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CRITICAL REVIEW

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Accessing biominerals from by-products wasted by the seafood processing industry

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Due to the global population's significant growth, the demand for seafood has increased exponentially over the last century. This has led to the generation of large amounts of inedible by-products from the seafood processing industry from a multitude of marine organisms, including finfish, mollusks, and crustaceans. In this review, the potential extraction and application of biominerals from waste bones and shells originating from the seafood processing industry are discussed. Existing reviews on fishery by-products often highlight only the potential of organic by-products and disregard the biominerals present. Shells always contain calcium carbonate, CaCO_3 , in the form of either calcite or aragonite and bones are primarily made of hydroxyapatite, $\text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2$. The conditions used in different extraction processes to isolate the mentioned biominerals and the resulting products are compared. Furthermore, we highlight the sustainability impact of sourcing natural biominerals for global applications (e.g., limestone industry) as this would prevent further environmental risks caused by improper disposal. Bio-derived minerals have also been used in other applications, such as environmental remediation and biomedicine. For calcium carbonate, the use of raw shells, their transformation to calcium oxide, CaO , and the production of precipitated calcium carbonate are presented. Next, we present in detail the isolation of hydroxyapatite from fish bones and scales, its transformation to other biphasic calcium phosphates, and its applications. The review concludes with a discussion on the strengths, weaknesses, opportunities, and threats to accessing biominerals wasted by the seafood processing industry.

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Sustainability spotlight

Seafood is a global requirement as an important source of protein as the human population grows significantly (UN SDG2). Currently, the seafood processing industry relies on landfill and/or ocean disposal to manage the generated by-products, but this had led to greenhouse gas emissions (UN SDG13), groundwater contamination (UN SDG6 and UN SDG15), eutrophication, and acidification (UN SDG14). The biominerals present offer the potential for the seafood processing industry to create other higher-value products for consumers, a step towards achieving a circular economy (UN SDG8, UN SDG9, and UN SDG12). Sustainable processes exist in the literature to isolate biominerals from bones and shells of aquatic organisms (UN SDG11) which have then been used in multiple applications, including biomedicine (UN SDG3), environmental remediation, and catalysis.

1. Introduction

Seafood is a necessary source of protein globally and its production has increased significantly in the last century to meet the demands of the growing human population.^{1,2} For example, in 1961 a person would consume on average 8.96 kg of seafood,³ but that has increased to 20.25 kg in 2020.³ Therefore it is not surprising that global seafood production has increased by nearly 11 000% from 1961 (1.96 million tonnes) to 2020 (214 million tonnes).⁴ The countries that consume the most fish are those that do not necessarily have access to food from land-based livestock. The Maldives consumes the greatest amount with 87.30 kg per person per year,³ followed by Iceland, Macao,

and Kiribati.³ Despite not being the largest consumers of fish, developing countries in Africa and Asia have experienced the greatest increase in seafood consumption. For example, Niger has seen the largest overall increase among all countries with an increase of over 3000% in seafood consumption while Rwanda, Nepal, and Iran have also seen significant increases of ~2400–2600%.³

Technological advances have contributed to the increased production of seafood. Historically, the only method of supplying fish and shellfish was by harvesting seafood directly from the ocean, from the place in which it was naturally grown, otherwise known as “capture fisheries”.^{3,5} However, since the 1980s, the global production from industrial capture fisheries has remained relatively stable at 90–95 million tonnes annually,⁵ and is limited by the ecosystems and legislation to control over-fishing. In many regions, the quantity of fish caught by industrial vessels has been declining due to geopolitical and/or

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commercial dynamics. Furthermore enormous progress has been achieved in aquaculture over recent decades.^{6,7} According to the Food and Agriculture Organization (FAO) of the United Nations (UN), aquaculture is defined as “the farming of aquatic organisms including fish, mollusks, crustaceans, and aquatic plants”.^{6,8} It was developed as a method to continue producing seafood products while alleviating some pressure on capture fisheries. In fact, aquaculture produced 123 million tonnes of seafood in 2020, surpassing quantities harvested *via* capture fisheries for the first time.⁴ Finfish species represent the majority of aquaculture production and account for 46.9% of processing, followed by algae (28.6%), mollusks (14.5%), crustaceans (9.2%), and other aquatic animals (0.9%).⁹ Aquatic organisms that are considered as the major aquaculture species by the FAO are listed in Table 1.

The seafood processing industries are critical for global food security, but face sustainability challenges. Industrial capture fisheries rely on fishing vessels that emit significant quantities of CO₂ (ref. 10) and can result in species depletion through overfishing. The industrial capture fish industry also causes food loss, waste, and ocean pollution during harvesting because management often encourages disposal of fish at sea (especially bycatch *i.e.* capture of unintended species),⁵ and fishing gear can be abandoned, lost, or discarded.⁵ Abandoned fishing gear has numerous environmental consequences including the introduction of microplastics and toxins into feed webs, habitat degradation, and the spread of invasive alien species.¹¹ While aquaculture was developed to overcome some of these challenges, it has also led to unintended problems that have been amplified with significant growth in this sector.† A lot of seafood is wasted due to poor harvesting practices that contribute to diseases that can cause entire pens of fish to die.^{5,12} Some of the fish can potentially escape the pens and interact with local species, altering the environment and also affecting the genetics of wild fish that are protected as species at risk. If some of the escaped fish are sick, they can spread disease,³ further affecting the local environment. Additionally, some companies have taken advantage of aquaculture and have attempted to breed too many fish in one area, thus creating anoxic zones and affecting local wildlife. Aquaculture has the potential to contribute to many of the UN's Sustainable Development Goals if care is taken to ensure economic, environmental, and social viability.⁷

Another source of waste that has yet to be mentioned is the processing of aquatic organisms post-harvest.¹³ In general, processing involves chemical and mechanical operations required to transform and also preserve seafood. The first step typically involves mechanically (frequently manually) removing the entrails (*e.g.*, gutting). Most of the time, processing takes place in a factory environment which can often be labor intensive and/or highly automated. The main objective of fish processing plants is to provide fish for human consumption and the by-products (frames, skins, viscera) are sometimes processed into secondary low-profit products such as fertilizer

and less frequently into higher value nutritional supplements. Processing plants situated in remote areas face challenges regarding management of by-products because alternative uses are not economically viable nor accessible. While landfills remain the most common method of disposal, remote areas are sometimes issued permits for disposal at sea when waste cannot be feasibly sent for reprocessing or to landfills.¹⁴ In emerging economic regions of the world and in places based on subsistence/traditional fisheries, disposal of waste at sea is most common.

Relying on landfills and oceans for the disposal of seafood discards has environmental consequences. According to the FAO, food loss and waste account for 8–10% of global greenhouse gas (GHG) emissions.¹⁵ Landfills have a significant global warming potential (GWP) because the organic content is decomposed by bacteria through aerobic and anaerobic digestion, releasing carbon dioxide (CO₂), methane (CH₄), sulfur dioxide (SO₂), and nitrogen oxides (NO_x).¹⁶ Discards, often containing P and N, are dumped in the ocean causing eutrophication due to the abundance of nutrients present.^{17,18} Eutrophication creates algae blooms that are potentially toxic, disrupt the ecosystem, taint drinking water, and eventually result in anoxic zones.^{17–19} Furthermore, once these bacteria die, they release more CO₂ and contribute to ocean acidification.²⁰ The ocean's decrease in pH will eventually be crucial for any capture fishery or aquaculture practice at sea because it slows the growth of fish and shellfish, thus reducing catch and harvest size.²⁰ Due to their contribution to global warming, seafood discards should be valorized rather than disposed of and the fishing industry should strive towards a more circular economy.²¹ For example, Iceland has been making significant strides in utilizing every aspect of fish towards creating higher-value products. In 2021, 80% of all Icelandic raw materials from fish were valorized as either food or other consumer products.²² Within Iceland, the 100% Fish Project seeks a wide range of products that could be created using every part of the fish (Fig. 1).^{22–24} Other than the edible seafood products, items such as cosmetics, medicinal products, and even leather have been prepared from wasted by-products that would have historically been unsustainably wasted.^{22–24}

Fish and shellfish waste offer the potential to extract many high-value compounds. Typically, researchers focus on extracting organic compounds and biopolymers from seafood waste. Fish waste has been processed into various products including collagen, non-collagenous proteins, omega-3 fatty acids, and biogas. Marine collagen is of particular interest among researchers because it is highly abundant not only in fish and sharks, but also in by-catch organisms (*e.g.*, jellyfish and sea stars – previously known as starfish) too. Collagen has been extracted from seafood waste with several processes such as chemical hydrolysis, enzymatic treatment, and fermentation using proteolytic bacteria.²⁵ This biogenic collagen has been used in several biomedical and pharmaceutical applications because it is biocompatible and biodegradable, and it has also been used as a food additive and incorporated into packaging materials.²⁵

Other biologically active compounds, including various fatty acids, have been isolated from fish by-products. For example,

† It is thought that the origins of aquaculture date back to 3000 BC *via* the construction of artificial fish ponds in China.



Table 1 Production of major aquaculture species in 2020, as reported by the FAO⁹

Group	Species	World production (10 ³ t, 2020)	% of group
Finfish	Grass carp (<i>Ctenopharyngodon idellus</i>)	5791.5	10.1%
	Silver carp (<i>Hypophthalmichthys molitrix</i>)	4896.6	8.5%
	Nile tilapia (<i>Oreochromis niloticus</i>)	4514.6	7.9%
	Common carp (<i>Cyprinus carpio</i>)	4236.3	7.4%
	Catla (<i>Catla catla</i>)	3540.3	6.2%
	Bighead carp (<i>Hypophthalmichthys nobilis</i>)	3187.2	5.5%
	<i>Carassius</i> spp.	2748.6	4.8%
	Atlantic salmon (<i>Salmo salar</i>)	2719.6	4.7%
	Striped catfish (<i>Pangasianodon hypophthalmus</i>)	2520.4	4.4%
	Roho labeo (<i>Labeo rohita</i>)	2484.8	4.3%
	Clarias catfish (<i>Clarias</i> spp.)	1249.0	2.2%
	Milkfish (<i>Chanos chanos</i>)	1167.8	2.0%
	Tilapias nei (<i>Oreochromis</i> spp.)	1069.9	1.9%
	Rainbow trout (<i>Oncorhynchus mykiss</i>)	959.6	1.7%
	Wuchang bream (<i>Megalobrama amblycephala</i>)	781.7	1.4%
	Black carp (<i>Mylopharyngodon piceus</i>)	695.5	1.2%
	Largemouth black bass (<i>Micropterus salmoides</i>)	621.3	1.1%
	Mulletts nei, (Mugilidae)	291.2	0.5%
	Gilthead seabream (<i>Sparus aurata</i>)	282.1	0.5%
	Large yellow croaker (<i>Larimichthys croceus</i>)	254.1	0.4%
	European seabass (<i>Dicentrarchus labrax</i>)	243.9	0.4%
	Groupers nei (<i>Ephinephelus</i> spp.)	226.2	0.4%
	Coho (silver) salmon (<i>Oncorhynchus kisutch</i>)	221.8	0.4%
	Japanese seabass (<i>Lateolabrax japonicus</i>)	196.9	0.3%
	Pompano (<i>Trachinotus ovatus</i>)	160.0	0.3%
	Japanese amberjack (<i>Seriola quinqueradiata</i>)	137.1	0.2%
	Barramundi (Giant seaperch, <i>Lates calcarifer</i>)	105.8	0.2%
	Red drum (<i>Scianops ocellatus</i>)	84.3	0.1%
	Subtotal of major species	45 388.1	79.0%
	Subtotal of other species	12 072.8	21.0%
Total	57 461.1	100.0%	
Algae	Japanese kelp (<i>Laminaria japonica</i>)	12 469.8	35.5%
	Eucheuma seaweed (<i>Eucheuma</i> spp.)	8129.4	23.2%
	Gracilaria seaweed (<i>Gracilaria</i> spp.)	5180.4	14.8%
	Wakame (<i>Undaria pinnatifida</i>)	2810.6	8.0%
	Nori (<i>Porphyra</i> spp.)	2220.2	6.3%
	Elkhorn sea moss (<i>Kappaphycus alvarezii</i>)	1604.1	4.6%
	Fusiform sargassum (<i>Sargassum fusiforme</i>)	292.9	0.8%
	Spiny eucheuma (<i>Eucheuma denticulatum</i>)	154.1	0.4%
	Subtotal of major species	32 861.5	93.7%
	Subtotal of other species	2216.0	6.3%
	Total	35 077.6	100%
	Mollusks	Cupped oyster (<i>Crassostrea</i> spp.)	5540.3
Japanese carpet shell (<i>Ruditapes philippinarum</i>)		4266.2	24.0%
Scallops nei (Pectinidae)		1746.4	9.8%
Sea mussel (Mytilidae)		1108.3	6.2%
Constricted tagelus (<i>Sinonovacula constricta</i>)		860.3	4.8%
Pacific cupper oyster (<i>Magallana gigas</i>)		610.3	3.4%
Blood cockle (<i>Anadara granosa</i>)		457.9	2.6%
Chilean mussel (<i>Mytilus chilensis</i>)		399.1	2.2%
Subtotal of major species		14 898.6	84.0%
Subtotal of other species		2843.6	16.0%
Total		17 742.2	100%
Crustaceans	Whiteleg shrimp (<i>Penaeus vannamei</i>)	5812.2	51.7%
	Red swamp crawfish (<i>Procambarus clarkii</i>)	2469.0	22.0%
	Chinese mitten crab (<i>Eriocheir sinensis</i>)	775.9	6.9%
	Giant tiger prawn (<i>Penaeus monodon</i>)	717.1	6.4%
	Giant river prawn (<i>Macrobrachium rosenbergii</i>)	294.0	2.6%
	Indo-Pacific swamp crab (<i>Scylla serrata</i>)	248.8	2.2%
	Oriental river prawn (<i>Macrobrachium nipponense</i>)	228.8	2.0%
	Green mud crab (<i>Scylla paramamosain</i>)	159.4	1.4%
	Subtotal of major species	10 705.3	95.3%
Subtotal of other species	531.8	4.7%	



Table 1 (Contd.)

Group	Species	World production (10 ³ t, 2020)	% of group
	Total	11 237.0	100%
Other	Chinese softshell turtle (<i>Trionyx sinensis</i>)	334.3	31.5%
	Japanese sea cucumber (<i>Apostichopus japonicus</i>)	201.5	19.0%
	Frog (<i>Rana</i> spp.)	147.8	13.9%
	Edible red jellyfish (<i>Rhopilema esculentum</i>)	90.4	8.5%
	River and lake turtle (<i>Testudinata</i>)	49.3	4.6%
	Subtotal of major species	823.3	77.5%
	Subtotal of other species	239.0	22.5%
	Total	1062.3	100%

Dave *et al.* investigated fatty acid yields in fish oil extracted from Atlantic salmon (*Salmo salar*) using enzymes and the extracted compounds included saturated fatty acids, monosaturated fatty acids, polyunsaturated fatty acids, omega-3 fatty acids, and omega-6 fatty acids.²⁶ Other sustainable methods for accessing omega-3 polyunsaturated acids from waste have been reported including extraction with supercritical CO₂.²⁷ This oil can either be used for nutritional supplementation, or it can be used as a starting material that is further reacted to create a product with other value-added applications. This has been achieved by Laprise *et al.* by epoxidizing waste-derived fish oil followed by reaction with CO₂ to yield an oil rich in cyclic carbonate groups, which could then be converted to a non-isocyanate polyurethane material.²⁸ Often, a sludge remains after the oil extraction process and is viewed as a by-product,²⁶ but it also has potential value. Paone *et al.* described their concept of an anchovy (*Engraulis* spp.) biorefinery and in their attempts to create a fully circular process, the remaining sludge after extraction was used for the production of biogas through anaerobic digestion.²⁹

Crustacean shells are also useful materials because they consist of ~40% protein, ~20–25% chitin, ~5–10% lipids, and pigments.³⁰ Chitin, a linear polysaccharide of *N*-acetyl- β -glucosamine, is a particularly valuable compound. While it only has a few applications as a raw material because of its hydrophobicity, it is widely used in its deacetylated form, chitosan.³¹ Because chitosan is more soluble in acidic and aqueous environments, it has been used in water treatment, nutraceuticals, biomedicine, and cosmetics.³² The standard protocol to extract chitin from crustacean shells is energy- and chemically intensive, relying on hydrochloric acid (HCl) and sodium hydroxide (NaOH) for the elimination of minerals and proteins.³² However, Burke and Kerton have recently investigated green extraction methods for the sequential extraction of carotenoid pigments, protein, and chitin from snow crab (*Chionoecetes opilio*) shells.³³ Astaxanthin was extracted with vegetable oil, a protein powder was yielded using citric acid, and chitin was recovered using enzymes.³³ Other methods that do not rely on acids and bases to extract chitin from crustacean shells include ionic liquids,³⁴ deep eutectic solvents (DESs),³⁵ supercritical fluids,³⁶ and mechanochemistry.³⁷

The isolation of biominerals has been less prevalent compared to organic compounds despite being present in large quantities. For example, mussel shells are made of

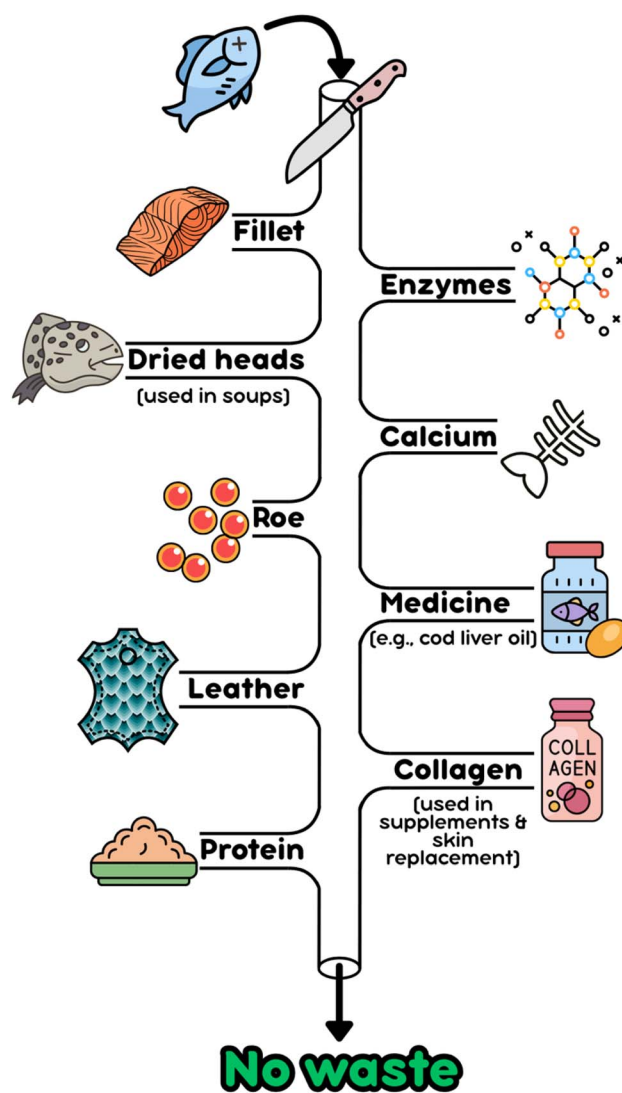


Fig. 1 Schematic representation of different value streams achieved by the Iceland Ocean Cluster's 100% Fish Project. Prepared with Canva.



approximately 95% calcium carbonate (CaCO_3) and fish bones consist of 60% hydroxyapatite (HAP). In this review, the two biominerals most abundant in seafood waste, CaCO_3 and HAP, will be discussed. Marine-derived CaCO_3 is compared to mined limestone and exists as several polymorphs. In Section 2, CaCO_3 polymorphs present in shells are compared between different species of bivalves, crustaceans, and gastropods. We also touch on the transformation of biogenic CaCO_3 to CaO and its further processing to precipitated calcium carbonate (PCC) or HAP. In Section 3, HAP isolated from fish is compared to stoichiometric HAP and its potential benefits for biomedical applications are discussed. Finally, different isolation methods such as calcination and alkaline deproteinization are discussed for obtaining HAP by treating fish backbones and scales. Articles describing processes to isolate CaCO_3 or HAP from waste biomass, but with limited characterization details on the resulting minerals are omitted from this review.

2. Calcium carbonate, CaCO_3

It is estimated that the average American uses 40 000 lbs of new minerals annually.³⁸ Among these minerals, limestone makes up 71% of all crushed stone produced by the United States.³⁹ To be officially considered as limestone, the rock must contain >50% CaCO_3 .⁴⁰ Limestone can be classified as impure, low purity, medium purity, high purity, and very high purity depending on the CaCO_3 content. Rocks with <85% CaCO_3 are considered impure limestone while very high purity limestone contains >98.5% CaCO_3 .⁴⁰ While limestone itself has several applications (e.g., filler in cements, concretes, plastics, paints, rubbers, and chalk),⁴¹ it is often converted to lime, CaO .

Lime is one of the most important materials globally because it is necessary for several sectors, including steel, construction, agriculture, environmental remediation, and chemical industries.⁴² For example, in 2023, 88 million tons of Portland cement were produced in the United States,⁴³ accounting for 95% of hydraulic cement production.⁴⁴ Portland cement is made of 60.2–68.7% CaO ⁴⁵ and therefore its production is very important industrially across the world. The transformation of CaCO_3 to CaO is carried out *via* calcination in kilns at very high temperatures of 700–900 °C.^{46–48}

Other products that can be formed from limestone: CaO , otherwise known as lime (or quicklime), can be prepared through calcination as described above. Hydrated lime (or slaked lime), $\text{Ca}(\text{OH})_2$, is easily prepared from lime by reacting it with water.⁴⁷ PCC is CaCO_3 that has been artificially prepared by reacting CO_2 with a slurry of hydrated lime to produce CaCO_3 purer than limestone.⁴⁹ Synthesizing PCC also ensures that the CaCO_3 product has a definite morphology, size, and structure that may be required for specific applications.⁴⁹ For example, PCC is preferred for pharmaceutical purposes because of its high purity that comes at a relatively high cost.⁴⁰

The limestone and lime industries are major contributors to GHG emissions, especially CO_2 . According to the International Energy Agency (IEA), cement production is responsible for 7% of global anthropogenic CO_2 emissions and is the third-largest industrial energy consumer.⁵⁰ This trend is expected to rise with

the growing global population, infrastructure development, and urbanisation patterns. By 2050, a 12% increase in cement production is expected. To limit the Earth's rising temperature to 2 °C, CO_2 emissions from cement manufacturing must decrease by an estimated 24%.⁵⁰ It is difficult to decarbonize the cement industry because of its inherent nature, although strides towards reducing the CO_2 emissions have been made utilizing clinker replacement, fuel alternatives, and carbon capture technologies.⁴⁶ The conversion of CaCO_3 to CaO will continue to be unsustainable because it is energetically demanding and releases CO_2 from both the fuel and decomposition process. Despite the difficulty in decarbonizing the limestone industry, other researchers have expressed optimism. Rissman *et al.* suggested that accepting and adopting new cement chemistries and alternative materials could help achieve a 50% reduction of industrial emissions by 2050.⁵¹

Another unsustainable aspect of the cement industry is obtaining limestone by mining in open quarries and blasting techniques. These methods are environmentally harmful and hazardous to the employees tasked with mining the mineral.⁵² Issues that arise from mining limestone include groundwater contamination, high energy usage that generally comes from fossil fuels, creation of sinkholes, reduced biodiversity, and dust-related hazards.⁵³ In 2021, Stepkin *et al.* published a population health risk assessment involving communities near an industrial site for ace quarrying. They detected several pollutants in the atmosphere that have concerning toxicological characteristics, including a (1) carcinogenic effect, (2) embryotropic action, (3) gonadotropic action, (4) teratogenic action, and/or (5) a mutagenic effect.⁵⁴ To overcome the environmental and health risks associated with quarrying, CaCO_3 can be prepared through synthetic processes to limit environmental impact, such as precipitation/crystallization from mixing calcium and carbonate salts, but the calcium salts likely originate from the mined/quarried limestone as a feedstock.⁵⁵ A more sustainable way to obtain CaCO_3 is by isolating it from calcifiers: animals that rely on CaCO_3 to form their shells.⁵⁶ After all, most carbonate deposits that are commercially viable were created by the sedimentation of carbonate-secreting organisms many years ago.⁵⁷ This method reduces the amount of waste shells being disposed of by the fish processing industry and could thus reduce CO_2 emissions, acidification, and eutrophication.⁵⁸ Overall, extracting CaCO_3 from waste shellfish could help meet demands more effectively and result in an ocean-based circular economy.^{59–61} Furthermore, bio-derived CaCO_3 is better than limestone-based CaCO_3 for applications that involve human use (e.g., CaCO_3 tablets).⁶² An example of this is Biocoral®, a CaCO_3 -based coral material used as a substitute for bone grafts.⁶³

CaCO_3 from shellfish exists as one or a combination of its three polymorphs: calcite, aragonite, and vaterite.⁶⁴ Calcite particles are found in various morphologies; it is the most thermodynamically stable polymorph and the most abundant polymorph,⁶⁵ and it has a trigonal structure with the most common form being a rhombohedron.^{66,67} Aragonite particles can be needle-like⁶⁶ or plate-like nacre,⁶⁸ and occur in an orthorhombic system.⁶⁷ Vaterite particles are typically



Table 2 CaCO₃ polymorphs present in various species of bivalves, crustaceans, and gastropods

Category	Species	Polymorphs in raw shells	Reference
Bivalves	Blue mussel (<i>Mytilus edulis</i>)	75% calcite & 25% aragonite	Murphy ^{68,79}
	Blue mussel (<i>Mytilus edulis</i>)	68% calcite & 32% aragonite	Cardoso ⁷⁸
	Sururu mussel (<i>Mytella falcata</i>)	94% aragonite & 6% calcite	Cardoso ⁷⁸
	Sururu mussel (<i>Mytella falcata</i>)	Mostly aragonite, little calcite	Araújo ⁸⁰
	Green mussel (<i>Perna viridis</i>)	100% aragonite	Ismail ⁷³
	Mediterranean mussel (<i>Mytilus galloprovincialis</i>)	Calcite & aragonite	Nam ⁸¹
	West Indian pointed venus clam (<i>Anomalocardia brasiliiana</i>)	100% aragonite	Cardoso ⁷⁸
	Spiny fileclam (<i>Lima lima</i>)	Aragonite	Anand ⁸²
	Brown clam (n.s. ^a)	Calcite & aragonite	Aguila-Almanza ⁸³
	Grey clam (n.s. ^a)	Calcite & aragonite	Aguila-Almanza ⁸³
	Ark clam (<i>Scapharca subcrenata</i>)	Aragonite	Nam ⁸¹
	Manila clam (<i>Ruditapes philippinarum</i>)	Mostly aragonite, little calcite	Nam ⁸¹
	Venus clam (<i>Meretrix petechialis</i>)	Mostly aragonite, little calcite	Nam ⁸¹
	True oyster (Ostreidae)	90% calcite & 10% aragonite	Ramakrishna ⁸⁴
	Pacific oyster (<i>Crassostrea gigas</i>)	Calcite & aragonite	Nam ⁸¹
	Oyster (n.s. ^a)	Calcite	Aguila-Almanza ⁸³
	Oyster (n.s. ^a)	Calcite	Nguyen Quang & Ta Hong ⁸⁵
	Yesso scallop (<i>Patinopecten yessoensis</i>)	Calcite & aragonite	Nam ⁸¹
	Pen shell (<i>Atrina pectinate</i>)	Calcite & aragonite	Nam ⁸¹
	Crustaceans	Egg cockle (<i>Fulvia mutica</i>)	Aragonite
Blue crab (<i>Callinectes sapidus</i>)		Calcite	Ogresta ⁸⁶
Mediterranean green crab (<i>Carcinus aestuarii</i>)		Calcite	Ogresta ⁸⁶
Mangrove crab (<i>Ucides cordatus</i>)		Calcite	Cardoso ⁷⁸
Mangrove crab (<i>Ucides cordatus</i>)		Calcite	Araújo ⁸⁰
Gastropods	Flower crab (<i>Portunus pelagicus</i>)	Calcite	Bayuseno ⁷³
	Sea snail (<i>Loittioidea</i>)	Aragonite	Anand ⁸²
	Netted olive sea snail (<i>Oliva reticularis</i>)	Aragonite	Anand ⁸²

^a Abbreviations – n.s., non-specified.

spherical⁶⁶ and belong to the hexagonal crystal system,⁶⁹ but there is ongoing discussion on the space group of vaterite, with several additional structural models being proposed such as monoclinic.⁶⁹ Calcite is the main polymorph of CaCO₃ found in limestone; therefore for specific applications it may be required to separate it from the other polymorphs present. Aragonite and vaterite are metastable polymorphs of CaCO₃ that readily transform to calcite,^{65,66,70} especially vaterite which recrystallizes to calcite when it is dissolved in water.⁶⁴ CaCO₃ from shellfish waste is present in the form of calcite and/or aragonite depending on the species (Table 2), but vaterite can also be synthesized based on shell treatment (Table 5).^{71–73}

Shellfish are made of two main groups of organisms: mollusks and crustaceans. Researchers have investigated waste shells from both groups as potential feedstocks for CaCO₃. The main difference between crustacean and mollusk shells is that while mollusk shells are made of ~95% CaCO₃, crustacean shells are more complex (e.g., crab shells are composed of ~20–50% CaCO₃, Fig. 2).⁷⁴ Therefore, as a raw material, mollusk shells are exploited more than crab shells as a source of CaCO₃. However, some applications require treating the shells with calcination, thus decomposing the protein (and also chitin in crustaceans) in the process. In the next section, we will discuss using raw shells as a source of different CaCO₃ polymorphs and subsequently transforming them to CaO and PCC. It should be noted that seashells often contain Mg, Mn, Zn, Y, Cu, Sr, Ba, and Pb that can significantly influence the performance of materials.⁷⁵

2.1. Raw shells

Raw shells from mollusks are made almost entirely of CaCO₃ in the form of calcite and aragonite. However, to utilize the shells, the organic residues must be removed; otherwise they will eventually decompose and result in a stench that is unwanted by seafood processing industries and surrounding communities. Herein, organic residues refer to all organic-containing material, including collagen. The process of cleaning waste shells is rarely discussed in the literature; however Murphy *et al.* optimized two enzymatic treatments to decompose the organic residues from raw and cooked blue mussel (*Mytilus edulis*) shells by using 1.0–2.0 μL g⁻¹ Multifect PR 6L for 4 h at 55 °C and 6.0 μL g⁻¹ Multifect 7L for 10 h at 25 °C, respectively.⁷⁶ A more recent preparation techniques used to isolate CaCO₃ from shells uses 55% v/v NaClO instead of enzymes to degrade the organic residues; however this process is less sustainable than using enzymes.⁷⁷ To isolate CaCO₃ from natural sources, calcination is typically avoided because it converts CaCO₃ to CaO at 800–900 °C.⁷⁸ Examples of CaCO₃ polymorphs that have been found in shells from bivalves, crustaceans, and gastropods are summarized in Table 2.

2.1.1. Bivalves. The CaCO₃ polymorphs present in raw shells of various bivalves have been investigated. In a 2015 study by Nam *et al.*, XRD was performed on eight species of bivalves.⁸¹ They demonstrated that the presence of specific polymorphs depends on the evolutionary taxonomy of the bivalve's species (e.g., shells within the same subclass have very similar



Natural Sources of CaCO₃

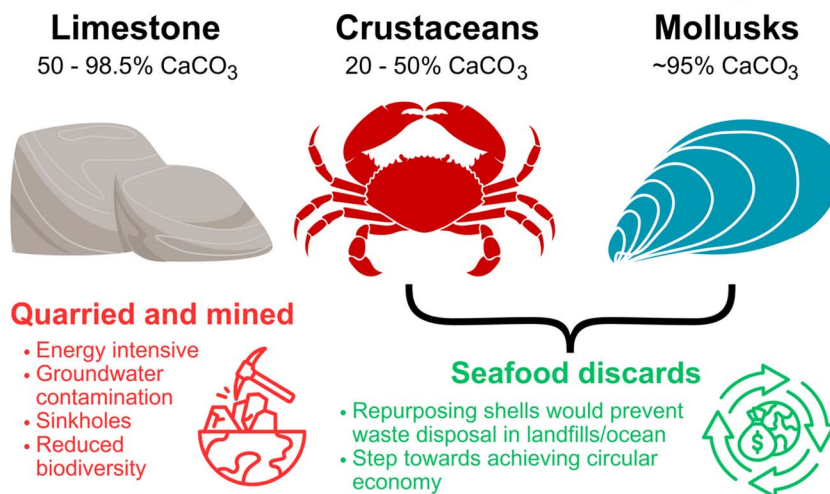


Fig. 2 Natural sources of CaCO₃ include limestone and calcifiers (e.g., crustaceans & mollusks). Prepared with Canva.

composition). A simplified phylogenetic tree of bivalves is shown in Fig. 3.

Species that evolved within the same subclass have similar properties and contain the same CaCO₃ polymorph or a mixture of polymorphs.⁸¹ For example, bivalves belonging to the Heterodonata and Pteriomorphia subclasses are almost entirely made of aragonite. Species belonging to the Heterodonata subclass include mud-dwelling clams (*Meretrix petechialis*), Manila clams (*Ruditapes philippinarum*), and cockles (*Fulvia mutica*) while lischke (*Scapharca subcrenata*) belongs to the Pteriomorphia subclass. The Eupteriomorphia subclass of bivalves includes species that have shells containing either

calcite alone or a mixture of calcite and aragonite. The Ostreoida subclass includes species that have only calcite in their shells, such as Pacific oysters (*Crassostrea gigas*) and yesso scallops (*Patinopecten yessoensis*). A combination of calcite and aragonite is found in the shells of species from the Pterioda (e.g., pen shells, *Atrina pectinata*) and Mytiloida (e.g., Mediterranean mussels, *Mytilus galloprovincialis*) orders of the subclass Eupteriomorphia; however the calcite content consistently greater than that of aragonite.

Mussel shells are the most studied source of CaCO₃ described in the literature. Murphy *et al.* characterized the mussel shells cleaned enzymatically and showed that calcite

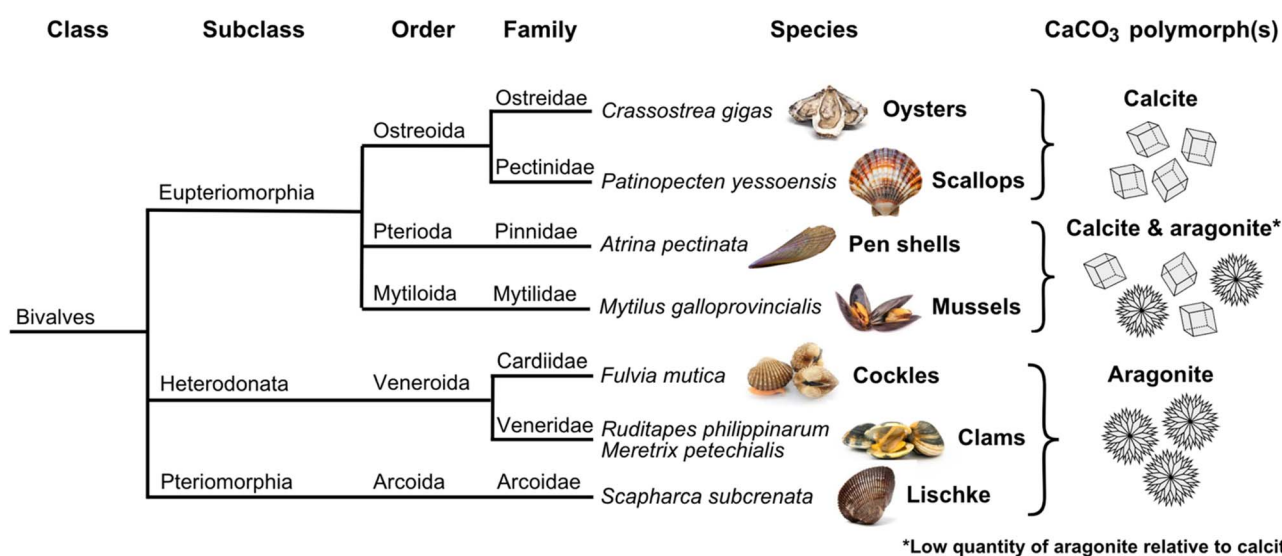


Fig. 3 Phylogenetic tree of bivalves. The subclasses Heterodonata and Pteriomorphia (e.g., cockles, clams, and lischke) have shells made almost entirely of aragonite. The bivalves from the Pterioda and Mytiloida orders of the Eupteriomorphia subclass have shells made of calcite and aragonite while those from the Ostreoida order have shells made solely of calcite. These findings demonstrate that there is a relationship between CaCO₃ polymorphs present in shells and the evolutionary path of specific species of bivalves. Prepared with Canva.



and aragonite were present by XRD analysis.^{68,77,79} Since that study, it has been consistently observed that blue mussel shells in particular contain a mixture of calcite and aragonite.⁷⁸ The same trend can be applied to the shells from sururu mussels, but in this case there is significantly more aragonite than calcite.⁷⁸ Interestingly, green mussel shells were observed by XRD to be composed entirely of aragonite,⁸⁷ which may be a geographical evolutionary trait.

Clam and oyster shells have also been investigated as potential feedstocks of CaCO₃, although they are less prevalent. Águila-Almanza *et al.* observed that clams were made of calcite and aragonite; however they did not specify the specific species of clams studied but rather categorized them based on their shell colour.⁸³ Cardoso *et al.* studied the shell composition of *Anomalocardia brasiliana* by XRD and reported that aragonite was the only polymorph present.⁷⁸ Oyster shells have been shown by multiple studies to consist mostly of calcite.^{81,83–85}

2.1.2. Crustaceans. Compared to bivalves, there is not as much research into utilizing raw shells from crustaceans as a source of CaCO₃ because of the overall composition of their shells. While mussel shells are made of 95% CaCO₃, crab shells only contain 20–50% CaCO₃. Among all crustacean species (crabs, lobsters, and shrimp), crabs have been investigated the most as potential CaCO₃ feedstocks. Often, crab waste utilization is achieved by sequential extraction of various bio-products to yield chitin, proteins, and pigments.³² This involves demineralization and results in other calcium-based products.^{32,33} For example, Burke and Kerton described demineralizing snow crab shells using citric acid, thus yielding calcium citrate as a by-product instead of CaCO₃.³³ Raw crab shells from various species have been analyzed as precursors for CaO transformation (to be discussed in Section 2.2). Compared to bivalves, crab shells are much more uniform in terms of polymorph composition. Blue crab (*Callinectes sapidus*),⁸⁶ Mediterranean green crab (*Carcinus aestuarii*),⁸⁶ mangrove crab (*Ucides cordatus*),^{78,80} and flower crab (*Portunus pelagicus*)⁷³ shells are all made solely of calcite – no aragonite was present in any of these species.

2.1.3. Gastropods. Marine gastropods, otherwise known as seasnails, are not studied often as a CaCO₃ source because they are not as popular for human consumption. In 2021, Anand *et al.* observed that two gastropod species, *Loittideoa* and *Oliva reticularis*, were made entirely of aragonite.⁸²

2.1.4. Applications. As mentioned previously, shells can be used as a replacement for cement manufacturing and they could decrease CO₂ emitted from this industry. In Section 2.2 we describe how shells are converted to CaO, but untreated shells can also be used as aggregates in mortar and plaster – an ancient technology that continues to be used in many areas of the world.⁸⁸ A dry mortar requires three components: a binder (*e.g.*, cement), an aggregate (*e.g.*, sand), and additives (*e.g.*, fibers). CaCO₃ does not react with cement and therefore has potential to act as an aggregate material for this industry.⁸⁸ Mo *et al.* reviewed several examples where the properties of different types of seashells (*e.g.*, oysters, scallops, periwinkle, and mussel) were investigated as aggregates in concrete.⁸⁸ While seashells could offer a beneficial effect because of the

filler effect, it should be noted that it has been reported in multiple instances that their incorporation decreases the overall strength. Therefore, the authors of the previous review focused on this specific application recommend keeping aggregate replacement below 20%. Table 3 describes some of these studies and the parameters considered.

In another study by Lertwattanaruk *et al.*, Portland cement incorporated with seashells from four species demonstrated comparable properties to those of typical plastering and masonry construction.⁹⁵ The shells were processed using wet ball milling and had average particle sizes of 13.56–29.87 μm, similar to that of Portland cement (22.82 μm). While the water requirements and setting time were improved by increasing the proportion of seashells incorporated (5–20 wt%), it should be noted that the compressive strength of mortar decreased, but the shell-based mortar continued to have adequate compressive strengths that are still higher than those needed for plastering purposes. Interestingly, the specific properties of shell-based cements were species-dependent. For example, short-necked clams (*Leukoma staminea*) and oysters decreased drying shrinkage of mortars while green mussels and cockles resulted in the opposite effect. According to the authors, mortars prepared with high volumes of short-necked clams were the best in terms of overall performance because of its relatively low mixing water requirement, good compressive strength, lower drying shrinkage, decrease in thermal conductivity, and increased setting time.

Outside of the cement industry, shell-derived CaCO₃ has been explored for other applications like catalysis. For example, Pasa and co-workers prepared a CaCO₃ catalyst from shellfish waste for biocrude production from sugarcane bagasse and ethanol.⁸⁰ Carapaces from mangrove crab and sururu mussels were crushed and sieved before being used as-is for catalysis. After 30 min at 300 °C and 10 bar N₂, biocrudes were prepared from sugarcane bagasse in high yields (72–83%). The shellfish catalysts increased the yields slightly; however there was a greater impact on the product's composition. When anhydrous ethanol was used in the presence of the catalyst, a biocrude rich in furans was observed, while hydrous ethanol instead produced ethyl esters.

Another application for shells involves dye adsorption and oil recovery. In 2020, following their work on enzymatically processing blue mussel shells,⁷⁶ Murphy *et al.* discovered that they can create an absorbent material from the shells termed “soft calcite” (SC).⁷⁹ Unlike calcite that is hard and ordered, SC has a nest-like morphology able to adsorb dyes from aqueous solutions and absorb crude oil from seawater. As mentioned previously, mussel shells are made of calcite and aragonite; however they are always present as two distinct layers. After heating the shells to 220 °C for 48 h, the outer prismatic layer (calcite) and inner nacreous layer (aragonite) can be separated mechanically. The calcite layer then undergoes two subsequent treatments with 5% acetic acid to yield SC. This novel material has a sponge-like morphology when moist, and when dried it has a cotton candy texture. It was demonstrated that SC can adsorb 1–24 wt% of common dyes (*e.g.*, crystal violet) and exhibits a significant absorption capacity of 977 g oil/g SC ± 84 g



Table 3 Summary of recommendations in the literature on using seashells as aggregates and cement

Shell	Replacement	Suggested replacement level reported in the literature	Parameter(s) studied	Reference
Oyster	Fine aggregate	<20%	• Compressive strength	Yang ⁸⁹
Oyster	Fine aggregate	<50%	• Compressive strength	Eo & Yi ⁹⁰
Scallop	Fine aggregate	<40%	• Workability	Varhen ⁹¹
Scallop	Aggregate	20–60%	• Mechanical properties	Cuadrado-Rica ⁹²
			• Compressive strength	
Cockle	Cement	<15%	• Durability	Othman ⁹³
	Fine aggregate	20–30%	• Porosity	
Cockle	Fine aggregate	20–30%	• Compressive strength	Muthusamy ⁹⁴

for crude oil with good recyclability over ten reuse cycles. A more detailed study on the adsorption of methylene blue and safranin-O using the same materials was recently published where adsorption capacities of 1.81 and 1.51 mg g⁻¹, respectively, were achieved.⁹⁶ Since these studies, calcite from blue mussel shells has been investigated for other uses including the production of calcium acetate that can be used as a de-icer⁹⁷ or as a replacement for calcium chloride to form hydrogels.⁹⁸ Other researchers have since used mussel shells to adsorb other industrial dyes, including Eosin Y.⁷⁷

Shells have also been explored as a possible source of CaCO₃ nanoparticles that can carry out drug delivery applications. In 2021, Ibiyeye and Zuki used cockle-shell-derived aragonite CaCO₃ nanoparticles to successfully co-deliver doxorubicin and thymoquinone to treat breast cancer.⁹⁹ The CaCO₃ nanoparticles were synthesized by simply grinding raw shells, adding SB-12, and then rolled on a roller mill for five days.¹⁰⁰ The drug-loaded aragonite nanoparticles were able to successfully destroy breast cancer stem cells and also suppress their properties. The authors describe that this combinational therapy could be useful as a potential curative strategy to manage metastasis and breast cancer recurrence.⁹⁹

2.2. Shells as a precursor to CaO

Rather than quarrying limestone and transforming it to lime, CaCO₃ from shells can serve as the precursor material. The most prevalent method described in the literature to transform shells from bivalves, crustaceans, and gastropods to CaO is by calcination at temperatures above 800 °C.^{78,82,101–104} CaCO₃ decomposes and releases CO₂ at 800 °C, resulting in CaO as the final product. A few authors, including Dampang *et al.*, use NaOH as a pre-treatment before calcination to remove organic residues;¹⁰¹ however this step is probably unnecessary as proteins would decompose during calcination. A list of calcination conditions and properties of the CaO obtained are described in Table 4.

2.2.1. Applications. CaO from calcined shells have been explored as an additive in cement mortar. Compared to using CaCO₃ from untreated shells, calcined shells can be used as an expansive cement additive as described by Jang and colleagues.¹⁰⁵ The current industrial expansive additives require clinkering processes (1450 °C).¹⁰⁶ It should be noted that while the authors claim using calcined oyster shells could be an

alternative, their process still requires 1000 °C for 3 h. Cement was substituted with 0–12 wt% calcined oyster shells and it was observed that it had a significant effect on compressive strength.¹⁰⁵ For example, using only 3 wt% of calcined oyster shells positively contributed to the compressive strength; however doses over 9 wt% had a significantly negative influence on the compressive strength. Therefore, while adding small quantities of calcined shell could enhance the overall properties, large quantities could lead to a lower compressive strength and loss of workability.

Shell-derived CaO has been investigated as a catalyst in the literature,¹⁰⁷ especially for the production of biodiesel. For example, shells from sururu, blue mussels, mangrove crab, and West Indian pointed venus clam were used to create CaO catalysts.⁷⁸ The catalyst was studied for the production of fatty acid methyl esters (FAMES) by transesterification of soybean oil with yields of 79–94% achieved after 3.5 h. The transformation of CaCO₃ to CaO was achieved by calcining selected shells at 900 °C and pre-activating it with methanol for 1 h before reactions. Significantly different results were achieved depending on the shell which the CaO was derived. CaO from sururu mussels had a FAME yield of 89% after 1 h, while the other catalysts had less than a 50% yield in the same time period. The sururu CaO was investigated further, producing FAME yields of 91% after 3 h following four consecutive reuse cycles. While this is lower than the 96% FAME required to meet biodiesel production, it is a step towards using recycled waste materials for more sustainable biodiesel.

Pal and co-workers prepared a CaO catalyst using unspecified “used mollusk shells” as the starting material.¹⁰⁸ Interestingly, rather than just using calcination to transform CaCO₃ to CaO, the authors used a sol-gel method to produce CaCl₂ before hydrolyzing it to Ca(OH)₂ and then finally calcining it at 850 °C to yield CaO. The resulting product had a very distinct cubic morphology with an average particle size distribution of 341 nm. This catalyst was studied for the adsorption of a neutral red dye and the transesterification of cotton seed oil and methanol to biodiesel. When compared to commercial CaO, the mollusk-derived catalyst had better removal efficiency after 140 min (49 vs. 78%). Similar results were observed for transesterification reactions as mollusk-derived CaO had a maximum 98% biodiesel yield compared to commercial CaO which reached 79% with the same catalyst loading after 4.5 h of reaction time.



Table 4 Calcination conditions and properties of CaO produced from shells of various species of shellfish since 2020

Species	Calcination conditions	CaO properties	Reference
Blue mussel (<i>Mytilus edulis</i>)	900 °C for 3 h	Irregular morphology, diameter <5 μm	Cardoso ⁷⁸
Sururu mussel (<i>Mytella falcata</i>)	900 °C for 3 h	Irregular morphology, diameter <2 μm	Cardoso ⁷⁸
Mussel (n.s. ^a)	950 °C for 4 h	Irregular morphology, diameter <10 μm	Nasir & Hazri ¹⁰²
West Indian pointed venus clam (<i>Anomalocardia brasiliana</i>)	900 °C for 3 h	Irregular morphology, diameter >5 μm	Cardoso ⁷⁸
Spiny fileclam (<i>Lima lima</i>)	900 °C for 4 h	Spherical nanoparticles, diameter: 17–31 nm	Anand ⁸²
Mangrove crab (<i>Ucides cordatus</i>)	800 °C for 2 h	Agglomerated microstructures with cauliflower morphology	Rezende ¹⁰³
Mangrove crab (<i>Ucides cordatus</i>)	900 °C for 3 h	Honeycomb morphology, diameter <5 μm	Cardoso ⁷⁸
Flower crab (<i>Portunus pelagicus</i>)	1000 °C for 5 h	n.s. ^a	Wibisono ¹⁰⁴
Sea snail (<i>Loittioidea</i>)	900 °C for 4 h	Spherical nanoparticles, diameter: 20–34 nm	Anand ⁸²
Netted olive sea snail (<i>Oliva reticularis</i>)	900 °C for 4 h	Spherical nanoparticles, diameter: 5–28 nm	Anand ⁸²
Seashell (n.s. ^a)	800–1000 °C for 2–4 h	Agglomerated irregular nanoparticles	Dampang ¹⁰¹

^a Abbreviations – n.s., none specified.

Biodiesel has also been produced from castor oil and methanol using a shell-based CaO catalyst doped with praseodymium.¹⁰⁹ Similarly to Cardoso *et al.*,⁷⁸ mussel shells were calcined at 950 °C for 4 h to yield CaO.¹⁰⁹ This was doped using praseodymium nitrate hexahydrate, Pr(NO₃)₃·6H₂O and further calcined at 700 °C to create 3–11 wt% Pr-CaO catalysts. A maximum FAME yield of 87.42% was achieved using a 7 wt% Pr-CaO catalyst with a 2.5 wt% catalyst loading and methanol to oil ratio of 8 : 1 at 65 °C for 4 h. It should be noted that compared to other methods discussed previously,^{78,108} the reusability of this catalyst decreased significantly to 52.46% after four cycles because of the reduced number of active sites.¹⁰⁹ While doping CaO with praseodymium improved the FAME yield by 7% when compared to using only calcined mussel shells, it has a negative impact on the catalyst's reusability and therefore may be impractical industrially.

Biodiesel production by transesterification of oils is not the only reaction studied using shell-derived CaO catalysts. In 2023, Eddy *et al.* used CaO from mangrove oyster (*Crassostrea gasar*) shells as a photocatalyst for the degradation of procaine penicillin in aqueous solutions.¹¹⁰ The catalyst was prepared following a similar sol-gel procedure to that of Thakur *et al.*¹⁰⁸ The resulting CaO nanoparticles were observed by TEM to be spherical, with an average diameter of 50 nm, but SEM showed that the size was relatively uneven and some particles were >100 μm.¹¹⁰ The photocatalyst had an average band gap of 4.46 eV and an absorption maximum at 280 nm, and it successfully degraded procaine penicillin under sunlight. A maximum of 97% degradation was realized experimentally after a 2 h reaction at a pH of 12 with a drug concentration of 0.0009 M. The catalyst could be reused five times while retaining efficiency above 86%. In a similar study, bromocresol green dye was degraded with efficiencies between 68 and 89% using the same CaO catalyst from oyster shells.¹¹¹

2.3. Transforming shells to precipitated CaCO₃

Precipitated CaCO₃ (PCC) is a purified and refined version of CaCO₃ that has been synthesized commercially since 1841.¹¹² A

typical process to make PCC using limestone involves (i) mining and crushing limestone, (ii) calcination to form lime, (iii) addition of water to form calcium hydroxide, Ca(OH)₂, (iv) combining Ca(OH)₂ with CO₂ to form CaCO₃, and (v) drying the slurry to yield PCC (eqn (1)).¹¹³ Other reactions that produce PCC include the lime soda process (eqn (2)) and the calcium chloride process (eqn (3)).¹¹²



CaCO₃ produced *via* PCC processes is purer than limestone and a specific polymorph formation can be achieved – including vaterite.⁷⁰ Limestone often contains impurities such as silica and clay (Si, Al, Fe, and Mg)¹¹⁴ that are avoided by using it as a precursor rather than a direct source of CaCO₃. Meanwhile, CaCO₃ must have a purity of >99% to be considered PCC.¹¹² Furthermore, the average size and distribution of particles formed by crushing limestone are much larger than PCC where nanoparticles can be synthesized.⁷⁰ For specific applications, such as filler pigments, PCC must be <2 μm.¹¹² This makes PCC superior to limestone for the preparation of impact resistant plastics and oil absorption.

As mentioned in Section 2.2, shells can also act as a source of CaO by calcining them at high temperatures to decompose CaCO₃. Therefore, they can be used as a precursor to synthesize PCC instead of relying on mined limestone. First, the shells are calcined at 800–1000 °C for several hours to yield CaO.^{71–73,84,115} After calcination, CaO can be treated with HNO₃ before being injected with CO₂.^{71,72} For example, Prihanto *et al.* reported treating CaO with HNO₃ to form an acidic solution of Ca²⁺ ions, and the pH was then increased using NH₄OH, and the desired PCC product was formed by CO₂ injection into the Ca(NO₃)₂ solution.⁷¹ Other methods report that CaO becomes hydrated to form Ca(OH)₂ during storage by absorbing water,⁷³ or the CaO is directly treated with water.⁸⁴



Table 5 Synthesis methods and properties of PCC from various species of shellfish since 2017^a

Species	PCC process	Polymorph(s)	Reference
Green mussel (<i>Perna viridis</i>)	(1) Calcined at 900 °C for 5 h (2) 2 M HNO ₃ (3) pH adjusted to 11 with NH ₄ OH (4) Injected with CO ₂ (0.25 L min ⁻¹) until pH reached 7 (5) Filtered and dried	55.20% vaterite 44.80% calcite	Prihanto ⁷¹
Green mussel (<i>Perna viridis</i>)	(1) Calcined at 800–900 °C for 5 h (2) 2 M HNO ₃ at 60 °C for 30 min (3) pH adjusted to 12 with NH ₄ OH (4) Injected with CO ₂ (5) Washed until pH reached 7 (6) Calcined at 800–900 °C for 5 h	91.26–92.16% vaterite 7.84–8.74% calcite	Ismail ⁷²
Green mussel (<i>Perna viridis</i>)	(1) Calcined at 900 °C for 5 h (2) Absorbed moisture during storage to form Ca(OH) ₂ (3) 0.6 M MgCl ₂ for 0.5–2 h (4) Injected with CO ₂ (50 mL min ⁻¹) until pH reached 7 (5) Filtered and dried	0.5 h stirring: 100% calcite 1 h stirring: 67.1% calcite and 32.9% aragonite 1.5 h stirring: 75.4% calcite and 24.6% aragonite 2 h stirring: 60.6% calcite, 10.0% aragonite, and 29.4% brucite	Irfa'i ¹¹⁵
True oyster (Ostreidae)	(1) Calcined at 1000 °C for 2 h (2) Hydrated with H ₂ O (40 g L ⁻¹) at 80 °C for 1 h (3) 0.6 MgCl ₂ at 80 °C for 3 h (4) Injected with CO ₂ (50 mL min ⁻¹) (5) Filtered and dried	Aragonite	Ramakrishna ⁸⁴
Flower crab (<i>Portunus pelagicus</i>)	(1) Calcined at 900 °C for 5 h (2) 2 M HNO ₃ at 60 °C for 30 min (3) Washed until pH reached 7 (4) Injected with CO ₂ (5) Filtered and dried	Mostly vaterite Aragonite Very low amount of calcite	Bayuseno ⁷³

^a Abbreviations – PCC, precipitated calcium carbonate.

One of the main advantages of using PCC rather than CaCO₃ is that the polymorph formation can be controlled. Naturally-derived CaCO₃ obtained in bulk is sometimes observed as mixture of calcite and aragonite particles with a wide particle size distribution of microparticles and nanoparticles. However, by modifying the conditions of the carbonation reaction to form PCC from Ca(OH)₂, the polymorph content can be tailored. Ramakrishna *et al.* studied the favourable formation of aragonite by optimizing the reaction conditions, such as the presence of MgCl₂, the temperature, and the time.⁸⁴ When using less than 0.1 M MgCl₂, calcite makes up 80% of the PCC formed; however at 0.2 M, Mg²⁺ ions start to incorporate into the Ca²⁺ ion sites of the calcite lattice to form magnesium calcite. At higher concentrations, 0.3 M, aragonite makes up 90% of the product, and at 0.6 M it makes up 100% of the PCC formed. The optimal temperature for the carbonation reaction was determined to be 80 °C by XRD as aragonite was the only polymorph present while calcite remained in small quantities at 60 and 70 °C. Finally, a reaction time of 3 h produced solely aragonite while 2.5 and 2 h resulted in a product containing calcite and unreacted Ca(OH)₂. Irfa'i *et al.* also investigated polymorph formation based on the duration of treatment of Ca(OH)₂ from green mussels with 0.6 M MgCl₂ prior to CO₂ injection.¹¹⁵

Stirring for 0.5 h resulted solely in calcite formation; at 1–1.5 h aragonite began to precipitate, and brucite (Mg(OH)₂) began to form when reacted for 2 h. Some examples of PCC processes and the resulting polymorphs formed are summarized in Table 5.

3. Hydroxyapatite (HAP), Ca₁₀(PO₄)₆(OH)₂

HAP is an inorganic mineral that makes up 60% of human bones¹¹⁶ and has been used extensively in biomedicine because of its biocompatibility,¹¹⁷ non-toxicity,¹¹⁸ and osteo-conductivity.¹¹⁹ Some examples include its incorporation into drug carriers,¹²⁰ surface coatings,¹²¹ anti-tumour drugs,^{122,123} and composites.¹²⁰ HAP has also been applied in dentistry for enamel remineralization and tooth restoration^{123,124} because it makes up 70–80% of human dentin and enamel.¹²⁵ Beyond biomedicine, HAP has also been used for the bioremediation of organic and inorganic pollutants from wastewater,¹²⁶ catalysis,¹²⁷ and energy storage materials.¹²⁸

HAP can be synthesized through a variety of methods that can be divided into three categories: dry, wet, and high temperature.¹²⁹ Solid state and mechanochemical routes are examples of dry methods in which the precursors are mixed in



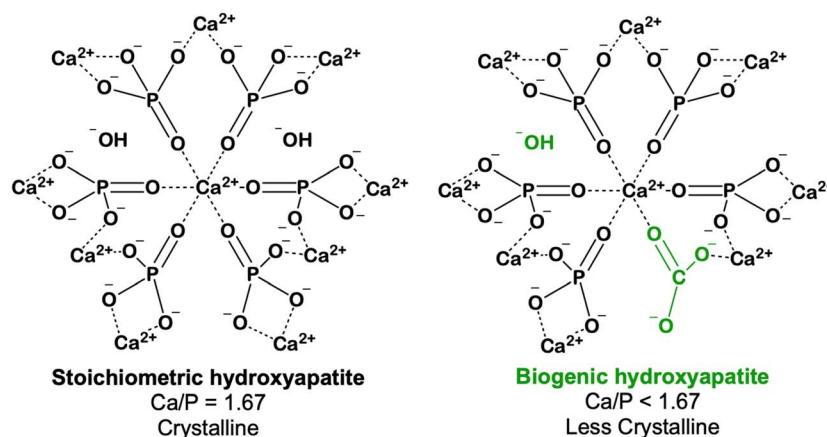


Fig. 4 Structures of stoichiometric HAP, $\text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2$ (left) and biogenic HAP, $\text{Ca}_{10-x}(\text{PO}_4)_{6-x}(\text{CO}_3)_x$ (OH or $\frac{1}{2}\text{CO}_3$) $_{2-x}$ $0 \leq x \leq 2$ (right, green).

a dry form to synthesize HAP. Wet methods, including chemical precipitation, hydrothermal, and hydrolysis reactions are useful for controlling the morphology and size of the produced particles. Pyrolysis and combustion, high-temperature methods, are more rarely reported for HAP synthesis because they can result in the production of secondary aggregates, such as β -tricalcium phosphate (β -TCP).

Biogenic HAP and stoichiometric HAP do not have identical properties because of the presence of impurities. While stoichiometric HAP has the formula $\text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2$ with a Ca/P ratio of 1.67, biogenic HAP, $(\text{Ca}_{10-x}(\text{PO}_4)_{6-x}(\text{CO}_3)_x$ (OH or $\frac{1}{2}\text{CO}_3$) $_{2-x}$); $0 \leq x \leq 2$, has a Ca/P ratio below 1.67 because of the presence of carbonate impurities and calcium and hydroxyl deficiencies (Fig. 4).¹³⁰ Other impurities that have been found in

biogenic HAP include the following metals: Na, Mg, Cr, Co, Cu, and Sr.¹³¹ Biogenic HAP also differs by having a lower degree of crystallinity, thus making it a more amorphous material. Stoichiometric HAP continues to be used for biomedical applications although biogenic HAP may prove to be superior because it is more biomimetic compared to the stoichiometric HAP. Stoichiometric HAP lacks the required carbonate substitutions that are critical for bones' growth, strength, and physiology. It is also a challenge to synthesize biomimetic HAP through a bottom-up process because the surface of biological apatite is rarely smooth and it contains many defects that affect crystallinity, crystal morphology, solubility, and thermal stability. Furthermore, some biomedical applications require HAP with

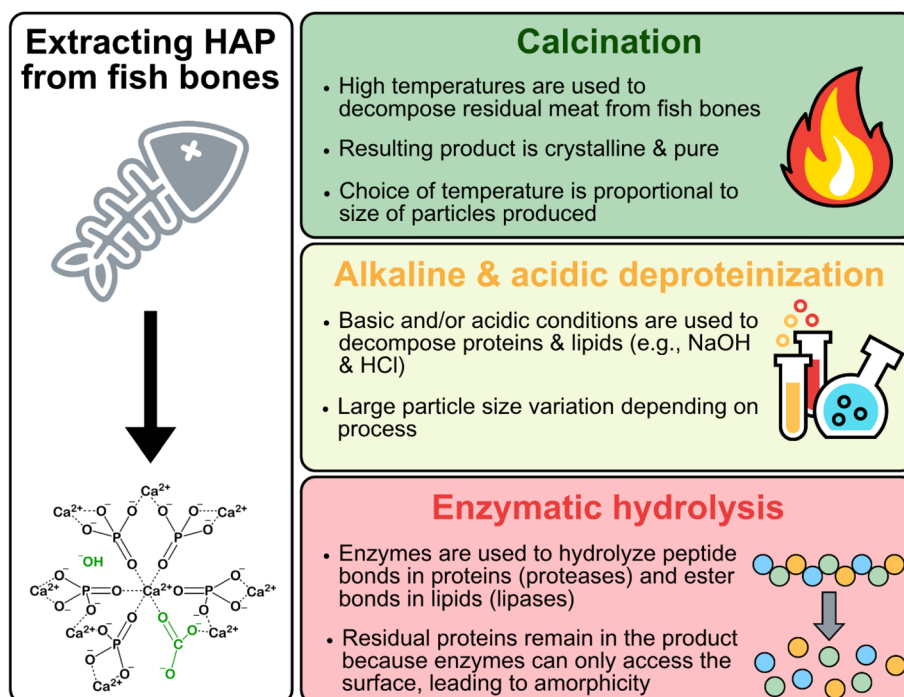


Fig. 5 Extraction techniques to isolate HAP from waste fish discards. Prepared with Canva.



low crystallinity, such as cell bone regeneration because it is more biosorbable.¹³²

Instead of synthesizing biomimetic HAP, this material can be obtained by isolating biogenic HAP from by-products of the seafood processing industry. Shells can be utilized as calcium precursors for HAP synthesis through bottom-up processes by converting the CaCO₃ in shells to CaO through calcination. The CaO can then be converted to HAP by reaction with a phosphate precursor (e.g., H₃PO₄ and NH₄H₂PO₄).¹³³ Cuttlefish bones, which are made of CaCO₃, have also been investigated to synthesize HAP by replacing the carbonate with a phosphate to preserve the porous CaCO₃ structure (note: despite the name, cuttlefish are mollusks).¹³⁴ Fish bones remain the most extensively researched source of HAP, followed by scales. Several forms of fish-based biomass can be exploited to directly isolate HAP, primarily bones, scales, and teeth.

3.1. Bones

As animal species evolved from primitive organisms, the existing exoskeletons based on CaCO₃ were replaced with apatite-containing endoskeletons.¹³⁵ HAP has been isolated from the bones of many species of fish, ranging from sardines to tuna, and there are no obvious differences in HAP content reported between species. There are three broad processes that have been reported in the literature to remove the organic residues from the discards to yield clean bones as a source of HAP: calcination, alkaline/acidic deproteinization (AAD), and enzyme hydrolysis (Fig. 5).

3.1.1. Calcination. Calcination is the most reported method in the literature to remove organic residues from waste fish by-products to produce clean bones. This process involves heating the waste to high temperatures for several hours (e.g., 1200 °C for 6 h)¹³⁶ in the presence of oxygen. It is a relatively simple method that does not require extensive sample preparation. Fish bones have been directly calcined as-is without pre-treatment (other than drying) in several studies.^{137,138} Boiling the fish waste ahead of calcination for a few hours is a common pre-treatment step prior to calcination; however as demonstrated in some studies, it is not essential.^{138–140} Furthermore, some methods report treating the by-products with acetone¹⁴¹ or sodium hydroxide¹⁴² as an additional (but not necessary) pre-treatment step. Grinding the fish bones (e.g., in mortar or ball-mill) is often performed before calcination,^{139–141,143–148} but this can be performed before or after exposing the fish bones to high temperatures (Table 6).

The choice of temperature has an impact on the HAP-containing material produced. In most studies reported to date, a range of temperatures up to 1200 °C have been explored. For example, Pal *et al.* studied the impact of different temperatures to treat barramundi (*Lates calcarifer*) waste by heating it to 200, 400, 800, 1,000, and 1200 °C with a heating ramp of 5 °C min⁻¹ and holding it isothermally for 1 h.¹⁴⁹ By XRD analysis, it was observed that the crystallinity of the product increases depending on the chosen temperature. The samples treated at 200 and 400 °C were amorphous; however those treated at 800 °C had increased crystallinity of 80.8%. The crystallinity further

increased to 89.2 and 92.9% when heated to 1000 and 1200 °C, respectively. A similar trend was observed for crystallite size, which increased from 5 nm when treated at 200 °C to 63 nm at 1200 °C. Similar results were achieved by Modolon *et al.* when they calcined tilapia (*Oreochromis niloticus*) bones at 600, 900, and 1200 °C for 2 h, observing crystallinities of 39.5, 98.3, and 99.6% and crystallite sizes of 14.4, 85.8, and 95.0 nm, respectively.¹⁵⁰

At higher calcination temperatures, biphasic calcium phosphates (BCPs) are formed from the decomposition of HAP, including β-TCP and/or α-tricalcium phosphate (α-TCP).¹⁵¹ This was reported by Castilho and co-workers after calcining black scabbardfish (*Aphanopus carbo*) bones at 400, 600, 800, and 1000 °C for 4 h, resulting in samples consisting of 0, 17.5, 29.7, and 40.1% β-TCP, respectively.¹³⁹ The formation of β-TCP is inconsistent across species. For example, Permatasari *et al.* observed that houndfish (*Tylosurus crocodilus*) bones calcined at 1000 °C for 4 h formed β-TCP while milkfish (*Chanos chanos*) and walking catfish (*Clarias batrachus*) did not.¹⁴⁰ This ultimately resulted in houndfish having much lower crystallinity (67.2%) compared to milkfish (98.3%) and walking catfish (95.0%).

The choice of calcination temperature also impacts size and morphology of BCP particles. In general, particles created at lower temperatures are smaller but less regular than those prepared at higher temperatures. For example, Carella *et al.* calcined round sardinella (*Sardinella aurita*) for 1 h at 300, 600, and 900 °C.¹⁵² The sample calcined at 300 °C had an aligned arrangement of char that remained from the incomplete combustion of organic matter, but the size of individual particles could not be established. At 600 °C, the particles were agglomerated and spherical with a diameter of 50–100 nm. When the temperature increased to 900 °C, the particle size increased drastically to 0.20–2.0 μm with rod shape geometry. Interestingly, the morphology of BCP particles from fish bones differs across studies. Mamun *et al.* calcined tilapia and surma (*Katsuwonus pelamis*) at 900 °C for 2 h and both species resulted in agglomerated spherical particles with diameters less than 1 μm.¹⁴⁸ Adam and co-workers reported that black tilapia (*Oreochromis placidus*) waste produced crystal plate-like particles that are 0.5–4 μm in diameter.¹⁵³ Finally, Renda *et al.* observed that calcining tambaqui (*Colossoma macropomum*) at 900 °C for 2 h yielded cubic particles with an average diameter of 620.2 nm.¹⁴⁴ More details regarding calcination procedures, including the Ca/P ratio, crystallinity, and size of yielded particles, are summarized in Table 6. Processes reported without sufficient characterization information have been omitted from the table.^{137,138,154–159}

3.1.2. Alkaline and acidic deproteinization (AAD). Deproteinization under acidic or basic conditions can remove protein residues from the waste fish frames. HCl and/or NaOH are typically used at concentrations between 0.1 and 1 N; however these processes often involve another treatment step to achieve the desired product, such as the use of more concentrated NaOH. Approaching this from a green standpoint, the use of acids and bases at high concentrations is undesirable because they pose a safety risk by causing severe burns and are highly



Table 6 Review of calcination treatments described in the literature to isolate BCPs^a from fish bones since 2015

Species	Optimal treatment	Ca/P	Crystallinity	Size	Morphology	Reference
Black scabbardfish (<i>Aphanopus carbo</i>)	(1) Ground with a hand mill (2) Calcined at 400–1000 °C for 4 h	1.47–1.54	n.s. ^a	<100 nm	Isotropic	Ideia ¹³⁹
Milkfish (<i>Chanos chanos</i>)	(1) Crushed using a ball mill (2) Calcined at 1000 °C for 4 h	1.66	^b 98.3%	n.s. ^a	n.s. ^a	Permatasari ¹⁴⁰
Belida (<i>Chitala lopis</i>)	(1) Crushed to fine powder (2) Carbonized at 500 °C for 2 h	n.s. ^a	^c 26.8%	<20 µm	Rough and irregular	Lestari ¹⁴³
Walking catfish (<i>Clarias batrachus</i>)	(1) Crushed using as ball mill (2) Calcined at 1000 °C for 4 h	1.68	^b 95.0%	n.s. ^a	n.s. ^a	Permatasari ¹⁴⁰
Tambaqui (<i>Colossoma macropomum</i>)	(1) Briefly boiled (2) Ground with a mortar (3) Calcined at 600/900 °C for 2 h	n.s. ^a	High	481.1–721.7 nm (average 602.3 nm)/ 561.7–666.6 nm (average 620.2 nm)	Agglomerates of spherical/cubical particles	Renda ¹⁴⁴
Common carp (<i>Cyprinus carpio</i>)	(1) Boiled for 2 h (2×) (2) 500 °C for 2 h (3) Calcined at 800–1200 °C for 2–6 h (4) Ground with a mortar	1.53–1.73	High	<1 µm	Agglomerates of uniform polyhedral grains	Maidaniuc ¹³⁶
Mackerel scad (<i>Decapterus macarellus</i>)	(1) Boiled for several hours (2) Ground with a grinder (3) Calcined at 900 °C for 5 h	n.s. ^a	High	100–500 µm	Spherical microstructures	Ishak ¹⁴⁵
South Asian carp (<i>Gibelion catla</i>)	(1) Manually blended (2) Boiled for 8 h (3) 900 °C for 4 h (4) Ground with a mortar	n.s. ^a	Polycrystalline	200–260 nm	Hexagonal particles	Sathiyavimal ¹⁴⁶
Asian stingray catfish (<i>Heteropneustes fossilis</i>)	(1) Boiled for 1.5 h (2) Crushed into pieces (3) Calcined at 900 °C for 2 h	2.82	High	<1 µm	Agglomerated spherical particles	Moullick ¹⁴⁷
Surma (<i>Katsuwonus pelamis</i>)	(1) Boiled (2) Crushed (3) Calcined at 900 °C for 2 h	3.16	^d 92%	<1 µm	Agglomerated spherical particles	Mamun ¹⁴⁸
Nile tilapia (<i>Oreochromis niloticus</i>)	(1) Calcined at 900 °C for 8 h (2) Ball-milled for 8 h	1.66	n.s. ^a	650–2000 nm (average 1200 nm)	Agglomerates of irregular and rounded particles	da Cruz ¹⁶⁰
Nile tilapia (<i>Oreochromis niloticus</i>)	(1) Boiled (2) Calcined at 800 °C for 5 h (3) Ground with a herb grinder for 10 min	1.64	Low	10–25 µm	Irregular-shaped particles	Khamkongkaeo ¹³²
Nile tilapia (<i>Oreochromis niloticus</i>)	(1) Boiled (2) Crushed in a knife-mill (3) Ball-milled (4) Calcined at 600–1200 °C for 2 h (5) Ball-milled for 1–3 h with isopropyl alcohol	1.85–1.93	^d 39–99%	1–2 µm	Irregular particles that become bigger with temperature	Modolon ¹⁵⁰
Nile tilapia (<i>Oreochromis niloticus</i>)	(1) Boiled (2) Crushed into small pieces (3) Calcined at 900 °C for 2 h	2.99	^d 94%	<1 µm	Agglomerated spherical particles	Mamun ¹⁴⁸
Black tilapia (<i>Oreochromis placidus</i>)	(1) Boiled (2) Calcined at 800 °C	n.s. ^a	High	0.5–4 µm (average 1.2 µm)	Plate-like structure	Hubadillah ¹⁵³



Table 6 (Contd.)

Species	Optimal treatment	Ca/P	Crystallinity	Size	Morphology	Reference
Pama croaker (<i>Otolithoides pama</i>)	(1) Boiled for 1.5 h (2) Crushed into pieces (3) Calcined at 900 °C for 2 h	2.82	n.s. ^a	<1 μm	Agglomerated spherical particles	Prosad Moulick ¹⁴⁷
Perch (<i>Perca</i> sp.)	(1) Boiled for 8 h (2) Calcined at 900 °C for 2 h (3) Ground with a mortar	~1.67	High	50–80 nm	Agglomerated spherical particles	Naga ¹⁶¹
Round sardinella (<i>Sardinella aurita</i>)	(1) Calcined at 300–900 °C for 1 h (2) Ground and sieved (50 μm)	1.54–1.60	^e <5–60%	50–100 nm (600 °C); 0.2–2 μm (900 °C)	Round particles (600 °C); regular and coarse rods (900 °C)	Carella ¹⁵²
Atlantic salmon (<i>Salmo salar</i>)	(1) Immersed in NaOH for 24 h (2) Calcined at 850 °C for 4 h (3) Crushed with a mortar (4) Ball-milled for 4 h	1.57	High	<5 μm	Agglomerated spherical particles	Popescu-Pelin ¹⁴²
Rabbitfish (<i>Siganus</i> sp.)	(1) Boiled (2) Boiled in water & acetone for 1 h (3) Ground with a mortar into 200-mesh powder (4) Annealed at 800–1000 °C for 2 h (5) Ball-milled for 30 min	1.696–1.876	n.s. ^a	1–5 μm	Agglomerated irregular particles	Fendi ¹⁴¹
Catfish (<i>Siluriformes</i> sp.)	(1) Boiled for 4 h (2) Sintered at 900 °C for 2 h (3) Ground with a mortar and sieved (300 μm)	1.58	^d 99%	<200 μm	Irregular flower-like microstructures with petal-shaped flakes	Akpan ¹⁶²
Gilt-head bream (<i>Spanus aurata</i>)	(1) Immersed in NaOH for 24 h (2) Calcined at 850 °C for 4 h (3) Crushed with a mortar (4) Ball-milled for 4 h	1.63	High	<5 μm	Agglomerated spherical particles	Popescu-Pelin ¹⁴²
Spanish mackerel (<i>Tenggiri scomberomorini</i>)	(1) Crushed to fine powder (2) Carbonized at 500 °C for 2 h	n.s. ^a	25.6% ^c	<20 μm	Rough and irregular particles	Lestari ¹⁴³
Houndfish (<i>Tylosurus crocodilus</i>)	(1) Crushed with a ball mill (2) Calcined at 1000 °C for 4 h	1.62	^b 67.2% ^a n.s.	n.s. ^a	n.s. ^a	Permatasari ¹⁴⁰

^a Abbreviations – BCP, biphasic calcium phosphate; n.s., non-specified. ^b Crystallinity calculated using an unspecified method. ^c Crystallinity calculated using software. ^d Crystallinity calculated by comparing the intensity of the largest peak and the valley between the largest peak and first peak on the right (eqn (S1)). ^e Crystallinity calculated by comparing the sum of peaks and the area between the peaks and background (eqn (S2)).

reactive. Furthermore, acids and bases can pose an environmental risk if not disposed of properly. That being said, they continue to be used to isolate HAP from fish by-products. For example, Surya *et al.* described boiling sardine (*Sardinella longiceps*) bones in 2% NaOH and acetone for 1 h, followed by another treatment with 5% NaOH for 5 h at 70 °C.¹⁶³ This was then further treated with 50% NaOH at 100 °C for 1 h to yield nano-hydroxyapatite (nHAP) particles.¹⁶³ In this reaction, not only is concentrated NaOH used that is corrosive and reactive, but acetone is also employed which can generate leachates. Rather than using 50% NaOH and acetone, Nag *et al.* immersed fish bones from various fish species in 30% H₂O₂ after an initial treatment with a commercial detergent solution.¹⁶⁴

Autoclaving and calcination are other treatment steps that are used in conjunction with AAD to ensure thorough removal of organic residues. Triwitono and co-workers treated fish bones from six species of fish by autoclaving them at 121 °C for 3 h before using 1 N HCl and 1 N NaOH.¹⁶⁵ Meanwhile, Hariani *et al.* calcined milkfish (*Chanidae* sp.) at 750 °C for 3 h after using 0.1 N HCl and 50% NaOH.¹⁶⁶ The method reported by Idowu *et al.* is quite different from others: Atlantic salmon bones were immersed in 2 M NaOH and soaked in hexane before being treated with 2.5% NaClO and finally calcined at 900 °C for 6–9 h.¹⁶⁷

AAD often yields a more amorphous product, whereas calcination increases crystallinity. For example, Horta *et al.* extracted HAP from tambaqui bones with 1% NaOH and calcined a portion of these bones at 800 °C.¹⁶⁸ They determined that the alkaline treatment decomposed a significant amount of organic material, but calcination was required to increase the crystallinity significantly. Based on SEM observations, samples not exposed to heat were made of agglomerated, irregularly shaped nanoparticles while calcination yielded larger, more defined particles. As with calcination, the HAP particles tend to agglomerate regardless of the method of preparation and it is often difficult to compare results since some studies report the size of individual nanoparticles while others describe the size of agglomerates. Deproteinization treatments involving acids and/or bases are summarized in Table 7 but reports that did not include sufficient characterization data have been omitted.^{164,169}

3.1.3. Enzymatic hydrolysis. Using enzymes is a relatively uncommon technique to isolate HAP from waste fish bones compared to calcination and alkaline/acidic deproteinization. It is the most sustainable and industry-friendly option because of the limited amounts of solvent, acid/base, and heat required for a successful process. Proteases are the most commonly studied enzymes for decomposing organic content in fish by-products, although other enzymes such as lipases and enzymatic cocktails have also been used. Fitri *et al.* used 8% papain enzyme for 10 h at 60 °C in water to treat milkfish bones; however some lipid content remained in the sample as indicated by IR data.¹⁷³ Furthermore, the product was highly amorphous based on the XRD diffractogram of the treated milkfish bones. Boudreau *et al.* overcame this challenge by using a protease, Neutrase, and a lipase, Lipozyme CALB L, simultaneously in water at 40 °C for 6 h.¹⁷⁴ While collagen remains present within the bone matrix, this is expected, as the enzymes do not penetrate the solid

material in which the collagen is encased. Compared to Fitri *et al.*,¹⁷³ the lipid content was significantly reduced by the addition of a lipase. Boudreau *et al.* continued this research by using mechanochemistry techniques to transform the cleaned fish bone macroparticles into HAP nanoparticles for further applications.¹⁷⁵ Furthermore, a simplified gate-to-gate life cycle analysis (LCA) showed that using enzymes and mechanochemistry to yield HAP nanoparticles is safer for human exposure and releases significantly lower quantities of CO₂ compared to using calcination and/or AAD.¹⁷⁵

To achieve a product with higher purity, Pou and co-workers calcined fish bones from horse mackerel (*Trachurus trachurus*), scorpionfish (*Scorpaena scrofa*), and Atlantic salmon at 750–950 °C for 10 h after treating them with 1% Alcalase for 4 h at 64.2 °C.¹⁷⁶ As expected, the extracted HAP is more crystalline and has no traces of organic residues (*e.g.*, lipids and proteins) compared to that obtained by Boudreau *et al.* because of the calcination step.^{174–176} The samples calcined at 950 °C were more crystalline than those calcined at 750 °C, but higher amounts of β-TCP were also detected. Similar results were achieved by enzymatically treating catfish (*Pangasius hypophthalmus*) waste with Alcalase and calcination for 3 h at 500–900 °C.¹⁷⁷ As the temperature increased, partial decomposition of HAP to β-TCP also increased.

Enzymes have the potential to create a more circular economy for the seafood processing industry. As mentioned above, not only are enzymes considered green, but every aspect of the fish waste could be utilized. While the focus of this review is on the inorganic fraction of the by-products, the protein hydrolysate remaining after enzymatic treatment contains beneficial amino acids that could be incorporated into fertilizers.^{178–180}

3.2. Scales

Researchers have successfully isolated HAP from waste fish scales despite having lower inorganic content (38–46%) compared to bones. While many follow processes previously reported for the treatment of bones (*i.e.*, calcination, AAD, and enzymes), other novel methods have also been used such as ultrasonic-assisted extraction (UAE) and treatment with DES. Among these methods, alkaline deproteinization is the most common treatment, followed by calcination, UAE, and finally DES. The isolated HAP is often observed as agglomerates of particles with sizes ranging from 5 nm to 167 μm. A detailed list of these methods is summarized in Table 8 but reports that did not provide sufficient characterization information are omitted.^{181–184}

3.2.1. Calcination and pyrolysis. To study the impact of temperature, Ideia *et al.* calcined scale-containing skins from grey triggerfish (*Balistes capricus*) at 400, 600, 800, and 1000 °C for 4 h.¹³⁹ Sardine (*Sardina pilchardus*) bones were also treated under the same conditions to compare the HAP obtained from bones to scales. At 400 °C, both samples were poorly crystalline; however crystallinity increased proportionally for scales and bones with temperature. Interestingly, while the bones calcined at 1000 °C were composed of 40.1 wt% β-TCP and 59.9 wt%



Table 7 Review of AAD^a treatments described in the literature to isolate HAP from fish bones since 2018

Species	Optimal treatment	Ca/P	Crystallinity	Size	Morphology	Reference
Milkfish (<i>Channidae</i> sp.)	(1) Crushed by fast milling (2) Sieved through a 200-mesh sieve (3) Washed with 0.1 N HCl (4) Treated with 50% NaOH for 5 h at 60 °C (5) Calcined at 750 °C for 3 h	1.65	n.s. ^a	<3 µm	Irregular and agglomerated particles	Hariani ¹⁶⁶
Tambaqui (<i>Colossoma macropomum</i>)	(1) Boiled in water for 1 h (2) Treated with 1% NaOH at 90 °C for 5 h (3) Crushed with a mortar (4) Calcined at 800 °C	1.67	n.s. ^a	<500 nm	Defined morphology with a faceted structure	Horta ¹⁶⁸
Grouper (<i>Epinephelus</i> sp.)	(1) Boiled in water for 1 h (2) Autoclaved at 121 °C for 3 h (3) Crushed with a mortar (4) Soaked in 1 N HCl for 1 h (5) Centrifuged for 15 min (6) Treated with 1 N NaOH for 1 h at 100 °C (3×) (7) Neutralized with 1 N HCl and centrifuged (8) Refined with a disc mill for 1 min (9) Sieved through a 200-mesh sieve	1.44	^b 69.8%	Ave. 281.4 nm	Agglomerates of asymmetrical particles	Kusumawati ¹⁶⁵
Barramundi (<i>Lates calcarifer</i>)	(1) Boiled in water for 1 h (2) Boiled in 1% NaOH (3) Calcined at 650 °C for 4 h (4) Crushed with a mortar	1.845	High	Ave. 39.42 nm long, 17.15 nm wide	Spherical and rectangular particles	Le Ho ¹⁷⁰
Snapper (<i>Lutjanus</i> sp.)	(1) Boiled in water for 1 h (2) Autoclaved at 121 °C for 3 h (3) Crushed with a mortar (4) Soaked in 1 N HCl for 1 h (5) Centrifuged for 15 min (6) Treated with 1 N NaOH for 1 h at 100 °C (3×) (7) Neutralized with 1 N HCl and centrifuged (8) Refined with disc mill for 1 min (9) Sieved through a 200-mesh sieve	1.38	^b 71.2%	Ave. 254.7 nm	Agglomerates of asymmetrical particles	Kusumawati ¹⁶⁵
Whitemouth croaker (<i>Micropogonias furnieri</i>)	(1) Washed with 1 N NaOH for 24 h (2) Washed in water overnight (3) Treated with 30% H ₂ O ₂ for 24 h (4) Calcined at 800 °C for 5 h (5) Milled	1.40	High	>10 µm	Agglomerates of asymmetrical particles	Yamamura ¹⁷¹
Tilapia (<i>Oreochromis niloticus</i>)	(1) Boiled in water for 1 h (2) Autoclaved at 121 °C for 3 h (3) Crushed with mortar (4) Soaked in 1 N HCl for 1 h (5) Centrifuged for 15 min (6) Treated with 1 N NaOH for 1 h at 100 °C (3×) (7) Neutralized with 1 N HCl and centrifuged (8) Refined with disc mill for 1 min (9) Sieved through a 200-mesh sieve	1.61	^b 70.2%	Ave. 87.4 nm	Agglomerates of asymmetrical particles	Kusumawati ¹⁶⁵



Table 7 (Contd.)

Species	Optimal treatment	Ca/P	Crystallinity	Size	Morphology	Reference
Catfish (<i>Pangasius</i> sp.)	(1) Boiled in water for 1 h (2) Autoclaved at 121 °C for 3 h (3) Crushed with mortar (4) Soaked in 1 N HCl for 1 h (5) Centrifuged for 15 min (6) Treated with 1 N NaOH for 1 h at 100 °C (3×) (7) Neutralized with 1 N HCl and centrifuged (8) Refined with a disc mill for 1 min (9) Sieved through a 200-mesh sieve	1.43	^b 72.2%	Ave. 239.9 nm	Agglomerates of asymmetrical particles	Kusumawati ¹⁶⁵
Atlantic salmon (<i>Salmo salar</i>)	(1) Boiled in water for 1 h (2) Treated with 1% NaOH (3) Calcined at 800 °C for 3 h (4) Ground in a centrifugal ball mill (5) Sieved through a 63 µm sieve	1.63	High	Ave. 122 nm	Agglomerates of grain-like particles	Bas ¹⁷²
Atlantic salmon (<i>Salmo salar</i>)	(1) Immersed in 2 M NaOH at 50 °C for 2 h (2) Ground with a crushing mill until particles were 3–4 mm long (3) Treated with hexanes for 1 h at 25 °C (4) Bleached with 2.5% NaClO for 30 min (5) Bleached with 2.5% H ₂ O ₂ for 1 h (6) Milled with a planetary ball mill for 2.5 h (7) Sieved with a 75 µm sieve (8) Calcined at 900 °C for 6–9 h (9) Milled with a planetary mill	1.66	High	22.2–27.5 µm	Nanoparticles agglomerates into large clusters	Idowu ¹⁶⁷
Indian oil sardine (<i>Sardinella longiceps</i>)	(1) Boiled at 200 °C (2) Boiled with 2% NaOH and acetone for 1 h (3) Grinded with a mortar (4) Treated with 5% NaOH at 70 °C for 5 h (5) Precipitants treated with 50% NaOH at 100 °C for 1 h (6) Sieved	n.s.	Low	<5 µm	Agglomerates of asymmetrical particles	Surya ¹⁶³
Kingfish mackerel (<i>Scomberomorus</i> sp.)	(1) Boiled in water for 1 h (2) Autoclaved at 121 °C for 3 h (3) Crushed with a mortar (4) Soaked in 1 N HCl for 1 h (5) Centrifuged for 15 min (6) Treated with 1 N NaOH for 1 h at 100 °C (3×) (7) Neutralized with 1 N HCl and centrifuged (8) Refined with a disc mill for 1 min (9) Sieved through a 200-mesh sieve	1.69	^b 64.9%	Ave. 105.2 nm	Agglomerates of asymmetrical particles	Kusumawati ¹⁶⁵



Table 7 (Contd.)

Species	Optimal treatment	Ca/P	Crystallinity	Size	Morphology	Reference
Tuna (<i>Thunnus</i> sp.)	(1) Boiled in water for 1 h (2) Autoclaved at 121 °C for 3 h (3) Crushed with a mortar (4) Soaked in 1 N HCl for 1 h (5) Centrifuged for 15 min (6) Treated with 1 N NaOH for 1 h at 100 °C (3×) (7) Neutralized with 1 N HCl and centrifuged (8) Refined with a disc mill for 1 min (9) Sieved through a 200-mesh sieve	1.36	^b 64.3%	Ave. 150.2 nm	Agglomerates of asymmetrical particles	Kusumawati ¹⁶⁵

^a Abbreviations – AAD, alkaline and/or acidic deproteinization; n.s., none specified; Ave., average. ^b Crystallinity calculated using software.

HAP, the scales were almost entirely made of HAP (95.8 wt%). β -TCP was present in scales only at 600 °C and in much lower quantities (4.6 wt%) compared to bones. Scales also contained minerals that were not observed in bone samples such as halite, NaCl, and rhenanite, NaCaPO₄. Halite was present in samples calcined at 200–800 °C and decreased with increasing temperature, while rhenanite was only observed in scales treated at 600 and 800 °C. Additionally, the remaining 4.2 wt% of scales calcined at 1000 °C was attributed to magnesium oxide, MgO, and cubic trisodium phosphate, γ -Na₃PO₄. The samples were studied by SEM and particulate size increased with temperature as a consequence of grain growth due to the added energy. The scale-derived particles were consistently larger than those created from bone, regardless of temperature. Furthermore, the particles prepared had completely different morphologies. While the calcination of bones at 600 °C resulted in agglomeration of spherical particles, scales yielded rod-like particles with a hexagonal prism morphology. Also, at 800 °C, there were other particles with a secondary flower-like morphology observed in the heated skin samples, which were attributed to rhenanite.

In 2022, Rattanakam and co-workers published a study on the dissolution performance of carbon/HAP nanocomposites from tilapia scales.¹⁹⁴ Instead of calcining, they pyrolyzed the discards at 450, 500, 550, and 600 °C for 5 h. The final material prepared was ultimately collected on a 0.5-mm sieve prior to analysis. The Ca/P ratio of each product was significantly higher (2.43–2.50) than those observed from other thermal treatments, but this can be attributed to the fact this was a pyrolysis treatment in the presence of nitrogen rather than calcination. Without oxygen, biochar was produced from the decomposition of protein, and therefore the carbon content was high (22.6–25.6 wt%). Broad and low intensity peaks were observed in the XRD diffractograms, indicating that the products were amorphous. The authors highlighted that the crystal size of the samples did not increase proportionally with temperature, unlike calcination, because the carbon in organic residues linked to HAP could block its nucleation sites, preventing

particle growth and recrystallization. The authors later noted that distinguishing between carbon and HAP particles during TEM analysis was not possible because of severe aggregation. Therefore, while pyrolysis may not increase the size of particles yielded, it does not prevent agglomeration.

3.2.2. Alkaline and acidic deproteinization (AAD). While researchers most often rely on calcination to isolate HAP from waste fish bones, AAD is much more common when treating scales because they have a higher lipid content.¹⁹⁹ Often these methods report using concentrated NaOH after pre-treating the scales with dilute acid or base. For example, Eswaran *et al.* used 0.1 M HCl to remove proteins from mullya garra (*Garra mullya*) scales, followed by 5% (w/v) NaOH for 7 h at 70 °C.¹⁸⁷ After being dried overnight and crushed using a mortar and pestle, samples were immersed in 50% (w/v) NaOH for 2 h at 100 °C for further deproteinization. Lee and co-workers followed a very similar procedure to synthesize HAP from black tilapia scales to study its adsorption efficiency for Cr(vi) removal.¹⁹¹ The particles prepared from the tilapia scales were slightly larger (50–60 nm wide and 30–200 nm long)¹⁹¹ than mullya garra (5–20 nm wide and 20–40 nm long).¹⁸⁷

The morphology of HAP particles extracted from natural sources is still not fully understood as the same technique can provide particles with different shapes. For example, Eswaran *et al.*¹⁸⁷ and Injorhor *et al.*¹⁸⁵ followed a method very similar to that reported by Kongsri *et al.*²⁰⁰ HAP from the Nile tilapia scales was in the form of hexagonal crystals with an average size of 20 nm. While the size is similar to those of other HAP particles prepared,²⁰⁰ scales from mullya garra were rod-like¹⁸⁷ and white seabass scales had irregular morphology.¹⁸⁵ While Eswaran *et al.* describe grinding their scales with a mortar and pestle during their multi-step alkaline treatment,¹⁸⁷ the other methods do not mention any further treatment prior to characterization.^{185,200} More research into the choice of mechanochemical processing method (*e.g.*, mortar and pestle, ball-milling, and planetary mill) could provide further insight into whether it has an impact on particle morphology.



Table 8 Review of calcination, AAD,^a DES,^a and UAE^a treatments described in the literature to isolate BCPs^a from fish scales since 2020

Method	Species	Optimal treatment	Ca/P	Crystallinity	Particle size & morphology	Reference
AAD ^a	White seabass (<i>Atractoscion nobilis</i>)	(1) Immersed in 0.1 M HCl for 1 h (2) Treated with 5% NaOH at 60 °C for 3 h (3) Treated with 50% NaOH at 80 °C for 3 h	2.01	^b 80.99%	Size: 100–500 nm (mean: 172.9 nm)	Injorhor ¹⁸⁵
	Catla (<i>Catla catla</i>)	(1) Soaked in 83 mL of 1 N HCl for 30 min (2) pH raised to 11 with 6 N NaOH (3) Vacuum filtered (4) Boiled in a bag in a water bath for 30 min (5) Frozen at –80 °C overnight (6) Incubated in a hot air oven at 160 °C overnight (7) Calcined at 800 °C for 2 h	1.66	n.s. ^a	Size: <1 μm Morphology: agglomerates of needles	Buraiki ¹⁸⁶
	Mullya garra (<i>Garra mullya</i>)	(1) Immersed in 500 mL 0.1 M HCl (2) 250 mL 5% NaOH added and stirred at 70 °C for 7 h (3) Dried precipitates crushed using a mortar and pestle (4) Stirred in 100 mL 50% NaOH at 100 °C for 2 h	1.67	High	Size: 20–40 nm long and 5–20 nm wide Morphology: interconnected nanostructured rods	Eswaran ¹⁸⁷
	Rohu (<i>Labeo rohita</i>)	(1) 100 g immersed in 0.6% KOH (×2) (2) Calcined at 550–1000 °C for 3 h	1.52–1.82	n.s. ^a	Size: 1–1000 μm, mean of 155–167 μm Morphology: surface layer with bony ridges of HAP in concentric rings & an inner fibrous layer of mainly collagen	Sarkar and Das ¹⁸⁸
	Asian sea bass (<i>Lates calcarifer</i>)	(1) Stirred in 1 L of 0.25 M HCl for 2 h (2) Calcined at 900 °C for 3 h (3) Ground and sieved through a cloth mesh	1.67	n.s. ^a	Size: 50–150 nm Morphology: spherical agglomerates	Wu ¹⁸⁹
	Nile tilapia (<i>Oreochromis niloticus</i>)	(1) Soaked in 83 mL of 1 N HCl for 30 min (2) pH raised to 11 with 6 N NaOH (3) Vacuum filtered (4) Boiled in a bag in a water bath for 30 min (5) Frozen at –80 °C overnight (6) Incubated in a hot air oven at 160 °C overnight (7) Calcined at 800 °C for 2 h	2.14	n.s. ^a	Size: <1 μm Morphology: spherical agglomerates	Buraiki ¹⁸⁶
	Nile tilapia (<i>Oreochromis niloticus</i>)	(1) Ground into a fine powder (2) Stirred in 1 L 0.75 M NaOH at 60 °C for 5 min (3) Solid portion added to 1.3 M HCl at 60 °C for 30 min (4) 5 M NaOH added dropwise (5) Ground	2.25	Low	Size: 250–2500 nm Morphology: dense agglomerated soft morphology with high surface roughness	Rashad ¹⁹⁰



Table 8 (Contd.)

Method	Species	Optimal treatment	Ca/P	Crystallinity	Particle size & morphology	Reference
	Black tilapia (<i>Oreochromis placidus</i>)	(1) Immersed in 0.1 M HCl for 12 h (2) Stirred in 5% NaOH at 70 °C for 5 h (3) Treated with 50% NaOH at 100 °C for 1 h	1.75	Low	Size: 50–60 mm wide, 30–200 nm long Morphology: irregular rod shape and crystal size	Selimin ¹⁹¹
	Sardine (<i>Sardinella longiceps</i>)	(1) Treated with 1 N HCl (2) Treated with 1 N NaOH (3) Calcined at 600–1000 °C	n.s. ^a	High	Size: <1 µm Morphology: spherical agglomerates	Ashwitha ¹⁹²
	n.s. ^a	(1) Soaked in 40% NaOH solution (2) Ground (3) Calcined at 800 °C for 2 h	1.18	High	Size: <5 µm Morphology: irregular and uneven-sized blocks	Wang ¹⁹³
Calcination	Nile tilapia (<i>Oreochromis niloticus</i>)	(1) Calcined at 450–600 °C for 5 h (2) Collected on a 0.5-mm sieve	3.06–3.17	n.s. ^a	Size: very small	Sittit ¹⁹⁴
UAE ^a	Pirarucu (<i>Arapaima gigas</i>)	(1) Calcined at 700 °C for 2 h (2) Sieved (325 mesh) (3) UAE ^a for 30 min (4) Matured for 24 h	1.58	^c 48%	Size: n.s. ^a Morphology: n.s. ^a	de Amorim ¹⁹⁵
	Nile tilapia (<i>Oreochromis niloticus</i>)	(1) UAE ^a with 0.8 M HCl 45 min at 60 °C (2) Added NaOH to increase pH to 12 (3) Sonicated for 30 min (4) Ground with a mortar and pestle	1.68	n.s. ^a	Size: 22.8 nm long and 8.6 nm wide Morphology: rice-shaped agglomerates	Sricharoen ¹⁹⁶
DES ^a	Bighead carp (<i>Aristichthys nobilis</i>)	(1) Ground and sieved (80 mesh) (2) Treated with DES ^a (choline chloride : glycerol = 1 : 2) at 70 °C for 2.5 h (3) Centrifuged for 7 min (4) Purified by stirring with 5% NaOH for 5 h at 70 °C	1.73	^c 43.1%	Size: 0.3–95 µm (median diameter 13.98 µm) Morphology: irregular and agglomerated morphology	Liu ¹⁹⁷
	Crucian carp (<i>Carassius carassius</i>)	(1) Ground and sieved (80 mesh) (2) Treated with DES ^a (choline chloride : 1,4-butanediol = 1 : 15) at 65 °C for 2 h (3) Centrifuged (4) Purified by stirring with 5% NaOH for 5 h at 70 °C	1.78	Low	Size: 0.235–117.1 µm (median diameter: 12.71 µm) Morphology: irregular and agglomerated morphology	Liu ¹⁹⁸

^a Abbreviations – AAD, alkaline and/or acid deproteinization; UAE, ultrasound assisted extraction; DES, deep eutectic solvent; BCP, biphasic calcium phosphate; n.s., none specified. ^b Crystallinity calculated by comparing the area of crystalline peaks to the total area of amorphous and crystalline peaks (eqn (S3)). ^c Crystallinity calculated using software.

Other researchers have used AAD as a pre-treatment before calcination. Compared to fish bones that are often directly calcined without the presence of acid or base, it is much more common to deproteinize fish scales prior to thermal treatment. These methods often use dilute base rather than concentrated base because calcination will ultimately remove a significant quantity of residual protein. Like fish bones, HAP particles from scales after calcination are larger than those not exposed to high temperatures.^{182,188}

3.2.3. Ultrasound assisted extraction (UAE). UAE is an extraction technique unique for fish scales that has not been extensively explored for bones yet. Note, Boudreau *et al.* have transformed fish bones to nanoparticles with UAE,¹⁷⁵ however this was after using solely enzymes to isolate HAP from bones.¹⁷⁴ UAE has been used to isolate compounds from other types of waste, specifically plant-based biomass, such as wheatgrass,²⁰¹ sugarcane bagasse,²⁰² coffee waste,²⁰³ wheat straw,²⁰⁴ tobacco waste,²⁰⁵ and wood waste.²⁰⁶ Extraction occurs when ultrasound



Table 9 Biomaterials prepared from the calcination and/or alkaline hydrolysis of fish bones/scales, published since 2019^c

Material	Preparation	Characteristics	Reference
HAP ^a nanoparticles	(1) Alkaline hydrolysis (silver carp bones)	<ul style="list-style-type: none"> Retains organic moieties to assist in better osseointegration Ca/P: 1.65 Haemolytic activity 2.24^b–3.35% at 1–5 mg mL⁻¹ 91% cell viability at 500 μL g⁻¹ Significant cell proliferation of 20% at 250 μg mL⁻¹ 	Acharya ²¹⁶
BCP ^a nanoparticles	(1) Calcination (silver carp bones)	<ul style="list-style-type: none"> Removal of organic content and more crystalline Ca/P: 1.45 Haemolytic activity 3.94^b–9.22% at 1–5 mg mL⁻¹ 86% cell viability at 500 μL g⁻¹ Cell proliferation of 15% at 250 μg mL⁻¹ 	Acharya ²¹⁶
HDPE ^a -BCP ^a composite	(1) Calcination (tilapia scales) (2) Spray-dried (3) 30 wt% HAP ^a mixed with HDPE ^a in extruder (4) Surface treated with MPTMS ^a	<ul style="list-style-type: none"> Density: 1.17 g cm⁻³ Melting point of 138.4 °C Tensile strength: 28.26 MPa Young's modulus: 1272 MPa Elongation at break: 43.6% Flexural strength: 21.4 MPa Flexural modulus: 796 MPa Impact strength: 46.90 kJ m⁻² 98.51% cell viability at 200 mg mL⁻¹ 	Aiza Jaafar ¹⁸³
BCP ^a bioceramics	(1) Calcination (salmon bones) (2) Compacted and sintered	<ul style="list-style-type: none"> Density: 2.96 g cm⁻³ Elastic modulus: 633 MPa Ca/P: 1.67 No cytotoxic effect compared to the control Suitable cytocompatibility No cell proliferation after 3 days 	Bas ¹⁷²
HAP ^a particles	(1) Alkaline hydrolysis (arowana scales)	<ul style="list-style-type: none"> Forms a bone-like apatite layer after immersed in McCoy medium for 3 days, characterizing its bioactivity 	Horta ¹⁸²
BCP ^a bioceramics	(2) Calcination (1) Calcination (tilapia bones) (2) Compacted and sintered	<ul style="list-style-type: none"> 102.8% cell viability after 48 h Ca/P: 1.64 Relative density: 90.92–96.43% Vickers hardness: 5.77 GPa Compressive stress: 89.16 MPa Young's modulus: 13.88 GPa 	Khamkongkaeo ¹³²
HAP ^a scaffold	(1) Calcination (tuna bones) (2) Mixed with 2% starch, compacted, and sintered	<ul style="list-style-type: none"> Ca/P: 1.87 Nontoxic behavior at concentrations <300 μg mL⁻¹ Enhanced cell proliferation over control 	Mondal ¹³¹
PLA ^a -HAP ^a filament	(1) Alkaline hydrolysis (sea bass scales) (2) Mixed with PLA ^a and eggshells (CaO) in extruder (3) Pressed into membranes	<ul style="list-style-type: none"> Ca/P: 1.67 Young's modulus: 3.52–3.95 GPa Tensile strength: 43.1–50.9 MPa Elongation at failure: 2.8–4.8% Decreased water resistance Cell viability was not different from control group at 18 days Significantly improved cell viability compared to PLA at 2 days Enhanced cell adhesion Good cytocompatibility Scavenging rate: 5–30% Generated inhibition zones for <i>E. coli</i> and <i>S. aureus</i> 	Wu ¹⁸⁹



Table 9 (Contd.)

Material	Preparation	Characteristics	Reference
HAP ^a nanoparticles	(1) Alkaline hydrolysis (Indian oil sardine bones)	<ul style="list-style-type: none"> • Cell proliferation 141% at 100 µg mL⁻¹ • Loss of contact with neighbouring cells at 250 µg mL⁻¹, suggesting higher concentrations cannot support osteoblastic growth • Good reposition of calcium in MG-63 cells at 7 days, which may be important for bone mineral density • Appropriate for cellular activities 	Surya ¹⁶³
HAP ^a -CMC ^a /SA ^a coating on Ti ₆ Al ₄ V alloy	(1) Calcination (rohu bones) (2) HAP ^a added to CMC ^a in ethanol/water for 12 h (3) SA ^a added and pH adjusted to 7 (4) Ti ₆ Al ₄ V added by electrophoretic deposition	<ul style="list-style-type: none"> • Induces growth of apatite on composite when immersed in SBF^a • Biomineralization evident • Microhardness value: 190 Hv • Enhanced antibacterial activity against <i>S. aureus</i> • Lower antibacterial resistance against <i>E. coli</i> • Cell viability up to 87% • No significant toxicity • Ca/P of targets: 1.57–1.63 	Sridevi ¹⁵⁷
BCP ^a coating	(1) Alkaline hydrolysis (sea bream & salmon bones) (2) Calcination (3) Pressed and sintered (4) Coatings prepared by PLD ^a	<ul style="list-style-type: none"> • Ca/P of films: 1.47–1.50 • Pull-off bonding strength sea bream: 49 MPa • Pull-off bonding strength salmon: 33 MPa • Mass gain of salmon after 7 days in DMEM^a: 3% • Mass gain of sea bream after 7 days in DMEM^a: 9% • Good biocompatibility • LDH^a release salmon after 24 h: 97% of control • LDH^a release sea bream after 24 h: 94% of control • Significantly increased anti-biofilm performances 	Popescu-Pelin ¹⁴²

^a Abbreviations – HAP, hydroxyapatite; BCP, biphasic calcium phosphates; HDPE, high-density polyethylene; MPTMS, 3-methacryloxypropyltrimethoxysilane; CMC, carboxymethylcellulose; SA, sodium alginate; SBF, simulated body fluid; PLD, pulsed laser deposition; DMEM, Dulbecco's Modified Eagle medium; LDH, lactate dehydrogenase. ^b Haemolytic activity <5% is labelled as non-haemolytic (non-toxic); haemolytic activity >5% is labelled as haemolytic (toxic). ^c Cell viability must decrease >30% to be considered cytotoxic.

(US) waves generate microbubbles containing gas and vapour within a liquid medium.²⁰⁷ When the microbubbles collapse, local pressures can reach 2000 atm and temperatures increase rapidly, a phenomenon known as cavitation.²⁰⁷ This process further dissolves and disintegrates solid materials into smaller particles.²⁰⁷

Compared to calcination and AAD, UAE remains relatively uncommon. In 2020, Sricharoen *et al.* applied UAE for the extraction of HAP from fish scales.¹⁹⁶ They performed UAE on Nile tilapia scales by immersing them in 0.2–1.2 M HCl and using ultrasonic power of 0.1–0.4 kW at 30–60 °C for 15–90 min. Based on results from ICP-AES, calcium and phosphorus extraction was optimized using 0.8 M HCl and 0.4 kW of

ultrasonic power at 60 °C for 45 min. Furthermore, the pH of the solution was adjusted between 8 and 12 with NaOH, followed by sonication at 0.4 kW at room temperature for 30 min. The material was then dried, and ground with a mortar and pestle. As the pH increased, the Ca/P ratio also increased from 1.44 to 1.68 which was considered ideal since it was closest to the stoichiometric ratio of HAP. Furthermore, HAP prepared at pH 12 was the most crystalline, and rice-shaped nanoparticles (22.8 nm long and 8.6 nm in diameter) were observed while particles synthesized at lower pH had more irregular and overlapping shapes.

In the same year, Chen and co-workers also isolated HAP from Mozambique tilapia (*Oreochromis mossambicus*) scales



using UAE; however they also used calcination and AAD as additional processing steps.²⁰⁸ The scales were first immersed in 1 N HCl for 24 h, 1 N NaOH for 24 h, boiled at 80 °C in water for 20 min, and dried. Next, the sample was sonicated in 50% (v/v) alcohol and calcined at 1000 °C before being ground with a mortar and pestle, and passed through a 200-mesh sieve. Without calcination, the scale powder showed broad peaks in the observed XRD pattern while the calcined product matched the diffractogram of a HAP standard. The mean diameter of particles was 5.96 μm, ranging from 4.2 to 7.8 μm, values that are significantly larger than those reported by Sricharoen *et al.*¹⁹⁶ These values are also higher than those observed after their initial extraction technique (not incorporating UAE) where sizes ranged from 500 nm–1.5 μm.²⁰⁸ This suggests that UAE might provide enough energy to grow HAP particles, similar to calcination, as similar results were observed during Boudreau *et al.*'s treatment on bones – larger particle sizes after prolonged US treatment.¹⁷⁵

That being said, the opposite effect has also been observed. For example, de Amorim *et al.* also used calcination with UAE, and the heat treatment was performed before sonication.¹⁹⁵ Scales from pirarucu (*Arapaima gigas*) were calcined at 700 °C for 2 h and sieved through a 325-mesh sieve. While a portion of this was doped with niobium ions during UAE, some scales were not in order to see whether doping was successful. These scales were sonicated in ethanol at 20 kHz for 30 min with an amplitude of 70% and dried prior to analysis. Unlike Chen and co-workers,²⁰⁸ the crystallite size decreased from 128.5 to 52.3 nm from the added UAE treatment after calcination. Furthermore, the crystallinity increased from 34 to 48%, suggesting that UAE successfully removed residual protein remaining after calcining for 2 h.

3.2.4. Deep eutectic solvents (DESs). While extraction with DESs remains the most under-researched topic discussed in this review, it is still important to mention because of its considered greenness. DESs are prepared by mixing two components that are typically solid at room temperature, but upon heating the components react to form the desired DES that remains liquid at room temperature. DESs are achieved by complexation between a hydrogen bond acceptor (HBA), typically a quaternary ammonium salt or halide salt, and a hydrogen bond donor (HBD).²⁰⁹ The melting point of the mixture is lower than the melting points of individual HBA and HBD. DESs have gained significant attention among green chemists because they are typically biodegradable, nontoxic, and inexpensive with low volatility and low vapor pressure.²¹⁰ It should be highlighted, however, that not all aspects of DESs are sustainable as the HBD and/or HBA are not always green, cheap, or nontoxic.²¹¹

Liu *et al.* have investigated different DESs for the extraction of HAP from scales of bighead carp (*Aristichthys nobilis*)¹⁹⁷ in 2020 and crucian carp (*Carassius carassius*) in 2021.¹⁹⁸ In both cases, the scales were ground and passed through a 80 mesh sieve before being treated with DES.^{197,198} For their initial study on bighead carp, three DESs were investigated – choline chloride:glycerol (1:2), choline chloride:citric acid (2:1), and choline chloride:acetic acid (1:2).¹⁹⁷ The DES that was selected

as the best extraction medium was choline chloride:glycerol (1:2) because it had high HAP solubility, moderate viscosity, and excellent stability, and it did not decompose HAP. An optimal extraction rate of 47.67% was achieved by treating scales at 70 °C in DES for 2.5 h with a solid:liquid ratio of 1:15 g g⁻¹. The particles were observed to have a high median particle size of 13.98 μm and tended to agglomerate, as observed in all other methods described. Because this process does not involve high temperatures, the product was amorphous and the relative crystallinity of the extracted HAP was 43.13%. Liu *et al.* screened other DESs for HAP extraction from crucian carp scales, including choline chloride:glycerol (1:2), choline chloride:triethylene glycol (1:4), choline chloride:glycol (1:2), and choline chloride:1,4-butanediol (1:2).¹⁹⁸ In this case, choline chloride:1,4-butanediol (1:2) was chosen since it had the best extraction rate of 40.58%. This is interesting because this extraction rate is lower than that in their initial study where choline chloride:glycerol (1:2) was the best.¹⁹⁷ The crystallinity and median particle size diameter (12.71 μm) of crucian carp were similar to those of bighead carp.^{197,198}

3.3. Applications of HAP and BCPs

3.3.1. Biomedical potential. There have been promising results using biologically sourced BCPs because of their biomimetic properties.²¹² For example, incinerated fish bones have been studied for enamel remineralization and occlusion of dentin tubules.²¹³ Several studies have prepared HAP and/or BCPs for biomedical applications by calcining fish bones at 600–1200 °C.^{172,176,214,215} One study compared the cell attachment and proliferation of two scaffolds created with synthetic HAP and tuna bones, respectively.¹³¹ Both scaffolds were nontoxic and, despite having a Ca/P ratio of 1.87, the scaffold created with fish bones demonstrated enhanced cell proliferation and attachment. This is believed to be caused by the presence of trace elements such as Na, Mg, Sr, and Co, which better mimic human bones. Acharya *et al.* prepared nano-BCP particles from silver carp (*Hypophthalmichthys molitrix*) bones using alkaline hydrolysis and calcination separately to study the effect of extraction technique on proliferation and toxicity.²¹⁶ The particles created by alkaline hydrolysis were observed to be made of HAP and had better biocompatibility than those obtained by calcination which yielded a product containing α-TCP and β-TCP. The MG63 osteoblast cell lines had a 70.1% proliferation efficiency and 91% cell viability in cytotoxicity studies. Another variable that has been demonstrated to play a significant role in the potential of bio-derived BCPs as a bone replacement material is the species of fish from which they originate.²¹⁷ The cell proliferation and differentiation of synthetic HAP and nanoparticles from calcined rainbow trout (*Onchornuchus mkiss*), cod (*Gadus*), and salmon (*Oncorhynchus keta*) bones were studied by Shi *et al.*²¹⁷ All three samples originating from fish had superior cell viability than the synthetic, commercially sourced HAP. Salmon and rainbow trout bones significantly enhanced the viabilities which was explained by the increased substitution of CO₃²⁻ in these samples.



Ultimately salmon bones were reported to be the most suitable material for bone regeneration because it stimulates cell proliferation due to its high CO_3^{2-} and Mg content.

HAP from fish bones and scales has also been shown to have enough mechanical strength for biomedical ceramics.¹⁶² In fact, Akpan *et al.*¹⁶² observed that BCPs from bones experienced a Vickers hardness value of 0.48 GPA which is within the range reported for human femoral cortical bone. The particles also had a fracture toughness of 5.72 MPa $\text{m}^{1/2}$ while maintaining a stable phase at 900 °C and a low brittleness index value of 0.084. Aiza Jaafar *et al.* created a high-density polyethylene (HDPE) composite incorporating BCPs obtained from the thermal degradation of fish scales at 1200 °C.¹⁸³ The size of resulting BCP particles was tailored by using spray drying and some were additionally treated with 3-methacryloxypropyltrimethoxysilane (MPTMS, $\text{C}_{10}\text{H}_{20}\text{O}_5\text{Si}$) to study its effect on the mechanical properties of composites. Young's modulus of HDPE was enhanced by 29.4% by adding 30 wt% HAP treated with MPTMS to the composite, reaching 1272 MPa, with a tensile strength of 28.26 MPa and a flexural modulus of 796 MPa. Furthermore the composite was shown by *in vitro* analyses to be non-toxic with potential to be used towards biomedical applications.

BCP materials derived from fish bones have been explored in other biomedical applications. For example, there have been several reports using BCPs from fish bones in sunscreen as an alternative to UV filters associated with health risks.²¹⁸

Adamiano *et al.* isolated BCPs from Atlantic salmon bones by calcining them at 800 °C and doped this material with Zn or Mn to increase sun protection factor (SPF)-boosting abilities.²¹⁹ Interestingly, the undoped BCP material was the most effective at increasing SPF. This demonstrates that bio-derived BCPs can be used to decrease the concentration of UV filters needed in sunscreens while maintaining high SPF values.

Table 9 lists several other example studies using BCPs sourced from fish bones and scales for biomedical purposes. Based on these reports, there is great potential for use of waste-derived HAP in bioceramics and implants, contributing towards a more sustainable industry and circular economy.

3.3.2. Environmental remediation. HAP has been studied extensively as a material for water and soil remediation because of its acid-base properties, ion-exchange capability, thermal stability, and non-toxic nature.¹²⁶ HAP's lattice is flexible and tolerant of substitutions, which explains why its carbonated version is found readily in biological systems. Cation exchange occurs through substitution of Ca^{2+} with Cu^{2+} , Mn^{2+} , Ni^{2+} , Zn^{2+} , Cd^{2+} , Co^{2+} , Mg^{2+} , Sr^{2+} , Ba^{2+} , Pb^{2+} , Al^{3+} or La^{3+} . Anion exchange can occur at OH^- and/or PO_4^{3-} sites. The OH^- has been substituted with F^- , Cl^- , Br^- , O^{2-} , and CO_3^{2-} while PO_4^{3-} has been exchanged with HPO_4^{2-} , AsO_4^{3-} , VO_4^{3-} , SO_4^{2-} , SiO_4^{4-} , and CO_3^{2-} .

Several industrial dyes have been removed from aqueous solutions using materials created with fish-derived HAP. For example, Mamun *et al.* recently prepared a photocatalyst from

Table 10 Industrial dyes treated with BCPs from fish bones/scales since 2019^a

Preparation	Dye	Optimal conditions	Degradation/adsorption	Reference
Calcined tilapia & surma bones	Congo red	240 min, ^b 20 ppm, ^c 40 mL, ^d 0.1 ^e g	65%	Mamun ¹⁴⁸
Pulverized silver carp bones	Congo red	240 min, ^b 200–300 ppm, ^c 100 mL, ^d 100–700 mg, ^e pH 2 ^f	91–97%	Parvin ²²⁰
Calcined shing and poa bones	Congo red	180 min, ^b 10 ppm, ^c 0.08 ^e mg	73–82%	Prosad Moulick ¹⁴⁷
Pulverized fish bones treated with Al_2O_3	Methyl green	60 min, ^b 80 mg L^{-1} , ^c 0.1 g, ^e pH 10.64, ^f 25 ^g °C	92%	Al-Kazragi ²²²
Calcined of catla bones	Congo red & crystal violet	75 min, ^b 50 mg L^{-1} , ^c 10 mg ^e	Congo red: 87% Crystal violet: 77%	Sathiyavimal ¹⁴⁶
Pulverized fish bones doped with copper and calcined	Crystal violet	35 min, ^b 20 mg L^{-1} , ^c 2 g L^{-1} , ^e pH 10, ^f 50 ^g °C	98%	Mejbar ²²³
Bleached and calcined fish bones	Brilliant green	20 min, ^b 50 mg L^{-1} , ^c 1 g L^{-1} , ^e pH 12 ^f	49.1 mg g^{-1}	Miyah ²²⁴
Calcined fish bones	Methylene blue	10 min, ^b 100 mg L^{-1} , ^c 250 mg, ^e pH 6.9 ^f , 30 ^g °C	>90%, 56.49 mg g^{-1}	Nurhadi ²²⁵
Calcined catfish bones mixed with chitosan	Methylene blue & methylene orange	Methylene blue: 60 min, ^b 35 mg L^{-1} , ^c 100 mL, ^d 0.1 g, ^e pH 8; ^f methylene orange: 180 min, ^b 35 mg L^{-1} , ^c 100 mL, ^d 0.1 g, ^e pH 6 ^f	Methylene blue: 84.89 mg g^{-1} Methylene orange: 44.05 mg g^{-1}	Trung ²²⁶
Rohu bones treated with NaOH	Methylene blue	12 min, ^b 50 mg L^{-1} , ^c pH 5, ^f 34.85 ^g °C	96.1%, 666.67 mg g^{-1}	Swamiappan ²²⁷
Rohu bones treated with NaOH	Melioderm HF brown G	120 min, ^b 200 ppm, ^c 2 g L^{-1} , ^e pH 2, ^f 24.85 ^g °C	98.33%	Hossain ²²⁸

^a Abbreviations – HAP, hydroxyapatite. ^b Reaction time for optimal dye degradation/adsorption. ^c Dye concentration for optimal dye degradation/adsorption. ^d Volume of dye solution for optimal dye degradation/adsorption. ^e Adsorbent mass/concentration for optimal dye degradation/adsorption. ^f pH for optimal dye degradation/adsorption. ^g Temperature for optimal dye degradation/adsorption.



Table 11 Heavy metals treated with BCPs^a from fish bones/scales since 2020

BCP source	Additives	Metal(s)	Optimal conditions	Removal rate/adsorption	Reference
Fish bones	Chitosan, Fe ₃ O ₄	Cd ²⁺	3 min, ^b 200 mg L ⁻¹ , ^c 0.05 g, ^d pH 5.4, ^e 25 ^f °C	25.134 mg g ⁻¹	Yang ²³¹
Fish bones	NA ^a	Pb ²⁺	672 h, ^b 10 ^d wt%	94–96%	Nag ²³⁵
Fish scales	Poly(lactic acid)	Pb ²⁺ , Cd ²⁺	12 h, ^b 5–100 mg mL ⁻¹ , ^c 15 ^d wt%	Pb ²⁺ : 112.6 mg g ⁻¹ Cd ²⁺ : 360.5 mg g ⁻¹	Fijol ²³⁶
Fish bones	NA ^a	Pb ²⁺ , Zn ²⁺	672 h ^b	Pb ²⁺ : 86.39% Zn ²⁺ : 63%	Saffarzadeh ²³⁷
Fish bones	NA ^a	Co ²⁺	400 min, ^b 10–1000 mg L ⁻¹ , ^c 0.001 g mL ^{-1d}	52 mg g ⁻¹	Renda ¹⁴⁴
Fish bones	NA ^a	As ³⁺	40 min ^b	1.4 mg g ⁻¹	Hubadillah ¹⁵³
Fish scales	MgCl ₂	Cu ²⁺ , Cd ²⁺ , Pb ²⁺	180 min, ^b 20–800 mg L ⁻¹ , ^c 1 mg mL ⁻¹ , ^d pH 3–7 ^e	Cu ²⁺ : 84.2% Cd ²⁺ : 74.2% Pb ²⁺ : 53.7%	Qi ²³²
Fish bones	Graphene oxide, chitosan	Cu ²⁺	120 min, ^b 100–1000 mg L ⁻¹ , ^c 1 mg mL ⁻¹ , ^d pH 5 ^e	256.41 mg g ⁻¹	Hoa ²³⁸

^a Abbreviations – BCP, biphasic calcium phosphate; HAP, hydroxyapatite; NA, not applicable. ^b Reaction time. ^c Heavy metal concentration.

^d Adsorbent mass/concentration. ^e pH for optimal dye degradation/adsorption. ^f Temperature for optimal degradation/adsorption.

calcined tilapia and surma bones for the degradation of Congo red dye,¹⁴⁸ an environmental pollutant. In a similar study, pama croaker and Asian stinging catfish were used to create the same photocatalyst, achieving degradation rates of 82% and 73%, respectively, towards Congo red.¹⁴⁷ That being said, much higher degradation rates of up to 97% were reported by Biswas and co-workers using raw, pulverized silver carp bones.²²⁰ This is interesting because the product contains significantly higher quantities of organic impurities compared to other photocatalysts due to the lack of heat or alkaline treatment. Congo red has anionic sites from sulfonate groups, and other anionic dyes have been studied (*e.g.*, acid blue 185),²²¹ while neutral and cationic dyes have also been successfully treated with fish bones such as methyl green,²²² crystal violet,^{146,223} brilliant green,²²⁴ and methylene blue.²²⁵ (Table 10).

Other pollutants besides dyes have been treated with HAP derived from fish discards. Synthetic HAP has been studied widely for the removal of a broad range of heavy metals.^{229,230} There have been several studies using bio-sourced HAP to remediate Cd²⁺,^{231,232} Pb²⁺,^{232–237} Co²⁺,¹⁴⁴ As³⁺,¹⁵³ Zn²⁺,²³⁷ and Cu²⁺.^{232,238} (Table 11). For example, Cd²⁺ and Pb²⁺ have been successfully adsorbed from water by Rashed *et al.* using calcined fish bones.²³⁹ A 99% removal rate of Cd²⁺ and Pb²⁺ was achieved by treating wastewater with 0.1 g HAP for 30 min at ~55 °C with a metal concentration of 10 ppm.²³⁹ Renda *et al.* discovered that HAP could also be used for the adsorption and desorption of Co²⁺ ions from aqueous solutions.¹⁴⁴ Co²⁺ is a controversial heavy metal because of its effect on human health, but it is also necessary in small quantities for certain bodily functions (*e.g.*, vitamin B₁₂, essential coenzyme for cell mitosis, metabolism, and N₂ fixation) and industrial applications (*e.g.*, mining, electronics, and electroplating). The desorbed Co²⁺ was then observed to have beneficial effects towards seed germination and root elongation.

4. Conclusions

There has been a significant increase in fish production over the last few decades to meet the demands of a growing human population. This has led to an abundance of seafood processing byproducts being disposed of unsustainably, contributing to greenhouse gas emissions, eutrophication, and ocean acidification. There have been several global examples that demonstrate that it is possible to process each portion of fish to higher value products instead of just focusing on the edible portion as being of sole value. Iceland's 100% Fish Project has created many different products from fish including fish leather. Norway has also been making great progress towards whole fish utilization. However, these examples face regulatory challenges.^{22–24} While successes have been achieved with new products based on marine collagen and materials from fish

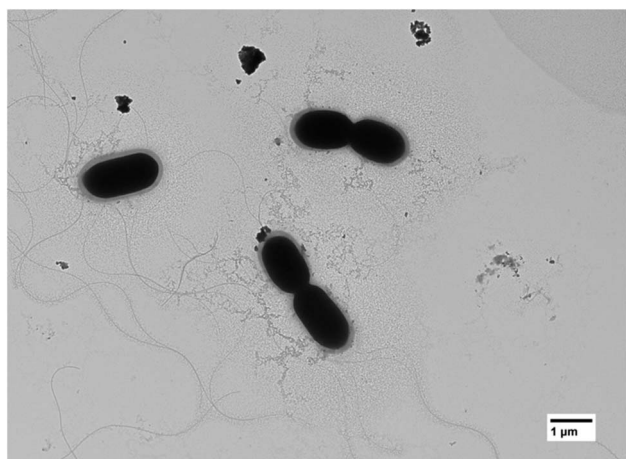


Fig. 6 TEM image obtained when HAP nanoparticles prepared in water were allowed to sit at room temperature for several days (SI). The black cylindrical particles approximately 1 μm in diameter are attributed to unspecified bacteria.



skin, there are, as far as we are aware, no commercial processes focused on the inorganic components present in seafood waste.

In this review, we have summarized the possibilities of isolating valuable biominerals from fish inedible discards. In general, it is more advantageous to source calcium-based materials naturally instead of producing them synthetically for sustainability and applicability purposes. Valorizing by-products from the seafood processing industry offers additional streams for financial gain, and prevents significant food waste from ending up in landfills and/or the ocean. This would limit GHG emissions, eutrophication, groundwater contaminations, and odor nuisances caused by the wasted discards. Specifically, this topic tackles many of the UN SDGs, including UN SDG2: Zero Hunger, UN SDG3: Good Health & Well-Being, UN SDG8: Decent Work and Economic Growth, UN SDG9: Industry, Innovation and Infrastructure, UN SDG11: Sustainable Cities and Communities, UN SDG12: Responsible Consumption and Production, UN SDG13: Climate Action, and UN SDG14: Life Below Water. While synthetic CaCO_3 and HAP continue to be used more prominently than biogenic materials for most applications, the biomimetic properties of natural HAP and CaCO_3 could make it more advantageous for biomedical purposes (e.g., enhanced biomineralization).

Shells from bivalves, crustaceans, and gastropods have the potential to be used as a feedstock for CaCO_3 which can then be converted to other value-added products, including CaO and PCC. An interesting area of research that could be investigated is the tailored morphology of polymorphs. While specific polymorphs have been achieved through PCC processes, calcite itself has a range of morphologies that could be explored for biomedical purposes. Khanjani *et al.* recently explored the factors that influence the morphology of microbially induced PCC, discovering that Ca^{2+} and functional groups present have an impact on the resulting polymorph and shape.²⁴⁰ While this study primarily focused on vaterite and calcite's spherical shape, it would be interesting to study other possible morphologies.

HAP has been isolated from fish bones and scales using a range of techniques including calcination, alkaline/acidic deproteinization, enzymatic hydrolysis, ultrasound assisted extraction, and the use of deep eutectic solvents. This bio-derived HAP product has been explored for various applications such as incorporation into biomedical composites and the treatment of heavy metals and industrial dyes. These processes have the potential to mitigate the large number of byproducts wasted in landfills and the ocean, although there are some drawbacks. While some reports have focused specifically on making their treatment sustainable and scalable (e.g., enzymes), many of the processes described are industrially inapplicable in seafood processing plants, requiring high temperatures for calcination and the use of hazardous chemicals such as concentrated base (e.g., KOH 50 wt%) for deproteinization. There are also inconsistencies in particle size and morphology of products originating from the same species of fish. One of the main drawbacks of studying biomass-derived products is the development of bacteria, especially in samples that contain trace amounts of residual protein. In fact, we have

observed bacteria in TEM images of some of our own HAP nanoparticle solutions from fish bone that were left at room temperature (Fig. 6). For future studies, it would be interesting to develop a low temperature method without the use of acids or bases that also prevents the growth of potentially harmful bacteria. This would ensure that CaCO_3 and HAP can be produced sustainably from seafood discards while also being suitable for biomedical purposes. Reduction in endotoxin contamination has been important in advancing the use of organic products such as chitosan from seafood waste streams.²⁴¹ As the global seafood industry continues to shift from wild catch to aquaculture practices, several threats have emerged. Some of these include the spread of diseases between wild fish and farmed fish and the risk of rising ocean temperatures from climate change affecting harvested species. By completely valorizing every aspect of caught marine organisms, industries would be able to continue making a profit despite challenges with the edible portion. Therefore, the fish itself would remain useful and valuable even if not being considered seafood. For food production, it is important to adapt to climate change and consider indoor recirculating aquaculture systems instead of open ocean systems to mitigate the variations in water temperature and composition.

Author contributions

Conceptualization, F. M. K. and S. B.; writing – original draft preparation, S. B.; writing – review and editing, S. B., E. L., and F. M. K.; supervision, E. L. and F. M. K.; funding acquisition, E. L. and F. M. K.

Conflicts of interest

There are no conflicts to declare.

Data availability

There is limited supporting data for this review article. The instrumental conditions for Fig. 6 are described in Boudreau *et al.*'s article in *RSC Sustainability* (2025) and in the supplementary information file (SI) for this review. Supplementary information is available. See DOI: <https://doi.org/10.1039/d5su00527b>.

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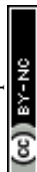
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