Review

Single Cell Protein: A Potential Substitute in Human and Animal Nutrition

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Abstract: Single cell protein (SCP) is the first product of the fermentation process and has proven to be a good protein alternative. Food competition is becoming more intense as the world’s population continues to grow. Soon, SCP may be able to compensate for a protein deficit. Various global businesses are focusing on SCP production, and the scope of its application is expanding as time and knowledge increases. High quantities of SCP can be produced by microorganisms, such as algae, yeast, fungi and bacteria, due to their fast development rate and the significant level of protein in their chemical structure. Beside proteins, SCP contains carbohydrates, nucleic acids, lipids, minerals, vitamins and several important amino acids. SCP has been an effective substitute for more expensive protein sources such as fish and soybean products. In conclusion, SCP can easily replace traditional protein sources in human and animal feed without detrimental effects. Potential substrate candidates and optimization strategies for SCP production have been extensively studied. This review article focuses on the various aspects of SCP, from its production, using different substrates, player microorganisms and nutritional benefits, to its economic aspects.

Keywords: single cell protein; algae; yeast; fungi; bacteria; mechanism of production; nutritional benefits

1. Introduction

The world population is continuously increasing. By 2050, the world population could increase to 9.3 billion [1,2] which at current consumption levels would cause the global demand for animal-derivative protein to reach 1250 million tons per year [3]. On the other hand, recent evidence has shown that almost 690 million people in the world (8.9 percent of the world population) are estimated to have been undernourished in 2019, and this is estimated to exceed 840 million by 2030. The most affected regions are Africa, Asia, Latin America and (to a lesser extent) the Caribbean [4]. Forecasting studies have shown that population increase will also be accompanied by economic progress, and consequently the increase in the living standard of about 3 billion people. It means that not only will water and food demands increase, but also the number of people who have more exigencies for food quality [5].

The United Nations Food and Agriculture Organization (FAO) describes proteins as macromolecules that make up the structural components of cells, tissues, muscles and organs. Proteins are required for metabolic function and are the nitrogen source for both humans and animals to build the structural and functional units required for living. The composition of a protein’s constituent amino acids reveals its nutritional worth. The most frequent are necessary amino acids, which humans and animals cannot synthesize. As a result, we consume meat or a protein sources to meet these needs [6].

Single cell proteins (SCPs) are isolated from the cells of microorganisms with high protein content as dried cells and/or as purified proteins. SCPs show very attractive features as a nutrient supplement for humans as they have a high protein content with...
wide amino acid spectrum, low fat content and a higher protein:carbohydrate ratio than forages [7]. SCPs contain vitamins, e.g., thiamine, riboflavin, pyridoxine, nicotinic acid, pantothenic acid, folic acid, biotin, cyanocobalamin, ascorbic acid, \(\beta\)-carotene and \(\alpha\)-tocopherol; essential amino acids, represented by lysine and methionine; minerals; nucleic acids and lipids [8,9]. Until now, SCPs have been used for a wide range of purposes, from food (aroma carriers, vitamin carriers, emulsifying acids, etc.) and feed (pigs, poultry, cattle, fish) production, to the paper and lead industry [9,10].

Common sources for SCPs are represented by waste and raw materials (starch, fruit, fruit waste, molasses, etc.), combustible and/or combustible waste and/or byproducts (natural gas, petroleum byproducts, ethanol, methanol, biomass, etc.) [8]. Methanol is soluble in its aqueous phase at all concentrations, and it can be easily removed from harvested biomass. Ethanol is a good substrate, but the process is not economically feasible. There are several advantages to utilizing waste for SCP production; these include the conversion of low-cost organic waste to useful products and a reduction in environment pollution. The cellulose, hemicellulose and lignin of natural waste wood originating from agriculture and forestry sources are attractive natural sources of SCP. However, they must be pretreated chemically (acid hydrolysis) or enzymatically (cellulases) to transform cellulose into fermentable sugars. Domestic sewage and waste retained from industrial cellulose processing, starch production and food processing can also be utilized. Agricultural waste has been reported to be an excellent substrate for cost-effective SCP production, the resulting protein-enriched product being of good quality and suitable for animal feed. It can even be consumed by humans after further processing [11].

The production of SCP from non-waste sources achieved industrial-scale production in the 1970s but was not economically competitive with other protein supplements Recently, interest in SCP has been renewed, partly because of the identification of new, less expensive production processes, but largely due to the realization that SCP production has vast potential environmental benefits over traditional protein supplements in animal feed [12].

1.1. Historical Background of SCP

Many microorganisms have been directly utilized as food. SCPs can be lifesaving in less privileged areas where malnutrition is a real and life-threatening problem. A species of alga called *Spirulina* was grown many years ago in Africa’s Lake Chad, and was subsequently used as food to compensate for local people’s protein shortfall [6]. Germans reportedly utilized a certain species of *Candida* in their meals during World War I, including sausages and soups. Proteins generated from bacterial, fungal and algal cultures were widely used in food and as food from then on. The concept of SCP arose from this method, and these proteins are now widely used [13].

1.2. Applications of SCP

SCP produced on commercial scale is used [13]:

- In animal feed and nutrition, for the stuffing and fattening of poultry, laying hens, calves and pigs.
- As food additives (vitamin and aroma carriers and emulsifying agents), to enhance nutritional value (of baked food items, ready-made meals, soups, etc.) and as starter cultures (baker’s, brewer’s and wine yeast).
- In industrial processes, as a foam-stabilizing agent and in paper and leather processing.

1.3. Mechanism of Production

There are different kinds of substrates, as categorized in [14]:

- High-energy resources: gas oil, natural gas, ethanol, methanol, \(n\)-alkanes and acetic acid
- Renewable plant resources: starch, sugar and cellulose
- Various wastes: sulfite waste liquor, molasses, whey, milk and fruit waste
- Carbon dioxide
The choice of substrate is made according to cost, availability, oxygen required during fermentation, quantity of heat produced and cooling capacity of the fermenter, but also the cost related to post-treatment processing [8,14]. Selected substrates are used as a growth medium by microorganisms such as bacteria, algae, fungi and yeast for increasing their cell mass, which is made up of SCP [15]. Fermentation is the main process responsible for SCP production [16], as shown in Figure 1.

Figure 1. General steps during industrial production of single cell proteins.

The available biomass is harvested when the fermentation process is completed and can be used as a protein source [17]. The biomass is processed further by purification, cell disruption, washing and protein extraction [18] to provide high production rates with high yields.

The demand for protein is increasing due to changes in food consumption patterns. However, the reliance on animal and dairy production to meet the growing demand for protein is ultimately unsustainable [3]. Hence, alternative sources should be explored to supply food for humans and animals [19].

SCP is widely viewed as a potential co-product that could boost the economic potential of a biorefinery process that is otherwise unprofitable and lower the downstream processing costs associated with the disposal of process waste. It is preferable to sell residual biomass as feed rather than as fertilizer. This can be seen in numerous publications in which specific waste products are converted to SCP and are assessed as food for specific animals [3].

The current review focuses on the various aspects of SCP, i.e., its application and the selection of a substrate and production method using different microbial sources (microalgae, yeast, fungi and bacteria), and also aims to identify future perspectives and challenges related to SCP. Data summarized herein can support researchers to create a food-grade SCP product with a high nutritional value (in terms of proteins, vitamins and lipids) by selecting the most suitable potential microorganism and a cost-effective technology.

2. Bacterial Metabolism

Bacteria have a wide range of metabolic processes characterized by enzymatic assimilation (the intake and use of organic and inorganic chemicals required for cellular growth and maintenance) and dissimilation reactions (the oxidation and breakdown of substrate). Assimilation reactions are endergonic, which means they require energy, whereas dissimilation reactions are exergonic, which means they generate energy. These reactions form the basis of the bacterial cell’s self-replication and are involved in critical functions of the cell [20].

Vasdekis and Stephanopoulos (2015) describe bacteria as cells that can value the energy they capture from the environment to accomplish their essential operations, thanks to their
particular energy-transforming ability [21]. Chemical energy is conserved in adenosine triphosphate (ATP), adenosine diphosphate (ADP) and/or molecules with a thioester link (e.g., succinyl SCoA and acetyl SCoA). These compounds have high energy phosphate bonds, which are used by enzymatic systems to synthesize new compounds needed for cell existence and development.

Bacterial enzymatic systems include B-complex vitamins as functional coenzymes that are involved in cell growth and energy transformation processes by catalyzing many oxidation–reduction reactions [22].

SCP metabolism involves the biological oxidation of organic compounds and yields simple organic and/or inorganic compounds together with ATP. The bacterial cell needs these compounds for anabolic pathways. Two modes of energy production are known in bacteria within heterotrophic metabolism: anaerobic respiration or fermentation (Figure 2A) and aerobic respiration (Figure 2B). Energy production may occur in both aerobic and anaerobic environments.

![Figure 2: Bacterial metabolism by fermentation (A) and aerobic aspiration (B). TCA, tricarboxylic acid.](image)

The fermentation is an anaerobic process, meaning that the terminal electron acceptor is other than O₂ (e.g., SO₄²⁻, NO₃⁻ or fumarate). Through glycolysis, glucose is broken down into pyruvate, yielding ATP and NADH (by the conversion of NAD). Pyruvate, in the presence of NADH, yields the end products of fermentation (Figure 2A). Aerobic respiration involves glucose catabolism. The pyruvate resulting from glucose breakdown produces acetic acid, carbon dioxide and NADH through mechanisms involving electron transport and chemiosmosis. Acetic acid together with coenzyme A yields acetyl SCoA, then acetyl radical enters in reaction cycle (Krebs or Glyoxylate) by detaching from CoA. The aerobic respiration is an exothermic process, yielding 380,000 calories when 1 mole of glucose is broken down by complete oxidation into carbon dioxide and H₂O, generating about 38 moles of ATP (Figure 2B).

**Bacterial Sources**

Bacterial SCP contains 50–80% protein (dry weight) [23], and is characterized by rich nucleic acid content, low density, small cell size and the capacity to easily multiply (within an interval of 20–120 min) on a wide range of substrates, such as sugar, starches, raw
materials, waste (i.e., organic waste and petrochemicals such as methanol and ethanol), and even water resources with a high nutrient and mineral content (Table 1) [8,24,25]. Plant-based products are mainly used as a substrate for SCP production. The most suitable waste for SCP production is agricultural waste, because when industrial waste is categorized it includes some of the agricultural waste that is generated by the food to be processed in industries [6]. Some bacteria (Methylococcus capsulatus, Methylococcus methanica, Methylobacterium soehngenii) can utilize methane for SCP production [26]. It can be observed from Table 1 that bacterial species like Methylobacterium methylophilum, Rhodopseudomonas palustris, Escherichia coli and Haloarcula sp. are very efficient in SCP production, so they have been most often exploited as protein producers.

Table 1. Bacterial protein content as single cell protein (SCP) on specific substrates.

| Bacteria                            | Substrate                                           | SCP (%) | Reference |
|-------------------------------------|-----------------------------------------------------|---------|-----------|
| Afifella marina STW181              | Commercial shrimp feed                               | >46     | [27]      |
| Bacillus cereus                     | Ram horn                                            | 68      | [28]      |
| Bacillus licheniformis              | Potato starch processing waste                       | 38      | [29]      |
| Bacillus pumilis                    | Potato starch processing waste                       | 46      | [30]      |
| Bacillus subtilis                   | Ram horn                                            | 71      | [28]      |
| Bacillus subtilis sp                | Soybean hull                                        | 26      | [31]      |
| Corynebacterium ammoniagenes        | Glucose + fructose                                  | 61      | [32]      |
| Cupriavidus necator                 | Synthetic growth medium                             | 40–46   | [33]      |
| Escherichia coli                    | Ram horn                                            | 66      | [28]      |
| Haloarcula sp. IRU1                 | Petrochemical wastewater                            | 76      | [34]      |
| Methylococcus capsulatus, Ralstonia sp., Bacillus agri, Anacarinibacillus sp., Methylococcus capsulatus, Methylophilus spp. Metilococ capsulatus | Methane (natural gas) | 67–73 | [25]      |
| Methylococcus capsulatus, Methylophilus spp. | Gas and liquid products of sewage                  | <41     | [19]      |
| Methylococcus methylophilus         | Methanol                                            | 81      | [25]      |
| Methylococcus capsulatus (bath)     | Methane                                             | 53      | [35]      |
| Methylococcus methylophilus         | Methanol                                            | 50      | [3]       |
| Methylococcus sp                    | Supernatant and biogas                               | 24      | [19]      |
| Methylocapsa acidiphila             | Methane                                             | 59      | [36]      |
| Methilomonas.sp                    | Natural gas                                          | 69.3    | [37]      |
| Methilomonas.sp                    | Biogas and supernatant of sewage sludge              | 56      | [19]      |
| Rhizobacter palustris               | Brewery wastewater                                   | >55     | [38]      |
| Rhodopseudomonas bisatica           | Wastewater from a latex rubber                       | 66.7    | [39]      |
| Rhodopseudomonas palustris          | Sludge and sago starch processing                    | 72–74   | [40]      |
| Rhodococcus gelatinosus             | Poultry slaughterhouse wastewater                    | 67.6    | [41]      |
| Rhodobacter sphaeroides P47         | Pineapple waste                                     | 66.6    | [42]      |
| Rhodopseudomonas as palustris P1    | Fermented pineapple extract                          | 65      | [39]      |
| Rhodobacter sphaeroides Z18         | Soybean wastewater                                  | 52      | [43]      |
| Rhodovulum sulphidophilum           | Glutamate malate medium                             | 15.6    | [44]      |
| Rhodococcus gelatinosus             | Pig farm waste                                      | 50.6    | [45]      |
| Rhodopseudomonas and R. falcaii     | Sugar refinery wastewater                            | 58      | [46]      |
| Rhodococcus gelatinosus             | Miso-like effluent medium                            | 63      | [47]      |
| Rhodobacter sphaeroides P47         | Dehydrated medium from pineapple peel waste         | 66.6    | [42]      |
| Rhodococcus gelatinosus             | Seafood processing wastewater                        | 50      | [48]      |
| Rhodopseudomonas sp.                | Synthetic medium                                    | 11      | [49]      |
| Rhodopseudomonas sp.                | Wastewater from noodle production                    | 50      | [50]      |
| Rhodopseudomonas sp.                | Municipal wastewater                                 | 60.1    | [51]      |
| Rhodococcus gelatinosus             | Tuna condensate                                     | 56      | [52]      |
| Rhodobacter capsulatus              | Synthetic medium                                    | 45      | [53]      |
| Rhodopseudomonas acidiphila         | Synthetic medium                                    | 23      | [54]      |
| Rhodococcus gelatinosus             | Cassava waste                                       | 56      | [55]      |
| Rhodopseudomonas palustris          | Simulated wastewater                                | 45      | [56]      |
| Rhodopseudomonas palustris          | Photosynthetic sludge                               | 74      | [57]      |
To be considered suitable for SCP production, bacterial strains must accomplish the requirements stipulated by several criteria: reaction conditions (heat and oxygen requirements during fermentation and foam generation); performance (yield, growth rate, pH and heat tolerance); behavior during fermentation (growth morphology and genetic stability); end product (in terms of bacterial protein composition and structure, purification yield and recovery rate) [58].

In a study by Kurbanoglu and Algur (2002), ram horn hydrolysate was processed in a batch system at 30 degrees Celsius in the presence of Bacillus cereus NRRL B-3711, Bacillus subtilis NRRL NRS-744 and Escherichia coli. The obtained bacterial cells had high total protein content (66%, 68% and 71% for Escherichia coli, Bacillus cereus and Bacillus subtilis, respectively). The obtained protein contained all amino acids for ruminant feed (alanine, aspartic acid, cystine, glutamic acid, glycine, serine, tyrosine, arginine, histidine, isoleucine, leucine, lysine, methionine, phenylalanine, proline, threonine, tryptophan and valine) [28]. Øverland et al. (2010) demonstrated that a bacterial culture (Methylococcus capsulatus, Ralstonia sp., Brevibacillus agri, Aneurinibacillus sp.) grown on natural gas as a carbon source and containing mainly the methanotroph Methylococcus capsulatus is a promising source of protein (67–73%) [25]. Imperial Chemical Industries developed a SCP (Pruteen) for animal feed from methanol using the bacterium Methylocapsulatus methylotrophus. Pruteen contained up to 70% protein and is used in pig feed [59]. Photosynthetic purple non-sulfur bacteria (PPNSB) (Rhodopseudomonas sp., Rhodobacter sp., Rhodocyclus sp.) cultivated on industrial wastewater containing heavy metals (Hg^{2+}, Cr^{6+}, Pb^{2+}) produced 70–72% crude protein. The strain proved to have efficient tolerance to those ions [60,61]. The amino acid profile of the protein composition (arginine, aspartic acid, cystine, glutamic acid, glycine, histidine, isoleucine, leucine, lysine, methionine, phenylalanine, proline, serine, threonine, tryptophan, tyrosine, valine) was similar to soybean protein (aspartic acid, glutamic acid, leucine, arginine, isoleucine, lysine, methionine, phenylalanine, proline, serine, threonine, tryptophan, tyrosine, valine) [23,62].

Kantachote et al. (2005) reported a protein percentage of 66.7% in fermented latex-rubber-sheet wastewater using bacteria Rhodopseudomonas blastica [63]. About 72–74% SCP was obtained by Getha et al. (1998) from sludge and sago starch processing with Rhodopseudomonas palustris [40]. The bacterial biomass was used as aquaculture feed. Ponsano et al. (2003) observed 67.6% SCP from Rhodococcus gelatinosus culture in poultry slaughterhouse wastewater. The bacterial biomass was used as a feed supplement [41]. Noparatnaraporn and Nagai (1986) used Rhodobacter sphaeroides P47 cultured in pineapple waste and reported 6.6% SCP [42]. Rhodopseudomonas palustris cultured in fermented pineapple extract contained 65% SCP and was used for the treatment of latex rubber sheet wastewater [39]. Rhodobacter sphaeroides Z08 produced 52% SCP when used soybean wastewater as a substrate [43]. Rhodopseudomonas capsulatus grown in glutamate malate medium produced 15.6% SCP [44]. Rhodococcus gelatinosus produced 50.6% SCP from pig farm waste [45]. Rhodopseudomonas and Rhodopseudomonas fulvum produced 58% SCP from sugar refinery wastewater [23]. 63% SCP was produced by Rhodococcus gelatinosus on miso-like effluent medium [47]. Rhodobacter sphaeroides P47 grown in dehydrated medium from pineapple peel waste produced 66.6% SCP [42]. Rhodococcus gelatinosus produced 50% SCP from seafood processing wastewater [48]. Rhodopseudomonas sp. grown in synthetic medium produced 11% SCP [49]. Rhodopseudomonas palustris and Rhodobacter blasticus grown in wastewater from noodle production produced 50% SCP [50]. Saejung and Thammaratana (2016) reported 60.1% SCP production from culturing Rhodopseudomonas sp. CSK01 cultured from municipal wastewater [51]. Rhodococcus gelatinosus R7 produced 56% SCP from tuna condensate [52]. Alexandre AJ et al. (2009) reported 45% SCP from Rhodobacter capsulatus cultivated from a synthetic medium [53]. Rhodopseudomonas acidophilus produced 23% SCP from a synthetic medium (ammonium sulfate and sodium acetate) [54]. Noparatnaraporn et al. (1987) reported 56% SCP from Rhodococcus gelatinosus cultivated in cassava waste [55]. Rhodopseudomonas palustris grown in simulated wastewater produced 45% SCP [56]. Rhodopseudomonas palustris produced 74% SCP from photosynthetic sludge and
the bacterial biomass was used as fish feed [57]. *Methylocapsa acidiphila* produced 59% protein when methane was used as a substrate [36]. Methanotrophic bacteria (*Methylophilus spp.* and *Methylomonas spp.*) produced 41% more protein content when sewage sludge was used as substrate. They also contain essential amino acids as histidine, valine, phenylalanine, isoleucine, leucine, threonine and lysine [19]. *Methylophilus*, widely used for SCP production [64], reached 69% protein when fed with natural gas. These were the growth conditions when methane salt broth and sodium nitrate were used as medium and nitrogen source, respectively [37]. *Rhodobacter sphaeroides* SS15 and *A. marina* STW181 (purple non-sulfur bacteria) mixed with commercial shrimp feed as a carbon source were explored by Chumpol Supaporn et al. (2018) [27]. Based on their protein content (53% and 46%, respectively) and essential amino acids, these biomasses are optimal for SCP. *Bacillus licheniformis* used waste potato as substrate to produce SCP. A temperature of 32.8 °C and pH of 6.67 were the optimal fermentation conditions, and the concentration of inoculum was 1.78%. These results demonstrate a potential application of this method in large-scale industrial production with 30% protein concentration [29].

SCP production was also reported by Kornochalert et al. (2014). Latex rubber sheet wastewater with added fermented pineapple extract was efficiently treated under microaerobic light conditions using *Rhodopseudomonas palustris*. The biomass contained 65% protein, 3% fat, 8% carbohydrate, 14% ash and 10% moisture. The SCP contained a higher content of protein, methionine and threonine concentration than soyabean meal (with 37% protein content) [39]. Kunasundari et al. (2013) cultivated *Cupriavidus necator* to produce a biomass high in both protein (40–46%) and polyhydroxyalkanoate (serving as a source of energy and as a carbon store). This biomass was used to feed rats [33]. Taran and Asadi (2014) showed that *Haloarcula sp.* IRU1 degrade petrochemical wastewater and use it as carbon source for single cell protein production in different conditions. They obtained SCP with 76.4% protein content [34]. Soya bean hull obtained from soya bean oil extraction is a cheap feed ingredient with a high fiber content. Two strains of *Bacillus subtilis* MR10 and TK8 were isolated from tua-nao, a traditional fermented soya bean in northern Thailand. The protein content after fermentation was 25.6% for MR10 and 26.6 for TK8 [31].

In terms of its high protein content and based on recent bacterial SCP research, it can be concluded that *Bacillus subtilis* (71%) together with *Rhodopseudomonas palustris* (72%–74%) and *Rhodopseudomonas spalustris* (74%) are the most exploited as protein producers, offering the advantage of high production rates and providing lipids and vitamins from the B group. However, bacterial SCP is disadvantaged by low familiarity and high nucleic acid content, which adds to the processing costs.

### 3. Algal Metabolism

The micro-algal metabolism refers to processes that include both biochemical mechanisms and nutrient transport. Intake nutrients are converted through a metabolic pathway into nutritional principles needed for vital processes, such as growth, reproduction and defense mechanisms. The mechanisms for the acquisition of carbon supplies, light capture, assimilation of nutrients (nitrogen and sulfur) and synthesis of the unique secondary metabolites make the difference between micro-algal metabolism and the metabolisms of other organisms [65].

The micro-algae display oxygen-evolving photosynthesis, which is a key feature that distinguishes them from other lower eukaryotes. This metabolic process is characterized by specific reactions often produced in the presence of light, at thylakoid membrane level. In algae, the basic chemo-organotrophic metabolism is similar to that encountered in bacteria. In algae, anabolic processes may occur both in the presence of light (photolithotrophic) or in the dark. The photorespiration process is catalyzed by specific enzymatic systems, including ATP synthases, the cytochrome b6–f complex or enzymes specific to the photosynthetic carbon-reduction cycle. In micro-algae, the photosynthetic reactions yield carbohydrates as phosphates. The algae photolithotrophic metabolism involves pathways that cannot take place in the dark. There are known linear pathways that produce the synthesis of
essential C skeletons, and they use photons, nitrogen, ammonium nitrate, ammonium sulfate, ammonium di-hydrogen phosphate and carbon dioxide for growth. Micro-algae that grow in the dark have only organic carbon as a source [66].

Figure 3 outlines the pathways of energy, carbon and oxygen in photosynthesis, photorespiration, dark respiration and the growth of an alga. No attempt is made to represent stoichiometries.

Figure 3. Algal metabolism [65]. Glyc, glycolate; PCOC, photorespiratory carbon-fixation cycle (or its equivalent); PCRC, photosynthetic carbon reduction cycle; PGA, 3-phosphoglycerate; Pglyc, phosphoglycolate; RuBP, ribulose bisphosphate; Rubisco, ribulose bisphosphate carboxylase–oxygenase.).

Algae Sources

Algae are autotrophic organisms characterized by wide genetic diversity. To sustain growth, besides inorganic nutrients (mainly nitrogen and phosphorus), they need water, carbon dioxide and light [67]. Environmental parameters such as pH and temperature also make influential contributions [68]. There are two algal categories: macro-algae (multicellular organisms) and micro-algae (unicellular organisms), commonly named phytoplankton and seaweed, respectively [69]. The macro-algae belong to four classes (phyla): i. cyanophyta (also known as the cyanobacteria group), are representative of the blue-green algae responsible for water blooms (e.g., Spirulina, Anabaena, Nostoc), ii. chlorophyta (green algae, i.e., Ulva, Enteromorpha), iii. haptophyta (brown algae, i.e., Fucus, Laminaria, Asphodelium, Macrocystis) and iv. rhodophyta/phodophyceae (red algae, the most diverse family, i.e., Porphyra, Rhodymenia) [67,68]. There is an enormous biodiversity of micro-algae species, with a high occurrence in marine systems, falling into five classes (phyla): i. chlorophyta (green micro-algae, i.e., Chlorella, Micractinium, Dunalella, Scenedesmus in freshwater systems, Chaetomorpha antennina, Phaeodactylum, Micractinium, Skeletonema, Ulva fasciata in marine systems), ii. rhodophyta/phodophyceae (red micro-algae, i.e., Rhodella reticulata, Porphyridium cruentum in freshwater systems, Gecarcinuc, Laurencia Palmaria palmata and Porphyra umbilicalis in seawater systems), iii. haptophyta (brown algae, i.e., Pavlova salina in marine systems), iv. stramenopiles (including brown micro-algae, Skeletonema costatum, Chaetoceros muelleri, Thalassiosira pseudonana in marine systems) and v. dinophyta (i.e., Cryptothecodium colinii in marine systems) [70].

The micro-algae have the capacity to produce cellular biomass with a major proportion of SCP (up to 70%), by converting solar energy [71]. Micro-algae mass cultivation produces high yields, 20- to 50-fold higher than soybean yields [72]. Micro-algae are single cell microorganisms, characterized by autotrophic growth, using light and carbon dioxide as energy and carbon suppliers. Heterotrophic growth is characterized by the use of molasses, manure or other cheap organic materials (e.g., industrial waste) as a carbon
source [73]. Some microalgae grow by combining both nutritional modes and are called mixotrophic algae.

Table 2. SCP production by algal species from different substrates.

| Algae                  | Substrate                  | SCP (%)             | Reference |
|------------------------|----------------------------|---------------------|-----------|
| *Chaetomorpha antennina* | Soda ash effluent          | 14.0–18.2%          | [74]      |
| *Ulva fasciata*        | Soda ash effluent          | 13.7–18.6%          | [74]      |
| *Chlorella sp.*        | Tofu waste                 | 52.32% ± 3.31       | [75]      |
|                        | Tempeh waste               | 52.00% ± 1.80       | [75]      |
|                        | Cheese waste               | 15.43% ± 2.55       | [75]      |
| *Chlorella salina*     | Saline sewage effluents    | 51%                 | [76]      |
| *Gracilaria domingensis* | Natural habit             | 6.2%                | [77]      |
| *Gracilaria birdiae*   | Natural habit              | 7.1%                | [77]      |
| *Laurencia filiformis* | Natural habit              | 18.3%               | [77]      |
| *Laurencia intricata*  | Natural habit              | 4.6%                | [77]      |
| *Palmaria palmate*     | Natural habit              | 8–35%               | [78]      |
|                        | Natural habit              | 8.0–35.0%           | [79]      |
|                        | Natural habit              | 11.9–21.9%          | [80]      |
|                        | Natural habit              | 13.5%               | [77]      |
|                        | Natural habit              | 12–21%              | [81]      |
| *Chondrus crispus*     | Natural habit              | 20.1%               | [82]      |
| *Porphyra umbilicalis* | Natural habit              | 15–37%              | [81]      |
| *Gracilaria verrucosa* | Natural habit              | 7.0–23.0%           | [79]      |
| *Chlorella sorokiniana*| Wastewater                 | 45%                 | [35]      |
| *Scenedesmus obliquus* | Wastewater                 | 52%                 | [83]      |
| *Spirulina*            | Salinated water            | 48.59%              | [84]      |
|                        | Desalinated wastewater     | 56.17%              | [84]      |

As shown in Table 2, several raw compounds are known as cultivation media and are used in micro-algae SCP production. For *Chlorella*, the literature mentions tempeh waste, with 52%; tofu waste, with 52.32% and cheese waste, with 15.43% total protein content [75]. Wastewater has been reported as a growing medium for *Scenedesmus obliquus* green alga with yields of 52% protein content [83], for *Arthrospira (Spirulina) platensis* cyanobacterium with SCP yields 48.59% or 56.17% by dry weigh, depending on the culture medium [84]. Jadeja and Tewari (2008) obtained 10–35% SCP yields using polluted water (soda ash industry effluent) as a carbon source for *Ulva fasciata* and *Chaetomorpha antennina* [74]. Different amounts of SCP were found in four red marine benthic species of algae collected from Espírito Santo State, Brazil: 18.3% dry weight in *Gracilaria birdiae*, 6.2% dry weight in *Gracilaria domingensis*, 4.6% dry weight in *Laurencia intricata* and 7.1% in *Laurencia filiformis* [77]. *Chlorella* produced high amounts of SCP, up to 45% of dry weight, which suggests the possibility of using algal biomass as substitute for conventional animal and vegetal protein sources [35]. Seasonal variations in micro-algae SCP production were also reported. *Palmaria palmata* gave the highest SCP yield (21.9% and 35% by dry weight) in winter/spring and lowest (11.9% and 8% by dry weight) in summer/early autumn [80,85]. Seasonal variations were reported of between 7% and 23% of dry weight for *Gracilaria verrucose* and between 15% and 37% of dry weight for *Porphyra umbilicalis* [85].

Based on recent Algal SCP research, *Chlorella sp* (up to 52%) and *Spirulina* (up to 56%) proved able to offer the highest protein content, while also providing healthy lipids and being seen as environmentally friendly and very “green.” It can be concluded that certain types of microalgae are successfully cultivated for animal and human consumption...
usually have healthy protein contents. Apart from the proteins, they are excellent sources of fats, mainly omega-3 fatty acids, mineral salts, vitamins and chlorophyll [86].

4. Fungal Metabolism

A series of specific reactions characterize fungal metabolism. The catabolic processes consist in the biosynthesis of a large number of compounds, usually divided into primary and secondary metabolites [87]. The primary fungal metabolites, such as alcohols, organic acids (citric and lactic acid) and amino acids (L-glutamate, L-lysine) are required for growth and reproduction. The secondary metabolites are not essential for cellular life, but have importance from an ecological point of view, because they are involved in versatile metabolic pathways and have the capacity to break down organic matter, which cannot otherwise be recycled [88,89]. These secondary metabolites are natural compounds, such as small peptides, amino acids, pigments and products with a potentially toxic action, such as mycotoxins and antibiotics [90–93].

The precursors of the primary metabolites of SCP are intermediate molecules involved in anabolic and catabolic pathways that may be used for the synthesis of macromolecular subunits (lipids, amino acids, nucleotides) and/or may be oxidized to generate ATP. Glucose and pyruvate (Figure 4) are used for the biosynthesis of fungal secondary metabolites. They are linked together as result of reactions catalyzed by enzymes such as dimethyl-allyl tryptophan synthetases (DMATSs), polyketide synthases (PKSs), terpene cyclases (TCs) or non-ribosomal peptide synthetases (NRPSs). Oligomers result from these reactions, which are often chemically modified by the action of tailoring enzymes controlled by transcriptional regulation [94]. Macromolecular biosynthesis consists in lipids and amino acids, which are the main nutritional components of biomass.

**Figure 4.** Fungal metabolism. TCA, tricarboxylic acid; ETC, electron transport chain.

**Fungal Sources**

Many fungal species are used to produce SCP (Table 3). Some fungal sources, such as *Pleurotus floria*, *Aspergillus niger* and *Fusarium venenatum*, are preferred due to their high protein content [95]. Fungi contain up to 63% protein when they are cultivated mainly for SCP production. Their amino acid profiles also meet the standards of the FAO concerning protein and amino acid in human nutrition [71]. Fungi proteins are rich in lysine and threonine but deficient in the sulfur-containing amino acids cysteine and methionine [71]. SCP obtained from fungi can also provide vitamin B-complex such as riboflavin, niacin,
thiamine, biotin, pantothenic acid, choline, pyridoxine, glutathione, amino benzoic acid, streptogramin and folic acid [96]. Fungi have a relatively high nucleic acid content (up to 10%) compared to algae (up to 6%) [71,97].

| Fungi                        | Substrate                  | SCP (%) | Reference |
|------------------------------|----------------------------|---------|-----------|
| Cladosporium cladosporioides | Rice bran                  | 10%     | [98]      |
| Penicillium citrinum         | Rice bran                  | 10%     | [98]      |
| Pleurotus florida            | Wheat straw                | 63%     | [99]      |
| Chrysonilia stophila         | Lignin                     | 39%     | [100]     |
| Aspergillus flaveus          | Rice bran                  | 10%     | [98]      |
| Aspergillus niger            | Apple pomace               | 17–20%  | [101]     |
| Aspergillus niger            | Banana waste               | 18%     | [102]     |
| Aspergillus niger            | Rice bran                  | 11%     | [98]      |
| Aspergillus niger            | Citrus pulp                | 25.6%   | [103]     |
| Aspergillus niger            | Potato starch processing waste | 38% | [29,30] |
| Aspergillus niger            | Waste liquor               | 50%     | [104]     |
| Aspergillus ochraceus        | Rice bran                  | 10%     | [98]      |
| Aspergillus oryzae           | Rice bran (deoiled)        | 24%     | [95]      |
| Fusarium semitectum and sp1 and sp2 | Rice bran         | 10%     | [98]      |
| Fusarium venenatum           | Glucose (Product: Quorn™)  | 44%     | [105]     |
| Monascus ruber               | Rice bran                  | 9%      | [98]      |
| Trichoderma harzianum        | Cheese whey filtrate       | 34%     | [106]     |
| Trichoderma virideae         | Citrus pulp                | 32%     | [103]     |

According to Baldensperger et al. (1985), the protein content of banana waste was raised from 6 to 18% by solid-state fermentation (SSF) using a strain of Aspergillus niger. Solar drying was used to make a green banana meal, and fermentation was carried out in a stirred reactor with a capacity of 15 kg (dry weight). Protein production was assessed to be 150% of the initial content because the substrate consumption was 24% of the starting weight after 43 h of fermentation. The fermented banana waste could be used as cattle feed because of its composition (50% total sugars, 13% reducing sugars and 18% proteins) [102]. In their study, Liu et al. (2013) converted potato starch processing waste using a two-step process, namely, the degradation of fiber in potato residue with Aspergillus niger and fermentation with wastewater using Bacillus licheniformis. Protein accumulation was also carried out to produce SCP as animal feed. The main objective was to convert potato starch processing waste into SCP as animal feed and to increase the SCP quality. A protein content of 28% SCP was obtained under optimized conditions [30].

Valentino et al. (2016) produced SCP from nine fungi using a rice bran substrate. The fungi used were Cladosporium cladosporioides, Aspergillus ochraceus, Aspergillus niger, Aspergillus flavus, Penicillium citrinum, Monascus ruber, Fusarium semitectum, Fusarium sp1 and Fusarium sp2. The SCP production potential was evaluated through the crude protein content (CPC) accumulated after 20 days of SSF. Results indicated that inoculation of fungi increased the crude protein content of rice bran. Aspergillus niger had the highest CPC (10.63%) followed by Aspergillus flavus (10.46%), Aspergillus ochraceus (10.25%), Fusarium semitectum (10.25%) and Cladosporium cladosporioides (9.69%). The uninoculated rice bran had the least CPC (9.53%). Aspergillus niger, Aspergillus flavus, Aspergillus ochraceus and Fusarium semitectum produced the highest percentage increases, of 11.51%, 9.48%, 7.59% and 5.25%, respectively. These results showed that endophytic fungi are good sources of SCP and enrich the CPC [98]. This study also evaluated the proximate composition of the rice bran. Results revealed a significant increase in the moisture, crude fiber, crude fat and ash content, while total energy and total carbohydrates decreased. Monascus-ruber-treated rice bran obtained the highest protein content (9%) [98]. Ahmadi et al. (2010) produced microbial protein by SSF, treating wheat straw with Pleurotus florida on a 2% NaOH substrate at 100 °C. The concentration of protein was 62.8% [99]. Chiou et al.
2001 reported 50% protein content from *Aspergillus niger* using waste liqueur [104]. This approach can also recycle food-processing-plant waste into animal feed resources. Bhalla and Joshi (1994) analyzed the co-culture of cellulolytic molds and yeasts on apple pomace in SSF and liquid-state fermentation. Results showed an increased protein content of apple pomace. The co-culture of *Candida utilis* and *Aspergillus niger* increased the protein content of dried and pectin-extracted apple pomace to 20% and 17%, respectively, under SSF conditions [101]. Ravinder et al. (2003) tested the growth of an industrially important fungus, *Aspergillus oryzae*, on deoiled rice bran through SSF technology. Among the various nitrogen sources tested, ammonium sulfate protein enrichment was 24.30%, followed by vegetable and fruit waste extract with a 23.50% protein content [95]. Rodriguez et al. (1997) confirmed that *Chrysonilia sitophilia* can degrade considerable amounts of lignin. High protein (39%) and low nucleic acid content indicates that *Chrysonilia sitophilia* SCP may be able to be used not only in animal fodder, but also as a potential food for humans [100]. Wiebe (2002) used glucose as a carbon source to produce SCP from *Fusarium venenatum*. The protein content was approximately 44% (w/w). This protein use-value is comparable to milk protein’s use-value [105]. Sişman et al. (2013) evaluated the nutritional characteristics and possible toxic effects of the SCP from *Trichoderma harzianum* grown on whey filtrate agar medium and obtained SCP with 34.21% protein [106].

SCP and crude pectinolytic enzyme production from citrus pulp is reported by De Gregorio et al. (2002). SCP and enzymes were produced by slurry-state flask cultivation of *Aspergillus niger* and *Trichoderma viride* on pulp collected from lemon juice clarification. “The highest protein level was reached after 14 days growth with Aspergillus niger and 25 days growth with Trichoderma viride, though the final amount was higher in Trichoderma viride (31.9%) than in Aspergillus niger (25.6%) [103].”

5. Yeast Metabolism

In yeasts, the basic metabolic mechanisms are identical for different cells, but the resulting metabolites are different. Complex enzymatic reactions are involved in yeasts’ metabolism. The metabolic process begins when the substrates penetrate the cell. Action-specific factors then produce alterations of their basic structure [107]. The enzymatic reactions catalyzing the metabolic processes are needed for different aims: production and development of other cells, oxidation–reduction reactions which generate carbon dioxide and alcohol. Carbohydrates are the main carbon and energy source for most yeast species. However, there are known yeast species that use other sources of carbon, such as *Cryptococcus aureus*, *Cryptococcus laurentii*, Hannaella aff. zeae, *Tremella encephala* and *Trichosporon coremiiforme* [108]. These microorganisms have the capacity to form proteins and amino acids from simple nitrogenous sources as presented in Table 4.

The glycolysis process uses simple carbohydrates as substrates, mainly fructose and glucose. If other, more complex carbohydrates (maltotriose or maltose) are involved, they are hydrolyzed to glucose by α-glucosidase. During the first step of the yeast’s metabolic process, glucose is degraded to pyruvate, through glycolysis. The pyruvate metabolic pathway leads to the release of energy. Fermentation generates small amounts of energy, while respiration produces a larger amount (Figure 5). The carbon substrate is totally oxidated through respiration. Oxidative phosphorylation and the Krebs cycle are the mechanisms involved in this process. The energy produced and accumulated in ATP contributes to the generation of intermediary metabolites. In presence of oxygen, pyruvate enters into mitochondria where, under the action of the pyruvate dehydrogenase multi-enzyme system, it is the subject of the oxidative decarboxylation to acetyl-CoA. When yeasts are grown on two carbon sources (e.g., ethanol or acetate), the glyoxylate cycle, which may be considered a shortcut for the Krebs cycle, is often the supplier of energetic compounds. The alcoholic fermentation of carbohydrates involves a process whereby yeasts re-oxidize NADH to NAD. It is a two-step reaction: (1) pyruvate decarboxylase catalyzes the process of pyruvate decarboxylation, (2) alcohol dehydrogenase catalyzes
the subsequent reaction of the reduction of acetaldehyde at the same time. Simultaneously, dihydroxyacetone phosphate generates glycerol.

![Figure 5. Glycolysis process.](image)

Besides glycolysis, the pentose phosphate cycle or hexose phosphate pathway is an alternative way to glucose oxidation. This pathway provides the yeast cell with cytosolic NADPH and pentose carbohydrates, needed for production of amino acids, alcohols or fatty acids through biosynthetic reactions. This pathway has the following steps: (1) dehydrogenation of glucose-6-phosphate to 6-phosphogluconate and generation of one NADPH molecule, which is a reaction catalyzed by glucose-6-phosphate dehydrogenase; (2) the action of phosphogluconate dehydrogenase, which catalyzes the decarboxylation of 6-phosphogluconate, resulting ribulose-5-phosphate and a second molecule of NADPH. This pathway has two major functions: NADPH generation and the production of ribose carbohydrates used in the biosynthesis of nucleic acid precursors and nucleotide coenzymes. NAD and FAD, which are oxidation–reduction carriers, become reduced during the breakdown of carbohydrates to NADH and FADH2, but they are re-oxidized in the electron transport chain located in the mitochondrial membrane. The ATP synthase enzymatic system is also located in the mitochondrial membrane and catalyzes ATP formation from ADP and inorganic phosphate, but also the process of oxidative phosphorylation. The energy released during the transfer of electrons is connected with oxidative phosphorylation [109].

Anaerobic alcholic fermentation involves the partial degradation of the substrate because it stops at the end of glycolysis, generating two molecules of pyruvate. It means that an incomplete oxidation of glucose occurs and this process provides low amounts of energy. Ethanol and carbon dioxide are the oxidation products of carbohydrates.

**Yeast Sources**

Yeast is a good source of SCP production and has been used for a long time, mainly due to its superior nutritional quality. In Germany during World War I, torula yeast (*Candida utilis*) was used in sausages and soups. At present, it is frequently used in animal feed as a fodder supplement (dog and fish feed) and used as a seasoning in vegetarian
foods [7]. Yeast cells are larger than bacterial cells, have high lysine and malic acid contents, low methionine and nucleic acid contents and are able to grow under acidic pH conditions [110]. Yeast has low growth rate and lower protein content (65%) compared to bacteria (80%) [71,111]. A variety of substrates and microbes have been used for SCP production; however, presence of toxic and carcinogenic compounds (aflatoxins of type B1, B2, G1 and G2, citrinin, trichothecenes and zearalenone) generated during the production process should always be considered. The two main limiting factors for yeast are the high nucleic acid content and low cell-wall digestibility [14,71,111].

Table 4. Yeast SCP production from different substrates.

| Yeast                 | Substrate                                      | SCP (%) | Reference |
|-----------------------|------------------------------------------------|---------|-----------|
| *Saccharomyces cerevisiae* | Orange pulp, molasses, brewer’s spent grain | 39      | [112]     |
| *Candida krusei*       | Cheese whey                                    | 48      | [113]     |
| *Candida tropicalis*   | Molasses                                       | 56      | [114]     |
| *Candida tropicalis*   | Bagasse                                        | 31      | [115]     |
| *Candida utilis*       | Waste capsicum powder                          | 29      | [116]     |
| *Candida utilis*       | Poultry litter                                  | 48      | [117]     |
| *Candida utilis*       | Potato starch industry waste                    | 46      | [30]      |
| *Candida utilis*       | Potato wastewater                               | 49      | [118]     |
| *Hanseniaspora uvarum* | Spoiled date palm fruit                        | 49      | [119]     |
| *Kefir sp.*            | Cheese whey                                    | 54      | [120]     |
| *Kefir sp.*            | Brewery’s spent grains (hemicellulosic hydrolysate) | 32    | [121]     |
| *Debaryomyces hansenii*| Cheese whey                                    | 43      | [113]     |
| *Kluyveromyces marxianus* | Orange pulp, molasses, brewer’s spent grain, whey, potato pulp, malt spent | 34 | [112] |
| *Kluyveromyces marxianus* | Cheese whey                                    | 43      | [113]     |
| *Zygosaccharomyces rouxi* | Spoiled date palm fruit                        | 49      | [119]     |
| *Yarrowia lipolytica*  | Inulin, crude oil, glycerol waste hydrocarbons | 48–54  | [122]     |

A series of studies conducted worldwide show promising results concerning the possibility of using yeast as SCP, as reflected in Table 4. Studies performed by Aggelopoulos et al. (2014) on the growth of *Kefir sp*, *Kluyveromyces marxianus* and *Saccharomyces cerevisiae* by SSF using substrates composed of various kinds of common food industry waste (whey, molasses, brewer’s solid waste and orange and potato residues) emphasize a protein content of 39% in *Saccharomyces cerevisiae*, 34% in *Kluyveromyces marxianus* and 23% in *Kefir sp* [112]. Another study analyzed a mixed culture of *Kluyveromyces marxianus* and *Candida krusei* to enhance chemical oxygen-demand removal efficiency, minimize contamination at extreme conditions (high temperature (40 °C) and low pH (3.5)) during batch and continuous aerobic fermentation, and to obtain improvements in the quality of the SCP using whey as a substrate. The results show that the SCP content was 47.53% for *Candida krusei* and 43% for *Kluyveromyces marxianus* [113]. An effective bioprocess to produce SCP from soy molasses using *Candida tropicalis* was developed by Gao et al. (2012). This SCP contained 56.42% crude protein and 5.28% nucleic acids. These results suggest that *Candida tropicalis* might be applied effectively to produce SCP using soy molasses as a low-cost substrate [114]. Jalasutram et al. (2013) reported 48% protein content from digested and undigested poultry litter by *Candida utilis* [117]. Research on sugar cane bagasse used as a substrate to grow *Candida tropicalis* in a 1.0-l bioreactor at 30 °C and a pH of 6.0. showed that the protein produced (31.3% of the total biomass) contained essential amino acids (lysine, histidine, arginine, aspartic acid, threonine, serine, glutamic acid, proline, glycine, alanine, cysteine, valine, methionine, isoleucine, leucine, tryptophan, tyrosine, phenylalanine) for animal feeding [115]. Capsicum powder medium contained sufficient nutrients and could be used as a good medium to produce SCP, while *Candida utilis* yeast used to produce microbial biomass yielded a protein content of 29%. Utilization of waste capsicum powder
can reduce environmental pollution produced by waste generation by converting it to use in animal feed [116]. Liu et al. (2013) conducted a study to reduce the pollution of the waste produced by the potato starch industry and transform the potato pulp and wastewater into SCP which could ultimately be used as animal feed. Results show that Candida utilis cultured in potato dextrose medium and potato wastewater at 28 °C for 48 h led to a protein content of 46% [30] and 49% [118], respectively. Hashem at al. (2014) used two yeast strains Hanseniaspora uvarum and Zygosaccharomyces rouxii from spoiled date fruits for single cell protein production. Both strains were assessed for SCP productivity in vitro and in a bioreactor. The highest production (48.9 g/L) of the two strains was achieved after 60 h [119]. Paraskevopoulou et al. (2003) produced SCP (53.9% protein) using the aerobic fermentation of cheese whey by Kefir microorganisms. The experiments were conducted under controlled pH (5.5) and temperature (30 °C) conditions [120]. Cui et al. (2011) reported that Yarrowia lipolytica is suitable for producing SCP from inulin and inulin-containing materials, obtaining a protein content of between 48–54% [122]. SCP production from Debaryomyces Hansenii yeast biomass using brewer’s spent grains (hemicellulosic) as a carbon source was demonstrated by Duarte et al. (2008). The total protein content obtained was 32% [121].

6. Nutritional Benefits of SCP

Wu et al. (2014) indicated that the worldwide human protein diet is 65% plant based and 35% animal based [123]. The average meat intake per capital is projected to rise by 29%, from 40.0 kg in 2013 to 51.5 kg in 2050. Global meat production is therefore expected to increase from 288 million tons in 2013 to 494 million tons by 2050. According to previous research, SCP could be a valuable solution for supplying the global demand of protein, due to its low cost of production, easy process and nutritional quality [8,11,71,124].

As previously stated, the primary goal of SCP production is to use it as a protein (meat) substitute to address food scarcity and hunger in the near future. SCP must meet nutritional requirements to be consumed as human food or animal feed, such as the required protein content, amino acid composition of the protein being generated and digestibility of that protein. According to the researchers, SCP must be produced to the highest standards to be safe and beneficial for food and feed [125].

Finnigan et al. (2017) defined some of the macro and micronutrients provided by SCP as follows: proteins, lipids, carbohydrates, β-carotene, vitamin A precursor, biotin, folic acid, niacin, pantothenic acid, pyridoxine, riboflavin, thiamine, vitamin B12 (cyanocobalamin), vitamin C and vitamin E [126].

Beside its protein content, the nutritional value of SCP depends on its chemical composition (amino acids, nucleic acids minerals, enzymes and vitamins) but it is relatively cheap compared to other plant and animal sources [127]. It was reported that dried cells of Pseudomonas spp. grown on petroleum-based liquid paraffin contain as much as 69% protein. SCP obtained by algae processing is about 40%. Though proteins obtained from microbes contain all essential amino acids, their composition depends on the type of substrate used (carbon or nitrogen) and the type of microorganism grown on a specific medium [128]. Microorganisms like bacteria and yeast have a very short multiplication time as they double their population in just 5–15 min, while algae and mold species double themselves in 2–4 h. The amino acid profile of SCP from bacteria shows a close resemblance to fish protein and protein from yeast resembles soya protein [129]. In addition, SCP is reported to be deficient in sulfur-containing amino acids such as methionine and cysteine, whereas high levels of lysine and other amino acids have been observed. Supplementation of methionine and cysteine is required for the use of SCP as a feed ingredient. Microorganisms normally contain vitamin B12 in significant quantities. Bacteria and algae are reported with high vitamin B12 and vitamin A content, respectively [130]. Most common vitamins present
in SCP are riboflavin, thiamine, pyridoxine, niacin choline, folic acid, pantothenic acid, biotin, para-amino benzoic acid, inositol and B12 [130]. Most microorganisms have very fast growth rate that yields a high amount of biomass (algae: 3–6 h, bacteria: 30 min to 2 h, yeast: 40 min to 3 h) [71]. These microorganisms can be used as a whole in contrast to most crop and animal protein sources which cannot be used entirely [131]. SCP produced from different microbes has high protein content (30–70%) as compared to different green plants and animal sources [86]. Moreover, these proteins have an excellent amino acid profile that makes them nutritionally more useful than conventional protein sources [132]. Nevertheless, scientists can add more valuable amino acids to proteins through genetic engineering, obtaining SCP with the desired composition. Some microorganisms during the course of SCP production produce significant amounts of vitamins which cannot be produced by the host individual in appropriate amounts. Production of SCP also requires low water content as compared to plant sources [133]. Unlike plant protein sources, SCP is independent of environmental and climatic variation and can be produced throughout the year as microbes are available round the clock. The shelf life of protein preparation is dependent on the nature of the protein and the storage conditions used, and can vary from a few days to more than a year. Optimal conditions for storage are distinctive to each protein.

Nutritive and food values of SCP vary with the microorganisms used. The microorganisms used for SCP production must be non-pathogenic, toxin-free, easy to handle and separate from the substrate and should also tolerate the scaling-up of the process. Fast-growing microorganisms are required for getting massive output (biomass weight produced per unit time). However, high output produces more quantities of RNA in the cell, and this is not desirable as it acts as anti-nutritional factor in the final product. The method of harvesting, drying and processing affects the nutritive value of the finished product [134–138]. The idea that SCP could help to overcome food shortages in developing countries has garnered interest among scientists and industry. For future success of SCP, food technology problems must be solved to make these foods familiar, while the production should compare favorably with other protein sources.

The raw materials for SCP production based on waste substances are cheap, readily available and their processing contributes to reducing environmental pollution. The origin of the feedstock must be carefully selected. Various types of raw materials are attractive sources for SCP production from a cost and sustainability perspective but may raise safety concerns. In addition to the safety requirements, the use of other unconventional waste-derived protein sources in human nutrition requires efforts to improve public perception and acceptance and increase consumer awareness of the benefits of SCP consumption in human diets.

Yeast SCPs have been used in aquaculture diets as partial replacement for fishmeal, thanks to their excellent nutrient profiles and cost-effective large-scale production [138,139], and have also been applied for the highly unsaturated fatty acid fortification of Artemia and rotifers [137]. Some yeast strains with probiotic properties, such as Saccharomyces cerevisiae [140] and Debaryomyces hansenii [141], increased larval survival by early maturation of the pancreas and intestine [141]. However, many of these yeast SCPs are deficient in sulfated amino acids, e.g., methionine [140], thus, cannot be used as sole protein sources. Micro-algal SCP may be used for both animal and human consumption, and their nutritional value is similar to, and sometimes higher than, values reported for conventional food/feed supplements. Besides their high protein content, they are a source of nucleic acids (up to 6%), minerals (sodium, magnesium, potassium, iodine) and vitamins (A, B group, D, C and E) and essential amino acids (leucine, valine, lysine, phenylalanine) [97] (Figure 6). Compared to many vegetable foods, micro-algae SCP contains higher amounts of vitamins such as riboflavin, thiamine, folic acid or pro-vitamins as carotene [73]. Micro-algae, as Spirulina and Chlorella, are sources of vitamin B12 (cyanocobalamin), which otherwise has animals as almost exclusive source [73]. Algal SCP has a nutritional value like other SCP sources. The crude protein content (N 6.25) varies between 45 and 73%,
while the lipid content is 2–20% and is rich in essential fatty acids, and the mineral content is 5–10%. The protein content of algae is higher than the value for soybean (70% and 40%, respectively), and its amino acid profile shows an adequate balance except, as for any other microbial biomass, for the sulfur-containing amino acids methionine and cysteine [71,73]. Algal SCP is a good source of vitamins A, B group, D, C and E; the content of some vitamins such as thiamine, riboflavin, folic acid and carotene is higher in algae than in many vegetable foodstuffs. Some microalgae, such as *Chlorella* and *Spirulina*, contain vitamin B12 (cyanocobalamin), which is found almost exclusively in animal origin foods. The content of nutrients, however, is highly dependent on cultivation and processing conditions [73]. In addition, the search for new insects as a source of protein and the related technology for processing requires further research.

**Figure 6.** Composition of algal species.

Due to its efficiency requirements (low-cost production, protein quality, easy process), SCP production is not an easy task, and the obtained end-product might also raise some potential limitations. A key challenge for the industry is sourcing a sustainable, renewable high-protein ingredient. SCP contains up to 16% nucleic acids [97]. This may be a problem when SCP is destined for human consumption because the recommended nucleic acid dose in human nutrition is below 2%. Purines resulting from nucleic acid breakdown during human metabolic processes are responsible for the potential harmful effects of high nucleic acid by raising the uric acid levels in plasma, which can lead to gout and kidney stones [71]. The process of converting SCP to a consumable food for humans also requires the additional development of aroma and taste, which is certainly not cost-effective and makes the process less efficient [76]. SCP can cause allergic reactions for some humans who have a sensitive digestive system or if their body refuses to recognize the biomass [3]. Moreover, waste materials used as substrates in SCP production may contain unknown substances which could raise other health issues [76].
7. Economic Analysis

SCP production can be a profitable business if certain tactics for economic production are used. The following categories/processes are included in the costs for its production [3,142]: the cost of raw materials and chemicals/enzymes needed for pretreatment of substrates, particularly lignocellulosic waste, the cost of reducing agents for strict anaerobes and the cost of scaling up the fermentation process. The cost of this scale-up is directly reflected in the cost of the final product. Thus, a continuous fermentation process is preferable for the industrial production of SCP, as this is the most profitable strategy. In a study conducted by Junaid et al. (2020), the product cost, capital investment and profit obtained from the product were used to determine the economic viability of SCP production. To achieve this goal an improvement of the strains of microorganisms, better fermentation methods and advanced down-streaming methods are needed.

Economic analysis performed by Liu et al. (2014) indicated that the processing of potato waste could simplify not only the pollution problem in the starch industry, but also the shortage of protein for animal feed in China [29]. SCP microorganisms incorporate most nutrients fed to them into the final harvested biomass. SCP could replace soybean meal and other protein supplements used in animal feed, such as meat meal and fishmeal. Offsets from these sources would create different environmental and economic incentives. Fishmeal is priced at almost five times the price of soybean meal, at $1500/ton versus $320/ton [143]. SCP production from food waste is another promising microbial technology that may yield better outcomes as a food-waste management strategy over anaerobic digestion.

8. Conclusions

Researchers and businesses from all over the world are interested in SCP production. Thanks to the multiple promising benefits that these proteins provide, various firms have sprouted up that claim to be able to commercialize SCP. However, a key challenge for the industry is sourcing a sustainable, renewable protein-enriched ingredient. Currently, fishmeal, along with terrestrial plant meals, makes up most of the protein content of diets. Among others, algae, fungi, yeast and bacteria can be utilized for SCP production but each have their own advantages and disadvantages. Bacteria possess higher growth rates, higher protein content and more sulfur-containing amino acids. From an industrial perspective, methane-oxidizing bacteria are the most advanced and market-ready bacteria for SCP production. Yeasts have been used as a source of SCP for a long time. SCP from fungi has been found to be useful in animal feed, while specific byproducts are employed in the beverage sector. Single cell algae have a high growth rate, protein content (up to 70%), and chlorophyll; contain bile pigments, fiber and mineral salts; and have a lower nucleic acid content (4–6%) compared with fungi (9.7%) and bacteria (16%). As well as providing lipids that are beneficial for human nutrition, algal SCP is a good source of vitamins, as the content of some vitamins such as thiamine, riboflavin, folic acid and carotene is higher in algae than in many vegetable foodstuffs. Some of these microalgae, like Chlorella and Spirulina, include vitamin B12 (cyanocobalamin), which is almost exclusively found in animal-derived diets. Nutrient content, on the other hand, is greatly reliant on growing and processing circumstances. Although it has very attractive characteristics, SCP also presents some challenges that must be considered, such as production cost and growth conditions, but most importantly, the higher concentration of nucleic acid in SCP than in other conventional protein sources, which is its main anti-nutritional factor. There is still a lot of work to be done on this subject to reap as many benefits of SCP as possible, but several challenges described above can be avoided by improving the strains of microorganisms, carefully optimizing the fermentation protocol during production and implementing advanced down-streaming methods. Moreover, the selection of active microorganisms along with a suitable substrate is helpful in counteracting the above limitations and makes the usage of SCP beneficial. Furthermore, anti-nutritional factors,
i.e., nucleic acids, can be removed by applying different physical and chemical treatments during processing.

SCP exhibits very attractive characteristics as a nutrient supplement for both animals and plants. It can be produced at any time in the year because of its independence from seasonal and climatic variations. It could be produced from various cost-free substrates. Therefore, it can easily replace conventional animal and protein sources in both humans and animal diets without any negative impact.

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