resistant nature and non-requirement of any specific growth conditions, much attention has been paid in the past 15-20 years to its agricultural aspects, for increasing its production all over the world, which has been well achieved. Cassava is a bushy plant producing tubers and is made up of an aerial part or stalk and an underground part. The stalk can be as high as 2-4 m with a trunk and branches on it. The underground part consists of two types of roots: the ones responsible for the plant nutrition, and the others with axial disposition surrounding the trunk. In recent years, there has been an increasing trend towards more efficient utilization of agro-industrial residues such as cassava bagasse, sugar cane bagasse, sugar beet pulp, coffee pulp/husk, apple pomace, etc. Several processes have been developed that utilize these as raw materials for the production of bulk chemicals and
value-added fine products, such as ethanol, single cell protein (SCP), mushrooms, enzymes, organic acids, amino acids, biologically active secondary metabolites, etc. [3-5]. Application of agro-industrial residues in bioprocesses on the one hand provides alternative substrates, and on the other hand helps in solving pollution problems, which their disposal may otherwise cause. In addition, a reduction of the cost of the fine products, such as ethanol, can be achieved by reducing the cost of the raw materials. With the advent of biotechnological innovations, mainly in the area of enzyme and fermentation technology, many new avenues have opened for their utilization.

Bioethanol, a renewable fuel is becoming increasingly important as a consequence of major concern for depleting oil reserves, rising crude oil prices and greenhouse effect [6]. Agro-industrial residue feedstock is considered as an attractive raw material not only for the liquid transportation fuel but also for the production of chemicals and materials, i.e. the development of carbohydrate-based biorefineries [7] because of its availability in large quantities at low cost [8]. Over the past few years, ever since the energy crunch began, there has been a tremendous interest in energy saving both on new and existing structures. Using certain materials and techniques can result in big savings. Today, the idea of utilizing biomass from agricultural and livestock wastes as a raw material for production of ethanol has attracted the interest of researchers especially in agricultural practicing countries. Thailand has an abundance of agriculture by-products available which are usually directly discharged as solid waste; causing environmental issues.

Thailand is an agricultural country. Each year the country produced not only agricultural product but also more than 50 million tons of agricultural residues. Cassava stalk is the fourth largest agricultural residues which accounted more than 4 million tons per year [9]. Cassava stalk was considered as useless agricultural residues. To fully utilize the cassava stem as a feedstock for ethanol production, pretreatment is required to render the cellulose fibers more amenable to the action of the hydrolytic enzymes.

In order to obtain fermentable sugars from the biomass, the biomass needs to be hydrolyzed. In an enzymatic process this calls for a pretreatment step, during which the biomass is made more accessible to enzymatic hydrolysis. The obtained monosaccharides from the enzymatic hydrolysis step, can be fermented to ethanol using microorganism. The fermentation can be made either together with the enzymatic degradation, known as simultaneous saccharification and fermentation (SSF), or as two separate steps, known as SHF (separate hydrolysis and fermentation). The advantage of the former process is a reduction in investment costs [10] and a release of end-product inhibition in the enzymatic hydrolysis. On the other hand, the SHF process allows the enzymatic hydrolysis to be carried out at a higher temperature than the fermentation, which is typically an advantage because of the higher temperature optimum for the enzymatic hydrolysis in comparison to the fermentation [11].

In this study, we intend to the biotechnological potential of cassava stalk for value addition of bioethanol product. The study is aimed to investigate ethanol production from cassava stalk acid and enzyme hydrolysate using mono-culture and co-culture of yeasts fermentation in SHF and SSF.

2. Materials and Methods

2.1 Substrate Preparation

Cassava stalks were collected and were washed manually using tap water to remove adhering dirt, dried at 45 °C in a hot-air oven for 4 days, milled, screened to select the fraction of particles with a size of 45-697 μm, homogenized in a single lot and stored until needed.

2.2 Hydrolysate Preparation

Hydrolysate was prepared by autoclaving under 15 lb/inch², the 1.5 g dried powder of cassava stalks with 100 mL of 0.1 M sulfuric acid, in conical flasks. Then, 250 mL filter-sterilize cellulase (Sumitime C; Shin
Nihon Chemical Co. Ltd., Japan) solution (cellulase activity: 20 Filter paper units (FPU) (g substrate)$^{-1}$, α-amylase 100 unit (g substrate)$^{-1}$, amylglucosidase 100 unit (g substrate)$^{-1}$, xylanase activity: 500 unit ((g substrate)$^{-1}$) and pectinase activity: 250 unit ((g substrate)$^{-1}$) in 0.1 M sodium phosphate (pH 5.0) was added to the flask and reacted at 50 °C and 120 rpm for 48 hours for hydrolysis. After the enzymatic reaction, the hydrolysate was centrifuged at 21,000 × g for 10 min. The supernatant was supplemented with additional nutrients to give a base medium composition of: 1 g/L yeast extract; 2 g/L (NH$_4$)$_2$SO$_4$; 1 g/L MgSO$_4$$\cdot$7H$_2$O.

2.3 Cassava Stalk Hydrolysate Medium

Fermentation medium composed of (g/L): yeast extract 1; (NH$_4$)$_2$SO$_4$ 2; MgSO$_4$$\cdot$7H$_2$O 1.

2.4 Batch Fermentation

Batch fermentation was conducted in a 250 mL conical flask with a working volume of 100 mL. The fermentation medium was inoculated with 5% v/v inoculum (20 hours culture, $1 \times 10^7$ cells/mL). The fermentation temperature was kept constant at 30 ± 0.2 °C in an incubation shaker. The broth was kept under agitation at 50 rpm. Samples were taken at regular time intervals during fermentations to determine the concentrations of cell mass, ethanol and residual sugars in the broth. All experiments were carried out in duplicate.

2.5 Analytical Methods

Total solids (TSs) moisture and crude protein in cassava stalk were determined according to standards [12]. Cellulose, hemicellulose and lignin contents were determined by the detergent extraction method [13].

2.6 Biomass Estimation

Culture dry weight was measured by centrifugation and drying at 105 °C, until no weight change between consecutive measurements was observed.

2.7 Sugar Estimation

Total reducing sugar was estimated by using dinitrosalicylic acid (DNS) reagent [14].

2.8 Ethanol Determination

The fermentation was carried out at 30 °C for 18 h. The fermentation broths were filtered through a 0.45 m Millipore filter. Ethanol in the samples was determined by gas chromatograph using a 60:80 Carbopack B: 5% Carbowax 20 M glass column. The injector was operated at 200 °C. The flame ionization detector (FID) was kept at 200 °C. Nitrogen gas was used as carrier gas at a flow rate of 30 mL/min. The temperature was programmed at 120 °C for 1.4 min, from 120 °C to 240 °C at 30 °C/min, then held 5 min at 240 °C.

3. Results and Discussion

3.1 Composition of Cassava Stalk

Production of ethanol through fermentation process from biomass is dependent on its quality. The average composition of cassava stalk is summarized in Table 1. Primarily, the major constituents of cassava stalks are relatively high carbohydrates (cellulose, hemicellulose and starch). The result indicated that cassava stalks could be a good source for bioconversion.

3.2 Cassava Stalk Acid and Enzyme Hydrolysate Preparation

Dilute sulfuric acid hydrolysis (0.1 M) under autoclaving under pressure of 15 lb/inch$^2$ at 135 °C for 10 min and enzyme hydrolysis as described in materials and methods was very effective in releasing a good amount of sugar from cassava stalks (Table 2). Higher temperature, higher yield of glucose and reducing sugars were released. Approximately 14% and 21% of glucose and reducing sugars were released at 120 °C less than that of at 135 °C, respectively. So, temperature at 135 °C was suitable to hydrolyse the cassava stalks for sugar production. Approximately 52.73% and 38.07% of the reducing sugars and glucose,
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Table 1  Average composition of cassava stalk.

| Constituents | % of wet weight |
|--------------|----------------|
| TSs          | 56.24-59.47    |
| Moisture     | 25.47-28.69    |
| Organic components (% TSs) | |
| Hemicellulose | 11.62 ± 0.24   |
| Cellulose    | 21.43 ± 0.17   |
| Lignin       | 22.64 ± 0.37   |
| Crude protein| 2.72 ± 0.29    |
| Starch       | 8.41 ± 0.32    |

Table 2  Effect of temperature on average sugar composition of cassava stalk acid and enzyme hydrolysate.

| Temperature (°C) | Glucose (g/L) | Reducing sugars (g/L) |
|------------------|---------------|-----------------------|
| 120              | 5.02 ± 0.16   | 6.78 ± 0.11           |
| 125              | 5.29 ± 0.12   | 7.04 ± 0.14           |
| 135              | 5.71 ± 0.07   | 7.91 ± 0.05           |

respectively, were released in the first 10 min of autoclaving and enzyme hydrolysis, at 30 min of autoclaving and enzyme hydrolysis, 74.12% and 54.27% of the reducing sugars and glucose, respectively, were released after which increase was observed (Table 3). After 60 min of autoclaving, sugar yield were 84.47% and 62.40% of the reducing sugars and glucose, respectively. These sugars were derived primarily from starch and cellulose component. The sugars yield (84.47%) was rather high, showing that starch and cellulose almost practically hydrolyzed.

3.3 Ethanol Production in SHF

The highest values of ethanol yield per unit biomass (Cₑ), the maximum ethanol production (Pₘₐₓ), ethanol production rate (Qₑ) and product (ethanol) yield coefficient (Yₑₚₛ) were found to be 0.415 g (g-biomass)⁻¹, 6.23 g/L, 0.593 g/L/hour and 0.477 g (g-total sugar)⁻¹, respectively, by the fermentation of S. cerevisiae KM1195. The lowest values of ethanol yield per unit biomass (Cₑ), the maximum ethanol production (Pₘₐₓ), ethanol production rate (Qₑ) and product (ethanol) yield coefficient (Yₑₚₛ) were found to be 0.393 g (g-biomass)⁻¹, 5.90 g/L, 0.911 g/L/hour and 0.422 g (g-total sugar)⁻¹, respectively, by the fermentation of mono-culture of S. cerevisiae TISTR5048. It was found that mono-culture of S. cerevisiae KM1195 could produce relatively higher ethanol yield than the co-culture (Table 4).

3.4 Ethanol Production in SSF

The highest values of ethanol yield per unit biomass (Cₑ), the maximum ethanol production (Pₘₐₓ), ethanol production rate (Qₑ) and product (ethanol) yield coefficient (Yₑₚₛ) were found to be 0.414 g (g-biomass)⁻¹, 6.22 g/L, 0.969 g/L/hour and 0.486 g (g-total sugar)⁻¹, respectively, by the fermentation of S. cerevisiae KM1195. The lowest values of ethanol yield per unit biomass (Cₑ), the maximum ethanol production (Pₘₐₓ), ethanol production rate (Qₑ) and product (ethanol) yield coefficient (Yₑₚₛ) were found to be 0.360 g (g-biomass)⁻¹, 5.42 g/L, 0.911 g/L/hour and 0.422 g (g-total sugar)⁻¹, respectively, by the fermentation of mono-culture of S. cerevisiae TISTR5048. It was found that mono-culture of S. cerevisiae KM1195 could produce relatively higher ethanol yield than the co-culture (Table 5).

The enzymatic response was evaluated as a function of the temperature and time of pretreatment. The experiments were carried out in order to find the optimal conditions. The temperature ranged from 120°C to 135 °C, and the acid concentration was 0.1 M in the optimal test.
Table 5  Ethanol production by SSF with mono-culture and co-culture.

| Strain                  | C_E | P_{max} | Q_E | Y_{p/s} |
|-------------------------|-----|---------|-----|---------|
| SSF with single-culture |     |         |     |         |
| S. cerevisiae TISTR5048 | 0.360 | 5.42   | 0.911 | 0.422   |
| S. cerevisiae KM1195    | 0.414 | 6.22   | 0.969 | 0.486   |
| S. cerevisiae KM7253    | 0.371 | 5.57   | 0.327 | 0.435   |
| SSF with co-culture inoculation of S. cerevisiae TISTR 5048 with C. tropicalis TISTR 5045 | 0.394 | 5.89   | 0.664 | 0.461   |

C_E: Ethanol yield per unit biomass (g (g-biomass)^{-1}),
Q_E: Ethanol production rate (g/L/hour),
P_{max}: Maximum ethanol production (g/L),
Y_{p/s}: Product (ethanol) yield coefficient (g (g-total sugar)^{-1}).

Fig. 1 and Fig. 2 show the time-course for growth, sugar utilization and ethanol concentration in the hydrolysate medium at initial pH 5.0 ± 0.2 of mono-culture and co-culture of S. cerevisiae and C. tropicalis. The fermentation parameters are summarized in Tables 4 and 5. Carbohydrate contents of cassava stalks (Table 1) are directly proportional to the yield of ethanol. The yield (C_E) and productivities (P_{max}, Q_E and Y_{p/s}) in SHF and in SSF of mono-culture of S. cerevisiae KM1195 relatively high ethanol yield almost equal to the co-culture of S. cerevisiae and C. tropicalis when grown in the hydrolysate medium. This showed that co-culture of S. cerevisiae and C. tropicalis fermentation employed for the treatment of the cassava stalk acid enzyme hydrolysate has partially used reducing sugars as substrate but not affected to improve the fermentability. The ethanol production rate (g/L/hour) of co-culture comparing to mono-culture of S. cerevisiae KM1195 was higher about 24% in SHF while ethanol production rate (g/L/hour) of co-culture reduced about 32% in SSF.

However, the ethanol yield of co-culture for the cassava stalk acid enzyme hydrolysate was rather similar to that obtained with S. cerevisiae KM1195.

Fig. 1  The time course of growth (x), reducing sugars (●), glucose (▲) and ethanol (●) concentration in SHF by S. cerevisiae TISTR5048 (a), S. cerevisiae KM1195 (b), S. cerevisiae KM7253 (c) and co-culture of S. cerevisiae TISTR5048 and C. tropicalis TISTR5045 (d) at 30 ± 0.2 °C and pH 5.0 ± 0.2 using simulated synthetic hydrolysate medium.
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For producing high quality ethanol, pretreatment of biomass is essential and this leads to better enzymatic hydrolysis fractionate, solubilize, hydrolysis components [15]. Several treatment technologies include concentrated acid [16], dilute acid, alkaline, steam explosion, wet oxidation and liquid hot water. Among these methods, acid hydrolysis is frequently used as a pretreatment because it can be tailored to a wide variety of feedstocks. This method not only exposes starch, cellulose for enzymatic saccharification but also solubilizes hemicellulose and converts it into a fermentable sugar, xylose [17]. However, rapid and efficient fermentation of fermentable sugars is limited because of toxic compounds such as furfural and HMF which are generated during high temperature pretreatment [18] and ultimately inhibits microbial growth. The rationale of dilute sulfuric acid pretreatment of cassava stalk in this study is to obtain a higher yield of monomeric fermentable sugars with an aim to minimize inhibitor generation.

The results showed that SSF with S. cerevisiae KM1195 performed similar to SHF. The temperature optima for the yeast and the enzymes used differ, which means that the conditions used in SSF cannot be optimal for both the enzymes and the yeast. However, advantage of SSF is that when combining the two process (saccharification and fermentation) steps, it results in a lower capital cost and reduces the risk of contamination [19]. However mixing the lignin residue
with yeast makes yeast recirculation very difficult [20].

Compared to the two-stage hydrolysis–fermentation process, SSF has the following advantages: increase of hydrolysis rate by conversion of sugars that inhibit the cellulase activity; lower enzyme requirement; higher product yields; lower requirements for sterile conditions since glucose is removed immediately and ethanol is produced; shorter process time; and less reactor volume because a single reactor is used.

However, ethanol may also exhibit inhibition to the enzyme activity in the SSF process. Wu and Lee [21] found that cellulase lost 9%, 36% and 64% of its original activity at ethanol concentrations of 9, 35 and 60 g/L, respectively, at 38 °C during SSF process. The disadvantages which are considered for SSF include: incompatible temperature of hydrolysis and fermentation; ethanol tolerance of microbes; and inhibition of enzymes by ethanol. Bioethanol production has been improved by new technologies, there are still challenges that need further investigations. These challenges include maintaining a performance of the optimal yeasts in commercial scale fermentation operations, developing more efficient pretreatment technologies for lignocellulosic biomass, and integrating the optimal components into economic ethanol production systems.

4. Conclusion

It can be concluded that bioconversion of cassava stalk could be economically useful for the production of ethanol. The maximum values of ethanol yield (C_e), productivity (P_{max}, Q_E and Y_{p/s}) and percent sugar utilization were obtained, when co-culture of S. cerevisiae TISTR5048 and C. tropicalis TISTR5045 or mono-culture of S. cerevisiae KM1195 was grown in treated cassava stalk acid enzyme hydrolysate medium both in SHF and SSF at temperature 30 ± 0.2 °C and pH 5.0 ± 0.2. However, SSF may hold promise and focus should be made on developing SSF technologies. Development of efficient microbial strains suitable for bioconversion of cassava stalk is still a largely unexplored area. Efforts should be also made for improving cassava stalk hydrolysis conditions; its effective conversion into fermentable sugars is an area which needs further inputs in terms of research and development. Cassava stalk could serve as a good substrate for production of value-added bioethanol product and the fermentation of cassava stalk for ethanol production was carried out in a high yield by optimum treatment and culture of yeast strains.

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