Stereo and scanning electron microscopy characteristics of poultry breeding beetle (*Alphitobius diaperinus*) – a filamentous toxigenic fungi carrier

Carlos Eduardo da Silva Soares1*, André Weber2, Vildes Maria Scussel1

1Laboratory of Mycotoxicology and Food Contaminants, Department of Food Science and Technology, Federal University of Santa Catarina, Brazil; 2Federal Institute Catarinense - Campus Araquari, SC, Brazil

**ABSTRACT**

This study isolated *Alphitobius diaperinus* (*live* and *dead*) insects from shed’s aviary bed to investigate their fungi spores distribution (that affects chicken health and meat production) and their accumulation sites (*dorsal* & *ventral*) characteristics by different microscopies (stereo and scanning electron). Despite *live* beetles being the main fungi spore carriers, the *dead* ones had far more spores attached on their body exoskeleton thereby being a focus of infection. That was due to the anatomical sites favoring spores trapping effect, together with beetles’ different moisture content. Regarding the spores distribution and so the hyphae presence & mycelia concentration on *dead* *A. w. diaperinus*, they were mainly detected at the (a) elytra, elytral suture and pronotum (on the *dorsal* side). Despite that, the highest spores/mycelia concentration was at the mouthparts, prosternum and legs (femur & tarsus) (on the *ventral* side). Indeed the beetle’s *ventral* anatomical microscopic structures (mouthparts & legs) sheltered the highest fungi spores concentration and colonies proliferation. Thus *dead* beetle colonies growth lead to spore multiplication, their dissemination throughout the aviary bed environment and so their contact to chicken feet and body, leading to discomfort and diseases development/mycotoxicosis. The filamentous fungi were most detected from the *Aspergillus* and *Penicillium* genera. Therefore *dead* beetles should be removed from aviary (at each 45 breeding cycle) to reduce contamination. They represent rich substrates for fungi development with possibility of toxin formation, apart from the chicken diseases exposure due to their insects eating habits.

**Key-words:** aviary bed, fungi, mycotoxins, poultry, vectors

**1. INTRODUCTION**

Insects of different genera and species infest poultry farming during breeding and can distribute fungi spores (deteriorating and toxigenic) throughout shed’s facilities including the aviary bed (Lacey et al., 1996; Chandler et al., 1997; Green, 2008; Soares et al., 2017). They can mechanically transport fungi spores on their bodies, thus increasing significantly the chicken infections rate (Kluth, 2002; Quirino, 2008). Aviary sheds’ temperature and humidity conditions play an important role in in insects infestation in poultry breeding there by resulting in fungi infections including other living organisms (mites, bacteria and virus), thus affecting chicken health (Scussel, 2002; Hazeleger et al., 2008; Bosly et al., 2014; Soares et al., 2017).

Infestation in aviary beds of insect such as of *Alphitobius diaperinus* (Coleoptera, Tenebrionidae), commonly called darkling beetle, is responsible for large losses in poultry farming (Skov et al., 2004; Rolf and Schiller, 2016; Soares et al., 2017). Apart from fungi also bacteria such as *Campylobacter jejuni*, *Salmonella enteric* among others, have been reported to contaminate aviary bed and chicks (Hazeleger et al., 2008). When there is shortage of food during rearing, chicks/chicken begin to eat those insects (living organisms contaminated) leading to diseases development. This also results in decreased chicken feed conversion, diarrhea, stress and reduces body weight (Matias, 1992; Despins and Axtell 1995; Skov et al., 2004). Other *Alphitobius* species such as *A. laevigatus* (Fabricius), *A. stephens* and *A. piceus* (Oliver) have been isolated from stored grains and flour (Hagstrum, 2017).

*Corresponding author:*
Carlos Eduardo da Silva Soares, Laboratory of Mycotoxicology and Food Contaminants, Department of Food Science and Technology, Federal University of Santa Catarina, Brazil. E-mail: c.ess@posgrad.ufsc.br.

Received: 18 July 2017; Accepted: 15 February 2018
During the day, those insects remain under the aviary bed surface, inside the shed structures (columns/walls) and equipment (feeders). In nocturnal period they become more active, attracted by the light (moving in and out of the aviary) and distributing fungal spores (Paiva, 2000). Those spores can be carried by adhesion on insect exoskeleton waxy surface and so under its cuticles (Noh et al., 2016; Dittmer et al., 2011, Bruns, 1995; Lenardon et al., 2010). Insect constant movement, within an ecosystem, contributes to Aspergillus, Penicillium, Mucor and Rhizopus (all storage fungi), including Fusarium (field fungi) spores dispersal (Bidoehka 1997; Saint Géronis-Grid, 1984; Scussel et al., 2011; (Slipinski & Escalona, 2016).

Regarding the effect of fungi presence in chicken breeding environment, they can lead to development of diseases in chicken/chicks (aspergillosis) and embryos (susceptibility to fungal effects during post-hatching) including spores inhalation (lung mycosis). Despite that, some of those fungi genera can produce mycotoxins. Mainly the filamentous fungus species (A. flavus & A. parasiticus, P. ochraceus, F. verticillioide, F. graminearum– aflatoxins, ochratoxin A, fumonisins, deoxinivalenol and zearalenon, respectively) and those fungi genera can produce mycotoxins. Mainly the filamentous fungi species (A. flavus & A. parasiticus, P. ochraceus, F. verticillioide, F. graminearum– aflatoxins, ochratoxin A, fumonisins, deoxinivalenol and zearalenon, respectively) and can contaminate aviary beds (high humidity/feast residues) and so the chicken (Placinta, 1999; Van Broeckevenson et al., 2017). In young birds, mycotoxin intoxication lead to high mortality (Wang, 2012; Arné et al., 2011; Andreatti Filho, 2000). It should be noted that storage fungi are found spread on the poultry farming machinery (equipments mills and hoppers) and in the storage environment (sheds, warehouses, silos) as long as good conditions (temperature and humidity) are present (Mallmann, 1994; Saleemi et al., 2010).

Considering the extensive poultry farming’s activities; the problems with insect infestation; the consecutive and high number (8x5) of chicken breeding/rearing cycles (45 days each) produced per year (same aviary bed) and the development of fungi infections that reduces chicken/meat sanitary conditions an investigation was carried out on the A. diaperinus (adult & larvae stages) microorganisms carrier characteristics and its main anatomical sites able to shelter and spread fungi spores (aviary bed & chicken environment contamination) by stereo (SM) and scanning electron (SEM) microscopies.

This is the first work investigating A. diaperinus detailed anatomic characteristics and fungi accumulation sites reported by SM & SEM.

2. MATERIAL AND METHODS

2.1 Material

(a) Samples: Insects (A. diaperinus), extracted from aviary bed (at the 45th day of chicken breeding, no insecticide application), both live and dead (adults and larvae stages).

(b) Equipment: tweezers, Prolab (Sao Paulo, SP, Brazil); sieve system, 9-16 mesh (2.00-1.00 mm/µm apert., 10-18 USM/ASTM) Beffer (Caias, SP, Brazil); a meter, model Aqua- Lab4TE, Decagon (Sao Jose dos Campos, SP, Brazil); drying oven, Olidef-cz (Ribeirao Preto, SP, Brazil); stereo microscope, model Opticam (Doral, FL, USA); scanning electron microscope (x5000), model JSM-6390LV, Jeol (Peabody, Mass, USA) and gold coating machine, model EM-Scd500, Leica (Leider, Ill., USA). Other materials: stubs (small metal blocks, 9 diameter and 10 mm height).

2.2 Method

(a) Aviary bed and insect collection: Prior to the procedure of insect isolation, an infested aviary bed sample (mc: 40.6%, a: 0.98) was collected as follows (a.1) aviary bed- a portion (100 g) was obtained from the shed’s floor (10 cm depth) after 45 days of chicken complete growth stage, as reported by Soares et al. (2017) and proceeded to (a.2) insects isolation – separated beetle samples (both growth stages: adult & larvae) were collected (live & dead) from the aviary bed as in (a.1), through sieving (9-16 mesh) by picking them with tweezers for the microscopy analysis preparation. *killed by applying a insecticide (cypermethrin) solution in acetonitrile.

(b) Insects preparation for microscopy: Insects (both growth stages) were prepared for SM and SEM, as follows (b.1) SM – the whole (isolated alive & dead) and different insects parts (dorsal e ventral) were separated in Petri dishes (and so the larvae and directly taken for SM observation; for (b.2) SEM – those samples from (b.1) were prepared by stubs mounting and their surfaces gold coated, as reported by Scussel et al. (2014a). Briefly, insects were fixed on stubs (containing carbon double-sided tape), then vacuum gold coated (by placing them onto a Planetary Gold Coater stubs holder, vacuum applied and coated with a 40 nm gold layer).

(c) Microscopy observation: Insects whole and parts samples prepared in (b) were taken for microscopy observation, (c.1) SM – they were directly taken (from b.1) for characteristics identification of head, thorax, abdomen, legs at different amplification (26 to x80) and so the special anatomical parts that shelter fungi spores and/or growing colonies; for (c.2) SEM – the stubs with the insects parts were taken to the SEM microscope to investigate the fungi presence/distribution/proliferation/genera identification (mycelia, hyphae, conidia) on insects’ parts. Also the sites where they are mainly adhered onto the insect body (25 to x1,100) were investigated.

3. RESULTS AND DISCUSSION

Data on the A. diaperinus microscropic characteristics (SM & SEM) regarding the anatomical parts that shelter fungi spores at high concentration, were identified (Martins
et al., 2016). The fungi spores distribution/accumulation, whether on beetles captured alive or found dead in the aviary bed, were registered, including their differences. Fig 1-6 present beetles (adult & larvae stages) characteristics and the main parts that most accumulate fungi spores (either on dorsal or ventral, and whether alive or dead).

3.1 Alphitobius diaperinus microscopic characteristics

In order to understand the A. diaperinus insect’s fungi spores transport through the poultry breeding chicken environment and their effect on chicken safety, it was necessary initially to identify its micro-morphological characteristics (adult & larvae stages) (Povaluk, 2017 and Faruk et al., 2005). The main SM & SEM beetle (both stages) different body parts (head/thorax/abdomen) characteristics are shown in Fig 1 and 2, respectively.

SM: regarding the beetle exoskeleton morphological parts at (a) adult stage, it was possible to identify some details by SM (up to x80) of the three different parts that comprise the whole insect, as follows. At head – the eyes, antennae & mouthparts (that comprises of different robust small structures – to be seen on SEM); thorax - pronotum (dorsal) & prosternum showing its hairiness distributed along its hypomeron with thoracic legs (ventral); abdomen – the two elytrons (with the suture between them) & hindwings (dorsal) and ventrites & legs (femur/tibia/tarsus) (ventral). Fig 1.a shows details of adult A. diaperinus by SM. On the other hand, the beetle at the (b) larvae stage, which is seen as a hyaline structure by SM, comprises also of three main parts, i.e., head - a pair of immature eyes, antennae & mouthparts; thorax – with the pro/meso/metathorax and abdomen – segments, legs & spines. Fig 1.b shows the dorsal view larvae structures.

SEM: the more detailed main beetles adult and its larvae stage micro-morphological SEM (up to x1,100) characteristics were as follows. For the (a) adult stage the head – presents one pair of compound eyes (i.e., many lenses/ommatidia pack into hexagonal array with dome format facets), mouthparts (set of robust maxillae, maxillary palp, labrum, labial palp, labium, mandible and hairiness) & the antennae (funicle and club, with pores and spines). At the thorax region – is the hypomeron (a broad pubescent area) & the prosternum (a strongly developed front at the medial part) that has a thick hairiness (sensitive structures near the hypomeron anterior edge) on ventral side. On the other side, is the pronotum (also hard structure, that some species use it in combat) at the (dorsal). The abdomen – comprises of a pair of elytra & its suture (containing intercostal cuticle layer) (dorsal) and ventrites with legs - femur, tibia (both with row of spines with different sizes & shapes, tibial spurs) & tarsus (comprises of a series of tarsomeres - tarsal segments, with a pair of claws at the end, each) (ventral). Figs 2a.1-a.4 show some adult A. diaperinus structures. Other beetle’s adult characteristics that also trap fungi are shown in the next Section (3.2). Regarding the SEM characteristics of the beetle at (b) larvae stage: it was possible to observe the head – with a comprehensive set of structures as follows, a pair of larval eyes (rather visible), mouthparts (similar to the adult beetle) and antennae (antennal segments I-III and a thin apical segment); the thorax - with the prothorax containing a pair of legs and abdomen - with its several segments called tergites (dorsal) and sternites (ventral) with spines. Figs 2b.1 and b.2 show the features on larvae dorsal side.

3.2 Alphitobius diaperinus fungi spores accumulation and colonies proliferation – anatomical sites

Utilizing both microscopy techniques, specially SEM, it was possible to clearly identify the beetle’s anatomical fungus sheltering special sites. Those that (a) most concentrate spores (adhered easily on insect body) and/or fungi colonies (mycelia/hyphae/reproductive structures) growth. Also those sites that (b) provide condition to allow fungi infection to take place (high humidity & rich substrate) (Butt et al., 1995). All of them, thus contribute to the spores/fungal transfer throughout the poultry environment and animals are shown in Figs 3 to 6.

SM: by searching fungi presence/accumulation on the (a) adult stage, when the beetle was captured (a.1) alive by SM, it was possible to slightly visualize spores adhered on them
(although they were, apparently, not visible by naked eye - not seemed to be infected). On the other hand, from all the \((a.2)\) dead beetles and their parts that were viewed (head/thorax/ abdomen) both, on the dorsal & ventral sides, it was registered high fungi infections with colonies growth throughout the insect’s body (Bleiker et al., 2009). Despite that, they were more concentrated in certain parts as follows (in decreasing order of spores - colonies concentration - infections): on the thorax (prosternum/pronotum), followed by the legs (tibia/ tarsus), head (mouthparts) and abdomen (elytra). (Fig 3). Fungi reproductive structures were clearly more visible, including hyphae, adhered on to the dorsal (both at elytra suture and pronotum) side (Fig 3 a.1, a.2). On the ventral side, the fungi colonies were mostly concentrated on the thorax (prosternum); legs (tibia/tarsus) and head (mouthparts) (Buett et al., 1992). At the \((b)\) larvae stage, the presence of spores was seen throughout larvae morphological structures, however more concentrated on its abdomen (segments) (Fig 3b). Those features and spores location observed by SM, will be seen more visible and detailed by SEM micrographs next.

**SEM:** when beetles were investigated for their anatomical sites fungi accumulation at higher magnification, their distribution were specified more accurately. At the \((a)\) adult stage, when beetle was captured \((a.1)\) alive, the spores (conidia) were mainly adhered (concentrated) on the abdominal ventrites. On the other hand, in the \((a.2)\) dead beetles, high fungal infections were observed in both, at the dorsal and ventral regions (colonies development). The presence of reproductive structures (was mainly on the elytral suture (abdomen), on the dorsal region (Fig 4.a-f). Next, they were seen on the head (mouthparts - mandible and hairiness) and thorax (prosternum hairiness) as high colonies infection with conidia spread all over, followed by the abdomen (ventrites) and legs (tibia/tarsus) (ventral region) (Fig 5a,b). The filamentous fungi most detected were from the Aspergillus and Penicillium genera (Fig 4.c/d and e/f, respectively). It is important to emphasize, that the dead \((b)\) larvae stage: micro-morphological sites of fungi accumulation were observed mainly on the head structures of mouthparts & antennae of the thorax, prothorax and abdomen segments (tergites and sternites) (Fig 6).

### 3.3 A. diaperinus proliferation in poultry rearing facilities versus chicken health and safe meat production

Considering the spores presence and fungi proliferation detected in the \(A.\) diaperinus in the current study and its obvious dispersal throughout the poultry breeding shed’s, one can conclude that they also disseminate diseases and affect chicken well-being during their growth, thus

---

**Fig 2.** Scanning electron micrographs of *Alphitobius diaperinus* (Panzer) anatomical special parts characteristics isolated from aviary bed: (a) adult stage - (a.1) head & prosternum (a.2) antenna (club); (a.3/a.4) abdomen and (b) larvae stage - (b.1/b.2) head and abdomen respectively [27 to x400].

**Fig 3.** Stereo micrographs of dead *Alphitobius diaperinus* (Panzer) isolated from aviary bed fungi contaminated/infected distribution: (a) adult stage (dorsal - a.1/a.2; ventral - a.3/a.4) and (b) larvae stage (dorsal view) [26 to x160].
interfering on meat formation and carcass production (standard size).

A. diaperinus beetles have been reported being ingested by chicken and were detected in poultry pro-ventricles and gizzards post-mortem (da Silva et al., 2001; Soares et al., 2017). That lead to certain concern, as that beetle is carrier of living organisms thus increasing fungal survival on their bodies and environment, thus transmission and reproduction (Moser et al., 2010).

Regarding chicken safety, apart from interfering on their well-being (by insects ingestion and/or discomfort by contact), diseases such as aspergillosis in adults, fungi high susceptibility in embryos (contamination during posture), bacterial infections, allergies (mites proliferation) can take place, apart from spores inhalation (lung mycosis) (Lambkin et al., 2007 and Banjo et al., 2005). Therefore, there is a need to apply control/prevention procedures such organic and/or chemical insecticide applications.

4. CONCLUSION

The A. diaperinus main anatomical sites that most concentrate fungi spores, leading to their distribution in live beetles were abdominal ventrites (especially in their sutures) and legs. On the other hand, the main sites on the dead beetles, were the elytral suture, mouthparts, legs and prosternum, which increase contamination spreading by the colonies development.

Regarding the larvae, it had spores adhering throughout the body, being the main contamination on its head.

The fungi most isolated were of Aspergillus and Penicillium genera.

As far as the chicken well-being is concerned, beetles presence in the aviary bed can be ingested causing diseases (bacterial contamination) and fungi infections in chicks and adult animals (that ingestion may be more intense whether the feed supply is short or ceased).

To minimize the problems caused by beetles infestation, chemical (ozone/diatomaceous earth/zinc compounds/insecticides) and/or biological (plant extracts/insects entomopathogenic species) control should be applied in poultry farms.

This is the first work investigating A. diaperinus detailed anatomic characteristics and fungi accumulation sites reported by SM & SEM.
Fig 6. Scanning electron micrograph of _Alphitobius diaperinus_ (Panzer) LARVAE (HEAD & ABDOMEN) isolated from aviary bed FUNGI INFECTED: (a.1/a.2) head, (a.3-a.6) abdomen [25 x to 700].

REFERENCE


Gereges-Gridelet, S. 1984. Effects of dietary lipids on the population...
growth of Dermatophagoides pteronyssinus. In: D. A. Griffiths and C. E. Bowman. (Eds.), Acarology VI, Ellis Horwood, Chichester, UK.


