THE HAEMOCYTES EFFECT OF THE *PAENIBACILLUS LARVAE* SPORES IN ADULT HONEY BEE WORKERS (*APIS MELLIFERA L.*)

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Abstract

This work aimed to study the pathogenic effect of the *Paenibacillus larvae* spores on adult honey bee workers to explain the death many of them in the communities affected by American foul brood disease. This work was done through two methods; first injected the microbial suspension in the intersegmental membrane of the back lymph cavity of adult bee workers. The blood films were done after 1, 2, 4&12 hrs. beginning of the injection method, while with the feeding method it carried out after one, two and three months. During the first period of the microbial injection one hour later an elongation and minor abnormalities were occurred in some blood cells. The second stage began after 2 hrs. is characterized by loss of the blood cells dye strength. After 4 hrs. the third stage is characterized by increase atrophies of some blood cells. The fourth and final stage after 24 hrs. abnormalities in most blood cells with sharp decline in their numbers clearly. This stage considers beginning end of the blood cells alive and the rapid death of most of bee workers was detected. This experiment affirms that adult bee workers can be affected by the *Paenibacillus larvae* spores. That may be one reason of death bee workers in infected bee colonies by the American foul brood disease.

Keywords: Honey bee, Haemocytes, *Paenibacillus larvae*.

INTRODUCTION

Bees live in the world in a complex interaction with their environmental condition. In order to understand one of these interventions between pathological and physiological process is indispensable particularly if their effects reflects on the honey bee colony alive. One of these interposition factors is the American foul brood disease (AFB) which is one of the most jeopardy bacterial attacking honey bees colonies causing a decline in the bee population and colony vigor (Generisch *et al.*, 2006). Although several antibiotics were used for controlling the microbial infection (Genersch and Otten, 2003), may several problems still affecting the honey bee lifespan and colony products (Genersch 2010). Despite the pathogen mainly infected honey bee larvae, it can seriously injured adult bees reflects on the cellular immunity responses and most physiological organs (Szynas and Jedruszuk, 2003). One of subjects that worry researchers and beekeepers is ambiguity of gradually death honey bee members and rapid replacement honey bee queens. May the circular system be injured by different pathogens particularly by the (AFB). Therefore, the present work
aims to clarify the pathogenic role of the *Paenibacillus larvae* spores on the blood cells as a general physiological factor in side and in another side to clarify other defensive behavior can the workers take during different attack methods. Through microbial feeding method a gradually decrease in the blood cells were detected in the haemolymph of infected honey bee workers during three months of completely death bee colonies.

**MATERIALS AND METHODS**

The present work was done at the department of Apiculture, Plant Protection Research Institute, Agriculture Research Center, Dokki, Giza, Egypt. Nine of honey bee colonies were conducted for this study and divided into three groups one for injection the *paenibacillus larvae* spores which caused the American foul brood disease to honey bees. The 2nd was feeding adult bee workers by the microbial through sugar solution. The last for the control not received any treatments.

1- **Isolation and purification of the *Paenibacillus larvae* spores**

This method was done according to Nordstrom and Fries (1995) and Anderson (1990) techniques at Fac. of Science. El-Mansoura Univ.

2- **Microbiological assays**

Bacterial smears were microscopically examined after treated with Gram stain according to Shimanuki and Knox (1991).

3- **The inoculation with *P. larvae* Spores**

The following techniques were carried out;

3.1. **Injection method**

10 µl of the of *P. larvae* spores suspension with concentration of (6x10⁶ /1ml) was added to 100 ml of the saline solution (Na Cl 0.09%), then 4 µl of them was injected into healthy nursing honey bee workers through back lymph cavity between intr segmental membrane (about fifty samples / colony) (Casteels et al.,1989). The blood films were done after 1, 2,4,24 hrs. of injection initiation from treated and non treated bee workers with the pathogen.

3.2. **Feeding method**

10 µl of the microbial suspension of *P. larvae* spores with concentration of (6x10⁶ spores /1ml) was added to 100 ml of the saline solution (Na Cl 0.09%) and sweated with the sucruse then feeding honey bee colonies using spraying method between bee combs (10 ml / colony / one time). The blood films of nursing honey bee workers were done after 1, 2 and 3 months of treatments.

4- **Blood smear films**

Blood films were prepared and pigments with Gimsa's stain for each treatment (Nappi and Streams, 1969).
RESULTS AND DISCUSSION

The classification of the blood cells in honey bee workers was done according to Zakaria (2011). The haemolymph of the honey bee workers inoculated with the Paenibacillus larvae spores showed different features towards the microbial infection as follows;

{I} – Microbial injection method

I- One hour later of treatment

Through injected the microbial suspension of the pathogen (Paenibacillus larvae) into back cavity lymph of adult honey bee workers a different features against the bacterial infection were detected as follows;

1- Neutrophile: Neutrophile cells take different shapes towards bacterial invasion in comparison to control one Fig. 1(B). Whereas expansion in the outer surface wad detected as shown in Fig. 1(C). Also laceration in the general cells components (Fig. 1-D).

2- Spindle shaped cells: Dwarf in the Spindle shaped cells due to bacterial infection as shown in Fig. (1-F) was detected in comparison to healthy one (Fig.1 E).

3- Macronucleocytes: They are the most famous as a phagocytes Fig.1(G). In Infected state this cell showed slight to sharp elongation shaped in their outer surface (Fig.1-H).

4- Plasmatocyte: Plasmatocyte cells lost parts of their components (Fig. 1-J) in comparable to healthy one (Fig.1-I).

II – Two hours later of treatment

1- Proleucocytes: Proleucocyte cells showed flabbiness in the outer cytoplasm tissue (Fig.2B) inverse to control one (Fig.2A).

2- Hyalinocytes: The cytoplasm of the Hyalinocyte cells showed more faint stain (Fig.2D ) adverse to control one (Fig. 2C).

3- Basophile cells: The Basophilic cells showed rupture in their outer membrane (Fig.2F ) and lost of force pigments (Fig. 2G) opposite with healthy one (Fig.2E).

4- Other blood cells: Other blood cells were not severe affected as the pervious mention. This period is characterized by increase the transparency of the cell cytoplasm stain.

III – Four hours later of treatment

1- Basophile cells: In spite of the damage of the Basophile cells was detected (Fig.2H). The most of them was still healthy as that of Fig. (2E).

2- Other blood cells: Other blood cells were affected by different degrees. This period is characterized by increase the initiation of dwarf cells.
IV– Twenty four hours later of treatment

After 24 hours of treatment a sharp deformation in most of the blood cells with force decline in their numbers was detected reach to 90% particularly with the Hyalinocytes. This is considered end of the blood cells alive. Most of the bee workers were dead after 24hrs. of the microbial injection period.

{II} – Microbial feeding method

Through microbial feeding method a gradually decrease in the blood cells were detected in the haemolymph of infected honey bee workers during three months of completely death bee colonies. The Spherulocyte cells showed more activity in state of the bacterial infection as shown in Fig.(3-A,B). The Oenocytoids cells showed abnormal state (Fig.3 C, D), whereas Spindle cells showed dwarf shape (Fig.3 E,F). Despite of a sharp decrease in the blood cells, little of infected bee workers were still alive before one month of the completely dead bee colony.

Generally, it could be concluded that during the first period of the one hour of the microbial injection an elongation and minor abnormalities were occurred in some blood cells. After 2 hrs. the second stage began, which is characterized by loss of the blood cell dye strength. After 4 hrs. the third stage is characterized by increase dwarfism of the blood cells. The 4th final stage after 24 hrs. abnormalities in most blood cells with sharp decline in their numbers was detected. The last stage consider beginning end of the blood cells and the rapid death of most of bee workers. This experiment affirm that adult bee workers can be affected by the Paenibacillus larvae spores. That may be one reason of death adult bee workers in infected bee colonies by the American foul brood disease.

Paenibacillus larvae spore’s effects on the blood cells of immature and mature honey bee workers were detected by Zakaria (2007). Papadopoulou et al., (2003) reported that blood cells of the honey bees artificially infected with Pseudomonas aeruginosa cause septicemia showed not significantly effect on the total and differential haemocyte counts. Also, deficient nutrition can impair immune function and increase the susceptibility of individuals to disease as suggested by Alaux et al., (2010). Julia et al., (2017) reported that the toxin Plx2A is an important virulence factor of Paenibacillus larvae, the etiological agent of American Foulbrood, the most destructive bacterial disease of honey bees. Plx2A induced actin cytoskeleton reorganization while in insect cells, vacuolization and the occurrence of bi-nucleated cells was observed. The latter is indicative of an inhibition of cytokinesis. All these cellular effects are consistent with Plx2A inhibiting the activity of RhoA by covalent modification. That review clarifies the rupture occurred to the blood cells resulted the exposure to the pathogen (Paenibacillus larvae). Michel (2018) reported that toxins are powerful pathogenicity factors produced by certain bacteria, fungi, animals, and plants which mediate drastic interactions of these pathogens on the organism host.
Notably, bacterial toxins were the first compounds which were identified as responsible for severe bacterial diseases in animals. Endotoxins are membrane compounds of Gram-negative bacteria which elicit an inflammatory response in host. Gilliam (1978), suggested that the microorganisms could have entered the queen during copulation.

![Fig.1 Paenibacillus larvae spores and blood cells of adult honey bee workers one hour later of the bacterial injection(x-400).](image)

A. *Paenibacillus larvae* spores.
B. Healthy Neutrophile cell.
C. Elliptical of the Neutrophile cell.
D. Indentation in the outer surface of the Neutrophile cell.
E. Spindle shaped cell as normal one.
F. Spindle cell a dwarf as bacterial infection.
G. Macronucleocytes in normal state.
H. Macronucleocytes with slight elongation due to bacterial infection.
I. Plasmatocyte cell in healthy state.
J. Damage in the Plasmatocyte cell resulted of *P. larvae* infection.

![Fig.2 Blood cells of adult honey bee workers in normal and bacterial infection state after 2,4 hrs. of the microbial injection.(x-400).](image)

A. Proleucocyte cells.
B. Flabbiness in the outer surface of the Proleucocyte cells in infection state.
C. Hyalinocytes of transparent cytoplasm in healthy one.
D. Faint stain of infected Hyalinocytes.
E. Basophile cells in normal state.
F. Basophile cells showed extensions in the outer membrane.
G. Basophile cells showed basophilic faint stain.
H. Damage the basophile cells after 4 hrs. of infection state.
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Fig. 3. Blood cells of adult honey bee workers through inoculation the Paenibacillus larvae spores in feeding syrup (x-400).

A. Spherulocyte cells in normal state.
B. Spherulocyte cells showed more activity in the bacterial infection.
C. Oenocytoids in healthy state.
D. Oenocytoids in the infection showed deformation.
E. Spindle shaped cells in normal state.
F. Dwarf Spindle shaped cells in the microbial infection.

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التأثير الخلوي لجراثيم *Paenibacillus larvae* شغالات نحل العسل البالغة (*Apis mellifera* L.)

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يهدف هذا البحث إلى دراسة التأثير الخلوي لجراثيم تعرف الحضنة الأمريكية لطوايف نحل العسل على خلايا الدم لشغالات نحل العسل البالغة والتي قد تكون أحد أسباب هالك الطائفة، وقد تم ذلك من خلال المدعم الصناعية بجراثيم المسبب المرضي عن طريقين الأول حق المسبب المرضي بين الحفاظات الظهرية في التجويف الظهري للنحل البالغ والثاني عن طريق تغذية النحل بالمحلول السكري المحتوي على زراعة الميكروب المختبر. أجريت أفلام الدم خلال فترات زمنية مختلفة بعد ساعة واحدة وإثنتين وأربع ساعات ثم بعد أربعة وعشرون ساعة بالنسبة لتجربة الحقن أما في الحالة الثانية فقد أخذت عينات الدم بعد شهر وثرين وثلاثة أشهر. وقد تلاحظ وجود استجابات مختلفة لخلايا الدم تجاه المسبب المرضي خصوصاً في الفترة الأولى بعد ساعة واحدة من الحقن الميكروبوي وجود استطالة وتشوهات في بعض خلايا الدم وبعد ساعتين تلاحظ فقدان خلايا الدم لقوة الصبغ بينما بعد أربع ساعات من بداية الحقن فقد تلاحظ وجود ضمور في خلايا الدم أما في المرحلة النهائية بعد أربعة وعشرون ساعة من المعاملة فقد تلاحظ عظم معتظم خلايا الدم ومتانة شغالات نحل العسل سرعان بعد تلك الفترة. أما في حالة العدوى بجراثيم الميكروب عن طريق التغذية بالمحلول السكري تتبخر وجود نقص تداجم في خلايا الدم خلال الثلاثة أشهر الأولى من بدء المعاملة حيث تدهورت وانتهت طوانة النحل المعاملة تماما. ومن نتائج هذه التجربة يتضح تأثر خلايا الدم النحل البالغ بالسبب المرضي الذي قد يكون أحد أسباب انهيار طوانة نحل العسل المصابة بمرض تعرف الحضنة الأمريكية.