Production of green juice with an intensive thermo-mechanical fractionation process. Part I: Effects of processing conditions on the dewatering kinetics

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PRODUCTION OF GREEN JUICE WITH AN INTENSIVE THERMO-MECHANICAL FRACTIONATION PROCESS. PART I: EFFECTS OF PROCESSING CONDITIONS ON THE DEWATERING KINETICS

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Abstract

The thermally assisted mechanical dewatering (TAMD) process is a new intensive solid/liquid separation device. When applied to ‘nature-wet’ biomass, the TAMD process significantly enhances the separation yield. The TAMD process couples in one stage a mechanical dewatering at low pressure ($P_{\text{applied}} = 300 \text{ kPa}$ in the present study) with a moderated heating ($T_{\text{wall}} \leq 90^\circ\text{C}$). An increment of pressure can be applied in a second stage
to further enhance the dry solid content of the press cake. In the present study, the TAMD process was used to dewater spinach leaves and alfalfa stems and leaves. The influence of cutting, pulping and temperature on the fractionation kinetics and the extraction yield were specifically investigated. Experiments were carried out on a laboratory compression cell, heated through the piston. Results shows that, at ambient temperature, pulping is an essential pre-processing unit operation to reach an extraction yield of 55%. Under moderate heating conditions (T_{wall} = 50 or 70°C), a thermally assisted mechanical dewatering, without any pre-processing stage, can remove 69% of the inherent water from alfalfa, that is to say an increase by 23% of the yield. But, compared with the conventional fractionation process, the duration of the mechanical fractionation must be at least twice longer. Beyond 70°C, the temperature does not have any influence on the extraction kinetics. Cutting has a very limited influence with an enhancement of the dry solid content from 2 to 5% at best.

Keywords: Pressing, Heating, Biomass, Press-cake, Maceration, Leaves.

1. Introduction

Sustainable development requires the use of renewables as alternative feedstocks for chemical, fuel and material productions. Share of renewable raw materials in the US feedstock consumption runs to approximately 5% in the chemical industry and is expected to increase notably and to reach 25% by 2030 [1]. The same tendency is impelled by the European governments, but targets were exclusively defined for energy purposes [2]. To face this challenge, the concept of biorefinery was set up [3]. The underlying idea is to maximize the value derived from renewable feedstocks by producing a variety of chemicals, fuels and intermediates or end-products, using well integrated, environment- and resource-friendly technologies [4]. To guarantee an optimal exploitation of the various components in biomass, processing routes are defined according to the organic compositions of the raw materials.
Taking this into account, five biorefinery systems are commonly considered [5]: whole crop, green, oilseed, forest-based and, more recently, waste biorefineries. Systems based on agricultural crops raised some social and ecological criticisms recently. To avoid any competition with food, the ideal feedstock is nowadays considered to be residual by-products accumulated in abundance and with low or no profit and nutritional value [6]. These includes classically food and feed crop residues, wood residues, herbaceous plants, grasses, leaves and stems. The latter are still hardly used in spite of their large availability, their high biomass profit per hectare and their good coupling with agricultural production.

These ‘nature-wet’ raw materials require a fast primary processing or the use of preservation methods, like silage or drying. But due to the energy consumption of thermal dryer, the biomass is seldom transformed after drying, so that mechanical fractionation of the wet material is usually the first unit operation in a green biorefinery plant. The vegetative biomass is separated into a fiber-free extract containing the protoplasmic fluid - the green juice - and a fiber-rich solid fraction - the press cake. Both fractions have an economic value. The green juice is a raw material for high quality fodder proteins, cosmetic proteins, human nutrition or platform chemicals like lactic acid and lysine [7, 8, 9, 10, 11] or can be used as substrate for biogas production [12]. The press cake can be used as fodder pellets after thermal drying [13], as silage feed [11] or as solid fuel [14]. Conventional technologies for juice extraction involve mechanical pulping to disintegrate the cell walls followed by pressing [15], possibly with multistage addition of water and expression following the first pressing [16]. Most of the time, pressing is carried out with a screw press to provide additional maceration of the cell walls in complement of the action of the pressure. For vegetative biomass like alfalfa, clover and grass, screw presses remove approximately 55-60% of the inherent liquid [11, 15].

To date, the economic viability of green biomass fractionation process, for instance the ProXan-Process [17] used to produce a marketed leaf protein concentrate, depends, to a large
extent, on the utilization of the solid fraction as a high quality ruminant feed [15, 18]. The emergence of biorefineries, with further alternative processing routes especially for the nutrient-rich liquid fraction, can make the wet fractionation process more profitable [11]. Consequently, there is a lot of research at present into methods that enhance the dewatering ability of conventional mechanical processes. Intensification of mechanical dewatering processes can take several forms: simultaneous application of a pulsed electric field [19, 20], superimposition of ultrasounds [21, 22] or with heat supply [23]. The underlying objectives are to obtain a juice in larger quantity [24] and of better quality. In the same time, other works relate to the processes of fractionation of the juice. The green juice is usually subjected to a differential coagulation to get the protein-rich chloroplast fraction, containing almost all the extracted lipids and lipid-associated pigments, and the cytoplasmic fraction containing the soluble proteins, like the Rubisco, with very small amount of lipids and pentosans. Conventional techniques include differential fractionation of the extracts by heat, centrifugation or pH adjustment, or with solvents, polyelectrolytes or other reagents, or by a combination of some of these techniques. Until now, there is no simple and economical method for isolating large quantities of plant enzymes even if nanofiltration and ultrafiltration [25] or expanded-bed chromatography [26, 27] seem to produce good performances on a laboratory scale.

At all events, the initial fractionation of the biomass remains an essential operation for biorefinery process. For a few years now, we specifically investigate a thermally assisted mechanical dewatering (TAMD) process [28], which couples a mechanical dewatering with a simultaneous moderate heating of the walls of the apparatus in contact with the product. Thermal mechanical dewatering method is not a brand new technology [23, 29-31] but the operating conditions of the TAMD process and its use for the mechanical fractionation of the herbaceous biomass are quite innovating. The compression is carried out in two step: a first
compression stage at low pressure (usually 300 kPa) and a second stage around 1500 kPa. The TAMD process was used to effectively dewater alfalfa biomass under a variety of processing conditions ranging from 21 to 90°C and from 300 to 3000 kPa [24]. It has been illustrated that it can remove up to 83% of the inherent liquid fraction from alfalfa under moderate processing conditions, which is a much higher extraction yield than in conventional mechanical dewatering process. The TAMD process is thus likely to extract a green juice in larger quantity than with conventional fractionation processes. Now, the operating conditions of the TAMD process, especially the influence of the temperature, on the extraction kinetics and the green juice quality has to be investigated, in order to evaluate its potential of valorization and optimize the TAMD process accordingly. This paper is the first of a series of two papers, which explores the TAMD biomass fractionation procedure when applied to typical ‘nature-wet’ biomass, addressing the following questions:

1. To what extent does the mechanical fractionation with the TAMD process result in a separation among a fiber-free liquid extract and a fiber-rich solid fraction?

2. Is pulping necessary to enhance the amount of green juice recovered?

3. Compared to conventional mechanical fractionation procedure, what is the efficiency of the TAMD process?

After a short description of the experimental device and the materials used for the mechanical fractionation experiments in section 2, the influence of a prior cutting and of the heating conditions are analyzed in section 3. The efficiency of the TAMD process is assessed in section 4 and compared to the one of a conventional mechanical fractionation process. Finally, the optimal conditions for the separation are given as concluding remarks.
2. Materials and methods

2.1 Experimental set up

At the laboratory scale, the experimental set up consists of a compression cell inserted in a CARVER® hydraulic press (Carver Inc., Wabash, United State), which has a maximum pressing capacity of 14800 kPa and provides the pressure required to progress a downward expression.

The cell (Figure 1) consists of a compressive piston, a hollowed cylindrical vessel and a filter medium. The filtration chamber has a diameter of 148 mm and a maximum height of 60 mm. In spite of its low mechanical resistance, Teflon™ was selected as constitutive material of the vessel walls to minimize the frictions with the piston. Consequently, a stainless steel external jacket is added to ensure the mechanical resistance of the unit. The cell is fitted with a planar medium, made of Teflon™, of 165.2 cm² area. To investigate the influence of a heat supply, three electric resistances are inserted in the upper part of the compressive piston, made of copper to reduce thermal inertia. Each resistor can supply up to 350W. Three thermocouples, including the one for the temperature regulation of the copper block, are introduced into the piston. The accuracy of these sensors with theirs acquisition line is estimated at ±0.03°C.
2.2 Feed materials

The composition of the liquid fraction is influenced by a multiplicity of factors operating during the whole process, beginning with the agronomic factors (plant species, spacing, fertilizers, climate…) and the pre-pulp conditions (maturity of the vegetation, harvesting and post-harvesting environment…). Moreover, it is well-known that the chloroplasts contribute to a large extent to the total leaf protein (up to 75% for photosynthetically active leaves) and that the Rubisco, which is located within the chloroplasts, is the most abundant soluble protein in photosynthetic organs. This enzyme catalyses the first step of the metabolic pathway found in the stroma of the chloroplast in which carbon enters in the form of CO$_2$ and leaves in the form of sugar. Partially purified powder of Rubisco from spinach leaves, available in large amounts as a lyophilised powder with stable activity, can be bought from the chemical trade. Thus, to develop the analytical methods for the green juice characterization [32] and to investigate the influence of the TAMD fractionation procedure on the green juice composition, we selected spinach as a reference biomass. Alfalfa was selected as a second feed material according to its industrial use [17].
2.2.1 Spinach

Fresh winter spinaches (*Spinacia oleracea*) were bought in a small independent store, having a short circuit of provisioning. Fresh spinaches in bulk were preferred to packaged one, whose water content is higher because of a pre-wash before packaging. The fresh leaves were stored in a tight container at 4°C to avoid the desiccation of the plant, in accordance with the storage protocol already developed [24]. After 7 days, the dry solid content of the biomass remained unchanged and, visually, neither yellowing nor fading were observed. From the beginning of February to mid-April, six batches of spinach leaves were bought. The wet-basis moisture content of the leaves ranged between 88.3 and 90.9%, according to the batch. Dewatering experiments were carried out in the week following the purchase. About 240g of fresh spinach leaves were introduced into the TMAD process. In a same batch, the size of the leaves was very variable, from 5 to 12 cm long and from 4 to 10 cm broad. To investigate the influence of the size on the fractionation kinetics, three physical “structures” were considered: the whole leaves, the leaves chopped into pieces of 3×3 cm and the leaves chopped into 0.5 cm broad bands.

2.2.2 Alfalfa

Alfalfa, also known as Lucerne, is a perennial legume, in the pea family Fabaceae, widely grown throughout the world as forage for cattle and harvested as hay. Alfalfa has the highest feeding value of all common hay crops, with high protein content and highly digestible fiber. Grown on well-drained soils with neutral pH, total yield is commonly around 8 tons per hectare but can reach up to 15-20 tons per hectare. The plant is very resilient to droughts, thanks to its deep root system. Moreover, its root nodules contain bacteria with the ability to fix nitrogen. In most climates, alfalfa is cut 3 to 4 times a year. Alfalfa (*Medicago sativa L.*) was field-chopped at the pre-blossoming stage. Approximately 300 to 500 mm long fresh stems with leaves were sampled at weekly intervals between mid-May and mid-June 2010. The
fresh alfalfa was transported to the laboratory and stored at 4°C in a tight container within 1h after the harvest. The wet basis moisture content of the fresh alfalfa ranged between 78.4% and 82.6%, according to the week of sampling. Dewatering experiments were carried out in the three days following the harvest. About 200g of fresh alfalfa were chopped into 5-cm pieces before introduction into the TAMD process.

2.3 Experimental procedures

Previous studies [24, 33] emphasized that the dewatering enhancement for biomass results only from thermal effects. As a consequence, the influence of the pressure applied was not investigated in the present study. The mechanical fractionation protocol was as follows. The biomass was introduced at room temperature into the TAMD device. At t=0, a pressure of 300kPa was applied through the compressive piston, previously heated at the selected operating temperature. Progressively, the biomass was separated into a green juice and a press cake. The filtrate recovered in the collector flowed out into a container laid on the computer interfaced balance. The applied pressure, the piston temperatures, the temperature of the face of the cake in contact with the filter medium and the mass of filtrate were recorded at set time intervals of 1 s.

At the end of the experiment, the press cake was weighed. Its dry solid content, defined as the weight of the dry sample divided by the weight of the wet sample times 100, was measured according to the AFNOR standard procedure N° X31-505. This procedure recommended a drying at 105°C for 24 hours. To check that the mass balance was preserved during the experiment, the dry solid content of the initial biomass was determined using the same protocol and the moisture content of the press cake at the end of the experiment was calculated knowing the mass of the juice fraction and compared to the experimental value. The dry solid content, S, and the dry basis moisture content, W, were calculated according to the following equations:
Observations were carried out to highlight the hydric state of the cells, before and right after the TAMD process has been applied. The plant tissues were fixed with 2.5% glutaraldehyde (25%) in a 0.05M sodium-cacodylate buffer for an hour. After rinsing and ambient-air drying, the samples were cut into 30μm-thick sections with a microtome (MICROM HM 255). Microscopic observations of the samples were then carried out thanks to the optic microscope LEICA DMRB whose magnifying power goes from x100 to x630. The microscope is equipped with a black and white camera (MICAM VHR 1000) linked to a television set that transmits the pictures to an acquisition software (Studio Version 8).

3. Results and discussion

3.1 Influence of the operating parameters on the TAMD kinetics

3.1.1 Evaluation of the process repeatability and estimate of the measure accuracy

Dewatering experiments were repeated at least 4 times with the same process operating conditions to evaluate the repeatability of the results. Dry solid contents achieved at the end of a dewatering experiment performed with a wall temperature of 50°C or 70°C are reported in Table 1 for spinach leaves chopped into 0.5 cm broad bands and alfalfa chopped into 5-cm pieces. The maximum standard deviation on the dry solid content of the dewatered biomass is ±0.8% for spinach and ±1% for alfalfa. This uncertainty is in agreement with former results [24]. As can be seen in Table 1, the dispersion introduced by the batch number is more important than that associated with the sampling and the repeatability of the experiment, that can be estimated ±0.55% for alfalfa.
Table 1 - Influence of sampling and processing repeatability on the initial and final dry solid contents of the biomass

The fractionation kinetics of spinach leaves measured for a wall temperature of 70°C are plotted in Figure 2. In spite of the heterogeneity of material (in a given batch) and the various batches, the kinetics are appreciably the same even if, around 750 seconds, the difference in the initial rate of fractionation, highlighted on Figure 2, results in a significant dispersion of the mass of filtrate (±22g). The dispersion decreases with time (±6.1g after 5000s).
3.1.2 Influence of the cutting on the fractionation kinetics

Three cuttings were considered for spinach leaves:
- none (whole leaves);
- coarse (leaves chopped into pieces of 3×3 cm);
- and fine (leaves chopped into 0.5 cm broad bands).

For each cutting, the fractionation experiments were performed for four wall temperatures, namely 30, 50, 70 and 90°C. Whatever the piston temperature used, the tendency is similar. For example, the average fractionation kinetics obtained with a wall temperature of 70°C for the three available cuttings are given on Figure 3. To reduce the measurement uncertainty, the feed material for all the experiments was sampled in the same batch. Thus, according to the process repeatability, the maximum standard deviation on the mass of filtrate is ±3g. The mass of green juice recovered during the mechanical fractionation increases with the degree.
of cutting, especially when the biomass is cut finely. This result was expected. Indeed, the cutting breaks some cell walls and plasma membranes, releasing the contents of the cytoplasm and more precisely the cytosol, as well as some tonoplasts, releasing the contents of the vacuole [34, 35]. The cytosol is the portion of a plant cell that is not enclosed within membrane-bound organelles. It is composed of water, salts and organic molecules. The vacuole is a membrane-bound organelle, which could be represented as an enclosed compartment, filled with water containing inorganic and organic molecules. Thus, the pre-processing treatment by cutting makes the liquid fractions in the cell plant more accessible and facilitates the dewatering.

![Figure 3](image)

*Figure 3 - Influence of cutting on the mechanical fractionation kinetics of spinach leaves (T_piston = 70°C).*

The influence of cutting on the press cake dry solid content is reported in Table 2. According to the measurement uncertainty, the changes observed with a wall temperature set to 30°C are not significant, whatever the degree of cutting. Furthermore, for all the wall temperatures investigated, a coarse cutting does not change much the dry solid content of the press cake compared to the value achieved with whole leaves. When the biomass is cut more finely, the
dry solid content increases more significantly. At best, an increase in the dry solid content from 2 to 5% can be expected. For a biomass with relatively large leaves, like spinaches, the rupture of the cell walls facilitates the mechanical fractionation of the material: the amount of green juice recovered is larger and consequently the press cake is drier.

![Table 2 - Influence of the cutting and the wall temperature on the final dry solid content (in %) of spinach leaves](image)

<table>
<thead>
<tr>
<th></th>
<th>None</th>
<th>Coarse</th>
<th>Fine</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_{wall} = 30^\circ C$</td>
<td>11.5</td>
<td>11.2</td>
<td>11.9</td>
</tr>
<tr>
<td>$T_{wall} = 50^\circ C$</td>
<td>19.6</td>
<td>21.6</td>
<td>23.2</td>
</tr>
<tr>
<td>$T_{wall} = 70^\circ C$</td>
<td>23.1</td>
<td>23.8</td>
<td>25.3</td>
</tr>
<tr>
<td>$T_{wall} = 90^\circ C$</td>
<td>31.7</td>
<td>31.9</td>
<td>37.6</td>
</tr>
</tbody>
</table>

1

### 3.1.3 Influence of heating on the fractionation kinetics

The influence of heating on the dry solid content of the spinach press cake can be analyzed through the values reported in Table 2. Whatever the degree of cutting, an increase in the piston temperature significantly enhances the dry solid content of the press cake compared to a conventional dewatering at room temperature. At 30°C, pressing enhances the dry solid content of spinach cake from 9% to 12%. The dry solid content reaches 25.3% for a wall temperature of 70°C and even 37.6% for a wall temperature of 90°C. Thus, the effect of the wall temperature is much more important than the degree of cutting. As can be seen in Figure 4, the dry solid content enhancement, defined as the difference between the dry solid content of the press cake after three hours of processing and the initial dry solid content of the biomass, is linearly proportional to the set temperature ($R^2=0.94$).
Figure 4 - Influence of the wall temperature on the dry solid content enhancement (compared to the initial dryness) after three hours of processing.

The average fractionation kinetics measured for the four wall temperatures, namely 30, 50, 70 and 90°C, are plotted in Figure 5 (a). Results emphasize that the higher the temperature is, the larger the recovered mass of green juice is (as previously mentioned) and the faster the dewatering is, even if the kinetics of fractionation observed with a wall temperature of 70°C or 90°C are practically identical. The same behaviours are observed with alfalfa (see Figure 5 (b) for the kinetics and Table 3 for the influence of the heating on the dry solid content). The amount of green juice produced is at least five times larger with a heat supply through the piston wall than with a conventional pressing at ambient temperature and the separation is faster.
Figure 5 - Influence of the wall temperature on the mechanical fractionation kinetics of

spinach leaves (a) or alfalfa (b)
Compared to the maximum amount available with spinach, the mass of green juice is a little lower but fresh alfalfa was initially drier than spinach leaves. This result is in agreement with the work of Savoie and Beauregard [36], which emphasized that the proportion of green juice increases (from 10 to 21%) when the initial wet basis moisture content increases (from 80 to 84%). For both materials, the difference between the fractionation kinetics obtained with a wall temperature of 50°C or 70°C lies primarily in the duration of separation, the total amount of juice recovered being identical according to the measurement repeatability.

<table>
<thead>
<tr>
<th>Initial dry solid content (%)</th>
<th>Final dry solid content (%)</th>
<th>Dry solid content variation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_{\text{wall}} = 30^\circ\text{C}$</td>
<td>17.1</td>
<td>19.4</td>
</tr>
<tr>
<td>$T_{\text{wall}} = 50^\circ\text{C}$</td>
<td>17.8</td>
<td>34.3</td>
</tr>
<tr>
<td>$T_{\text{wall}} = 70^\circ\text{C}$</td>
<td>17.8</td>
<td>36.8</td>
</tr>
</tbody>
</table>

*Table 3 - Influence of the wall temperature on the variation of the dry solid content variation for alfalfa*

One possible explanation of the role of temperature lies in the phenomenon of osmosis. Osmosis is the movement of water molecules across a partially permeable membrane, here the plasma membranes and the tonoplasts, down a water concentration gradient. This is the principal mechanism by which water is transported into or out of plant cells. The osmotic pressure of a dilute solution is conventionally calculated by the van’t Hoff equation [37]. This formula connects the osmotic pressure to the product of the molarity of the solute, the dimensionless van’t Hoff factor, the gas constant and the absolute temperature. Thus, a difference in temperature between two solutions on either side of a semi-permeable membrane will result in an osmotic gradient and a flow of water, from the colder towards the warmer area, until the balance in osmotic pressure is reached. As can be seen on Figure 6, this
osmoregulation causes the plasma membranes to peel away from the cell walls, a process known as plasmolysis.

\[\text{Figure 6 - Photograph of the vegetable cells after fractionation in the TAMD process, x400}\]

In experiments, such local temperature gradients cannot be highlighted. Only the temperature gradient at the press cake boundaries can be measured. To follow the rise in temperature of the press cake, a thin temperature sensor was inserted and placed on the top of the filter media. The variation of the temperature over time at the base of the alfalfa press cake, dewatered with a piston temperature set to 55°C, is plotted in Figure 7. The temperature of the bottom part of the press cake increases to a maximum value of 50°C after two hours and a half. It is also noticed that, in the same time, the green juice mass tends towards an asymptote. With a wall temperature of 70°C, the increase in temperature is faster and the bottom part of the press cake reaches 50°C after 15 minutes and the maximum temperature of 66°C after only 20 minutes. This information is also important for the juice quality. Indeed, some of its components, like the green “hydrophobic” proteins, have a low resistance to thermal treatments [38].
3.2 Assessment of the TAMD process efficiency

The optimal operating conditions for an efficient separation known, the performances of the TAMD process could be compared to those of the conventional process. Traditionally, juice extraction involves mechanical pulping followed by pressing. At the lab scale, the chops were macerated in a blender. 200gr of pulp was then introduced in the compression cell and dewatered at 300 kPa and ambient temperature.

3.2.1 Fractionation kinetics with a conventional process

The fractionation kinetics of the minced and chopped alfalfa, dewatered at ambient temperature, are reported in Figure 8. As expected, pulping affects the amount of green juice extracted from alfalfa: the mass is enhanced by 717% after one hour of processing. Moreover, the juice flows out almost immediately: for instance, 85% of the juice is recovered after 15 minutes. The duration of this dwell time is in accordance with former results [39] and common practices.

Figure 7 - Evolution of the press cake temperature and green juice mass during the mechanical fractionation of alfalfa (T$_{piston}$ = 55°C)
3.2.2 Comparison of the extraction yields

To assess the performances of the TAMD process, alfalfa cut in 5cm pieces was dewatered with the TAMD process, whose piston temperature was heated to 70°C. The corresponding fractionation kinetic is plotted in Figure 8. The maximum amount of green juice extracted with these operating conditions is higher than that produced with the conventional process even if the initial extraction velocity is slower. Nevertheless, after 15 minutes of processing, the recovered mass of green juice is identical. To compare the results, an extraction yield, defined as the ratio of the mass of green juice produced after one hour of pressing to the mass of the liquid intrinsically introduced, is reported in Table 4. As expected, the extraction yield for a pressing, preceded by a mechanical pulping, is close to 55%. For the TAMD process operating at 300 kPa with a piston temperature set to 70°C, the extraction yield amounts to 69%, that is to say an increase of 23%. For the TAMD process operating at 300 kPa during one hour with a piston temperature set to 50°C, the extraction yield is 53%. If the dewatering time is increased to one hour and a half, the extraction yield rises to 69%. Indeed, the mass of green juice produced is roughly the same as shown in Figure 5. For a wall temperature lower
than 70°C, the time of dehydration must be significantly longer than that conventionally used to obtain a larger quantity of juice.

Table 4 - Extraction yield for pressing at ambient temperature, for pulping prior to pressing at ambient temperature and for the TAMD process set to 70°C

<table>
<thead>
<tr>
<th></th>
<th>Extraction yield at 1 h (%)</th>
<th>Final solid content of the cake after 1h (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pressing at 30°C</td>
<td>7.8</td>
<td>18.3</td>
</tr>
<tr>
<td>Pulping prior to pressing at 30°C</td>
<td>55.9</td>
<td>31.8</td>
</tr>
<tr>
<td>TAMD process with T_{wall}=70°C</td>
<td>69.0</td>
<td>41.1</td>
</tr>
</tbody>
</table>

4. Summary

The TAMD process was used to efficiently dewater ‘nature-wet’ biomass like spinach leaves and alfalfa stems and leaves. The influence of cutting, pulping and heating conditions on the dewatering kinetics and the extraction yield were investigated. The influence of the pressure applied was not considered in the present paper. It has been illustrated that a mechanical pulping, performed at the lab scale using a blender, has a much greater influence both on the maximum amount of green juice extracted and on the extraction velocity than cutting. Only a relatively fine cutting (0.5 cm broad bands in the present study) increases significantly the quantity of green juice recovered. As a result, the dry solid content is enhanced from 2 to 5% at best, for an applied pressure of 300 kPa. The heating conditions have much more influence on the extraction. For an applied pressure of 300 kPa, the TAMD process can be used to remove 69% of the inherent water from alfalfa under moderate heating conditions, the enhancement being of 55% in a conventional dewatering process with prior pulping. With a piston temperature set to 70°C, the duration of the mechanical fractionation must be at least 30 minutes. If the piston temperature is decreased to 50°C, a longer pressing time, at least one
hour and a half, is required. Beyond 70°C, the temperature does not have any influence on the extraction kinetics. The temperature gradient in the press cake is supposed to generate a difference in osmotic pressure in the biomass, which induces a flow of green juice through the membrane walls, until the balance in osmotic pressure is reached. This assumption will have to be confirmed by future work. The first question, which arises at this stage of the process design, relates to the heat supply on the composition of the green juice produced. Indeed, the face of the press cake in contact with the filtering medium increases rapidly to 50°C and it is well-known that some of the proteins precipitate at low temperature. The second question concerns the specific energy consumption of the TAMD process and the cost of the separation. These two items will be discussed in future publications.

Acknowledgments

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