Research Article

Confirmation on Status of *Chaetocnema basalis* (Coleoptera: Chrysomellidae) as A Vector of Stewart Wilt Disease

Konfirmasi Status Chaetocnema basalis (Coleoptera: Chrysomellidae) sebagai Vektor Penyakit Layu Stewart

Heri Widodo^{1)*}, Arman Wijonarko²⁾, Witjaksono²⁾, & Suputa²⁾

¹⁾Agricultural Quarantine Station of Tanjung Balai Karimun, Jln. Jend. Sudirman Tanjung Balai Karimun, Kepulauan Riau 29661 ²⁾Department of Crop Protection, Faculty of Agriculture, Universitas Gadjah Mada Jln. Flora 1, Bulaksumur, Sleman, Yogyakarta 55281

*Coresponding author. E-mail: heri.widodo@ymail.com

Submitted March 14, 2017; accepted May 16, 2017

ABSTRACT

Chaetocnema pulicaria and C. denticulata are recognized as vectors of Stewart wilt disease caused by Pantoea stewartii on maize. These insects have not been reported yet in Indonesia, but Stewart wilt disease has been reported in Java and Sumatera Islands. Genus Chaetocnema which presented in Indonesia is C.basalis. It is not cleared whether C. basalis is a vector for Stewart wilt disease like C. pulicaria and C. denticulata. This reseach was aimed to conduct the confirmation on status whether C. basalis have a role as vector of Stewart wilt disease on maize or not. C. basalis imago were collected from maize growing areas in Yogyakarta, and then starved for 24 h. Treatments were applied by placing imago of C. basalis on infected-P. stewartii plants for 72 h. Five insects were then transferred to each plot of healthy plant (1 plot consisted of 5 plants) for 72 h. For control, imago of C. basalis were put on healthy plants for 72 h and five insects were then transferred to other healthy plant (1 plot consisted of 5 plants) for 72 h. Each treatment was repeated three times. On the fifteenth days after transmission, PCR assays were carried out on leaf samples and isolates of bacteria. All sampled leaves analysis showed that there were no Stewart wilt disease transmission based on PCR assay and bacterial isolates. This concluded that C. basalis is not a vector for Stewart wilt disease on maize.

Keywords: Chaetocnema basalis, Stewart wilt, vector

INTISARI

Chaetocnema pulicaria dan C. denticulata merupakan serangga vektor penyakit layu stewart yang disebabkan oleh bakteri Pantoea stewartii pada tanaman jagung. Kedua serangga ini belum pernah dilaporkan keberadaannya di Indonesia tetapi penyakit layu stewart telah ditemukan di pulau Jawa dan pulau Sumatera. Serangga Genus Chaetocnema yang ada di Indonesia adalah Chaetocnema basalis. C. basalis belum diketahui secara pasti sebagai vektor penyakit layu stewart seperti halnya C. pulicaria dan C. denticulata. Penelitian ini bertujuan untuk melakukan konfirmasi status apakah C. basalis berperan sebagai vektor penyakit layu stewart pada tanaman jagung atau tidak. Serangga uji berupa imago C. basalis yang dikoleksi dari pertanaman jagung di Yogyakarta, lalu dilaparkan selama 24 jam. Pengujian perlakuan dilakukan dengan menempatkan imago C. basalis pada tanaman terserang P. stewartii selama 72 jam. Kemudian dipindahkan pada tanaman sehat sejumlah 5 ekor per plot tanaman bersungkup (1 plot terdiri dari 5 tanaman) selama 72 jam. Perlakuan kontrol dilakukan dengan menempatkan imago C. basalis pada tanaman sehat selama 72 jam, kemudian dipindahkan pada tanaman sehat yang lain sejumlah 5 ekor per plot tanaman bersungkup (1 plot terdiri dari 5 tanaman) selama 72 jam. Masing-masing perlakuan diulang sebanyak tiga kali. Pada hari ke-15 setelah penularan, dilakukan uji PCR daun tanaman sampel dan isolat bakteri. Hasil pengujian semua sample daun menunjukkan negatif sehingga dipastikan bahwa C. basalis bukan merupakan vektor penyakit layu stewart pada tanaman jagung.

Kata kunci: Chaetocnema basalis, layu stewart, vektor

INTRODUCTION

Chaetocnema is a grass flea beetlewhich belongs to family of Chrysomellidae and distributes almost throughout the world. It is approximately 437 of about 630 species had been identified (Konstantinov *et al.*, 2011). In Indonesia, Kalshoven (1981) reported the presence of genus Chaetocnema populations on paddy, maize and grasses which were assumed as *Chaetocnema basalis*.

Stewart wilt disease caused by bacterium Pantoea stewartii has generated large problems for maize producing countries such as United States of America (USA), resulting yield loss up to 95%. Currently, this disease has been reportly distributed in all over the world such as: Austria, Bolivia, Brazil, Canada, Costa Rica, Guyana, Mexico, Peru, Puerto Rico, USA, China, India, Malaysia, Thailand, Vietnam, and Indonesia (Anonymous, 2016a). In Indonesia, it was considered as new disease on maize and had been reported distributing in Pasaman Barat, West Sumatera (Rahma, 2013), Bogor (Rahma et al., 2014), and Yogyakarta (Anonymous, 2015). Rahma (2013) detected the presence of this disease in maize producing center of West Sumatera with disease incidence ranging 1-15%.

Stewart wilt disease dispersed through seed and vector (Pataky, 2004). *Chaetocnema pulicaria* and *C. denticulata* were reported as vectors of this disease (Poos & Elliot, 1936). These two spesies have not been found yet in Indonesia. Genus Chaetocnema which was found associating with maize in Indonesia is *C. basalis*. This research was aimed to investigate whether *C. basalis* that grouped into same genus with *C. pulicaria* and *C. denticulata* is able to transmit Stewart wilt disease on maize.

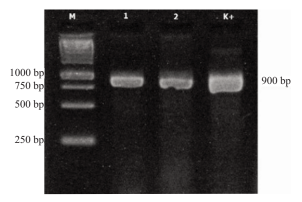


Figure 1. Molecular test of artificially inoculated of *Pantoea* stewartii to plant: 1=plant 1, 2=plant 2, M=1 kb DNA ladder, K+ = positive control

MATERIALS AND METHODS

Research was conducted in a green house, Faculty of Agriculture, Universitas Gadjah Mada, from Desember 2016 until Januari 2017. Imago of *C. basalis* were collected from maize growing areas in Sleman, Yogyakarta (110,4000 BT; 7,7099 LS). Bonanza variety of sweet corn was used as a test plant.

The 3-days old bacterial isolate on yeast dextrose calcium carbonate (YDC) medium was then diluted into sterile aquadest to obtain bacterial suspension. Turbidity level of suspension was measured using spectrophotometer at wavelength of 600 nm and OD of 0.3 which was estimated containing bacterial cell about 1.6×10⁹ cell ml⁻¹. The suspension was then injected on 8-days old maize plant, and confirmed to be infected with Stewart wilt through molecular assay (Figure 1), then 24 h-starved *C. basalis* were put on infected plant for 72 h. Five *C.basalis* were then transferred to plot of 8-days healthy plant (each plot consisted of 5 covered plants) for 72 h, furthermore considered as Treatment 1 (P1).

The collected and 24 h-starved *C. basalis* were placed on healthy plant for 72 h. Five insects were then transferred to plot of 8-days plant (each plot consisted of 5 covered plants) for 72 h, here in after used as control (P0), each treatment was repeated three times (Figure 2).

On the fifteenth day after transmission, leaves of each plant were sampled and analysed using two methods, i.e. direct plant extraction and isolation of Nutrient Agar (NA) medium. DNA extraction was performed using method developed by Goodwin *et al.* (1994).

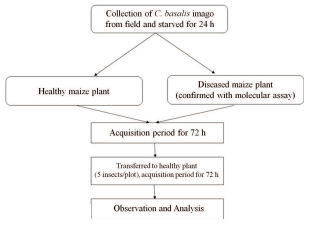


Figure 2. Flow chart of transmission assay of *Chaetocnema* basalis against *Pantoea stewartii* on maize

Leaf samples were washed with running water, surface sterilized with 70% ethanol for 1 min, rinsed 3 times with sterile water, and dried on sterile paper towels. Symptomatic leaf samples were aseptically cut into 10 mm×10 mm pieces and put into NA medium, incubated over night to allow release of bacteria into the medium. The bacterial suspension was then transferred and purified on YDC, the DNA of 3-days pured cultures were then extracted using CTAB method of Goodwin et al. (1994). Extracted pellet was checked to find out the presence of DNA by mixing the pellet with loading dye in 1:4 ratio, and then put into well of agarose 1% and electrophoreted (50 Volt for 40 min). Afterwards, DNA was amplified according to Coplin et al. (2002), molecular assay was carried out with Polymerase chain reaction (PCR) using primer of HRP3c 5'-GCG GCA TAC CTA ACT CC-3' and HRP1d 5'-CA CTC ATT CCG ACC AC-3', in following mixed composition: 1 µl of DNA pellet, 1 µl of primer HRP3c, 1 µl of primer HRP1d, 12.5 μ l of go taq reaction, and 9.5 μ l of dd H₂O. The protocol were as follows: predenaturation of 94°C for 1 min, denaturation of 94°C for 15 s, annealing of 55°C for 15 s, extension of 72°C for 30 s, and final extension of 72°C for 5 minutes. Steps of denaturation, annealing, and extension were cycled for 25 times.

PCR products were then put into well of 1.5% and electrophoreted (50 Volt for 45 min). Electrophoresis product visualized on UV transillluminator. Percentage of disease incidence was analysed with *t*-test with α level of 5% (p = 0.05).

RESULTS AND DISCUSSION

Transmission test showed that *C. basalis* could not transmit *P. stewartii*, pathogen of Stewart wilt. Molecular assay as shown on Figure 3, 4, and 5, in which all of samples and analyses methods showed negative results. Figure 3 on test control plants (P0),

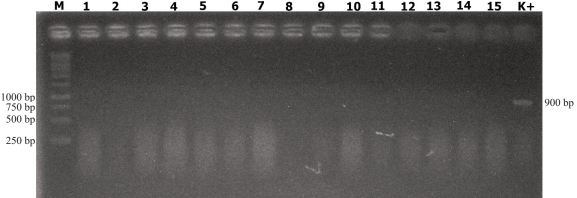


Figure 3. Molecular test of extracted leaves directly on control (P0): 1=P011, 2=P012, 3=P013, 4=P014, 5=P015, 6=P021, 7=P022, 8=P023, 9=P024, 10=P025, 11=P031, 12=P022, 13=P023, 14=P024, 15=P025; M=1 kb DNA ladder, K+ = positive control

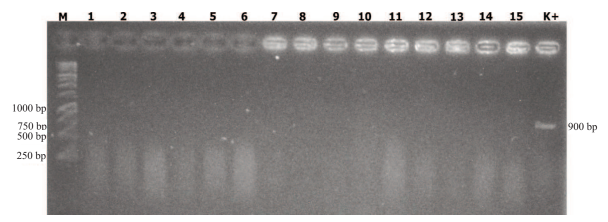


Figure 4. Molecular test of extracted leaves directly on treatments 1 (P1): 1=P111, 2=P112, 3=P113, 4=P114, 5=P115, 6=P121, 7=P122, 8=P123, 9=P124, 10=P125, 11=P131, 12=P122, 13=P123, 14=P124, 15=P125; M=1 kb DNA ladder, K+ = positive control

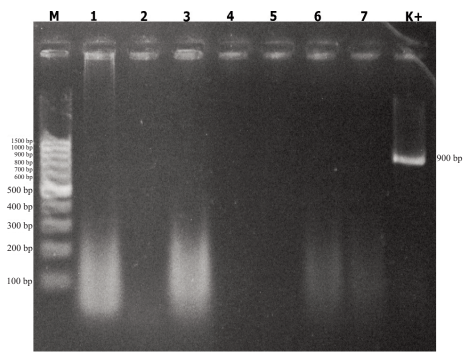


Figure 5. Molecular test of bacterial isolates from leaf isolation, 1=P115, 2=P122, 3=P124, 4=P131, 5=P132, 6=P133, 7=P135; M=100 bp DNA ladder, K+ = positive control

indicating negative results of all repetition and tested plant have been extracted and molecular analysed. Figure 4 showed that on treated plants (P1) also revealed negative results in all repetition and tested plant has been extracted and molecular analysed. Figure 5 showed that transmission test on all treatments and isolated plant samples, leaf samples exudating bacterial colony and then purified and extracted. Molecular assay expressed negative results on all sampled isolates.

Disease transmission through vector was a complicated interaction between insect, plant and pathogen (Martini *et al.*, 2015). Feeding manner of insect, imunity reaction of insect and plant, and characteristics of pathogen were some considerating factors. *C. basalis* was beetle with mouthparts type of bitting and chewing, acquition of pathogenic bacteria occured through its mouthpart during feeding process. Transmission happened through saliva containing pathogen which would penetrate via lession occuring for feeding process, so that the pathogen could spread through xylem vessels and cell cavities (Orlovskis *et al.*, 2015). However, feeding process of *C. basalis* did not generate the infection of Stewart wilt, the occurred lession was the enter for other pathogen.

The strange objects which entered into insect bodies could be pathogenic or symbiotic (Ammar *et al.*, 2014). They would establish defense systems

when foreign objects penetrated into their bodies (Chapman, 2013). The same reaction also occurred on plants when extraneous particles got into their tissues, i.e. utilizing induced resistance systems which were considered as a normal mechanism functioning in restriction of pathogen growth and spread on plant (Agrios, 2005).

P. stewartii associated with digestive tracts of insect vectors (foregut, midgut, hindgut, and malpigian tubule). Pathogen could be observed in midgut and hindgut until 12 days after acquisition (Ammar *et al.*, 2014). Pathogen would be brought by blood circulation and dissolved in saliva. Such condition took place when the pathogen was persistent in vector.

P. stewartii used two type III of secretion systems (T3SS) to establish colony on host plant and its vector. Colony establishment on host plant was initiated by producing injectisomere or pili to transfer effector protein to host plant. T3SS was important factor for pathogenicity of P. stewartii on maize plant. Secretion system of second T3SS was the establishment of Pantoea secretion island 2 (PSI-2) which was required by pathogen to be persistent in insect bodies. Mutagenesis of PSI-2 psaN gene was very influential in reducing the persistency of P. stewartii in gut of vector and the ability of beetle in transmitting the pathogen on host plant (Correa et al. 2012). The incompetence of Chaetocnema basalis in transmitting

Stewart wilt was much affected by the inability of *P. stewartii* to survive in *C. basalis*.

Generally, interaction of pathogen and vector was specific, in this case, *C. pulicaria* was the specific vector for *P. stewartii*. The presence of other *Chaetocnema* species in pathosystem of Stewart wilt was not necessarily vector for dispersal of *P. stewartii*. The existence of *C. basalis* on maize was also restricted as herbivores on those plants.

Stewart wilt dispersed through seed and vector, the seed dispersal intensity was less than 0.3%, so that insect vector was considered as main factor in distribution of this disease. *C. Pulicaria* and *C. denticulata* were effectively recognized in dispersing of this disease, and the spread by other vector was assumed less effective (Pataky & Ikin, 2003).

In several areas reporting the incidence of Stewart wilt disease, pathogen could not survive and establish in outside of North America (Pataky & Ikin, 2003). Austria, Greece, Poland, Rumania, dan Russia documented the incidence of this disease, however the pathogen could not survive in those regions. China, Vietnam, Thailand, and Malaysia had also recorded the prevalence of this disease, but the pathogen could not survive as well (Anonymous, 2016b). Pathogen survives in insect bodies and host tissue, those factors should be simultaneously for pathogen to be able to survive and establish in certain areas. Yet found of C. pulicaria and C. denticulata can be considered in evaluation the status of Stewart wilt disease. Although the disease prevalence has been reported, pathogen can not survive and establish in Indonesia. So that it is required sustainable monitoring of this disease and its vector. Moreover, the improvement of supervision on pathways at entry points should be performed in order to ensure that Indonesia could be free from Stewart wilt disease.

ACKNOWLEDGEMENT

This article is part of author's thesis. Author would like to express a great thank to Masanto and Bambang Trianom for the assistance in DNA extraction, as well as to Indonesia Agricultural Quarantine Agency as the donatur of scholarship.

LITERATURE CITED

Agrios, G.N. 2005. *Plant Pathology*. Fifth edition. Academic Press, New York. 922 p.

Ammar, E.D, S.A. Hogenhout, V. Correa & M.G Redinbough. 2014. Immunofluorescence Localization and Ultrastructure of Stewart's Wilt Disease Bacterium *Pantoea stewartii* in Maize Leaves and in its Flea Beetle Vector *Chaetocnema pulicaria* (Coleoptera: Chrysomelidae). *Journal of Microscopy and Ultrastructure* 2: 28–33.

Anonymous. 2015. *Laporan Pemantauan Organisme Pengganggu Tumbuhan Karantina 2015*. Balai Karantina Pertanian Kelas II Yogyakarta, Yogyakarta. 155 p.

Anonymous. 2016a. http://www.plantwise.org/Knowledge Bank/PWMap.aspx? speciesID=31363&dsID=21939 & loc=global, modified 05/02/17.

Anonymous. 2016b. *Pantoea stewartii* subsp. *stewartii*. *EPPO Bulletin* 8: 30–35. https://www.eppo.int/QUARANTINE/data_sheets/bacteria/ERWIST_ds.pdf., modified 04/09/16.

Chapman R.F. 2013. *The Insect "Structure and Function"*. Cambridge University Press, Cambridge, UK. 929 p.

Coplin, D.L, & D.R. Majerczak. 2002. Identification of *Pantoea stewartii* subsp. *stewartii* by PCR and Strain Differentiation by PFGE. *Plant Disease* 86: 304–311.

Correa, V.R, E.D. Ammar, S.A. Hogenhout, & M.G. Redinbough. 2012. The Bacterium *Pantoea stewartii* Uses Two Different Type III Secretion Systems to Colonize its Plant Host and Insect Vector. *Applied and Environmental Microbiology* 78: 6327–6336.

Goodwin, P.H. B.G. Xue, C.R. Kuske, & M.K. Sears. 1994. Amplification of Plasmid DNA to Detect Plant Pathogenic Mycoplasmalike Organisms. *Annals of Applied Biology* 124: 27–36.

Konstantinov, S.A, A. Baselga, V.V. Grebennikov, & S.W. Lingafelter. 2011. *Revision of the Paleartic Chaetocnema Species (Coleoptera: Chrysomellidae: Geluricinae: Alticini)*. Pensoft Publisher, Moscow. 366 p.

Martini, X, M. Hoffman, M.R. Coy, L.L. Stelinski, & K.S. Pelzstelinski. 2015. Infection of an Insect Vector with a Bacterial Plant Pathogen Increases its Propensity for Dispersal. *PLoS ONE* 10: 1–16.

Orlovskis, Z, M.C. Canale, V. Thole, P. Pecher, J.R.S. Lopes, & S.A. Hogenhout. 2015. Insect-Borne Plant Pathogenic Bacteria: Getting a Ride Goes beyond Physical Contact. *Current Opinion in Insect Science* 9: 16–23.

Pataky, J. & R. Ikin. 2003. *Pest Risk Analysis: The Risk of Introducing* Erwinia stewartii *in Maize Seed*. The International Seed Federation, Nyon, Switzerland. 79 p.

Pataky, J. K. 2004. *Stewart's Wilt of Corn. The Plant Health Instructor*. http://www.apsnet.org/edcenter/intropp/lessons/prokaryotes/Pages/StewartWilt.aspx. modified 09/05/17.

Poos, F.W & C. Elliott. 1936. Certain Insect Vectors of Aplanobacter Stewarti. *Journal of Agricultural Research* 52: 585–608.

Rahma, H. 2010. *Penyebaran Penyakit Stewart oleh Bakteri* Pantoea stewartii *sebagai Penyakit Baru pada Tanaman Jagung (Zea mays) Studi Kasus di Pasaman Barat.* http://repository.unand. ac.id/4112/1/ARTIKEL_Haliatur.doc. Project Report. Universitas Andalas, Padang. 11 p. (Unpublished).

Rahma, H, M.S. Sinaga, M. Surahman, & Giyanto. 2014. First Report of Stewart's Wilt of Maize Caused by *Pantoea stewartii* subsp. *Stewartii* in Bogor District, Indonesia. *Journal of the International Society for Southeast Asian Agricultural Sciences* 20: 131–141.