Bioprocess development of 2, 3-butanediol production using agro-industrial residues

Sulfath Hakkim Hazeena¹,² · Narasinha J. Shurpali³,⁴ · Henri Siljanen³ · Reijo Lappalainen⁵ · Puthiyamdam Anoop¹ · Velayudhanpillai Prasannakumari Adarsh¹ · Raveendran Sindhu¹ · Ashok Pandey⁶,⁷,⁸ · Parameswaran Binod¹,²

Received: 25 April 2022 / Accepted: 18 July 2022 / Published online: 12 August 2022 © The Author(s) 2022

Abstract
The valorization of agricultural and industrial wastes for fuel and chemical production benefits environmental sustainability. 2, 3-Butanediol (2,3-BDO) is a value-added platform chemical covering many industrial applications. Since the global market is increasing drastically, production rates have to increase. In order to replace the current petroleum-based 2,3-BDO production, renewable feedstock's ability has been studied for the past few decades. This study aims to find an improved bioprocess for producing 2,3-BDO from agricultural and industrial residues, consequently resulting in a low CO₂ emission bioprocess. For this, screening of 13 different biomass samples for hydrolyzable sugars has been done. Alkali pretreatment has been performed with the processed biomass and enzyme hydrolysis performed using commercial cellulase. Among all biomass hydrolysate oat hull and spruce bark biomass could produce the maximum amount of total reducing sugars. Later oat hull and spruce bark biomass with maximum hydrolyzable sugars have been selected for submerged fermentation studies using Enterobacter cloacae SG1. After fermentation, 37.59 and 26.74 g/L of 2,3-BDO was obtained with oat hull and spruce bark biomass, respectively. The compositional analysis of each step of biomass processing has been performed and changes in each component have been evaluated. The compositional analysis has revealed that biomass composition has changed significantly after pretreatment and hydrolysis leading to a remarkable release of sugars which can be utilized

* Narasinha J. Shurpali
narasinha.shurpali@luke.fi

¹ Microbial Processes and Technology Division, CSIR-National Institute for Interdisciplinary Science and Technology (CSIR-NIIST), Thiruvananthapuram 695019, Kerala, India
² Academy of Scientific and Innovative Research (AcSIR), Ghaziabad 201002, India
³ Department of Environmental and Biological Sciences, University of Eastern Finland, Kuopio campus, Kuopio, Finland
⁴ Natural Resources Institute Finland (Luke), Halolantie 31 A, 71750 Maaninka, FI, Finland
⁵ Biomaterials Technology, Dept. of Applied Physics & SIB-Labs, University of Eastern Finland (Kuopio Campus), Yliopistonranta 1 F, 70211 Kuopio, FI, Finland
⁶ Centre for Innovation and Translational Research, CSIR-Indian Institute of Toxicology Research, Lucknow 226 001, India
⁷ Sustainability Cluster, School of Engineering, University of Petroleum and Energy Studies, 248 007, Dehradun, India
⁸ Centre for Energy and Environmental Sustainability, Lucknow 226 029, India
by bacteria for 2,3-BDO production. The results have been found to be promising, showing the potential of waste biomass residues as a low-cost raw material for 2,3-BDO production and thus a new lead in an efficient waste management approach for less CO₂ emission.

**Graphical Abstract**

- 13 different agro-industrial residues were tested for fermentable sugars.
- Oat hull and spruce bark biomass hydrolysate were used for 2,3-Butanediol production.
• Batch fermentation with oat hull hydrolyzate yielded 37.59 g/L of 2,3-butanediol.
• Fermentation using spruce bark hydrolyzate yielded 26.74/L of 2,3-butanediol.

**Keywords** 2, 3-Butanediol · Biomass · Fermentation · Bioprocess

**Introduction**

Despite current global regulations for climate change, atmospheric greenhouse gas levels are skyrocketing. According to the International Energy Agency (IEA), global energy-related CO₂ emission was around 33 gigatonnes in 2019 (IEA (2020)). This number reveals that our regulatory approaches are not impressive enough to cope with real-world emissions. Climate-related risks are adversely affecting health, food security, water supply, and thereby economic growth. More effective strategies and policies have to be shaped to tackle this issue precisely.

Dependence on fossil fuel is the major cause of anthropogenic emissions, not only for fuel and power but also for the manufacture of platform chemicals. Many value-added platform chemicals that are currently being synthesized via petrochemical routes can be derived from renewable biomass. 2, 3-Butanediol (2,3-BDO) is currently produced from petroleum routes and potentially could be produced from biomass [1].

2, 3-BDO has wide applications in agriculture, pharmaceuticals, and polymer industry. It can be derivatized to high-value fuel additive due to its high heat of combustion [2]. Optically pure 2,3-BDO is used in the synthesis of chiral compounds [3]. Levo (2R,3R (−)) form of 2,3-BDO has been used as a potential antifreeze agent in the pharmaceutical industry because of its very low freezing point (−60 °C) [4]. According to Ameco market research, 2,3-BDO market is expected to reach 10300 million US dollars by 2025.

Biological synthesis of 2, 3-BDO has a long history from 1906. It was first reported in *Klebsiella pneumoniae* and later in other species such as *Klebsiella, Serratia, Enterobacter, Bacillus, and Paenibacillus polymyxa* [5]. Along with these native producers, a couple of heterologous hosts such as *S. cerevisiae* and *E. coli* were successfully demonstrated to produce 2, 3-BDO [6] [7]. Different renewable feedstock such as lignocellulosic biomass, biodiesel-derived glycerol and non-crop plants were tested for 2, 3-BDO production. Studies shows that sugarcane bagasse pretreated with green liquor, containing Na₂CO₃ and Na₂SO₃, followed by enzymatic hydrolysis can be used as a carbon source for producing 2,3-butanediol. The yield of 0.395 g/g sugar was reached after 72 h of fermentation, indicating that the lignocellulosic biomass could be used to produce 2,3-BDO affordably using metabolically modified *Enterobacter aerogenes* [8]. Saratale and coworkers successfully performed the pretreatment of kenaf core biomass with inorganic salts and calcium peroxide along with their use in the synthesis of 2,3-BDO [9]. Biologically derived 2, 3-BDO was commercialized by a few industries [5].

Current biological production yields are not sufficient enough with wild-type strains. Substrate cost is another major limiting factor in industrial bioprocess for 2, 3-BDO production. Its production from agro-industrial residues of the Finnish ecosystem has relevance in terms of waste management system for less CO₂ emission. Integrating a waste management system for value-added chemical production will have potential benefits. The current study covers a wide spectrum of biomass wastes from Finnish agricultural and industrial sector and evaluates their ability for generating fermentable sugars. Later the study demonstrates a renewable methodology for production of 2,3-BDO from biomass hydrolysat via fermentation. To decrease the current anthropogenic emission, an attempt has been made by adopting a renewable route for the production of 2,3-BDO, and in future for the commercial synthesis of 2,3-BDO this will be a relevant reference. The study aims to develop a renewable route for the production of 2,3-BDO using agricultural waste from Finnish agricultural sector. For this purpose, different agricultural residues were chosen and different pretreatment strategies were employed for the degradation of cellulose structure. Later, enzymatic hydrolysis of cellulose was performed using commercial cellulase and the resulting hydrolysate has been used for 2,3-BDO fermentation. Among all biomass, one with highest release of total reducing sugars has been chosen for further experiment. *Enterobacter cloacae* SG1 has been used for 2,3-BDO fermentation and the efficiency of the process evaluated. Moreover, the compositional analysis of biomass after each step of pretreatment, hydrolysis, and fermentation was performed to evaluate the effectiveness of the process.

**Materials and methods**

**Media and chemicals**

(2S, 3S)−(−), (2R, 3R)−(−) and meso-2,3BDO and acetoin (>98%) were procured from Merck (Germany). All other chemicals of analytical grade were used in this study. Fermentation media components were (in gram per Liters) yeast extracts-5.0, KH₂PO₄-6.0, K₂HPO₄-14.0, Sodium citrate dehydrate-1.0, ammonium sulphate-2.0, and magnesium sulphate heptahydrate-0.2.
Biomass samples

Thirteen agro-industrial residues were selected for the study. The agro-residual biomass samples and biogas digestate were received from Maaninka Research Station, Kuopio, Finland. Leaf samples were collected from Municipal Sewage Waste collection facility, Kuopio, Finland. Wood bark and chip samples were local industrial samples used for bioenergy production. Hemp hurd was obtained from Futura 75 fiber hemp grown in Northern Savo. Paper mill effluent samples were received as frozen from Mondi Powerflute Oy, Kuopio, Finland. The raw biomass, except oat hull and barley hull, were milled to a particle size of 3–5 cm length, 2–3 cm breadth and 1 cm thickness, dried and stored at room temperature until used. Oat hull and barley hull were processed for pretreatment and hydrolysis as such from industrial residues normally used for bioenergy.

Pretreatment of biomass

The biomass samples (15% w/w) were pretreated in 250 mL Erlenmeyer flasks with 1.5% NaOH (w/w) at 121 °C, 15 lbs for 20 min. After cooling, excess alkali was washed thoroughly with water and filtered and dried at 65 °C for 12 h and subjected to enzymatic hydrolysis. Liquid samples such as paper mill effluents and biogas digestate were tested for reducing sugar availability without any treatment.

Enzymatic hydrolysis

Enzyme hydrolysis of biomass was performed in 1 M citrate buffer (pH 4.8) using Trichoderma reesei cellulase (Sigma Aldrich). The hydrolysis conditions were as follows: biomass loading 10% (w/w), enzyme loading 20 FPU (filter paper unit) and antibiotic loading 0.001% (w/w) incubation at 50 °C, 200 rpm. Water and buffer were added to the biomass and allowed to equilibrate at 50 °C. After this, the antibiotic solution and enzyme were added so that the fermentation reaction could proceed. Samples were collected in every 24 h and checked for total reducing sugars.

Preliminary screening of the biomass for sugar yield

For the preliminary screening of the biomass materials, the pretreated and hydrolysed samples were analyzed for reducing sugars by DNS method [9].

Compositional analysis

The composition of selected native and alkali pretreated samples were determined according to National Renewable Energy Laboratory (NREL) analytical methods for biomass (Sluiter et al. 2011).

Culture maintenance

Enterobacter cloacae sp. SG1 was maintained as glycerol stocks and sub cultured regularly in nutrient agar plates. Peptone, beef extract**** and NaCl were used for preparing seed media.

Fermentation for 2, 3-BDO production

Fermentation medium was inoculated with 24 h-old inoculums. Media components and biomass hydrolysate were sterilized by autoclaving separately and added during the time of inoculation. 20 g/L purified glucose was added to the media separately. Fermentation conditions were pH 6.5, temperature at 37 °C, and 200 rpm for 120 h. Samples were withdrawn periodically and analyzed for 2, 3-BDO and sugar [10].

Analytical methods

Total reducing sugar concentration was estimated by DNS method [11]. Compositional analysis of biomass at various experimental stages was done according to NREL protocol [12]. Bacterial growth was monitored spectrophotometrically by checking optical density at 620 nm (Shimadzu, Japan).

Statistical analysis

The experiments were repeated for a minimum of three times. All data were expressed as means ± SD. Statistical differences between control and treated groups were evaluated using Student’s t test, and differences between groups were considered statistically significant at p-values < 0.05.

Results

Preliminary screening of the biomass

Preliminary screening of agricultural and industrial biomass for 2, 3-BDO fermentation was done based on the ability to release reducing sugars for microbial growth and fermentation. For this purpose, alkali pretreatment was done at high temperature and pressure which would eventually lead to lignin removal and disrupt the crystallinity of cellulose. Then enzymatic hydrolysis of cellulose was performed using cellulase from Trichoderma reesei. Cellulase enzyme
cleaves cellulose fibers to individual glucose molecules and finally, this can be utilized by bacteria for growth and 2, 3-BDO production. Upon alkali pretreatment and hydrolysis, the sugar release from the biomass at 24 h of hydrolysis was estimated (Table 1).

Among all biomass tested except paper industry wastes such as wood chip wash water and paper mill condensate possess a little amount of reducing sugars in it. Biogas digestate samples also contain a small amounts of free sugars. Birch bark biomass was also detected with zero amount of reducing sugars. All other biomass including leaf wastes, hemp, oat, barley and aspen biomass contains a significant amount of sugars that can be valorized for energy purposes or can be used as the low-cost feedstock for fermentation. Maximum sugar release during 24 h of hydrolysis was found from oat husk (84.27 g/L) followed by hemp (46.36 g/L) and spruce bark biomass (46.21 g/L). The sugars released from leaf samples were also found to be promising. The liquid samples such as paper mill effluent and biogas digestate contain very little or no reducing sugars. Oat hull and spruce bark biomass which gave a higher amount of reducing sugars were selected for 2,3-BDO fermentation.

### Compositional variability of biomass

The composition of native, alkali-pretreated biomass and the residue after enzymatic hydrolysis are presented in Table 2. The composition analysis provides detailed picture of the native composition of individual components in biomass and the changes that occurred in each step of pretreatment and hydrolysis. Composition analysis thus assists in calculating the efficacy of the whole process from pretreatment to fermentation.

The cellulose content in the native oat hull was higher (61.52) than that of spruce bark (47.15). After alkali pretreatment, hemicellulose fraction was found to increase in its percentage, mainly because of the lignin removal. Lignin removal was efficient in oat hull and close to 50% lignin removal was observed in alkali-pretreated biomass (12.7) compared to the native (22.24). The lignin content in spruce (44.55) is almost twofold higher than oat hull (22.24). A notable amount of lignin was reduced in spruce biomass (from 44.55 to 29.75) in pretreatment. A corresponding increase was observed with cellulose fraction (47.15 to 64.96) in spruce biomass at the same time. In enzymatic hydrolysis 33% of cellulose was hydrolyzed in oat hull while it was 31% in spruce bark. After enzyme hydrolysis, lignin fraction was reduced from 22.24% to 12.98% in oat hull biomass while it is 44.5 and 25.32, respectively, in spruce. The relative removal of lignin concentration was prominent in spruce biomass.

### Pretreatment and enzymatic hydrolysis

The cellulose in biomass was degraded to glucose and finally, utilized by the bacteria for growth and 2, 3-BDO production. DNS analysis of oat hull and spruce bark biomass showed 31.56 g/L and 25.45 g/L of total reducing sugars, respectively. Detailed HPLC analysis revealed individual sugars present in hydrolysate. 11.32, and 9.62 g/L glucose was present in oat hull and spruce bark hydrolysate fraction. Additionally, arabinose (2.6 g/L) was also found present in oat hull hydrolysate. The hydrolysate was a mixture of both

### Table 1 Total reducing sugar concentration in different biomass

| Sl No | Biomass          | Total reducing sugars (g/L) at 24 h of hydrolysis |
|-------|------------------|-----------------------------------------------|
| 1     | Hemp hurd        | 46.36 ± 7.65                                  |
| 2     | Aspen bark       | 20.93 ± 0.03                                  |
| 3     | Oat hull         | 84.27 ± 0.09                                  |
| 4     | Barley           | 22.57 ± 0.30                                  |
| 5     | Spruce bark      | 46.21 ± 0.92                                  |
| 6     | Wood chips       | 22.9 ± 0.00                                   |
| 7     | Birch bark       | 0.00                                          |
| 8     | Leaf type A      | 46.8 ± 2.8                                    |
| 9     | Leaf type B      | 27.49 ± 8.8                                   |
| 10    | Digestate sample 1 | 1.72                                         |
| 11    | Digestate sample 2 | 2.04                                         |
| 12    | Paper mill condensate | Nil                                    |
| 13    | Wood chip wash water | 1.71 ± 0.98                                 |

### Table 2 Compositional variation of the biomass used in the study

| Sample description                  | Cellulose (%) | Hemicellulose (%) | Lignin (%) | Ash (%) | Total (%) |
|-------------------------------------|---------------|------------------|------------|---------|-----------|
| Oat hull biomass native             | 61.52 ± 0.95  | 7.31 ± 0.37      | 22.24 ± 0.58 | 1.53 ± 0.15 | 92.6 ± 1.93 |
| Oat hull biomass pretreated         | 64.59 ± 1.78  | 11.19 ± 1.21     | 12.7 ± 1.87 | 0.399 ± 0.033 | 88.88 ± 2.54 |
| Oat hull biomass residue after hydrolysis | 31.92 ± 1.2  | 19.68 ± 1.03     | 12.98 ± 0.3 | 0.42 ± 0.011 | 64.57 ± 22.41 |
| Spruce bark biomass native         | 47.15 ± 1.64  | 11.80 ± 0.43     | 44.55 ± 0.26 | 0.166 ± 0.033 | 103.67 ± 2.06 |
| Spruce bark biomass pretreated     | 64.96 ± 0.28  | 6.78 ± 1.07      | 29.75 ± 1.02 | 0.099 ± 0.033 | 101.59 ± 2.34 |
| Spruce bark biomass residue after hydrolysis | 34.06 ± 0.92 | 13.17 ± 1.34     | 25.32 ± 0.44 | 0.37 ± 0.003 | 72.49 ± 0.79 |
pentoses and hexoses with slight amount of inhibitors such as acetate.

**Fermentation of biomass hydrolysate for 2, 3-BDO production**

2, 3-BDO fermentation was initiated by the inoculation of *Enterobacter cloacae* SG1 into the hydrolysate medium. HPLC analysis reveals the individual concentration of major products during 2, 3-BDO fermentation. Apart from 2, 3-BDO, acetate and acetoin were produced predominantly and their concentrations depicted in Figs. 1 and 2. After 24 h of each batch fermentation, 37.59 g/L of 2,3-BDO from oat hull hydrolysate were obtained. Also, 20.72 g/L of acetoin was found to be co-produced with 2, 3-BDO (Fig. 1). Around 26.74 g/L of 2,3-BDO was produced from spruce bark biomass along with 20.36 g/L of acetoin (Fig. 2). Since the reaction between 2, 3-BDO and acetoin is reversible it is clear from the figure that in 24 h, 2, 3-BDO concentration decreases while that of acetoin increases. The direction of reaction could be changed depending upon the concentration of each of the involved products. When the medium became more anaerobic there was a tendency to accumulate acetoin while 2,3-BDO prefers microaerophilic conditions. This can be noted by the accumulation of acetoin along the time of fermentation. A gradual decrease in the oxygen level in the medium would result in acetoin accumulation. The maximum 2,3-BDO production was achieved in 24 h of fermentation and almost 90% sugar utilization was also noticed within this period. Even though acetoin concentration was found to be increasing after 24 h, the maximum 2,3-BDO was obtained at 24th hour itself. So prolonging the fermentation time does not have any remarkable effect on product yield. In oat hydrolysate, 0.39 g/L of acetate was present initially after pretreatment. Acetate is formed in the hydrolysate mainly because of the hydrolysis of acetyl groups in hemicellulose [13]. Acetate is one of the major products of mixed acid fermentation pathway. Acetate concentration gradually increased and reached a maximum of 2.0 g/L in 48 h of fermentation.

**Growth of *E. cloacae* SG1 in biomass hydrolysate**

Growth of *E. cloacae* SG1 was monitored spectrophotometrically in each 24 h of fermentation. The growth pattern of *E. cloacae* SG1 in Fig. 3 was a prototype of bacterial growth curve. The logarithmic phase was achieved at 24 h of incubation and later it proceeded to the stationary phase. Growth in spruce biomass hydrolysate was slightly retarded compared to oat hull biomass. This may be because of the presence of pigments and phenolic compounds present in the bark.

![Fig. 1 Production of 2,3-BDO using oat hull hydrolysate](image-url)
Discussion

2,3-BDO fermentation using different low-cost substrates, both lignocellulosic and non-lignocellulosic feedstock, has been well studied in the past decades. The substrates include food industry wastes, wood hydrolysate, algal biomass, molasses, and many other non-crop substrates like Jerusalem artichoke tubers [14] [15] [16] have successfully

Fig. 2 Production of 2,3-BDO using spruce bark biomass hydrolysate

Fig. 3 Growth pattern of \textit{E. cloacae} SG1 in oat hull and spruce bark hydrolysate
been evaluated for its ability to produce 2,3-BDO. This study attempted to valorize a broad-spectrum agricultural and industrial waste for 2,3-BDO production especially from Finnish agricultural and industrial area. This comprehensive evaluation of agricultural waste residues, particularly for the Finnish agricultural sector, was reported for the first time. All the biomass tested were available abundantly and at cheap costs. Since one of the challenges in bioenergy production is substrate cost [17] and availability, these factors should be considered essential criteria for selecting feedstock.

Almost a major number of biomasses tested contain a significantly higher amount of reducing sugars. However, the splendid availability of oats hull and spruce bark biomass raises them for special attention. The oat cultivation status of Finland for the past years showed a relative abundance of oat availability. According to Luke (Natural resources Institute, Finland), 1200 million kilograms of oat were cultivated in Finland in 2019. Accordingly, a significant amount of oat hull has been generated after its processing. Burning up this much biomass will bring environmental concerns as we mentioned in the introduction. Oat hull comprises 28% of grain weight and contains 45% cellulose [18]. Among the biomass, oat and spruce bark biomass have been found to be the maximum reducible sugars and were selected for further studies. Even though studies explaining the cellulose composition of different biomass including spruce [19] and oat [20] the composition can vary due to different factors including species variation and climatic conditions [21].

Composition analyses have revealed the amount of each component of the lignocellulose biomass. Lignin can act as hindering the binding of cellulose degrading enzyme by contributing non-specific binding [22]. It can also act as a potential source of origin of phenolic compounds that eventually leads to retardation of bacterial growth [23]. A significant amount of lignin could be removed by alkali pretreatment [24]. Alternate pretreatment strategies such as alkaline and organic solvent treatment on biomass such as corn stover, poplar and Douglas fir produced significant amount of sugars and on subsequent fermentation of the sugars resulted 2,3-BDO [25]. 2,3-BDO production has also been reported using Jatropha hulls after ionic liquid pretreatment followed by dilute acid hydrolysis [26]. Similarly, in oat and spruce biomass, after efficient lignin removal, enzymatic hydrolysis has been reported to lead to an efficient conversion of polysaccharides to monosaccharides.

As the spruce bark biomass was composed of pigments and phenolics, and the hydrolysate was itself darker in appearance. Because of such inhibitory compounds, a retarded growth, and corresponding less 2,3-BDO production were noted with spruce biomass hydrolysate compared to oat hull. Similar observation was noted with Strizincova et al., in spruce bark biomass [27]. Phenolic compounds and degradation products of sugars are inhibitory compounds for microbial growth and fermentation. The inhibitory phenolic compounds are generally removed by overliming [28]. Here no treatment for inhibitor removal was performed because it will cause significant removal of sugars and reduce the final product yield. In the case of oat hull biomass, even though lignin removal was happening to a good extent, cellulose fraction was not increasing accordingly. While analyzing the sugar composition of oat biomass hydrolysate along with glucose, arabinose was also found to be present. The strain E. cloacae SG1 is known for its ability of using both hexoses and pentoses [29]. While using biomass-derived sugars it is important to check the organism’s ability to utilize different sugars as the carbon source for fermentation. Recently a newly isolated Cronobacter sakazakii was reported for its ability to utilize both glucose and xylose for 2,3-BDO production [30].

Other than monosaccharides a smaller amount of cellobiose was also found to be present in the hydrolysate. This indicates the incomplete hydrolysis reaction that occurred in cellulose fraction. The incomplete hydrolysis can reduce the efficiency of fermentation as the bacteria cannot use partially hydrolyzed cellulose [31]. The effectiveness of hydrolysis of biomass has to be addressed in an industrial bioprocess. While using oat and spruce biomass hydrolysate significant amount of 2,3-BDO was produced. Okonkwo and coworkers observed similar 2,3-BDO production from non-detoxified wheat straw hydrolysate using Paenibacillus polymyxa DSM 365 [32]. Similarly, 32.7 g/L 2,3-BDO was produced from nondetoxified sugarcane bagasse hydrolysate-derived sugars by Enterobacter ludwigi [33]. It has been demonstrated that, while conducting a non-sterile fermentation using non-sterile food waste using a thermophilic Bacillus licheniformis YNPS-TSU, the 2,3-BDO production was less in comparison with sterile conditions [34]. An improvised study using the same bacteria showed a significant increase in 2,3-BDO production because of the increase in initial sugar concentration in the hydrolysate [35]. It showed the effect of hydrolysis and media components on the diol titers. Different biomass and the corresponding 2,3-BDO titers using microbial fermentation are depicted in Table 3. Other than this biomasses, Brewers’ spent grain hydrolysate, bakery waste, and bread waste have also been found as potential source for 2,3-BDO production [36] [37] [38].

2,3-BDO can be recovered from fermentation broth by aqueous two phase extraction system using an organic solvent. Aqueous two phase extraction has been successfully used in the separation of 2,3-BDO produced using biomass hydrolysate [39]. The results were promising within a scale-up possibility since the 2,3-BDO titers using oat hull and spruce bark biomass were optimal. Valorizing lignocellulose waste material could be beneficial by reducing the CO₂ emission and utilizing the reserved carbon as fuel and
Biomass Pretreatment if any  | Microorganism  | 2,3-BDO (g/L)  | Reference  
---|---|---|---
Non-detoxified wheat straw  | Dilute acid hydrolysis  | *Pseudomonas polyaeroxidans* DSM 365  | 32.0  | [32]  
Jatropha hull  | Ionic liquor pretreatment  | *Klebsiella oxytoca*  | 33.49  | [26]  
Sugarcane bagasse  | Hydrothermal pretreatment  | *Enterobacter ludwigi*  | 32.7  | [33]  
Non sterile food waste  | NA  | *B. licheniformis* YNP5-TSU  | 5.9  | [34]  
Food waste hydrolysate  | NA  | *B. licheniformis* YNP5-TSU  | 36.7  | [35]  
Oat hull  | Alkali pretreatment  | *Enterobacter cloacae* SG1  | 37.59  | This study  
Spruce bark  |  |  | 26.74  |  

Table 3  Biomass hydrolysis for 2,3-BDO fermentation

Conclusion

The biomass samples from Finnish agricultural and industrial sector were tested for 2,3-BDO production of fermentable sugars. Among the 13 biomass samples tested, oat hull, spruce bark biomass, hemp hurd, and leaf waste produced significant amount of fermentable sugars (84.27, 46.21, 46.36, and 46.8 g/L, respectively) and these biomass were found to be promising low-cost substrates for valorization to fuel and chemicals. Alkali pretreatment followed by enzyme hydrolysis significantly alters the composition of biomass by releasing the sugar moiety from cellulose fraction which could finally be used in 2,3-BDO fermentation. Results on 2,3-BDO fermentation was found to be promising and have the potential to be upscaled to industrial level. 2,3-BDO fermentation with oat hull and spruce bark biomass could produce 37.59 and 26.74 g/L, respectively, in submerged fermentation. The successful production of high-value 2,3-BDO from comparably cheaper biomass helps in developing efficient strategies for commercialization of biomass-derived fuels and chemicals. Increasing our knowledge about unexploited biomass wastes for value-added chemicals and fuels will have a bright future in renewable energy generation.

Acknowledgements  Sultaf Hakim Hazeena acknowledge SERB Overseas visiting doctoral fellowship program 2018-19 supported by Science & Engineering Research Board (SERB), Department of Science and Technology (DST), Govt. of India for financial assistance for this study (SERB/F/5367/2019-20). SH Hazeena would like to acknowledge University Grants Commission (UGC), New Delhi for granting financial support for doctoral research. The authors would like to acknowledge Mondi Powerflute Oy, Kuopio, Finland for providing paper mill effluent samples for the study. The authors acknowledge the financial support by the Department of Science and Technology (DST), New Delhi under INNO-INDIGO/INDO-NORDEN project (Sanction No. DST/IMRCD/INNO-INDIGO/INDO-NORDEN/2017(G), Narasinha Shurpali acknowledges the financial support from the Project NC-GRASS (VN/28562/2020-MMM-2) and Project WoodPro (VN/17097/2022) funded by the Finland Ministry of Agriculture and Forestry.

Funding  Open access funding provided by Natural Resources Institute Finland (LUKE).

Declarations

Conflict of interest  The authors declare that there is no conflict of interest.

Open Access  This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

1. Hakizimana O, Matabaro E, Lee BH (2020) The current strategies and parameters for the enhanced microbial production of 2,3-butanediol. Biotechnol Report 25:e00397. https://doi.org/10.1016/j.btre.2019.e00397
2. Lee Y-G, Seo J-H (2019) Production of 2,3-butanediol from glucose and cassava hydrolysates by metabolically engineered industrial polyploid *Saccharomyces cerevisiae*. Biotechnol Biofuel 12:204. https://doi.org/10.1186/s13068-019-1545-1
3. Yang Z, Zhang Z (2018) Production of (2R, 3R)-2,3-butanediol using engineered Pichia pastoris: strain construction,
characterization and fermentation. Biotechnol Biofuel 11:35. https://doi.org/10.1186/s13068-018-1031-1
4. Yang Z, Zhang Z (2019) Recent advances on production of 2,3-butanediol using engineered microbes. Biotechnol Adv 37:569–578. https://doi.org/10.1016/j.biotechadv.2018.03.019
5. Song CW, Park JM, Chung SC, Lee SY, Song H (2019) Microbial production of 2,3-butanediol for industrial applications. J Ind Microbiol Biotechnol 46:1583–1601. https://doi.org/10.1007/s10295-019-02231-0
6. Erian AM, Gibisch M, Pflügl S (2018) Engineered E. coli W enables efficient 2,3-butanediol production from glucose and sugar beet molasses using defined minimal medium as economic basis. 06 biological sciences 0605 microbiology. Microb Cell Fac 17:1–17. https://doi.org/10.1186/s12934-018-1038-0
7. Kim S-J, Kim J-W, Lee Y-G, Park Y-C, Seo J-H (2017) Metabolic engineering of Saccharomyces cerevisiae for 2,3-butanediol production. Appl Microbiol Biotechnol 101:2241–2250. https://doi.org/10.1007/s00253-017-8172-1
8. Um J, Kim DG, Jung M-Y, Saratate GD, Oh M-K (2017) Metabolic engineering of Enterobacter aerogenes for 2,3-butanediol production from sugarcane bagasse hydrolysate. Bioresour Technol 245:1567–1574. https://doi.org/10.1016/j.biortech.2017.05.166
9. Saratate RG, Shin HS, Ghodake GS, Kumar G, Oh MK, Saratate GD (2018) Combined effect of inorganic salts with calcium peroxide pretreatment for kenaf core biomass and their utilization for 2,3-butanediol production. Bioresour Technol 258:26–32. https://doi.org/10.1016/j.biortech.2018.02.115
10. Hazeena SH, Nair Salini C, Sindhu R, Pandey A, Binod P (2019) Simultaneous saccharification and fermentation of oil palm front for the production of 2,3-butanediol. Bioresour Technol 278:145–149. https://doi.org/10.1016/j.biortech.2019.01.042
11. Miller GL (1959) Use of dinitrosoalicylic acid reagent for determination of reducing sugar. Anal Chem 31:426–428. https://doi.org/10.1021/ac60147a030
12. Adney B, Baker J (2008) Measurement of cellulase activities. Technical Report NREL/TP-510-42628 National Renewable Energy Laboratory, Colorado, USA (2008)
13. Jönsson LJ, Alriksson B, Nilvebrant N-O (2013) Bioconversion of lignocellulose: inhibitors and detoxification. Biotechnol Biofuel 6:16. https://doi.org/10.1186/1754-6834-6-16
14. Perego P, Converti A, Del Borghi A, Canepa P (2000) 2,3-Butanediol production by Enterobacter aerogenes: selection of the optimal conditions and application to food industry residues. Bioprocess Eng 23:613–620. https://doi.org/10.1016/s004900002010
15. Dai J-Y, Zhao P, Cheng X-L, Xiu Z-L (2015) Enhanced production of 2,3-butanediol from sugarcane molasses. Appl Biochem Biotechnol 175:3014–3024. https://doi.org/10.1007/s10210-015-1481-x
16. Qi Y, Lei P, Zhang Y, Sha Y, Zhan Y, Xu Z, Li S, Xu H, Ouyang P (2018) Recent advances in bio-based multi-products of agricultural Jerusalem artichoke resources. Biotechnol Biofuel 11:151. https://doi.org/10.1186/s13068-018-1152-6
17. Zheng Y, Yu X, Zeng J, Chen S (2012) Feasibility of filamentous fungi for biofuel production using hydrolysate from dilute sulfuric acid pretreatment of wheat straw. Biotechnol Biofuel 5:50. https://doi.org/10.1186/1754-6834-5-50
18. Skiba EA, Budavka VY, Ovchinnikova EV, Gladysheva EK, Kashcheyeva EI, Pavlov IN, Sakovich GV (2020) A technology for pilot production of bacterial cellulose from oat hulls. Chem Eng J 383:123128. https://doi.org/10.1016/j.cej.2019.123128
19. Zhu YJ, Zhuang XS (2012) Conceptual net energy output for biofuel production from lignocellulosic biomass through biofueling. Prog Energy Combust Sci 38:583–598. https://doi.org/10.1016/j.pecs.2012.03.007
20. de Oliveira JP, Bruni GP, Lima KO, El Halal SLM, da Rosa GS, da Dias ARG, Zavareze ER (2017) Cellulose fibers extracted from rice and oat husks and their application in hydrogel. Food Chem 221:153–160. https://doi.org/10.1016/j.foodchem.2016.10.048
21. Waliszewska B, Młeczek M, Zborowska M, Golinski P, Rutkowski P, Szentner K (2019) Changes in the chemical composition and the structure of cellulose and lignin in elm wood exposed to various forms of arsenic. Cellulose 26:6303–6315. https://doi.org/10.1007/s10570-019-02511-2
22. Haarmeyer CN, Smith MD, Chandawat SPS, Sammond D, Whitehead TA (2017) Insights into cellulase-lignin non-specific binding revealed by computational redesign of the surface of green fluorescent protein. Biotechnol Bioeng 114:740–750. https://doi.org/10.1002/bit.26201
23. Qin L, Li W-C, Liu L, Zhu J-Q, Li X, Li B-Z, Yuan Y-J (2016) Inhibition of lignin-derived phenolic compounds to cellulase. Biotechnol Biofuel 9:70. https://doi.org/10.1186/s13068-016-0485-2
24. Yan X, Cheng J-R, Wang Y-T, Zhu M-J (2020) Enhanced lignin removal and enzymolysis efficiency of grass waste by hydrogen peroxide synergized dilute alkali pretreatment. Biosour Technol 301:122756. https://doi.org/10.1016/j.biortech.2020.122756
25. Guragai VN, Bastola KP, Madl RL, Vadlani PV (2016) Novel biomass pretreatment using alkaline organic solvents: a green approach for biomass fractionation and 2,3-butanediol production. BioEnergy Res 9:643–655. https://doi.org/10.1007/s12151-015-9706-y
26. Qun Jiang L, Fang Z, Li XK, Luo J (2013) Production of 2,3-butanediol from cellulose and Jatropha hulls after ion liquid pretreatment and dilute-acid hydrolysis. AMB Express 3:1–8. https://doi.org/10.1186/2191-0855-3-48
27. Strzżyczkova P, Ház A, Burčová Z, Feranc J, Kreps F, Šurina I, Jablonský M (2019) Spruce bark-A source of polyphenolic compounds: optimizing the operating conditions of supercritical carbon dioxide extraction. Molecules 24:4049. https://doi.org/10.3390/molecules24224049
28. Zhang Y, Xia C, Lu M, Tu M (2018) Effect of overliming and activated carbon detoxification on inhibitors removal and butanol fermentation of poplar prehydrolysates. Biotechnol Biofuel 11:178. https://doi.org/10.1186/s13068-018-1182-0
29. Hazeena SH, Pandey A, Binod P (2016) Evaluation of oil palm front hydrolysate as a novel substrate for 2,3-butanediol production using a novel isolate Enterobacter cloacae SG1. Renew Energy 98:216–220. https://doi.org/10.1016/j.renene.2016.02.030
30. Keo-oudone C, Phommachan K, Sulyo I, Nurcholis M, Bouphanmy S, Kosaka T, Yamada M (2022) Highly efficient production of 2,3-butanediol from xylose and glucose by newly isolated thermostolerant Cronobacter sakazakii. BMC Microbiol 22:164. https://doi.org/10.1186/s12866-022-02577-z
31. Chen R (2015) A paradigm shift in biomass technology from complete to partial cellulose hydrolysis: lessons learned from nature. Bioengineered 6:69–72. https://doi.org/10.1080/21655979.2014.1004019
32. Okonkwo CC, Ujor VC, Mishra PK, Ezeji TC (2017) Process development for enhanced 2,3-butanediol production by Paeoniae cortex polysaccharides DSM 365. Fermentation. https://doi.org/10.3390/fermentation3020018
33. Amraoui Y, Nairsetty V, Coulon F, Agrawal D, Chandel AK, Maina S, Koutinas A, Kumar V (2021) Integrated fermentative production and downstream processing of 2,3-butanediol from sugarcane bagasse-derived xylose by mutant strain of Enterobacter ludvigii. ACS Sustain Chem Eng 9:10381–10391. https://doi.org/10.1021/acssuschemeng.1c03951
34. OHair J, Jin Q, Yu D, Wu J, Wang H, Zhou S, Huang H (2021) Non-sterile fermentation of food waste using thermophilic and alkaliphilic Bacillus licheniformis YNP-5TSU for 2,3-butanediol...
production. Waste Manag 120:248–256. https://doi.org/10.1016/j.wasman.2020.11.029
35 Yu D, O’Hair J, Poe N, Jin Q, Pinton S, He Y, Huang H (2022) Conversion of food waste into 2,3-butanediol via thermophilic fermentation: effects of carbohydrate content and nutrient supplementation. Foods. https://doi.org/10.3390/foods11020169
36. Amraoui Y, Prabhu AA, Narisetty V, Coulon F, Kumar Chandel A, Willoughby N, Jacob S, Koutinas A, Kumar V (2022) Enhanced 2,3-butanediol production by mutant Enterobacter ludwigi using Brewers’ spent grain hydrolysate: process optimization for a pragmatic biorefinery loom. Chem Eng J 427:130851. https://doi.org/10.1016/j.cej.2021.130851
37. Maina S, Schneider R, Alexandri M, Papapostolou H, Nychas G-J, Koutinas A, Venus J (2021) Volumetric oxygen transfer coefficient as fermentation control parameter to manipulate the production of either acetoin or D-2,3-butanediol using bakery waste. Biore- sour Technol 335:125155. https://doi.org/10.1016/j.biortech.2021.125155
38. Narisetty V, Zhang L, Zhang J, Sze Ki Lin C, Wah Tong Y, Loke Show P, Kant Bhatia S, Misra A, Kumar V (2022) Fermentative production of 2,3-Butanediol using bread waste—a green approach for sustainable management of food waste. Biore sour Technol 358:127381. https://doi.org/10.1016/j.biortech.2022.127381
39. Narisetty V, Amraoui Y, Abdullah A, Ahmad E, Agrawal D, Parameswaran B, Pandey A, Goel S, Kumar V (2021) High yield recovery of 2,3-butanediol from fermented broth accumulated on xylose rich sugarcane bagasse hydrolysate using aqueous two-phase extraction system. Biore sour Technol 337:125463. https://doi.org/10.1016/j.biortech.2021.125463
40. Gil A (2021) Current insights into lignocellulose related waste valorization. Chem Eng Adv 8:100186. https://doi.org/10.1016/j.cejadv.2021.100186
41. Scown CD, Baral NR, Yang M, Vora N, Huntington T (2021) Technoeconomic analysis for biofuels and bioproducts. Curr Opin Biotechnol 67:58–64. https://doi.org/10.1016/j.copbio.2021.01.002

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.