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Introduction

The global population requires sustainable and innovative protein sources as the demand for food protein is estimated to double by 2050 with the growing population. Insects are recognized, among all alternative sources, to have the potential to fulfill the huge demands for protein in the future. Insects are widely consumed; it is estimated that 1900 insect species are eaten by different groups of the population from around 80 countries in Asia, Africa and America. The most consumed and well-researched insects in recent years are *A. domesticus*, *T. molitor* and *A. assamensi* pupae. Insects are nutritious; the average protein content in insects is around 40% and for some species like *A. domesticus* and *T. molitor* larvae it is even higher.^{1–3} The essential amino acid requirement recommended by the FAO/WHO is also covered by insect pro-

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Current state of insect proteins: extraction technologies, bioactive peptides and allergenicity of edible insect proteins

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This review aims to provide an updated overview of edible insect proteins and the bioactivity of insectderived peptides. The essential amino acid content of edible insects is compared with well-known protein sources to demonstrate that edible insects have the potential to cover the protein quality requirements for different groups of the population. Then the current methodologies for insect protein extraction are summarized including a comparison of the protein extraction yield and the final protein content of the resulting products for each method. Furthermore, in order to improve our understanding of insect proteins, their functional properties (such as solubility, foaming capacity, emulsifying, gelation, water holding capacity and oil holding capacity) are discussed. Bioactive peptides can be released according to various enzymatic hydrolysis protocols. In this context, the bioactive properties of insect peptides (antihypertensive, antidiabetic, antioxidant and anti-inflammatory properties) have been discussed. However, the allergens present in insect proteins are still a major concern and an unsolved issue for insect-based product consumption; thus, an analysis of cross reactivity and the different methods available to reduce allergenicity are proposed. Diverse studies of insect protein hydrolysates/peptides have been ultimately promoting the utilization of insect proteins for future perspectives and the emerging processing technologies to enhance the wider utilization of insect proteins for different purposes.

> teins.⁴ Moreover, the vitamin, mineral (calcium, iron, and zinc) and monounsaturated and polyunsaturated fatty acid contents are high in edible insects.⁵ Insect protein was reported to be more digestible (usually between 76% to 98%) than plant proteins like peanuts and lentils (usually around 52%), but slightly less digestible than animal protein like beef (89%) and egg white (100%).⁶ Furthermore, the energy content of edible insects varies from 1821 to 1896 kJ per 100 g which is comparable to most meats except for pork (67 750 kJ per 100 g).⁵ Other notable advantages compared to existing protein sources are related to positive environmental impacts such as less water use, less land occupied, and fewer gas emissions.⁷ Besides these benefits, sensory quality and acceptance are other important aspects that should be considered when evaluating alternative proteins. Although the willingness to consume insects is still an ongoing issue, earlier studies showed that a promising option was to use yellow mealworm flour as a supplement to fortify tortillas, muffins and beef/ mealworm burgers, which were found to be acceptable by consumers.8-11 As a result of ongoing efforts, insects have received more attention in recent years.

> Compared to whole insects, insects as ingredients might be an easier alternative for people to consume if in an unrecog-

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nizable form. Del Valle *et al.* (1982) further indicated that insect protein extraction for future uses in the food industry and novel foods was extremely relevant for countries that did not consume insects, such as North American and European countries.¹² As for insect protein extraction, the most commonly used protein extraction method is alkaline solubilization coupled to isoelectric precipitation, despite some publications also applying alkaline solubilization or acid solubilization separately. Other novel methods like dry fractionation and ultrasound-assisted extraction are gradually being researched.^{13,14}

The bioactive properties of insect proteins can be improved through enzymatic hydrolysis either by in vivo or in vitro means, as these peptides within the sequence of the parent proteins are inactive.^{15,16} Currently, the widely demonstrated bioactivities of insect-derived peptides have been antihypertensive, antidiabetic and antioxidant properties. In recent years, numerous bioactive peptides were isolated from a wide variety of food proteins; significant ACE inhibitory activity has been observed in some common protein sources like egg white, rice, sweet potato, and cod.¹⁷⁻¹⁹ Similarly, ACE inhibitory activity has also been assessed in some insect species, such as B. mori, S. littoralis, S. gregaria, B. terrestris, A. assamensis pupae and G. sigillatus.^{20,21} As for diabetes, the WHO estimates that the number of people with diabetes will double by 2030; type 2 diabetes is the main contributor to cardiometabolic diseases.²² Insect proteins have presented antidiabetic properties, for example, whole G. sigillatus showed increased bioactivity by both simulated gastrointestinal digestion and enzymatic hydrolysis by food-grade enzymes such as alcalase.²³ Other bioactivities such as anti-inflammatory activities are gradually studied in insects. But to date, no in vivo study has been performed on the antidiabetic and antioxidant properties of insect-derived peptides.

Furthermore, the new antigens and potential allergens that are released at the same time as bioactive peptides should also be taken into careful consideration. The allergenicity of insects can be caused in several ways including 'sting, inhalation, direct contact and ingestion'. Cross reactivity with allergens in crustaceans and house dust mites can also be triggered after insect ingestion.²⁴ Among the potential allergens found in insects, tropomyosin has been identified as a major one.^{25,26} Another allergen found in insects is arginine kinase. However, studies regarding the cross reactivity of arginine kinase are still limited. Some publications reported that hydrolysis under the assistance of microwave could reduce the immunoreactivity of insect protein. A possible explanation is that the compounds present may have some thermal and non-thermal interactions with microwave treatment, which is not the case for convection heating.^{21,27} Similar results were observed in other food products like dairy whey protein hydrolysates and fish frame protein hydrolysates.²⁸

This review summarizes the current state of insect proteins including their nutritional value in terms of essential amino acid content and digestibility. Then, the identification and functional properties of insect proteins are discussed for various species. Protein extraction methods and their corresponding protein extraction yields and protein contents of the final products are also reported. Furthermore, the main methods and common enzymes used for protein hydrolysates production and release of bioactive peptides are outlined. Several bioactive properties including antihypertensive, antidiabetic, antioxidant and anti-inflammatory properties are also described along with potential mechanisms of action. Finally, the allergenic properties of insect proteins are reviewed and the potential use of some novel technologies to reduce their allergenic properties are discussed.

2. Proteins in edible insects

2.1. The nutritional properties of insect proteins

Legume proteins have been widely applied in many commercial food products, however insect protein is reported to have enhanced nutritional value, as it contains all the essential amino acids (EAAs) like Leu, Ile, Lys, Met, Thr, Trp, Phe, and Val, and the amount of EAAs prove to be higher than an adult's daily amino acid FAO requirement.⁷ A comparison of EAA contents of diverse insect sources and well-known protein sources is displayed in Fig. 1. The EAA levels of a diverse set of insects were comparable to soybean proteins, while being lower than casein.³ H. illucens seems to be the insect containing the highest EAA amounts, reaching 45 g/ 100 g protein, comparable to beef (41.9 g/100 g protein). Widely researched T. molitor, A. domesticus and A. mellifera also contain a high content of EAAs: 33.84 g/100 g protein, 36.39 g/100 g protein and 37.7 g/100 g protein, respectively. Moreover, the protein content in common edible insects such as A. domesticus (72.45%), T. molitor (45%) and A. assamensis pupae (38.05%) are higher than that in common legumes such as lentils (26.7%), beans (23.5%) and soybean (41.1%).^{1,2,29,30}

Another advantage of insect proteins is their relatively high digestibility (as measured by in vitro assays). In vitro protein digestibility assays are rapid, easy to achieve, relatively cheap, give results similar to in vivo data and have been applied as a useful method to monitor protein digestibility.31 Several factors such as thermal unfolding, aggregation, carbonylamine reactions, cross-linking and interactions with other compounds such as carbohydrates or polyphenols have an impact on protein digestibility; furthermore, the polypeptide chain's accessibility and flexibility also affect protein digestibility. Digestibility values for insect proteins were first reported for edible insects from Mexico, for honeybees (89%), Liómetopum ants (82%), Sphenarium grasshoppers (75%), bees (71.7%), and ants (77.63%).³² Similar studies reported the digestibility values of S. ricinii pupae (86%) as well as T. molitor (54%), M. subhylanus (76.3%) and R. differens (90.5%).³³⁻³⁵ The digestibility of some insect proteins can be comparable to some common food proteins such as whole beef (89%), pork (90%), turkey (78%) and salmon (85%).

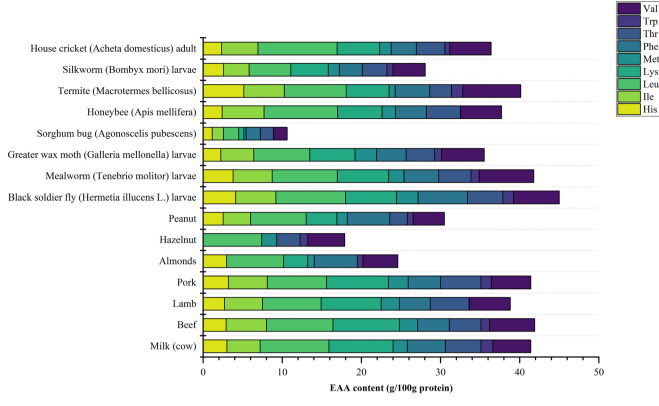


Fig. 1 Comparison of essential amino acid contents of insect sources and common food sources.

2.2. Insect protein characterization

To date, SDS-PAGE has been widely applied to separate insect proteins with different molecular weights. Then bands are selected for in-gel extraction, trypsin digestion and LC-MS/MS analysis in order to identify proteins. Insect protein identification nowadays is based on multiple sequencing and alignments tools to find regions of local similarities between protein sequences.³⁶ Although it is difficult to identify the proteins or peptides from most edible insects, as many of the expressed proteins have not been fully sequenced yet, there is a lack of complete insect proteomes, which are still needed. The proteins in insects such as D. melanogaster flies were fully identified, including 13 protein categories mainly classified as 'gene expression transcription, protein metabolism, muscle structure, cytoskeleton organization and cell function'.³⁷ More specifically, proteins in T. molitor have been gradually identified, but only a 10% match of mass data was obtained.³⁵ For example, proteins with molecular weights less than 14 kDa in T. molitor are mainly proteins with anti-freezing properties, such as hemolymph proteins ranging from 8.5 to 13 kDa. Cuticle proteins range from 14 to 30 kDa and vitellogenin-like proteins are mainly over 95 kDa.³ Also, actin-like (42 kDa), α-actinin-4 (107 kDa), myosin heavy chain (262 kDa), myosin-2 essential light chain (16.8 kDa), tropomyosin 1 (75.2 kDa) and 2 (32.5 kDa), troponin I (23.8 kDa), troponin T (47.3 kDa), and putative troponin C (18.3 kDa) have been identified. Muscle proteins like α -actinin-4 (107 kDa), tropomyosin 1 and 2, and calponin (20.3 kDa) were present in significant amounts in the water-insoluble protein fraction.³⁵

Proteins in different parts of B. mori have also been gradually characterized. Zhou et al. identified five proteins in the midgut of B. mori, including myosin 1 light chain, tropomyosin 1, profilin, serpin-2 and glutathione peroxidase.³⁸ Other studies reported that the most abundant proteins in B. mori heads were myosin and actin, and other identified proteins were related to the neurological system such as cuticle and chemosensory proteins.³⁹ The proteins in the skeletal muscle of B. mori such as 'contractile proteins, metabolic proteins, regulatory proteins and signal transducing proteins' were identified during the change from larvae to pupae.⁴⁰ More interestingly, insect protein expression was found to be linked to the insect's diet in B. mori. Comparing B. mori fed with fresh mulberry leaves and B. mori fed with an artificial diet, a decrease in proteins related to immunity and energy metabolism was found in those fed with an artificial diet.38

However, current proteomic studies mainly focus on specific parts of insects such as midgut proteins or insect secretions such as venom.^{41,42} Only limited studies revealed the intriguing traits behind whole protein groups in specific insects; among the proteomic studies available,^{37,38} the main focus has been on water soluble and water insoluble proteins, so potentially different protein groups such as those that are

soluble in salts solutions, organic solvents or alkaline/acid solutions are of interest for further studies as these have been rarely characterized.

2.3. Functional properties of insect proteins

It is important to evaluate the functional properties of insect proteins such as solubility, foaming and emulsifying properties, gelation, water holding capacity and oil holding capacity.^{43,44} Table 1 shows the functional properties of insect derived ingredients.

Among all the functional properties, solubility is certainly the most important one as many functional properties are influenced by the extent of protein solubilized in aqueous solution.45 The solubility of insect derived ingredients is influenced by protein structure, protein size, protein charge and pH. Less compact proteins are more likely to interact with water and have a higher solubility; also lower protein size improves solubility. In addition, enzymatic hydrolysis is believed to release more ionized groups, thus producing smaller proteins and peptides, and as a result, increasing solubility. As for the pH, the highest solubility tends to be observed at extreme pH values as more hydrophobic groups are exposed as a result of protein unfolding.⁴⁶ The highest solubility among T. molitor, S. gregaria and G. sigillatus derived ingredients was observed for mealworm protein with 97% solubility at pH 11. S. gregaria protein and G. sigillatus protein also showed high solubility values of 90% and 96%, respectively, at pH 11.47 However, when the pH is close to the isoelectric point (pI) of insect proteins, these tend to aggregate and they become less soluble. For example, T. molitor protein showed approximately 15% solubility and H. illucens defatted flour also showed only 11% solubility at pH 4 (close to the pI; pI around 4).48

Foams can be defined as air bubbles imprisoned in liquid and stabilized by proteins at the interface of air and liquid. Foam formation is also affected by a variety of factors including protein structure. As the formation of foam requires protein unfolding, globular and compact proteins were demonstrated to be less effective for foam formation than other fibrous proteins.⁶ The main fibrous proteins found in insects are from muscle, keratin and connective tissues, and they are mainly responsible for the foaming properties of insects.⁴⁹ So, except for the proteins commonly used due to their high foaming capacity like egg white protein, whey protein, caseins and soy protein, as well as insect protein sources extracted under the optimum conditions could be applied for foaming purposes although some insect species such as *B. mori* have a relatively low foaming capacity.^{6,50} The low foaming capacity of some insect proteins can be due to their conformational characteristics. For some insects, globular proteins are the main protein type. Since globular proteins are less unfolded at the interface between air and water, it limits their air bubble encapsulation ability.⁴⁷ Also, the low foaming capacity of insect proteins is attributed to the large variation of protein molecular weights, which negatively impacts the formation of protein films.⁵¹ Besides the foaming

 Table 1
 Insect protein functional properties

	Type of				Emulsifying properties	perties	Foaming properties	ties	÷
Insect	ingredient and protein content	Protein solubility	water holding capacity	Oil holding capacity	Capacity	Stability	Capacity	Stability	Gening
Mealworm $(Tenebrio molitor)^a$	Protein ND	97% (pH 11)	$3.95\pm 0.2~g~g^{-1}$	$2.74\pm 0.06~g~g^{-1}$	$66.6 \pm 2.16\%$	$51.31 \pm 0.46\%$	$32.67 \pm 0.94\%$	$30.33 \pm 0.47\%$	ND
Locusts (<i>Schistocerca gregaria</i>) ^a	Protein ND	90% (pH 11)	$2.31\pm 0.19~g~g^{-1}$	$3.22 \pm 0.16 \ g \ g^{-1}$	$67.78 \pm 1.6\%$	$50.41 \pm 1.99\%$	$32.0 \pm 1.88\%$	$6.17\pm0.71\%$	ND
Crickets (Gryllodes sigillatus) ^a	Protein ND	96% (pH 11)	$3.44\pm0.13~g~g^{-1}$	$3.33 \pm 0.11 \mathrm{~g~g^{-1}}$	$72.62\pm1.9\%$	$38.3 \pm 0.8\%$	$99.0 \pm 1.41\%$	$92.0 \pm 1.88\%$	ND
Mealworm $(Tenebrio\ molitor)^a$	Insect flour 52.35%	ND	$1.29\pm0.19~g~g^{-1}$	$1.71\pm 0.13~gg^{-1}$	$65.96 \pm 1.5\%$	$27.59 \pm 1.18\%$	$31.0 \pm 1.41\%$	$26.0 \pm 0.94\%$	ND
Locusts (<i>Schistocerca gregaria</i>) ^a	Insect flour 76%	ND	$2.18\pm 0.07~g~g^{-1}$	$1.98\pm 0.16~gg^{-1}$	$69.17 \pm 0.59\%$	$48.11 \pm 0.57\%$	$22.33 \pm 1.41\%$	$19.33 \pm 0.94\%$	ND
Crickets (<i>Gryllodes sigillatus</i>) ^a	Insect flour 70%	ΟN	$2.34\pm 0.28~g~g^{-1}$	$2.82\pm 0.08~g~g^{-1}$	$62.0 \pm 1.25\%$	$31.65 \pm 0.92\%$	$41.0 \pm 1.41\%$	$34.67 \pm 2.82\%$	ND
Cirina forda ^b	Insect flour 20%	90% (pH 5.5)	$248.3 \pm 2.4\%$	$178.7\pm0.0\%$	$135.7 \pm 2.3\%$	ND	0%0	ND	$14 \pm 0.1 \text{ w/v}$
Oryctes owariensis c	Insect flour 50.64%	ΟN	$220.33 \pm 1.5\%$	$265.9 \pm 1.31\%$	$29.97\pm0.12\%$	$104.84 \pm 1.23\%$	$17.87 \pm 0.16\%$	$12.36 \pm 0.03\%$	ND
Cirina forda ^d	Insect flour 55.5%	55% (pH 11)	$300.0 \pm 0.15\%$	$358.44 \pm 0.21\%$	$36.67 \pm 0.11\%$	$45.36 \pm 0.21\%$	$7.10\pm0.20\%$	$3.00\pm0.00\%$	$6.0\pm0.0\%$
House cricket (Acheta domesticus) ^e	Protein concentrate 71.7% (DW)	ΟN	$201.99 \pm 10.01\%$	ND	$7.52 \pm 0.24\%$	$97.36 \pm 1.13\%$	$26.00 \pm 2.00\%$	$86.79 \pm 2.21\%$	ND
Field cricket (<i>Gryllus bimaculatus</i>) ^e	Protein concentrate 60.7% (DW)	ND	$201.22 \pm 5.65\%$	ND	$7.19 \pm 0.38\%$	$97.38 \pm 0.44\%$	$16.67 \pm 1.15\%$	$92.58 \pm 0.91\%$	ND

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capacity, the foaming stability of *B. mori* protein concentrates (93.46%) was reported to be higher than for other insect protein concentrates such as *A. mellifera* (44.4–55.5%), *T. molitor* (30.33%) and *S. gregaria* (6.17%) and even higher than milk protein concentrates (55.86%).⁶ This is due to the amino acid composition of *B. mori* protein concentrates; they contain a high ratio of non-polar/polar side amino acid chains, which seems to enhance the stability and cohesive nature of the membrane. On the other hand, the high sugar content in protein flour is another explanation for the low foaming stability for the aforementioned species, as it can decrease the protein–protein interactions and interrupt the formation of firm interfacial membranes, and as a result, reduce the ability to stabilize foams.⁴⁷

Emulsions are homogenous mixtures of two immiscible liquids, either water droplets in an oil phase or oil droplets in an aqueous phase. Although whole Gryllidae sp. flour is found to have poor emulsifying properties, proteins are known to reduce the surface tension of the water-oil surface due to their amphiphilic character. Recent studies also showed that the emulsifying stability of T. molitor protein (51.3%) was higher than milk protein concentrates (33.50%). Furthermore, the hydrolysis of parent proteins is a valuable approach for improving the insect protein emulsifying properties, as it can facilitate the exposition of inaccessible amino acids and thus these residues can increase hydrophobic interactions and increase the formation of emulsions. For example, hydrolyzing G. sigillatus derived proteins with alcalase for 60 s resulted in an improvement in both emulsion capacity and emulsion stability after 60 min.⁵²

Another key functional property is the oil holding capacity (oil binding capacity or oil holding capacity), which refers to the ability of the protein to absorb lipids. Small, low-density hydrophobic proteins were reported to be able to preserve more lipids, thus providing a tender structure and more palatable textures.^{53,54} Zielińska *et al.* (2018) reported that the lowest oil holding capacity was 2.74 g oil per g protein for *T. molitor*, due to its lower protein content and higher fat content than other insects.^{5,47} Zhao *et al.* (2016) also reported a similar oil holding capacity value for *T. molitor* protein with 2.33 g oil per g protein.⁵⁵ In comparison, *G. sigiilatus* protein and *S. gregaria* protein showed greater oil holding capacities, with values of 3.33 g oil per g protein and 3.22 g oil per g protein, respectively.⁴⁷

Protein gels are able to retain a large amount of water as they are a structured protein network. Gelation depends on electrostatic interactions and temperature (the most important factors) as heat promotes the unfolding and denaturation of protein and causes their rearrangement and aggregation.⁵⁶ However, studies focusing on the gelling properties of insect proteins are scarce. Yi *et al.* (2013) evaluated the relationship between the gelation properties and pH of five edible insects and found that no gel formation was observed at a concentration of 3% w/v at pH 3, 5, 7 and 10, except for *A. domesticus* protein, which formed a firm gel at pH 7. For 30% w/v systems, all the insect proteins led to firm gels at pH 7 and pH 10. The transparency was related to the pH applied, samples heated at pH 3 and 10 led to a greater transparency than those at pH 5 and 7. As for the heating time, after 10 minutes, the gels formed were stable, and no impact on the gelation properties was observed following additional heating.3,57 Another functional property highly related to gelation is the waterholding capacity (or water-binding capacity or water adsorption capacity), which reflects the ability of proteins to retain water. This functional property is generally increased by heatinduced denaturation. This functional property of insect proteins has been characterized mainly for T. molitor; the water holding capacity in T. molitor flour was 129%, while that in T. molitor protein concentrate was 395%, much higher than what was observed for soy protein concentrate (227%), cowpea flour (124.6%), soy flour (130%) and chickpea flour (131.6%).47,58

2.4. Current insect protein extraction methods

Protein extracted from insects as food ingredients is suggested to be a useful way to increase the acceptance of eating insects.⁵⁹ Lipid removal before insect protein extraction was the most common step reported in the literature. Organic solvents like n-hexane have been applied to A. assamensis, T. molitor, Z. morio, A. diaperinus, A. domesticus, and B. dubia for lipid removal.^{3,30} Petroleum ether is another common solvent currently applied for defatting T. molitor powder and *B. mori* pupae.^{60,61} Other defatting protocols have used combinations of ethanol and isopropanol or hexane and isopropanol (3:2 (v/v)) to defat *T. molitor*.⁵⁵ However, no comparison of the effectiveness of different defatting methods has been reported. The major drawback of applying organic solvents is protein loss, as proteins like prolamins may have affinity for the solvent and be washed away during defatting, but this has not yet been quantified. Another disadvantage of using organic solvents is safety issues. Another solvent of interest for defatting is ethanol, which is considered as GRAS (generally recognized as safe), but the efficiency of ethanol is lower than that of other organic solvents.⁶² Therefore, a more versatile method that extracted proteins from whole insect meals was carried out as follows, the whole insect meal was mixed with demineralized water and sieved through a 500 µm sieve, then the lipid layer was removed from the top of the supernatant after centrifugation (at 4 °C), thus this method could be employed as an alternative way to remove lipids from insect homogenates.³

To date, insect protein extraction protocols are mainly based on wet fractionation (Fig. 2). The most common/traditional wet fractionation method is alkaline solubilization coupled to isoelectric precipitation, although some research may apply alkaline solubilization or acid solubilization separately (Table 2). In brief, for alkaline solubilization coupled to isoelectric precipitation, the insect meal is solubilized at alkaline pH (8–11) to solubilize proteins, then centrifugation is applied to separate soluble proteins in solution, then the soluble protein solution is collected and the pellet discarded, afterwards the pH is adjusted to the isoelectric point (pI) to precipitate proteins. The protein content of final products with

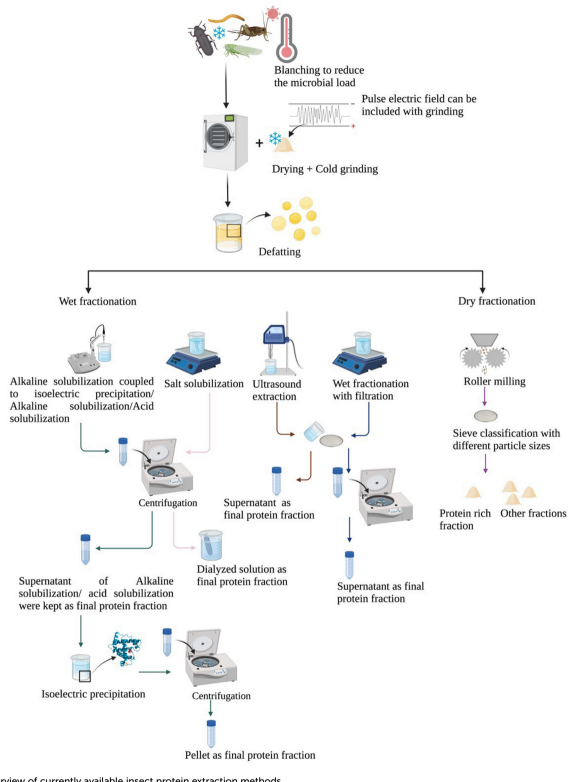


Fig. 2 Overview of currently available insect protein extraction methods.

this type of extraction method was between 34.7% and 89.05% according to different insect species or individual differences observed for the same insect species (Table 2). The precipitation pH is vital to alkaline solubilization coupled to isoelec-

tric precipitation, as it determines the protein content of the products and protein recovery. The pI distribution of different insect proteins has been reported for various insect species, and was first measured in *Sarcophaga falculata* larva; the outer

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Table 2 Studies of insect protein extraction

Source	Life stage	Protein content (starting material)	Method of extraction	Conditions	Protein content (final product)	Protein extraction yield/ protein recovery	Ref.
Muga silkworm (<i>Antharaea assamensis</i>)	Pupae	38.05% (wet basis)	Alkaline solubilization coupled to isoelectric precipitation	Ratio 1 : 20 defatted powder : deionized water, at PH 9.0 for 2 h, centrifuged ($8000g$, 15 min at 20 °C). The pellets were freeze dried, supermatant PH 4.5, centrifuged ($8000g$, 15 min at 20 °C). Wash twice and freeze dry. Store at 4 °C.	$65.16 \pm 0.53\%$	28.07 ± 0.34% (protein recovery)	30
Cricket (Acheta domesticus)	Adult	72.45 ± 1.30 g/ 100 g (dry basis)	Alkaline solubilization	Ratio 1: 50 cricket powder: 0.5 M NaOH, stir for 60 min; 2 mL of mixture are withdrawn at 15 min intervals and centrifuged (10000g, 10 min at 20 °C). The supernatant was isolated and stored at 4 °C.	About $46 g/100 g$	63.49% (protein extraction yield)	1
House cricket (Acheta domesticus)	QN	53%	Alkaline solubilization coupled to isoelectric precipitation.	Ratio 1:15 defatted powder: 0.25 M NaOH, stir for 60 min at 40 °C, centrifuged (3500g, 20 min at 4 °C) twice. Adjust the pH of supernatants to 4.3 4.5 before centrifugation (250g, 15 min at 4 °C) the residue was usafied twice with distilled water at pH 4.5 and centrifuged (250g, 10 min, at 4 °C). The residue and supernatant	For different defatting methods: Hexane: $74.7 \pm$ 0.3%; Petroleum ether: $74.7 \pm 0.2\%$; Ethyl acetate: $74.4 \pm 1.2\%$; Ethanol: $78.5 \pm 2.0\%$	For different defatting methods: Hexane: 32.4 \pm 3.8%; Pertoleum ether: 33.1 \pm 1.0 %; Ethyl acetate: 31.6 \pm 3.0%; Ethanol: 31.0 \pm 4.0%	0
Mealworm (Tenebrio molitor)	DN	45%	Alkaline solubilization coupled to isoelectric precipitation.	were neces-uncurve Ratio 1:15 defatted powder: 0.25 M NaOH, stir for 60 min at 40 °C, centrifuged (3500g, 20 min at 4 °C) twice. Adjust the pH of supernatants to 4.3-4.5 before centrifugation (2500g, 15 min at $^{\circ}$ C). The residue was washed twice with distilled water at pH 4.5 and centrifuged (2500g, 10 min, at were freeze-dried.	For different defatting methods: Hexane: 74.0 \pm 2.2%; Petroleum ether: 7.2.7 \pm 1.5%; Ethyl acetate: 75.4 \pm 0.5%; Ethanol: 75.3 \pm 0.8%	For different defatting methods: Hexane: 33.7 $\pm 1.6\%$; Pertoleum ether: $33.5 \pm 1.2\%$; Ethyl acetate: $33.2 \pm 0.6\%$; Ethanol: $33.9 \pm 3.7\%$ (Protein recovery)	0
Mealworms (Tenebrio molitor)	Larvae	QN	Alkaline solubilization coupled to isoelectric precipitation	Ratio 1 : 10 ground insects : 0.2% NaOH, stir for 1 h at room temperature. Centrifugation (8000g, 20 min at 4 °C). Precipitation at isoelectric point. Then centrifuged (8000g, 20 min at 4 °C) and washed at 4 °C) and	QN	QN	26
Locusts (Schistocerca gregaria)	Adult	ДИ	Alkaline solubilization coupled to isoelectric precipitation	utucu at - 10 °C. Ratio 1: 10 ground insects: 0.2% NaOH, stir for 1 h at noom temperature. Centrifugation (8000g, 20 min at 4 °C). Precipitation at isolectric point. Then centrifuged (8000g, 20 min at 4 °C) and washed with distilled water. Freeze-	QN	QN	97
Crickets (Gryllodes sigillatus)	Adult	QN	Alkaline solubilization coupled to isoelectric precipitation	drucd at - 18°C. Ratio 1 : 10 ground insects : 0.2% NaOH, stir for 1 h at noom temperature. Centrifugation (8000g, 20 min at 4°C). Precipitation at isoelectric point. Then centrifuged (8000g, 20 min at 4°C) and washed with distilled water. Freeze- dried at -18°C.	QN	Q	97

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Table 2 (Contd.)

Source	Life stage	Protein content (starting material)	Method of extraction	Conditions	Protein content (final product)	Protein extraction yield/ protein recovery	Ref.
Crickets (Gryllidae)	Q	63.43%	Alkaline solubilization or acid solubilization	Ratio of $6:1$ defatted powder : 0.5 M aqueous NaOH or ascorbic acid. Vortex and centrifuge (3500g, 20 min at 4 °C). Then the second extraction step was on the insoluble pellet. Vortex and centrifuge again. Keep the pellet and freeze dry.	61.75% (alkaline); 69.69% (acid)	82.95% (alkaline); 67.66% (acid) (protein extraction yield)	131
Tenebrio molitor	Larvae	19.1 ± 1.3%	Wet fractionation with filtration	Ratio 1: 3: 0.005 N ₂ frozen insects : demineralized water : ascorbic acid (w/v/ w), blend for 1 min. The suspension was sieved through a stainless-steel filter sieve (pore size 500 µm). The fil- trates and residues were collected. After centrifugation (15 000g, 30 min, at 4 °C) the supermatant, the pellet, and the fat fraction were obtained. The residue, the pellet and the supermatant fractions	Supermatant (d.b.) : 50% to 61%; Pellet (d.b.): 65% to 75% ;Residue (d.b.): 58% to 69%	103% (protein extraction yield)	m
Zophobas morio	Larvae	$20.7 \pm 0.3\%$	Wet fractionation with filtration	were freeze dried. Watio 1: 3: 0.005 N ₂ frozen insects : demineralized water : ascorbic acid (w/v/ w), blend for 1 min. The suspension was sieved through a stainless-steel filter sieve (pore size 500 µm). The fil- trates and residues were collected. After centrifugation (1500g, 30 min, at 4 °C) the supermatant, the pellet, and the fat fraction were obtained. The residue, the pellet and the supermatant fractions	Supermatant (d.b.) : 50% to 61%; Pellet (d.b.): 65% to 75%; Residue (d.b.): 58% to 69%	88.1% (protein extraction yield)	е
Alphitobius diaperinus	Larvae	$20.6 \pm 0.1\%$	Wet fractionation with filtration	where freeze dried. Ratio 1 : 3 : 0.005 N ₂ frozen insects : demineralized water : ascorbic acid (w/v/ w), blend for 1 min. The suspension was sieved through a stainless-steel filter sieve (pore size 500 µm). The fil- trates and residues were collected. After centrifugation (15000g, 30 min, at 4 °C) the supermatant, the pellet, and the fat fraction were obtained. The residue, the pellet and the supermatant fractions	Supermatant (d.b.) : 50% to 61%; Pellet (d.b.): 65% to 75%; Residue (d.b.): 58% to 69%	91.8%(protein extraction yield)	m
Acheta domesticus	Adult	$21.5 \pm 0.5\%$	Wet fractionation with filtration	were increase dired. Ratio 1: 3: 0.005 N ₂ frozen insects : demineralized water : ascorbic acid (w/v/ w), blend for 1 min. The suspension was sieved through a stainless-steel filter sieve (pore size 500 µm). The fil- trates and residues were collected. After centrifugation (15 000g, 30 min, at 4 °C) the supernatant, the pellet, and the fat fraction were obtained. The residue, the pellet and the supernatant fractions	Supernatant (d.b.) : 50% to 61%; Pellet (d.b.): 65% to 75%; Residue (d.b.): 58% to 69%	89.5%(protein extraction yield)	n
Blaptica dubia	Adult	19.3 ± 0.9%	Wet fractionation with filtration	were treeze arted. Ratio 1: 3: 0.005 N ₂ frozen insects : demineralized water : ascorbic acid (w/v/ w), blend for 1 min. The suspension was sieved through a stainless steel filter sieve (pore size 500 µm). The fil- trates and residues were collected. After centrifugation (1500g, 30 min, at 4 °C) the supermatant, the pellet, and the fat fraction were obtained. The residue, the pellet and the supermatant fractions were freeze dried.	Supernatant (d.b.) : 50% to 61%; Pellet (d.b.): 65% to 75%; Residue (d.b.): 58% to 69%	86.5% (protein extraction yield)	n

Table 2 (Contd.)

od & Fur	nction								Rev
Ref.	132	133	134	134	134	70	20	70	14
Protein extraction yield/ protein recovery	Precipitate: 32.72 ± 1.34%; Residue: 48.32 ± 0.92 (protein extraction yield)	Q	Ŋ	Q	QN	Ŋ	ND	ND	35% (protein extraction yield)
Protein content (final product)	Precipitate: 66.66 ± 0.82% (crude protein); Residue: 65.79 ± 1.3% (crude protein)	$82.26 \pm 0.62\%$	QN	CIN	Q	ND	QN	QN	Q
Conditions	Ratio 1 : 3 : 0.005 cricket meal : distilled water : ascorbic acid, blended. The suspension was steved through a stainless steel filter sieve (pore size 500 μ m). The residue was freeze dried, and stored at 4 °C. The filtrate was centifuged (10 000 rpm, 30 min at 4 °C), then the precipitate was freeze dried.	Ratio 1:1 locust meal: deionized water at pH 9.0, stirred for 2 h. Chitin was separated by filtering the slurry through a fine-mesh cheese filtering cloth (por size 0.2–0.5 mm, 100% cotton), then the filtrate was freeze-dried.	Ratio 1:2 defatted insect powder :0.02% ascorbic acid, homogenized and filtered by medical gauze. The filtered solution was centrifuged (15 000g 30 min); the sumen atapt was collected	Ratio 1:2 defatted insect Ratio 1:2 defatted insect powder :0.02% ascorbic acid, homogenized and filtered by medical gauze. The filtered solution was centrifuged (15 000g 30 min); the current attact was collected	Particular function of the sector of the sector of the sector powder : 0.02% ascorbic acid, homogenized and filtered by medical gauze. The filtered solution was contribuyed (15 000g 30 min); the supernatant was collected.	Defatted sample was used for sonication: 20 kHz for 15 min, with 75% amplitude and pulse every 3 s with	Defatted sample was used for sonication: 20 kHz for 15 min, with 75% amplitude and pulse every 3 s with a 1 s interval.	Defatted sample was used for sonication: 20 kHz for 15 min, with 75% amplitude and pulse every 3 s with a 1 s interval.	Ratio 1 : 16 defatted insect coarse meal : distilled water containing 9.46 mM ascorbic acid, sonication was carried out for 20 min on ice at 20 kHz with a 75% AMP and pulsed every 3 s. Then the aliquots were collected at 1, 2, 5, 10, 15, and 20 min. Samples were sieved through a stainless-steel filter (pore size of 1 mm) and filtrates were collected and freeze-dried.
Method of extraction	Wet fractionation with filtration	Wet fractionation with filtration	Wet fractionation with filtration	Wet fractionation with filtration	Wet fractionation with filtration	Ultrasound extraction	Ultrasound extraction	Ultrasound extraction	Ultrasound extraction
Protein content (starting material)	$59.84 \pm 1.64\%$	$65.87 \pm 0.42\%$	$58.44 \pm 2.89 \text{ g/}$ 100 g	$49.68 \pm 0.54 g/$ 100 g	$51.08 \pm 0.40 \text{ g}/100 \text{ g}$	ND	ЛЛ	ŊŊ	Q
Life stage	QN	QN	Larvae	Larvae	Larvae	Adult	Larvae	QN	Larvae
Source	Crickets (Acheta domesticus)	Migratory locust (Locusta migratoria)	Tenebrio molitor	Allomyrina dichotoma	Protaetia brevitarsis seulensis	Crickets (G. Bimaculatus)	Mealworm (T. molitor)	Silkworm (<i>B. mort</i>) pupae	Yellow mealworm (<i>Tenebrio molitor</i>)

Food & Function

Review

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Table 2 (Contd.)

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Ref.	14	14	135	136	48	48
Protein extraction yield/ protein recovery	37% (protein extraction yield)	94% (protein extraction yield)	59.9% (protein extraction yield) (larvae); 26.4% (protein extraction yield) (larvae meal)	(protein extraction yield) pH 3: 10.3%; pH 4: 10.5%; pH 5: 19%; pH 6: 13%; pH 9: 47%; pH 10: 53.6%	CN .	QN
Protein content (final product)	Q	Q	83.7 ± 3.8% (larvae); 80.4 ± 0.9% (larvae meal)	pH 3: 0.85 mg mL ⁻¹ ; pH 4: 0.9 mg mL ⁻¹ ; pH 5: 1.5 mg mL ⁻¹ ; pH 6: 1.7 mg mL ⁻¹ ; pH 7: 1.8 mg mL ⁻¹ ; pH 9: 2.2 mg mL ⁻¹ ; pH 9: 4.6 mg mL ⁻¹ ; pH 10:	$68.2 \pm 0.3 \text{ g g}^{-1} \text{ DM}$	$34.7 \pm 0.2 \text{ g g}^{-1} \text{ DM}$
Conditions	Ratio 1: 16 defatted insect coarse meal: distilled water containing 9.46 mM ascorbic acid, sonication was carried out for 20 min on ice at 20 kHz with a 75% AMP and pulsed every 3 s. Then the aliquots were collected at 1, 2, 5, 10, 15, and 20 min. Samples were sieved through a stainless-steel filter (pore size of 1 mm) and filtrates were collected and freezed-rise	Ratio 1: 16 defatted insect coarse meal : distilled water containing 9.46 mM ascorbic acid, sonication was carried out for 20 min on ice at 20 kHz with a 75% AMP and pulsed every 3 s. Then the aliquots were collected at 1, 2, 5, 10, 15, and 20 min. Samples were sieved through a stainless-steel filter (pore size of 1 mm) and filtrates were collected and freeze-dried.	Ratio 1 : 40 larvae meal/distilled water, pH 10. The mixture was stirred for 1 h at 45 °C. After stirring, distilled water at pH 10 was added to reach 250 mL, then the mixture was centrifuged (10 000g for 30 min) and the supernatants were freeze-dried.	Ratio 1 : 10 larvae meal : buffer solution with pH varied from 3.0–10.0 (a) 0.1 M citric acid/Na ₂ HPO ₄ / > 0); (b) 0.1 M Na ₂ HPO ₄ / Na ₄ PO ₄ buffers (pH 9.0 1 M Na ₅ CO ₃ /NaHCO ₃ buffers (pH 9.0 and 10.0) Centrifugation and separation of fat fraction (upper), aqueous fraction (intermediate) and insoluble com- ponents (lower slurry). Then the protein content in the aqueous fraction (inter- mediate) was measured by BCA assay.	Ratio 1: 25 defatted meal : distilled water at pH 10, stirring (300 rpm, 30 min, at 60 °C.) The extract was centrifuged (4000; 20 min at 20 °C), then the pH of the supernatant was adjusted to 4. Re-extraction (PH 2, 60 °C, 30 min) and precipitation again. Both extracts were centrifuged (4000; 20 min at 20 °C). Foreins and insoluble resolves unsoft for a 20 °C).	Ratio 1: 25 defatted meal: distilled water at pH 10, stirring (300 rpm, 30 min, at 60 °C.) The extract was centrifuged (4000g, 20 min at 20 °C), then the pH of the supernatant was adjusted to 4. Re-extraction (pH 2, 60 °C, 30 min) and precipitation again. Both extracts were centrifuged (4000g, 20 min at 20 °C). Proteins and insoluble residues were freeze dried at -80 °C.
Method of extraction	Ultrasound extraction	Ultrasound extraction	Alkaline solubilization	Salt solubilization	Alkaline solubilization coupled to isoelectric precipitation	Alkaline solubilization coupled to isoelectric precipitation
Protein content (starting material)	QN	â	65.6% (larvae) 71.6% (larvae meal) blanched, dried and ground	8.2 g/100 g	30.0 ± 1.2 g per g DM	31.7 ± 0.5 g per g DM
Life stage	Adult	QN	Larvae	Larvae	Larvae	
Source	Field cricket (Gryllus bimaculatus)	Silkworm pupae (Bombyx mori)	T. molitior	Rhynchophorus phoenicis	Black soldier fly (<i>Hermetica illucens</i>)	Yellow mealworm (<i>T. molitor</i>)

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Table 2 (Contd.)							1
Source	Life stage	Protein content (starting material)	Method of extraction	Conditions	Protein content (final product)	Protein extraction yield/ protein recovery	Ref.
Tenebrio molitor	Larvae	$58.44 \pm 2.89\%$	Acid solubilization and salt solubilization	Ratio 1 : 2 defatted powder : buffer, stepwise extraction was carried out (a) Water-soluble protein : 0.056 (w/v) ascorbic acid; (b) Salt-soluble protein : 0.58 M saline solution (0.49 M NaCl, 17.8 mM Na ₂ P ₃ O ₁₀ , and 1 mM NaCl, 17.8 mM Na ₂ P ₃ O ₁₀ , and 1 mM NaVl, 17.8 mM Na ₂ P ₃ O ₁₀ , and 1 mM NaVl, 17.8 mM na ₂ P ₃ O ₁₀ , and 1 mM redical gauze. The filter through medical gauze. The filter through tuged (15 000g; 30 min at 2 °C), the water-soluble and salt-soluble protein solutions were obtained.	Q	Q	137
Allomyrina dichotoma	Larvae	$49.68 \pm 0.54\%$	Acid solubilization and salt solubilization		ND	ND	137
Protaetia brevitarsis seulensis	Larvae	$51.08 \pm 0.40\%$	Acid solubilization and salt solubilization		ND	ND	137
Grasshopper (<i>S. gregaria</i>)	Adult	38.1 ± 1.4%	 Alkaline solubilization coupled to isoelectric precipitation (ALK) 2. Alkaline and sonication extractions (SON) 	ALK: ratio of 1: 15 insect powder: distilled water at pH 10, stirred $(40 \circ C, 1$ h). Then centrifuge (30009, 20 min at $4 \circ C$) and separate the insoluble residues. Adjust the supernatant pH to 4 (grasshopper) and 5 (honeybee) for 1 h (room temperature), then centrifuge solutions (15 009, 20 min at 4 °C). Precipitates were neutralized to pH 7. All fractions were freeze-dried; SON: ratio of 1: 15 insect powder: distilled water at pH 10 (using 0.1 M NOH), sonication (amplitude – 70, pulse-on time – 30 s, pulse-off time – 30 s, process	52.9 ± 1.3%(ALK); 57.5 ± 1.0%(SON)	24.2 ± 1.2% (ALK); 30.6 ± 1.6% (SON) (ALK: Protein recovery; SON: Protein extraction yield)	66
Honeybee (A. meltifera)	Larvae and pupae	$22.1 \pm 0.1\%$	 Alkaline Alkaline solubilization coupled to isoelectric precipitation (ALK) 2. Alkaline and sonication extractions (SON) 	Alk: ratio of 1:15 insect powder: distilled water at pH 10, stirred $(40^\circ C_1)$ h). Then entrifuge (300g; 20 min at 4 °C) and separate the insoluble residues. Adjust the supernatant pH to 4 (grasshopper) and 5 (hneybee) for 1 h (room emperature), then centrifuge solutions (15 000; 20 min at 4 °C). Freipiates were freeze-dried; SON: ratio of 1:15 insect powder: distilled water at pH 10 (using 0.1 M NOCH), sonication amplitude – 70, pulse-on time – 30 s, pulse-off time – 30 s, process time – 6 min).	39.6 ± 2.1%(ALK); 55.2 ± 1.3%(SON)	27.5 ± 1.1% (ALK); 30.8 ± 0.8% (SON) (ALK: Protein recovery; SON: Protein extraction yield)	66
Yellow mealworm (<i>Tenebrio molito</i>)	Larvae	$51.5 \pm 0.5\%$	Alkaline solubilization coupled to isoelectric precipitation	Ratio 1: 15 defatted powder : 0.25 M NaOH solution at 40 °C for 60 min, and vortex every 15 min, centrifuge (3500g, 20 min at 4 °C. The supernatart and gel were adjusted to pH 4.3–4.5 (room temperature), centrifuged (2500g, 20 min at 4 °C). Washed twice and centrifuged twice (2500g, 20 min at 4 °C). and freeze dried.	79.0% (true protein)	53% (protein recovery)	55

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(Contd) Table 2

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Table 2 (Contd.)

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Source	Life stage	Protein content (starting material)	Method of extraction	Conditions	Protein content (final product)	Protein extraction yield/ protein recovery	eview
Silkworm (<i>Bombyx mori</i>)	DN	$48.20 \pm 0.84\%$	Alkaline extraction coupled to isoelectric precipitation	Ratio 1:10 sample: distilled water, stirred for 1 h and vacuum filtered. The pH of the crude protein extract was adjusted to 3.5 or 4.5, stirred for 10 min and centrifuged (10 000g, 10 min at 1 eV. beso in for 000 s 10 min at	0.462 ± 0.005 mg per mg defatted insect (Soxhlet method at pH 3.5)	QN	138
Bamboo caterpillar (Omphisa fuscidentalis	QN	$47.52 \pm 0.55\%$	Alkaline extraction coupled to isoelectric precipitation	with 100 cm $(9.5 - 0.5, 1.11)$. Ratio 1:10 sample: distilled water, stirred for 1 h and vacuum filtered. The pH of the crude protein extract was adjusted to 3.5 or 4.5, stirred for 10 min and centrifuged (10 000g, 10 min at 4 °C). Collect the pellets and diy them	0.18 ± 0.008 mg per mg defatted insect (Folch method at pH 4.5)	QN	138
Field cricket (Gryllus bimaculatus	QN	$52.08 \pm 0.13\%$	Alkaline extraction coupled to isoelectric precipitation	with note at ($3>-0^{-0}$, 1 , 1). Ratio 1:10 sample: distilled water, stirred for 1 h and vacuum filtered. The pH of the crude protein extract was adjusted to 3.5 or 4.5, stirred for 10 min and centrifuged (10 000g, 10 min at 4 °C). Collect the pellets and dry them with hot air (55–60 °C, 1 h).	0.734 ± 0.019 mg per mg defatted insect (Folch method at pH 3.5)	Q	138
Mealworm (<i>Tenebrio molitor</i> , L.)	Larvae	$38.64 \pm 1.74 g/$ 100 g	Alkaline solubilization	Deionized water was added to a concentration of 600 g L^{-1} and the slurry was mixed for 60 s, pH 10.5 and stirred for 2 h, then centrifuged (4000g, 10 min). The supernatant fraction was separated, frozen at -30 °C and freezedred at 0.2 mbar for 72 h.	$62.6 \pm 1.3 \text{ g/100 g}$	58.7 ± 1.2% (protein extraction yield)	13
Black soldier fly (<i>Hermetia ülucens</i>)	Larvae	42.00 ± 0.33%	Alkaline solubilization coupled to isoelectric precipitation	Ratio 1: 24.85 defatted meal : 0.25 M NaOH, heated (52.23 °C, 59.43 min) and stirred (100 rpm). The shurty was centrifuged (4500g, 10 min, at 4° C), and the supernatant pH was adjusted 44. The slurry was centrifuged (4500g, 10 min, at 4° C), and the precipitate was washed and centrifuged twice (4500g, 10 min, at 4° C). The pH was adjusted to 7, stored at -20° C, and freeze-dried.	$80.42 \pm 0.90\%$	64.44% (protein recovery)	139
Black soldier fly (<i>Hermetia ülucens</i>)	Larvae	37.3%	Alkaline solubilization coupled to isoelectric precipitation	Ratio 1 : 15 defatted meal : 0.25 M NaOH, stirred (300 rpm, 1 h, at 40 °C) and centrifuged (2493, 20 min, at 4 °C), same procedure was repeated as above. The solid fraction was collected, and the pH of supermatants was adjusted to 4.3–4.5, then the solution was centrifuged (1272g, 15 min, at 4 °C). The precipitate was washed twice with distilled water and freeze-dried.	61.1%	Q	140

ND: not determined.

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layer proteins of Sarcophaga falculata had an isoelectric point of 5.1, and the chitin-protein complex (inner layer) had a different pI of 3.5.63 Later, Huang et al. (1984) categorized the insects in various groups according to the pI values of their proteins, "Coleoptera, Orthoptera, Odonata, Hemitera and Lepidoptera" were included in the first group where the pI ranged between 5.56 and 5.69, with no difference between thoracic and leg muscle actins. The pI of cicada's sound muscle (5.57) and thoracic (5.65) and leg muscle actins (5.68) were also in the same range without significant difference. Another group included Hymenoptera and Diptera (pl = 5.75-5.87). In this group, the thoracic muscles may contain two or three actin isoforms (α -actin, β -actin, γ -actin). So for instance, the pI values of honey bee thoracic muscle were 5.75 and 5.87, while the ones of fruitfly were 5.70, 5.77 and 5.84.64 More recently, the pI values of different insects such as A. mylitta, A. pernyi, A. yamamai, G. mellonella, B. mori and B. mandarina were reported to be quite similar with values of 5.6, 5.82, 5.35, 4.67, 4.59 and 4.64, respectively.⁵⁷ Another widely used protocol is wet fractionation with mechanical separation, which is effective at obtaining different fractions such as fat and protein in separate layers. In brief, insect meal is mixed with ascorbic acid or sodium hydroxide, and the stainless filter sieve with a suitable pore size is applied to separate proteins, then the fat fraction is collected from the top of the supernatant after centrifugation (Table 2). The last novel protocol is ultrasound-assisted extraction, which results in extracts with protein contents varying between 35% and 94% (depending on the sample under study).⁶⁵ Ultrasound-assisted extraction does not change the amino acid profile of the resulting extract,¹⁴ although its final protein content is higher than that from alkaline solubilization coupled to isoelectric precipitation. Mishyna et al. (2019) observed that only 39.6% of protein was found in the final product extracted by alkaline solubilization coupled to isoelectric precipitation, compared to 55.2% obtained by ultrasound-assisted extraction. But this result may have limitations, which may be due to the specific species studied and/or the insect life stage, as no further studies are available.⁶⁶ Other novel technologies such as pulsed electric fields, which is a type of non-thermal technology, can enhance protein extraction from A. domesticus. Psarianos et al. demonstrated that the increase in protein extraction yield exceeded 18% when treated with pulsed electric fields, also, other components like fat also showed an increase (exceeded 40%) in extraction yield.¹

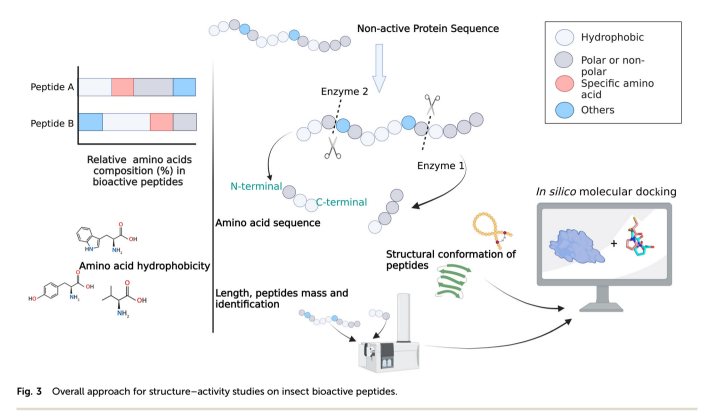
Another protein extraction protocol is dry fractionation, which avoids generating wastewater and energy-intensive processing. Similar to wet fractionation, defatting is also carried out before dry fractionation.^{13,67} So far, dry fractionation has not been widely applied to insect fraction separation. Dry fractionation produces different fractions with various nutrients; the protein content of *T. molitor* protein rich fraction (particle <500 μ m) can reach 58%.¹³ Among dry fractionation was reported as an efficient method for the generation of three fractions (lipid fraction, chitin-rich fraction and protein-rich

fraction) from *T. molitor* and *A. domesticus*; the protein-rich fraction was reported to contain less chitin than other fractions. However, the water solubility of protein in the defatted meal can be decreased by air classification; the *A. domesticus* defatted meal showed 13–23% solubility and the *T. molitor* defatted meal showed 14–27% solubility according to different pH values. This may be attributable to the fact that the large particle size hinders protein extraction.⁶⁷ Table 2 summarizes the protein content of various insect species at different research stages and also a comparison of the final protein content and extraction yield when applying different insect protein extraction protocols.

3. Insect protein hydrolysis and bioactive properties of insect-derived peptides

Generally, bioactive peptides consist of 2 to 20 amino acids as a non-active sequence encoded in parent proteins.⁶⁸ The amino acid sequence, length and other structural features such as secondary structure (β -sheets), cyclic structure (via disulfide bonds), relative ratio of a specific amino acid or group of amino acids, hydrophobic properties, molecular weight, type of residues at C- and N-terminals, among others, are linked to the structure-activity relationships of bioactive peptides and these are fundamental for the biological activities exerted (Fig. 3). Bioactive peptides can be released/produced through various processes, such as enzymatic hydrolysis, in vivo/in vitro digestion, microbial fermentation, chemical hydrolysis (not GRAS), peptide synthesis and novel processing technologies, such as high hydrostatic pressure, microwave and pulsed electric fields. Enzymatic hydrolysis has been identified as a good means to enhance or control bioactivity such as antihypertensive, antioxidant and antidiabetic properties.^{69–71} To date, farmed *B. mori*, which is the main byproduct of the silk reel industry, appears to be identified as a potential medicinal source and has gradually become the most frequently studied insect to generate insect-derived bioactive peptides.69,72

In general, enzymatic hydrolysis is applied to insect protein concentrates/isolates or directly to flours.^{73,74} Different studies outlined several available enzymes used in the generation of insect-derived peptides. For example, pepsin, as an aspartic acid endoprotease, has an active site with dual aspartic acids to hydrolyse peptide bonds. Trypsin is used mainly for cleaving arginine and lysine amino acid residues.⁷⁵ Enzymes have different and specific excision sites that generate different peptides, thus displaying various bioactivities. Currently, mammalian digestive enzymes such as amylase, lipase and trypsin appear to be the most frequently employed ones and are used to simulate gastrointestinal digestion. *S. littoralis, B. mori, S. gregaria*, and *B. terrestris* were reported to be hydrolyzed by *in vitro* gastrointestinal digestion.^{20,76} Farmed insects such as *G. sigillatus, S. gregaria* and *B. mori* have also been successfully



hydrolyzed through gastrointestinal digestion and new bioactive peptides have been isolated.5,77,78 Besides, microbial enzymes including alcalase, thermolysin, flavourzyme and acid protease were used for the hydrolysis of A. assamensis pupae and S. littoralis and T. molitor proteins.^{30,70,79} However, only limited research reported the use of plant-derived enzymes like papain for producing insect protein hydrolysates.⁸⁰ One recent study reported that the utilization of papain mixed with alcalase at 3.0% enzyme-substrate ratio and incubation for 3 h achieved the highest ACE inhibition, and the highest inhibition of DPP-IV was achieved when mixing papain and flavourzyme at 2.3% enzyme-substrate ratio for 3 h incubation.³⁰ Another innovative alternative is the combination of enzymatic hydrolysis with novel technologies: microwave-assisted enzymatic hydrolysis of whole crickets is demonstrated to be an advantageous method to hydrolyze whole insects as protein unfolding can be increased by microwave energy, and microwave energy can transport the peptide fragments hydrolyzed that are not by proteolytic enzymes.28,81,82

For different studies describing enzymatic hydrolysis of insect proteins, the degree of hydrolysis of insect protein has been assessed, varying from 3% to 100%. The large variation may be caused by the differences in hydrolysis conditions used such as the type of enzyme, E/S ratio, enzymatic activity, temperature, pH, the presence of protease inhibitors, and the insect species studied. For instance, when applying the same hydrolysis protocol (*in vitro* digestion with α -amylase, pepsin and pancreatin) to different insects, the degrees of hydrolysis of *A. annulipes* and *L. migratoria* were 15.8 and 36.3%, respect-

ively.⁷⁷ Results reported by Zielińska *et al.* (2015),⁵ for *G. sigillatus*, *T. molitor* and *S. gregaria* after *in vitro* digestion were approximately 32%, 14% and 30%, respectively. Also, the degree of hydrolysis was affected by different processing methods; boiling and baking increased the degree of hydrolysis of *S. gregaria* and *T. molitor* compared to raw samples. However, different results were observed for *G. sigillatus*: the degree of hydrolysis decreased from approximately 32% to 27% when applying boiling. But baked *G. sigillatus* still showed a higher degree of hydrolysis (37.5%) compared to the untreated sample. Table 3 summarizes the enzymatic conditions currently applied in some studies and the bioactivity of insect derived peptides from different insect species.

3.1. Antihypertensive properties

ACE inhibition is associated with antihypertensive properties; peptides with ACE inhibiting activity have already been extracted from various food proteins like soy, milk, fish and egg white.83-86 Studies have demonstrated that insect protein hydrolysates bioactive have properties including antihypertensive.87-89 Enzyme selection is a key factor for the generation of ACE inhibitory peptides.^{76,90} To achieve the highest ACE inhibition, recent studies reported different enzymes and optimum hydrolysis conditions for different insect proteins. Enzymes such as pepsin, trypsin and α-chymotrypsin were commonly used to simulate stomach and gastrointestinal digestion; α -amylase and pancreatin were applied to simulate saliva and intestinal digestion; and alcalase, thermolysin and flavourzyme were frequently applied to generate ACE-inhibiting hydrolysates. Other enzymes like neu-

Table 3 Studies on insect protein hydrolysis and bioactivity of insect-derived peptides

Source	Life stage	Protein content (starting material)	Enzymatic conditions	Main highlights	Re
Silkworm (Bombyx mori); Cotton leafworm (Spodoptera littoralis); Locust (Schistocerca gregaria); Buff-tailed bumblebee (Bombus terrestris)	Larvae; Larvae; Adults; Adults	ND	1. Simulated gastric phase: pepsin (enzyme/substrate: $1/250$ w/w for 2 h at 37 °C, pH 2). And simulated intestinal phase: trypsin and α -chymotrypsin (1 : 1) (enzyme/substrate: $1/250$ w/w for 2.5 h at 37 °C, pH 6.5); 2. Alcalase (enzyme/substrate: 48 U kg ⁻¹ for 3 h at 55 °C, pH 8) and thermolysin (enzyme/ substrate: $1/1600$ w/w for 5 h at 37 °C, pH 8)	1. ACE inhibitory activity increased when hydrolyzed by enzymes such as gastrointestinal proteases, alcalase, and thermolysin; 2. Insect protein contains antihypertensive components and can be applied to functional foods and nutraceuticals.	20
Cotton leafworm (<i>Spodoptera</i> littoralis)	Larvae	ND	1. Simulated gastric phase: pepsin (enzyme/substrate: 1/250 w/w for 2 h at 37 °C, pH 2). And simulated intestinal phase: trypsin and α -chymotrypsin (1 : 1) (enzyme/substrate: 1/250 w/w for 2.5 h at 37 °C, pH 6.5); 2. Alcalase (48 U kg ⁻¹ for 3 h at 55 °C and at pH 8) and thermolysin (enzyme/substrate: 1/1600 w/w for 5 h at 37 °C, pH 8)	1. A new tripeptide Ala-Val-Phe was identified to have ACE inhibitory ability ($IC_{50} = 2123 \mu$ M); 2. The cotton leafworm (<i>S. littoralis</i>) hydrolysates (hydrolyzed with gastrointestinal enzymes) showed ACE inhibition activity; 3. New applications as functional foods, dietary supplements and antihypertensive agents.	79
Silkworm (<i>Bombyx mori</i>)	Larvae	ND	1. Pepsin (1750 U mg ⁻¹ for 90 min at 37 °C, pH 2.0), followed by hydrolysis with trypsin (2500 U mg ⁻¹ for 2.5 h at 37 °C, pH 6.5) and α -chymotrypsin (1000 U mg ⁻¹ for 2.5 h at 37 °C, pH 6.5)	1. The EAA composition of silkworm larvae protein isolate is balanced, and it is a high-quality protein source; 2. Silkworm larvae protein isolates displayed ACE inhibitory activity ($IC_{50} = 8.3 \ \mu g \ mL^{-1}$), DPPH scavenging activity ($IC_{50} = 57.91 \ \mu g \ mL^{-1}$) and ferrous ion chelating capacity ($IC_{50} = 2.03 \ m g \ mL^{-1}$).	143
Silkworm (<i>Bombyx mori</i>)	Pupae	ND	1. Pepsin (1000 U mg ⁻¹ for 1.5 h at 37 °C, pH 2.0). Trypsin and α -chymotrypsin (1000 U mg ⁻¹ for each enzyme for 2.5 h at 37 °C, pH 6.5).	1. A novel ACE-inhibitory tripeptide (Ala-Ser- Leu) (IC ₅₀ = 102.15 μ M) was isolated, showing competitive inhibition behavior; 2. The docking complex was stabilized by amino acids (Lys453, Asp415, His383, Val380, Val397, Ala354, His353, Gln281) in the ACE active site. 3. Strong hydrogen bonds formed by Ala354 and Gln281 and His353 are the main contributors to the ACE inhibition.	112
Cricket (Gryllodes sigillatus); Mealworm (Tenebrio molitor); Locust (Schistocerca gregaria)	Adult; Larvae; Adult	70.0 ± 1.7 (dry basis); 52.35 ± 1.1 (dry basis); 76.0 ± 0.9 (dry basis)	1. Stimulated saliva: α -amylase (50 U mg ⁻¹ , enzyme : substrate 1 : 10 w/w for 10 min at 37 °C, pH 6.75); 2. Simulated gastric phase: pepsin (250 U mg ⁻¹ , enzyme : substrate 1 : 100 w/w for 2 h at 37 °C, pH = 2.5); 3. Simulated intestinal phase: 0.7% pancreatin (for 1 h at 37 °C, pH 7.0) and 2.5% solution of bile extract (1 : 2.5 v/v)	1. Baked cricket (<i>Gryllodes sigillatus</i>) showed the highest degree of hydrolysis (37.76%); 2. The hydrolysates of raw, cooked and baked crickets, mealworm, and locust showed inhibition of human skin fibroblasts CRL-2522 (cytotoxicity).	5
Silkworm (<i>Bombyx mori</i>)	Рирае	ND	1. Stimulated saliva: α -amylase (50 U mg ⁻¹ , enzyme: substrate 1: 10 w/w for 10 min at 37 °C, pH 6.75); 2. Simulated gastric phase: pepsin (250 U mg ⁻¹ , enzyme:substrate 1: 100 w/w for 2 h at 37 °C, pH = 2.5); 3. Simulated intestinal phase: 0.7% pancreatin (for 1 h at 37 °C, pH 7.0) and 2.5% solution of bile extract (1: 2.5 v/v)	1. The peptides from silkworm pupa showed antiradical activity (<i>via</i> ion chelation); 2. The highest peptide concentration before digestion was 3.13 mg mL^{-1} for locust, but after digestion locust peptide concentration was still highest at 5.88 mg mL ⁻¹ .	77
Mealworms (Tenebrio nolitor); Locusts (Schistocerca gregaria); Crickets (Gryllodes sigillatus)	Larvae; Adult; Adult	ND	1. Stimulated saliva solution: 7 mM NaHCO ₃ and 0.35 mM NaCl, pH 6.75, for 10 min; 2. Stimulated gastric phase: pepsin (250 U mg ⁻¹ for 2 h at 37 °C, pH 2.5); 3. Simulated intestinal phase: 0.7% pancreatin and 2.5% bile extract (1 : 2.5, v/v) and incubated at 37 °C for 1 h.	1. Heat treatment increased LOX and COX-2 (referred to as anti-inflammatory) inhibition in mealworm, locusts and crickets; 2. Anti-inflammatory properties were shown in both whole insect hydrolysates and insect protein hydrolysates. The highest anti-inflammatory activity was shown in cricket (LOX: $IC_{50} = 0.13 \ \mu g mL^{-1}$; COX-2: $IC_{50} = 0.26 \ \mu g mL^{-1}$); 3. The highest antioxidant activity: highest Fe ²⁺ chelating (EC ₅₀ = 2.21 \ \mu g mL^{-1}) shown in mealworm protein hydrolysates. Highest ABTS and DPPH scavenging shown in cricket protein hydrolysates (EC ₅₀ = 2.75 \ \mu g mL^{-1} and 6.91 \ \mu g mL^{-1}, respectively).	104
Mealworms (Tenebrio molitor); Locusts (Schistocerca gregaria); Crickets (Gryllodes sigillatus)	Larvae; Adult; Adult	ND	1. Stimulated saliva solution: 7 mM NaHCO ₃ and 0.35 mM NaCl, pH 6.75 for 10 min; 2. Stimulated gastric phase: pepsin (250 U mg ⁻¹ for 2 h at 37 °C, pH 2.5); 3. Simulated intestinal phase: 0.7% pancreatin and 2.5% bile extract (1: 2.5, v/v) and incubated at 37 °C for 1 h; 4. Simulated absorption process: a membrane (molecular weight cut-off 3.5 kDa) was used for dialyzing hydrolysates, for 1 h at 37 °C without light.	1.5 gregaria peptide fractions showed the highest enzyme inhibitory activities: ACE ($IC_{50} = 3.95 \ \mu g$ mL ⁻¹) for boiled samples, lipase ($IC_{50} = 9.84 \ \mu g$ mL ⁻¹) for baked, and α -glucosidase ($IC_{50} = 1.89 \ \mu g$ mL ⁻¹) for raw <i>S. gregaria</i> , respectively; 2. The sequences of synthesized peptides with highest inhibitory activity were KVEGDLK, YETGNGIK, AIGVGAIR, IIAPPER, and FDPPPK; 3. When apply- ing heat treatment to edible insects, it led to posi- tive effects on the enzyme's inhibitory activity of peptides produced.	97

Table 3 (Contd.)

Source	Life stage	Protein content (starting material)	Enzymatic conditions	Main highlights	Rei
Muga silkworm (Antheraea assamensis)	Pupae	ND	1. Alcalase (enzyme/substrate: 0.5% v/w at 50 °C, pH 8.0), flavourzyme (enzyme/substrate: 1.5% v/w at 50 °C, pH 7.0), papain (enzyme/substrate: 2.3% v/w at 60 °C, pH 6.0) and thermolysin (enzyme/substrate: 3.0% v/w at 70 °C, pH 8.0). Aliquots were taken at 0, 1, 2, 3, 5, 8, 12, 24 h to measure bioactive properties.	1. The highest ACE inhibition activity was obtained when alcalase and papain were used at 3% enzyme-substrate ratio and incubated for 3 h; 2. 3% enzyme-substrate ratio and incubated for 5 h resulted in the highest DPPH scavenging activities; 3. Flavourzyme and papain enzymatic system at 2.3% enzyme- substrate ratio and incubated for 5 and 3 h, respectively, resulted in the highest DPP-IV inhibition activity.	30
Mealworm (<i>Tenebrio molitor</i>)	Larvae	ND	1. Alcalase (enzyme/substrate: 1 : 100 w/w at 50 °C, pH 8.5)	1. Tenebrio molitor (L.) larvae can be used as a source of ACE inhibitory peptides after hydrolysis with alcalase; 2. Single oral administration of ACE inhibitory peptide (Tyr- Ala-Asn) showed a significant reduction of systolic blood pressure in rats; 3. RP-HPLC was applied to purify Tyr-Ala-Asn (novel ACE inhibitory peptide); 4. The <i>in vivo</i> antihypertensive activity of Tyr-Ala- Asn needs to be further confirmed.	60
Cricket (Gryllodes sigillatus)	ND	ND	1. Establish nine conditions to test different enzyme-substrate concentrations and hydrolysis time on protein functionality. Alcalase (enzyme : substrate: 0.5, 1.5, and 3% w/w for 30, 60, and 90 min at 50 °C, respectively)	1. Whole cricket can produce protein hydrolysates with improved functionality; 2. The protein solubility can be improved over a wide range of pH, so the cricket protein hydrolysates have potential to be applied in acidic foods; 3. Higher foamability was observed for protein hydrolysates, but the foam stability was not as good as the non- hydrolyzed protein.	52
Cotton leafworm (<i>Spodoptera</i> <i>littoralis</i>)	Larvae	ND	1. Alcalase (48 U kg ⁻¹ for 3 h at 55 °C, pH 8); thermolysin (enzyme/substrate: 1/1600 w/w for 5 h at 37 °C, pH 8); 2. To simulate the human gastrointestinal digestion process as follows: pepsin (enzyme/substrate: 1/250 w/w for 2 h at 37 °C, pH 2) and small intestine: trypsin and α -chymotrypsin (enzyme/ substrate: 1/250 w/w for 2.5 h at 37 °C, pH 6.5); 3. Hydrolysis with mucosal peptidases (enzyme/substrate: 1/500 w/w for 2 h at 37 °C); 4. Different com- binations of hydrolysis were used: a. Gastrointestinal digestion + mucosal peptidases; c. Digestion with thermolysin; d. Digestion with thermolysin + gastrointestinal digestion; e. Digestion with thermolysin + gastrointestinal digestion + mucosal peptidases; f. Digestion with alcalase; g. Digestion with alcalase + gastrointestinal diges- tion; h. Digestion with alcalase + gastro- intestinal digestion + mucosal peptidases.	1. The S. littoralis hydrolysates showed both ACE inhibitory activity and antioxidant activity (<i>in vitro</i>), but these showed no correlation; 2. The S. littoralis hydrolysates showed a relatively low antioxidant activity; 3. Gastrointestinal digestion ($IC_{50} = 320 \ \mu g \ mL^{-1}$) and digestion with mucosal enzyme ($IC_{50} =$ 211 $\mu g \ mL^{-1}$) showed higher ACE inhibitory activity compared to alcalase hydrolysis ($IC_{50} =$ 827 $\mu g \ mL^{-1}$), thermolysin hydrolysis ($IC_{50} =$ 1392 $\mu g \ mL^{-1}$).	76
Lesser mealworm (A. diaperinus)	ND	ND	peptidases. 1. Alcalase and the enzyme corolase PP (enzyme : substrate: 0.5, 1.5 and 3.0% v/w, 50 °C, aliquots at different time points were taken 0, 30 min, 1, 2, 4, 6, 8 and 24 h, pH 8.0).	1. The enzymatic hydrolysis process increased the nutritional value of insect powder such as improving the free amino acid content and small peptides; 2. The hydrolysates have antioxidant (TEAC value: $95 \pm 0.8 \mu$ mol TE per g) and antihypertensive properties (IC ₅₀ = $55.5 \pm 6.2 \mu$ g mL ⁻¹); 3. No antimicrobial activity or α -glucosidase inhibition activity were found.	15
Cricket (Gryllodes sigillatus)	Adult (6 weeks old)	ND	1. Establish nine hydrolysis conditions: alcalase (enzyme : substrate: 0.5, 1.5, and 3% w/w for 30, 60, and 90 min, respectively at 50 °C, pH 8.0); 2. Pepsin (enzyme : substrate: 4% w/w for 2 h at 37 °C, pH 2), then bile salts and pancreatin (enzyme : substrate: 4% w/w for 2 h at 37 °C, pH 6.8)	1. Hydrolysates displayed good ACE ($IC_{50} = 0.062 \text{ mg mL}^{-1}$), DPP-IV inhibition (69%), and antioxidant activity (ABTS assay: 799.4 ± 8.2 µmol TE per mg sample; DPPH assay: 1926.3 ± 4.6 µmol TE per mg sample); 2. Bioactivity increased after simulated gastroin- testinal digestion; 3. A degree of hydrolysis between 60 and 85% can completely remove allergenicity.	23

Food & Function

Table 3 (Contd.)

for 5 h at 55 °C, PI 8.0); 2. Plavourzyme (enzyme: substrate: 4% w/w for 5 h at 55 °C, PI 7.0); 3. Protamex (enzyme: substrate: 4% w/w for 5 h at 40 °C, PI 7.0); 4. Trypsin (enzyme: substrate: 4% w/w for 5 h at 40 °C, PI 7.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, PI 7.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, PI 7.0); 6. Pepsin (enzyme: substrate: 4% w/w for 5 h at 50 °C, PI 7.0); 6. Pepsin (enzyme: substrate: 4% w/w for 5 h at 40 °C, PI 2.0).i. The α-Pa fraction showed the highest ACE inhibition before purification; after purifi- cation with RP-HPLC and PHLC, α-Pa-6-b showed the highest ACE inhibition; 2. Good solubility, heat and acid resistance was shown in purified pupa angiotensin 1-converting enzyme inhibitory activity of silkworm pret and acid resistance was shown in purified pupa angiotensin 1-converting enzyme inhibitory activity of silkworm pret and acid resistance was shown in purified pupa angiotensin 1-converting enzyme inhibitory activity of silkworm pupae and into silve for 50 min, pH 9.0. The ulfrasound-pretreated (20 kHz, 250 W to 600 W, 24 min) protein solution was adjusted with distilled water to 1% (w/v) at 50 °C for 10 min before hydrolysis.1. Microwave-assisted enzymatic hydrolysis by alcalase. Alcalase (7.54 U); enzyme is ubstrate: 3% for 10 min before hydrolysis.1. Microwave-assisted enzymatic hydrolysis by alcalase. Alcalase (7.54 U); enzyme: substrate: 3% for 10 min before hydrolysis.1. Microwave and enzyme is ubstrate: 3% for 10 min penzyme is ubstrate: 3% for 10 min or 20 min at 55 °C, pH 8.0. Microwave uni with warping power (600 W; 80% power1. Microwave and pret-1000000	Source	Life stage	Protein content (starting material)	Enzymatic conditions	Main highlights	Re
Silkworm (Bombyx mori) Pupae 66.83% 1. Acid protease (enzyme: substrate: 1% w/ for 4.92 h at 35 °C, pH 2.18). b) Molumin was the cassister protein fractions (g10)(s)s (17.32%) and ACE inhibitory activity (g13)(s) (and head there in the protein fractions (g10)(s)s (17.32%) and ACE inhibitory activity (g13)(s) (and head there in the protein fractions (g10)(s) (and fractions) and the protein fractions (G2m 0.047 m gmt ⁻¹). Silkworm (Bombyx mori) Pupae ND 1. Acid protease (protein/water, 1: 7 (g10), 3% acid protease (EX = 3000) b) Molecone (G2m 0.047 m gmt ⁻¹). Silkworm (Bombyx mori) Pupae ND 1. Acid protease (protein/water, 1: 7 (g10), 3% acid protease (EX = 3000) b) Molecone protein hydrolysates (G2m 0.047 m gmt ⁻¹). Silkworm (Bombyx mori) Pupae ND 1. Acida protease (enzyme: substrate: 4% wire (enzyme: substrate: 4% wire for 5 h at 5 °C, pH 1.0.0 b) Molecone (G2m 0.047 m gmt ⁻¹). Silkworm (Bombyx mori) Pupae ND 1. Acalase (enzyme: substrate: 4% wire for 5 h at 5 °C, pH 1.0, 10, 2. Flavours (M or 5 h at 5 °C, pH 1.0, 10, 2. Flavours (M or 5 h at 40 °C, pH 1.0, 10, 10, 20	Cricket (Gryllus bimaculatus);	Adult;	ND	and 60 U per g protein; b) Alcalase with 12 and 72 mU per g protein; c) Neutrase with 4 and 24 mU per g protein; d) Protamex with 7.5 and 45 mU per g protein 2 A mixture group a) Flavourzyme 30 U per g protein with Alcalase 12 mU per g protein; b) Flavourzyme 60 U per g protein with Alcalase 72 mU per g protein All hydrolysis were performed at 55 °C for	solubility than the unhydrolyzed proteins. The highest solubility was presented for the sample cotreated with two enzymes (flavourzyme 60 U g ⁻¹ + alcalase 72 mU per g protein). The solubility of mealworm larvae showed a significant increase (30%–50%); 2. Alcalase hydrolysates were the best emulsifiers because of the high emulsifying stability. But the combination of flavourzyme and alcalase led to a decreased emulsifying property; 3. The greatest inhibition of ACE was shown for the alcalase-treated group; 4. The highest α -glucosidase inhibitory activity was shown for silkworm pupae treated with alca- lase and mealworms treated with the flavour- zyme and alcalase; 5. Anti-inflammatory activity was only observed for silkworm pupae,	70
Silkworm (Bombyx mort) Pupae ND 1. Acid protease (protein/water, 1: 7, welv, 3% ocid protease (ES = 3000 U g^-1) for 5.0 h at 35 °C, pH 2.0 5. Silkworm (Boodyx mort) Silkworm (Bombyx mort) Pupae ND 1. Alcalase (enzyme : substrate: 4% welve for 5 h at 35 °C, pH 8.0); 2. Flavourphytersites in a long-term test. Silkworm (Bombyx mort) Pupae ND 1. Alcalase (enzyme : substrate: 4% welve for 5 h at 35 °C, pH 8.0); 2. Flavourphytersites in a long-term test. 1. Alcalase (enzyme : substrate: 4% welve for 5 h at 35 °C, pH 8.0); 2. Flavourphytersites in a long-term test. Silkworm (Bombyx mort) Pupae ND 1. Alcalase (enzyme : substrate: 4% welve for 5 h at 35 °C, pH 8.0); 2. Flavourphytersites in a long-term test. 1. Algi degree of hydrolysis sex secssary to reach a high antioxidant activity of the protein hydrolysates in a long-term test. Silkworm (Bombyx mort) Pupae ND 1. Trye in (1, 7, 7), 8, 10; 2, 10; 10; 10; 10; 10; 10; 10; 10; 10; 10;	ilkworm (<i>Bombyx mori</i>)	Pupae	66.83%		1. Albumin was the easiest protein fraction to hydrolyze showing a higher degree of hydrolysis (17.32%) and ACE inhibitory activity (81%) than the other three protein fractions (prolamin, globulin and glutelin) obtained by the Osborne method; 2. A peptide sequence APPPKK, which inhibits the angiotensin I-converting enzyme activity, was identified in the albumin fraction; 3. The highest ACE inhibitory activity was in silkworm albumin	14
Silkworm (Bombyx mori)PupaeND1. Alealase (nzyme : substrate: 4% w/w for 5 h at 55 °C, pl f 3.0); 2. Flavourzyme (enzyme: substrate: 4% w/w for 5 h at 55 °C, pl f 3.0); 2. Flavourzyme (enzyme: substrate: 4% w/w for 5 h at 55 °C, pl f 3.0); 3. Protamex (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl f 3.0); 5. Papain (enzyme: substrate: 4% w/w for 5 h at 50 °C, pl 6.0)1. The α high antoxidant activity was 66.1%.Silkworm (Bombyx mori)PupaeND1. Ultrasound-assisted enzymatic hydrolysis by alcalase. Al	iilkworm (<i>Bombyx morî</i>)	Pupae	ND	w/v), 3% acid protease (E/S = 3000 U	1. Silkworm pupae protein hydrolysates showed <i>in vivo</i> ACE inhibition (blood pressure of rats was 139.0 ± 10.4 mmHg after consumption and after 6 h, blood pressure was recovered); 2. Acute toxicity research showed that protein hydrolysates were safe for human consumption; 3. The systolic blood pressure of non-hypertensive rats is not	14
Silkworm (Bombyx mori)PupaeND1. 7% alkaline protease (for 4 h at 50 °C, pH 9.5).1. The α-P ₃ fraction showed the highest ACE inhibition before purification; after purifi- cation with RP-HPLC and HPLC, α-P ₃ -6-b showed the highest ACE inhibition; 2. Good solubility, heat and acid resistance was shown in purified pupa angiotensin 1-converting enzyme inhibitory peptides.Silkworm (Bombyx mori)PupaeND1. Ultrasound-assisted enzymatic hydrolysis by alcalase. Alcalase (3500 U per g protein) at 50 °C for 50 min, pH 9.0. The ultrasound-pretreated (20 kHz, 250 W to 600 W, 24 min) protein solution was adjusted with distilled water to 1% (w/v) at 50 °C for 10 min before hydrolysis.1. Microwave-assisted enzymatic hydrolysis by alcalase. Alcalase (7.54 U): enzyme : substrate: 3% for 10 min or 20 min at 55 °C, pH 8.0. Microwave uni with varying power (600 W; 80% power1. Microwave-assisted enzymatic hydrolysis by alcalase. Alcalase (7.54 U): enzyme: substrate: 3% for 10 min or 20 min at 55 °C, pH 8.0. Microwave uni 	ilkworm (<i>Bombyx morî</i>)	Pupae	ND	for 5 h at 55 °C, pH 8.0); 2. Flavourzyme (enzyme:substrate: 4% w/w for 5 h at 55 °C, pH 7.0); 3. Protamex (enzyme : substrate: 4% w/w for 5 h at 40 °C, pH 7.0); 4. Trypsin (enzyme : substrate: 4% w/w for 5 h at 40 °C, pH 8.0); 5. Papain (enzyme : substrate: 4% w/w for 5 h at 50 °C, pH 7.0); 6. Pepsin (enzyme : substrate: 4% w/w for 5 h at	1. A high degree of hydrolysis was necessary to reach a high antioxidant activity of the protein hydrolysates; 2. The optimum conditions for silkworm protein alcalase hydrolysis were E/S 7.38%, pH 7.97 at 60 °C, and the maximum	69
Silkworm (Bombyx mori)PupaeND1. Ultrasound-assisted enzymatic hydrolysis by alcalase. Alcalase (3500 U per g protein) at 50 °C for 50 min, pH 9.0. The ultrasound-pretreated (20 kHz, 250 W to 600 W, 24 min) protein solution was adjusted with distilled water to 1% (w/v) at 50 °C for 10 min before hydrolysis.1. The ACE inhibitory activity of silkworm pupae can be improved by ultrasound pretreatment; it showed a 39.9%–67.3% increase (ultrasonic power 250–600 W) compared to non-ultrasonic treatment; 2. A novel peptide (Lys-His-Val) resistant to gastrointestinal proteases exhibited ACE inhibitory activity (IC 50 = 12.82 µM); 3. The active site of ACE could effectively interact with Lys-His-Val pretide as result of molecular docking analysis.Crickets (Gryllodes sigillatus)Adults (6 weeks old)ND1. Microwave-assisted enzymatic hydrolysis by alcalase. Alcalase (7.54 U): hydrolysis by alcalase. Alcalase (7.54 U): enzyme : substrate: 3% for 10 min or 20 min at 55 °C, pH 8.0. Microwave unit with varying power (600 W; 80% power1. Microwave-assisted enzymatic hydrolysis.	ilkworm (<i>Bombyx mori</i>)	Рирае	ND	1. 7% alkaline protease (for 4 h at	inhibition before purification; after purifi- cation with RP-HPLC and HPLC, α - β_3 -6-b showed the highest ACE inhibition; 2. Good solubility, heat and acid resistance was shown in purified pupa angiotensin I-converting	14
$ \begin{array}{c} \mbox{Crickets} (Gryllodes sigillatus) \\ \mbox{Crickets} (Gryllodes sigillatus) \\ \mbox{weeks old} \\ \mbox{Weeks old} \\ \mbox{Weeks old} \\ \mbox{H} \\ H$	ilkworm (<i>Bombyx morî</i>)	Pupae	ND	hydrolysis by alcalase. Alcalase (3500 U per g protein) at 50 °C for 50 min, pH 9.0. The ultrasound-pretreated (20 kHz, 250 W to 600 W, 24 min) protein solution was adjusted with distilled water to 1% (w/v) at 50 °C for 10 min	1. The ACE inhibitory activity of silkworm pupae can be improved by ultrasound pretreatment; it showed a 39.9%–67.3% increase (ultrasonic power 250–600 W) compared to non-ultrasonic treatment; 2. A novel peptide (Lys-His-Val) resistant to gastrointestinal proteases exhibited ACE inhibitory activity (IC $_{50}$ = 12.82 µM); 3. The active site of ACE could effectively interact with Lys-His-Val peptide as result of molecular	61
maximum).	Crickets (Gryllodes sigillatus)		ND	hydrolysis by alcalase. Alcalase (7.54 U): enzyme : substrate: 3% for 10 min or 20 min at 55 °C, pH 8.0. Microwave unit	1. Microwave-assisted hydrolysis led to highest inhibition of ACE ($IC_{50} = 0.096 \text{ mg mL}^{-1}$) and DPP-IV ($IC_{50} = 0.27 \text{ mg mL}^{-1}$); 2. Low immuno- reactivity peptides can be generated through	21

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trase, protamex, acid protease and alkaline protease were also applied. Some studies combined different enzymes to achieve increased bioactivity. Vercruysse *et al.* (2009) combined gastrointestinal digestion and thermolysin, alcalase or mucosal peptidases and found that the combination of these enzymes increased ACE inhibition, for example, gastrointestinal digestion (with mucosal peptidases) significantly lowered the IC_{50} value to 211 µg mL⁻¹ and showed the best ACE inhibitory ability, compared to gastrointestinal digestion ($IC_{50} = 320 \ \mu g$ mL⁻¹).⁷⁶ Similarly, insect protein hydrolyzed by gastrointestinal proteases, alcalase, and thermolysin led to an increase of ACE inhibitory activity with IC_{50} values of 1.26 mg mL⁻¹ for *B. terrestris* and 0.77 mg mL⁻¹ for *S. littoralis*, compared to 43.22 mg mL⁻¹ and 2.58 mg mL⁻¹ before hydrolysis.²⁰

Combining conventional hydrolysis with techniques such as microwave and ultrasound treatment can achieve a lower IC₅₀.⁷⁶ Hall and Liceaga (2020) reported that the microwavehydrolyzed G. sigillatus protein had the highest ACE inhibition $(IC_{50} = 0.096 \text{ mg mL}^{-1})$ compared to enzymatically hydrolyzed protein $(IC_{50} = 0.20 \text{ mg mL}^{-1})$.²¹ Similar to microwave treatment, ultrasound treatment is another good alternative for hydrolyzing proteins and increasing ACE inhibition, as the treatment induces the molecular unfolding of proteins and increases the surface hydrophobicity, which promotes the release of peptides with an ACE inhibitory function. However, parameters for ultrasound hydrolysis must be optimized as sonication with high power and long duration leads to the formation of a stable structure and hinders the release of ACE-inhibiting peptides.^{61,91} Table 3 summarizes the common enzymes and suitable conditions applied to insect protein hydrolysis.

More specifically for ACE inhibitory peptides (ACEIPs), their inhibitory activity is related to peptide properties such as hydrophobicity, charge, and structural conformation. ACE inhibition activity is improved by the presence of hydrophobic rings in structural amino acids.⁹² Similarly, another study also demonstrated the importance of aromatic amino acids in ACE inhibitory peptides.⁹³ Peptides such as IIe-Pro-Pro and Val-Pro-Pro with a proline ring at the C-terminus were recognized as potent ACEIPs. Zhang *et al.* (2020) also observed that a C-terminal hydrophobic amino acid was more effective than an N-terminal one in ACE inhibition.^{92,94} Known antihypertensive peptides AF, GW, GY and PH in *M. domestica* and GKDAVIV and VAPEEHPV in *S. gregaria* also agreed with the fundamental features of ACEIPs, and thus displayed great antihypertensive activities.^{95,96}

However, only limited *in vivo* studies confirmed the antihypertensive activity of insect-derived peptides. Dai *et al.* (2013) reported a significant decrease in blood pressure when applying multiple dose oral administration of protein hydrolysates of *T. molitor* to spontaneously hypertensive rats; the systolic blood pressure decreased to 27 mm Hg at 400 mg per kg weight after 4 h post-administration.⁶⁰

3.2. Antidiabetic properties

High postprandial blood glucose is caused by DPP-IV, but to date, the application of DPP-IV inhibitors is expensive and

clinicians lack experience.97 Some publications reported the DPP-IV inhibition activity of insect protein hydrolysates obtained with various enzymes. For enzymatic hydrolysis by simulated gastrointestinal digestion, G. sigillatus protein hydrolysates showed low inhibition between 0 to 50% (depending on the enzymatic hydrolysis conditions used) before simulated gastrointestinal digestion, but after simulated gastrointestinal digestion the inhibition activity increased and reached values between 62 and 69%.²³ In other insect species, DPP-IV inhibitory properties were reported in A. assamensis protein hydrolysates (in gastric fluid $IC_{50} = 5063 \ \mu g \ mL^{-1}$ and intestinal fluid IC₅₀ = 5221 μ g mL⁻¹), but the highest inhibition was found when applying flavourzyme and papain and hydrolyzing at a 2.3% enzyme to substrate ratio for 3 h.³⁰ This may suggest that for A. assamensis, flavourzyme combined with papain shows higher efficiency at generating antidiabetic peptides due to the specific cleavage of hydrophobic amino acid residues. Similar to the antihypertensive properties, antidiabetic properties can also be increased by microwave treatment; a low IC50 value was observed for the microwave plus alcalase treated group (IC₅₀ = 0.27 mg mL^{-1}) when compared to the group subjected to alcalase hydrolysis only (IC₅₀ = 0.65 mg mL⁻¹).^{21,98} However, a limited number of publications have reported the α-glucosidase inhibitory activity of insect protein hydrolysates or insect-derived peptides, which helps in delaying digestion of carbohydrates, thereby reducing the levels of glucose in blood. Yoon *et al.* (2019) reported α -glucosidase inhibitory activity for T. molitor, G. bimaculatus and B. mori hydrolysates and found that the most effective inhibitors were those from silkworm pupae treated with alcalase and mealworms treated with flavourzyme coupled to alcalase hydrolysis, which showed approximately 33% and 35% inhibition, respectively.⁷⁰ But to date the in vivo antidiabetic properties of insect protein hydrolysate have not been assessed.

In general, antidiabetic peptides contain 2 to 17 amino acid residues, with most of them being hydrophobic amino acids. However, antidiabetic peptides with hydrophilic amino acids such as 'Thr, His, Gln, Ser, Lys, Arg' can also be found. For example, antidiabetic dipeptides presenting 'Trp/Thr/Met at the N-terminus' and 'Ala/Leu/His at the C-terminus' and peptides containing 'Pro/Leu/Arg at the C-terminus' have been identified previously.⁹⁹ The *G. assimilis* derived peptides identified as α -glucosidase inhibitors were 'LAMVEA, LPPPP, ALLVVW, DSYPL and YPGDV'. Thus, these specific traits correspond to the general features of antidiabetic peptides (over 4 amino acids), with hydrophobic amino acids as dominant residues.

3.3. Antioxidant properties

Antioxidant peptides can be released in several ways, including chemical hydrolysis, enzymatic hydrolysis, novel technologies coupled to assisted enzymatic hydrolysis (*i.e.* ultrasound, pulsed electric fields, *etc.*), fermentation and other food processing methods (*i.e.* steaming, boiling, *etc.*).⁹⁵ For instance, insect studies have demonstrated the potential of enzymatic hydrolysis to release antioxidant peptides.^{89,99,100} The release

of antioxidant peptides by alcalase from B. mori protein was reported; the enzyme-substrate ratio, pH and temperature were tailored to 7.38%, 7.97 and 60 °C, respectively, to achieve the maximum antioxidant activity (66.1%).⁶⁹ However, Hall et al. (2018) evaluated the antioxidant activity of G. sigillatus protein hydrolysates, but found that simulated gastrointestinal digestion had little effect on the antioxidant activity, as this was similar before and after digestion. For example, for the ferric ion reducing antioxidant power assay and ABTS assay, cricket protein alcalase hydrolysates showed 516.8 ± 13.5 and 906.6 \pm 4.1 µmol TE per mg sample, compared to 689.0 \pm 13.8 and 1137.4 \pm 2.8 µmol TE per mg sample after gastrointestinal digestion.²³ For G. sigillatus, a possible explanation for the stable antioxidant activity is that antioxidant fragments mainly have β -sheet and random coil structures rather than α -helices, although the current study does not include the secondary structure analysis of G. sigillatus antioxidant fragments. Another study showed that G. sigillatus water-soluble proteins were dominated by the α -helix secondary structure, which was hard to break down during digestion, thus, resulting in the negligible generation of peptides with antioxidant properties.101

The relationship between antioxidant and antihypertensive properties has also been studied. Vercruysse *et al.* (2009) showed that the hydrolysates of *S. littoralis* exerted dual activity as both *in vitro* ACE inhibition and antioxidant activity were observed, but no relationship was found between ACE inhibition and antioxidant activity.⁷⁶ The inexistent link between antioxidant and antihypertensive activities was also found in *A. diaperinus* when hydrolyzed by alcalase coupled to corolase PP.¹⁵ However, as for the antidiabetic properties, the *in vivo* antidiabetic properties of insect protein hydrolysates have not been extensively studied.

The amino acid composition is an important decisive factor for the release of antioxidant peptides with enhanced bioactivity. Peptides with a high presence of hydrophobic amino acids are regarded as peptides with higher radical scavenging potential than peptides with a lower occurrence of hydrophobic amino acids.¹⁰² Known antioxidant peptide sequences such as APVAVAHAAVPA and ASVVEKLGDY in A. diaperinus and LAPSTIK in G. sigillatus also align with this feature.97,103 Regarding these known sequences, the high content of hydrophobic amino acids may explain why edible insects have a higher antioxidant activity than the hydrolysates obtained from some animal products or leafy plants after digestion.⁷⁷ Also, the content of hydrophobic amino acids in insects such as H. illucens, A. domesticus and M. bellicosus is comparable to that of animal products such as milk, beef and lamb, and is higher than for plant materials (Fig. 1).

3.4. Anti-inflammatory properties

The anti-inflammatory properties of insects have not been widely researched. The lipoxygenase (LOX) and cyclooxygenase (COX-2) inhibitory activities are target anti-inflammatory biomarkers. According to limited studies, the anti-inflammatory activities seem to be highly dependent on the hydrolysis conditions.⁷⁰ For example, G. sigillatus protein hydrolysates showed LOX and COX-2 inhibitory activity with IC50 values of 0.13 and 0.26 $\mu g m L^{-1}$, respectively, compared to T. molitor protein hydrolysates (LOX: $IC_{50} = 0.17 \ \mu g \ mL^{-1}$; COX-2: $IC_{50} =$ 0.35 μ g mL⁻¹) and *S. gregaria* protein hydrolysates (LOX: IC₅₀ = 0.18 µg mL⁻¹; COX-2: IC₅₀ = 0.26 µg mL⁻¹).¹⁰⁴ However, different results were reported by Yoon et al. (2019).⁷⁰ Neither T. molitor protein nor T. molitor protein hydrolysates showed anti-inflammatory properties when hydrolysed with flavourzyme and alcalase. Furthermore, heat processing such as boiling and baking can increase the anti-inflammatory activity of the whole insect hydrolysates produced by *in vitro* digestion. T. molitor, G. sigillatus and S. gregaria showed a significant increase of LOX and COX-2 inhibition after heat treatment, for example, when boiling was applied to G. sigillatus flour, the IC_{50} value (LOX) decreased to 16.9 µg mL⁻¹ (compared to raw cricket flour IC₅₀ = 20.74 μ g mL⁻¹).¹⁰⁴

Several studies have shown that low molecular weight peptides are more resistant to digestion, and therefore, can easily cross the intestinal barrier and be absorbed in the intestine. Longer peptides such as lunasin, containing 43 amino acids, are also anti-inflammatory peptides, but their mechanism of action is not clear so far.¹⁰⁵ A limited number of anti-inflammatory peptides were reported in *S. gregaria*, including IIAPPER, LAPSTIK and AIGVGAIER.¹⁰⁴ Interestingly, these peptides also have antioxidant properties. This may be attributable to the fact that similarly to antioxidant peptides, hydrophobic amino acids are also present in anti-inflammatory peptides. Normally, they are located at the N-terminal position, but more studies are needed to understand the underlying mechanisms.

4. Bioinformatic (*in silico*) tools for insect peptides prediction

Bioinformatic (in silico) analysis is based on the use of a peptide database for the identification, selection and prediction of potential bioactive peptides released from proteins during enzymatic hydrolysis.¹⁰⁶ The databases available contain various detailed information about peptides, including chemical structures, toxicity, allergenicity, sequences and amino acid contents (Fig. 3). The most common and widely used database is BIOPEP-UMV, while other databases such as UniProtKB and NCBI are mainly used to retrieve known sequences for analysis and comparative and homology purposes.¹⁰⁶ Currently, the in silico analysis of insect proteins and insect-derived peptides is mainly used as an aid/guidance for in vitro enzymatic hydrolysis. Most of the studies available simulate the in silico hydrolysis of proteins to mimic individual or sequential enzyme action, and then the selection of the most promising fraction/peptide (obtained from enzymatic hydrolysis) is proposed, and afterwards chemical synthesis of the selected peptide(s) is carried out to verify the activity in vitro. Finally, molecular docking analysis is used to confirm the mechanisms behind specific bioactivities.107 Insect-

derived peptides with antihypertensive and/or antidiabetic properties are the most studied ones by *in silico* analysis. Other peptides with antithrombotic and anti-SARS-CoV-2 properties were also reported with *in silico* tools.^{108,109} Currently, most of the insect-derived peptides discussed in section 3 in this review have been identified using the aforementioned databases.

Software like ExPASy-peptidecutter can be applied as a prediction tool before *in vitro* enzymatic hydrolysis. The substrate (protein sequence) and specific enzymes can be selected to simulate the cutting sites and generate the potential peptide profiles.¹⁰⁶ For example, eight major proteins from house fly larvae (scientific name not provided) were considered for simulated gastrointestinal digestion while the allergenic proteins from *B. mori* were simulated for pepsin and bromelain hydrolysis by *in silico* approaches to select functional peptides.^{96,107} Besides, *A. mellifera* peptides (generated by neutrase hydrolysis) with ACE inhibition activity were subjected to *in silico* digestion to simulate the effectiveness of peptides after human digestion.¹¹⁰

Then the potential mechanisms behind the bioactivities are determined by molecular docking, which is based on simulating the interaction between receptors and ligands (bioactive peptides) or small molecules.¹¹¹ For insects, molecular docking studies have mainly focused on the purified insectderived peptides and ACE; the inhibition of ACE is mainly due to hydrogen bond formation and Zn(II) interactions between peptides and ACE. An ACE inhibitor tripeptide (Ala-Ser-Leu) from B. mori was found to complex with ACE through more than fifteen hydrogen bonds, with the main binding sites being between Gln281 and His353 in the ACE S2 pocket and Ala354 in the S1 pocket, but this tripeptide has no interaction with the Zn(II) in ACE.¹¹² A similar study also reported Lys-His-Val as a potent and more effective ACE inhibition peptide, which formed eighteen hydrogen bonds with ACE, and similarly to lisinopril (a well-known ACE inhibitor used to treat high blood pressure), Lys-His-Val showed interactions at Gln281 and His 383.⁶¹ Another reported ACE inhibitor, *B. mori* peptide (GAMVVH), showed competitive coordination with Zn(II) in ACE; the stable structure of ACE was distorted when combining with GAMVVH, besides that, it could also form hydrogen bonds with Ala354, Lys511, and Gln281 in the ACE S1 and S2 pockets, which further helped to stabilize binding.¹¹³ For antidiabetic peptides purified from insect sources, they are known to inhibit DPP-IV by mainly forming hydrophobic interactions with the DPP-IV binding pocket. For example, Leu-Pro-Pro-Glu-His-Asp-Trp-Arg (from B. mori) is known to show hydrophobic interactions with Tyr510 and Phe320 in the DPP-IV S1 and S3 pockets, respectively. Similarly, hydrophobic interactions are the main interactions present between Leu-Pro-Ala-Val-Thr-Ile-Arg and the DPP-IV S1 binding pocket.¹⁰⁷ Additionally, for the antioxidant B. moriderived peptide (AKPGVY), it is known to be stabilized by hydrogen bonds, van der Waals interactions and hydrophobic interactions within the human Prxs binding site.¹¹⁴

Moreover, for antidiabetic insect peptides, an *in silico* modeling method, QSAR (quantitative structure-activity relationships), was applied to predict peptide bioactivities according to their sequence/structure.106 QSAR modeling was first applied to identify the α -glucosidase inhibiting peptides in silkworm (scientific name not available). The four silkwormderived peptides with strong inhibition activity were Gln-Pro-Gly-Arg (IC₅₀ = 65.8 μ mol L⁻¹), Ser-Gln-Ser-Pro-Ala (IC₅₀ = 20 μ mol L⁻¹), Gln-Pro-Pro-Thr (IC₅₀ = 560 μ mol L⁻¹) and Asn-Ser-Pro-Arg (IC₅₀ = 205 μ mol L⁻¹).⁴⁰ They are known to mainly complex with Lys776 through hydrogen bonds, and Lys776 is potentially the target amino acid for α -glucosidase inhibition peptides, but more studies are required. Nonetheless, in silico analyses are still needed in order to understand and elucidate the possible mechanisms of insect peptides and simulate the interactions between peptides and targeted proteins/ligands (through molecular docking) to predict the behaviour of peptides in regulating enzyme activities such as ACE, DPP-IV, α -glucosidase, inflammatory biomarkers, *etc.* There is a lack of published studies using bioinformatic tools to determine insect peptides released during digestion, evaluation of peptide stability or changes in biological activity due to different enzyme actions, and allergenicity prediction, among others.

5. Allergenic properties of insect proteins

Possible allergenic responses are becoming a major concern in insect consumption. Although allergens for novel foods are not unique, it is necessary to evaluate the potential risks of edible insect proteins.^{23,115}

A major panallergen, tropomyosin (TM), was identified in insects. Vertebrate-TM and invertebrate-TM shared 55% homology in terms of amino acid composition, and invertebrate-TM was not recognized as an allergen at first. However, as tropomyosin was gradually identified as an insect allergen, the cross-reactivity hypothesis among TMs from various species was supported.¹¹⁶ Thus, in order to reveal the potential allergenicity and mechanisms behind tropomyosin present in insects, a "bottom-up" approach reported by Zhao et al. (2023) should be carried out in order to analyze the potential allergic reactions among different insect species (from amino acid sequences to physicochemical properties, protein structures, epitopes and allergic peptides).¹¹⁷ Recently, the TM amino acid sequences of crustaceans and mollusks were compared; interestingly, 65% homology among the amino acid sequences was observed, suggesting cross-reactivity between the two species. The authors reported that allergenic reactions for crustacean allergic individuals could be triggered by some insects.^{100,118} So, potentially, insect TMs should also share, to higher or lower extents, homology in terms of amino acid sequence with crustaceans and/or mollusks. Research already proved that the cricket-TM shared over 60% homology in terms of amino acid sequence with known shellfish-TM.¹¹⁹ Furthermore, potentially, the allergenicity level of various insect species may depend on the amino acid sequence hom-

ology between the different tropomyosin isoforms that have shown highest allergenicity (species are unknown) and specific insect-TMs, as well as the amount of allergen-specific IgE present, the route by which the allergen is introduced and the dose of allergen. Regarding the physicochemical properties, both insect-TMs and invertebrate-TMs are resistant to heat and proteolysis. Although the tropomyosin helical secondary structure collapses when heated above 80 °C, it may reform upon cooling.^{25,26,120} However, the differences between the physicochemical properties of TMs from different species are not available. Additional studies regarding the epitopes and protein secondary structures of insect allergens are needed, in order to elucidate how these are related to potential allergenicity. For G. sigillatus-TM, 31 peptides were identified as potential linear epitopes, among them, the sequence 'RSQQDEERM' shared by peptides such as 'RSQQDEERMDQ, was RSQQDEERMDQLTNQ and NRSQQDEERMDQLTNQ'.120

Another allergen reported in invertebrates is arginine kinase (AK), an enzyme involved in energy metabolism and muscle mobilization.¹²¹ AK is resistant to heat and digestion. A recent study showed limited AK cross reactivity present between crickets and mealworm allergic subjects.^{122,123} This proved the non-systematic cross-reactivity of AK, thus providing knowledge on the extent to which these different antigens

appear similar or different to the immune system. However, the amino acid sequence of AK still shows high homology between different insect species, highlighting the importance of further studies on AK cross reactivity.

Conventional enzymatic hydrolysis and other processing methods such as microwave assisted-hydrolysis were investigated to lower the immunoreactivity and generate hypoallergenic peptides from insects.^{21,23,100,124} The decrease in allergenicity can be explained by the changes in the protein secondary structure; α -helices and β -sheets are stable structures and are hard to modify, and therefore, it makes it difficult for antibodies to combine.¹¹⁷ For example, aiming to lower the allergenicity of tropomyosin, 15 min microwave treatment (1000 W, 2.45 GHz) at 125 °C significantly reduced allergenicity by 75%. This decrease is associated with secondary structure modifications, including a decrease in α-helices and turns and an increase in β-sheets. The associated structural changes result in fewer recognition opportunities by the IgE of sensitized subjects.¹²⁵ In other work, Hall et al. (2020) also reported the differences in the tropomyosin-IgG binding capacity between conventional enzymatic hydrolysis and microwaveassisted hydrolysis. They found that the lowest tropomyosin-IgG binding was obtained with microwave-hydrolyzed G. sigillatus protein (Fig. 4).²¹ Although the epitopes associated

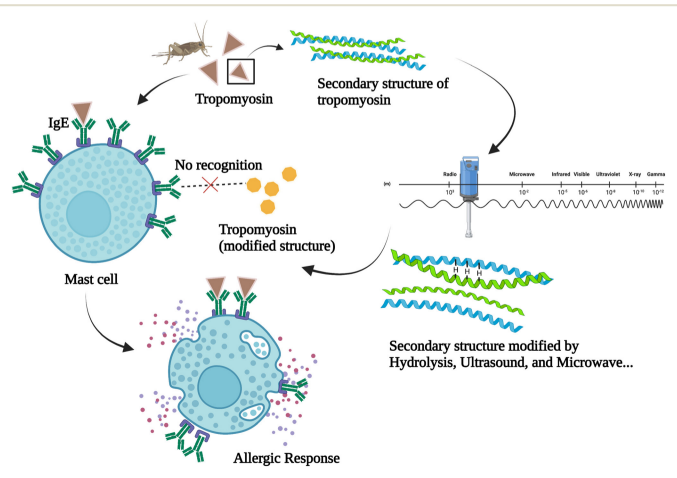


Fig. 4 Overview of the procedure to reduce the allergenicity of insect tropomyosin by hydrolysis, microwave and ultrasound treatment.

with reactivity are altered by processing and therefore are associated with decreased allergenicity, it is worth noting that new binding sites can be created and then induce potential sensitization and allergic responses.¹²⁶

6. Conclusion and future perspectives

This review has provided relevant evidence that insect proteins can be used as an alternative protein source. From the nutritional perspective, in terms of protein quality, it can be comparable to common protein sources (plant/animal) as the EAA content of well-studied insects is higher than that in plant materials such as peanuts and almonds and similar to that in beef and cow milk. Also, it is worth mentioning that the protein digestibility of edible insects has shown high values in *in vitro* assays and similar values to those of beef and pork.

Insect protein extraction is considered to be an effective method to improve the acceptance of insect intake. Wet fractionation is the most widely used method including traditional alkaline solubilization coupled to isoelectric precipitation or salt extraction and wet fractionation combined with some novel extraction techniques such as ultrasound and microwave treatments. Novel techniques not only have the potential to increase protein quality and the purity of protein ingredients, but also increase the bioactive traits of hydrolysates/peptides. Dry fractionation is less applied and additional studies are required to fully determine its potential to produce insect fractions.

Insects such as B. mori, T. molitor and G. sigillatus have potential for generating bioactive peptides, and more edible insects should be included in further studies to unlock more insect-derived peptides that could benefit human health. The main bioactivities that have been studied and identified in peptides and/or hydrolysates (in vitro) so far are antihypertensive properties, antidiabetic properties and antioxidant properties. Enzyme selection, amino acid composition/sequence, hydrophobicity and the structure of peptide fragments are commonly related to the potential bioactivity of peptides. Only a few anti-inflammatory properties have been researched and most of them have been studied at the protein hydrolysate stage instead of identifying the specific peptides exerting such activities. Also, mechanistic studies on anti-inflammatory insect-derived peptides are still missing, so more in silico studies are needed to reveal the mechanisms behind insect bioactive peptides and their specific inhibition sites of action. It is worth mentioning that current studies on protein identification and peptide bioactivities mainly focus on water-soluble proteins; the bioactivities of other protein groups with different solubility (salt soluble, organic solvents and alkali/ acid diluted solutions) could provide key knowledge to characterize and classify insect proteins in more depth and generate insights on novel peptide bioactivities.

To date, the allergenicity of insect proteins is still an urgent matter to study and explore. The side effects of these allergenic reactions can be decreased through enzymatic hydrolysis, and the use of other technologies such as ultrasound and microwave, which have the capacity to disrupt the secondary structure of certain allergens and as a result decrease their recognition by IgE, but it is important to note that new active sites can be generated during processing.¹²⁶ Also, the cross reactivity mechanisms of other less studied allergens found in insects, such as AK, still need to be elucidated. Edible insects may be safe for human consumption as they have been part of the human diet for many centuries in some countries. However, scientific research is still needed to protect consumer health and increase innovation and cutting-edge research on insect proteins, peptides and bioactive compounds that may exert health benefits for the wider population.

Author contributions

Zidan Ma: writing and editing. Alan Javier Hernández-Álvarez: supervision, reviewing and editing. Martin Mondor and Francisco Goycoolea Valencia: reviewing and editing.

Conflicts of interest

The authors declare no conflict of interest.

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