

Gross Chemical Profile and Calculation of Nitrogen-to-Protein Conversion Factors for Five Tropical Seaweeds

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ABSTRACT

Despite decades of research on marine algae, there are still significant gaps in basic knowledge about chemical composition of these organisms, especially in tropical environments. In this study, the amino acid composition and contents of total nitrogen, phosphorus, lipid, carbohydrate and protein were determined in Asparagopsis taxiformis, Centroceras clavulatum, Chaetomorpha aerea, Sargassum filipendula and Spyridia hypnoides. The seaweeds showed low lipid contents (lower than 5.5% d.w. in all species) and were rich in carbohydrates (more than 16% d.w. in all seaweeds). The percentage of nitrogen, phosphorus and protein varied widely among species, which red algae showed the highest concentrations. The amino acid composition was similar among the seaweeds, which glutamic acid, aspartic acid and leucine as the most abundant. All species are poor in histidine. An average of 24.2% of the total nitrogen is non-proteinaceous. From data of total amino acid and total nitrogen, specific nitrogen-to-protein conversion factors were calculated for each species. The nitrogen-to-protein conversion factors calculated ranged from 4.51 to 5.21, with an overall average of 4.86. These findings show that the traditional conversion factor of 6.25 should be avoided for seaweeds, since it overestimates the actual protein content.

Keywords: Nitrogen-to-Protein Conversion Factors, Protein, Amino Acid, Seaweeds, Element Composition, Tropical Environment

1. Introduction

Utilization of algae has increased considerably over the past years as a consequence of growth of research in various fields [1]. Because seaweed species are rich in beneficial nutrients, in countries such as China, Japan and Korea, they have been commonly utilized in human nutrition for centuries [2]. They have been found to be good sources of vitamins, carbohydrates, minerals and proteins [3], but show wide variations among species. Seaweeds have been increasingly viewed as potential sources of bioactive compounds with immense pharmaceutical, biomedical and nutraceutical importance [4]. Moreover, they have been used in agricultural and industrial research due to their high content of carbohydrates, proteins, vitamins, and minerals [5], but show wide variations among species. However, data on the bioavailability of these components are limited.

Protein data of marine algae presents many applications, involving both basic and applied research. However, comparisons of protein content among species are difficult because of methodological differences [6,7]. Extraction is one of the main problems in algal protein analysis, which is performed with variable efficiency by different methods [8]. Differences in algae cell wall composition and in procedures used for protein extraction establish remarkable influence on final results [9].

The most common methods used for protein determination in algae, Lowry's method [10] and Bradford method [11] assays, are subject to interferences from many factors [12], which are independent of the problems related to the protein extraction. The interferences are a consequence of the effects of some substances on specific amino acids, since that the chemical reactions which produce the protein quantification depends on the reactivity of the amino acid side groups [13].

By contrast, total nitrogen analysis is relatively simple and easy to perform, and nitrogen-to-protein conversion factors (N-Prot factors) can be used to estimate crude protein content. The use of N-Prot factors to determine protein content has some important advantages if compared to other methodologies. Total nitrogen analysis, carried out by Kjeldhal's method [14], Hach techniques [15] or CHN analysis, eliminates the necessity of extracting the protein content of the sample to be analyzed, the major problem in protein analysis of algae [8]. Barbarino and Lourenço (2005) [7] showed that algal proteins associated with cell membranes are hardly extracted, confirming the difficulty of reproducing extraction figures and increasing differences in values found with different methods. Thus, the use of N-Prot factors also allows better comparisons of results among researchers, since protein is estimated without a tricky previous extraction.

The use of specific N-Prot factors is widely recommended in order to get more accurate estimates of protein content [16]. The nitrogen:protein ratio does vary according to the source considered [17]. The use of N-Prot factors is particularly wide in food science. Except for a list of specific N-Prot factors available for certain cereals (e.g. 5.26 for rise, 5.47 for wheat; [18]), legumes (e.g. 4.75 - 5.87 for cassava root; [19]), mushroom (4.70; [20]), Cheddar cheese (6.38; [21]) and milk (5.94, [22]) among other products, the factor 6.25 calculated by reference [23] is still used for most plant and animal sources. The use of the traditional factor 6.25 is based on the assumption that samples contain protein with 16% nitrogen and an insignificant amount of non-protein nitrogen (NPN) [24]. However, the amino acid composition varies from one protein source to another, existing different N content in each amino acid. Moreover, this assumption is invalid for organisms that contain high concentrations of other nitrogenous compounds, such as nucleic acids, amines, urea, inorganic intracellular nitrogen (ammonium, nitrate and nitrite), vitamins and alkaloids [25].

Plant materials, fungi and algae commonly show significant amounts of NPN [7,20,26]. In addition, it is common to find plant materials showing total protein with less than 16% nitrogen in total amino acid [27]. The same trends may be applied to the nitrogen distribution in seaweeds, and the use of the factor 6.25 tends to overestimate the protein data [28,29]. Despite this, several authors continue to use the factor 6.25 to estimate seaweed protein content (e.g. [30-33]).

To compensate the influence of NPN, specific N-Prot factors must be calculated. Specific N-Prot factors have already been proposed for 12 marine microalgae [34], with an overall average N-Prot factor of 4.78. Studies in this field are needed for seaweeds, since very limited information is available (e.g. [28,29]).

In a broader sense, data on chemical composition of seaweeds are predominantly obtained with species from temperate (e.g. [28,35,36]), warm temperate (e.g. [37,38]) and subtropical coastal environments (e.g. [39-41]). By comparison, information on chemical composition of algae from tropical environments is relatively scarce [42-44] and more data are needed from those regions. Compared with land plants, the chemical composition of seaweeds has been poorly investigated and most of the available information only deals with traditional edible seaweeds [32,45,46].

The purpose of our study was to determine specific N-Prot factors for five tropical marine seaweeds, based on the ratio of amino acid composition to total nitrogen (TN) content. In addition, we also characterized and compared the seaweed species regarding hydrosoluble protein, carbohydrate, lipid, nitrogen and phosphorus contents.

2. Materials and Methods

2.1. Algae

In this study five macroalgae species were analyzed. The identification of the species was carried out following the checklist of reference [47] and with experts' supervision. Chlorophyta: *Chaetomorpha aerea* (Dillwyn) Kützing; Rhodophyta: *Asparagopsis taxiformis* (Delile) Trevisan de Saint-Léon; *Spyridia hypnoides* (Bory de Saint-Vincent) Papenfuss; and *Centroceras clavulatum* (C. Agardh) Montagne; Ochrophyta: *Sargassum filipendula* (Agardh).

2.2. Sampling

C. aerea, C. clavulatum and S. filipendula were collected in June 2007 and S. hypnoides was collected in September 2007 at Arraial do Cabo (22°57'S 42°01'W). A. taxiformis was sampled in June 2007 at Angra dos Reis (23°00'S 44°19'W). Both sites are located in Rio de Janeiro State, southeastern Brazil (Figure 1) and show oligotrophic characteristics and minor anthropic influence. Whole thalli of adult plants were collected and washed in the field with local seawater in order to remove epiphytes, sediment and organic matter. Plants were packed in plastic bags and kept on ice until returned to the laboratory (ca. 150 km, Figure 1). In the laboratory samples were gently brushed under running seawater, rinsed with distilled water, dried with paper tissue and frozen at -18° C. Subsequently, the samples were freeze dried in a Terroni Fauvel, model LB1500TT device. The dried material was powdered manually using a mortar and pestle, and it was kept in desiccators containing silica-gel, under vacuum at room temperature, until the chemical analyses were carried out.

2.3. Tissue Analysis

The Lowry's method [10] was used to analyze hydro-

soluble protein in the samples, with bovine serum albumin as a protein standard. Spectrophotometric determinations were performed at 750 nm, 35 min after the start of the chemical reaction.

Total carbohydrate was extracted with 80% H₂SO₄, according to reference [48]. The carbohydrate concentration was determined spectrophotometrically at 485 nm, 30 min after the start of the chemical reaction, by the phenol-sulfuric acid method [49], using glucose as a standard.

Total lipid was extracted according to reference [50], and determined gravimetrically after solvent (chloroform) evaporation.

Total nitrogen and phosphorus were determined in algal tissue after peroxymonosulphuric acid digestion, using a Hach digestor (Digesdhal[®], Hach Co.) [15]. Samples were digested with concentrated sulfuric acid at 440°C and treated with 30% hydrogen peroxide. Total nitrogen and phosphorus contents in the samples were determined spectrophotometrically after specific chemical reactions. See reference [51] for analytical details.

Total amino acid was determined by high performance liquid chromatography with pre-column derivatization with AccQ.Fluor[®] reagent (6-aminoquinolyl-*N*-hydroxysuccinimidyl carbamate), reverse phase column C₁₈ AccQ.Tag[®] Nova-Pak (150 × 3.9 mm; 4 µm), ternary mobile phase in gradient elution composed by sodium acetate 140 mM + TEA 17 mM pH 5.05 (solvent A), acetonitrile (solvent B) and water (solvent C), flow 1 ml·min⁻¹ [52]. A Waters, model Alliance 2695 chromatograph was used, equipped with a fluorescence detector Waters[®] 2475 ($\lambda_{ex.}$ 250 nm, $\lambda_{em.}$ 395 nm). Analytical conditions were suitable to determine all amino acids, except tryptophan, cysteine + cistine and methionine. The percent of nitrogen in each amino acid was used to calculate nitrogen recovered from total amino acid analysis. Aspartic acid, threonine, serine, glutamic acid, proline, glycine, alanine, valine, isoleucine, leucine, tyrosine, phenylalanine, histidine, lysine, and arginine contents were multiplied by 0.106, 0.118, 0.134, 0.096, 0.123, 0.188, 0.158, 0.120, 0.108, 0.108, 0.078, 0.085, 0.271, 0.193, and 0.322, respectively [16].

2.4. Calculation of N-Prot Factors

N-Prot factors were determined for each species by the ratio of amino acid residues (AA-Res) to total nitrogen (TN) of the sample: N-Prot factor = AA-Res/TN. Thus, for a 100 g (dry weight) sample having 16.21 g of amino acid residues and 3.48 g of TN, an N-Prot factor of 4.66 is calculated.

The amino acid residues of the samples was calculated by summing up the amino acid masses retrieved after acid hydrolysis (total amino acids), minus the water mass (18 H_2O/mol of amino acid) incorporated into each amino acid after the disruption of the peptide bonds [53].

2.5. Statistical Analysis

The results were analyzed by one-way analysis of variance (ANOVA) with significance level $\alpha = 0.05$ [54] followed, where applicable, with a Tukey's multiple comparison test.

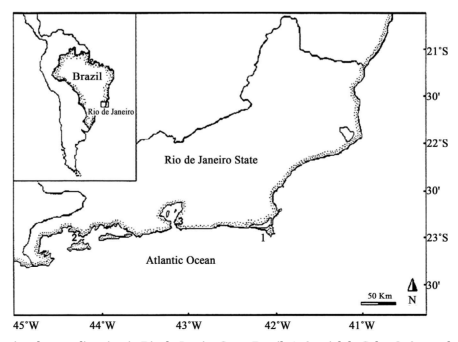


Figure 1. Map showing the sampling sites in Rio de Janeiro State, Brazil: 1. Arraial do Cabo; 2. Angra dos Reis. Location of the laboratory at Fluminense Federal University is indicated as "3".

3. Results

The hydrosoluble protein contents ranged from 8.7% (S. filipendula, brown alga) to 16.1% (C. aerea, green alga) of the dry weight with intermediate and similar concentrations (p > 0.05) in red algae (**Table 1**). Carbohydrates were the most abundant substances measured in all species, ranged from 16.8% (S. filipendula) to 29.4% (C. aerea) of the d.w. The values tended to be higher and similar in green and red algae, except for the rhodophyte A. taxiformis, which was significantly lower than the other three species (p < 0.001). All species studied contained low concentration of total lipid. The highest value was recorded in C. aerea (5.5%, d.w.) and the lowest concentration was found in C. clavulatum (2.8%). The red algae S. hypnoides and A. taxiformis showed significantly higher lipid concentration than C. clavulatum (p < 0.001) (Table 1).

The percentage of nitrogen showed wide variations among species, ranging from 1.75% (S. filipendula) to 5.56% (A. taxiformis) of the dry weight. Red algae showed higher total nitrogen concentrations in the thalli, with significant differences (p < 0.001) to others groups (Table 1). The green alga C. aerea showed an intermediate concentration of N in comparison to red and brown algae. The concentrations of phosphorus also varied widely among species. C. clavulatum showed the highest value (0.54%, p < 0.001) and S. filipendula and A. taxiformis showed the lowest concentrations (0.27% and 0.30%, respectively) (p > 0.05). The green C. aerea and red algae S. hypnoides showed intermediate and similar values (Table 1). The tissue N:P ratios were low for all species (N:P < 11:1), except for A. taxiformis, which showed the highest N:P ratio (18.9:1, p < 0.001), significantly higher than the other red algae. The lowest N:P ratios were recorded in the green and brown algae analyzed (5.83:1 and 6.56:1, respectively).

The amino acid profiles of seaweeds samples are pre-

sented in Table 2. Glutamic acid was the most abundant amino acid in all species studied. The highest concentration of glutamic acid (16.3% of total amino acids) was found in S. filipendula, while A. taxiformis had the lowest (10.3%) concentrations. Aspartic acid was the second most abundant amino acid in seaweeds. These values varied from 9.59% (A. taxiformis) to 12.7% (C. aerea). The percentage of histidine was the lowest in all species, and only the brown algae S. filipendula achieved values close to 2%. The red alga A. taxiformis showed higher concentrations of valine, phenylalanine and arginine than the others species studied. The highest concentrations of tyrosine were observed in the red algae S. hypnoides and C. clavulatum with values close to 5%. Percentages of leucine and threonine were similar among all species and the lowest concentrations of glycine were observed in red algae S. hypnoides.

The total protein content of the samples is showed in **Table 3** as total amino acid residues. The seaweeds showed a wide range of total protein concentration, varying from 8.62% (*S. filipendula*) to 25.1% (*A. taxiformis*) of the d.w. The red algae recorded the highest values of total protein. Nitrogen mass within total amino acid ranged from 1.36% (*S. filipendula*) to 4.14% (*A. taxiformis*). The relative percentage of protein nitrogen was estimated as the ratio of nitrogen recovered from amino acid to total nitrogen (**Table 1**). Protein nitrogen ranged from 69.5% (*C. clavulatum*) to 81% (*S. hypnoides*) and the red algae tended to show higher percentages of NPN, except *S. hypnoides*.

From the ratio of the mass of amino acid residues to total nitrogen we calculated specific N-Prot factors for the seaweeds. The N-Prot factors ranged between 4.51 (*A. taxiformis*) to 5.21 (*S. hypnoides*). The others three species recorded intermediate values of N-Prot factors. An overall average N-Prot factor = 4.86 was calculated from the data for all species.

Species	Hydrosoluble protein	Total carbohydrate	Total lipid	Total nitrogen	Total phosphorus	N:P ration (by atoms)
	***	***	***	***	***	***
Asparagopsis taxiformis	$11.7\pm0.58^{\text{b}}$	$22.9\pm1.35^{\text{b}}$	$4.80\pm0.24^{\text{b}}$	5.56 ± 0.29^{a}	$0.30\pm0.04^{\rm c}$	$18.9\pm1.66^{\rm a}$
Centroceras clavulatum	$11.3\pm0.64^{\text{b}}$	$27.1\pm1.73^{\rm a}$	$2.78\pm0.23^{\text{d}}$	$4.63\pm0.15^{\text{b}}$	0.54 ± 0.03^{a}	8.55 ± 0.22^{b}
Chaetomorpha aerea	16.1 ± 0.25^{a}	$29.4\pm0.78^{\rm a}$	5.49 ± 0.09^{a}	$2.56\pm0.13^{\text{d}}$	0.43 ± 0.04^{b}	$5.83\pm0.16^{\rm c}$
Sargassum filipendula	$8.72\pm0.54^{\rm c}$	$16.8\pm0.97^{\rm c}$	$2.92\pm0.13^{\text{d}}$	$1.75\pm0.03^{\text{e}}$	$0.27\pm0.02^{\rm c}$	$6.56\pm0.41^{\circ}$
Spyridia hypnoides	$10.7\pm0.92^{\text{b}}$	$27.7 \pm 1.47^{\rm a}$	$4.20\pm0.36^{\rm c}$	$3.98\pm0.14^{\rm c}$	0.39 ± 0.02^{b}	10.3 ± 0.85^{b}

Table 1. Gross chemical composition of five species of seaweeds sampled in a tropical site of Brazil. Values are expressed as percentage of the dry mass and represent the mean of four replicates \pm standard deviation (n = 4)[#].

[#]Mean values significantly different: ***p < 0.001, a > b > c > d > e. Identical superscript letters (a, a; b, b) or absence of letters indicate that mean values are not significantly different.

Amino acid	Asparagopsis taxiformis	Centroceras clavulatum	Chaetomorpha aerea	Sargassum filipendula	Spyridia hypnoides
Aspartic acid	9.59 ± 0.82	11.1 ± 0.49	12.7 ± 1.65	11.3 ± 0.26	11.8 ± 0.19
Threonine	5.33 ± 0.24	5.24 ± 0.11	4.89 ± 0.09	4.76 ± 0.08	4.73 ± 0.02
Serine	5.46 ± 0.29	5.15 ± 0.06	4.43 ± 0.05	4.54 ± 0.06	5.57 ± 0.10
Glutamic acid	10.3 ± 0.29	11.8 ± 0.39	12.8 ± 0.45	16.3 ± 1.74	13.4 ± 0.21
Proline	4.17 ± 0.60	4.95 ± 0.18	5.06 ± 0.32	4.38 ± 0.11	4.87 ± 0.13
Glycine	4.77 ± 0.24	5.18 ± 0.09	6.22 ± 1.14	5.48 ± 0.30	3.94 ± 0.12
Alanine	6.95 ± 0.18	6.76 ± 0.14	5.97 ± 0.31	6.16 ± 0.19	6.73 ± 0.09
Valine	7.00 ± 0.46	6.18 ± 0.17	6.31 ± 0.32	5.85 ± 0.15	6.13 ± 0.09
Isoleucine	5.91 ± 0.21	5.46 ± 0.14	4.88 ± 0.12	5.15 ± 0.15	5.34 ± 0.08
Leucine	8.43 ± 0.55	7.39 ± 0.18	8.11 ± 0.49	7.97 ± 0.24	8.16 ± 0.13
Tyrosine	$3.58 \pm .044$	4.96 ± 0.14	3.82 ± 0.26	3.67 ± 0.05	4.95 ± 0.20
Phenylalanine	6.60 ± 0.28	5.02 ± 0.09	5.48 ± 0.05	5.41 ± 0.25	5.74 ± 0.31
Histidine	1.15 ± 0.10	1.84 ± 0.17	1.75 ± 0.19	1.91 ± 0.08	1.00 ± 0.15
Lysine	5.91 ± 0.15	6.80 ± 0.10	7.33 ± 0.21	6.02 ± 0.12	7.26 ± 0.21
Arginine	8.78 ± 0.87	7.31 ± 0.09	6.69 ± 0.50	6.07 ± 0.16	5.65 ± 0.14
Total	93.9 ± 6.60	95.1 ± 2.52	96.4 ± 6.16	95.0 ± 3.93	95.3 ± 2.16

Table 2. Total amino acid composition of five seaweeds. Results are expressed as grams of amino acid measured in 100 g of algal protein and represent the actual recovery of amino acids after acid hydrolysis. Values are the mean of three replicates \pm SD (n = 3).

Table 3. Calculation of nitrogen-to-protein conversion factors for five seaweeds based on the amino acid residues to total nitrogen ratio. Values are expressed as percentage of the dry matter. Results represent the mean of three replicates \pm SD (n = 3).

Species	Total amino acid	Amino acid residues	Amino acid-N	Protein-N	N-Prot factors
Asparagopsis taxiformis	29.3 ± 2.41	25.1 ± 2.07	4.14 ± 0.34	74.5 ± 6.14	4.51 ± 0.37
Centroceras clavulatum	26.8 ± 0.79	23.0 ± 0.68	3.22 ± 0.09	69.5 ± 2.04	4.98 ± 0.15
Chaetomorpha aerea	14.0 ± 0.96	12.0 ± 0.82	1.94 ± 0.13	75.8 ± 5.20	4.69 ± 0.32
Sargassum filipendula	10.1 ± 0.38	8.62 ± 0.33	1.36 ± 0.05	78.0 ± 2.96	4.93 ± 0.19
Spyridia hypnoides	24.1 ± 0.58	20.7 ± 0.50	3.22 ± 0.08	81.0 ± 1.95	5.21 ± 0.13

4. Discussion

Carbohydrates are the most abundant substances in most seaweeds, since they occur in cell wall (ex.: agar, cellulose) and as storage products (ex.: starch, laminaran). Brown algae tend to show lower carbohydrates concentration than others groups of seaweeds [33] and the presence of less reactive carbohydrates may generate underestimates of total carbohydrate [55]. This might contribute to increase differences in comparison to both green and red algae. Kumari *et al.* (2010) [4] investigated the carbohydrate contents in eighteen species of seaweeds with the content ranged from 15 to 43% and the value reported for *Chaetomorpha* spp. (30%) was similar to our study. Instead, the reference [56] found 33.5% of carbo-

hydrate in *Sargassum polycystum*, higher than the value measured by us in *S. filipendula*.

The metabolism of benthic seaweeds typically involves the production of large amounts of carbohydrates as storage products [55]. The production of lipids is greater in planktonic algal species, in which they contribute for floating mechanisms. The total lipid contents vary with all species and this may reflect the difference capability of accumulating lipids. The fat content of seaweeds is generally low and accounts for 1% - 6% d.w. [57,58]. Altogether, seaweeds species were low in fat and high in carbohydrate. Both lipid and carbohydrate contents agree with previous studies [45,46]. Studies that used Folch's method to measure total lipid content are especially useful for comparison, such as [33,57,59]. These studies reported crude lipid values in most seaweed predominantly lower than 5% of d.w. The same trend was confirmed in our study, in which all species showed less than 5.5% of lipids.

The seaweeds show variable N and P tissue concentrations. Differences are related to taxonomic traits and species-specific differences of seaweeds in taking up dissolved nutrients [60]. Red algae tend to show higher N tissues concentrations than green and brown algae [29]. Red algae contain phycoerithrin, an N-rich pigment that increases the nitrogen budget of these species [55]. In addition, the three red algae tested are fast-growing species, which account for a higher N content in comparison to other species. Conversely, *S. filipendula* has a complex thallus and low rate of growth, showing typically low content of nitrogen [61]. A similar trend dwells a sandy substrate and it is partially burred.

S. filipendula showed the lowest P concentrations in its thallus, and this trend can also be interpreted as a consequence of its low growth rates. *Chaetomorpha aerea* showed the second higher P concentration, and this may be related to its contact with sediments. The occurrence of high tissue concentrations of phosphorus was recorded in three seaweeds that also occur partially burred in sediments (*Chaetomorpha crassa, Gracilaria cervicornis* and *Gracilariopsis tenuifrons*) in a seasonal study in Araruama Lagoon, a hypersaline coastal environment [51].

According to the reference [62] classification of macroalgal nutrient status based on N:P ratio of tissues, a N:P ratio < 16 indicates N limitation; a N:P ratio 16 - 24 indicates N-sufficiency and P-sufficiency—*i.e.* no limitation and N:P > 24 indicates P-limitation. According with this classification, the low N:P ratios found for all species (N:P < 11:1), except for *A. taxiformis* (18.9:1), suggests that these species trend to be N-limited. This is in accordance to the characteristics of Brazilian coastal waters, typically oligotrophic [51,63], with low availability of nitrogen to algal populations. However, this interpretation must be taken with care, since the amount of data in our study is small, and does not allow for conclusive remarks on this subject.

Proteins are composed of the one or more chains of amino acids and the nutritional quality of a protein is basically determined by the content, proportion and availability of its amino acids [64]. The main findings of amino acid composition of algal proteins described here are in agreement with previous studies [8,9,29,31,33,40]. In general, all species are rich in the acidic amino acids, glutamic and aspartic acid and poor in histidine. All seaweeds samples exhibits similar amino acid patterns, in which aspartic and glutamic acid constituted a substantial amount of total amino acids, ranged from 19.9% (*A*.

taxiformis) to 27.6% (*S. filipendula*). These two amino acids contribute to the flavour-related properties characteristic of the marine products and are responsible for the special taste of the seaweeds. The concentrations of these two amino acids were higher in brown algae than in red algae, as previously described by reference [33]. The level of glutamic and aspartic acid together can represent up 26% and 32% of the total amino acids of the green species *Ulva rigida* and *Ulva rotundata* [8]. The reference [29] showed that values for aspartic and glutamic acid together varied from 20.8% to 31.1% in 19 species of seaweeds. The highest value of lysine was observed in *Chaetomorpha aerea* a green alga, in contrast to reference [31] who found in red algae higher value for lysine.

Protein content of macroalgae from tropical and subtropical coastal environments frequently show low protein concentrations [40,65]. Our data indicate low protein concentrations in the algae studied, and this agrees with the predominantly oligotrophic condition of the Brazilian coastal waters. According to the literature, in general the total protein of brown seaweeds is low (3% - 15% of d.w.) compared to green and red seaweeds (10% - 47% of the d.w.) [9,46]. Variations in the protein content of seaweeds can be due to differences in species composition and seasonal periods [9]. The level of total and hydrosoluble proteins recorded in Sargassum filipendula agrees with the resuts generally found for the protein content of Sargassum species [29,59]. S. filipendula showed the lowest concentrations of both hydrosoluble protein and total protein, what agrees with the low TN found in its thalli. In addition, results for total and hydrosoluble proteins in S. filipendula were similar, which indicates that the extraction of protein was efficient with this species. On the other hand, the red algae showed both total protein content and TN concentration higher than the other species tested here. This suggests the presence of both high concentrations of non-protein nitrogen and variable degrees of efficiency in the extraction of protein.

In the present study can be observed a remarkable difference between the protein concentrations obtained with Lowry's method and the sum of AA-Res. The protein concentration estimated with Lowry's method achieved only about 50% in red algae in comparison protein concentration estimated with total AA-Res. The both method to protein quantification registered similar values for green and brown algae. This could suggest a lower efficiency on extraction in red algae, than brown and green algae. The inefficiency of the protein extraction in seaweeds has been discussed by reference [7], especially in freeze-dried samples. On the other hand, total amino acid analysis involves an acidic hydrolysis of the samples, which eliminates problems with protein extraction.

The best estimation of protein content was the sum of

AA-Res, which represents the true protein in each sample. The reference [28] analyzed the protein content from two species of *Porphyra* by different methods and found that the most accurate estimation of protein would be obtained with the knowledge of the molecular weights of the sequence of the amino acids. These authors affirm that the sum of amino acids appeared to be the most accurate method of determination of protein. Otherwise, the protein contents obtained by N-Prot factors were in good agreement with those of their AA-Res.

A usual way to determine N-Prot conversion factors is based in the sum of amino acid residues and determination of the amount of the total protein nitrogen, considering the individual contribution of each amino acid [16, 22]. Therefore, organisms that have proteins rich in highly nitrogenous amino acids (e.g. arginine) tend to have lower N-Prot conversion factors. In contrast, if the total protein contains large amounts of amino acids with a low proportion of nitrogen (e.g. tyrosine), the corresponding factors is likely to be higher. Thus, variation of the total amino acid concentrations may markedly influence the calculation of N-Prot factors [16]. The reference [25] indicated that the N-Prot factors calculated for many Japanese vegetables by total nitrogen could give a more accurate protein value than N-Prot value calculated by total amino acid nitrogen. This trend results from the presence of significant amounts on NPN in vegetables.

The use of the ratio of amino acids residues to total nitrogen to calculate N-Prot factors was described by reference [29]. The total amino acid content of seaweeds represents not only amino acids derived from proteins but also those in the free form. Thus the presence of free amino acids contributes to an overestimation of the total protein. However, according to reference [53], the use of data of total amino acid, without determination of free amino acids, is a widely accepted procedure to estimate protein, since in acid hydrolysis some amino acids are partially or totally destroyed (e.g. tryptophan, cystine, methionine and serine). The loss during acid hydrolysis might compensate for the influence of free amino acids in the quantification of protein by the sum of the total amino acid residues.

The overall mean N-Prot factors calculated in this report was 4.86. In general a remarkable similarity was observed with the current overall N-Prot factor proposed by reference [29]. These authors reported an average N-Prot factor of 4.92 for 19 seaweeds studied, with average specific factors for groups: 5.13 for green algae; 5.38 for brown algae and 4.59 for red algae. Reference [28] proposed mean N-Prot factor of 5.0 obtained for two species of *Porphyra*, the seaweed used to make Japanese sushi. The average N-Prot factors calculated for red algae in this study was 4.9, with highest value of 5.21, calcu-

lated for S. hypnoides.

Red algae tend to show larger amounts of NPN (30.5% in *C. clavulatum* and 25.5% in *A. taxiformis*), with exception of *S. hypnoides* (19%), than brown and green algae (22% and 24.2%, respectively). As a consequence of a high NPN in red algae, the reference [29] found that the N-Prot factors calculated for these algae tended to be lower than for the other algal groups. This could not be assessed in the present study because of the small number of seaweeds studied.

5. Conclusions

The seaweeds assessed here are poor in lipid and rich in carbohydrate. Results for hydrosoluble protein indicated that the extraction and reaction of protein occurred in variable degrees, with lower efficiency with A. taxiformis and maximum with S. filipendula. The present results showed that seaweeds have relatively high non-protein nitrogen concentrations and reinforce that the calculation of total protein content by the use of the traditional factor 6.25 overestimates the protein data. From the current data set it is clear that the factor 6.25 is unsuitable for estimating seaweeds protein contents. The present study establishes lower N-Prot factor than the traditional factor for all species. We recommend that the specific N-Prot factors calculated in this work are used in researches involving the species assessed here. We currently are assessing the effects of temporal variations on N-Prot factors calculated for seven seaweeds throughout two years of sampling. These new results will be published soon.

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