Extraction of laminarin from *Saccharina latissima* seaweed using cross-flow filtration

Martin Sterner · Fredrik Gröndahl

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**Abstract**

Laminarin is a low-molecular-weight polysaccharide found in seaweed (kelp), often in equal concentrations to that in the commercially important hydrocolloid alginate. However, while alginate can be easily recovered by dissolution followed by acid precipitation, for laminarin, there is no such straightforward way of recovering it. Laminarin can be used as dietary fiber and, if efficiently extracted, it may be used for functional food/feed applications and as a component in plant defense stimulants for agriculture. One way of concentrating laminarin from dilute solutions is to press the solution through ultrafine membranes that the molecules cannot pass through. When alginate is extracted, an acid pretreatment step is used and the dilute acid residue from that process also contains laminarin. We used cross-flow filtration to concentrate laminarin from *Saccharina latissima*, retrieving it from the dilute acid solution of the acid pretreatment of an alginate extraction. Three ceramic membranes with 5, 15, and 50 kDa molecular weight cutoffs were used, and the pressure, temperature, and feed velocity were altered to reveal which parameters controlled the flow through the membrane and how efficiently laminarin was concentrated. The effects on laminarin extraction for fresh vs. frozen biomass were evaluated showing that frozen biomass releases more laminarin with a similar biomass homogenization technique. Thermal and microbial degradation of the feed components was studied during the course of the filtrations, showing that microbial degradation can affect the laminarin concentration, while the temperature of the process ~ 65 °C had little impact on laminarin. The techniques used to monitor the components in the feed and permeate during filtration were nuclear magnetic resonance, $^1$H-NMR, and size exclusion chromatography. The filtrations were performed in a pilot-size filtration unit with ceramic membranes (ZrO$_2$/TiO$_2$, TiO$_2$/Al$_2$O support, 0.08 m$^2$). To be able to operate without quick membrane fouling, the most important parameter was to have a high liquid velocity over the membrane, 4.7 m s$^{-1}$. A good technique to concentrate laminarin was to prefilter it through a 50-kDa membrane using 2 bar liquid pressure and to concentrate it over a 5-kDa membrane using 5-bar liquid pressure. With these settings, the liquid flux through the filter became 60–80 and 30–40 L m$^{-2}$ h$^{-1}$ over the 50-kDa and 5-kDa membrane.

**Keywords** Cross-flow filtration · Laminarin · Mannitol · Saccharina latissima · Kelp · Extraction

**Introduction**

Mariculture of seaweed (kelp) has the potential to generate large volumes of new biomass. Seaweed cultivation at sea requires no fertilization, irrigation, or maintenance during its growth, and the production can give a high yield per cultivated area in a highly sustainable way (Thomas et al. 2020).

One species of seaweed that is suitable for farming in cold saline water is *Saccharina latissima*, a brown alga in the order Laminariales. It grows at a rapid pace and thrives on the Swedish west coast where it is cultivated but also grows wild. Seaweed cultivation in Sweden has been developed in two research projects called “Seafarm” and “Agri Aqua Kelp” during 2014–2020, but from 2020 on, seaweed farming has become commercial via a startup company (KosterAlg). An average yearly harvest of 19 t ha$^{-1}$ is achieved for the KosterAlg cultivation (Hasselström et al. 2020). It has also been estimated that it is possible to sequester more CO$_2$ in the seaweed than the total CO$_2$ equivalents required for cultivation and preservation (Thomas et al. 2020).

Since *S. latissima* contains more than one utilizable component, it can be advantageous to find ways to extract these
components in sequence and enable the use of a greater share of the biomass (Sterner and Edlund 2016; Sterner et al. 2017).

The main components in *S. latissima* are laminarin ~ 7–40%, alginate ~ 15–35%, mannitol ~ 10–15%, protein ~ 4–13%, and cellulose ~ 5–15% (Obluchinskaya 2008; Jard et al. 2013; Schiener et al. 2015; Vilg et al. 2015; Sterner and Edlund 2016; Sterner et al. 2017; Stévant et al. 2017). The share of algal components varies between years and seasons and by the local growth conditions.

Laminarin is the primary energy reserve of the seaweed that is built up during spring and summer to be spent during the winter (Schiener et al. 2015; Vilg et al. 2015). It is a low-molecular-weight (Mw) polysaccharide that usually has a Mw of 3–5 kDa and consists of glucose units joined mainly by 1,3-beta bonds alternated with a low amount of 1,6-beta bonds, with some 1,6-beta bound branches of single glucose units (Rinaudo 2007). Laminarin can be used as dietary fiber, as a stimulant of the innate immune system of mammals, and as a plant pathogen defense stimulant, the first application, since it is not digested by the human small intestine enzymes and the two later applications due to the resemblance of the laminarin glucose 1,3-beta-bond motif to that of certain fungal polysaccharides to which plants and mammals respond. (Yvin et al. 1998; Klarzynski et al. 2000; Aziz et al. 2003; Brown and Williams 2009; Myklestad and Granum 2009; McDonnell et al. 2010). Alginate is a high-Mw linear charged polysaccharide consisting of two alternating uronic acids, mannuronic acid, and guluronic acid (Rinaudo 2007). It is extracted for its use as a viscosity modifier, gel material, or texturizer in a vast range of products, including food cosmetics and medical products (Draget et al. 2005; Rinaudo 2014; Porse and Rudolph 2017). Mannitol is a sugar alcohol that regulates the osmotic balance in seaweed and can be used as a sweetener similar to xylitol and sorbitol (Pawl 2014).

While alginate can be solubilized in the extraction process and later retrieved again by precipitation with acid, this is not possible for laminarin (Black et al. 1951; McHugh 2003). Laminarin is soluble in water at all pH values, and it cannot be precipitated without the addition of a high amount of precipitant, such as ethanol or acetone at more than 60% by weight (Black et al. 1951; Sterner and Edlund 2016). A common technique used to extract alginate is to preheat the seaweed with dilute acid to leach out calcium ions, which makes it easier to dissolve the alginate at a later stage (McHugh 2003). When using acid pretreatment in the alginate extraction process, if laminarin follows the dilute acid leaching solution, this may allow a combined process to extract both compounds and possibly make the extraction of laminarin easy enough to enable commercial applications.

A way to concentrate laminarin is to use membrane filtration that can separate laminarin according to the size of the molecule (Yvin et al. 1998). One such technique is cross-flow filtration, in which the solution to be filtered (feed) flows at high velocity over the membrane, so that small molecules follow through the membrane with the solution as it goes through (permeate), and the constant high flow hinders the buildup of a filter cake that limits the flux through the membrane (Bhave 1996). For ultrafiltration, the thin layer of ultrafine particles and large molecules that precipitate in a gel-like manner close to the membrane is referred to as “the gel layer,” but more generally, it is called the “concentration polarization layer” (Bhave 1996; Ripperger et al. 2013).

To have a successful cross-flow filtration, a setting needs to be found to maximize the liquid volume going through the membrane per unit of time (flux), not only at the start of the filtration but also in a longer time perspective by minimizing the gradual clogging of the membrane (membrane fouling).

If the solution is pressed through the membrane under high pressure, the initial flux is increased, but this triggers the formation of a thicker gel layer and, eventually, more membrane fouling, which soon decreases the flux again. To decrease the thickness of the gel layer, the feed velocity can be increased, but this requires a more powerful pump and generates a greater pressure drop in the membrane filter channels, which allows shorter filters in the filtration setup.

Choosing a membrane with a higher Mw cutoff and thus larger pores allows for a greater flux, but it cannot allow a great share of the targeted compound to pass through the filter (passage ratio). The passage ratio, which is the share of the targeted compound that goes through the filter, is the opposite of the retention ratio or retention, which is the share of the targeted compound that does not go through the filter. The real membrane cutoff also depends on the thickness and nature of the formed gel layer, and therefore, the cutoff also depends on the feed velocity and pressure in the filtration (Ripperger et al. 2013).

The general theories on cross-flow filtration indicate which factors can affect the performance, but in the end, every kind of feed is unique, and it is hard to predict, for instance, the permeability of the gel layer and the progression of membrane fouling without real testing. Cross-flow filtration has been used as a method for the separation of laminarin before (Yvin et al. 1998), but we found no study that has examined which filtration settings should be used.

The aim of this study was to investigate which cross-flow filtration settings, in the concentration of laminarin from the solution from acid pretreatment of homogenized *S. latissima*, that lead to fast filtration with high yields. The explored parameters were the feed velocity and pressure over the membranes, choice of membrane Mw cutoff, time of filtration, temperature, and effect of prefiltering with a membrane with a higher Mw cutoff. The major part of the study was on cross-flow filtration, but a minor part covered homogenization and extraction, answering the questions regarding whether freezing the biomass affected the homogenization, if longer extraction times allowed more laminarin to be released from...
homogenized seaweed and if the studied homogenization technique leaves seaweed in a state suitable for alginate extraction.

The gained knowledge may increase the utility of *S. latissima* biomass by providing efficient ways to recover its second-most abundant component, laminarin.

**Materials and methods**

**Materials**

The bulk cultivated *Saccharina latissima* seaweed was harvested on July 17, 2018, from a cultivation site close to the Sven Lovén Center for Marine Science (University of Gothenburg) on the Swedish west coast. The seaweed was frozen, kept at $-20 \, ^\circ\text{C}$ and transported to KTH (Royal Institute of Technology in Stockholm) (Scheme 1, step 1). To pool the material, 450-kg frozen seaweed was sawed into smaller pieces, blended thoroughly in a big plastic container, and put in smaller bags of approximately 2.5 kg each (Scheme 1, step 2). A smaller part of seaweed was harvested fresh close to the Sven Lovén Center on May 2, 2019.

Alginic acid from *Macrocystis pyrifera*, laminarin from *Laminaria digitata*, sodium citrate dihydrate ACS reagent $\geq 99\%$, sodium hydroxide $\geq 97\%$ ACS reagent, and hydrochloric acid 37% ACS reagent were from Sigma Aldrich. D-(-)-Mannitol purum was from KEBO. Deuterium oxide ($\text{D}_2\text{O}$) (DLM-4-100) 99.9% was from Cambridge Isotope Laboratories, Inc., and $\text{D}_2\text{O}$ (#87156.0100) 99.96% was from VWR. Ethanol 99.6% (v/v) was from VWR.

**Processes**

**Pre-process**

Approximately 5 kg of frozen mixed and pooled seaweed was thawed in the plastic bags and mixed in a food processor of the type BOSCH MultiTalent 8 in batches of 300 g of seaweed at a time, with 1.2 L added deionized water to each, for 5 min (Scheme 1, step 3). For every two runs with the food processor, the mixed seaweed was moved to a beaker for further homogenization. The mixed seaweed in the beaker, which had a volume of approximately 3 L, was processed by an Ultra-Turrax (X-1020 Intern. Laborat. App. Dottingen, GmbH), with its tip centrally 5 cm from the bottom in the beaker and run for 10 min (Scheme 1, step 4). All homogenized seaweed from the Ultra-Turrax treatment was poured into a bigger container, with a volume of $\sim 30 \, \text{L}$, in which it was mixed with a propeller blade of 10-cm diameter placed close to the bottom of the container via stirring at 300 rpm for 4 h (Scheme 1, step 5a). Part of the slurry containing homogenized seaweed was weighed on a Petri dish with a known weight, dried in a desiccator, and weighed again to determine the dry weight of each batch of processed seaweed. Part of the slurry was centrifuged for 5 min at $3850 \times g$, and the supernatant was saved for analysis. In the next step, 800 mL HCl (1 M) was added to the container, and it was stirred for 10 min more at 300 rpm, giving a final pH of approximately 2.65 (Scheme 1, step 5b). The mixed and acidified algal slurry was poured onto a nylon mesh attached over a box with a length of 45 cm and a width of 37 cm in a hanging fashion and left for 1 h to let the liquid drip through the mesh (Scheme 1, step 6). The nylon mesh was from Derma AB and had 71 $\mu\text{m}$ aperture width, 45 $\mu\text{m}$ thread diameter, and 37% open screening area. The filter cake containing seaweed that did not go through the mesh, which typically totaled 8 kg, was saved and used for alginate extraction, and approximately 620 mL of NaOH (1 M) was used to neutralize it (Scheme 1, step 7). The permeate that did go through the mesh (typically 18 kg) was used for cross-flow filtration, and approximately 180 mL NaOH (1 M) was used to neutralize it (Scheme 1, step 8).

**Small-scale laminarin and alginate extractions using fresh, frozen, and freeze-dried material**

To test the impact of frozen vs. nonfrozen material, small-scale experiments were performed on algal material that was...
freshly harvested, cut in pieces with scissors, and mixed, with half of it frozen and the other half processed fresh. These seaweeds were mixed in a food processor for 5 min, just as in the pre-processing, but with 300 g seaweed to 1.8 L water. The mixed seaweed was further homogenized with an Ultra Turrax for 10 min, as described in the pre-process (Scheme 1, step 4), but with a volume of 2.1 L. The mixing was followed by homogenization of the seaweed in three replicates both for fresh and frozen seaweed. The homogenized seaweed was then transferred to Falcon tubes, with 2 tubes containing 40 mL and 2 tubes containing 20 mL and 20 mL added sodium citrate (0.2 M). The tubes were put on a shaking board for 16 h at 150 rpm and a shake radius of 0.5 cm (Scheme 2, step 2A/B). The sodium citrate was added to dissolve the alginate. The Falcon tubes were then centrifuged for 10 min at 3850×g (Scheme 2, step 3A/B). The supernatant from the Falcon tubes without sodium citrate (31–35 mL) was transferred to small glass vials, into each of which 3 mL of sample was transferred and then dried under airflow for less than 6 h (Scheme 2, step 4A).

The supernatant from the Falcon tubes with sodium citrate (31–35 mL) was transferred to other Falcon tubes, and then, 10 mL of HCl (1 M) was added to precipitate the alginate for 1 h (Scheme 2, step 4B). The Falcon tubes with the precipitated alginate were centrifuged for 5 min at 3850×g to form an alginate pellet in the bottom (Scheme 2, step 5B). The liquid above the pellet was discarded, and the pellet was dispersed in deionized water to a total volume of 45 mL in the tubes, shaken for 30 min, and centrifuged again. The procedure was performed once with water and then four times with an ethanol/water-mix (60%/40% (w/w)) (Scheme 2, step 6B). Afterwards, the alginate pellets were transferred to weighed small plastic Petri dishes, on which it was dried to measure its dry weight (it was dried in the form of acidified alginate, alginic acid) (Scheme 2, step 7B).

An experiment was also made to compare the Ultra-Turrax-based extraction technique for the extraction of laminarin (Scheme 2, step 1–4A), with that of a harsher protocol when the biomass was powdered and dissolved chemically to remove alginate, and laminarin measured in the solution after alginate removal. This method followed the route of (Scheme 2, step 1–5B), but with the exception that it started with freeze-dried powder (of the seaweed used for the main cross-flow filtration study), and that it used sodium carbonate (0.1 M) instead of sodium citrate. The freeze-dried powder was added 0.40 g to 40 mL sodium carbonate solution. In this method the supernatant from the first centrifugation of (Scheme 2, step 5B) was not discarded but instead neutralized with hydrochloric acid and dried for the measure of laminarin and mannitol as in (Scheme 2, step 4A).

Cross-flow filtration setup

The cross-flow filtration is schematically described in Scheme 3. The cross-flow filtration setup had a 60-L tank for the feed solution positioned 30 cm higher than the pump to give a certain elevation to simplify the filling of an empty pump. Two pumps were used—an internal gear pump from Micropump, Inc., for the lower feed flow speeds and a horizontal multistage centrifugal pump, Grundfos CM-G 3-8, for the higher feed flow speeds. The membrane housing was of the type KERASEP Mini, with a cylindric ceramic membrane sealed in position to make sure that the filter solution only
 flowed through the axial channels enclosed in the ceramic membrane, while the permeate was pressed out through the filter in the radial direction through the membrane. The volume of the membrane house outside the filter was 0.5545 L. The ceramic membranes (ZrO$_2$/TiO$_2$, monolithic TiO$_2$-Al$_2$O$_3$ support) were of the brand Kerasep BW from Novasep with stated membrane $M_w$ cutoffs at 5 kDa, 15 kDa, and 50 kDa (5-, 15-, and 50-kDa membranes). The membranes were 38 cm long with a diameter of 2.5 cm and had 19 axial channels each with a diameter of 3.5 mm, giving them an inner area of 0.08 m$^2$. A pressure-control handle was placed after the membrane housing to control the pressure of the feed entering the membrane housing. A pressure gauge was positioned on both sides of the membrane housing to measure the pressure in the feed passing through the channels of the filter. The pressure gauges positioned on each side of the membrane house were also used to record the pressure drop due to friction in the channels of the membrane. All parts of the filtration setup were attached to each other by tubing clamps, bolts with PTFE thread tape, and gaskets with rubber O-rings that were resistant to pressure up to 10 bar and temperatures up to 90°C. The tubing, which was 5 m in total, had a diameter of 2.54 cm and was made of fiber-reinforced synthetic rubber. For some samples, a cooling system was installed with a 30-m-long 0.8-cm diameter rubber tube in 0°C water, through which the feed was led before it entered the feed tank. In other experiments, the permeate was led directly back to the feed tank, thus keeping the feed volume constant. Between every process, the equipment was thoroughly washed with NaOH solution of at least pH 12.5 with a temperature of 60°C for more than 2 h. This was followed by thorough washing with deionized water.

**Cross-flow filtration procedure**

The permeate from the pre-process (Scheme 1, step 8) was loaded into the feed tank and flowed through the cross-flow filtration unit. At several times during the filtration, a sample was gathered from both the feed solution and the permeate solution, with the sample gathered from the tube from which the permeate dripped. At these times, the flow of the permeate per minute was measured, and the total weight of the permeate, the temperature of the feed solution, and the pH of the feed solution were also recorded. At the end of the filtration, samples were taken from both the feed and the permeate, and a sample was taken from the permeate tank where the permeate from the whole process was mixed. For each sampling, two small glass beakers were filled with 1 mL and 3 mL of sample and dried under airflow, drying in under 6 h. The small beakers with dried samples were used for NMR and SEC analyses.

**Large-scale alginate extraction**

To demonstrate that the pre-process leaves a filter cake that is useful for alginate extraction, the filter cake was tested as visualized in Scheme 4. The filter cake from the pre-processing of 5-kg frozen algae was used for alginate extraction. The filter cake was neutralized with NaOH, diluted with water to 60 L with the addition of sodium citrate, and stirred for 16 h at 300 rpm with a propeller with a diameter of 10 cm in the solution (Scheme 4, step 1). The solution was diluted to double the volume, totaling 120 L, with water and left for 20 h to sediment (Scheme 4, step 2). The liquid over the sedimented material was pumped to another container, into which acid was added to reach pH 1.6, which was acidic enough to let the alginate precipitate (Scheme 4, step 3). The liquid with precipitated alginate was poured onto a nylon mesh (the same mesh as in Pre-process), with mechanical scraping by a rotating cylinder with a diameter of 35 cm rotating at approximately 30 rpm to let the water through but not the acidified alginate (Scheme 4, step 4). After removal, the acidified alginate was dissolved again with the addition of NaOH to reach a neutral pH (Scheme 4, step 5). In Scheme 4, steps 3–5 were repeated three times and a final step 4 was performed, and after as much water as possible had dripped through the mesh, the acidified alginate was mixed with an equal volume of ethanol, which enabled far more liquid to be squeezed out (by hand). After the final neutralization via Scheme 4, step 5, the neutralized alginate solution was added to plastic trays with a total surface area of 0.7 m$^2$ and left to dry with airflow overnight (Scheme 4, step 6).

**Analytical techniques**

$^1$H-NMR

$^1$H-NMR was used as the main technique to determine the amounts of laminarin and mannitol in a sample. It was also used to monitor possible bacterial activity during the cross-flow filtration process, which was seen as a decrease in mannitol with a simultaneous buildup of acetic, formic, succinic, and lactic acid. NMR samples were prepared by adding 1 mL of D$_2$O to vials with dried material inside, capping the vials and putting them on a shaking board for 5 h, shake frequency of 150 rpm, and shake radius of 0.5 cm. To make sure that
dried material further up on the vial walls also had a chance to dissolve, the vials were manually rotated one by one to let the D₂O reach the full area of the glass wall, which was a process done three times per samples and initiated after approximately 3 h of shaking time. In the cases in which particles of solid material were visible after shaking, the vials were centrifuged at 2000×g for 5 min to simplify the filling of NMR tubes with mainly liquid contents. The samples were transferred to NMR tubes with 5 mm diameters. ¹H-NMR spectra were recorded on an Advance 400 (Bruker, USA) spectrometer with the standard ¹H-NMR program and 128 scans for each measurement. MestReNova software was used for data acquisition. To determine the amounts of laminarin and mannitol, peaks were chosen based on the criterion of as little overlap as possible between laminarin and mannitol at the position of the peak. The peak at 3.54 was chosen for laminarin, and the peak at 3.91 was chosen for mannitol. After baseline correction by the Splines-method from MestReNova, the intensity of each peak was recorded. The intensity was normalized by dividing the recorded intensity by the receiver gain value (RG) and the number of scans in each measurement. To correlate the measured normalized intensity with the actual concentration, a calibration curve was made for each compound by making five measurements with different concentrations of the pure compound. For laminarin, the calibration curve was slightly bent since the laminarin peaks became a little thicker with higher concentrations. For mannitol, the curve was straight. The calibration curves for laminarin and mannitol, as well as the spectra, are seen in Table 1.

On one occasion, fractions from SEC (Size exclusion chromatography) were used for NMR analysis. These samples were treated like the other samples with the exception that 0.45 mL D₂O was used for dissolution and 4100 scans for analysis.

### Table 1 Processing results of biomass used for cross-flow and filtration, as well as fresh/frozen biomass. The processes are described in Scheme 1 and Scheme 2. The biomass was diluted with water to 5 times its weight in the homogenization step of the main cross-flow filtration study and 7 times its weight in the pre-study on fresh vs. frozen biomass. ± is the standard deviation, of 23 samples for the frozen biomass of the main cross-flow study, and of 6 samples for the fresh/frozen biomass of the pre-study and the dissolutions with sodium carbonate. The viscosity was measured with at least 3 replicates per temperature.

| Processing before cross-flow filtration (Scheme 1) | Prestudy: fresh vs. frozen biomass (Scheme 2) |
|--------------------------------------------------|---------------------------------------------|
| Frozen biomass                                   | Frozen biomass                              |
| Dry weight of biomass average (% w/w)            | 11.00 ± 1.02                                | 5.70 ± 0.15  | 4.88 ± 0.18 |
| Laminarin solubilized from non-dried biomass (% w/w) | 2.70 ± 0.467                  | 0.965 ± 0.149 | 0.083 ± 0.046 |
| Alginate extracted from non-dried biomass (% w/w) | 4.60                                       | 2.39 ± 0.19  | 2.42 ± 0.07  |
| Permeate through the nylon mesh average ratio (% w/w) | 70.2 ± 6.3       | x            | x            |
| Other tests on the biomass used for the cross-flow filtration study |                                |
| Laminarin solubilized from biomass at 25 °C for 0, 1, 4, 20 h and at 65 °C for 20 h (% w/w) | 25 °C: 2.96, 2.91, 3.03, 3.02 | 65 °C: 2.95  |
| Laminarin solubilized from freeze-dried pulverized biomass with sodium carbonate (% w/w, of the frozen biomass) | Laminarin: 2.699 ± 0.055 |
| Dynamic viscosity of feed solution determined by capillary viscometry at 14, 20, 25 and 37°C (mm² s⁻¹) | 14 °C: 1.395 ± 0.020 | 20 °C: 1.201 ± 0.008 |
| Particles in feed solution determined by centrifugation, 3850 ×g, and by comparing the dry weight of pellet and supernatant (% w/w) | 25 °C: 1.072 ± 0.001 | 37 °C: 0.851 ± 0.001 |
| 0.122                                            | 0.122                                       | 0.122         |

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**Scheme 4** Alginate extraction from the filter cake after processing
Size exclusion chromatography

SEC was used to determine the $M_w$ and eventual change in the $M_w$ of laminarin during processing by cross-flow filtration. SEC samples were prepared by letting 1 mL of a sample dry in a small vial, as assisted by airflow. For every new filtration, a sample was also prepared for the solution right before the start of filtration, and one sample was taken before the acid addition and filtration in Scheme 1, steps 5b–6. The samples taken before filtration were centrifuged for 5 min at 3850×g to remove debris, and then, the supernatant was dried in a vial, as were the other samples. To the vials containing dry material, 5 mL of deionized water was added, and the vials were put on a shaking board for 4 h. The estimation of the $M_w$ distribution of laminarin during processing by cross-flow filtration. The kinematic viscosity was measured with a capillary tube viscometer (Xylem Analytics Germany GmbH, prod. nr: 501 10/1, k-factor 0.009701 mm² s⁻¹) placed in a tank of 100 L water with temperature controlled to a precision of 0.1 °C. Prior to measurement, the solution was centrifuged for 5 min at 3850×g and only the part of the particle-free supernatant was used for viscosity measurement.

Viscosity measurement

The kinematic viscosity was measured with a capillary tube viscometer (Xylem Analytics Germany GmbH, prod. nr: 501 10/1, k-factor 0.009701 mm² s⁻¹) placed in a tank of 100 L water with temperature controlled to a precision of 0.1 °C. Prior to measurement, the solution was centrifuged for 5 min at 3850×g and only the part of the particle-free supernatant was used for viscosity measurement.

Results

Pre-processing results and results from the study on fresh vs. frozen biomass

*Saccharina latissima* can serve as a source of both laminarin and alginate. We previously showed that a solution of sodium citrate works well for the extraction of alginate due to the strong chelation of the citric acid anion, which also occurs at a neutral pH (Sterner and Edlund 2016). A common technique used to extract more alginate with a more dilute extraction solution is to first apply an acid pretreatment step on the algae. In this paper, we focus on the extraction of laminarin from the acidic solution that is left after acid pretreatment of seaweed, with the help of cross-flow filtration. Most of the experiments were performed on frozen biomass, but since freezing can make a difference in extractions, a test was performed to determine to what extent alginate and laminarin can be extracted from fresh and frozen biomass. The results from the processing of the biomass that was later used for cross-flow filtration, as well as the comparison between the processing of fresh and frozen seaweed biomass, are presented in Table 1.

The extraction of laminarin was done with the same homogenization equipment and similar concentrations, extraction times, and amounts of stirring in all cases. The amounts of extracted compounds varied greatly between the two different batches of frozen biomass. The batch of biomass used mainly for this study had a higher laminarin content compared to the biomass used for the comparison of fresh and frozen biomass. More laminarin was extracted from the frozen biomass than from the fresh biomass. When the time and temperature of stirring the solution was set between 0-20 h and 25°C and 65°C it was seen that the amount of laminarin extracted did not vary greatly with these setting. It was seen that the amount of laminarin extracted did not vary greatly with these settings. Also, when laminarin was solubilized by dissolving freeze-dried and pulverized biomass in sodium carbonate and measuring laminarin in the solution, after the removal of alginate by acid precipitation, the amount of extracted laminarin was similar to the ordinary method of solubilizing laminarin. Alginate was extracted from frozen and fresh biomass with little difference, especially when accounting for the higher dry weight of the frozen biomass.

Cross-flow filtration study

The goal of the study was to determine which settings are suitable for cross-flow filtration for the extraction of laminarin. To evaluate this, a number of batches with processed homogenized biomass were produced and used for the study. In Table 2, there is a summary of which process parameters were evaluated.

The lower liquid velocity generates less heat, which simplifies the cooling of the system. For several of the...
experiments with 0.2 m s\(^{-1}\) feed velocity, we applied cooling by circulating the liquid through an ice bath, as a way of delaying the biological attack of the feed. A higher pump speed with a high feed velocity, which is the signum for cross-flow filtration, reduces the amount of membrane fouling that occurs on the ceramic membrane. In later experiments, a higher feed velocity and higher temperature were used since it was shown to be more effective; in these experiments, no cooling was used. The ceramic membranes separate molecules based on their \(M_w\), which is related to their hydrodynamic volume. Three different membranes were evaluated, 5, 15, and 50 kDa, in order to find membranes with two functions—a membrane with a high passage ratio of laminarin but that filter of particles and larger molecules such as proteins (prefiltering) and the other membrane with low laminarin passage ratio to make sure laminarin becomes concentrated in the feed. Except at times when the temperature increased rapidly, the flux through the membrane decreased gradually because of membrane fouling. To study the long-time effects of filtration settings, some experiments were done with reflux of the permeate directly back to the feed to achieve a steady state.

The pressure is written as 2, 3, or 5 bar as the pressure before the filter (feed pressure), while for the higher feed velocity (4.7 m s\(^{-1}\)), there was a pressure drop caused by the resistance in the ceramic membrane filter channels, so the feed pressure before the filter was 0.5 bar higher than the pressure after. In the cases of a lower feed velocity, the pressure drop was considered neglectable. The average pressure over the membrane (transmembrane pressure) was calculated as:

\[
\text{transmembrane pressure} = \frac{(\text{permeate pressure} – \text{pressure drop})}{2}\quad(1)
\]

### Schematic Table of Parameters That Were Tested for the Filtrations

| Pump                  | Pump speed | Feed velocity | Temperature | Membrane | Pressure | Volume  | Mode         |
|-----------------------|------------|---------------|-------------|-----------|----------|---------|--------------|
| Gear pump             | 2 L min\(^{-1}\) | 0.2 m s\(^{-1}\) | 10–13 °C | 5, 15, 50 kDa | 5 bar   | 17–21 L | Filtration   |
| Gear pump             | 16 L min\(^{-1}\) | 1.7 m s\(^{-1}\) | 34–36 °C | 50 kDa   | 2 bar   | 15–17 L | Filtration   |
| Centrifugal pump      | 52 L min\(^{-1}\) | 4.7 m s\(^{-1}\) | 40 °C     | 5, 15, 50 kDa | 2, 3, 5 bar | Standard: 15–17 L | Filtration, reflux |

Filtrations with a lower feed velocity and cooling

At a lower feed velocity, the permeate flux was low, causing the filtration to take a long time. During the course of the filtration, the laminarin passage ratio was gradually reduced, measured by 1H-NMR in the feed and permeate solution (Figs. 2a and 2a–d). The logarithmic scale of laminarin concentration was chosen since, at a steady passage ratio, the feed and permeate lines keep a constant distance between them. The laminarin passage ratio was 37% for the 50-kDa membrane (Fig. 2a). Even less, 30% went through the 15-kDa membrane, and 14% went through the 5-kDa membrane. The average laminarin passage ratio through the 5-kDa membrane increased somewhat to 17% when the temperature was increased to 36 °C. The probable cause of the low amount of laminarin that passed through the filter is that the thick gel layer and membrane fouling effectively reduced the pore size of the membrane so that it did not separate molecules of the size for which it is designed. The membrane fouling increased during the course of the filtration, and it was seen as a constant decrease of the laminarin passage ratio. Since data on passage ratio was gathered at different times and with different amounts of remaining feed for each filtration experiment, a question arose concerning how to best report the average laminarin passage ratio. Two ways to report were tested for this—the passage ratio was measured as a linear trend of either the remaining feed volume or the filtration time. Then, the linear trends were used to report the passage ratio at half of the filtration time or at half of the filtrated volume. These two measures gave fairly similar results, and these results were also similar to that from taking the plain average of the laminarin passage ratio.

The membrane permeates flux decreased rapidly during the course of the filtration, as shown by the individual permeate flux curves in Fig. 2a–d and shown as the difference from the starting flux to the average flux for the first half of filtration (100–50% remaining feed), which was a reduction by more than a third of the value. For the third quarter of the filtration (50–25% remaining feed), the flux was reduced yet again by more than one-third of the value.

**Filtrations with a higher feed velocity**

With a higher feed velocity over the membrane, the flux increases greatly. The laminarin passage ratio was high when high feed velocities were used together with a 50-kDa membrane. The medium-high feed velocity of 1.5 m s\(^{-1}\) gave an ~57% laminarin passage ratio, and with a 4.7 m s\(^{-1}\) feed velocity, a passage ratio of 75% was achieved, in contrast to the passage ratio of ~37% with the low feed velocity of 0.2 m s\(^{-1}\) (Fig. 3e–f). The flux was also heavily increased with an increased feed velocity. The flux increases at the beginning of the filtrations are due to the increases in temperature. The highest temperature stated in the

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upper right of Fig. 3e–h was achieved in close connection with the point of the highest flux. The small drop in flux that occurred at the beginning of the filtration with a 50-kDa membrane was most likely due to initial filter membrane fouling that was soon
overtaken by the greater effect of the increased temperature (Fig. 3f). The flux dropped greatly at the end of the filtrations due to the problem of pumping too little liquid with a high pump speed, which forms bubbles that reduce the ability of the pump to pump well. It is seen in Fig. 3 h that the laminarin passage ratio was similarly low for the 5-kDa membrane at a lower feed velocity (~16%). The laminarin passage ratio of the 15-kDa membrane (Fig. 3 g) is in the middle, with a small increase compared to the values from the low feed velocity, changing from 30 to 40%. With a system that has only a low amount of membrane fouling, it is seen from the flux curves and the max flux of Fig. 3f–g that for similar settings, the flux approximately doubles for filtrations with 50-kDa membranes compared to the 15-kDa membrane, which is in turn twice as high as the 5-kDa membrane. Long-term filtering effects can be studied either by having a great amount of solution to filter or by refluxing the permeate back into the feed, and this was studied to further test which settings were best for a longer filtration.

The effect of pressure on 50-kDa filtrations (with reflux circulation or a high initial volume)

The experiments with reflux were performed for 48 h+, during which time the permeate was led back into the feed by a thin tube. The experiment without reflux that could compare in terms of time was a big filtration of 65 L solution (Fig. 4 l). From the start of the filtration, the laminarin passage ratio decreased steadily with time, and it decreased faster with increased feed pressure, as seen in Fig. 4i-j. An immediate difference between the feed pressure at 5 bar and 2 bar and, 3 bar was seen, but after a longer time, a difference was also seen between 2 bar and 3 bar. The observed decrease in the laminarin passage ratio was most likely caused by membrane fouling. After 48 h, biological activity in the 2-bar filtration had attacked the laminarin in the system (see below). More on biological activity is found later in Fig. 6 and Table 3. The filtrations at 3 and 5 bar showed a distinct maximum flux peak before decreasing again due to membrane fouling, which was not seen for the filtration at 2 bar. An increase in the temperature increased the permeate flux, as seen in Fig. 4i–l (lower right plot). More information on flux changes with temperature is found in Table 4.

The effect of pressure on 5-kDa filtrations (with reflux circulation or a high initial volume)

The experiments with reflux were performed for 48 h+, during which time the permeate was led back into the feed by a thin
In Table 4, the relation between the temperature and the flux is calculated as the average of the individual log-linear regressions for each filtration (a linear curve fit with ln(m) as y-axis and t as the x-axis). The degradation of mannitol per hour is equal to the exponent of the mannitol ratio expression. ± is the standard deviation.

| Filtrations | Mannitol concentration (g L⁻¹) | Mannitol ratio t (h) |
|-------------|--------------------------------|----------------------|
| Homogenized solutions before filtrations | 1.96 ± 0.36 | |
| Degradation with cooling, 10–13 °C | 2.25 ± 1.44 | e⁻⁰.⁰²²⁵ t (R² 0.62⁺) |
| Degradation without cooling, 37–66 °C | 8.64 ± 3.43 | e⁻⁰.⁰⁸⁸⁴⁴ t (R² 0.88⁺) |
| Degradation after prefiltering, 57–61 °C | -0.16 ± 1.50 | e⁺⁰.⁰⁰⁰¹⁶ t (R² 0.91⁺) |

*Average R² for the fitted curve of each individual filtration in the category.

higher permeate flux (Fig. 6q). Initially, the change from a 5-kDa to a 15-kDa membrane roughly doubled the permeate flux, and the change from 15 to 50 kDa roughly doubled it again. Prefiltering the solutions through a 50-kDa membrane showed the potential to increase the permeate flux for filtrations with 5-kDa membranes (Fig. 5p). When comparative reflux filtrations were performed with 5-kDa membranes with ordinary feeds and feeds that had been prefiltered through either 15-kDa or 50-kDa membranes, it revealed a permeate flux increase for the two prefiling cases but also showed that the laminarin passage ratio increased drastically in the 15 kDa prefiling case but not in the other cases (Fig. 6r). During the course of the extractions, on a few occasions, the laminarin content decreased, as seen in Fig. 6 s–t. This is attributed to biological activity. Several indications of this effect were observed, such as a smell, a decreased pH, and the formation of organic acids, including acetic, formic, lactic, and isovaleric.

**Filtrations with previously filtered samples and signs of biological activity**

For comparable filtrations only differing in terms of the filter membrane used, it is seen that a higher M_w cutoff gave a higher permeate flux. Initially, the change from a 5-kDa to a 15-kDa membrane roughly doubled the permeate flux, and the change from 15 to 50 kDa roughly doubled it again. Prefiltering the solutions through a 50-kDa membrane showed the potential to increase the permeate flux for filtrations with 5-kDa membranes (Fig. 5p). When comparative reflux filtrations were performed with 5-kDa membranes with ordinary feeds and feeds that had been prefiltered through either 15-kDa or 50-kDa membranes, it revealed a permeate flux increase for the two prefiling cases but also showed that the laminarin passage ratio increased drastically in the 15 kDa prefiling case but not in the other cases (Fig. 6r). During the course of the extractions, on a few occasions, the laminarin content decreased, as seen in Fig. 6 s–t. This is attributed to biological activity. Several indications of this effect were observed, such as a smell, a decreased pH, and the formation of organic acids, including acetic, formic, lactic, and isovaleric.

**Table 3** Starting concentration of mannitol, the degradation of mannitol per hour as a percentage of the starting concentration, and the average exponential curve that describe the mannitol ratio per time t in hours (mannitol ratio being mannitol divided by starting mannitol). The curve is calculated as the average of the individual log-linear regressions for each filtration (a linear curve fit with ln(m) as y-axis and t as the x-axis).

| Filtrations | Mannitol concentration (g L⁻¹) | Mannitol ratio t (h) |
|-------------|--------------------------------|----------------------|
| Homogenized solutions before filtrations | 25 | 1.96 ± 0.36 | |
| Degradation with cooling, 10–13 °C | 12 | 2.25 ± 1.44 | e⁻⁰.⁰²²⁵ t (R² 0.62⁺) |
| Degradation without cooling, 37–66 °C | 10 | 8.64 ± 3.43 | e⁻⁰.⁰⁸⁸⁴⁴ t (R² 0.88⁺) |
| Degradation after prefiltering, 57–61 °C | 3 | -0.16 ± 1.50 | e⁺⁰.⁰⁰⁰¹⁶ t (R² 0.91⁺) |

*Average R² for the fitted curve of each individual filtration in the category.

The experiment without reflux that could compare in terms of time was a big filtration of 55 L solution (Fig. 5p). In the reflux filtrations with 5-kDa membranes, the laminarin passage ratio decreased steadily with time and decreased faster with a higher feed pressure. The filtration at 2 bar with a 5-kDa membrane (Fig. 5 o) showed signs of a biological attack on laminarin after 48 h. More on biological activity is found later. An increase in temperature increased the flux, especially for laminarin after 48 h. More on biological activity is found later. During the course of the extractions, on a few occasions, the laminarin content decreased, as seen in Fig. 6 s–t. This is attributed to biological activity. Several indications of this effect were observed, such as a smell, a decreased pH, and the formation of organic acids, including acetic, formic, lactic, and isovaleric.

**Table 4** Permeate flux minimum, maximum and “steady state” (24-h value) for the different pressures and temperatures and the flux normalized to 1 bar pressure and the viscosity at 20 °C temperature. In the lower part, flux is shown as a function of temperature and viscosity for the linear stage when the temperature increase occurs faster than membrane fouling.

| Filtrations | 5 bar (4.75⁺) | 3 bar (2.75⁺) | 15 bar (3.75⁻) | 2 bar (1.75⁻) |
|-------------|---------------|---------------|----------------|---------------|
| Flux initial min (L m⁻² h⁻¹) at the temperature (°C) | 28.7 30 29.9 26 78.9 38 10.8 22 14.6 26 39.8 20 85.9 36 | 12.1 32 54.2 22 | 1.73² | 2.00 |
| Flux at max (L m⁻² h⁻¹) at the temperature (°C) | 34.1 51 44.2 54 111.2 61 20.9 55 26.9 60 64.5 55 111.8 56 | 15.0 51 80.5 52 | 1.00 1.1 0.88 c 1.06 |
| Flux 24-h value (L m⁻² h⁻¹) at the temperature (°C) | 27.5 65 101.1 64 13.0 60 18.5 60 60.0 59 78.0 59 | 13.2 58 61.5 58 | 2.70 1.12 1.00 0.88 c |
| Flux initial min (L m⁻² h⁻¹) normalized to 1 bar, 20 °C | 4.8 5.5 11.3 3.7 4.6 14.5 21.9 | 5.3 | 29.6 |
| Flux at max (L m⁻² h⁻¹) normalized to 1 bar, 20 °C | 3.9 4.8 10.8 3.8 4.5 11.8 20.2 | 4.6 | 24.3 |
| Flux 24-h value (L m⁻² h⁻¹) normalized to 1 bar, 20 °C | 2.5 9.4 22.2 3.1 10.3 13.4 | 3.6 | 16.9 |

*a Transmembrane pressure in bars
*b Prefiltered sample before 5-kDa membrane filtering
*c The time and temperature ranges were low for this sample, and the value should be considered approximate.
and succinic acids (seen by H-NMR but data not included). A very good signal of biological activity was the decrease of the compound mannitol, which is always present initially in the seaweed. Mannitol is a small enough molecule to pass through all the membranes unhindered, so the only reason that mannitol should change in concentration is degradation. Table 3 is a summary of the degradation of mannitol. For every filtration, the mannitol concentration was observed at the start and over time and fitted to an exponential function, and the average of the exponent was calculated. The average degradation of mannitol was approximately 2.2% h$^{-1}$ for experiments with cooling and 8.6% h$^{-1}$ for an experiment at higher temperatures and almost zero for samples that had been prefiltered.

| Laminarin passage ratio (%): Permeate/feed | i | j | k | l |
|--------------------------------------------|---|---|---|---|
| 0.5 h                                      | 48.5 | 61.1 |
| 1 h                                        | 43.5 | 63.7 | 57.2 | 63.0 |
| 3 h                                        | 41.5 | 54.8 | 58.6 | 57.7 |
| 6 h                                        | 43.2 | 50.9 | 62.1 | 7 h | 52.5 |
| 24 h                                       | 26.6 | 37.6 | 50.2 | 10.5 h | 57.9 |
| 48 h                                       | 28.6 | 31.0 | 89.8 |

| Permeate Flux (L m$^{-2}$ h$^{-1}$) | i | j | k | l |
|-------------------------------------|---|---|---|---|
| 0.5 h                               | 81 | 86 | 80 | 54 |
| 1 h                                 | 87 | 84 | 58|
| 3 h                                 | 109 | 112 | 95 | 72 |
| 6 h                                 | 110 | 106 | 95 | 7 h | 78 |
| 24 h                                | 101 | 78 | 86 | 10.5 h | 62 |
| 48 h                                | 104 | 67 | 78 |

Fig. 4 Filtrations at a higher feed velocity with a 50-kDa membrane and varied pressure during filtration. i–k have the permeate refluxed into the feed, while l has a sufficient volume to reach a steady state of 65 L. Top plots: Laminarin concentration and permeate flux plotted versus the time of the filtration. The membrane M$_w$ cutoff, max temperature, feed velocity, and feed pressure are specified over each plot. The altered setting in i–k is the feed pressure. k–l are comparative filtration experiments—one with the permeate refluxed back into the feed and one with a large but finite feed volume. Lower right plot: Permeate flux and temperature for the first 10 h of filtrations i–l. Straight, striped, dotted, and dotted straight lines are the curves of i–l, as indicated at the top of the plot.

The effects of pressure and temperature on the flux

A number of filtrations were performed with the reflux of the permeate back to the feed or with a large feed volume. The results of these filtrations show the long-time effects on membrane fouling but are also interesting in the start stages since the feed composition was more or less constant, while the temperature increased rapidly. To give a good overview, the permeate fluxes of the initial minimum, the maximum, and the “steady state” of 24 h are given for these filtrations with the temperature specified (Table 4). A normalized flux was calculated by first dividing the original permeate flux by the transmembrane pressure, which gave the flux as it would be.
for an equally blocked membrane but with a transmembrane pressure of 1 bar (pressure-normalized flux). As the next step, the pressure-normalized flux was divided by the dynamic viscosity of water at 20 °C (1 mm² s⁻¹) and multiplied by the dynamic viscosity of water at the temperature of the measurement, which gave the permeate flux that would result if the feed temperature was 20 °C and the transmembrane pressure was 1 bar. The logic behind this is that since our feed is a dilute low viscous solution, the viscosity depends linearly on the viscosity of water, while the permeate flux is inversely proportional to the viscosity of the feed. When the viscosity was measured between 14 °C and 37 °C for our solution, presented in Table 2, this linear relation between pure water viscosity and the viscosity of our solution was true to 1.5% accuracy, and the viscosity of our solution was 1.208 times higher than of pure water.

\[
\text{Normalized permeate flux} = \frac{\text{permeate flux}}{\text{transmembrane pressure}} \times \frac{\text{H}_2\text{O visc. at 20°C}}{\text{H}_2\text{O visc. at temp}}
\]

(2)

From the results of the normalized permeate flux (Table 4), it is seen that for all filtrations, while the flux increased from the minimum at the start of the filtration to the maximum value...
in the early hours of the filtration, the normalized flux decreased and after 24 h it decreased further.

At the specific interval when the temperature of the feed and, thus, the permeate flux increased very rapidly, there was a faster flux increase than the flux decrease due to membrane fouling or changes in the gel layer. This interval was called the range of linearity. A linear trend line was fitted to the data within the range to calculate the percent impact of a 1 °C temperature change or, similarly, a 1% viscosity change of the feed (Table 4). A 1% decrease in the feed viscosity resulted in an approximately 1% increase in the permeate flux.

Changes in molecular weight and SEC data

As seen in Fig. 7 a, the majority of laminarin is found in one peak at around 9 kDa (peak II), which was seen by running fractions from the SEC by NMR, this from a sample in which the laminarin content had been concentrated in the feed by filtering through a 5-kDa membrane for 63 h, 10–13 °C, and to one-third of the original feed volume. It is also seen that peak II has a smaller extra peak. The composition of the extra peak was not identified but it was found that this peak went through the 5-kDa membrane at a higher percentage than laminarin, giving that after longer concentration experiments, it was reduced in comparison to laminarin. This is clearly shown in Fig. 7 a, when a sample was concentrated to one-third of its volume twice, with a refill of water in between and over the course of 63 and 72 h, making the extra peak almost disappear. In Fig. 7 a, the change in peak shape of peak II in the permeate of the different membranes is visualized, showing both a decrease in peak area and a position at slightly lower Mw. In Fig. 7 b, the ratio of permeate vs feed for Mw and area of peak II is presented. The Mw in the permeate decreased compared to the feed with ratios down to around 60% for the extended filtrations with permeate reflux. In Fig. 7 b, data are presented for the ratio of peak area II between permeate and feed, and the ratio of laminarin between permeate and feed was previously determined by NMR (laminarin passage ratio). The values of these ratios followed each other rather well. In total, also including the filtrations now shown in Fig. 7 b, the difference of peak area II ratio minus laminarin passage ratio for 50-, 15-, and 5-kDa membranes was $4.0 \pm 3.6\%$ ($n = 22$), $-0.3\% \pm 4.9\%$ ($n = 8$), and $-3.3\% \pm 4.9\%$ ($n = 15$) (where $n$ is the number of compared pairs of ratios and ± the standard deviation).

The peak with the highest Mw compounds, peak I, decreased in molecular weight during the course of the filtrations.
Concentration of laminarin in feed at 5 bar, 0.2 m s\(^{-1}\), 10-13°C

Fig. 7 a Peak I–II in SEC chromatograms for some filtrations. M\(_w\) marks on the axis are calculated from a pullulan standard calibration curve. Shown sections are from 14 to 34 min out of 40 min. a\(_i\) filtrations with a 5-kDa membrane in which laminarin becomes concentrated in the feed. a\(_i\) One sample fractionated by SEC to measure laminarin by NMR (Size exclusion chromatography). a\(_{ii}\) Feed and permeate after a filtration with feed volume reduced to 1/3, taking 63 h, and the repeated procedure taking 72 h, after refilling with water between. a\(_{iii}\) Filtrations with permeate reflux.

Discussion

Initial tests

Regarding the initial tests on fresh and frozen biomass (beginning of Cross-flow filtration study), the amounts of extracted M\(_w\) of peak II of Fig. 7 a\(_i\) from which the extra peak was almost removed had a M\(_w\) of 9.7 kDa.
compounds varied between the two different batches of frozen biomass. That laminarin content varies greatly with the season and growth site is a known trait and was expected (Black 1950; Schiener et al. 2015). That great difference in extracted laminarin between fresh and frozen biomass, with almost ten times as much laminarin coming from the frozen biomass is a factor to consider if a similar disruption technique as ours is to be used. This is probably since freezing helps break cell walls and make homogenization and extraction easier. That neither changing the time and temperature of the laminarin extraction step nor changing to a technique with sodium carbonate-based dissolution of the freeze-dried and powdered seaweed managed to extract more laminarin is the reason to believe that our laminarin dissolution was extracting a major part of the laminarin.

Cross-flow filtration

The higher feed velocity of 4.7 m s$^{-1}$ as compared to 1.7 m s$^{-1}$ or 0.2 m s$^{-1}$ was beneficial for all filtrations since it increased the permeate flux, especially in the long run, by reducing membrane fouling (Filtrations with a lower feed velocity and cooling, Filtrations with a higher feed velocity, The effect of pressure on 50-kDa filtrations (with reflux circulation or a high initial volume), The effect of pressure on 5-kDa filtrations (with reflux circulation or a high initial volume)). The reason to choose a lower pressure is that even though a higher pressure such as 5 bar increase the permeate flux, it also reduces the laminarin passage ratio, and as seen on the comparably low value on pressure/temperature-normalized permeate flux (at max: 10.8 and 24.3 L m$^{-2}$ h$^{-1}$, for 5 and 2 bar), meaning that the energy demanding buildup of pressure has low effect on the flux (The effects of pressure and temperature on the flux). (In the ideal case, the buildup of double the pressure requires double the energy.)

On the contrary for the filtrations with the 5-kDa membrane, the higher feed pressure of 5 bar was preferred since it gave the lowest laminarin passage ratio and since increasing the pressure did not increase the permeate flux much. A good permeate flux at this setting was 60–80 L m$^{-2}$ h$^{-1}$ at approximately 60 °C (The effect of pressure on 50-kDa filtrations (with reflux circulation or a high initial volume)). The reason to choose a lower pressure is that even though a higher pressure such as 5 bar increase the permeate flux, it also reduces the laminarin passage ratio, and as seen on the comparably low value on pressure/temperature-normalized permeate flux (at max: 10.8 and 24.3 L m$^{-2}$ h$^{-1}$, for 5 and 2 bar), meaning that the energy demanding buildup of pressure has low effect on the flux (The effects of pressure and temperature on the flux). (In the ideal case, the buildup of double the pressure requires double the energy.)

To check whether thermal or microbial activity degraded laminarin its M$_w$ over time was measured by SEC (Changes in molecular weight and SEC data). Our laminarin samples were measured to 9 kDa in M$_w$, but in a peak that included a small overlapping extra peak so it should be considered approximate value. Laminarin, as most (1,3)-β-glucans, is a stiffer molecule than pullulan, that was used as SEC standard, which makes laminarin occupy a larger hydrodynamic volume than pullulan and have its M$_w$ somewhat overestimated by the SEC (Gidley and Nishinari 2009). In other studies, laminarin is also often determined to slightly lower M$_w$ values, such as 3–5
kDa (Rinaudo 2007) or 4–6 kDa (Stone 2009). However, both laminarin and pullulan occupy a larger hydrodynamic volume than the more globular molecules (typically dextran or proteins) that are used for calibration of the ceramic membranes, meaning that they pass the membrane to a lower degree than the molecule of which the Mw cutoff refers to (Ripperger et al. 2013). The substance, that showed up as a small extra peak that overlapped the laminarin peak in the SEC chromatograms, was more prone to pass the 5-kDa membrane compared to laminarin, seen as a gradual decrease in its concentration relative to laminarin in the feed during filtrations with 5-kDa membrane. The laminarin containing fraction showed no sign of decreasing in Mw over time but instead increased a little. The increase can be related to the fact that the fraction with the highest Mw decreased in Mw and that the fragments of larger molecules joined in with the fraction containing laminarin. That fractions of higher Mw compounds get degraded can possibly have a positive effect on filtrations since larger molecules drive the membrane fouling in these filtrations (Bhave 1996); however, if these fragments join in with the laminarin fraction, it could complicate the final purification of the product. In the filtrations at the highest pressure and temperature, an increase in permeate flux at the end of the filtrations was observed (The effect of pressure on 5-kDa filtrations (with reflux circulation or a high initial volume)) which could indicate that thermal (or shear) induced changes in the feed make it easier to filter. The final step of purification was not investigated in this study.

Comparison to other studies

A few publications and three patents can be found in which cross-flow filtration is suggested for the concentration of laminarin. In a patent by Yvin et al. (1998), a 5-kDa membrane is used for the concentration of laminarin. In a patent by Yvin et al. (1999) a 50-kDa membrane is used for pre-filtration and a 0.5–1-kDa membrane is used for the concentration of laminarin, and in a patent Hjelland et al. (2–12), a membrane with Mw cutoff of 10 kDa or greater is used for prefiltering and a membrane with Mw cutoff of 1 kDa or smaller is used for laminarin concentration (Yvin et al. 1998, 1999; Hjelland et al. 2012). In a paper by Klarzynski et al. (2000), a 300-kDa membrane is used for pre-filtration and a 1-kDa membrane is used for the concentration of laminarin. In a paper by Becker et al. (2020), laminarin from microalgae is concentrated using a 1-kDa membrane while laminarin is also consumed by microorganisms during the process.

Besides describing the filtration membranes used, we found no paper or patent that describe the cross-flow filtration for the concentration of laminarin in terms of filtration settings, permeate flux, or laminarin passage/retention ratio. The comparisons are done versus studies that have similar Mw cutoff span (5–50 kDa), similar ZrO2/TiO2 surface structure as the Kerasep filter in this study are used and similar low viscous water-based feed solutions as in our study. These studies are found in Table 5. When using a high feed velocity (4.7 m s⁻¹) for pre-filtration of our extract solutions (by using a 50- or 15-kDa membrane), the permeate fluxes at comparable temperature correspond fairly well to the fluxes for clarification of apple juice (Table 5. Ref. D) and filtration of vanillin lignin solutions for the 50-kDa membranes (Table 5. Ref. C) for the 50-kDa membrane, but our studies permeate flux values are lower than that of filtering oligosaccharides from caprine milk (Table 5. Ref. B). The trend when filtering our feed solution was that the permeate flux was higher with a 50-kDa membrane compared to a 15-kDa membrane, but seemingly with other feed such as the feed composed of lignin and vanillin in (Table 5. Ref. C), it was instead lower. As a general rule and in most studies, the choice of a membrane with greater Mw cutoff was however increasing the permeate flux. At the low feed velocity (0.2 m s⁻¹), our extract solution got a lower permeate flux than that of apple juice, with comparable settings (Table 5. Ref. A). The separation of laminarin with the 5-kDa membrane can be compared to that of filtrating xylan oligosaccharides from rice husk autohydrolysis product since these were also polysaccharides of low Mw, 1–3 kDa (Table 5. Ref. E). With the lower Mw in mind, the comparison between our results and the results of xyloglucan hydrolysis product, 7–16% vs 30% passage ratio, is reasonable. The permeate flux of the xyloglucan hydrolysis product was comparatively higher than in this paper which, when they were also using a lower temperature during the filtration. When comparing to a study in which Kerasep membranes were used as in our study, in which a model lignin degradation product is filtered, our permeate flux is also comparatively lower for the 5-kDa membrane, both when comparing our high and low feed velocity results (Table 5. Ref. F).

Summary and conclusion

In conclusion, laminarin can be concentrated from the solution from the acid pretreatment of homogenized *Saccharina latissima* seaweed using cross-flow filtration as the method. Based on the filtrations in the study, a good method is to prefilter through a 50-kDa membrane at a high feed velocity of ~ 5 m s⁻¹ and a low feed pressure of 2 bar and use the permeate from that filtration as the feed for a 5-kDa membrane filtration. In the 5-kDa membrane filtration, the laminarin is concentrated in the feed, and it should have a high feed velocity of ~ 5 m s⁻¹ and a high feed pressure of 5 bar. With these settings, the laminarin passage ratio began at 15% but reached 7% after 24 h of filtration with a 5-kDa membrane. The laminarin Mw was determined approximately to 9 kDa which motivates the use of a 5-kDa membrane to retain the majority of the product. Cooling the filtrations decreased the microbial
Table 5  Values from our study and values from comparable cross-flow filtration studies from references A-F. A: (Vladisavljević et al. 2003), B: (Martinez-Ferez et al. 2006), C: (Žabková 2006), D: (De Bruijn et al. 2003), E: (Vegas et al. 2008), and F: (Busse et al. 2016). 15–50-kDa filters are for filtration and 5 kDa for concentration

| REF | Feed solution | Product | Filter**, cutoff M<sub>w</sub> (kDa) | Feed velocity (m s<sup>−1</sup>) | Temp. (°C) | Transmembrane pressure (bar) | Permeate flux (L m<sup>−2</sup> h<sup>−1</sup>) | Passage ratio (%) |
|-----|---------------|---------|---------------------------------|-------------------------------|------------|-------------------------------|---------------------------------|------------------|
|     |               |         |                                 | Initial | Steady | Initial | Steady* | Initial | Steady* |
| A   | Apple juice   | Clear juice | Carbosep (50) | 0.2     | 10–13   | 5       | 7, 6   | 4, 4    | 37, 30  |
|     | This study    | Laminarin | Kerasep (50, 15) | 4.7     | 22, 30  | 4       | 54, 79 | 60, 101 | 50–60  |
| B   | Caprine milk  | sugars ~ 0.8 kDa | INSIDE CéRAM (50) | 3.3     | 30      | 2       | 115    | x       | 100    |
|     | Lignin & vanillin 5, 22, 66 g L<sup>−1</sup> | Vanillin | Kerasep (50) | 0.5, 1, 1.2 | 25 | 1.3, 1.3, 1.3 | 60, 40, 30 | 35, 20, 23 | x |
| C   | Apple juice   | Clear juice | Carbosep (50) | 2, 2, 7, 7 | 50–55 | 1.5, 1.5, 4 | 150, 175, 150, 240 | 50, 80, 65, 115 | x |
|     | This study    | Laminarin | Kerasep (15) | 4.7     | 20      | 59      | 110, 60, 28 | 80, 37, 18 | x |
| D   | Apple juice   | Clear juice | Carbosep (15) | 2, 2, 7, 7 | 50–55 | 1.5, 1.5, 4 | 80, 240, 100, 275 | 20, 75, 30, 100 | x |
|     | This study    | Laminarin | Kerasep (5) | 4.7     | 32, 30 | 65, 1.75, 4.75 | 12, 29, 13, 28 | 7–16 | x |
| E   | Hydrolysed rice husk | Xylan, 1–3 kDa | Membralox T1-70 (5) | 2–3     | 26      | 6       | 47     | x       | 30     |
| F   | Model lignin product | Guaiacol 0.104 kDa | Kerasep (5) | 2, 2, 0.2 | 25 | 1, 1.5, 1.3 | 34, 50, 15 | 32, x, x | 64, x, x |

*The 24-h value was taken from our study and the value of the latest time in each study was taken if the curve showed signs of reaching an equilibrium

**All filters being from Novasep, except Membralox T1-70 500 from Pall Corp. and Inside Céram from TAMI Industries
degradation of the feed while filtrating but also made the filtration go slower which makes us not recommend it. Instead, the increase of the permeate flux with temperature and the fact that laminarin showed no detectable sign of thermal degradation motivates testing and increasing the temperature even further.

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Conflict of interest The authors declare no conflict of interest.

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