Entomophagy and Public Health: A Review of Microbiological Hazards

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Abstract
The world population’s constant increase and the continuous need to ensure food safety and security are among the major challenges to be faced in the next 30 years. In addition, human diet is evolving with a decreased inclusion of proteins from animal sources. In this context, consumption of insects by humans (entomophagy) could be an alternative solution to the intake of protein derived from conventional livestock, due to the lower environmental impact of insect rearing compared to traditional farming. Furthermore, various insect species have promising nutritional profiles regarding both macro and micronutrients. Nowadays, it is recognized that about 2 billion people consume insects at a worldwide scale, with more than 2000 different species to have been reported. Since the beginning of the 2000s, mass rearing of insects for human consumption has been developing all over the world. Nevertheless edible insects are foodstuffs of animal origin and are usually consumed in their entirety, including the digestive tract, meaning that they may contain biological agents with hazardous potential (e.g. bacteria, parasites, viruses, prions, yeasts, molds, mycotoxins, histamine, and antibiotic resistance genes) and they must undergo a thorough analysis. Therefore, establishing the synthesis of the current knowledge on entomophagy and the related biological hazards is the main purpose of this review.

Keywords
Entomophagy, Microbial Hazards, Public Health, Food Safety

1. Introduction
The future of food in the next decades will be highly influenced by the popula-
tion growth, the need to ensure sustainable food safety and food security and consumer trends like the decrease of intake of proteins from animal sources. The current model of production of animal proteins, mainly from cattle, swine, poultry and fish has been shown to have limitations and negative impacts on the environment. In this context, consumption of insects by humans (entomophagy) could be an alternative solution as insects appear to be a still under-exploited food source [1] [2] [3]. In fact, this relatively “new” source of animal proteins for western countries appears to have promising nutritional values and requires, for example, two to ten times less agricultural lands to produce one kilogram of insect protein compared to the production of one kilogram of protein from pigs or cattle, respectively [4] [5]. Nowadays, it is recognized that about 2 billion people consume insects in the world with more than 2000 different species to have been reported [3]. In some countries, insects are a culturally highly desirable food, especially in the tropics [6]. In other countries, particularly in occident, entomophagy is not yet fully accepted, neither culturally nor socially [7] [8]. At this stage, it is obviously necessary to differentiate between occasional or even personal insect consumption with consumption at the population level, which requires the development of mass rearing to produce animal proteins rapidly, at a lower cost and with a lower environmental impact [5].

This industrialization phase of production is underway in several developed countries, including some belonging to the European Union (e.g. the Netherlands, France, and Belgium). Careful choice of the insect species to be reared among the existing ones has to be done, in order to ensure the sustainability and profitability of these farms [8]. For mass rearing, the main criteria of choice are: the nutritional profile of the insect species, a high rate of reproduction and synchronization of nymphosies, an adapted ethology (no cannibalism) and great ease of handling with insects small in size that should not fly or jump when possible [9]. They must also have a high feed conversion rate (*i.e.* biomass gain per kg feedstock), with inexpensive but non-waste feed [10] [11] [12]. Once the choice is made, zootechnical parameters for rearing are adapted and controlled (temperature and ventilation of the premises, circadian rhythm or not, lighting...) [10].

Foods contaminated by biological or chemical hazards can cause outbreaks or sporadic cases in consumers. Thus, in many countries, foodstuffs are subject to various regulations and monitoring of their production conditions [13]. As such, it is very important to be able to quantify the relative contributions of different sources to the human disease burden; it is the role of source attribution studies that are currently developed [14]. In fact, several studies have shown in recent years, all over the world, that food of animal origin was responsible for more than 50% of reported food-borne infectious diseases outbreaks [15] [16] [17] [18]. If, in the coming years in industrialized countries, insects are to become an increasing part of the population’s diet, it is crucial to identify and analyze biological hazards (*e.g.* bacteria, parasites, viruses, prions, yeasts, molds, mycotoxins,
histamine, antibiotic resistance genes) likely to be transmitted, especially since insects are generally consumed, or used for the manufacture of powders, whole, with their digestive tract [19]. This is the main objective of this review.

2. Parasites

It is well known that insects can carry specific internal and external parasites, which can cause diseases to insects. For example, it has been reported that nematodes belonging to the genera Steinernema and Heterorhabditis, can negatively affect Acheta domesticus [20], while Spinochordodes tellinni causes high mortality in crickets and grasshoppers [21]. Insects can also be intermediate or accidental hosts, or even simple vectors, as in the case of Trypanosoma and Leishmania transmission [22] [23].

The transmission of parasites to humans through the consumption of insects was for instance illustrated by [24], who studied the occurrence of intestinal flukes in Southeast Asia. Among the parasite species discussed in the study with the potential to infect humans, around 10% (6/65) of the species had as a host larvae or nymphs of aquatic odonates insects (e.g. dragonfly, damselfly):

- Phaneropsolus bonnei in naiads of dragonfly, in damselfly;
- Prosthodendrium molenkampi: in naiads of dragonfly, in damselfly;
- and 4 species of Plagiorchis (harinasutai, javensis, muris and philippinensis) in insect larva, naiads, nymphs...

Metacercariae of these parasites, known to cause intestinal parasitosis in humans, have been identified in these odonates in Thailand, Indonesia or Laos. The authors linked the cases of human intestinal infection by these flukes to the consumption of the aforementioned insects, since these insect species are consumed in these countries. Regarding Plagiorchis philippinensis, transmission to humans through the consumption of insect larvae has been proven [25].

The case of Gongylonema pulchrum, parasite of the esophagus and mouth, is of particular interest, since some insect species can act as mandatory intermediate host, the final host being most often livestock (domestic cattle). Humans are accidental host by consuming the intermediate host. Following consumption (voluntary or not) of raw insects, patients with nausea and vomiting describe the “Zigzagging” sensation in the mouth or throat [26]. Several cases have been reported in the United States [26] [27].

The consumption of insects (ants) containing metacercariae of Dicrocoelium dendriticum led to dicrocoeliasis in humans. Generally, this food-borne zoonosis is caused by consumption of bovine or mainly ovine contaminated meats [28].

Regarding the main parasites of interest when assessing the microbiological safety of food or water, the bibliography is scanty, probably showing a certain species barrier. However, some protozoa such as Entamoeba histolytica and Giardia lamblia have been isolated from some species of cockroaches. These insects may also carry Toxoplasma sp. and Sarcocystis sp. In some Diptera, various protozoa potentially dangerous to human health have been very rarely isolated (Tox-
oplasma gondii, Sarcocystis sp., Isospora sp., Giardia sp., and Cryptosporidium sp.) [29].

3. Viruses & Prions

3.1. Viruses

Like parasites, there are many entomopathogenic viruses specific for invertebrates that cause Animal Health problems, but their presence and virulence in vertebrates have never been described [30]. The most well-known are the Densovirus (AdDV) and the Cricket Paralysis Virus (CrPV), with the latest to cause particularly high mortality rates for A. domesticus populations [19]. These viruses are a severe problem in mass rearing for the production of food for humans or animals [12]. Notwithstanding, some of these viruses belong to families taxonomically related to human viruses (Iridoviridae, Paroviridae, Flaviridae, Dicistroviridae and Reoviridae), and the question of a possible crossing of the vertebrate/invertebrate border would require to be further investigated. Tests carried out with Densovirus (AdDV) did not lead to multiplication of this virus on vertebrate cells cultures, suggesting that, in that case, this border is efficient [31].

Nevertheless, in some cases, insects allow the multiplication and transmission of viruses to humans or livestock, as active viral vectors [12]. Such is the case of Arboviruses (Arthropod-born viruses) which are responsible among others for manifestations of Dengue fever, Chikungunya disease, and West Nile disease in humans and manifestations of Schmallenberg virus disease in cattle. In fact, these viruses multiply both in vertebrates and invertebrates (mosquitos, ticks). However, mosquitos and ticks are not considered as edible insects and, to date, detection of such viruses in edible insects have not be done yet.

Insects can also transmit vertebrate viruses, with an example being the transmission of H5N1 influenza virus to chickens via infected flies, under experimental conditions [32]. The ability of some viruses to maintain infectivity, for several days, in different matrices, including manure, should draw attention to the use of such animal materials as a substrate for insect rearing [11]. It should be noted that in the European Union the use of waste for the rearing of farmed animals is forbidden. Regarding foodborne viruses of main concern, such as norovirus, rotavirus and Hepatitis virus [17], there is a lack of information on their possible transmission via consumption of insects. In addition, cooking practices and processing methods (e.g. boiling, frying, drying...) can potentially mitigate the risk of transmission [12].

3.2. Prions

So far, the existence of prion or prion-related protein encoding genes in insects has not been reported, meaning also that insects lack the potential to act as biological vectors and amplifiers of animal/human prions [12] [19] [33]. Nevertheless, it has been demonstrated that some insect species can act as mechanical
vectors of prions that are present in the rearing substrate, as highlighted for scrapie in sheeps [34]. Reference [19] summarized studies that report the high stability of prions in the environment, and the prions’ ability to maintain their infectivity for long timespan in soil and water. Use of substrates of non-human and non-ruminant origin [12], as well as control of the substrate’s quality and feed quality [19] could act as preventive measures to avoid occurrence of prions in the edible insects.

4. Fungi (Yeasts, Molds—Mycotoxins Thereof)

It is know that several hundred species of entomopathogenic fungi, often with host specificity, cause significant mortality rates among farmed insect populations [30]. Few studies are available on the contamination of edible insects with yeasts and molds pertinent to food safety. Quantitative levels of these microorganisms serve as hygiene indicators during the manufacturing process of insect-based foodstuffs, usually with no further identification of species or genera. Samples (insect powder or insects) with levels of yeast and molds around $10^4$ cfu/g of powder or insects are not uncommon [35]. Generally, yeasts are not the dominant flora, despite [36] recently demonstrated the presence of *Trichosporon asahii*, an opportunistic pathogenic yeast that causes superficial skin infections in immunocompromised individuals, in edible insects. Reference [37] detected *Debaryomyces hansenii* in dried *Tenebrio molitor* larvae produced in Belgium and the Netherlands.

In an exhaustive review [35], it has been reported that different fungal species were detected on mopane worm (*Imbrasia belina*) in the 90s [38] [39]. *Aspergillus, Penicillium, Fusarium, Cladosporium* and Phycomycetes were found both in the gastrointestinal tract of fresh insects and in whole dried insects [40]. *Cladosporium* and *Fusarium* are food contaminants with fast growing rates, with the ability to deteriorate the product quickly if it is not dehydrated, and to produce certain mycotoxins. Ubiquity and ability of *Aspergillus* and *Penicillium* species to grow even on substrates with low moisture content highlight the need to monitor the levels of these species post-treatment [41]. For example, *Aspergillus flavus* is known to produce aflatoxins, with carcinogenic, mutagenic and teratogenic potential in animals and humans. More recently, several studies have detected species of mycotoxin-producing molds in such as:

- *Aspergillus niger* and *Aspergillus flavus/parasiticus* (aflatoxin producers);
- *Aspergillus ochraceus* (ochratoxin producer);
- *Penicillium citrinum* (citrinin producer);
- *Penicillium verrucosum* (citrinin and ochratoxin producer).

Others insects than mopane worms can harbor these molds:

- *A. niger* was detected in *Bunae alcinoe* and pallid emperor moth (lepidopters), *Oryctes monoceros* and *Rynchophorus phoenicis* (coleopters);
- *A. flavus* was detected in *Macrotermes bellicosus* (isopter), *Rynchophorus phoenicis* and *Alphitobius diaperinus* (coleopters).
Other *Aspergillus* species (*cristatus*, *chevalieri* and *intermedius*) were isolated from *Buyus domesticus* purchased in the Netherlands [37]. It is important to note that the presence of toxigenic molds does not necessarily mean the presence of mycotoxin in products, especially in insect powders. Conditions for mycotoxinogenesis are not always present in insects, although it is known that it is possible, as discussed by [39] in mopane worms from 12 different locations in Botswana. Seven of the twelve samples had higher aflatoxin concentrations, up to 50 μg per kg, than the tolerable dose of 20 μg per kg [39] [42]. Recently, [43] have shown that mycotoxins added in rearing substrate can contaminate lesser mealworms and black soldier fly.

5. Bacteria (and Their Toxins), Histamine and Other Concerns

Once again, several authors wrote that entomopathogen bacteria are not pathogen for human because they are phyllogenetically far from human pathogens [3] [12] [30]. Generally speaking this could be true, but some studies bring some nuances to that truism. For example, *Klebsiella pneumoniae*, an opportunistic human pathogen occurs frequently in intestinal tract of oriental migratory locust [44]. *Spiroplasma* sp., a well known insect pathogens, can cause neurodegenerative diseases in humans and animals. Systemic infection in a human by *Spiroplasma* sp. has also been reported [45] [46]. *Wohlfahrtiimonas chitiniclastica* (a fly pathogen) caused a fulminant sepsis in a 70-year-old man in Argentina [47]. The case of *B. thuringiensis* (BT) is very interesting too because of production of intracellular protein crystals, which are toxic for a large amount of insect larva and it’s used in biological control of insect pest. Nevertheless, recently, role of BT in food borne disease is alleged [48].

Biological hazards likely to be transmitted during the consumption of insects are [3]:

- related to the intrinsic microflora of insects (digestive tract and other anatomical compartments);
- related to extrinsic origin from the environment and rearing conditions.

The biological risk assessment related to the consumption of insects requires a further insight into the probability of transmission of the main foodborne bacterial pathogens (spore forming and non-spore forming). In several studies, various hygiene indicators of insects rearing and insect products thereof have been investigated, in order to evaluate the impact of processing steps (e.g. fasting, heating, freezing, drying, frying...) on the microbiological status of the final product. All these different points will be further discussed below.

5.1. Hygiene Indicators

One of the first microbiological indicators used is the Total Mesophilic Aerobes (TMA) or Total Aerobic Count (TAC) that gives information on the total aerobic microbial load of the under-analysis product. Current literature indicates
that the microbial loads of insect-based foodstuffs are generally high, with the presence of the gastrointestinal tract. Reported TAC values for crickets ranged from $10^1$ to $8.9 \times 10^8$ cfu/g according to type of insect product analyzed, (hard processed or fresh), respectively [49] [50]. A fasting step of 24 - 48 h is usually implemented before insect slaughtering, in order to allow the insects to empty their bowel content and decrease the total microbial load. However, the efficiency of this measure has not been clearly demonstrated yet [19]. Microbiological criteria (hygiene or safety) for insects and products thereof as food at a European Union level do not exist yet. Nevertheless, for instance, the competent authorities of the Netherlands and Belgium, have already proposed 6 log cfu/g of lyophilized insects and 5.7 to 6.7 log cfu/g, respectively, as hygiene criteria [51] [52].

Lactic Acid Bacteria (LAB) and Enterobacteriaceae are the two other indicators discussed in several studies. LAB is a large group of Gram-positive aero-anaerobic bacteria involved in fermentation process. They are generally not pathogenic for human but they can cause spoilage of many foods, including edible insects [53] [35]. High counts of LAB in unfermented products could be a sign of a malfunction of the production process and deviation from the hygiene procedures. For example, in red meat industries the ratio LAB/TAC (number of Lactic Acid Bacteria on number of Total Aerobic Count) has to be around $10^{-2}$. If the ratio is near 1, Quality Management Programs (QMP) have to be checked, and the products are retained [54].

Enterobacteriaceae is a group of Gram-negative bacteria whose main reservoir is the digestive tract of humans and warm-blooded animals. Enterobacteriaceae are sometimes detected in edible insects, especially in insects collected from the wild [55]. However, their presence in products is a sign of poorly hygiene. Since a simple heating step is sufficient to reduce number of Enterobacteriaceae, their presence after such a step is a sign of re-contamination of fecal origin and/or handling error [53]. In Netherlands a threshold of $10^3$ cfu/g of Enterobacteriaceae has been recommended for edible insects [56].

In addition, these biological indicators are used to evaluate the microbicidal efficacy of specifics production steps, namely fasting period, boiling, freezing, drying, frying, grinding, packaging and storage [49] [57]. Not surprisingly, it appears that heat treatments (boiling better than frying or heat drying) permit drastic reduction of load of vegetative cells, specially, but not only, Enterobacteriaceae [35] [58]. Reference [58] has compared TAC reduction (Total Aerobic Count) of Tenebrio molitor larvae after:

- 10 min at 90°C and 45°C (boiling);
- 10 min at 600 MPa (high hydrostatic pressure);
- 10 min at 90°C (dry oven).

The most efficient TAC reduction of larvae was obtained after boiling at 90°C (4 logs) and after High Hydrostatic Pressure treatment (3 logs). Treatment for ten minutes at 90°C in a dry oven or at 45°C in water achieved the same result: around 1 log reduction [58].
5.2. Histamine

Histamine, a heat stable toxin, is a metabolite resulting from the decarboxylation of histidine. A recent outbreak investigation that occurred in 2014 in Thailand was described by [59]. During a seminar, 41 students amongst 227 bring snacks for dinner. Snacks, from a unique vendor in vicinity, were fried grasshoppers, silkworm pupae and green frogs fried as well as crickets and meatballs. Among the 41 students who consumed the snacks, 28 developed symptoms with: flushing, pruritus, urticarial rashes, headache, nausea and vomiting, two were hospitalized. The attack rates were highest (82.6% and 85%) among students who ingested fried grasshoppers and silkworm pupae and lowest (4.4% and 5.3%) among those who did not consume them (RR of 18.7 at 95% CI). The authors postulated that histidine, found in high levels in grasshoppers and silkworm pupae, was converted to histamine via decarboxylation by the bacterial populations present in insects, during storage. Since histamine is heat stable, it was not eliminated during frying, leading to the observed clinical outbreaks [59].

5.3. Non Spore Forming and Spore Forming Bacteria

In Europe, the main food-borne bacterial pathogens are Campylobacter sp., Salmonella, STEC (Shiga Toxin E. coli), L. monocytogenes, C. perfringens (and botulinum), S. aureus, B. cereus group, Yersinia sp. Outside Europe, it would be necessary to add Vibrio sp. to these bacteria. This list contains both non spore forming and spore forming bacteria (such as Clostridium and Bacillus), which are resistant to a wide range of stress conditions related to environment or industrial processes.

In the last few years, several studies have been conducted to analyze further this list of aforementioned microorganisms, with regard to farmed insects and products thereof, and in order to better know if these bacteria are only potential or significant hazards [49] [57] [60]-[68].

Concerning Campylobacter jejuni/coli and STEC (E. coli O157:H7) it seems that insects are not a primary reservoir for these bacteria. Nevertheless, precedent studies have shown that arthropods, like flies, can be a vector of dissemination of Campylobacter in poultry production [69], and a vector of amplification for E. coli O157:H7 [70]. Campylobacter genera have been already identified in edible insects [63]. More recently, [71] identified OTUs (Operational Taxonomic Unit) with a high 16S rRNA similarity with Campylobacter rectus and Campylobacter concisus. Despite this, it seems that Campylobacter jejuni/coli and E. coli O157:H7 are not of critical concerns [34]. Even Yersinia enterocolitica does not seem to be a critical concern for edible insects for [36].

Regarding Vibrio sp., its presence in edible insects is very rare. Only V. hangzhouensis and V. diazotrophicus, which are not pathogenic, have been detected in processed giant water bug [67]. Thus, Vibrio sp. is not considered as a safety concern with regard to edible insects [35].

To our knowledge, Listeria monocytogenes has not been isolated yet from ed-
ible insects by cultural method. Furthermore, processed insects are unable to support the growth of *L. monocytogenes* [68]. Despite this, [65] have detected *Listeria* spp. in cricket powder and processed mealworm, then [68] have isolated *Listeria* sp. from salted mealworm by MPN method. These authors found that the species *L. monocytogenes* is not a critical hazard in edible insects but they recommended further studying.

Regarding *Salmonella*, an important foodborne pathogen, several studies have shown absence in 25 g samples of fresh and processed edible insects [65] [61] [67] [68]. Reference [35] indicates in their review that *Salmonella* has been occasionally detected in tenebroidid beetles, flies, cockroaches [38] [72]. In his review it has been reported that only [73] isolated *Salmonella* from fresh and fried grasshoppers in the North of Cameroon. Notwithstanding, this bacteria is still a major concern since [74] have shown that *Salmonella* can survive in the substrate used during rearing of mealworms and can been further transmitted to the larvae.

Concerning *Clostridium*, as far as we know, detection of *C. botulinum* in edible insects has not been reported yet, although *Clostridium* spp. have been reported in fresh *Tenebrio molitor*, processed whole crickets and grasshoppers [60] [66] [71]. Reference [63] reports that spores of *C. perfringens* have been counted in processed (boiled and dried) whole crickets, grasshoppers and mealworms as well as in cricket powder. The results were around 2 log cfu/g, which is under the ID50 (Infectious Dose) for *C. perfringens*. It is important, however, to draw attention to the conservation conditions of processed insects which should not favor the increase in *C. perfringens* cells. For [75] rehydration and use of *C. perfringens* contaminated insect powders in other preparations (e.g. baby porridge) is a potential risky practice.

Regarding *Staphylococcus aureus*, it seems that the genera *Staphylococcus* is very abundant in the microflora of numerous edible insects [35] [37] [76] [77] [78]. Identification at the species level has not been done in all cases but sometimes *S. aureus* has been detected in fresh and processed insects [61]. When species-level identification is done there is no information on the enterotoxigenic ability of the strain, as far as we know. Presence of *S. aureus* is due to his omnipresence in nature, frequent human carriage and/or processing error. *Staphylococcus aureus* is sensitive to heat but is also halophilic and resistant to low Aw, its toxin is heat-resistant. Physico-chemical properties of insect’s powders are compatible with survival or growth of *S. aureus*. Many authors believe that with respect to this hazard, without overestimating it, we must remain vigilant, especially when taking into consideration the conditions of production and conservation of edible insects and products thereof.

Concerning *Bacillus cereus* group and other *Bacillus* sp., it seems that they are frequently detected in numerous fresh and processed edible insects, wild or reared [35]. Particularly, two studies have focused on *B. cereus* group, with quantitative data on edible insects (mealworms, crickets and silkworms) [37] [68]. *B. cereus*
counts of 4 to 6.6 logs cfu/g have been found by [68] while, [37] reported B. cereus counts of around 5 logs cfu/g in marketed cricket powder. Such counts are compatible with the Toxic Dose of B. cereus [75]. Once again, presence of high number of cells doesn’t mean that B. cereus heat stable toxin (cereulid) is present. Emetic syndrome caused by cereulid toxin of B. cereus needs toxinogenesis in the food. Fresh or processed edible insects are not the ideal substrate for this toxinogenesis. Notwithstanding, B. cereus spores are capable to survive during very long period of time in insect powders. Several authors draw attention to the risk for consumers when rehydrated insect products. Due to its abundance in soils and insects, and due to its resistance to industrial treatments and other stress, B. cereus group is a major concern in consumption of edible insects [68].

5.4. Others Concerns

In addition to these major food-borne pathogens, other publications draw attention to less common (Cronobacter) and/or non-foodborne (Pseudomonas) pathogens. Recently, two studies focused on the dissemination of antibiotic resistance genes through the microbiota of edible insects.

Concerning Cronobacter spp. (formerly known as the single species Enterobacter sakazakii), it’s a Gram-negative rod shaped bacteria present in different environment, causing bacteremia, meningitis and necrotizing enterocolitis especially in babies via the consumption of infant formula [75]. These bacteria can survive in very dry media, it have been detected in dry foods, industrial environment and in edible insects [63] [66]. Reference [75] have pointed out the potential risk using cricket powder, which have texture and physico-chemical properties similar to infant formula, for enrich the nutritional quality of children porridge in Cambodia. The authors concluded that Cronobacter spp. (and C. sakazakii) risks in insect powders have to be evaluated more thoroughly.

Reference [35] pointed out that Pseudomonas sp. and P. aeruginosa are present in several habitats (soil, water, plants...) and have been regularly detected in edible insects [76] [79] [80] [81]. P. aeruginosa, a strictly aerobic Gram-negative bacillus, is considered as an opportunistic pathogen involved in many nosocomial infections, with the ability to contaminate humans via multiple routes (eye, wounds, urinary tract, mouth...). Food is a potential dissemination medium for Pseudomonas aeruginosa, which is sensitive to moderate heat treatment and its presence in edible insects can be therefore controlled by practices such as boiling and heat drying. Nevertheless, it is also known to be very difficult to treat because it is resistant to disinfectants and antibiotics, and can thus participate in the dissemination of resistance genes [82].

Dissemination of antibiotic resistance genes is not a microbiological hazard sensu stricto; it’s an indirect hazard coming from microorganisms. Antibiotic resistance is an emerging threat to public health and microbiota of foods has a significant role to that. The use of antibiotics in edible insects rearing is an under-investigated field. In case of emergency, treatments to eradicate entomopathogen sensitive agent from rearing site are possible, alongside implementation
of others cleaning procedures, in order to save some adults for reproduction [30]. On the other hand, edible insects reared on manure may be exposed predominantly to veterinary drug residues and also to fecal bacteria that may have antibiotic resistance genes [56]. A recent study failed to show the bioaccumulation of antibiotic residues in larvae of black soldier flies reared on a substrate containing them [56], other recent studies have focused on potential detection of antibiotic resistance genes in insect samples [82] [83] [84] [85] [86]. It is known now that genes resistance to tetracycline, erythromycin and β-lactams can been detected in edible insects’ samples with more or less high frequencies depending on the type of insects and their geographical origin. Rearing practices may also have a role in its frequencies (use of antibiotics, quality of feed for insects rearing, quality of substrate use for rearing). These authors agree to say that this phenomenon deserves a thorough evaluation in the coming years [35].

6. Conclusions

The consumption of insects has always been a widespread social and cultural practice in tropical countries [87]. Today, this practice seems to gradually gain the interest of the western world. Insect-derived proteins could have in the next years a quite significant presence in the human diet [88]. Insect consumption could be beneficial from an environmental point of view, since insect rearing, compared to the farming of conventional livestock, requires the use of less agricultural land and water, and has a lower carbon footprint in total.

Insects (and products thereof) can then be perceived as a food of animal origin, with the particularity that insects are consumed with their digestive tract, which, in most cases, is not a natural reservoir for the main and well known foodborne microbiological hazards (e.g. Salmonella, Campylobacter, Listeria, etc.). However, in poorly controlled and hygienic rearing conditions, insects can be vectors of these microbiological hazards. Main sources of contamination of insects by these hazards are the rearing substrate (water, soil, litter, feed) and the human handling during farming. It is therefore very important to supervise the activities of mass rearing of insects with the contribution of good farming practices, and manufacturing and hygienic guidelines [89] [13]. These guidelines should be complemented by the establishment of a food safety assurance system using the principles of the HACCP method in the sites of mass production of insects [90]. For these reasons, production of insects for self-consumption at a personal level and consumption of insects collected from the wild, raw or cooked, shall not be encouraged [3] [12].

It is therefore crucial to be able to mass rearing edible insects in high hygienic conditions, with a steady control. But how to manage that? The question of establishing microbiological criteria (safety or hygiene) for edible insects is an important issue to address. Facing this matter with efficiency would require more quantitative data. Through the present work, it has been highlighted that several authors drew attention to use Total Aerobic Count and spore-forming bacteria
count, as indicators of a proper management of the rearing process. We have also seen that *B. cereus* group (spore forming bacteria) could be a significant hazard related to consumption of processed insects, and that better knowledge of the microbial reduction effects of some key steps of the process is needed. It seems today that these particular points are the priorities for future studies on biological hazard related to the consumption of edible insects.

**Disclosures**

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**Conflicts of Interest**

The authors declare no conflicts of interest regarding the publication of this paper.

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Abbreviation List

BT: *Bacillus Thuringiensis*

CFU: Colony Forming Unit

CI: Confidence Interval

HACCP: Hazard Analysis Critical Control Point

ID50: Infectious Dose 50

LAB: Lactic Acid Bacteria

MPN: Most Probable Number

OTU: Operational Taxonomic Unit

QMP: Quality Management Program

RR: Risk Ratio or Related Risk

rRNA: ribosomal Ribonucleic Acid

STEC: Shiga Toxin *Escherichia coli*

TMA: Total Mesophilic Aerobes

TAC: Total Aerobic Count

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