Study of Biogas Production from Cassava Industrial Waste by Anaerobic Process

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Abstract. Biogas production processes from tapioca wastewater have several problems that cause the biogas production is not optimal, such as pH drop at beginning of the process because the rate of acid formation is too fast and the rate of starch wastewater degradation is too slow. Therefore, to obtain optimal biogas production it is required two-stage reactor. The purposes of this research were to (i) study the influence of one stage fermentation and two stage fermentation on biogas production, (ii) study the effect of buffer Na2CO3 on biogas production, and (iii) study the effect of methanogenic bacteria concentration on biogas production from cassava starch effluent. The first method of our research was hydrolysis process by “Saccharomyces cerevisiae” as substrate activator. The second is the arrangement of pH and the last is process of methane production. The results showed that the highest biogas production is achieved at concentration of methanogenic bacteria 20% (v/v) that is equal to 2458 ml. At concentration of 8% (v/v) and 15% (v/v), biogas production was 2105 ml and 2117 ml. The addition of Na2CO3 can extend to 16 days with accumulation of 372 ml. While without the addition of buffer, biogas production period was only 9 days with accumulation of 620 ml. In semi continuous process, the analysis carried out every 3 days. Highest biogas production achieved in the variable addition of yeast with the accumulation 9329 ml. Without yeast, accumulation of biogas was 6831 ml. Yeast is use as substrate activator so it can accelerate the hydrolysis process and increased biogas production. The addition of Na2CO3 is increase the alkalinity so the pH drop did not occur early in the process.

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1 Introduction

In the processing of cassava into wet tapioca starch, the tapioca miss rate that comes with wastewater during deposition slurry process slurry is large enough. It’s about 1% of total cassava tapioca [1] which directly lost upon environment. Large contain of tapioca in these wastes can be used as raw material for biogas making.

The process of biogas production from tapioca wastes have several problems that led the biogas production not optimal, among others, the pH drop at beginning process because the rate of acid formation is too fast that causing death of microbial methanogens. Besides that, slow degradation rate of tapioca waste caused by the solids content of tapioca waste is still a polysaccharide molecule. Therefore, to obtain optimal biogas production need two stages reactor. The first stage is hydrolysis to produce simpler saccharide compounds (disaccharide and monosaccharide) by Saccharomyces cereviceae. The second stage is the formation of methane from the output of first reactor by methanogens bacterial. Meanwhile, pH drop in early process can be prevented by Na2CO3 which acts as buffer to maintain the pH. With the addition of Na2CO3 and the hydrolysis of polysaccharides is expected to produce a lot of biogas in a short time. Similar with tapioca, many researchers have conducted research on biogas from waste such as agricultural waste is able to produce a lot of gas through an anaerobic process [2-12].

In general, the purpose of the study was to evaluate biogas production with two-stage fermentation of waste water from tapioca industrial with substrate activator. Specifically, the purpose of the research was to study the effect of anaerobic microbial concentration on the rate of biogas formation, study the effect of Na2CO3 buffer addition on the rate of formation of biogas, study the effect of fermentation 1 stage and 2 stage in rate of biogas production from starch waste.

2 Materials and Methods

Materials used in the experiment is synthetic wastewater of tapioca manufacture that made by mixing 20 grams of tapioca in 2 litters of water. Yeast (Saccharomyces cereviceae) NKL brand. Water is obtained from Chemical Process Laboratory, Chemical Engineering, Diponegoro University. Urea was obtained from Chemical Process Laboratory, Chemical Engineering, Diponegoro University, and Buffer Na2CO3 non-PA obtained from chemical shops in Semarang.

The tools used in the experiments is a two-stage anaerobic biogastage which consists of waste and nutrient storage tank, mixing tank, airtight stirred tank, airtight stirred tank and pressure resistant, pH meters, water pump, valve, pipe, pressure gauge, hose with diameter 1 cm, 250 ml measuring cup, water tank, and electric motors. Overall design of equipment used is presented in Figure 1.

The fixed variables tested to several respond variables. These variables include: fermentation 1 stage and 2 stage in continuous conditions with Na2CO3 buffer, rumen concentration is 10% (v/v), and yeast. In two stage fermentation, when yeast variables of 0.08 w/v, the rumen concentrations is 8%, 15%, and 20% v/v. When yeast variables of 0.15% w/v, the addition of Na2CO3 buffer is batch with variables the presence or absence of and rumen concentration of 15% (v/v) and yeast 0.15% (w/v) in batch conditions.

The process begins by assembling the equipment like in Figure 1. Then mix the tapioca wastes 1% (w/v) with 0.4% urea (w/v). In the second stage of fermentation the mixture is then poured into hydrolysis tank and then add the Saccharomyces cerevisiae 0.08% (w/v), and let stand for 1 day, so polysaccharides will be degraded into disaccharides and monosaccharides. Furthermore disaccharide and monosaccharide produced in hydrolysis tank flowed by pumped to the pH regulator tank. The tank was in the range of pH 7 using Na2CO3, and then pumped it into biogas formation tank. In this biogas formation, disaccharide and monosaccharide was mixed with methanogenic microbial starter 10% (v/v) and tapioca wastewater as substrate of 100 ml every 12 days.

Basic experimental of 1 stage and 2 stage fermentation be the same. In 1 stage fermentation, the wastes directly neutralized with buffer and mixed with rumen bacteria with a concentration of 10% (v/v) without yeast, one of its pH tank is not regulated and without addition of substrate, and in each tank was added rumen bacteria with different concentration of 8%, 15%, 20% (v/v) without addition of substrate.

Test results were carried out by analysis amount of methane by using a water displacement technique, temperature and pH measurements was carried out every day. Observations were made continuously until the methane was not formed anymore.

3 Results and Discussions

Biogas is produced depends on composition of feed material. Organic materials that can be used as raw material for biogas production are animal waste, solid waste, sewage sludge, etc. This study used tapioca...
wastewater with batch and semi-continuous processes. Characteristics of the biogas produced in each process are described below. Experiments were carried out to raise the levels of C/N of 2 liters tapioca waste water tapioca 1% (w/v) in each tank with urea 0.4% (w/v). Experiments with rumen concentration (8%, 15%, 20% v/v), yeast concentration 0.15% (w/v), with Na₂CO₃ buffer batch in 2 stage fermentation system obtained the following results in Figure 2.

**Figure 2. The Effect of Substrate Activator to Biogas Production**

Figure 2 describes the relationship of methanogenic bacteria (rumen) concentration in anaerobic fermentation process. It can be seen that the daily biogas production rate is best obtained when the addition of methanogenic bacteria by 8% [1, 13]. This is due to the process of biogas production with tapioca waste material optimum obtained when the number of methanogenic bacteria by 20%.

Bacteria which first worked out in the process of changing such complex carbohydrate polymer is cellulolytic bacteria or other hydrolytic bacteria. Cellulolytic bacteria break or cut the cellulose molecule, the molecule with high weight into cellulobiose (glucose) and free glucose. Glucose is then fermented anaerobically to produce a variety of fermentation products such as acetate, propionate, butyrate, H₂ and CO₂. H₂ results from primary fermentation immediately used by methanogenic bacteria (methanogens) which is the last bacteria used in anaerobic fermentation process. In addition, it is also required for the conversion of acetate to methane in anaerobic fermentation by some methanogenic bacteria.

At this stage of methane gas formation, methanogens bacteria plays the role. Methanogens bacteria will utilize the results of second stage, such as formats, carbon dioxide, and hydrogen as substrates to produce methane, carbon dioxide, trace of gases such as H₂S and water. Data shows that the highest accumulation of biogas production obtained when addition of 20% (v/v) methanogenic bacteria is 2458 ml or 122.9 ml/g TS. While the production of biogas for variable concentrations of methanogenic bacteria by 8% (v/v) and 15% (v/v) are 2105 ml or 105.25 ml/g TS and 105.85 ml or 2117 ml/g TS. Biogas production on addition of 20% (v/v) methanogenic bacteria is lower in the beginning of process than the addition of 15% (v/v) methanogenic bacteria. It happened until the fourth day. The characteristics of resulting curve above in accordance with growth curve of bacteria which consisting of four stages, such as lag stage, exponential stage, logarithmic stage, stationary stage, and death stage.

Methanogens bacteria growth in the beginning of process is still going through adjustment to the raw material circumstances in which will be broken down into the biomass, in terms of nutrients, pH, or temperature that corresponds to its place. Furthermore, these bacteria will experience rapid growth process that will produce maximum biogas because the use of good nutrition by methanogens bacteria. The next stage, bacteria begin lack of nutrition in which the number of bacteria that grow as much as the bacteria die so that the biogas produced is relatively constant (fixed). And at the end of the fermentation process the bacteria have started to die, so that biogas production has begun to decline.

There are five generation of methanogenic bacteria that play a role in the formation of methane, namely: Methanobacterium, Methanobacillus, Methanopyrales, Methanococcus, and Methanosarcina. Methanogens bacteria are the most important bacteria in the decomposition anaerobic process. This bacterium is a group of organisms that are very sensitive to oxygen and pH changes in the medium. Bacterial growth rate was slower and more sensitive to changes in the environment when compared with bacterial non-methanogens. Special ability of these bacteria is to remove the excess electrons in an effective way by converting hydrogen into methane which causes the hydrogen concentration in medium remained low. It is important for the presence of cytogenesis bacteria and ultimately led the decomposition process can continue effectively [14].

In the process of anaerobic fermentation for biogas formation used cow rumen as microbes decomposer. There are three groups of bacteria that play a role in anaerobic fermentation, such as:

1. Polymers and monomers bacteria decomposer contained in the waste material and produces mainly acetate and hydrogen, and a number of volatile fatty acids like propionic, butyric and alcohol.
2. Acetogenic obligate bacteria produced hydrogen that covert propionate and butyrate become acetate and hydrogen.
3. Groups of methanogenic bacteria producing methane from acetate or hydrogen.

The final product of cellulose decomposition by cellulolytic bacteria were succinate, acetate, formate or butyrate. Below is a chart of pH and temperature characteristics at various feed composition for variable addition of methanogenic bacteria.

From Figure 3-5, it can be seen that for variable concentrations of methanogenic bacteria (rumen bacteria) showed that in the tank I have temperature range between 27 to 31.20°C, tank II between 26 to 32.50°C, and tank III between 26 to 33.8°C. From the graph above can be seen temperature fluctuates in each tank for each variable with different ranges. All temperature range is still in the mesophilic temperature range between 28 – 45°C. In addition to mesophilic temperature, anaerobic fermentation process can also occur in the thermophilic temperature, in temperature
range 45 – 65°C. When the temperature decreased, the activity of microorganisms also decreased so that the production of biogas will be decreased.

Fig 3. Methanogenic Bacteria Concentration of 8%

Fig 4. Methanogenic Bacteria Concentration of 15%

Fig 5. Methanogenic Bacteria Concentration of 20%

In the terms of pH, it can be seen that for variable concentrations of methanogenic bacteria (rumen bacteria) showed that for tank I have pH range between 5.31 to 7, tank II between 5.2 to 7, and tank III between 5.26 to 7. At initial fermentation, acid-forming bacteria will produce acids that can quickly cause pH drop. That acid formation will produce acetic acid, H₂ gas, and some VFA, such as butyric acid and propionate. When the pH value is low, the microorganisms will be in-active state in order to influence the rate of biogas formation, especially methanogenic bacteria. However, it can be handled with the addition of Na₂CO₃ buffer to improve its alkalinity. Methanogenic bacteria were able to grow well in pH range from 6.8 to 7.2 [15]. A good relationship between asidogenesis and methanogens stage is in neutral pH (7) and no dramatically increase in acidity and alkalinity [16].

From Figure 4 above can be seen that the addition of Na₂CO₃ buffer is different each day on each tank. In the tank I, the highest addition buffer is 3.13 grams that achieved on fourth day. Tank II, the highest addition of buffer is 4 grams which occurred on first day. This is because in those days, the pH in the tank II was drop quite dramatically, reaching 4.97 so that the required buffer is much to return it to neutral pH of around 7. While for the tank III, the highest buffer addition is 3.65 grams on first day. In this tank, on first day already showed pH drop reached 5.35.

The addition of Na₂CO₃ buffer every day is different on each tank. This is because the pH drop in each tank also vary every day. Tank with high pH drop will require a lot of buffer, and vice versa. In experiment in variable with or without addition of Na₂CO₃ buffer in 2 stage batch fermentation obtained following results in Figure 6.

Figure 6 describes the effect of buffers on the rate of biogas formation. It can be explained that the production of biogas by using buffer Na₂CO₃ much less than not using buffer. On day 4, the process of biogas formation by using buffers on feed achieving the highest number of 372 ml with microbial life during 16 days. Rapid formation of acid can cause a number of methanogens bacterial be dead because it is not resistant to acidic conditions. In this case, using Na₂CO₃ aims to maintain pH range so bacteria can survive longer.

While fermentation without buffer, the biogas production is 620 ml. In this pH rumen bacteria have died because the life span of methanogens bacterial was lowest at pH 5, so it can be concluded that the gas methane was not formed, but CO₂. Cessation of biogas production is due to the pH getting down (pH drop) until it reaches 4. These pH conditions cause all the organisms die, so the biogas production ceased.

Figure 7 shows that the highest accumulation of biogas production obtained on the addition of buffer 2117 ml or 105.85 ml/g TS. On the 3rd day of biogas production on fermentation with buffer less rapidly with fermentation without buffer, this is because the gas generated in variable without buffer is not CO₂ but methane. At this time the pH of medium was around 4-5 which optimum pH for yeast growth is. It happened until the 9th day. However, in variable with addition of buffer
has higher accumulation. The addition of buffer resulted in pH medium being in the range of methanogens bacteria growth, so biogas production can continue until the substrate discharged on 16th day. In variable without buffer the production stopped 9th day. This is due to all microbes die because the pH is too low.

In semi-continuous process, which is the feed pumped into tank I consists of 1% (w/v) tapioca waste, urea 0.04% (w/v), and rumen bacteria 10% (v/v). Tank 2 consists of tapioca waste 1% (w/v), urea 0.04% (w/v), yeast 0.08% (w/v), and rumen bacteria 10% (v/v). The addition of 8 grams tapioca waste fresh feed done every 12 days.

Figure 8 shows that tank I consist of tapioca waste 1% (w/v), urea 0.04% (w/v), and rumen bacteria 10% (v/v). Biogas production rate in tank I is still inferior to tank II that consist of tapioca waste 1% (w/v), urea 0.04% (w/v), rumen bacteria 10% (v/v), and yeast 0.08% (w/v). The peak of production in tank II reached on 3rd day that is equal to 625 ml, while tank I was on 6th day that equal to 642 ml. It is because the tank II hydrolyse polysaccharide molecules into simpler molecules is faster with the help of Saccharomyces cerevisiae. So that the substrates used by methanogens bacterial is substrate with a shorter chain so that the process is also more rapid degradation. Therefore, biogas production rate in the tank II is faster than tank I.

Figure 9 showed the accumulation of biogas production in vary feed composition variable. It can be seen that the accumulation of biogas production in tank 2 is higher than in tank 1. It happens because in the tank 2 contained of yeast (Saccharomyces cerevisiae) which acts as substrate activator. Substrate activator can increase the rate of biogas production because it can speed up hydrolysis process by hydrolytic fermentative bacteria that produce simpler compounds for methanogens bacterial substrate. Total production of biogas in tank I is 6831 ml or 155.25 ml/g TS, and in tank II is 9329 ml or 212.02 ml/g TS.
butyrate, H₂, and CO₂. H₂ resulted from primary fermentation immediately used by methanogenic bacteria which is the last bacteria used in the anaerobic fermentation process. In addition, acetate is also needed for conversion to methane in anaerobic fermentation by some methanogenic bacteria. On the other hand, methanogenic bacteria were able to grow well by addition of buffer solution to increase its alkalinity. The addition of buffer used if the pH drop was above 1. Magnitude of the addition of Na₂CO₃ buffer varies in each tank.

The growth rate of the acid-forming bacteria run faster than the growth of methanogenic bacteria, so the population of methanogenic bacteria is not sufficient to consume the amount of acid production. Excess of acid production in the beginning of the process could lead to die some methanogenic bacteria. On the other hand, methanogenic bacteria were able to grow well on the medium that was initially in range of pH 6.8 to 7.2 with a pH drop is not greater than 1 [16]. From Fig. 10 can be seen that pH drop was greater than 1 at day 21. Good relationship between asidogenesis and methanogens stage is currently in a state of pH drop below 1, because in this state methanogens bacteria do not die in acidity medium levels that drop too fast.

In terms of temperature, from Fig. 10 can be seen that for the variable concentrations of methanogenic bacteria (rumen bacteria) in biodigester has temperature range between 28 – 30°C. The temperature fluctuates in each tank for each variable with different ranges. All of the temperature range is still in the mesophilic temperature range, it is between 28 – 45°C. When the temperature is decreased below 28°C, the microorganism activity decreases which resulting in decreasing biogas production. However, when the temperature rose again, microbial activity will takes place as it should.

4 Conclusions

Biogas production is determined by the concentration of rumen bacteria, the addition of Na₂CO₃ buffer, and the type of fermentation. Greater concentration of rumen will increase accumulation of biogas production because there are more decomposer and rate of acid formation could be offset by formation of methane. The addition of Na₂CO₃ buffer able to providing biogas production period longer and the volume of biogas was greater than without addition of Na₂CO₃ buffer because pH drop in early process can be prevented. More complex fermentation type (2-stage) produce more biogas volume due process hydrolysis is faster.

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