Emerging synergies on the co-treatment of spent coffee grounds and brewer's spent grains for ethanol production

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Abstract

**Purpose:** Given that the depletion of fossil fuels is an important issue that should be tackled urgently, alternative routes are being sought for biofuels production. This paper, under this context, studied the potential synergies of co-treatment of two kinds of agro-waste (brewers' spent grains, BSG and spent coffee grounds, SCG) towards the direction of biodiesel and bioethanol production, alleviating at the same time their treatment and management issues.

**Methods:** The technical feasibility of oil extraction and ethanol production was studied and optimized in both laboratory and bench scale. A treatment train including alkaline pretreatment of SCG and dilute acid pretreatment of BSG, neutralization of the solutions by mixing them, enzymatic saccharification and ethanolic fermentation led to the production of bioethanol.

**Results:** The co-treatment of BSG and SCG proved beneficial and optimum experimental conditions were defined by setting the bioethanol production cost as optimization parameter. The lowest ethanol production cost (3.9€/kg) was observed for scenario 4 (SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (400 μL/g cellulose, 16h)) and scenario 5 (SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (800 μL/g cellulose, 5h)). Further upscaling of the process verified the technical feasibility of process units leading to a further increase of ethanol yield and thus a decrease in the production cost.

**Conclusion:** Conclusively, it was demonstrated that the recovery and extraction of the oil content of BSG and SCG and the achievement of high ethanol yields is technically feasible. Within this context, an integrated biorefinery was presented, making evident that these substrates could reverse the eminent energy crisis via biofuels production.

**Statement Of Novelty**

The novelty of this work is the co-treatment of two agro-wastes (spent coffee grounds and brewers’ spent grains) towards biodiesel and bioethanol production. To our knowledge, the experimental set-up including the acidic pretreatment of BSG and the alkaline pretreatment of SCG followed by the enzymatic hydrolysis of the mixture was examined for the first time during this study. The main benefits of the co-treatment are the absence of a neutralisation stage avoiding the respective cost and the improved efficiencies of the bioprocesses. Based on the lab and bench scale experiments and a preliminary cost analysis presented, a biorefinery conceptual design is proposed where the synergistic effect of the co-treatment was made evident. Thus, this innovative process will allow to mitigate the adverse environmental impacts of fossil fuels without affecting with the entire food chain.

1. **Introduction**

In the concept (“reduce reuse recycle recover”) of circular economy and energy security, waste is considered as a valuable source for further exploitation. Current industrial activity generates a substantial amount of biomass bioproducts that are left unused, despite their multi-dimensional contribution potential to the development of value-added products.

Through the entire food supply chain, from harvesting and processing to the final consumable foodstuff, edible and inedible parts of the original organic materials are discarded. The agro-food industry itself is responsible for the 19% of the overall food waste generated. Approximately 17*10^6 tons of waste are accumulated from the processing and manufacturing of beverages and food products per year in the EU [1].

Taking into account the emerging demand for renewable energies on European level, agro-food wastes have been attracting much attention as potential biomass feedstock for bioenergy and biofuel production.

The EU statistical report 2020 determines the overall fuel demand in Europe to 405 Mtoe, implying that at least 56 Mtoe of energy should be supplied by biofuels to fulfil the EU commitment. Currently, blends of fossil fuels with liquid biofuels are used as renewable energy sources in transportation. Among the most widely produced biofuels stands biodiesel and
biogasoline, which in 2017 represented respectively 12 Mtoe and 2.5 Mtoe of produced energy in the EU [2]. Bioethanol is primarily produced by first generation feedstocks, based on sugars and starch, such as maize, wheat and sugarbeet. However, the competition with land, water and the human food supply that has arisen from their extended use, has rendered its continuing exploitation unsustainable. Alternative feedstocks based on lignocellulosic biomass (second generation feedstocks) have been attracting much attention as a viable substitute of food crops. Abundant in different forms (agricultural, wood, forest residues, industrial wastes, organic fraction of municipal solids wastes), produced in large quantities, easily accessible and available with promising compositional characteristics, when undergone specific processes can yield sufficient amounts of valuable components to be valorized into sustainable pathways [3].

The production of lignocellulosic ethanol is often reported in literature, based mainly on the valorization of single feedstocks, while mixed feedstocks have been poorly explored[4, 5]. Since the former is reliant upon the continuous availability and supply of the raw materials, its full-scale production is highly stimulated by extraneous factors. Such bottlenecks constitute the seasonal fluctuation of substrates influencing their availability, storage and transportation costs, intrinsic characteristics of the biomass demanding special handling or processing (low bulk density, recalcitrance) but also the competitive uses of the available feedstocks (soil enhancement, domestic fuel) [4].

The mixed biomass concept, by simultaneously utilizing two or more different types of biomass, of similar or different origin, in varying proportions, can provide an efficient alternative both from a practical and technical viewpoint [4]. The combination of feedstocks can outbound the insufficiencies of individual substrates, promote the synergetic supplementation of nutrients, minimize the need of external supplements addition, improve the quality of the overall feedstock (carbohydrates content, ash)[6], ensure the continuous availability of feedstock, thereby increasing the throughput of existed installations and their cost savings. While only a few laboratory-scale studies have investigated the feasibility of the mixed feedstock approach, benefits reported include: enhanced removal of lignin, lower concentration of inhibitors in the hydrolysate mixtures, increased sugar recovery in pretreated mixtures and occasionally higher ethanol yields achieved, compared to single substrate trials [7–9].

Given the high popularity of their marketable products, beer and coffee industries are two of the most active waste generators in the agro-food sector.

Beer brewing rejects different types of waste among which brewers’ spent grains (BSG) have the highest accumulation factor [10]. The overall brewing process includes several biochemical reactions (malting of barley grains, mashing, boiling, fermentation to ethanol), that break down the malt starch to fermentable sugars. BSG is the solid fraction separated from the wort after mashing and it represents 30% of the original malt weight. Approximately 20kg of BSG are produced per 100L of beer indicating that about 8*10^6 tons of BSG are produced per year only in the EU [11].

On the other hand, spent coffee grounds (SCG) are the primary residue generated from the brewing of coffee. They represent the 45% of the initial coffee bean, after it has been treated to release its constituents (aromas, flavors) to the consumable coffee liquid [12].

Agro-wastes from industrial food processing, have been reported to present fluctuating characteristics. BSG and SCG are high in organic content and other nutritional valuable compounds ranging from fibers and sugars to protein, vitamins, minerals and antioxidants, rendering them promising raw materials for food and non-food applications. The chemical composition of both BSG and SCG is directly linked to the initial barley grains and coffee beans respectively. They are principally lignocellulosic materials, with high fiber content in the form of cellulose (12-26% w/w for BSG and 7-15% w/w for SCG), hemicellulose (19-42% w/w for BSG and 28-42% w/w for SCG), lignin (7-28% w/w for BSG and up to 33% w/w for SCG), lipids (0-11% w/w for BSG and 2-24% w/w for SCG) and proteins (15-30% w/w for BSG and 6-18% w/w for SCG) [13–15].
Their composition varies with the grain or bean variety, harvesting time, brewing conditions applied or secondary raw materials involved in the process of brewing. Table 1 presents the variations in structural components reported in literature for both grain substrates.

Until now, spent grains have been used as animal feed given their high nutritional value (fibers, protein, vitamins and aminoacids). Given their increased production, incorporation to fertilizing products, composting or landfilling practices have also been reported as alternatives in handling the large volumes accumulated[12]. However, problems related to their disposal (biological instablity, odors generation, autoxidation susceptibility, spoilage and potential toxicity of several constituents) have led the updated environmental legislation, to introduce sustainable management practices aiming to prevent waste generation and minimize their environmental footprint[12]. By reclaiming the compositional interest of the recovered biomass streams, valuable compounds can be extracted thereof or directly be transformed to new marketable products and bioenergy. In recent years, extended research has been dedicated to either BSG or SCG streams as potential feedstocks for biodiesel and bioethanol production.

While the biodiesel potential of agro-food wastes is often mentioned in literature, few studies have experimentally proven their relevant suitability. SCG has been more extensively investigated with a few studies reporting successful oil extraction and biodiesel production thereof [16, 17], whereas only one recent study reveals the feasibility for BSG [18].

The bioethanol route involves an alternated sequence of processes including pretreatment of the lignocellulosic biomass, hydrolysis of polysaccharides to their monomer sugars (mainly glucose, xylose, arabinose) and fermentation. The final product is received by distillation of the fermented liquor. BSG and SCG hydrolysates have been widely investigated in literature as for their bioethanol potential with variations regarding the pretreatment conditions, the types and dosage of saccharifying enzymes for hydrolysis as well as the fermenting agents applied. Pretreatment is an essential step in the bioconversion route, that degrades the recalcitrant lignocellulosic fibers and facilitates the subsequent step of enzymatic hydrolysis. Among the pretreatment techniques most commonly reported for BSG and SCG (alkaline, acidic, steam explosion, ionic liquid [19, 20] alkaline pretreatment appears more effective in conditioning the lignocellulosic biomass and partially decrystallizing cellulose, thereby increasing the accessible surface area for hydrolytic enzymes [21], whereas acid pretreatment has also been reported as a fast and efficient pretreatment method in some cases [22]. Regarding fermentation of both substrates, an investigation of a variety of yeasts (Saccharomyces cerevisiae, Pichia stipites, Kluyveromyces fragilis) [23] has outlined S. cerevisiae as the strain providing generally the highest ethanol yields.

According to our previous work SCG and BSG has the potential to be valorised for bioediesel and bioethanol production [20, 24, 25]. It has been proved that this is feasible when chemical pretreatment is applied; alkaline pretreatment for SCG and dilute acid for BCG. However, the high amount of chemicals used for both the chemical pretreatment and neutralization prior to enzymatic hydrolysis render the whole process problematic in economic and resource-efficient terms. Thus, this study aims to investigate possible synergies in the co-treatment of SCG and BSG towards a sustainable biodiesel and bioethanol production.

2. Materials And Methods

2.1 Feedstocks

Brewers’ Spent Grains (BSG) were provided by the Hellenic Breweries of Atalanti (EZA), located in Atalanti, Fthiotida province, Greece. The BSG were received in the premises of UEST (Unit of Environmental Science and Technology, School of Chemical Engineering, National Technical University of Athens, Greece), and were subjected to further processing. Processing involved simultaneous milling and dehydration using GAIA GC-100, a rotary drum dryer suitable for food waste, which operated for 14 hours at 105°C. The substrate was dried up to 3.3% moisture, was physico-chemically characterized and stored at room temperature prior to further processing.
Spent Coffee Grounds (SCG) samples were periodically collected from the coffee-shop of the School of Chemical Engineering, in the National Technical University of Athens, Greece, over a period of 3 months. Each batch was dried for 24h at 105°C to minimize the occurrence of microbial growth. All samples were physico-chemically characterized and stored at room temperature prior to further processing.

2.2 Experimental set-up

The oil extraction, chemical pretreatment, enzymatic saccharification and fermentation were the main processes that were studied in laboratory and bench scale. Lab scale experiments were carried out in autoclavable bottles (200 mL) using an Incubator Shaker (IKA-KS 3000i), while bench scale experimenting in a 4L mechanically stirred reactor with double walls.

2.3 Oil extraction

For laboratory scale experiments, the conventional Soxhlet extraction process was adopted and operated for 4 hours with n-hexane as solvent at a solid to solvent ratio 1:20 (g/mL), based on previous studies optimizing the process [26].

While Soxhlet is a widely accepted and efficient method for extraction of oils from a range of materials, its long-time operation needs, special equipment and energy requirements render its application in industrial scale less viable. In this framework, liquid solid extraction was investigated as an alternative and more easily applicable method for bench scale experimenting. N-hexane 95% was used as solvent. More specifically, separate batches of BSG and SCG were mixed with n-hexane, at a solid-to-solvent ratio of 1:10, loaded into the 4L reactor, with continuous mechanical stirring at 30°C for 4 hours. The insoluble matrix was separated from the solute oil enriched extract with the use of conventional sieve and was dried at 105°C for 24h so as to remove the residual hexane from the grains. A BÜCHI Rotavapor - R KRvr 65/45, rotary evaporator was used to recover the oil from which the residual moisture and solvent were removed at 105°C for 24h. Dried defatted solids served as reference feedstock for further experimental trials.

2.4 Chemical pretreatment

Alkaline pretreatment with NaOH at 50°C was chosen as pretreatment method for SCG and dilute acid pretreatment (dilute H₂SO₄) was applied for BSG samples at 50°C in line with our previous work [20, 24]. In both cases, the severity of pretreatment (concentration of chemicals (0.5, 1N), pretreatment duration (5h, 16h)) was studied. All experiments were performed thrice.

2.5 Enzymatic Hydrolysis

After the oil extraction and chemical pretreatment of separate batches of BSG and SCG, 50:50 (w/w) mixtures of BSG and SCG were prepare aiming to reach the optimum pH for the enzyme operation without extra addition of chemicals. Enzymatic hydrolysis of defatted and chemically pretreated BSG, SCG or their mixtures was carried out at 50°C by adding the cellulase CellicCTec2 (Novozymes, Denmark), which contained 2620 β-glucosidase U/mL and 120 FPU/mL. Acid (H₂SO₄) or alkaline (NaOH) solutions were used to set the pH at 5.0-5.5, when necessary. The effect of enzyme dosing (400 and 800μL/g cellulose) and time duration (5h, 16h) was studied. All experiments were performed thrice.

2.6 Ethanol fermentation
Saccharified hydrolysates were directly subjected to fermentation by 2\% w/w Saccharomyces cerevisiae (baker’s yeast) at 32°C for 24h. When the fermentation process was completed, slurries were separated to their solid and liquid fractions by centrifugation and were subjected to physico-chemical characterization. Ethanol concentrations were determined in the liquid phase so as to estimate the ethanol yield. The latter was calculated as (Equation 1):

\[
\eta_{eth}(\%) = \frac{\text{Ethanol produced (g)}}{\text{Theoretical ethanol maximum}} \times 100\%
\]

where

\[
\text{Theoretical ethanol maximum} = \text{mass of sample (g)} \times \text{percentage of cellulose} (\% w/w) \times 1.11 \times 0.51
\]

taking into account the water hydrolysis (factor 1.11) and the stoichiometric glucose to ethanol conversion during fermentation (factor 0.51, the remainder transforms to CO₂).

### 2.7 Analytical methods

Characterization of raw materials, pretreated and fermented substrates was carried out in accordance with the NREL laboratory analytical protocol by Sluiter et al. [27]. Prior to any quantitative determination, the samples received were centrifuged for 10 min at 3000 rpm (IEC/Centra CL2). In the solid fraction, total solids, water soluble and volatile solids, lignin (soluble and insoluble), cellulose, hemicellulose and starch were determined. Glucose, total reduced sugars, total organic carbon (TOC), total nitrogen (TN), volatile fatty acids (VFA) as degradation by products and ethanol were assessed in the liquid fraction. Glucose measurement was carried out using a commercial kit (Biosis SA) through the Glucose oxidase-peroxidase method (GOD/PAP). Ethanol concentration was estimated by Ethanol Assay Kit (K-EtoHLOQR, Megazymes). Starch was determined as total starch, using AOAC Method 996.11 Total Starch Assay Kit (K-TSTA-100A, Megazymes) and it was measured as glucose released after digestion with thermostable a-amylase and amyloglucosidase. Phenolic compounds were measured by the kit Spectroquant Phenol Test 100856 and VFA quantification was carried out by the Spectroquant Volatile Organic Acids Test kit (Merck Millipore). Total reducing sugars were determined by the 3,5-dinitrosalicylic acid method [28]. Samples assessment was held in duplicate.

In addition, total organic carbon (TOC) was determined by the Solid Sample Module (SSM-5000A) of the SHIMADZU TOC-VCSH (Total organic carbon analyzer), nitrogen (TN) by the KJELDAHERM block digestion unit and the Gerhardt - Vapodest 30s device and pH by the Mettler-Toledo MPC227 pH/Conductivity Meter.

### 3. Results And Discussion

#### 3.1 Composition of BSG and SCG

BSG and SCG are lignocellulosic substrates and are mainly constituted by lignin, cellulose, hemicellulose, proteins and lipids. Their composition varies according to the variety of beans, the harvesting time but also process parameters involved in the brewing of beer or coffee. Table 1 summarizes the composition of the two substrates investigated in this work along with the respective values from literature. The composition of the BSG falls in the ranges determined by Rojas-Chamoro et al., Kemppainen et al. [29, 30], while Mussato et al. [31] have determined higher lignin content up to 27.8\%. Regarding SCG, similar composition for cellulose has been reported by Mussatto et al. [32] and Cruz Lopes [33] but with lower hemicellulose fraction (36-39\%) and lignin below 10\%. Higher lignin has been observed in SCG by Kwon et. [34] and Ballesteros et al. [35] which ranged between 23-29\%. It must be noted that approximately the 53\% of the BSG and the 59\% of the SCG (dry basis) consists of carbohydrates with the capacity to be degraded into their monomers and undergo fermentation, while their oil content, especially for SCG indicates their capacity for oil extraction and subsequent biodiesel production.

| Table 1. Composition of BSG, SCG substrates used |
3.2 Oil extraction

The oil recovery of BSG and SCG substrates was evaluated in laboratory scale experiments with the standard Soxhlet method. The ranges of oil recovered were 6.0% ± 0.8% for BSG and 16.9% ± 0.9% for SCG on a dry basis. Oil content for BSG falls into the range of literature references (Table 1), while SCG content is among the highest reported. Faulds et al. [37] reported about 5% lipids in BSG while Kemppainen et al. [30] observed oil contents up to 11.4%. Literature findings for SCG vary from 2.5-6% [35, 38], but also up to 10-15% [39], depending on the coffee variety and brewing process.

3.3 Chemical Pretreatment

Aiming to enlighten the impact of chemical pretreatments on the structure of agro-waste (alkaline pretreatment on SCG and acid pretreatment on BSG), the chemically pretreated substrates were physico-chemically analysed. Table 2 presents the composition of the liquid and solid phase of alkaline pretreated SCG and acid pretreated BSG.

| SCG NaOH (N) | Time (h) | Glucose | TRS | VFA | Phenolic compounds | TOC | TN | Cellulose | Hemicellulose | Starch | AIL | ASL |
|--------------|----------|---------|-----|-----|-------------------|-----|----|-----------|-------------|--------|-----|-----|
| 0.5          | 5        | 9.5±0.7 | 23.5±5.7 | 111.7±10.4 | 3.4±0.9 | 166.0±10.8 | 9.7±2.3 | 12.5±0.3 | 40.7±1.5 | 0.04±0.03 | 7.4±2.3 | 1.0±0.2 |
| 1            | 5        | 12.3±0.5 | 19.8±2.4 | 131.6±11.2 | 3.2±0.7 | 133.4±11.6 | 8.3±1.7 | 12.8±1.3 | 33.9±1.2 | 0.07±0.02 | 6.6±1.3 | 0.9±0.1 |
| 0.5          | 16       | 8.1±0.3 | 19.6±3.1 | 121.4±9.7 | 3.1±0.4 | 141.6±14.0 | 9.7±1.9 | 15.8±1.7 | 49.1±1.9 | 0.05±0.01 | 21.9±2.0 | 1.1±0.3 |

| BSG H₂SO₄ (N) | Time (h) | Glucose | TRS | VFA | Phenolic compounds | TOC | TN | Cellulose | Hemicellulose | Starch | AIL | ASL |
|---------------|----------|---------|-----|-----|-------------------|-----|----|-----------|-------------|--------|-----|-----|
| 0.5           | 5        | 0.9±0.05 | 1.3±0.1 | 15.9±1.7 | 0.4±0.03 | 57.4±1.9 | 6.9±0.9 | 12.5±1.0 | 6.6±2.3 | 0.7±0.3 | 42.0±1.9 | 1.4±0.2 |
| 1             | 5        | 1.8±0.03 | 25.8±1.7 | 86.2±2.4 | 0.3±0.02 | 32.9±1.1 | 1.0±0.05 | 2.3±0.3 | 0.2±0.06 | 0.0±0.01 | 49.1±2.1 | 10.5±0.03 |
| 0.5           | 16       | 0.9±0.02 | 54.6±2.4 | 22.3±1.8 | 0.5±0.02 | 61.8±2.2 | 2.6±0.8 | 20.0±1.6 | 10.8±1.3 | 0.1±0.01 | 30.3±1.6 | 2.5±0.2 |

During alkaline pretreatment of SCG, the solid degradation ranged from 19.2 to 22.9%, presenting a positive effect for both parameters under investigation. It was revealed that alkaline pretreatment resulted in high lignin degradation (53.7-71.2%) with moderate degradation of cellulose and hemicellulose. Similarly, Girotto et al. [40], who pretreated SCG under different NaOH loadings only achieved a 4.8% hemicellulose degradation with 7.5% NaOH, but similar hemicellulose degradations with 2% w/w NaOH which increased as the NaOH concentration increased. They also reported increased lignin removal with increased NaOH concentration when pretreating SCG.
Delignification by-products (phenolic compounds and volatile fatty acids) were also present in all prehydrolysates, as a side effect of the pretreatment method and could be attributed to the lignin degradation. Under alkaline conditions, phenolic compounds from the degradation of lignin and volatile fatty acids (VFA) from the hydrolyzed acetyl groups of hemicellulose are released [41]. Such compounds have been reported to act inhibitory to downstream biochemical processes, causing deterrence of cellulosic enzymes and inhibition of fermenting microorganisms [42, 43]. A pretreatment method is efficient when it accomplishes high delignification and simultaneous alleviation of inhibition problems. Regarding the utilized yeast S. cerevisiae, it has been reported that VFA concentration above 6g/L can inhibit ethanol fermentation, causing up to 100% inhibition for 10g/L (inhibition accessed as reduction in cell mass) [44]. Similarly, according to Colombi et al. [45], phenolic compounds at a concentration between 0.1-1g/L can be toxic for S. cerevisiae causing up to 46% drop in ethanol yield. Yang et al. (2016) also reported that 1.5g/L of phenols can cause 5.4% decrease in ethanol yield, while higher inhibitory levels are reported by Gu et al. (2019) (3-5g/L). Phenolic compounds and VFA concentrations fluctuate according to the degradation of lignin and hemicellulose respectively, in accordance with previous researches [23]. Similar concentrations of degradation products in the liquid phase have also been studied by Passadis et al. [24], although the concentration of glucose reported was lower (1.8–4.9 mg/g SCG). Increase of the pretreatment time did not particularly differentiate polysaccharides degradation but slightly increased the inhibitors concentrations in pretreated liquors. The phenolic compounds detected after alkaline pretreatment of SCG were below 300mg/L and thus below the ones reported by Mussato et al. and M. Fernandez-Delgado et al. [31, 46], probably due to the lower delignification percentages achieved. Hence, they are not expected to induce severe inhibition to microbial growth. On the other hand, high VFA concentrations were detected in individual SCG (10.5± 0.9g/L) prehydrolysates, that fluctuate within the potentially toxic levels for S. cerevisiae.

Dilute acid pretreatment on BSG, on the other hand, did not affect the lignin structure, while cellulose and hemicellulose degradation reached 89% and 99% respectively when 1N H_2SO_4 was used. In other words, elevated concentration of H_2SO_4, from 0.5N to 1N, while it degraded more acid soluble lignin and hemicellulose, it was linked with significantly higher accumulation of VFA compared to the other acid pretreatment levels examined. The high VFA concentrations that were detected in individual BSG (8.3± 0.8g/L) prehydrolysates could be considered toxic for S. cerevisiae. In general, the degradation results are in accordance with our previous study [20] where low solubilization of BSG, low lignin degradation and high polysaccharides degradation were observed when acid pretreatment was applied on BSG. Similar results have also been reported elsewhere in literature. Rojas-Chamorro et al. [29] indicated that pretreatment of BSG with H_2SO_4 resulted in high hemicellulose degradation between 70-89.4% and lower toxic compounds concentration. Dimos et al. [47] and Carvalheiro et al. [48] have also concluded that the dilute acid pretreatment results in the hemicellulose hydrolysis. Despite the high carbohydrates degradation, free sugars were not released in the liquid phase, implying that the depolymerization of carbohydrates was partial and did not result in monomeric sugars.

### 3.4 Ethanol production

BSG, SCG and their mixtures were examined as for their bioethanol potential via a biochemical conversion route. Table 3 presents the characteristics of the liquid phase of the fermentation broth and Table 4 presents the efficiencies achieved in terms of structural components degradation and ethanol yield.

**Table 3. Characteristics of liquid phase of fermentation broth**
### Table 4. Degradation efficiencies of structural components and ethanol yields achieved

| Substrate | Pretreatment | Enzymatic Hydrolysis | % degradation | Ethanol Yield (%) |
|-----------|--------------|----------------------|---------------|------------------|
|           | Concentration (N) | Time (h) | Loading (µL/g cellulose) | Time (h) | TS | Cellulose | Hemicellulose | Starch | AIL | ASL |               |
| SCG       | 0.5          | 5       | 400 | 5 | 16.4±0.7 | 15.3±0.3 | 14.9±2.7 | 45.1±2.2 | 30.7±2.2 | 68.4±2.2 | 19.9±1.8 |
| BSG       | 0.5          | 5       | 400 | 5 | 15.1±0.7 | 21.3±1.1 | 27.8±2.5 | 87.9±7.7 | 9.4±1.3 | 4.2±2.1 | 39.5±2.3 |
| SCG/BSG   | 0.5          | 5       | 400 | 5 | 27.2±0.9 | 41.3±2.8 | 39.1±5.6 | 23.5±1.7 | 2.0±0.1 | 20.5±1.7 | 31.4±1.4 |
| SCG/BSG   | 0.5          | 16      | 400 | 5 | 14.1±0.3 | 28.8±2.2 | 61.3±4.8 | 48.5±1.7 | 7.3±1.3 | 1.2±1.5 | 51.6±2.1 |
| SCG/BSG   | 0.5          | 5       | 400 | 16 | 26.9±1.2 | 24.8±2.2 | 33.3±3.2 | 95.1±2.1 | 3.5±0.9 | 1.7±0.1 | 44.3±1.1 |
| SCG/BSG   | 0.5          | 5       | 800 | 5 | 26.3±2.1 | 35.5±1.9 | 49.0±2.7 | 92.8±2.1 | 0.8±0.1 | 17.8±1.5 | 57.3±2.7 |
| SCG/BSG   | 1           | 5       | 400 | 5 | 23.7±2.3 | 24.7±2.0 | 46.8±3.7 | 95.0±2.0 | 11.5±1.1 | 16.2±1.1 | 10.0±1.9 |

The alkaline pretreatment, enzymatic hydrolysis and ethanolic fermentation of SCG achieved moderate degradation degrees of cellulose (15.3±0.3%), and were followed by low ethanol yields (19.9±1.8%). In literature, defatted SCG have been utilised for bioethanol production providing a range of operational configurations and conditions and thus a range of efficiencies [25]. Alkaline pretreatment is often suggested as the most suitable pretreatment method, given the milder operational conditions, the lower environmental impact, the lower corrosivity of NaOH solutions and the higher delignification yields [24].

Although in literature high ethanol yields for BSG have been reported [49], they significantly vary depending on the processes applied. In general, it is well established that harsh pretreatment conditions favour ethanol production [20], but at the same time they induce severe cost and safety issues.

From Table 4, it is evident that in almost all cases the application of mixtures of BSG and SCG provided higher ethanol yields (40.0-57.3%) than each substrate alone (19.9% for SCG and 39.5% for BSG). Furthermore, the degradation of starch is almost complete and the degradation of hemicellulose is significantly increased. Thus, the synergistic effect of co-treatment is obvious.

Several references in literature report better performance of mixtures, produced from different combinations of biomass feedstocks against plain batches, after fermentation with different microorganisms [4], although no paper has been
published using mixtures of BSG and SCG as substrates. Chen et al. [50] processed corn stover and corn at a 1:1 ratio with acid pretreatment and fermentation with Z. mobilis at 30% solid loading, and achieved up to 94.1g/L ethanol (74.2% theoretical yield), against 0.89 g/L for corn stover alone. Erdei et al. [9], after simultaneous fermentation with ordinary baker’s yeast of separately pretreated mixtures of wheat straw and wheat meal, reported an increased theoretical ethanol yield (up to 99%) compared to those obtained by either of the substrates alone (68% and 91% respectively). In another study, Nguyen et al. [5] subjected different mixtures of coffee bean husks, cassava stems and coconut coir to simultaneous saccharification and fermentation and detected up to 12 g/L ethanol after 24h (47% ethanol theoretical yield). Mixtures of ricotta whey and sugarcane bagasse produced 49.7g/L ethanol after fermentation with Kluyveromyces marxianus, which was higher than the ones produced from sugarcane bagasse alone [7]. Moreover, acid pretreated residues from sugarcane industry (straw, bagasse, tops) when were studied in mixtures as for their bioethanol potential achieved up to 25% higher ethanol yield and 55% higher enzymatic conversion than bagasse alone [8]. Also, Imamoglu et al. [51] by combinedly pretreating mixtures of rice hulls (70%) and cotton stalks (30%) at different ratios and by subsequent fermentation with E. coli produced 20.7 g/L ethanol compared to 7.2g/L for corn stalks alone. All the references above resulted in comparable or higher ethanol yields than the present study.

3.5 Selection of conversion pathway based on ethanol production cost

As optimization parameter of the bioethanol production process, the bioethanol production cost was set, since the ethanol yields should be considered taking into account the consumption of energy and additives.

According to Ge et al. [52], the cellulosic ethanol production cost is the sum of 5 parameters: water, energy, enzymes bacteria and chemicals. The total bioethanol production cost includes a fixed fraction associated with labor, maintenance, insurance etc. and a variable fraction related to additives consumption. The latter could be described as (Equation 2):

\[ C_{production} = C_{Energy} + C_{Water} + C_{Chemical} + C_{Enzyme} + C_{Bacteria} \]  

Equation 2

The energy cost includes two aspects: the respective cost of the chemical pretreatment and of the enzymatic hydrolysis and fermentation processes. Mafe et al. [53] developed a model in order to estimate the energy demands during the acidic pretreatment of lignocellulosic biomass. It was proved that the highest energy requirements are related to biomass heating prior to its pretreatment, while the energy demands for reactions and maintenance could be neglected. Staying in line with this approach, the energy needs were estimated by the following equation (Equation 3):

\[ Q_{heating} = m_w \int_{T_0}^{T_H} C_p dT + m_b \int_{T_0}^{T_H} C_p dT + m_a C_p (T_H - T_0) \]  

Equation 3

\( C_p \) is the specific heat capacity (cal/kg/°C); \( m_w, m_b, m_a \) is the mass (kg) of biomass, water and chemicals respectively; \( T_H \) is the processes’ temperature (°C) and \( T_0 \) is the ambient temperature of 25 °C.

Given the narrow range of the operational temperature (20 °C -50 °C), the specific heat capacities of water and feedstocks were considered constant. The SCG and BSG specific heat capacities were measured according to the ASTM D2766 – 95.

The heat loss during the process was calculated using the following equation (Equation 4) in accordance with Mafe et al. approach [53].

\[ Q_{loss} = 0.05 \times \left( \frac{T_{maintenance}}{T_{heating}} \right) \times Q_{heating} \]  

Equation 4

where that \( T_{maintenance} \) is the residence time for the process and \( T_{heating} \) is the heating up time.
Water consumption rate is a function of the selected solid loading in the process. Chemical compounds are consumed during the pretreatment process (alkaline and acidic) and as additives for pH correction prior to enzymatic hydrolysis. The price of chemicals depends on the pretreatment type. A cellulolytic enzymatic formulation is utilised for the enzymatic hydrolysis and bacteria/yeast for ethanolic fermentation. Their unit prices are functions of saccharification and fermentation operational conditions.

Table 5. Values adopted for the construction of the model for bioethanol production cost

| Parameter | Value |
|-----------|-------|
| $C_{p,SCG}$ | 0.63 cal/g/°C |
| $C_{p,BSG}$ | 0.83 cal/g/°C |
| $C_{p,Water}$ | 1.00 cal/g/°C |
| $C_{p,NaOH}$ | 0.35 cal/g/°C |
| $C_{p,H2SO4}$ | 0.22 cal/g/°C |

Unit prices

| Chemical | Price |
|---------|-------|
| NaOH | 0.35 €/kg |
| $H_2SO_4$ | 0.25 €/kg |
| Water | 0.83 €/m$^3$ |
| CellicCTec2 | 1.00 €/L |
| Yeast | 1.20 €/kg |
| Energy | 0.0647 €/kWh |

According to the experimental results, the mass flows were set up and the ethanol production cost for each case was estimated. Table 6 presents the estimated ethanol production costs for the different scenarios. Scenario 1 considered separated and independent production line for each substrate, while scenarios 2 to 6 separate oil extraction and chemical pretreatment and subsequent treatment as a mixture.

Table 6. Estimated ethanol production costs for the different mixtures of BSG and SCG

| Scenario | Substrate | Chemical Pretreatment | Enzymatic Hydrolysis | Ethanol production cost (€/kg) |
|----------|-----------|-----------------------|----------------------|-------------------------------|
| 1        | SCG       | 0.5                   | 5                    | 9.31                          |
|          | BSG       | 0.5                   | 5                    | 7.32                          |
| 2        | SCG/BSG   | 0.5                   | 5                    | 4.20                          |
| 3        | SCG/BSG   | 0.5                   | 16                   | 3.88                          |
| 4        | SCG/BSG   | 0.5                   | 5                    | 7.15                          |
| 5        | SCG/BSG   | 0.5                   | 5                    | 3.89                          |
| 6        | SCG/BSG   | 1                     | 5                    | 3.89                          |

Fig. 1 presents the ethanol production cost for each experimental set-up along with the respective break down of the cost.

From Fig. 1, it is evident that the contribution of water cost is very low for all cases since it accounts for just 3-4%, while the yeast contributes 10-12% to the total cost. On the contrary, energy, chemicals and enzyme cost are the major contributors in the final ethanol cost, as was anticipated. Scenario 1 presents the highest cost and this fact validates the advantage of the co-treatment of SCG with BSG. In this scenario, the cost of chemicals was over 50%, since chemicals are used for both the pretreatment and the pH regulation before enzymatic hydrolysis. Scenarios 4 (SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (400 μL/g cellulose, 16h)) and 5 (SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (800 μL/g cellulose, 5h)) led to the lowest ethanol production cost (3.9€/kg).
3.6 Sensitivity analysis

In order to determine the critical factors in the present study, the effect of the assumed unit prices on the optimum results was studied. The factors that were considered are: (1) NaOH price, (2) H₂SO₄ price, (3) water price, (4) enzyme price, (5) yeast price and (6) energy price. They were altered by ±20% to assess the impact.

In Fig. 2, it is obvious that the cost of bioethanol has non-linear correlations with the six factors. The minimum unit bioethanol cost is noted when the enzyme cost is decreased by 20% under the operational conditions of Scenario 5, that includes pretreatment with chemicals concentration 0.5N for 5h and enzymatic hydrolysis with elevated enzyme loading (800μL/g cellulose) for 5h. It is noted that for scenario 5, enzyme cost has the highest impact on ethanol production cost, while the rest of the parameters (apart from water) present the same weighted impact. Enzyme cost is also the most significant parameter when studying Scenario 4, but its impact is closer to the other parameters.

3.7 Bench scale

Applications of mixed lignocellulosic biomass have shown that combined substrates can achieve comparable or even greater fermentation efficiencies than single substrates. However, these results are based on laboratory evaluations and ethanol production from lignocellulosic mixtures is still an underdeveloped practice. Thus, the optimum scenarios were up-scaled from laboratory conditions to bench scale.

By solid-liquid extraction, the oil recovered amounted to 4.4% ±0.8% for BSG and to 10.2% ±0.6% for SCG. In other words, the oil content recovery efficiencies were 73.6% and 60.2% for BSG and SCG respectively. These efficiencies could be assumed satisfactory and thus, the scheme examined could be incorporated within a biorefinery that aims at the production of biodiesel. In the study of Mallen and Najdanovic-Visak [18], the oil extracted from BSG was converted by the use of sulfuric acid to biodiesel with an efficiency of 83.3% through a single-step transesterification, while the transesterification yields that have been reported in literature for SCG oil were higher than 56% and in some cases they even reach 100%. Such yields are comparable with those that have been reported for other vegetable oils [25].

The defatted substrates were treated according to the operational conditions of Scenarios 4 (SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (400 μL/g cellulose, 16h)) and 5 (SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (800 μL/g cellulose, 5h)) in the 4L mechanically agitated reactors. The ethanol yields achieved reached up to 79.7% and 84.7% respectively.

Hence, the upscaling led to a significant increase of ethanol yield in both cases. This fact can be mainly attributed to the mechanical stirring, which in turn led to a more efficient action and diffusion of chemicals and enzymes. The increased ethanol yields, as expected, led to a further decrease of ethanol production cost reaching 2.13€/kg and 2.63€/kg for scenarios 4 and 5, respectively.

Thus, an integrated multi-feedstock biorefinery that utilizes both SCG and BSG in combination as feedstocks producing both used oil suitable for biodiesel production and bioethanol could stand as a promising alternative. Fig. 3 presents the basic mass flows of the proposed biorefinery applying the operational conditions of Scenario 4.

4. Conclusions And Further Perspective

SCG and BSG, biowaste streams from beer and coffee industries, are feedstocks with high perspectives for valorization given their abundance, the simplicity of their collection network (point sources) and their composition. In this context, a multi-feedstock biorefinery that would use and upgrade these feedstocks to added value products was set as main goal of this study. In this study, the technical feasibility of oil extraction and ethanol production was validated and optimized in both laboratory and bench scale experiments. The ethanol production process unit included alkaline pretreatment of SCG and
dilute acid pretreatment of BSG, neutralization of the chemically pretreated substrates by mixing them, enzymatic hydrolysis and ethanolic fermentation. The co-treatment of the two substrates proved to be advantageous compared with the independent parallel treatments, given the savings from the neutralization chemicals and possible emerging synergies during bioprocessing. The production cost of ethanol was calculated and set as a driver for the choice of the operational conditions. Scenarios 4 (SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (400 μL/g cellulose, 16h)) and 5 (SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (800 μL/g cellulose, 5h)) led to the lowest ethanol production cost (3.9€/kg). From the sensitivity analysis, it was noted that the bioethanol unit price has non-linear correlations with the unit prices of the additives (chemicals, enzymes, yeast), with the enzyme cost being the most significant parameter. The upscaling of the process proved the technical feasibility of oil extraction with high efficiencies and led to an increase of ethanol yield, with subsequent decrease of its production cost.

Thus, the perspective of a multi-feedstock biorefinery that would valorise SCG and BSG producing oil and bioethanol seems to be promising. Nevertheless, the ethanol production cost is still elevated if the market price of cellulosic ethanol is taken into account. Therefore, in future work, this study could be extended to take into account technical issues such as minimization and/or recycling of enzymes, increase of solids loading and valorization of the fermentation broth via anaerobic digestion. In terms of sustainability, carbon and water footprint, energy integration, recycle of waste should also be considered. The outmost goal could be set as the design of a sustainable supply chain for the multi-feedstock biorefinery that could be simultaneously optimized in terms of its economic and sustainability performance.

Declarations

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Figures

Figure 1

Scenario 1: SCG (NaOH pretreatment (0.5N, 5h), enzymatic hydrolysis (400 μL/g cellulose, 5h)) BSG (H2SO4 pretreatment (0.5N, 5h), enzymatic hydrolysis (400 μL/g cellulose, 5h)) Scenario 2: SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (400 μL/g cellulose, 5h) Scenario 3: SCG/BSG mixture, chemical pretreatment (0.5N, 16h), enzymatic hydrolysis (400 μL/g cellulose, 5h) Scenario 4: SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (400 μL/g cellulose, 16h) Scenario 5: SCG/BSG mixture, chemical pretreatment (0.5N, 5h), enzymatic hydrolysis (800 μL/g cellulose, 5h) Scenario 6: SCG/BSG mixture, chemical pretreatment (1N, 5h), enzymatic hydrolysis (400 μL/g cellulose, 5h)
Figure 2
Sensitivity analysis results

Figure 3
Integrated multi-feedstock biorefinery

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