Valorization of Seafood Processing Discards: Bioconversion and Bio-Refinery Approaches

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The seafood industry generates large volumes of waste. These include processing discards consisting of shell, head, bones intestine, fin, skin, voluminous amounts of wastewater discharged as effluents, and low-value under-utilized fish, which are caught as by-catch of commercial fishing operations. The discards, effluents, and by-catch are rich in nutrients including proteins, amino acids, lipids containing good proportions of polyunsaturated fatty acids (PUFA), carotenoids, and minerals. The seafood waste is, therefore, responsible for loss of nutrients and serious environmental hazards. It is important that the waste is subjected to secondary processing and valorization to address the problems. Although chemical processes are available for waste treatment, most of these processes have inherent weaknesses. Biological treatments, however, are environmentally friendly, safe, and cost-effective. Biological treatments are based on bioconversion processes, which help with the recovery of valuable ingredients from by-catch, processing discards, and effluents, without losing their inherent bioactivities. Major bioconversion processes make use of microbial fermentations or actions of exogenously added enzymes on the waste components. Recent developments in algal biotechnology offer novel processes for biotransformation of nutrients as single cell proteins, which can be used as feedstock for the recovery of valuable ingredients and also biofuel. Bioconversion options in conjunction with a bio-refinery approach have potential for eco-friendly and economical management of seafood waste that can support sustainable seafood production.

Keywords: seafood by-products, seafood waste treatment, valorization, bioconversions, bio-refinery, marine biotechnology

INTRODUCTION

Global food production is growing significantly to meet rising consumer demands. The seafood industry, a major segment of the food industry, provides finfish and shellfish of choice to consumers worldwide. Marine seafood includes finfish (pelagic, anchoveta, pollock, tuna, herring, mackerel, whiting, and others), and shellfish, which include crustaceans such as shrimp, krill, crab, lobster, and, mollusks, consisting of bivalves (mainly mussels, oysters, clams, and scallops), cephalopods (squid and cuttlefish), and gastropods (mainly abalone and snails). In 2018 global
seafood production was 178.5 million tons (MT), consisting of 96.4 MT of capture fisheries (FAO, 2020). Of the 71.9 MT of finfish, the most popular marine species were anchoveta (7 MT), followed by Alaska pollock (3.4 MT), skipjack tuna (3.2 MT), herring (1.8 MT), and blue whiting (1.7 MT). Aquaculture production was 82.1 MT in 2018 consisting of finfish, mollusks, and crustacea at 7.3, 17.3, and 5.7 MT, respectively. Seafood amounting to 67.1 MT consisting of marine and farmed species including shrimp, prawns, salmon, mollusks, tilapia, catfish, sea bass, sea bream, and others constituted international trade. It is projected that utilization of fishery products for human consumption will reach 204 MT by the year 2030. It should be noted that percentage of fish stocks within biologically sustainable levels in 2017 was only 65.8% against 90% in the year 1990 (FAO, 2020). This suggests efforts by stakeholders for sustainability of the seafood resources are necessary.

SEAFOOD PROCESSING DISCARDS AND EFFLUENTS

The processing of food generates enormous amounts of waste, which include both solid discards as well as wastewaters containing portions of food being processed, which are released as process effluents. It has been estimated that, on average, about one-third of food produced globally, amounting to 1.3 billion tons, is wasted (Gustavsson et al., 2011). In the case of seafood, all the species harvested are not adequately used as food. Consumers prefer only a few select seafood items. A significant portion of the total harvest, therefore, remains unused or poorly used due to inherent problems related to unattractive color, smaller size, and high fat content. These result in a sizeable amount of catch being treated as by-catch, which is usually a combination of several species, particularly from tropical shrimp fisheries, and is unused or poorly used as food (Venugopal and Shahidi, 1998). Gustavsson et al. (2011) observed that food loss and waste for the whole fisheries sector amounted to 35% of global catches; 9–15% of these losses arise from by-catch.

The seafood industry processes about 80% of total harvest into chilled, frozen, smoked, dried, fermented, or marinated products. The centralized pre-processing operations, which include beheading, de-shelling, skinning, gutting, removal of fins and scales, fileleting, washing, and others, leads to significant amounts of solid wastes and wastewater as effluents. The waste, on a wet weight basis, constitutes as high as 50% of whole shellfish, such as shrimp, krill, and crab. Shrimp waste contains about 70% head and 30% shell (Yan and Chen, 2015). Argentine red shrimp (ARS) is a highly popular shellfish, the industrial processing of which yearly yields 18,000 MT of shell waste, which is responsible for environmental pollution and ecological imbalances in Argentine Patagonia (Cretton et al., 2020). Lobster processing generates 50–70% of the shellfish as by-products such as heads, shells, livers, and eggs, which annually amounts to more than 50,000 MT (Nguyen et al., 2017). India produces up to 80,000 MT of shellfish waste (Chandrasekharan, 2015). Discards from finfish constitute 25–50% of the raw material, and is comprised of entrails, heads, skeletal frames, skin, scales, and viscera. Processing of freshwater fish such as trout, carp, pike, and bream generates 40–60% of the fish as waste (Venugopal, 2006). Love et al. (2015) reported that during the period 2009–2013 about 47% of the edible seafood supply was not available in the US for human consumption. This also included 16–32% of the harvest discarded as by-catch. In Europe, for each ton of seafood consumed an almost equal amount is estimated to be discarded as waste. The processing of shrimps and crabs in the EU alone results in more than 100,000 MT of shell waste each year (Sieber et al., 2018). Further, it is cautioned that large amounts of new fish biomass will be generated in European ports following the Landing Obligation Guidelines issued by the EU (Uhlmann et al., 2019).

The seafood industry, apart from solid discards, generates voluminous amounts of wastewater as process effluents as a result of operations such as washing, chilling, Blanching, fileleting, cooking, marination, and others. It has been estimated that ~10–40 m^3 water is required for processing each ton of raw seafood (Arvanitoyannis and Kassaveti, 2008). One of the largest herring-processing factories in Europe, with an annual production of about 50,000 MT, releases ~1,500 m^3 of wastewater daily (Steinke and Barjenbruch, 2010). Surimi production, which involves repeated washing of fish mince, uses more water than canning, curing, or freezing (Park, 2013). Water requirements for farmed production per metric ton of fish range from 1.5 to 6 m^3 for generic fish (Hall et al., 2011).

Problems Associated With Seafood Discards and Effluents

Loss of Nutrients

Seafood items are known to be rich in nutritionally valuable proteins, essential fatty acids, particularly long chain n-3 polyunsaturated fatty acids (omega-3 PUFAs), mainly eicosapentaenoic acid (EPA, C20:5; n-3) and docosahexaenoic acid, (DHA,C22:6; n-3), and vitamins and minerals (James, 2013; Venugopal and Gopakumar, 2017; Venugopal, 2018). Analysis of the compositions of more than 40 types of seafood processing discards showed that the discards have average contents of 60% proteins, 19% fat, and 22% ash, as shown in Table 1 (Islam et al., 2004). Shrimp head waste, on a dry weight basis, contains up to 65% proteins, 21% ash, and 18% chitin (Yan and Chen, 2015). Comparative dry-basis proximal analysis of shells of Argentine red shrimp (ARS) and southern king crab (SKC) showed both had 19–20% chitin. However, there were significant differences in contents of proteins and ash (18 and 48% for SKC and 26 and 55% for ARS, respectively). Analysis of shell and heads of ARS showed the highest lipid content (11%), with 5 mg and 158.8 µg of n-3 PUFAs and carotenoids per g, respectively. This suggested potential for recovering n-3 PUFAs and carotenoids from the ARS waste (Cretton et al., 2020). The raw heads, shells, and tails of Northern pink shrimp and spotted shrimp have crude proteins, which were rich in aspartic acid, glutamic acid, phenylalanine, lysine, and arginine. Their lipid contents ranged from 9.3 to 11.6%, while the contents of calcium, phosphorous, sodium, and magnesium were of 3,000, 400, 270, and 100 mg%, respectively. The contents of free amino acids (taurine, threonine, leucine,
**TABLE 1 |** Major components of seafood processing discards.

| Nutrients          | Composition |
|--------------------|-------------|
| **SOLID DISCARDS*** |             |
| Crude protein (%)  | 57.9 ± 5.3  |
| Fat (%)            | 19.1 ± 6.1  |
| Crude fiber (%)    | 1.2 ± 1.2   |
| Ash (%)            | 21.8 ± 3.5  |
| Calcium (%)        | 5.8 ± 1.3   |
| Phosphorous (%)    | 2.0 ± 0.6   |
| Potassium (%)      | 0.7 ± 0.1   |
| Sodium (%)         | 0.6 ± 0.1   |
| *Effluents*        |             |
| Total suspended solids* | 27–1,201 mg/ml |
| Ammoniacal nitrogen | 50 mg%     |
| Nitrate nitrogen   | 50 mg%      |
| Phosphate          | 95 mg%      |
| BOD$_5$*           | 179–276 mg/ml |
| COD*               | 458–1,717 mg/ml |

*Adapted from Islam et al. (2004). The values represent average values of 43 samples on dry weight basis, corrected to single decimal. *Typical values for effluents released from a seafood processing plant (Jamieson et al., 2017).

Tryptophan, tyrosine, and phenylalanine) of the processing by-products and edible parts were 2.0 and 1.7 g%, respectively (Heu et al., 2003). Lobster processing waste is responsible for appreciable loss of nutrients. Lobster liver may contain up to 41% proteins, on a dry weight basis, while its head contains meat up to 20% of the shellfish weight (Nguyen et al., 2017).

The characteristic features of seafood processing effluents are contents of total suspended solids (TSS), fats, oils, and grease (FOG), and pigments and minerals. The TSS includes proteinous matter (myofibrillar proteins, collagen, gelatin, enzymes, soluble peptides, and amino acids) in soluble, colloidal, or particulate form (Islam et al., 2004). Ching and Ghufran (2017) reported 2.2% total solids (consisting up to 550 mg% TSS and 260 mg% dissolved solids), 50 mg% each of ammonia and nitrate nitrogen, and up to 100 mg% of phosphate typical effluent. Tuna processing effluents contained TSS, fat, chemical oxygen demand (COD), and biochemical oxygen demand (BOD) at 1,570, 450, 11,100, and 6,600 mg per kg, respectively (Achour et al., 2000). These losses deprive the consumers of significant amounts of nutrients. It has been estimated that annual seafood discards in the US represent a loss of about 208 billion g of proteins and 1.8 trillion mg of n-3 PUFA (Love et al., 2015). These indicate that processing effluents, in addition to solid discards and by-catch, contribute to losses of nutrients.

**Environmental Impacts**

The food system has generally been considered a threat to the environment. The seafood industries consider huge volumes of by-catch, solid waste, and effluents a burden because of their potential to become environmental hazards. The industry dumps enormous amounts of by-catch in the ocean, while good amounts of solid wastes are disposed off as landfill or subjected to incineration. Ocean dumping causes reduced oxygen levels at the ocean bottom, burial or smothering of living organisms, and introduction of disease to the ecosystem of the sea floor (US EPA, 2017). Nguyen et al. (2017) observed that the disposal of lobster processing costs annually upward of about $7.5 million, and also presents an environmental burden to the lobster processors. Composting and ensilage of waste has been practiced, but they have limitations because of their longer process time, higher costs, and emission of volatile organic compounds. Anaerobic decomposition of seafood in landfill causes formation of methane (CH$_4$), ammonia (NH$_3$), and hydrogen sulfide (H$_2$S), which are detrimental to the environment (Xu et al., 2018). Landfill contributes to climate change, about 10 times larger than other waste disposal options, while composting has the largest impact on carcinogens (Gao et al., 2018). The TSS and FOG of process effluents are responsible for high BOD and COD values, indicative of their adverse influence on the oxygen balance and, in turn, the flora (Gonzalez, 1995). Furthermore, shortage of drinking water, eutrophication (growth of unwanted biota), biotic depletion, algal blooms, habitat destruction, water acidification, disease outbreaks, and extensive siltation of corals are other environmental hazards (Hall et al., 2011). The environmental problems and loss of nutrients associated with seafood process effluents have been pointed out recently (Venugopal and Sasidharan, 2021).

**Measures for Sustainable Seafood Processing**

Food sustainability demands optimal uses of resources for maximum benefits, including economic viability. The challenges of sustainable seafood processing are linked to reducing environmental pollution, conservation of water, and prevent losses of nutrients. Improving waste utilization is essential for a sustainable industry to prevent or minimize the environmental impact (López-Pedrouso et al., 2020). The problems of environmental hazards and nutrient losses facing the seafood industry can be addressed by measures such as selective trawling to reduce by-catches, appropriate treatments of wastes and effluents, and valorization of wastes by recovery of useful ingredients. The major advantages with respect to waste treatments are reduction of environmental hazards, conservation of water, and isolation of commercially valuable ingredients and improvement of the economy. In view of seafood-related environmental hazards, a need for a biological solution for the disposal of the seafood processing discards has been recognized (Pal and Suresh, 2016). Failure in these efforts not only leads to loss of potential revenues but also increases the cost of waste disposal and also public health problems (Etemadian et al., 2021). The recent United Nations Conference on Sustainable Development acknowledges the global importance of food losses and food waste. The Conference aims to halve per capita global food waste and reduce food losses along production and supply chains, including post-harvest losses, by 2030 (FAO, 2020).

**Conventional Processes for Waste Valorization and Their Limitations**

Conventional processes for valorization of seafood discards and effluents are based on chemical and physical methods. These...
processes invariably have several limitations. For instance, the process of chitin extraction from crustacean shells involves initial alkali treatment generally with 5 M sodium hydroxide for deproteinization followed by hydrochloric acid treatment for demineralization, which involves decomposition of calcium carbonate in the shells. Alkali treatment can result in the hydrolysis of chitin and also partial decetylation of chitin. The process is also corrosive. Further, it requires large volumes of fresh water to wash off alkali and acid from the treated shells, releasing harmful wastewater (Mao et al., 2017; Yadav et al., 2019). Alkali extraction of proteins can lead to the loss of certain amino acids (Venugopal, 2006). Similarly, the traditional solvent extraction of fish oil can cause its oxidation, which is rich in unsaturated fatty acids. de Oliveira et al. (2016) extracted oil from tuna by employing chemical refining. The process consisted of degumming, neutralization, washing, drying, bleaching, and deodorization. Although chemical refining was successful, temperature and chemical reagents favored the removal of PUFA from the oil. In view of these drawbacks, interest in alternate green processes is growing. Biological processes can have minimal environmental impacts, be cost-effective and safe, and have minimum adverse impacts on the properties of the isolated components. These processes, therefore, offer an economic and versatile way to transform and concentrate waste and wastewater into valuable products.

Biotechnology holds promise for novel waste treatment and resource recovery processes, diversification of value-added products, and in quality assurance (Pleissner and Lin, 2013; Chandrasekharam, 2015). Such processes are supported by novel green techniques for industrial recovery of biomolecules from seafood by-products and discards. The extraction efficiency can vary highly depending on the food matrix, the target compounds, and methods of extractions. Therefore, the choice of green extraction technique depends essentially on the matrix and the target compound features (Bruno et al., 2019). Such approaches can also lead to a bio-based economy (Puyol et al., 2017). This article will examine potentials of biological processes depending on bioconversions of components in seafood discards, including effluents. Prospects of algal biotechnology and bio-refinery approaches will also be discussed.

**BIOCONVERSION REACTIONS FOR VALORIZATION OF FOOD WASTES**

Bioconversion reactions are biological methods, which are carried out to initially detach food components from their matrices. The detached components can be recovered and purified by suitable techniques. Two major bioconversion processes have been recognized, which employ either microorganisms or externally added enzymes. These help detach the components from the food matrices. Both microbial and enzymatic processes are environmentally friendly, cost-effective, and safe, unlike most conventional chemical extraction processes. Microbial fermentation of the waste results in the production of hydrolytic enzymes by the organism, which causes bioconversions of the food components. Exogenous enzymes, on the other hand, cause a direct release of the components from the food matrix. A number of microorganisms, particularly lactic acid bacteria (LAB), have been used for fermentation. The use of algae is a novel approach in this respect. The algae-induced biotransformations help not only with the treatment of wastewater but also production of nutrient-rich biomass, which is useful for a variety of applications. In addition to microbial and enzymatic methods, biophysical processes such as modifications of pH and temperature can favor the release of components such as proteins and lipids from the food matrices. Figure 1 depicts a schematic representation of the bio-conversion processes for valorization of seafood discards. These aspects will be discussed in detail.

**Microbe-Mediated Bioconversions**

The microbe-mediated bioconversion is termed as fermentation, which is safe, environmental-friendly and energy savvy. The process makes use of live microorganisms (bacteria, fungi, mycelium, or microalgae) to convert raw materials into products with desired qualities. Fermentation can be traditional, biomass or precision types. Traditional fermentation has been practiced for a few centuries. Since the 1980s biomass fermentation has emerged in the food industry for the production of cell mass for further use as sources of enzymes, flavors, food, biomaterials, therapeutics, fuels and in recent times, as sources of alternative proteins to develop cultivated meat formulations. Precision fermentation is intended to produce specific functional ingredients using tailor-made microbial hosts (GFI, 2020). Fermentation by lactic acid bacteria (LAB) has been used as a popular method for the development of fermented fishery products (Anihouvi et al., 2012). The efficiency of lactic acid fermentation depends on the type of organism, inoculums size, initial pH and pH attained during fermentation. The lactic acid formed during sugar breakdown creates low pH, which, in turn, suppresses growth of spoilage causing microorganisms and enhances activity of acid proteases, optimally acting on seafood proteins, many of which remain bound to chitin, lipids and carotenoids.

The bioremediation using microorganisms and their aggregates is recognized to be an efficient low-cost green process. The technology of microbial conversion also provides a potential way to isolate and exploit compounds of biotechnological potential (Wang et al., 2019). The microorganisms used for the purpose may be aerobic, anaerobic, or facultative including bacteria, fungi, and protozoa. Microbial fermentation processes can be under solid state, submerged or liquid state, anaerobic, batch, continuous, or fed batch conditions. The process is influenced by factors such as the nature of the starter culture, time, pH, and substrate composition. Fed-batch is a commonly used means for the production of microbial biomass, ethanol, organic acids, antibiotics, vitamins, enzymes, and other compounds in which the culture medium is added continuously or in pulses to reach the maximum volume. The advantages of fed-batch over the conventional batch operation include a higher biodegradation rate, higher productivity, higher dissolved oxygen in the medium, and decrease in fermentation time (Chandrasekharam, 2015; Puyol et al., 2017).

Microbial growth results in the production of hydrolytic enzymes such as proteases, lipases, and chitinases. Proteases and chitinases cause demineralization, deproteination, and
proteolysis in the substrate. Chitinases catalyze the cleavage of the β-1,4-O-glycosidic linkages in chitin. Lipases function as triacylglycerol hydrolases and also catalyze synthesis of ester compounds. Fish fermentation, which is traditionally used to increase fish shelf-life, results in the formation of bacteria metabolites of interest. Lactic acid bacteria (LAB) have been used for a long time for the development of fermented fishery products (Anihouvi et al., 2012). Fermentation by LAB produces lactic acid; the low pH enhances acid proteases which act on proteins bound to chitin, lipids, and carotenoids. Fermentation can be applied for the production of cell mass, enzymes, flavors, food additives, and a range of other high value-added products. For example, fermentation of shrimp shell waste by symbiotic LAB such as S. thermophilus, L. acidophilus, and L. bulgaricus rapidly decreased pH to about 4.2 and promoted the removal of calcium and protein, with 91.3% calcium, 97.7% protein, and 32.3% carotenoid removed from shrimp waste after 168 h fermentation activated acid proteases gave bioactive peptides of size between 1,000 and 10,000 Da (Shan et al., 2011). LAB-induced fermentations bring about diversity into foods, make otherwise inedible foods products edible, enhance nutritional value, decrease toxicity, preserve food, and decrease cooking time and energy requirements. The technology is safe, environmentally friendly, and does not consume much energy.

Microbe-assisted bioconversions are ideal for the bioremediation of seafood processing waste and production of aquafeed and fertilizer. Applied to fish by-products, fermentation gives rise to quality protein hydrolysates and oil and produces antioxidant compounds (Marti-Quijial et al., 2020). Fish offal and a mixture of sawdust and wood shavings in equal proportions were subjected to composting by placing them in an open structure with passive aeration. Solid state fermentation converted the waste into a highly nutritive fertilizer with a nitrogen content as high as 12% (Wang et al., 2019). Dried skipjack tuna waste (red meat, gills, viscera, fins, etc.) was mixed with 25% wheat flour and fermented with L. plantarum and B. licheniformis for 14 days. The proximate analysis showed significant changes in the composition of L. plantarum. The fermented product can be used as a nutritive aquafeed ingredient (Hena et al., 2009). Conversion of fish waste to liquid fertilizer was achieved with mixed microorganisms, resulting in about 28% degradation of fish waste. The product was stable against putrefaction for 6 months at ambient temperature (Dao and Kim, 2011). The microbe-assisted aerobic bioprocess of aquaculture solid waste for 15 days at 35 °C and at pH 6.0–6.5 maximized nitrogen bioconversion in the form of ammonium ions (NH\textsubscript{4}\textsuperscript{+}) (Khiari et al., 2019). Fermentation is a viable alternative to chemical treatment for the extraction of collagen (Song et al., 2021).

Rashid et al. (2018) fermented shrimp-shell powder with B. cereus to produce sugar, antioxidant, and DNA protective compounds. The fed-batch biodegradation was operated in a 5-L bioreactor for 96 h according to three time pulse-feeding strategy. On the basis of the equal working volume of 3 l, the fed-batch biodegradation showed a better production of the target compounds than the batch biodegradation, with higher cell density and a shortened biodegradation period. The maximum values of the target compounds were about 0.3 mg per ml of reducing sugar and 92 to 98% antioxidant activities. Fed batch fermentation gave ~3–12% higher values compared with batch biodegradation.
Microorganisms can treat seafood industry process effluents in reaction systems such as activated sludge, aerobic lagoons, trickling filters, and rotating disc contactors. In the commonly used activated sludge system, the sludge consisting of an optimized, mixed flora of microorganisms degrades the organic materials in the presence of dissolved oxygen, thereby decreasing the BOD of the effluent (Gonzalez, 1995; Choudhury et al., 2010). An aerobic continuous bioreactor treated high saline fish processing wastewater for 8 h, which removed the offensive odor of the effluent (Ching and Ghufran, 2017). Anaerobic digestion (AD), a popular green technology for waste treatment, involves fermentation of the material in the absence of molecular oxygen with the formation of CO₂, hydrogen, and/or acetic acid; reduction of the CO₂ and acetate leads to the production of methane. AD of tuna processing effluents involved a decanter to remove the fats and the TSS, an anaerobic digester, and an activated sludge aerated bioreactor. The integrated system helped with the removal of up to 95% of the COD (Achour et al., 2000). AD of seafood industry effluents in a dissolved air flotation system (DAF) removed organic contents. The process flow consists of the separation of effluent in DAF and treatment of clarified water in a double nitrification-de-nitrification stage. AD of solids separated during the DAF process produces biogas and significantly reduces sludge volume (Fluneau, 2019).

Algae-Based Bioconversions

Microalgae such as Chlorella, Spirulina, Dunaliella, diatoms, and cyanobacteria, commonly referred to as blue green algae, are the main algae grown commercially as sources of functional materials in natural foods. These organisms are bestowed with a high growth rate in nutrient media under phototrophic (light and CO₂) conditions. Their digestive actions allow the degradation of organic contents, unused food, and excretory products together with the removal of CO₂, NH₃-N, CO₂, and H₂S, thereby ameliorating environmental pollution (Payul et al., 2017; Gifuni et al., 2019). The phototrophic algae can be cultivated in open ponds or in closed photo-bioreactors, or heterotrophically in closed systems. One of the major advantages associated with the open ponds is its low production and operating costs. However, the limitations include uneven light availability and distribution within the pond. Heterotrophic cultivation in closed systems eliminates the requirement of light, but heterotrophic culture is prone to contamination by other microbial species (Nigam et al., 2020).

Microalgae are promising agents for bioconversions of food, fishery, and agricultural waste into biomass rich in bioactive compounds (Das, 2015). The algal mass (referred as single cell proteins, SCP) can contain up to 60% proteins, good amounts of oil, and also polysaccharides, minerals, and pigments including chlorophylls, carotenoids, and phycobiliproteins. Stringent nitrogen limitations stimulate algae to produce more lipids, as high as 75% with high n-3 PUFA contents (Stengel and Connan, 2015). SCP has found wide applications as sources of bioactive peptides, plant growth stimulants, animal feeds, food additives, cosmeceuticals, drugs, and as probiotics in aquaculture. SCP can also replace expensive soy meal and fishmeal in animal and aquaculture feeds (Sharma and Sharma, 2017; Caporgno and Mathys, 2018; Smarason et al., 2019). Cultivation of microalgae in wastewater offered the highest atmospheric carbon fixation rate (1.83 kg CO₂/kg biomass) and rapid biomass productivity—40–50% higher than terrestrial crops (Shahid et al., 2020). Growth of microagal biomass has been estimated to require 200–1,000 liter of water per kg of dry biomass. This suggests comparable volumes of seafood industry effluents could be treated for producing equivalent amounts of microalgae as SCP (de Farias and Barbera, 2018). Batch cultivation of Chlorella sp. in seafood processing water gave a biomass yield of 896 mg per l (Gao et al., 2018). 



Enzyme-Assisted Bioconversions

Processing with an enzyme holds enormous potential in waste management. Enzymes can mitigate hazards of conventional chemical transformations for resolution of food waste-related environmental problems, help production of novel compounds, and function as analytical tools for food quality assessment. Enzymes can be included as additional processing aids to conventional processes or can be exclusively used to upgrade existing technologies in seafood processing. The advantages of enzymes are their low energy requirements, safety, and low-cost (Venugopal, 2006; Chandrasekharan, 2015; Fernandes, 2016;
Hydrolases, which include carbohydrases, proteases, and lipases, are popular enzymes in biotechnology. Enzymatic hydrolysis of protein from aquatic by-products and livestock, poultry, and plants offer novel products with applications in foods, pet feed, pharmaceutical, and other industries (Etemadian et al., 2021). Specific, energy-efficient, and easily controllable enzymatic techniques using proteases, glycoside hydrolases, lipases, transglutaminases, and other enzymes are emerging as bio-processing techniques for seafood processing (Shah et al., 2016; Huang et al., 2017).

DOWNSTREAM PROCESSING FOR THE RECOVERY OF SEAFOOD COMPONENTS

The by-catch and the various seafood processing discards are rich in balanced proteins, collagen, enzymes, lipid, carotenoids such as astaxanthine and β-carotene, polysaccharides including chitin, glycosaminoglicans, and various minerals. These can be recovered by coupling the various bioconversion processes, discussed above, supported by marine biotechnology-based downstream processes. These processes are mostly mild, energy-efficient, safe, and environmentally friendly. Green techniques include pressurized liquid, sub-critical, super-critical, enzyme-mediated, microwave-, and ultrasound-assisted extractions (Muffler and Ulber, 2005; Freitas et al., 2012; Chavez et al., 2013). Membrane bioreactors integrate reaction vessels with membrane separation units for producing materials such as peptides, chito-oligosaccharides, and PUFA from seafood discards (Kim and Senevirathne, 2011).

The past few years have seen notable interests in seafood-derived compounds for varied applications including food, pharmaceutical, agriculture, and other industries. For example, bioactive compounds from lobster processing by-products can be extracted using microwave, ultrasonic, and supercritical fluid extraction. The proteins, chitin, lipids, minerals, and pigments recovered from lobster processing by-products possess several functionalities and bioactivities, useful for their applications in water treatment, agriculture, food, nutraceutical, pharmaceutical products, and biomedicine (Nguyen et al., 2017). The diverse compounds that can be extracted from seafood processing discards can be grouped into four classes, depending upon their chemical nature. The major classes include nitrogenous, lipid, polysaccharide, and mineral-based compounds. Some of the individual compounds under each class are given in Table 2. In addition, a multitude of derivatives can also be developed from many of these components. These include bioactive peptides, gelatin, n-3 PUFA, glucosamine, chitosan, and its various derivatives. Detailed aspects of preparations, properties, and applications of components of seafood processing discards will not be discussed here; nevertheless, a very brief mention may be made. Fish proteins and protein hydrolyzates have the potential to be used as a protein supplement, in the fortification of foods, and as sources of bioactive peptides. Fish oil has numerous health benefits and can also be used to impart functional properties to food products. Chitosan and their oligosaccharides are applied as antioxidants, antibacterial and antifungal agents, and functional compounds in food, pharmaceutical, and other industries. Gelatin is used for the purposes of gelling, edible coating, emulsification, and microencapsulation. The reader is referred to a few recent articles for details (Karim and and Bhat, 2009; Yu and Gu, 2013; Venugopal and Lele, 2014; Vidanarachchi et al., 2014; Muxika et al., 2017; Sasidharan and Venugopal, 2019; Shahidi et al., 2019; Ashraf et al., 2020). Nawaz et al. (2020) recently observed a need to focus on bioavailability, interaction with other ingredients, nutritional, biotechnological, and sensorial aspects, and other factors of seafood-derived compounds that can significantly favor valorization of fisheries by-products. The following discussion will focus on bioconversion processes for extraction of major classes of compounds seafood discards and effluents.

| TABLE 2 | Major seafood-derived industrial components and nutraceuticals. |
|----------|------------------|
| **Finfish** | **Shellfish (Crustaceans and mollusks)** |
| **Nitrogenous compounds** | **Nitrogenous compounds** |
| Proteins (myofibrilar) | Proteins |
| Collagen | Collagen |
| Gelatin | Gelatin |
| Protein hydrolyzate | Protein hydrolyzates |
| Bioactive Peptides | Bioactive peptides |
| Enzymes | Enzymes |
| Proteases, Collagenases, Transglutaminases, Chitinases, β-1,3-Glucanase, β-Galacto-sidase, Lysozyme, Catalase, Glutathione, Peroxidase, and others | Lysozyme from crab, scallop, others |
| **Lipid-based:** | Lipid-based: |
| PUFA rich oil | PUFA rich oil |
| Squalene, Squalamine | Carotenoids |
| **Mineral-based:** | Astaxanthin |
| Biological/calcium | B-β-carotene |
| Phosphopeptides | Polysaccharide-based: |
| Hydroxyapatite from tuna bone | Chitin, Chitosan, Chitosan derivatives, Glucosamine, oligosaccharides, novel polysaccharides from mussel |
| **Polysaccharide-based:** | Mineral-based: |
| Glycosaminoglycan, Chondroitin, Dermatan sulfate, Hyaluronic acid | Calcium carbonate, Calcium lactate |

**Nitrogenous Compounds**

Diverse proteins, peptides, amino acids, and their co-products constitute the nitrogenous fraction. Scarcity of nutritional proteins in many parts of the world necessitates novel and economical processes to recover them from unexplored sources (Henchion et al., 2017). Seafood processing discards, having a maximum of about 60% proteins, can be good sources of proteins, which can be recovered while retaining most of its native properties. Bioconversion processes for protein
recovery from the discards have been developed. These make use of biophysical changes that induce their coagulation and precipitation of the macromolecules. These are discussed below.

**Isoelectric Solubilization Precipitation (pH Shift Process)**

Isoelectric solubilization precipitation (ISP) is a gentle bioprocess. This involves homogenization of underutilized fish or processing discards with either dilute acid (pH 2.5–3.5) or alkali (pH 10.8–11.5). The treatment dissolves sarcoplasmic and myofibrillar proteins, while insoluble impurities such as bone, skin, oil, and membranes are removed. Up to 90% of the dissolved proteins are precipitated by raising the pH of the solution to their isoelectric pH of 5.2–6.0. The proteins are concentrated by centrifugation or filtration. The process, as depicted in Figure 2, is ideally performed at 10°C or below, to avoid denaturation of the protein (Hultin et al., 2005). Recovery of proteins by the ISP process can be enhanced by coupling it with high intensity sonication, electro-flocculation, and ultrafiltration. Sasidharan and Venugopal (2019) summarized studies on ISP-based protein recovery from various species of finfish and shellfish, their discards, by-catch, and also process effluents. Some of the fishery sources included mackerel, catfish, rockfish, Pacific whiting, rainbow trout, Atlantic croaker, channel catfish, bullhead catfish, shrimp, crab, mussel, and squid. The fish protein isolates (FPIs) have protein contents of at least 65% and fat below 2%. FPIs differ from conventional surimi, which is a concentrate of fish myofibrillar proteins obtained by repeated washing of fish meat mince (Park, 2013). Unlike surimi, which is only myofibrillar proteins, particularly myosin and actomyosin, FPIs contain sarcoplasmic proteins along with the refined concentrate of myofibrillar proteins. The FPIs generally retain biochemical, nutritional, and functional properties of the native proteins, which make them valuable raw materials for applications such as the development of restructured food products, protein supplements, and bioactive peptides (Sasidharan and Venugopal, 2019).

Apart from myofibrillar proteins, collagen is another protein from marine sources. Marine collagen is a promising biocompatible alternative to mammalian collagen, particularly in biomedical and food applications. Collagen-based novel functional food ingredients contain a nutritional benefit, such as essential and non-essential amino acid, to improve the quality of different food products. It can also be used as a natural antioxidant and texturizing agent that can reduce the utilization of chemical food additives and may be able to fulfill the consumer demands for safe and green food products (Pal and Suresh, 2016).

Collagen and collagen hydrolysate (CH) were recovered from the bone and skin containing residues emerging during the ISP process from silver carp. Isolated collagen maintained their triple-helical structure and was characterized as type I collagen. Pepsin-hydrolysis and sequential hydrolysis by pepsin and trypsin degraded all heavy molecular weight chains of collagen; sequential enzyme treatment yielded a higher degree of hydrolysis. When CH was added to silver carp protein isolate prior to gelation, the gel properties were dependent on the molecular weight of the added CH. More hydrolyzed collagen emerging from sequential hydrolysis improved the water holding capacity of the gel while reducing its breaking force. The results suggest that residue from pH-shift processing of fish can be used for isolation of functionally active collagen and CH (Abdollahi et al., 2018). Tilapia type I collagen is biocompatible and can be used as an effective biodegradable scaffold biomaterial for regenerative medicine (Hayashi, 2020). Gelatin is extracted from collagen generally by pre-treatment with dilute NaOH, followed by swelling with dilute acetic acid and then by warm (45°C) water (Vázquez et al., 2019). Gelatin prepared from collagen of skins and bones of various marine and freshwater fishery sources have good gelling properties. Gelatin extract of big eye tuna skin had glycine, up to 32% of total amino acids, and hydroxyproline together with proline and alanine. Rheological studies revealed Newtonian and shear thickening properties of the gelatin. The tuna gelatin could be useful for the formulation of functional foods and nutraceutical and biomedical applications (Dara et al., 2020).
Mild Acid Induced Gelation
This process makes use of the ability of muscle structural proteins to undergo gelation under mild acidic conditions when water is strongly bound to the protein matrices. The process of extraction involves initial mechanical deboning of by-catch or fish discards such as heads and frames. The meat mince is washed twice with chilled (0–5°C) water, followed by homogenization of the washed mince in equal amounts of fresh chilled water. The pH of the homogenate is lowered to 3.5–4.0 by drop-wise addition of weak acid, such as acetic acid, which induces gelation of the proteins. The gelation process is associated with a fall in viscosity of the homogenate; the viscosity fall can be enhanced by mild heating to 50°C. Proteins in the low-viscous dispersion are highly stable as they cannot be precipitated by heating even at temperatures as high as 100°C. Such thermo-stable dispersions have been prepared from Atlantic herring, Atlantic mackerel, threadfin bream, and shark. Shark meat, however, is an exception. The homogenate of washed shark meat in water exhibited an increase in viscosity during the course of acidification to pH 4 to 5 (Venugopal, 2017). In the case of capelin, thermo-stable dispersion could be prepared without the need for acidification. The thermo-stable protein dispersions prepared from different fishery products can have varied applications, such as preparation of fish protein powder, as protein coating of fresh fish to extend their refrigerated shelf life, preparation of fermented sauce, or for the development of edible packaging (Venugopal, 1997). Meat recovered from by-catch fish can be used for protein dispersion or can be a resource for value added food products such as urimi, sausages, or fermented products, among others (Venugopal and Shahidi, 1998).

Flocculation
Proteins, which are present in suspended or dissolved states in process effluents, can be flocculated and precipitated by food grade polysaccharides such as carrageenan, alginate, and carboxy methylcellulose, which are then subjected to concentration by filtration, sedimentation, and/or centrifugation (Forghani et al., 2020). Proteins from herring industry processing effluents were recovered using electroflocculation (EF) and ultrafiltration (UF). EF and UF recovered up to 80% proteins. The highest protein and fatty acid contents of the effluent were 12.7 and 2.5 g per kg, respectively. Leucine and glutamic acid/glutamine were the dominating amino acids while calcium and magnesium were the dominating trace elements. The proteins had good foaming and emulsifying properties, which make them good functional additives (Gringer et al., 2015). Biomass from cooking wastewaters of snow crab was concentrated by membrane filtration. The concentrate had 59% proteins and contained desirable flavor compounds. The extract can be a natural aroma for the food industry (Tremblay et al., 2020).

Enzymatically Hydrolyzed Proteins
Proteolytic enzymes from various sources, including microorganisms (such as alcalase, flavourzyme, and protamex), animal (collagenase, proteinase, serine-protease, neuramidase, and trypsin), and plants (papain, bromelain, and ficin), can extract proteins from seafood processing discards as fish protein hydrolyzates (FPHs). The ideal treatment conditions are incubation temperature, 35–37°C; enzyme to substrate ratio, 1–50; and incubation up to 24 h. The degree of hydrolysis determines the properties of the hydrolyzate, such as solubility, water-holding capacity, emulsification, and foam-forming ability, and the contents and chemical nature of peptides formed. The FPH can be concentrated by spray drying or ultrafiltration. They generally show a beneficial effect on growth performances and feed utilization at low inclusion levels (Chalamaiah et al., 2012; Vijaykrishnaraj and Prabhasankar, 2015). Fish frames without heads from Atlantic salmon and Atlantic cod were treated with commercial proteases for 2 h. Salmon treated with alcalase and cod treated with pepsin yielded 64 and 68% proteins, respectively (Lisaset et al., 2003). Proteins, together with chitin and astaxanthin, were extracted from shrimp using enzymatic treatment with alcalase and pancreatin. Alcalase was more efficient than pancreatin, which increased recovery of proteins from 57.5 to 64.6% and of astaxanthin from 4.7 to 5.7 mg astaxanthin per 100 g of dry waste, at a degree of hydrolysis of 12%. An increase in the DH from 6 to 12% resulted in 26% to 28% protein recovery (Routray et al., 2019). Alcalase hydrolysis of the industrial waste from Xiphopenaeus kroyeri shrimp allowed 65% protein recovery in the form of hydrolysates (Holanda and Netto, 2006).

Bioactive peptides are specific protein fragments that can have high nutraceutical potentials and may be able to address important public health issues like obesity, stress, hypertension, and more. Such peptides have been produced from hydrolysates of several fish and shellfish. Their potential functions include antimicrobial, antiviral, antitumor, antioxidative, antihypertensive, cardioprotective, anti-ammesic, immunomodulatory, analgesic, anti-diabetic, antiaging, appetite-suppressing, and neuroprotective activities (Chalamaiah et al., 2012; Vijaykrishnaraj and Prabhasankar, 2015). These activities are related to the sequence, composition, and type of amino acids in the peptides. FPHs can be used as a source of bioactive peptides with potentials for use as functional food ingredients industry. Tonon et al. (2016) prepared protein hydrolysate from the shrimp cooking effluents by enzymatic hydrolysis and ultrafiltration. The hydrolyzate prepared at 75°C and at pH of 9.0 have essential amino acids that can satisfy people’s recommended daily needs. The preparation had significant antioxidant activities. Hypertensive and antioxidant peptides were prepared by enzymatic hydrolysis of proteins from cuttlefish wastewater. The proteins were initially concentrated by ultrafiltration (Amado et al., 2013). Pepsin soluble collagen (PSC) was enzymatically hydrolyzed, and the resultant hydrolysates were ultrafiltered and characterized. Electrophoretic patterns showed the typical composition of type I collagen, with denaturation temperatures ranging between 23 and 33°C. In terms of antioxidant capacity, results revealed significant intraspecific differences between hydrolysates, retentate, and permeate fractions when using β-Carotene and DPPH methods (Blanco et al., 2017). The presence of both omega-3 fatty acids and ACE-inhibitory peptides in squid hydrolyzate suggested its nutraceutical potential (Apostolidis et al., 2016). In order to use and commercialize bioactive hydrolysates and peptides as
Enzymes from seafood processing discards for seafood processing.

| Enzymes                        | Discard source                                                                 | Seafood processing applications                                                                 |
|--------------------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|
| Proteases (pepsin, trypsin,    | Seafood processing discards (finfish intestines, shrimps heads, etc.)          | Tenderization of squid, Isolation of seafood flavor Recovery of proteins including collagen and chitin |
| chymotrypsin, collagenases,    |                                                                                  | Pigments from shell discards, Production of fish protein hydrolyzates                                |
| acid and alkaline proteinsases,|                                                                                  | Enhancement of digestibility of aquafeed protein ingredients                                        |
| cathepsins, peptidases, and    |                                                                                  | Production of omega-3-enriched triglycerides, Improvement of flavor                                |
| others)                       |                                                                                  |                                                                                                  |
| Lipases                       | Discards of Atlantic cod, seal, salmon, sardine, Indian mackerel, red sea bream.|                                                                                                  |
| Chitinases                    | Shellfish, squid liver, octopus                                                | Deacetylation of chitin to chitosan                                                               |
| Amylase                       | Abalone                                                                        | Peeling and deveining of shrimp                                                                  |
| β-1,3-Glucanase               | Abalone, scallop, tilapia, sea cucumber                                         | Removal of anti-nutritive effects of non-starch polysaccharides in aquafeed                       |
| β- galactosidase              | Arctic scallop shell, crab shell                                               |                                                                                                  |
| Lysozyme                      | Marine mussel and other organisms                                              | Bacteriostatic agent.                                                                             |
| Catalase, glutathione peroxidase|                                                                                   | Antioxidants, Preserve fish quality, in combination with lysozyme                                |
| Transglutaminases             | Various fishery sources                                                         | Texturization, encapsulation of nutraceuticals, development of protein sheets and surimi-analogs, edible films |

Summarized from Venugopal (2016).

The global production of fish oil is around one million tons, predominated by cod liver oil (Bimbo, 2007). Fish oils are rich in omega-3 PUFA and vitamins A and D. The oil of fish species, such as Atlantic mackerel, shark, anchovies, menhaden, and Atlantic sardine, can have up to 35% omega-3 fatty acids, with EPA and DHA at around 10% of the oil (Venugopal, 2009). Conventional, fish oil is extracted by wet reduction method involving cooking, pressing, and filtration. The extracted oil is purified by carbon treatment, degumming, alkali refining, deodorization, and stabilization by antioxidants for prolonged storage, while the protein-rich press liquor is used as animal feed (Venugopal, 2009). There have been concerns fueled by ominous predictions of depletion of several oil-rich oceanic fishes, which necessitates better exploitation of fish discards as oil resources.

LIPIDS AND CO-PRODUCTS

The enzymatic process disrupts the tissue and membranes under mild conditions to release the oil from fish by-products, such as liver and roe (Dumay et al., 2004). Treatment of salmon heads and other byproducts by commercial proteases (alcalase, neutrase, and flavourzyme) for 2 h released 17% oil, which contained 11.6 and 5.6% of DHA and EPA, respectively (Routravy et al., 2019). Oil from sardine was obtained at 5.5, the iso-electric
pH the fish meat, which was adjusted by citric acid. The separated oil had good n-3 PUFA contents and exhibited high oxidative stability (Okada and Morrissey, 2007). Alcalase-based extraction of oil from tuna was conducted for 120 min at 60°C and pH 6.5 at an enzyme–substrate ratio of 1:200. The enzyme-extracted oil had lowest acidity and peroxide values and higher levels of EPA and DHA contents than chemically refined oil (de Oliveira et al., 2016). Hydrolysis of shrimp waste with alcalase gave an oil yield of 28.6 μg per g waste (Sachindra and Mahendrakar, 2011). The hydrolyzate of squid processing by-products had EPA and DHA at 16.9 and 29.2% of oil, respectively. About half of the oil was comprised of phospholipids (Apostolidis et al., 2016).

Lipases have grown in importance due to their ease of availability and possibilities for product modifications. *Candida rugosa* lipase was used to concentrate fatty acids in the glyceride fraction of the oil. By controlling the degree of hydrolysis, two products were obtained, one having 50% n-3 PUFA, and the other having 40% DHA and 7% EPA. The glyceride from these reactions was converted back to triglycerides using *Rhizomucor miehei* lipase catalyzing partial hydrolysis and esterification (Moore and McNeill, 1996). Lipolysis of salmon oil by a commercial lipase gave a mixture of free fatty acids and acylglycerols. A hydrophobic membrane was used to separate high melting saturated fatty acids from low melting acylglycerols. The sum of total PUFA increased from 42% in the crude oil to 47% in the filtrate with increase of DHA and EPA contents from 9.9 to 11.6%, and from 3.6 to 5.6%, respectively (Linder et al., 2005).

**Carotenoids**

Carotenoids provide red and orange colors to some foods. The pigments are present in shellfish, krill, shrimp, crab, crayfish, and also in salmon and trout. The red-orange color of cooked crustaceans is attributed to partial or complete separation of astaxanthin from the protein moiety to which it is attached in the native state. Carotenoids may be hydrocarbons, such as β-carotene or xanthophylls, or oxygenated derivatives, such as astaxanthin, astacene, canthaxanthin, cryptoxanthin, lutein, neoxanthin, violaxanthin, and zeaxanthin. Astaxanthin (3,3′-dihydroxy-β, β′-carotene-4,4′-dione) and canthaxanthin (β-carotene-4,4′-dione) have been used in aquafeed for many years in order to impart the desired flesh color in farmed salmonids. The shells of shrimp, prawn, crayfish, krill, crab, and lobster are important sources of astaxanthin, bound to free protein or chitin, and range from 40 to 200 μg per g, dry weight. Canthaxanthin is present in crayfish, mytiloxanthin in mussel, and macractanin and fucoxanthin are present in clams (Sowmya and Sachindra, 2015; de Carvalho and Caramujo, 2017).

Microbial fermentation-based bioconversion methods can extract carotenoids from crustacean shells, giving better yields than conventional solvent extraction. During bacterial fermentation processes, the proteins and minerals present in the shrimp shells are effectively removed, thereby increasing the extraction efficiency of the pigments without any change in quality. Current extraction methods make use of proteolytic enzymes such as trypsin and alcalase and fermentation by LAB and other microorganisms (Prameela et al., 2017; Routray et al., 2019). Fermentation of shrimp shell waste using the lactic acid bacterium *Pediococcus acidolactici* under optimal conditions resulted in 98% deproteination, 72% demineralization, and carotenoid recovery of up to 78% (Bhaskar et al., 2007). Shrimp waste was hydrolyzed with alcalase at optimal conditions of 0.75% of enzyme for 150 min at 37°C. The recovered carotenoids were extracted in sunflower oil at an oil to hydrolyzed waste ratio of 2:1 at 70°C for 90 min (Sachindra and Mahendrakar, 2011). Caroteno-protein from pink shrimp (Parapeneaus longirostris) waste was extracted by trypsin treatment for 1 h at 25°C. The recovered caroteno-protein fraction after freeze-drying contained about 71% protein, 16% lipid, 8% ash, 2% chitin, and 87 μg astaxanthin per g of the sample. Enzymatic hydrolysis of the protein–pigment complex allows studies on pigment absorption, stability, and application (Silé et al., 2012). Trypsin from bluefish was used to extract carotenoids from black tiger shrimp shells. The extract also contained 70% protein, 20% lipid, 6.6% ash, 1.5% chitin, and 87.9 μg with total astaxanthin per g sample (Klomklao et al., 2009). Autoalysis of shrimp heads resulted in the recovery of 195 μg carotenoids per g wet shells (Cahu et al., 2012).

Processing effluents can also be used as a medium for the production of carotenoids. Employing non-sterilized mussel processing wastewater as a low-cost substrate for yeast fermentation, the green microalga (*H. pluvialis*) was cultivated in fish effluents for the production of astaxanthin with significantly greater antioxidant capacity than the synthetic one (Shah et al., 2016). Similarly, mussel processing water was used for the production of astaxanthin by *Xanthophyllumyces dendrorhous* (Amado and Vázquez, 2015). With shrimp waste, being highly perishable and seasonal, the fermented carotenoid-rich liquor can be prepared as per the availability of the waste and can be stored up to 75 days under normal storage conditions. Carotenoids from the liquor can be extracted by ultrasonic or supercritical CO2 extraction, or can be extracted in palm oil or other vegetable oils or by high pressure chemical extraction (Sowmya and Sachindra, 2015). Astaxanthin is stable at 70–90°C in rice bran, ginger, and palm oils. Astaxanthin has important applications in the nutraceuticals, cosmetics, food, and aquaculture industries (Ambati et al., 2014; Shah et al., 2016).

**POLYSACCHARIDE-BASED COMPOUNDS**

The dry shell discards of crab, shrimp, and lobster may contain up to 70% chitin; dry squid skeleton pen and krill shells have a lower chitin content of 40%. The commercial process of extraction of chitin from crab and shrimp shells involves three steps: demineralization of dried and pulverized shells by dilute hydrochloric acid; deproteinization by dilute alkali; and decoloration, washing, and drying. Chitin is deacetylated to chitosan using 30 to 60% (w/v) sodium or potassium hydroxide at 80–140°C. The yield of chitin is about 25% of dry shell, and the yield of shrimp chitosan about 77% of the crude chitin (Dima et al., 2017). The chemical methods and the high treatment temperatures have influence on molecular weight, degree of deacetylation, and the functional properties of chitosan (Venugopal, 2009). Biological processes give products of better
quality, require less energy, and consume less fresh water, unlike chemical processes (Arbia et al., 2013; Kaur and Dhillon, 2015; Mao et al., 2017; Lopes et al., 2018).

Microbial Extractions of Chitin
Fermentation is beneficial for extraction of chitin from seafood processing discards (Yadav et al., 2019). Fermentation by lactic acid bacteria (LAB) has advantages over conventional methods for chitin extraction. One beneficial LAB is Lactobacillus plantarum. Other LABs include L. paracasei, L. acidophilus, L. lactis, L. paracasei, S. marcescens, and T. turnirae. Non-LAB organisms can also be used for fermentation (Vázquez et al., 2013, 2019). An epiphytic L. acidophilus is isolated from rapidly fermented shrimp waste. The chitin released in the fermented product can be easily transformed by a bleaching treatment. The product had better quality than chemically extracted chitin (Duan et al., 2012). Fermentation of shrimp head by a consortium of LAB for 48 h gave chitin and protein rich liquor; the latter can be used as aquafeed supplement (Ximenes et al., 2019). Jung et al. (2007) employed L. acidophilus for chitin extraction from crab shell waste by two-step fermentation, involving L. paracasei in the first step, followed by a protease producing bacterium Serratia marcescens. The process removed 94% CaCO₃ and 70% proteins. The highest deproteinization of 96% and demineralization (68%) were achieved through the combination of two-stage solid state culture by Lactobacillus brevis and Rhizopus oligosporus. Lactic acid was the main organic acid produced along with acetic, succinic, and oxalic acids. The purified chitin presented a molecular weight of 1,313 × 10³ Da, preserving a high crystalline index and acetylation of 94% (Aranday-Garcia et al., 2017). Sieber et al. (2018) suggested the use of natural microbial isolates as well as Serratia spp. and Lactobacillus spp. in fermentations that can realize a demineralization of 97%. Younes et al. (2016) used fermentation to extract highly acetylated chitin from crustacean shells, which were initially subjected to demineralization and enzymatic deproteinization prior to the treatment.

Fermentation by LAB for a maximum period of 7 days resulted in extensive deproteination and demineralization of crustacean shells, facilitating chitin recovery. The process can be conducted under conditions such as anaerobic, solid-stat, semi-continuous, or co-fermentation (Vázquez et al., 2020). Ghorbel-Bellaaj et al. (2011) optimized fermentation variables in accordance with Plackett–Burman design, which resulted in 96% demineralization and removal of 89% protein. Fermentation of shrimp head by Bacillus licheniformis released appreciable amounts of polysaccharides and other compounds in the fermented medium (Mao et al., 2017). Bacillus cereus and Exiguobacterium acetylicum accomplished 90% demineralization and deproteination during chitin extraction from shrimp waste (Sorokulova et al., 2009). Autolysis of shrimp heads could recover not only chitin and chitosan, but also protein hydrolyzate, carotenoids, sulfated-, and amino-polysaccharides. An amount of 25 mg of chitin and 17 mg chitosan (60–80% deacetylated) per g of wet shells were recovered (Cahu et al., 2012). A pilot plant study by Vázquez et al. (2017) employed a combination of enzymatic, acid, and alkaline processes for the recovery of chitin and also protein and carotenoprotein from the cephalothorax of Penaeus vannamei. The sequential treatment yielded 30% chitin with 92% acetylation. In another combination process, protease was used to remove Ca²⁺ and protein, followed by fermentation by Bacillus coagulans to extract chitin from crayfish shell waste, which resulted in recovery of chitin as high as 94% (Dun et al., 2019). Most of these studies reported deproteination and demineralization in the range of 45–90%.

Enzymatic Extractions of Chitin
Extraction of chitin from shellfish waste is facilitated by initial deproteinization and demineralization. One of the biological alternatives proposed is the use of proteases for deproteinization of crustacean shells, avoiding alkaline treatments. When using enzymatic deproteinization, previous demineralization is more convenient since it increases enzyme permeability of the tissues and reduces the presence of potential enzyme inhibitors. Crude proteases from fish discards such as viscera can be used for deproteination, which can lower the cost of treatment (Shah et al., 2016). Instead of the conventional hydrochloric acid, the use of organic acids (lactic and acetic) produced by cheese whey fermentation has been suggested for demineralization of shrimp shells. Organic acids were comparable to hydrochloric acid but less harmful when helping to maintain the integrity of chitin (Mahmoud et al., 2007). Proteolytic enzymes remove up to 90% of the protein and carotenoids from demineralized shrimp waste. Hamdi et al. (2017) extracted chitin from blue crab and shrimp shells by crude crab viscera alkaline protease digestion at pH 8.0 and 60°C. The treatment caused up to 91% deproteinization, facilitating the release of chitin.

In recent times there has been increased interest in using broad-specificity chitinases for chitin extraction. These novel enzymes possess two or three different catalytic activities, functioning as exochitinase, endochitinase, and N-acetyl-glucosaminidase. Endochitinases cleave chitin at internal sites, thereby generating low-molecular weight chitin oligosaccharides (COG), while exochitinases or chitobiases catalyze the progressive release of chitin dimmers by cleaving the polysaccharide at external sites (Suresh, 2012). Recently, a cold-adapted chitinase from a marine bacterium was characterized by broad pH stability, high thermo-stability, low Km value, and optimal activity at 30 °C, with 35% activity at 0°C. The enzyme completely degraded colloidal chitin into N-acetyl glucosamine (GlcNAc). The enzyme was suggested to be a superior candidate for producing bioactive oligosaccharides (Fu et al., 2020). A mass production of chitonolytic enzymes by cultures of micro-organisms, such as Trichoderma hamatum, T.viride, Aspergillus niger, and Carica papaya, will be beneficial for large scale extraction of chitin and its transformation into valuable commercial products as a solution to waste management (Yadav et al., 2019). Chitin-degrading enzymes from Serratia marcescens, Amanantichitinus ursilacus, and Andreporevia ripae have been used on a pilot scale to degrade chitin into monomers with yields up to 95% (Sieber et al., 2018).

Chitin deacetylation from fungi such as such as Mucor rouxii, M. mechei, and Aspergillus niger catalyzes the hydrolysis of N-acetylamido linkage of chitin to give chitosan. The crystallized chitin, after pretreatment with 18% formic acid, is amenable to
90% deacetylation by the fungal deacetylase (Suresh, 2012). To enhance the accessibility of chitin deacetylase to acetyl groups of natural crystalline chitin, pretreatment may be needed with physical or chemical methods such as sonication, grinding, heating, and derivatization (Yadav et al., 2019). The various applications of chitin and chitosan are not topics of discussion here.

Crude enzyme from Bacillus cereus was used to hydrolyze chitosan having 66% deacetylation in a membrane reactor, operated at 45 °C and pH 5. The major oligomers were chitobiose, chitotriose, chitotetraose, chitopentaose, and chitohexaose. The system could be operated for 15 h and still maintained a stable product composition (Kuo et al., 2004). Chito-oligomers (COS), the depolymerized products of chitosan, have attracted considerable interest due to their biocompatible, biodegradable, non-toxic, and non-allergenic natures, and potential applications in biomedical, food, pharmaceutical, agricultural, and environmental industries (Ngo et al., 2020). These suggest a need for commercial production of chinases and chitosanases (Suresh, 2012; Zhou et al., 2019). Chitosan and its derivatives have been reported to possess various biomedical activities including free radical scavenging, antihypertensive, anticoagulant, anti-diabetic, anti-inflammatory, antimicrobial, anticancer, and anti-Alzheimer effects. The antibacterial and antifungal properties of chitosan qualify it for use in food packaging films. Its mechanical, gas, and water vapor permeability properties can be enhanced by blending chitosan with other natural polymers such as starch, essential oils, and clay (Venugopal, 2011). Figure 3 depicts the biological process for extraction of chitin and its conversion to chitosan.

Glycosaminoglycans (GAGs) are hetero-polysaccharides defined by a repeating disaccharide unit without branched chains, in which one of the two monosaccharides is an amino sugar (N-acetyl-galactosamine or N-acetyl-gluconosamine) and the other one is a uronic acid. Based on the disaccharide composition, linkage type, and presence of sulfate groups, GAGs may be chondroitin sulfate (CS), hyaluronic acid (HA), dermatan sulfate, heparin, or keratan sulfate. CS chains have an important function in central nervous system development, wound repair, infection, growth factor signaling, morphogenesis and cell division, differentiation, and migration in addition to osteoarthritis and their conventional structural roles. CS from terrestrial and marine sources contains diverse chain lengths and sulfation. Shark cartilage may contain up to 29% CS, having a molecular weight of 40 kDa. Hyaluronic acid is a linear, high molecular weight linear, and non-sulfated GAG made by alternating disaccharide units of N-acetyl-D-glucosamine and D-glucuronic acids, linked by β-(1→3) and β-(1→4) glycosidic bonds. Autolysis of shrimp head waste gave about 8 mg sulfated GAGs per g that exhibited electrophoretic migration similar to mammalian standards. The degradation products of the GAGs suggested the presence of C6-sulfated heparan sulfate (Cahu et al., 2012). Vázquez et al. (2013) reviewed environmentally friendly processes combining microbial, enzymatic, and other strategies to produce CS, HA, chitin, and chitosan. Bacterial production of HA using Streptococci has been industrially developed (Vázquez et al., 2013). Chondroitin sulfate and hyaluronic acid are commercially valuable. The structural similarity of microbial capsular polysaccharides to these biomolecules makes bacteria ideal candidates as non-animal sources of glycosaminoglycan-derived product GAGs because of their high bioactivities and physiological functions. Fish cartilage products, such as shark cartilage and chondroitin sulfate, glucosamine, and other glycosaminoglycans, are able to alleviate rheumatoid arthritis (Venugopal, 2009).

**Mineral-Based Components**

Finfish discards, which contain significant amounts of bone, are rich sources of minerals. The bone is composed of up to 70% minerals, followed by collagen, certain carbohydrates, and lipids. Hydroxyapatite and calcium phosphate have attracted attention for biomedical applications such as implant materials. Grass fish bones were subjected to flavouryme treatment followed by fermentation with Leuconostoc mesenteroides, giving a preparation with a high content of soluble calcium lactate, calcium acetate, and also small peptides containing calcium. The calcium is bioavailable and therefore can promote growth, as shown by animal studies, suggesting its use as a calcium supplement (Tang et al., 2018). Salmon frames were hydrolyzed by a mixture of commercial proteases. After the procedure, the frames were separated by centrifugation into a bone fraction, which contained 62% of total ash present in the salmon frames. The fraction was high in calcium, phosphorus, and magnesium and also in various trace elements such as copper, iron, selenium, and zinc (Liaset et al., 2003). A fish bone phosphopeptide (FBP) containing up to 24% of phosphorus has a molecular weight of 3.5 kDa and a high calcium-binding activity. The FBP has potential nutraceutical value as a calcium binding agent (Jung et al., 2005). A combination of micro and nano-structured structures, developed with the help of microbial and enzymatic processes is aimed to develop bone products having a high content of calcium.
TABLE 4 | Bioconversions of seafood processing discards by microorganisms.

| Seafood discards       | Microorganism                  | Products                                      | References          |
|------------------------|--------------------------------|-----------------------------------------------|---------------------|
| Shell waste            | Serratia marcescens            | Chitinases, proteases, prodigiosin pigment     | Wang et al., 2020   |
| Shrimp head            | Bacillus licheniformis         | Antioxidant                                   | Mao et al., 2017    |
| Shrimp waste           | Lactobacillus acidophilus       | Chitin, protein hydrolyzate                   | Duan et al., 2012   |
| Shell waste            | Symbiotic lactic acid bacteria  | Calcium, carotenoids peptides                 | Shan et al., 2011   |
| Shrimp head            | Lactic acid fermentation        | Chitin and aquafeed                           | Ximenes et al., 2019|
| Shrimp waste           | Pedicococcus acidilactici       | Carotenoids                                   | Bhaskar et al., 2007|
| Crab shell             | L. paracasei, and S. marcescens | Chitin                                        | Jung et al., 2007   |
| Fish waste             | Mixed microorganisms            | Liquid fertilizer                              | Dao and Kim, 2011   |
| Shrimp shell waste     | Pseudomonas aeruginosa          | Chitin                                        | Ghorbeli-Belaaq et al, 2011|
| Shell waste            | Fermentation                    | Astaxanthin                                   | Routray et al., 2019|
| Shrimp waste           | Bacterial fermentation          | Protein and chitin                            | Sorokulova et al., 2009|
| Shell waste            | P. monocerium                  | a-Chitin                                      | Suresh, 2012        |
| Crayfish waste         | Fermentation with protease      | Chitin                                        | Dun et al., 2019    |
| Seafood effluents      | Microalgae                      | SCP                                           | Shahid et al., 2020 |
| Mussel processing effluents | S. zoeepidemicus fermentation | Hyaluronic acid                               | Vázquez et al., 2013|
| Aquaculture solid waste | Aerobic microbial bioconversion | Liquid fertilizer                              | Khiri et al., 2019  |
| Squid pens waste       | Paenibacillus sp.               | Chitosanas, Chitosan oligosaccharides         | Doan et al., 2020   |
| Tuna waste             | L. plantarum; B. licheniformis | Aquafeed                                      | Hena et al., 2009   |
| Tuna waste             | Staphylococcus epidermidis      | Lipase                                        | Esakikraj et al., 2010|
| Cod by-products        | Proteolysis                     | Oil                                           | Durnay et al., 2004 |
| Fresh water viscera    | Pediococcus acidilactici        | Calcium supplement                            | Rai et al., 2010    |
| Grass fish bone        | Proteolysis and L. mesenteroides fermentation | Glutamic acid, GABA | Sanchart et al., 2018 |
| Tuna condensate        | Candida rugosa and Lactobacillus futsai | Peptones                                      | Vázquez et al., 2020|
| Fish discards          | Various lactic acid bacteria    | Chitin                                        | Aranday-Garcia et al., 2017|
| Shrimp waste           | Successive Lactobacillus brevis and Rhizopus oligosporus | Reducing sugar, DNA protective compounds antioxidants | Rashid et al., 2018 |
| Shrimp shell powder    | B. cereus                       |                                               |                     |

hydroxyapatite (HAp) was isolated from tuna bone. The isolated HAp had comparable physicochemical characteristic with that of standard HAp and was also less toxic (Pallela et al., 2011). Pepsin hydrolyzate of channel catfish bones has antibacterial activity, suggesting that the fish bones are promising resources for generating antibacterial components (Ren et al., 2012).

The various bioconversion processes employing microorganisms and enzymes for the recovery of components from seafood processing discards and effluents are summarized in Tables 4, 5. Table 4 summarizes bioconversion processes using LAB and other microorganisms, while Table 5 summarizes enzymatic bioconversion processes.

THE BIOREFINERY CONCEPT

Bio-Refinery Approach for Valorization of Seafood Discards

The above discussions pointed out various biological processes for the extractions of important compounds from seafood discards. While individual processes may not be economically feasible, an integrated refinery-type process for the extractions of multiple products is more practical. A “refinery” generally means conversion of raw materials into products of higher values, petroleum oil refinery being the most popular example. The International Energy Agency defined bio-refinery as the “sustainable processing of biomass into a spectrum of bio-based products (food, feed, chemicals, and materials) and bio-energy (de Farias and Barbera, 2018). The bio-refinery concept visualizes bio-waste as a potential renewable feedstock that can be valorized through a cascade of various biotechnological processes to produce marketable products and bioenergy on par with petrochemical refineries. It involves stepwise refining processes using biological methods for the extraction of various high value biomolecules. Such downstream strategies could reduce overall production costs (Das, 2015; Mohan et al., 2016; Mitra and Mishra, 2019; Dineshkukar and Sen, 2020). Bio-refineries, which aim at valorizing biomass from agriculture and aquaculture, into a wide spectrum of products and bio-energy, have been recognized as part of a sustainable economy (Dragone et al., 2020).

Fish processing waste can be a promising renewable biomass for bio-refineries. The bio-refinery approach envisages conversion of fish waste into value-added products such as
TABLE 5 | Bioconversions of seafood processing discards by enzymes.

| Seafood discards           | Enzyme                        | Components                                      | References                                                                 |
|----------------------------|-------------------------------|-------------------------------------------------|---------------------------------------------------------------------------|
| Fin, head, scales          | Collagenase, trypsin          | Collagen                                        | Pal and Suresh, 2016; Shah et al., 2016                                   |
| Fish waste                 | Proteases                     | FPH                                             | Vijaykumar and Prabhasankar, 2015                                          |
| Salmon frame               | Proteases                     | Oil, peptides                                   | Liss et al., 2003                                                          |
| Salmon heads               | Proteolysis, lipase           | PUFA-rich oil                                   | Linder et al., 2005                                                        |
| Fish oil                   | C. rugosa lipase              | PUFA-rich oil                                   | Moore and McNeill, 1996                                                   |
| Tuna head                  | Alcalase                      | Deodorized oil                                  | de Oliveira et al., 2016                                                   |
| Chitin waste               | Chitinase                     | Oligosaccharides                                | Fu et al., 2020                                                            |
| Lobster waste, shrimp waste| Papain, alcalase, pancreatin  | Astaxanthin                                     | Routray et al., 2019                                                       |
| Shrimp shell               | Enzymatic, acid, alkaline     | Protein, chitin, carotenoprotein                | Vázquez et al., 2017                                                       |
| Marine waste               | Enzyme and microorganism      | astaxanthin                                     | Vázquez et al., 2013                                                       |
| Crude chitin               | Chitinase                     | N-acetyl-D-glucosamine                          | Zhang et al., 2016                                                         |
| Crude chitin               | Chitinase                     | Chitin oligomers                                | Suresh, 2012                                                               |
| Crab shell                 | Crab viscera protease         | Chitin                                          | Hamdi et al., 2017                                                         |
| Shrimp heads               | Brevibacillus alkaline protease| Chitin                                          | Doan et al., 2019                                                          |
| Shrimp, crab shell, squid pen| P. aeruginosa protease       | Protein and chitin                             | Wang et al., 2019                                                          |
| Shrimp waste               | Bacillus cereus protease      | Chitin, chitosan, FPH                           | Manni et al., 2010                                                         |
| Black tiger shrimp shells  | Bluefish trypsin              | Carotenoprotein, lipid                         | Klonikado et al., 2009                                                     |
| Fish bone                  | intestine crude proteinase    | Bone oligo-phospho-peptide                      | Jung et al., 2005                                                          |
| Shrimp shell               | Bacterial protease            | Astaxanthin                                     | Sachindra and Mahendrakar, 2011                                            |
| Shrimp shell               | Alcalase                      | Protein, chitin, astaxanthin                    | Holanda and Netto, 2006                                                    |

Biofuels, industrial chemicals, animal feed, organic fertilizer, nutraceuticals, and others. Low cost and simplicity of operation by reducing the cost of material, energy consumption, and labor, but maintaining high productivity are some of the important attributes of the process (Sahu et al., 2016). An example is shell refinery, where crustacean shell waste is subjected to sequential treatment to recover chitin, proteins, lipids, carotenoids, calcium carbonate, and chitin monomers (Hülsey, 2018). Vázquez et al. (2019) coupled alcalase hydrolysis with bacterial fermentation to extract gelatin, oils, fish protein hydrolysate including bioactive peptides, and fish peptides from heads, skin, and bones of fish discards. Cahu et al. (2012) reported an integrated process employing autolysis of shrimp heads to recover chitin and chitosan, protein hydrolyzate, and chitin oligomers (Vázquez et al., 2017; Routray et al., 2019). Similarly, anaerobic fermentation of fish waste resulted in methane and liquid fertilizer as primary products. The purchase price of methane is a crucial factor influencing the economics of the bio-refinery (Ratky and Zamazal, 2020). Another bio-refinery deals with extraction of oil from fish waste, its transesterification with ethanol, and concentration of n-3 PUFA. Fishmeal, glycerol, and saturated and short chain unsaturated fatty acids as liquid bio-fuel are the other products of the refinery. The process can significantly supply thermal energy and reduce CO2 discharge (Fiori et al., 2017). A bio-refinery developed within an EU-funded project combines chitin demineralization by Serratia spp. and Lactobacillus spp. and an enzymatic degradation of chitin by chitin-degrading enzymes from Serratia marcescens, Amanitchitinus ursilacus, and Andreprevotia ripae. The resulting N-acetylglucosamine monomers could be used for novel bio-based polymers. Proteins and lipids could be used as feed for biogas production (Sieber et al., 2018). Eurofish processes roughly 200 tons of tuna a day with discharge of at least 1,300 m3 effluents. The company coupled seafood waste-to-energy technology, generating 1,300 m3 methane daily. This reduced wastewater treatment costs by 50% and energy consumption by 35–40%. The plant has been in operation since March 2016, suggesting economic feasibility of bio-refiners based on seafood waste valorization (Fluence, 2019).

Algal biotechnology can be a promising platform for bio-refining of seafood discards. The cultivation of microalgae in bio-wastes is known to produce SCP, as mentioned earlier. The recovery of products from algal biomass is a matter of constant development and progress (Sosa-Hernández et al., 2018). Whereas, the exploitation of SCP for a single product such as biofuel is not economically viable, multiple products such as pigments, antioxidants, and n-3 fatty acids can be extracted from SCP to make the process cost-competitive. The various possibilities are depicted in Figure 4. The approach offers novel ways to utilize wastewater and also help in the promotion of microalgae in the commercial market (Koyande et al., 2019; Mitra and Mishra, 2019). The increasing resource limitations are expected to drive SCP production and improve the economic feasibility in the future (Puyol et al., 2017). On the basis of increasing demands, recently seaweed has been cultivated with improved traits to harvest more than one...
product through a bio-refinery. Three combination routes have been suggested for production of microalgal-based biodiesel, bio-hydrogen, and SCP (Banu et al., 2020). Cultivation of the microalga \textit{H. pluvialis} for both SCP and astaxanthin can be an economically sustainable process (Shah et al., 2016; Khoo et al., 2019). Food producers must now address environmental concerns, social responsibility and economic viability when designing their food processing techniques. Systematic improvement of the technology readiness level (TRL) could be successful if applied to microalga cultivation and processing (Caporgno and Mathys, 2018). Recent text mining tools on articles and patents published on algal biotechnology during the period 2012–2017 identified \textit{Reinhardtius} sp. for wastewater treatment and a \textit{Chlorella} strain for biofuel and fatty acids (Parkavi et al., 2020). At present, downstream processing, and in particular the fractionation of microalgal components, remains the most expensive step of the algal processes, demanding novel technologies for SCP processing (Gifuni et al., 2019). Cho and Park (2018) observed that commercialization of the microbial route for fuel production remains uncertain due to the high cost of feedstock or low lipid yield. However, considering the low cost of seafood discards and effluents and potentials for enhancing fuel production through algal technology, as mentioned earlier, this observation may not be realistic. Table 6 summarizes recent bio-refinery approaches for seafood valorization.

### CONCLUSIONS

The article pointed out the major problems associated with discards and process effluents, generated during industrial seafood processing. Besides being responsible for environmental pollution, the discards and effluents represent heavy losses of nutrients and other valuable compounds. There problems can be addressed by biological treatment processes, involving bioconversions of components of the waste by microorganisms and enzymes. Unlike conventional chemical treatments, biological processes are environmentally friendly, safe and economical. Further, biological processes do not adversely affect functional properties of isolated compounds, unlike the chemical processes. Biomass fermentations using microalgae are emerging as green and economical processes to recover functionally active compounds and also biofuel. Fermentation is ideal to recover functionally active chitin from crustacean shell. Similarly, fermentation or lipase-based processes can replace hazardous solvent extraction techniques for the recovery of fish oil. Microorganisms-mediated processes are highly desirable due to ease of handling, lower energy requirements and costs. With the development of tailor made bio-catalysts and advances in

| Bio-refinery | Products | References |
|--------------|----------|------------|
| Cultivation of alga, \textit{Haematococcus pluvialis} | Astaxanthin, SCP | Shah et al., 2016; Khoo et al., 2019 |
| Lactic fermentation | Astaxanthin, hydrolyzed protein and chitin | Routray et al., 2019 |
| Sequential treatment of crustacean shells | Chitin, proteins, lipids, carotenoids and CaCO3. | Hüseyin, 2018 |
| Demineralization and enzymatic degradation of N-acetylglucosamine | Chitin monomers | Sieber et al., 2018 |
| Sequential enzymatic, acid–alkaline extraction of shrimp cephalothorax | Chitin, chitosan, protein and astaxanthin | Vázquez et al., 2017 |
| Integrated autolysis of shrimp head | Chitin, protein hydrolyzate, sulfated glycosaminoglycans | Cahu et al., 2012 |
| Sequential extraction by ISP followed by enzyme | Collagen, myofibrillar proteins | Abdollahi et al., 2018 |
| Sequential treatment of marine cartilage | Chondroitin sulfate, fish meal | Vázquez et al., 2013 |
| Oil extraction, ethanol transesterification, n-3 PUFA concentration | Fish proteins, Glycerol, Liquid biofuel | Fiori et al., 2017 |
| Hydrolysis of fish waste by proteolytic enzymes | Food, Feed, Fertilizer ingredients | Sahu et al., 2016 |
| Anaerobic fermentation with cow dung | Methane, Liquid mineral fertilizer | Ratko and Zaman, 2020 |
| Algal bio-refinery | Various products | Mitra and Mishra, 2019 |
| Coupled alcalase hydrolysis and bacterial fermentation | Gelatin, oils, FPH, bioactive peptides, and fish peptones | Vázquez et al., 2019 |
green extraction techniques, it is possible to take up challenges of successful bio-processing of seafood discards and effluents. Genetic engineering of microorganisms, enzyme engineering, reactor designs and process optimizations offer strategies leading to a new manufacturing paradigm for successful valorization of seafood waste. The interesting features of microalgae such as their rapid growth, their photosynthetic ability, nutrient-rich sea- side waste. The interesting features of microalgae such as their rapid growth, their photosynthetic ability, nutrient-rich sea- products, and their potential use in food and feed industries. Fish protein hydrolysates: proximate composition, amino acid composition, antioxidant activities and applications: a review. Food Chem. 135, 3020–3038. doi: 10.1016/j.foodchem.2012.06.100

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Although process identification, scaling up and implementation of resource recovery solutions at an industrial level is a challenge, developments in these fields can favor successful waste utilization for sustainable fish processing towards a circular bioeconomy.

AUTHOR CONTRIBUTIONS

The author is the sole contributor of this work and has approved it for publication.
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Conflicts of Interest: The author declares that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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