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Published in:
Future Foods

DOI (link to publication from Publisher):
[10.1016/j.fufo.2022.100189](https://doi.org/10.1016/j.fufo.2022.100189)

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Publication date:
2022

Document Version
Accepted author manuscript, peer reviewed version

[Link to publication from Aalborg University](#)

Citation for published version (APA):

Gregersen Echers, S., Abdul-Khalek, N., Mikkelsen, R. K., Holdt, S. L., Jacobsen, C., Hansen, E. B., Olsen, T. H., Sejberg, J. J. P., & Overgaard, M. T. (2022). Is Gigartina a potential source of food protein and functional peptide-based ingredients? Evaluating an industrial, pilot-scale extract by proteomics and bioinformatics. *Future Foods*, 6, Article 100189. <https://doi.org/10.1016/j.fufo.2022.100189>

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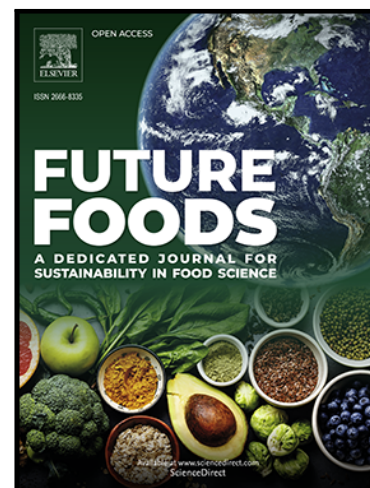
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PII: S2666-8335(22)00076-4
DOI: <https://doi.org/10.1016/j.fufo.2022.100189>
Reference: FUFO 100189



To appear in: *Future Foods*

Received date: 10 June 2022
Revised date: 14 September 2022
Accepted date: 20 September 2022

Please cite this article as: Simon Gregersen Echers , Naim Abdul-Khalek , Rasmus Kranold Mikkelsen , Susan Løvstad Holdt , Charlotte Jacobsen , Egon Bech Hansen , Tobias Hegelund Olsen , Jimmy J.P. Sejberg , Michael Toft Overgaard , Is Gigartina a potential source of food protein and functional peptide-based ingredients? Evaluating an industrial, pilot-scale extract by proteomics and bioinformatics, *Future Foods* (2022), doi: <https://doi.org/10.1016/j.fufo.2022.100189>

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Is Gigartina a potential source of food protein and functional peptide-based ingredients?

Evaluating an industrial, pilot-scale extract by proteomics and bioinformatics

Simon Gregersen Echers^{1,*}, Naim Abdul-Khalek¹, Rasmus Kranold Mikkelsen², Susan Løvstad Holdt², Charlotte Jacobsen², Egon Bech Hansen³, Tobias Hegelund Olsen⁴, Jimmy J. P. Sejberg⁵,
Michael Toft Overgaard¹

¹ Department of Chemistry and Bioscience, Aalborg University, Frederik Bajers Vej 7H, DK-9220 Aalborg, Denmark;
sg@bio.aau.dk (SG), nakg@bio.aau.dk (NAK), mto@bio.aau.dk (MTO)

² Research Group for Bioactives - Analysis and Application, National Food Institute, Technical University of Denmark, Kemitorvet, 201, Kgs Lyngby, 2800, Denmark; rkmik@food.dtu.dk (RKM), suho@food.dtu.dk (SLH),
chja@food.dtu.dk (CJ)

³ Research Group for Gut, Microbes and Health, National Food Institute, Technical University of Denmark, Kemitorvet, 202, 1250, Kgs Lyngby, 2800, Denmark; egbh@food.dtu.dk (EBH)

⁴ Department of Health Technology, Technical University of Denmark, Ørsteds Plads, 345C, Kgs. Lyngby, 2800, Denmark; tobiasheol@gmail.com (THO)

⁵ CP Kelco ApS, Ved Banen 16, Lille Skensved, 4623, Denmark; Jimmy.Sejberg@cpkelco.com (JS)

*Corresponding author: sg@bio.aau.dk (SG)

Highlights

- A seaweed protein extract (SPE) from *Gigartina sp.* was obtained in pilot-scale
- SPE displayed low aq. solubility but good nutritional quality
- Proteomics revealed SPE to be rich in particularly phycobiliproteins
- Predicted emulsifier peptides were validated by protein modelling
- *Gigartina sp.* is a promising source of natural emulsifiers

Abstract

Seaweeds attract substantial interest as novel sources of sustainable food protein, as they are established sources of industrial hydrocolloids with reasonable protein content. In this study, we investigate the protein composition and nutritional quality of a seaweed protein extract (SPE) from *Gigartina sp.* The SPE displayed low (<2%), but pH-dependent, aqueous solubility likely due to the harsh conditions employed during extraction. Solubility improved using alkaline buffering and detergent addition to facilitate proteomic characterization by quantitative LC-MS/MS. Proteomics analysis revealed that SPE was dominated by proteins related to light harvest and particularly phycobiliproteins (44%), where phycoerythrin was most abundant (28%). Based on subcellular localization, the extraction method was evaluated as good for release of cellular protein. SPE was rich in essential amino acids (36-41%) and particularly branched chain amino acids (22-24%), and thereby a potential source of nutritional food protein. Using bioinformatic prediction and structural modelling, we found abundant SPE proteins contained novel peptides with the amphiphilic properties required to stabilize an oil/water interface, and thereby high probability of being potent emulsifiers. Based on this study, *Gigartina sp.* could serve a good candidate for extraction of sustainable, nutritious food protein, with the possibility of further processing into hydrolysates with strong emulsifying properties for use as natural food ingredients.

Keywords: Rhodophyta, seaweed protein, valorization, proteomics, bioinformatics, peptide emulsifiers

1. Introduction

Global food systems are responsible for more than one third of the total anthropogenic greenhouse gas emission, where animal-based food systems account for more than half (Xu *et al.*, 2021).

Meanwhile, the global population continues to increase, thereby also increasing food and protein demand significantly in the future. It is estimated that within the period from 2010 to 2050, global food demand may increase up to 62% (van Dijk *et al.*, 2021). To accommodate this demand, and meanwhile reducing environmental footprint of food systems, development and application of alternative and sustainable protein sources is pivotal. While plant-based protein (green biomass) have received tremendous attention (Aschemann-Witzel *et al.*, 2020), aquatic/marine protein (blue biomass) is making headway in academia and industry as a means to solve global challenges related to sustainable food supply (Lange and Meyer, 2019). Although not by definition a plant (Sudhakar *et al.*, 2018), seaweeds (macroalgae) are regarded as a high potential aquatic source for sustainable and animal-free protein production, thereby also adhering to e.g. vegan dietary preferences (Raja, Kadirvel and Subramaniyan, 2022). In addition, seaweeds are often highlighted for their nutraceutical (Tanna and Mishra, 2018) and health beneficial properties (Holdt and Kraan, 2011). The primary industrial application of seaweeds is for production of hydrocolloids such as alginate, agar and carrageenan, which are used as gelling agents in the food industry (Brown *et al.*, 2014). With approximate 25 million MT cultivated in 2019, where upwards of 70% are mineral- and

protein-rich side- and/or waste-streams (Cai *et al.*, 2021), the global seaweed market is in rapid development and estimated to exceed US\$ 20 billion by 2024 (Custódio *et al.*, 2017). Within the hydrocolloid industry, red seaweeds (Rhodophyta) belonging to the *Eucheuma*, *Chondrus*, *Kappaphycus*, *Hypnea*, and *Gigartina* genera are commonly used as sources for carrageenan extraction (Colusse *et al.*, 2022; Singh *et al.*, 2022). In general, red seaweeds are considered of high potential for food application due to their high protein content, reaching upwards of 47 % dry matter basis (Biancarosa *et al.*, 2016) and therefore also highlighted as a promising source of alternative proteins in the quest to increase global food sustainability (Rawiwan *et al.*, 2022). As red seaweed furthermore constitute more than half of the global seaweed market (Cai *et al.*, 2021), side streams from hydrocolloid production represent a high potential protein source which still remains underutilized (Bleakley and Hayes, 2017; Naseri, Holdt and Jacobsen, 2019). *Gigartina* is a genus belonging to the *Gigartinaceae* family under the *Gigartinales* order and comprises 43 species (Cotas *et al.*, 2022). *Gigartina* sp are considered to be edible seaweed and have been highlighted for their nutraceutical potential (Cotas *et al.*, 2022) due to high levels of minerals, dietary fibers, and protein (Gómez-Ordóñez, Jiménez-Escrig and Rupérez, 2010). In addition to the many direct applications of *Gigartina* carrageenan, the genus has been reported to show antioxidant, skin protective, wound-healing, and anti-proliferative activity (Jiménez-Escrig, Gómez-Ordóñez and Rupérez, 2012; Chrapusta *et al.*, 2017; Cotas *et al.*, 2020). Nevertheless, *Gigartina* protein remains largely unutilized.

Mass spectrometry (MS)-based proteomics analysis has developed into a cornerstone of biotechnological and biomedical research (Armengaud, 2016; Noor *et al.*, 2021), but is still in its infancy for food protein analysis and characterization. In the food context, MS-based proteomics has primarily been employed to investigate food safety, food quality, allergenicity, as well as organism and cellular responses to exogenous stimuli in relation to e.g. crop breeding (Agrawal *et*

al., 2013; Andjelković and Josić, 2018; Mora, Gallego and Toldrá, 2018; Lexhaller, Colgrave and Scherf, 2019; Marzano *et al.*, 2020). Such studies have also been performed for red seaweeds (Beaulieu, 2019). However, the ability of MS-based proteomics to supply qualitative and quantitative insight on the single protein level also opens new possibilities for its application. For instance, MS-based proteomics and downstream bioinformatic data analysis has been employed as a tool to evaluate extraction efficiency from red seaweeds (Gregersen *et al.*, 2022) and how integrated extraction methods such as application of cell wall degrading enzymes can improve recovery of intracellular protein from *Eucheuma denticulatum* (Gregersen *et al.*, 2021).

Food protein is traditionally evaluated based on a range of bulk characteristics related to functional, nutritional, sensory, and safety aspects. Among these, a suitable amino acid (AA) profile is crucial for a protein being compatible with human nutritional requirements (Mæhre *et al.*, 2014). In this context, the content of particularly essential amino acids (EAAs) and the ratio to non-essential amino acids (NEAAs), $\sum\text{EAA}/\sum\text{NEAA}$, are central evaluation criteria. For a representative subsample of red seaweeds, this ratio (0.63) is slightly higher than for green (0.61) and brown (0.59) seaweeds. Compared to other alternative protein sources such as soy (0.91), pea (1.02), potato (1.09), and corn (0.78), this makes seaweed protein overall of a somewhat lower nutritional quality (Donadelli *et al.*, 2019). Nevertheless, some red seaweed species such as *E. denticulatum* have been shown to have comparable nutritional potential with $\sum\text{EAA}/\sum\text{NEAA} = 0.86$ (Naseri, Holdt and Jacobsen, 2019). In this respect it is also important to keep in mind that the AA profile for a whole organism may not at all reflect the profile in a protein extract or isolate. Traditionally, the AA composition is evaluated by means of conventional AA analysis, where protein is fully hydrolyzed in hot hydrochloric acid and subsequently individual amino acids are quantified relative to a standard curve (Rutherford and Gilani, 2009). However, conventional AA analysis comes with limitations. For instance, due to deamidation it is not possible to distinguish Asn and Gln from Asp

and Glu, respectively, Trp is fully degraded, and substantial losses of hydroxyl- (Thr and Ser) and sulfur- (Cys and Met) containing AAs is observed (Rutherford and Gilani, 2009). To alleviate this, the use of peptide- and protein-level MS data has been suggested as a promising alternative to estimate AA composition, and showed good overall correlation with conventional AA analysis (Jafarpour, Gomes, *et al.*, 2020; Gregersen *et al.*, 2021). Moreover, this approach is able to quantify Trp and free Cys as well as distinguish Asn/Gln from Asp/Glu.

A good nutritional quality is important for alternative proteins, but having functional properties increases applicability as food ingredients and thereby also product value (Granato *et al.*, 2010). For many alternative proteins, extraction methodology has the unwanted effect of being denaturing, thereby reducing or completely eliminating functionality and aqueous solubility (Sari *et al.*, 2015). This is particularly relevant when designing a scalable and economically feasible process in the transfer from small-scale lab experiments to full-scale industrial processes. While significant effort is being put into development of novel, green extraction methods that are both environmentally friendly and has potential to produce a non-denatured protein (Picot-Allain *et al.*, 2021), enzymatic hydrolysis offers an alternative solution to the problem. Hydrolysis has been reported to improve solubility, recovery, and functionality for a wide range of alternative proteins such as chickpea (Mokni Ghribi *et al.*, 2015), soy (Sun, 2011), potato (Gregersen Echers *et al.*, 2022), pea (García Arteaga *et al.*, 2020), just to name a few. While this approach holds great potential, the process is usually not well understood or characterized on the molecular level. Rather, hydrolysis is performed by trial-and-error, where specific protease(s) and process conditions are screened and subsequently evaluated based on functionality of the bulk hydrolysate. As the liberated peptides are responsible for the observed functionalities, molecular insight and peptide-level understanding may add value by the possibility of process design based on desired functionality, working more from the bottom-up rather than the top-down.

Recent development in bioinformatic prediction of peptide functionality, has facilitated *de novo* computational screening of proteins for embedded peptides with specific functionalities such as emulsifying (García-Moreno, Gregersen, *et al.*, 2020; García-Moreno, Jacobsen, *et al.*, 2020) and antioxidant (Olsen *et al.*, 2020) activity. Moreover, several web-tools have been developed to predict peptide functionality and bioactivity within multiple classes such as Peptide Ranker (Mooney *et al.*, 2012), MultiPep (Grønning, Kacprowski and Sché Ele, 2021), and BIOPEP-UWM (Minkiewicz, Iwaniak and Darewicz, 2019). While the specific predictors are highly flexible and able to use both protein and peptide sequence input to predict functional potential, the multi-label classifiers come with some limitations. Peptide Ranker and MultiPep are restricted to only peptide input. While BIOPEP-UWM has many flexible features, it is restricted to database searching against known functional and bioactive peptides. These restriction limit their applicability in the discovery phase of novel peptides using protein-level input data, whereas the specific *de novo* predictors have been applied to identify several novel peptides with emulsifying (Yesiltas *et al.*, 2021) and antioxidant (Yesiltas *et al.*, 2022) activity from various alternative protein sources, including seaweeds.

In this study, we present the first proteomic characterization of a protein-rich extract from *Gigartina*. The seaweed protein extract (SPE) was obtained at pilot-scale using a process that is both scalable and industrially relevant, and is implementable as a direct downstream process following initial carrageenan extraction. Using MS-based proteomics, we present a qualitative and quantitative analysis of the protein composition to obtain insight on protein extractability and the nutritional value of the extract. Moreover, using *in silico* sequence analysis for prediction of potential emulsifier peptides embedded in the most abundant proteins, we relate predictions to the peptide conformation within the native proteins. Through this analysis, we construct a protein- and peptide-level understanding of the proteomic composition which is an important step for *ex vitro*

evaluation of a protein source, in relation to its potential as a substrate for production of a functional hydrolysate.

2. Materials and Methods

2.1. Materials

The seaweed protein extract (SPE) was supplied by CP Kelco (Lille Skensved, Denmark) in dry form with a protein content of 42.3% (by Kjeldal analysis and an N-to-protein conversion factor of 6.25, CP Kelco supplied information) as a by-product from mild carrageenan extraction in pilot-scale. Extraction of SPE is described below. All used chemicals were of HPLC-grade.

2.2. Protein extraction

For the extraction of SPE, CP Kelco uses *Gigartina radula* which is commonly used industrially as a joint description of mixed *Gigartina sp.*, more precisely *Gigartina broad leaf*, *Gigartina narrow leaf*, *Gigartina skottsbergii*, *Gigartina chammisoi*, and *Gigartina pistillata*. Aqueous (aq.) carrageenan extraction was performed at 95°C and pH 9 for two hours under constant stirring and subsequently centrifuged at 8,221 g for 20 min. The sediment (initial byproduct) was subsequently washed with aq. HCl (pH 2) for 30 min to degrade residual carrageenan and centrifuged as above. Lastly, the remaining filter cake was washed with dH₂O until a neutral pH was obtained, centrifuged as above, and lyophilized to yield the final SPE.

2.3. Protein solubility

Protein solubility was investigated as pH-dependent aq. solubility and by the addition of buffer and/or detergents. For aq. solubility, ddH₂O was adjusted to six pH values (2, 4, 6, 8, 10, and 12) using dilute hydrochloric acid or sodium hydroxide and 1 mL was added to 22.1 mg SPE corresponding to a total protein concentration of 10 mg/mL (based on supplied protein content) at full solubilization. To investigate the effect of buffer and detergents on protein solubility, 22.1 mg SPE was added 5 mL (final protein concentration 2 mg/mL at full solubilization) of five different solvents: A) 200 mM Ammonium bicarbonate (ABC, Sigma Aldrich, USA) pH; 8.0. B) 2% Sodium dodecyl sulfate (SDS, Applichem, Darmstadt, Germany) in 200 mM ABC pH 8.0. C) 0.2% SDS in 50mM ABC, pH 8.2. D) ddH₂O (no pH adjustment). E) 1% Sodium deoxycholate (SDC, Sigma Aldrich, USA) in 50 mM Triethylammonium bicarbonate (TEAB, Sigma Aldrich, USA), pH 9.5. In all experiments, SPE was mixed with solvent, vortexed for 30 s, sonicated for 30 min, and centrifuged at 3095 RCF and 4 °C for 10 min in a 5810 R centrifuge (Eppendorf, Germany) to precipitate solids prior to aliquoting the supernatant. The total soluble protein (TSP) was determined using both Qubit protein assay (Thermo Scientific, Germany) and A₂₈₀ by Nanodrop (ND, Thermo Scientific) according to the manufacturer guidelines. For Qubit measurements, blank solvents were measured as reference, while for ND, the respective solvent were used to blank the instrument prior to sample analysis. All measurements were performed in triplicates and solubility calculated as % of full solubilization based total protein from Kjeldal analysis (N x 6.25).

2.4. SDS-PAGE

Supernatants from all solubilization experiments were analyzed by SDS-PAGE using precast 4-20% gradient gels (Genscript, USA) in a Tris-MOPS buffered system under reducing conditions and according to manufacturer guidelines. As molecular weight marker, 5 µL PIERCE Unstained

Protein MW Marker P/N 26610 (ThermoFisher Scientific, USA) was used. Visualization was performed by Coomassie Brilliant Blue G250 (Sigma-Aldrich, Germany) staining and gel imaging conducted with a ChemDoc MP Imaging System (Bio-Rad, USA).

2.5. In-solution digest

Proteins were digested in-solution as technical triplicates, according to (Zhou *et al.*, 2015) with minor modifications. In brief, aliquots of 50 µg protein (based on nanodrop) in solvent E (1% SDC in 50 mM TEAB) was heated to 99°C for 10 minutes and subsequently cooled to room temperature (RT). Next, proteins were reduced using tris(2-carboxyethyl)phosphine (TCEP, Sigma-Aldrich) by adding 1 µg TCEP per 25 µg protein and incubating at 37°C for 30 min. Then, proteins were alkylated using iodoacetamide (IAA, Fluka Biochemika, Switzerland) by adding 1 µg IAA per 10 µg protein and incubating for 20 min at 37°C in the dark. Lastly, proteins were digested with sequencing grade modified trypsin (Promega, USA) by adding 1 µg trypsin per 50 µg protein and incubating overnight at 37°C. Following digestion, samples were acidified by adding formic acid (FA, VWR, Denmark) to a final concentration of 0.5% and incubated at RT for 5 min to precipitate SDC. The samples were centrifuged at 13,000 g for 20 min at 4°C, whereafter the supernatants were transferred to new tubes, and centrifuged again (same conditions). Final supernatants were purified using in-house prepared C-18 StageTips, dried down, and resuspended in 0.1% FA in 2% acetonitrile (ACN, VWR) for analysis, as previously described (García-Moreno, Gregersen, *et al.*, 2020).

2.6. Bottom-up proteomics by LC-MS/MS

Protein digests were analyzed by shotgun, bottom-up proteomics, as previously described (Gregersen *et al.*, 2021). Briefly, LC-MS/MS analysis was performed using a system composed of an EASY nLC 1200 (Thermo Scientific, Germany) coupled in-line to a Q Exactive HF (QE-HF) mass spectrometer (Thermo Scientific) via a Nanospray Flex ion source (Thermo Scientific). Peptides were loaded on a reverse phase Acclaim PEPMAP NANOTRAP column (C18, 100 Å, 100 µm. × 2 cm, (Thermo Scientific)) in solvent A (0.1% FA) followed by separation on a reverse phase ACCLAIM PEPMAP RSLC analytical column (C18, 100 Å, 75 µm × 50 cm (Thermo Scientific)). Samples were loaded using 0.1% FA and separated through a gradient from 5-100% of 0.1% FA in 80% ACN over 60 min. The QE-HF was operated in top-20, positive ion, DDA mode, using 28 eV HCD fragmentation and a 1.2 m/z isolation window for MS2. MS1 scans were obtained from 400-1200 m/z with a resolution of 60,000 (at 200 m/z) and MS2 was performed with a resolution of 15,000 (at 200 m/z). Maximum ion injection time was set to 50 for MS1 and 45 for MS2 scans. ACG target was set at 1e6 and 1e5 for MS1 and MS2, respectively. The underfill ratio was set to 3.5% and a dynamic exclusion of 30 sec was applied. During acquisition, “peptide match” and “exclude isotopes” were enabled.

2.7. Protein database construction and LC-MS/MS data analysis

As no specific proteomic, genomic, or transcriptomic databases are available for *Gigartina sp.*, a custom database was created for analysis of LC-MS/MS data. For this, all entries from Uniprot (UniProt Consortium *et al.*, 2021) annotated to the order of *Gigartinales* (16,072 entries) were retrieved. This custom database also includes a references proteome for the related species *Chondrus crispus* (UP000012073, 9598 entries), thereby presumably ensuring sufficient depth and coverage of the *Gigartina sp.* proteome.

LC-MS/MS data were subsequently analyzed in MaxQuant v.1.6.10.43 (Cox and Mann, 2008) against the custom protein database using standard setting for tryptic digest, as previously described (Gregersen *et al.*, 2021). Up to five modifications and two missed cleavages were allowed and a false discovery rate (FDR) of 1% was applied on both peptide (PSM) and protein level. For quantification, both the label-free quantification, MaxLFQ (Cox *et al.*, 2014), and intensity-based absolute quantification, iBAQ (Schwanhüusser *et al.*, 2011), were employed. Quantitative reproducibility for relative iBAQ (riBAQ) quantification was evaluated by the coefficient of variation (CV), determined as the standard deviation of the replicates divided by the mean for each protein (group). MaxQuant output data were further analyzed in Perseus v.1.6.1.3 (Tyanova and Cox, 2018) to filter contaminants and false positives and investigate reproducibility between technical replicates. Overall quantitative reproducibility by riBAQ was evaluated using the stochastic mean CV for all quantified protein groups and by determining an abundance-weighted mean CV, CV_{mean}^w , calculated using riBAQs as weights according to:

$$CV_{mean}^w = \sum_{pro=1}^m CV^q * riBAQ^q$$

where CV^q is the CV for protein q of m total proteins and $riBAQ^q$ is the riBAQ of the same protein.

Data was visualized using Perseus and Venny v.2.1 (freely available at

<https://bioinfogp.cnb.csic.es/tools/venny/>) before being exported for further downstream

bioinformatic analysis.

The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium (<http://proteomecentral.proteomexchange.org>) via the PRIDE partner repository (Perez-Riverol *et al.*, 2019) with the dataset identifier PXD034435 and DOI 10.6019/PXD034435.

2.8. Subcellular localization and signal peptide prediction

The subcellular localization of identified proteins was predicted by deepLoc (Almagro Armenteros *et al.*, 2017) using the open source web-interface (<http://www.cbs.dtu.dk/services/DeepLoc/index.php>) with the application of BLOSUM62 encoding. The predicted compartment was validated for the most abundant (riBAQ > 1%) by application of LocTree3 (Goldberg *et al.*, 2014), as well as annotated subcellular localization in Uniprot (UniProt Consortium *et al.*, 2021). Potential signal peptides in abundant protein (riBAQ > 1%) was predicted by SignalP (v.6.0) (Armenteros *et al.*, 2019) using the open-source web-interface (<http://www.cbs.dtu.dk/services/SignalP-5.0/index.php>). Predicted signal peptides were used to further validate subcellular localization prediction (if applicable).

2.9. Amino acid composition by peptide- and protein-level MS data

Amino acid composition was estimated using peptide- and protein-level MS data, as previously described (Jafarpour, Gomes, *et al.*, 2020; Gregersen *et al.*, 2021, 2022). For peptide-level MS data, the approach of intensity-weighted, peptide-level AA frequency was applied. Following filtering of reverse and contaminant peptides, peptide MS1 intensities were used as weights to estimate molar AA frequency, $f_{AA}^{peptide}$, as:

$$f_{AA}^{peptide} = \sum_{pep=1}^n f_{AA}^p * I_{rel}^p$$

where f_{AA}^p is the relative frequency of a given AA in peptide p and I_{rel}^p is the relative MS1 intensity of peptide p (i.e. the intensity of p divided by the sum of intensities for all n peptides).

For protein-level data, AA composition was estimated using the relative molar abundance (iBAQ). For all identified proteins, the relative AA frequency was computed using ProtrWeb (Xiao *et al.*, 2015) (freely available as webtool at <http://protr.org/>). Using the relative molar frequency of each AA, $f_{AA}^{protein}$, AA composition was calculated as:

$$f_{AA}^{protein} = \sum_{pro=1}^m f_{AA}^q * riBAQ^q$$

where f_{AA}^q is the relative frequency of a given AA in protein q and $riBAQ^q$ is the iBAQ of protein q divided by the sum of iBAQs for all m proteins.

AA composition was evaluated against the minimum adult dietary requirements according to the World Health Organization/Food and Agricultural Organization of the United Nations/United Nations University (WHO/FAO/UNU, 2007), following conversion to relative molar abundance, as previously described (Gregersen *et al.*, 2021).

2.10. Prediction of embedded emulsifier peptides and quantitative visualization

For abundant proteins ($riBAQ > 1\%$), protein sequences were analyzed using EmulsiPred (García-Moreno, Gregersen, *et al.*, 2020; García-Moreno, Jacobsen, *et al.*, 2020) to predict embedded, emulsifying peptides. Peptides were predicted as potential emulsifiers if given an amphiphilic score > 2 when projected in either α -helical or β -strand conformation (facial amphiphiles) or in an undefined (γ) conformation (axial amphiphilicity). Predicted (and Uniprot annotated) signal peptides were used to ensure predicted emulsifier peptides did not have overlap with a potential signal region, which would not be present in the mature protein.

2.11. Protein modelling and visualization

The structure of selected proteins was modelled using templated homology modelling using the Swiss-Model Workspace (Waterhouse *et al.*, 2018), as previously described (García-Moreno, Gregersen, *et al.*, 2020). Phycoerythrin alpha chain (Uniprot AC# I4DEE4) and Phycocyanin, alpha chain (Uniprot AC# M5DDK1) were modelled using the electron microscopy (EM) structures of Phycoerythrin alpha subunit (STML ID 6kgx.453.A) and C-phycocyanin alpha subunit (STML ID 6kgx.59.A) from the red algae *Porphyridium purpureum* as template, respectively. Ribulose biphosphate carboxylase (RuBisCO) small subunit (Uniprot AC# A0A7M3VH79) was modelled using the X-ray crystal structure of RuBisCO small subunit from the red algae *Galdieria partita* (STML ID 1iwa.1.B). Photosystem II reaction center protein L (Uniprot AC# M5DD55) was modelled using the crystal structure of oxygen-evolving photosystem II from the red algae *Cyanidium caldarium* (STML ID 4yuu.1.K). Cytochrome b6-f complex subunit 5 (Uniprot AC# M5DDH2) was modelled using the cryo-EM structure of cytochrome b6f from spinach (*Spinacia oleracea*) (STML ID 6rqf.1.G). Models were visualized using The PyMOL Molecular Graphics System v. 1.5.0 (Schrödinger, LLC.) and colored using the Swiss-Model hydrophobicity color scheme (from hydrophilic (blue) to hydrophobic (red)).

2.12. Statistical analysis

Statistical analysis was performed for experimental data in GraphPad Prism (v.9.3.0) using multiple comparisons of means by ordinary one-way ANOVA and Tukey's HSD test with a 95% confidence level. For proteomics data, technical replicability was evaluated by computing the Pearson correlation coefficient (PCC) between replicates using both riBAQ and LFQ quantitative data in

Perseus. Protein-level reproducibility was evaluated by computing the coefficient of variation (CV) as the standard deviation of the triplicate analysis divided by the mean.

3. Results and Discussion

Initially, the aqueous solubility of the seaweed protein extract (SPE) was investigated in a pH-dependent manner (Fig. 1, Table A.1). In general, SPE showed low aq. solubility (total soluble protein, TSP) regardless of pH and quantification method, although Nanodrop indicated significantly higher solubility (1.5%-11.6%) than Qubit (0.69-2.1%). Nanodrop is an unspecific quantitative estimate relying only on sample absorbance, and high A260/A280 ratios (1.61-2.08) from Nanodrop measurements (Table A.1) for aq. samples indicate low protein purity in the supernatants due to presence of e.g. nucleic acids (Schultz *et al.*, 1994), thereby likely making measurements overestimates. Consequently, these measurements can be considered unreliable due to sample complexity, and for that reason Qubit measurements are considered more accurate, being a protein-specific fluorescence-based assay.

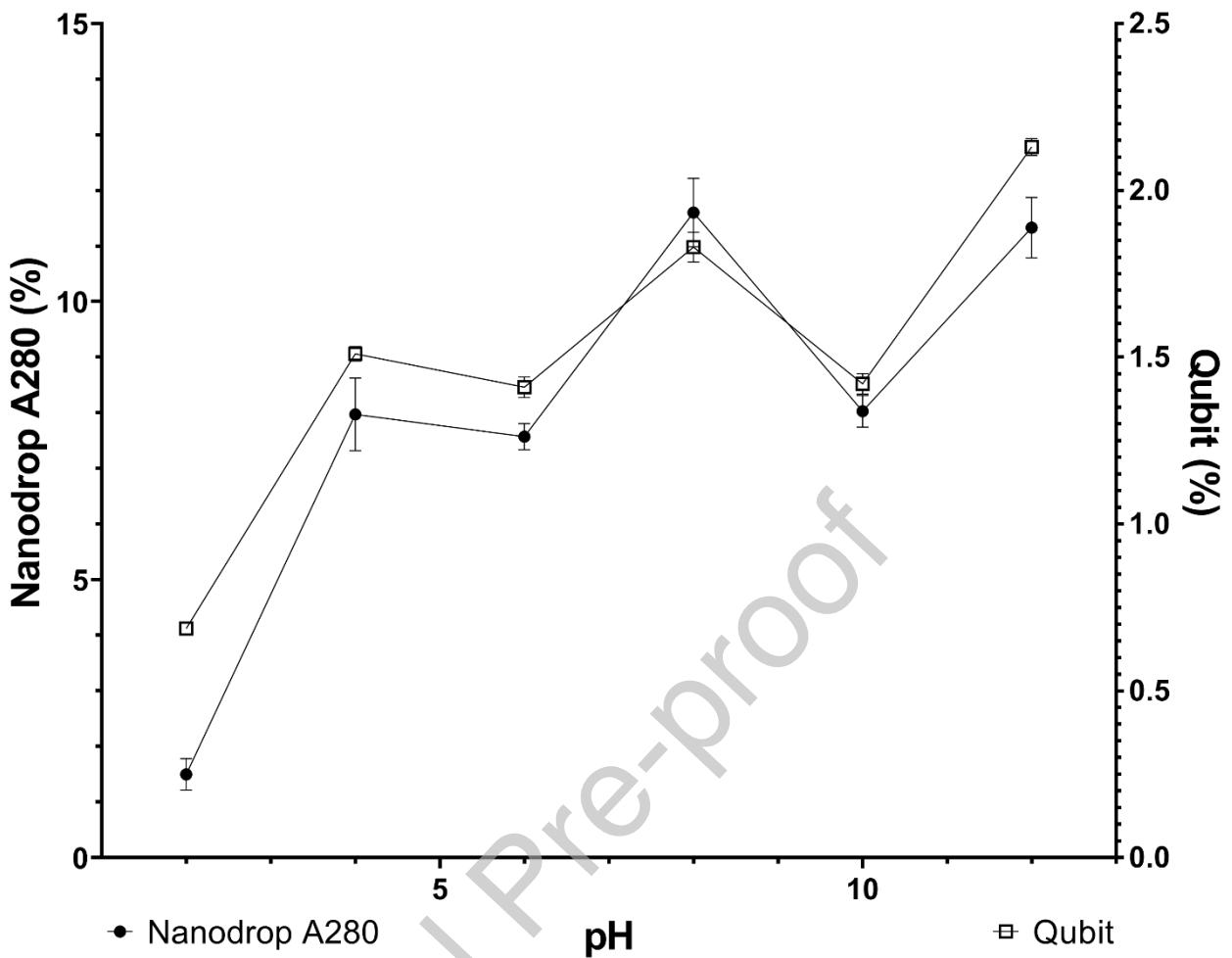


Fig. 1: pH-dependent aqueous protein solubility of SPE. Aqueous solubility of protein in SPE (% relative to supplied protein content by Dumas-N) as a function of pH and quantified using Nanodrop A280 (filled circles, left axis) and Qubit Protein Assay (open squares, right axis). Data points where no error bars are shown is a result of low SD (see Table A.1)

Both Nanodrop and Qubit TSP measurements show similar trends in terms of pH-dependent aqueous solubility of the SPE. While the solubility is significantly lower ($P < 0.05$) at pH 2, the solubility increases towards pH 8, drops at pH 10, and increases again at pH 12. Based on the upstream processing, these observations are not surprising. Initial carrageenan was performed under

slightly alkaline conditions, whereby proteins with high solubility in this range may have been extracted alongside carrageenan, explaining the small drop in solubility at pH 10. Next, the sediment was treated under highly acidic (pH 2) to degrade residual carrageenan, thereby also removing acid-soluble protein from the final SPE, explaining the low solubility at this pH. While the low aqueous solubility of SPE may limit its direct applicability in foods (Amagliani *et al.*, 2017), further processing, such as enzymatic hydrolysis, may significantly improve solubility as well as functionality of the predominantly insoluble protein (Ashaolu, 2020; Gregersen Echers *et al.*, 2022).

To improve protein solubility in order to facilitate deeper proteomic characterization, a range of different solvents were applied to the SPE. These solvents included commonly employed detergents within protein research (SDS and SDC) as well as a slightly alkaline buffer (ABC) at different concentrations. Detergents, buffer components, and their concentration were selected based on compatibility with other employed analytical methods. Moreover, a slightly alkaline pH improved aqueous solubility of SPE in accordance with previous studies on seaweed protein (Veide Vilg and Undeland, 2017; Juul *et al.*, 2022). In contrast to aq. solubilization, Qubit proved inferior due to low compatibility with applied detergents and buffers, where particularly solvent B (2% SDS in 200 mM ABC) and solvent E (1% SDC in 50 mM TEAB) resulted in higher response for solvent references than for the samples (Table A.2). This resulted in a negative protein solubility and is regarded unreliable. Lower detergent and buffer concentrations, i.e. solvent C (0.2% SDS in 50 mM ABC), appeared compatible with Qubit, and significantly ($P < 0.05$) improved protein solubility (7.4%) compared to both aqueous solubilization with no pH adjustment (solvent D. 2.2%) and slightly alkaline buffer in solvent A (3.0%). Solvent A improved SPE solubility slightly, but significantly ($P > 0.05$) compared to aq. solubility (solvent D) based on Qubit, but both were within range of pH adjusted aq. solubility (Table A.1).

Based on ND analysis of TSP, the addition of detergent, both SDS and SDC, improved protein solubility substantially ($P < 0.05$). While SDC facilitated lower solubility (19.5%) of SPE compared to SDS (24.7-27.2 %) (Table A.2), in line with previous studies (Lin *et al.*, 2014), the low content of solubilized protein, evaluated by SDS-PAGE (Fig. A.1, lane 10) was surprising. Nevertheless, the SDC/ABC solvent system was selected for further processing due to compatibility with enzymatic digest and the possibility to selectively precipitate the detergent prior to LC-MS/MS proteomics analysis. This is not possible with e.g. SDS, which is notoriously incompatible with LC (Lin *et al.*, 2008). Moreover, SDC-assisted protein solvation has been reported to improve the number of identified proteins and peptides compared to dilute SDS (Lin *et al.*, 2014); particularly for membrane-associated proteins (Zhou *et al.*, 2006). This could be of particular interest in the characterization of seaweed proteins, where many of the presumably abundant phycobiliproteins (Tandeau De Marsac *et al.*, 2003; Francavilla *et al.*, 2013; Lage-Yusty, Caramés-Adán and López-Hernández, 2013) are reported to be associated with e.g. thylakoid membranes (Hess *et al.*, 1999). Regardless of the method used to quantify TSP, the solubility was determined relative to the total protein determined by total-N and a N-to-protein conversion using the universal Jones Factor (Jones, 1931) of 6.25. Although the Jones factor is routinely used in food protein science and industry, species dependent variations have been determined (Mariotti, Tomé and Mirand, 2008). For seaweeds in general, this has been speculated to be substantially lower and a global seaweed conversion factor of 5 has been proposed (Angell *et al.*, 2016). Moreover, the conversion factor is based on the whole organism, and an extract may not be representative hereof. It was previously demonstrated how N-to-protein can substantially overestimate protein content (Gregersen *et al.*, 2021), and the obtained solubility of SPE here may therefore be substantially higher than what has been determined using both Qubit and ND.

3.1. Proteomics analysis of SPE

Using bottom-up proteomics by LC-MS/MS, we investigated the protein composition of the SPE. Following removal of contaminants and false positive (decoy) hits, we identified 298 protein groups composed of 1,380 potential proteins from the *Gigartinales* protein database, across three technical replicates (Table A.3). The protein groups were identified based on 602 peptide identifications across replicates, following filtering of contaminant and reverse peptides. The vast majority of the peptides (590) were identified with Andromeda scores >40 , indicating an overall high level of confidence (Table A.4). The proportion of Andromeda scores >40 were substantially lower on the protein level, where only 68 of the 298 groups had a score >40 (Table A.3). This can primarily be ascribed to low number of peptides within each group (and thus low sequence coverage), as only 84 of the identified protein groups contained >2 peptides (including razor peptides) and only 59 groups contained >2 unique peptides. Of the 298 identified protein groups, 216 (72%) were identified in all three replicates. 38 (13%) in two of three replicates, 28 (9%) in only one replicate, while 16 (5%) were not quantified (Fig. 2A). Overall, there was good reproducibility between replicates with a Pearson correlation coefficient (PCC) >0.98 based on riBAQ quantification (Fig.2B) and PCC >0.97 based on \log_2 LFQ intensities (Fig 2C). Although the analysis was not performed using a *Gigartina* reference proteome, as such is not currently available, the completeness of the reference proteome for the highly related *Chondrus crispus* (UP000012073), is expected to facilitate sufficient coverage in the applied protein database. The two Rhodophyta species have been co-examined in previous studies showing similar traits (Mathieson and Burns, 1971; Chen *et al.*, 1973; Lipinska *et al.*, 2020) as well as being within close phylogenetic distance, particularly for *Gigartina skottsbergii* (Hommersand, Fredericq and Freshwater, 1994; Chopin, Kerin and Mazerolle, 1999).

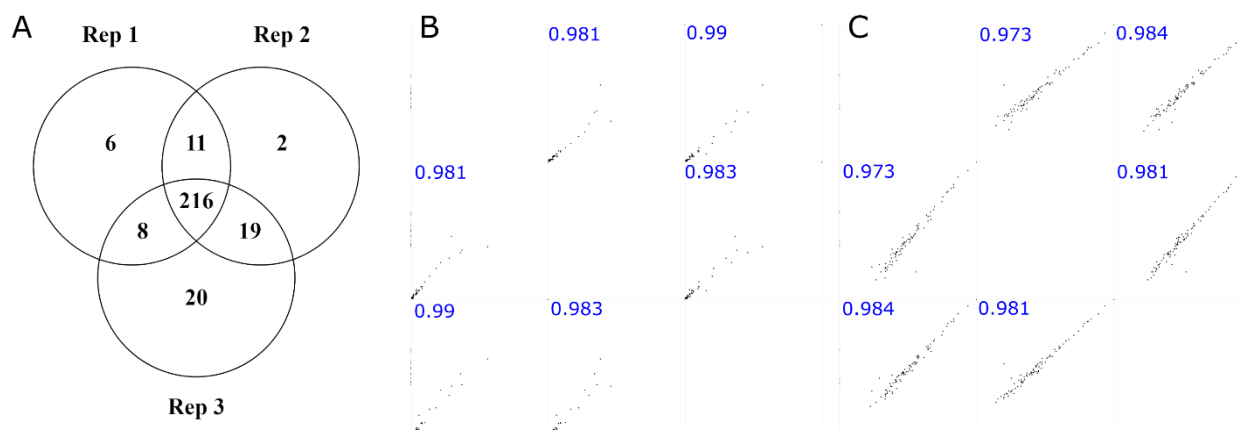


Fig. 2: Reproducibility between technical replicates for LC-MS/MS. Venn diagram showing overlap of identified proteins between sample replicates (A) and scatter plots of quantitative reproducibility using riBAQ (B) and MaxLFQ (C) quantification. For scatter plots, Pearson correlation between individual replicates is indicated in blue.

Eighteen protein groups are considered highly abundant (riBAQ > 1%) and together constitute 79% of the mean, molar protein content in the SPE (Table 1). With the exception of one protein group (M5DD57), the abundant protein groups are found with good quantitative reproducibility between replicates (CV < 23%). Overall, the quantitative reproducibility for the analysis is high, with a stochastic mean CV of 45% and an abundance-weighted mean CV of 15% (Table A.3), illustrating that the higher stochastic variability can primarily be ascribed to lower abundance protein groups.

Among the abundant proteins, representing 79% of the identified protein, phycoerythrin (both subunits) constitute 27.9% of the total protein, while phycocyanin and allophycocyanin constitute 10.0% and 6.3%, respectively (Table 1). RuBisCO accounts for 9.2%, while photosystems proteins and cytochromes constitute 15.6% and 9.9%, respectively. That the majority of proteins are related to light harvest is not surprising, as phycobiliproteins are reported to be highly abundant in red seaweeds (Tandeau De Marsac *et al.*, 2003; Francavilla *et al.*, 2013; Lage-Yusty, Caramés-Adán

and López-Hernández, 2013), and RuBisCO is found abundantly in most autotrophic organisms (Andersson and Backlund, 2008). Moreover, our data incorporate the general understanding that of the phycobiliproteins, phycoerythrin is the most abundant (Glazer, 1994; Tandeau De Marsac *et al.*, 2003; Pina *et al.*, 2014). In addition, phycoerythrin γ -subunit (R7Q9W8) as well as phycobilisome linker-protein (M5DET7) and anchor-proteins (A0A7M3VH69 and M5DBU8) were identified in lower abundance, strengthening that phycobilisomes were successfully extracted (Table A.3). The high abundance of phycobiliprotein subunits, which all have similar molecular weight (Table 1), correspond with the high intensity band observed in SDS-PAGE analysis (Fig. A.1).

Phycobiliproteins are considered heterodimers of the α - and β -subunits assembled into larger complexes within the phycobilisomes (Ritter *et al.*, 1999), but the equimolar stoichiometry is not reflected in our data. Rather, the α -subunit seems to be substantially enriched compared to the β -subunit (Table 1). This is found for all major phycobiliproteins, where the α : β ratio can be determined for phycoerythrin (1.96:1), phycocyanin (2.23:1), and allophycocyanin (5.20:1). While non-equimolar stoichiometry (3:2) has been suggested for allophycocyanin (Reuter, Nickel and Wehrmeyer, 1990), this does not change that α -subunits appear enriched, suggesting some degree of selectivity in the extraction method or different solubilities of the subunits. This is not observed for Ribulose biphosphate carboxylase (RuBisCO), which is known to exist in a heterohexadecamer form but also with equimolar stoichiometry between the large and small subunits (Hauser *et al.*, 2015; Gruber and Feiz, 2018). Here we find the large-to-small ratio for RuBisCO subunits to be 0.91:1 and thus very close to the expected.

Table 1: Summary of proteomics data and subcellular localization for the most abundant (>1%) protein (groups) identified in the *Gigartina* SPE. Protein (groups) are annotated by their lead protein ID (Uniprot AC#), name, molecular weight (MW), relative abundance (riBAQ), reproducibility (CV), number of proteins in the group (PIG), identified peptides, experimental sequence coverage (Seq. Cov., %) as well as subcellular localization as predicted by deepLoc and LocTree in addition to Uniprot-annotated compartment.

Lead Protein ID	Protein Name	MW	riBAQ	CV	PIG	Peptides	Seq. Cov.	DeepLoc	LocTree	Uniprot
I4DEE4	Phycocerythrin alpha chain	17.7	18.4%	8%	5	5	34.8	CYT	NUC	CHLM
I4DEE3	Phycocerythrin beta chain	18.5	9.4%	22%	11	8	42.9	CYT	CYT	CHLM
M5DDH2	Cytochrome b6-f complex subunit 5	3.96	7.3%	10%	9	1	16.2	MIT	MITM	THYM
M5DDK1	Phycocyanin, alpha chain	17.6	6.9%	20%	10	7	45.1	CYT	NUC	CHLM
M5DD55	Photosystem II reaction center protein L	4.36	6.8%	8%	9	2	36.8	GLG	ERM	THYM
A0A6H1U7F2	Photosystem II protein Y	3.82	5.4%	5%	6	1	20.6	MIT	EXT	CHLM
A0A7M3VH79	RuBisCO small subunit	16.1	5.2%	14%	4	4	35.5	CYT	CHL	CHL
A0A6H1U732	Allophycocyanin, alpha subunit	17.5	3.8%	9%	6	7	52.8	CYT	NUC	CHLM
A0A6H1U964	Phycocyanin, beta subunit	18.2	3.1%	11%	6	8	58.1	CYT	NUC	CHLM
Q9TN12	RuBisCO large chain	53.2	2.0%	15%	37	10	23.5	PER	CHL	CHL
M5DD57	Photosystem I reaction center subunit XII	3.30	1.8%	52%	2	1	23.3	EXT	EXT	THYM
M5DDA6	Cytochrome c6	11.5	1.7%	4%	3	1	8.3	EXT	CHL	CHLM
M5DD47	Photosystem I iron-sulfur center	8.77	1.6%	8%	7	2	22.2	EXT	EXT	THYM
M5DDG4	Allophycocyanin, alpha chain	17.4	1.5%	7%	4	6	41.6	CYT	NUC	CHLM
Q1AP03	RuBisCO large subunit (Fragment)	1.57	1.0%	9%	1	1	100	PLA	N/A	CHL
M5DDI6	Allophycocyanin, beta chain	17.6	1.0%	23%	10	6	45.3	CYT	NUC	CHLM
A0A7M1VMM8	RuBisCO large subunit	54.2	1.0%	22%	197	9	20.5	CYT	CHL	CHL
A0A342RZ93	Cytochrome c-550	18.1	1.0%	8%	9	3	16	ER	EXT	THYM

Abbreviations for subcellular localization annotation: Cytosol (CYT), mitochondria (MIT), mitochondrial membrane (MITM), golgi apparatus (GLG), peroxisome (PER), extracellular/secreted (EXT), plastid (PLA), endoplasmic reticulum (ER), endoplasmic reticulum membrane (ERM), nucleus (NUC), chloroplast (CHL), chloroplast membrane (CHLM), thylakoid membrane (THYM).

3.2. Subcellular localization and evaluation of protein extractability

To investigate if the proteomic composition was indeed indicative of intracellular, and thereby efficient, protein extraction, we investigated the subcellular localization of the proteome using both annotated compartments from Uniprot and predictive bioinformatics. From subcellular localization prediction (Table 1), a large discrepancy between prediction algorithms and the annotated compartments in Uniprot is observed for the most abundantly quantified proteins. This may be ascribed to the different architectures and approaches employed in the different models. Nevertheless, as identified proteins are imputed from related seaweeds with proper Uniprot annotation, this is considered a more viable interpretation of the origin of the abundant proteins. As the thylakoid system is not explicitly included in deepLoc, and because the phycobiliproteins have been suggested not to be directly incorporated into the thylakoid membrane, but rather exist in phycobilisomes with larger mobility (Stadnichuk and Tropin, 2017), this may be the reason as to why deepLoc annotates such a large proportion of the protein as cytosolic.

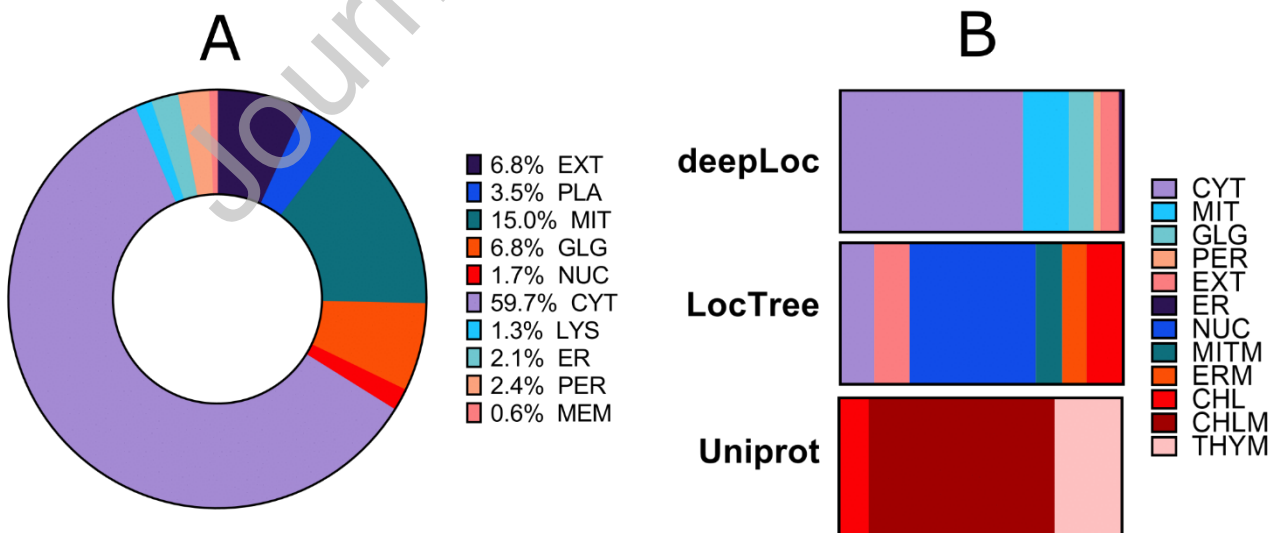


Fig. 3: Subcellular distribution of all identified proteins by deepLoc (A) and abundant (riBAQ > 1%) proteins using different localization prediction algorithms and annotated localization from Uniprot protein homologues (B). Abbreviations for subcellular localization annotation: Cytosol (CYT), mitochondria (MIT), mitochondrial membrane (MITM), golgi apparatus (GLG), peroxisome (PER), extracellular/secreted (EXT), plastid (PLA), endoplasmatic reticulum (ER), endoplasmatic reticulum membrane (ERM), nucleus (NUC), chloroplast (CHL), chloroplast membrane (CHLM), thylakoid membrane (THYM).

Although prediction therefor may be subject to uncertainties, summarizing across all predictions using deepLoc, the amount of predicted extracellular protein account for only 6.8% of the total protein identified (Fig. 3.A). Moreover, looking only at the most abundant proteins (Fig.3.B), extracellular proteins constitute only a minor part of the extracted proteins according to both deepLoc (5.1%) and LocTree (9.8%) while none are annotated as extracellular or secreted in Uniprot. In a previous study of extracts from *E. denticulatum*, a similar extraction principle was employed, where the soluble fraction following high temperature aq. carrageenan extraction was characterized (Gregersen *et al.*, 2022). In that study, the soluble fraction comprised almost exclusively (97%) extracellular protein. This is in agreement with our findings, where extracellular protein content in the residual solid fraction is depleted and the SPE is constituted almost exclusively by non-extracellular protein regardless of considering subcellular prediction or annotated compartments in Uniprot. Ultimately, this indicates that the SPE is a good source of extractable cellular protein. In particular, phycobiliproteins, RuBisCO, and other light harvest-related proteins.

3.3. Amino acid distribution and nutritional potential

To assess nutritional potential of the SPE, protein- and peptide-level data was used to compute the relative abundance of individual amino acids based on the mean riBAQ and mean relative MS1 peptide intensity, as previously described (Jafarpour, Gomes, *et al.*, 2020; Gregersen *et al.*, 2021, 2022).

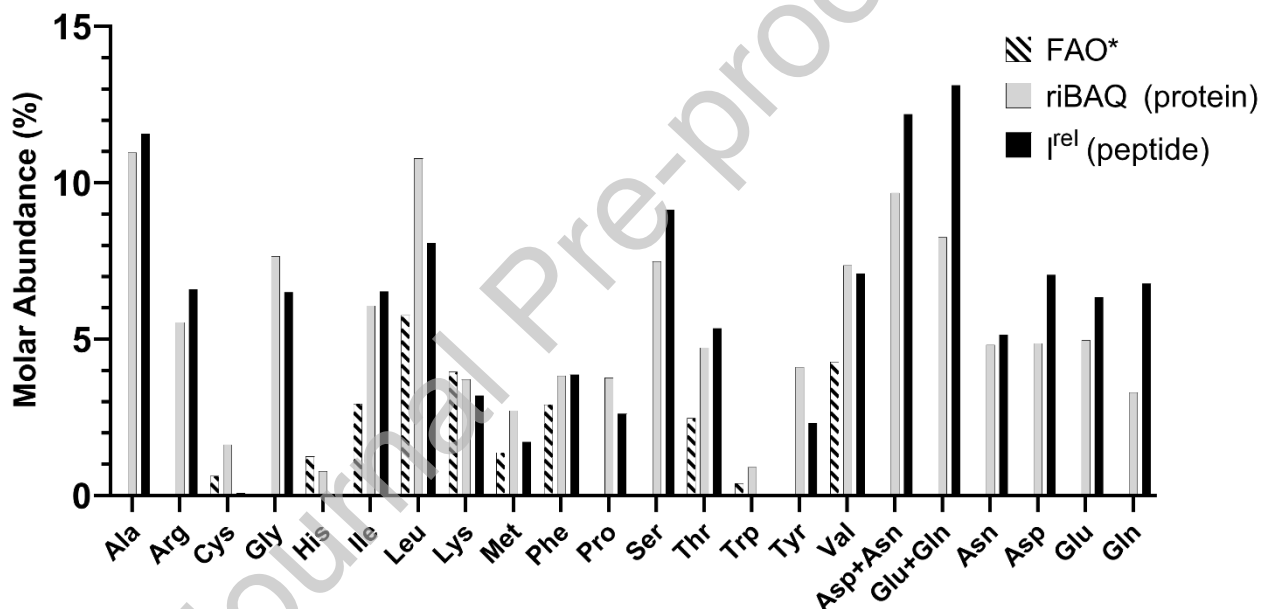


Fig 4: Estimation of amino acid (AA) composition (as relative molar abundance) using peptide- and protein-level MS data for SPE. For reference, minimum requirements according to the FAO* (WHO/FAO/UNU, 2007) are indicated.

Overall, we found good agreement between the use of protein- and peptide-level data for estimating AA composition (Fig. 4). Moreover, we find that, with the exception of His, the SPE contains

sufficient EAA levels relative to the minimum requirements by the FAO (WHO/FAO/UNU, 2007). The calculated EAA:NEAA ratio for SPE is 0.69 and 0.56 based on protein-level and peptide-level data, respectively. This is in line with red seaweeds in general (0.63), but still somewhat lower than other alternative protein sources (0.78-1.09), thereby decreasing the nutritional potential (Mæhre *et al.*, 2014; Donadelli *et al.*, 2019). The SPE contains high levels of branched chain AAs (BCAAs), namely Leu, Ile, and Val, corresponding to 24.2% and 21.7% based on protein- and peptide-level data, respectively. High levels of individual EAAs, in particular the BCAAs, is highly desired in fields such as sports nutrition (Campbell *et al.*, 2007). Moreover, BCAAs are linked with decreased food intake (Laeger *et al.*, 2014) and linked to numerous metabolic and physiological functions such as mammary and embryonic health, intestinal development, and immune response regulation (Zhang *et al.*, 2017), and SPE may therefore also be relevant in pregnancy and infant nutrition.

Although the chemical composition of *Gigartina* has been reported not to vary greatly over season (Amimi *et al.*, 2007), the seasonal variation of the proteomic composition has not been investigated. Based on investigations of other red seaweeds such as *Palmaria palmata*, substantial seasonal variation in both protein content and composition are likely (Fleurence, 1999). Quantitative variations of the *Gigartina* proteome will also affect the AA distribution, and should therefore be investigated if moving forward with the seaweed as a source of food protein. As MS-based proteomic determination of AA distribution is still in its infancy and thus related to some uncertainty (Jafarpour, Gomes, *et al.*, 2020; Gregersen *et al.*, 2021, 2022), supporting the analysis with more conventional AA analysis is advisable.

3.4. Abundant *Gigartina* proteins as source of emulsifier peptides

While the *Gigartina* SPE displays some potential as food protein, the high content of phycobiliproteins may also directly imply other potential applications. Phycobiliproteins are membrane-associated (Hess *et al.*, 1999) and furthermore have a very high content of α -helical domains (Apt, Collier and Grossman, 1995). Such proteins are known to have highly amphiphilic structural features (Fernández-Vidal *et al.*, 2007), and previous studies have shown that helical and amphiphilic peptides have enormous potential as natural emulsifiers (Malcolm, Dexter and Middelberg, 2007; Dexter and Middelberg, 2008; García-Moreno *et al.*, 2021; Yesiltas *et al.*, 2021). While RuBisCO is primarily localized in the chloroplast stroma (Gruber and Feiz, 2018), it has been shown to possess emulsifying properties in the native form (Di Stefano *et al.*, 2018; Tan *et al.*, 2022). To investigate this potential further, we applied the *in silico* predictive tool, EmulsiPred (García-Moreno, Gregersen, *et al.*, 2020; García-Moreno, Jacobsen, *et al.*, 2020) on the highly abundant SPE proteins.

Table 2: Summary of bioinformatic prediction of emulsifier peptides in abundant (riBAQ > 1%) SPE proteins. Each protein (group) is annotated with the lead protein ID (Uniprot AC#) and name along with the relative molar abundance in SPE (riBAQ). For each protein (group), the number of potential (score > 2) emulsifier peptides with facial (α and β) or axial (γ) amphiphilicity, as well as total potential emulsifier peptides are indicated. For very high abundance (riBAQ > 5%), the highest scoring peptides within each class (two for α and γ , one for β) are shown by score, type, and sequence.

Lead Protein ID	Protein Name	riBAQ	Predicted peptides				Selected highest scoring peptide		
			α	β	γ	Total	Score	Type	Sequence
I4DEE4	Phycocerythrin alpha chain	18.4%	354	0	214	568	3.90	α	EKVNKCYRDIDHYMRLINYAL
I4DEE3	Phycocerythrin beta chain	9.4%	214	0	460	674			
M5DDH2	Cytochrome b6-f complex subunit 5	7.3%	3	0	183	186	5.16	γ	IVLGLIPVTLAGLLVAAYLQYRRGNQ
M5DDK1	Phycocyanin, alpha chain	6.9%	458	0	66	524	3.88	α	EAAKSLTNNAQRLITGAAQAV
M5DD55	Photosystem II reaction center protein L	6.8%	0	0	293	293	5.84	γ	NPNKEPVELNRTSLFWGLLLIFVLAVLF
A0A6HIU7F2	Photosystem II protein Y	5.4%	57	0	41	98			
A0A7M3VH79	RuBisCO small subunit	5.2%	20	82	588	690	3.95	β	NVYIKIN
A0A6HIU732	Allophycocyanin, alpha subunit	3.8%	162	0	72	234			
A0A6HIU964	Phycocyanin, beta subunit	3.1%	175	0	190	365			
Q9TN12	RuBisCO large chain	2.0%	467	29	1052	1548			
M5DD57	Photosystem I reaction center subunit XII	1.8%	0	0	122	122			
M5DDA6	Cytochrome c6	1.7%	236	0	163	399			
M5DD47	Photosystem I iron-sulfur center	1.6%	0	36	34	70			
M5DDG4	Allophycocyanin, alpha chain	1.5%	266	0	141	407			
Q1AP03	RuBisCO large subunit (Fragment)	1.0%	0	0	0	0			
M5DDI6	Allophycocyanin, beta chain	1.0%	143	0	120	263			
A0A7M1VMM8	RuBisCO large subunit	1.0%	430	30	1113	1573			
A0A342RZ93	Cytochrome c-550	1.0%	139	0	456	595			
	Multiple target proteins		500	25	586	1111			

For the most abundant SPE proteins (Table 1), we predicted a total of 7,684 potential emulsifier peptides (score > 2), whereof 1,111 map to multiple target proteins (Table 2). Not surprisingly, the majority of predicted emulsifier peptides are found in α -helical or random (γ) conformation, as light harvest proteins are highly helical by nature. The highest amount of emulsifier peptides (>3000) were predicted from the large subunit of RuBisCO (Q9TN12 and A0A7M1VMM8), with the majority being in γ conformation although a substantial amount were also predicted in helical conformation (ratio γ : α ~2:1). The large number of RuBisCO-derived peptides should also be considered in the light that this protein is also approximately three times the size of phycobiliprotein subunits. Phycobiliproteins were predicted to contain 263-674 potential emulsifier peptides (ratio γ : α ~1:1) per subunit, with the lowest amount found in allophycocyanin, which is also the lowest abundance phycobiliprotein. Photosystem-related proteins and cytochromes are generally enriched in γ -peptides. To investigate high potential emulsifier peptides with a potentially high downstream yield, we focused only on very high abundance (riBAQ > 5%) proteins (Table 2). Looking at the highest scoring α -peptides (Table A.5), specific regions of the α -subunit in phycoerythrin (I4DEE4) and phycocyanin (M5DDK1) stand out (scores 3.6-3.9). For β - peptides, the small RuBisCO subunit (A0A7M3VH79) shows a particularly interesting domain with multiple peptides having scores in the range 3.1-4.0 (Table A.6). In the case of γ -peptides (Table A.7), particular regions from Photosystem II reaction center protein L (M5DD55) and Cytochrome b6-f complex subunit 5 (M5DDH2) are predicted to contain many peptide isoforms of very high emulsification potential (scores 4.5-5.8). Using EmulsiPred, γ -peptides generally have higher prediction scores (García-Moreno, Gregersen, *et al.*, 2020; Jafarpour, Gregersen, *et al.*, 2020; Yesiltas *et al.*, 2021; Gregersen Echers *et al.*, 2022). This can be attributed to the axial mode of emulsification compared to α - and β -peptides, which exert emulsification through facial amphiphilicity, thereby making the score difficult to compare directly. Compared to previous studies using EmulsiPred prediction (with the

same version) and peptide-level *in vitro* functional validation, these scores indicate that the SPE could indeed have very high potential for production of functional, protein-based ingredients (Yesiltas *et al.*, 2021). As all these proteins also have very high abundance (riBAQ > 5.2%), they could serve as sources for release of emulsifier peptides in high total yields.

To better understand and evaluate if predicted peptides are indeed potential emulsifiers, we applied templated homology-modelling of the native protein structure. The conformation of particularly amphiphilic α -peptides within the native protein, has previously been shown to be a good model for oil/water interfacial structure (García-Moreno *et al.*, 2021; Yesiltas *et al.*, 2021). Modelling was performed for the five proteins indicated above to contain particularly high scoring peptides (Fig. 5) yielding high quality models for all protein (Table A.8).

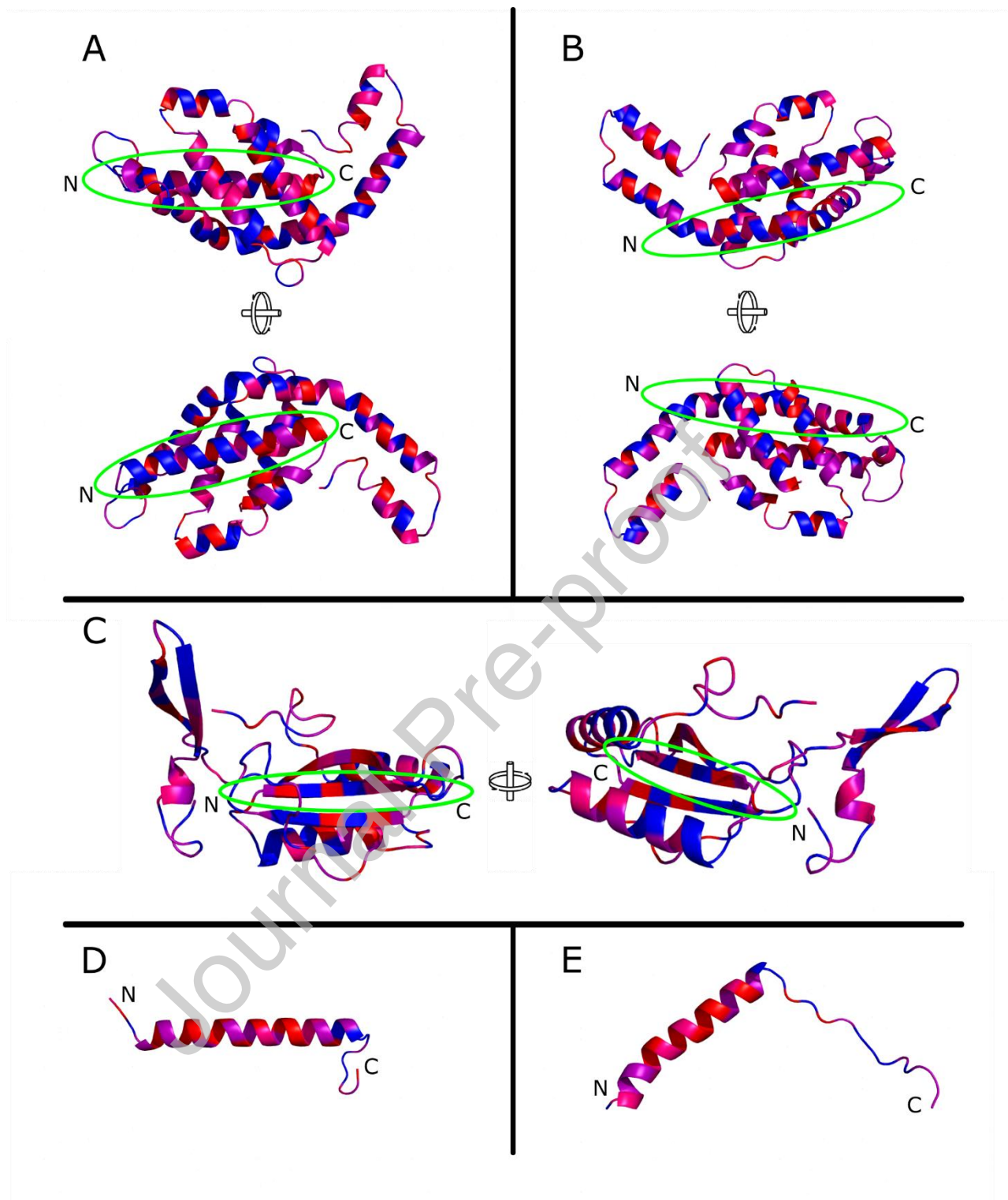


Fig. 5: Structural modelling of proteins embedding particularly high scoring emulsifying peptides.

A) Phycoerythrin alpha subunit, B) Phycocyanin alpha subunit, C) RuBisCO small subunit, D) Photosystem II reaction center protein L, and E) Cytochrome b6-f complex subunit. For structures

A-C, the region containing the highest scoring predicted emulsifier peptides has been highlighted and the N-, and C-termini of the regions are indicated. For structures D and E, predicted emulsifier peptides span the entire (short) length and the N- and C-termini are indicated for the full length proteins.

For phycoerythrin (Fig. 5A), the region containing high scoring peptides is indeed also a highly amphiphilic, surface-exposed helix in the native protein, underlining the emulsifying potential of this protein region. Moreover, the C-terminus of the region is highly hydrophobic, which has been speculated to improve the emulsification potential on amphipathic helices by serving as an anchor for the oil-phase (Gregersen Echers *et al.*, 2022). In Phycocyanin (Fig. 5B), a highly amphiphilic, surface-exposed helix is also found within the region of interest. Moreover, the region includes a kink within the helix, which is known to play a role in e.g. membrane interactions of antimicrobial peptides (Tuerkova *et al.*, 2020) and have been proposed to also facilitate improved interfacial stabilization in emulsions (García-Moreno, Jacobsen, *et al.*, 2020). For the small RuBisCo subunit (Fig. 5C), the structure consists of more β -strand, why the large number of β -peptides here from could be expected. High scoring (>3) β -peptides are rather short (7-12 AAs), which may be insufficient to produce a stable emulsion (García-Moreno, Gregersen, *et al.*, 2020). Photosystem II reaction center protein L and Cytochrome b6-f complex subunit 5 (Fig. 5D, 5E) are both short, helical proteins involved in larger quaternary complexes. Nevertheless, both were predicted to contain high scoring γ -peptides spanning all of the sequence length. γ -peptides represent axial amphiphilicity, and thus their secondary structure is not crucial for functionality (García-Moreno, Gregersen, *et al.*, 2020; García-Moreno, Jacobsen, *et al.*, 2020), although helical structure in γ -peptides has been seen to facilitate formation of highly stable emulsions (García-Moreno *et al.*, 2021; Yesiltas *et al.*, 2021). Moreover, both proteins are annotated to contain transmembrane

helices (UniProt Consortium *et al.*, 2021), indicating that the hydrophobic helix is indeed capable of inserting into a lipid phase. The *in vitro* functionality of the peptides should be further investigated in model emulsion systems to fully determine their potential. Recently, we presented a workflow, where using a basis of quantitative proteomics and bioinformatics, we designed a targeted hydrolysis process where we were able to release emulsifier peptides from potato protein (Gregersen Echers *et al.*, 2022). Such an approach could be valuable in exploring if the *Gigartina* SPE is indeed a good source of emulsifying peptides, releasable through enzymatic hydrolysis. As high scoring peptides were not only restricted to the modelled proteins, other high abundance proteins may also contribute to overall emulsifying properties of a hydrolysate by release of amphiphilic peptides. For instance, RuBisCO large chain had both high scoring α -peptides (3.37) and γ -peptides (4.68) (Tables 2, A.6, A.8), which should also be investigated further. As the cumulative abundance of RuBisCO large chain isoforms approaches 5% (Table A.3), this subunit may also be a source of potential emulsifier peptides with high yield.

4. Conclusion

Seaweeds, in particular Rhodophytae, are massively exploited for industrial hydrocolloid production. Side-streams from this production remains underutilized and may be used as a source of food protein and functional ingredients. In pilot scale, a seaweed protein extract (SPE) from *Gigartina radula* was isolated and investigated for its protein composition and quality. The aqueous solubility of SPE was very low (<2%), but did still display pH-dependent solubility, which could be related to the conditions employed during seaweed processing and protein extraction. The addition of detergent improved protein solubility, allowing for visualization by SDS-PAGE and bottom-up shotgun proteomics analysis by LC-MS/MS. Using all known proteins within the *Gigartinales* order

as reference database, 298 protein groups were identified and through relative quantification, the most abundant proteins were determined. In general, proteins related to light harvest dominate the SPE and particularly phycobiliproteins (44%) and RuBisCO (9%) were abundant. Within the phycobiliproteins, phycoerythrin constituted the majority of the quantified protein (28%). The expected equimolar stoichiometry of α - and β -subunits of phycobiliproteins was not observed (α -subunit favored), indicating a potential bias in the extraction, whereas the large and small subunits of RuBisCO were identified in equimolar stoichiometry as expected. Investigating the subcellular localization of identified proteins, the extraction method was deemed sufficient for release of intracellular protein. The SPE generally showed sufficient levels (36-41%) of essential amino acids (with the exception of His), indicating that it may be of potential use as a nutritional food protein, although an EAA:NEAA ratio of 0.56-0.69 is somewhat lower than other alternative proteins. High levels of branched chain amino acids (22-24%) indicate potential relevance in e.g. sports and infant nutrition. From the most abundant proteins in the SPE, almost 8,000 potential emulsifier peptides were predicted, including many very high potential peptides based on amphiphilic score. Through templated homology modelling of the native protein structure, this potential was verified as the peptides also have similar amphiphilic structure within the native proteins. With this study, we provide fundamental new knowledge within seaweed protein science and the first study of the *Gigartina* proteome. Moreover, we expect that *Gigartina* could be a valuable source of protein-derived natural emulsifiers for foods and this route should be explored further, also investigating the functional properties of the crude extract for comparison.

Data Availability

The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium (<http://proteomecentral.proteomexchange.org>) via the PRIDE partner repository (Perez-Riverol *et al.*, 2019) with the dataset identifier PXD034435 and DOI 10.6019/PXD034435. Data not supplied in PRIDE or appended as supplementary information, may be obtained upon request to the corresponding author.

CRedit Author Contribution

Simon Gregersen Echers: Conceptualization, Methodology, Investigation, Validation, Formal analysis, Data Curation, Writing - Original Draft, Writing - Review & Editing, Visualization, Supervision. **Naim Abdul-Khalek:** Investigation, Formal analysis, Writing - Review & Editing. **Rasmus Kranold Mikkelsen:** Formal analysis, Writing - Review & Editing. **Susan Løvstad Holdt:** Conceptualization, Resources, Writing - Review & Editing. **Charlotte Jacobsen:** Conceptualization, Resources, Writing - Review & Editing, Funding acquisition. **Egon Bech Hansen:** Conceptualization, Resources, Writing - Review & Editing, Funding acquisition. **Tobias Hegelund Olsen:** Methodology, Investigation, Formal analysis, Data Curation, Writing - Review & Editing. **Jimmy J. P. Sejberg:** Methodology, Investigation, Validation, Formal analysis, Resources, Writing - Review & Editing. **Michael Toft Overgaard:** Conceptualization, Resources, Writing - Review & Editing, Funding acquisition, Supervision.

Acknowledgements

The authors would like to thank CP Kelco (Lille Skensved, Denmark) and in particular, Laboratory Technician Helle Bech Olsen, for supplying the seaweed protein extract.

Funding

This work was supported by Innovation Fund Denmark (Grant number 7045-00021B (PROVIDE)).

Conflicts of Interest

Jimmy J. P. Sejberg is employed by CP Kelco ApS. All other authors declare no conflicts of interest.

Ethical Statement

There are no ethical aspects applicable for the study

Conflicts of Interest

JS is employed by CP Kelco ApS. All other authors declare no conflicts of interest.

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