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Molecular Identification and Sequence Analysis of *Tobacco Leaf Curl Begomovirus* from Jember, East Java, Indonesia

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Begomovirus had been proved as the causal agent of leaf curl disease in tobacco in Indonesia, or commonly in Indonesia called as penyakit krupuk tembakau. Association of Begomovirus with the disease was further confirmed by sequence analysis. Amplification of the virus was conducted following whitefly (Bemisia tabaci Genn.) transmission. Fragment of DNA 1.6 kb was amplified by polymerase chain reaction (PCR) located within the replication initiator protein gene and coat protein gene (top region). Conserved sequence of stem loop region was found, included nonanucleotide sequence TAATATTAC present in all geminiviruses. Begomovirus associated with leaf curl disease in tobacco showed the closest relationship with Ageratum yellow vein virus - Zimbabwe, a strain of Tobacco leaf curl virus from Southern Africa. It was also known that Begomovirus associated with leaf curl disease in tobacco from Jember, East Java was different from other Indonesian Begomoviruses reported earlier.

Key words: Begomovirus, leaf curl disease

INTRODUCTION

Geminivirus is a group of plant viruses with a distinct morphological characters. Its twinned isometric particles consists of circular single-stranded (ss) DNA genomes (Bock 1982). They are classified into four genera, i.e. Mastrevirus, Curtovirus, Begomovirus, and Topocuvirus, based on their vector relationship, host range and genome organization (Van Rogenmortel 2000). Members of the genus Begomovirus are transmitted by the whitefly Bemisia tabaci Genn. (Hemiptera:Aleyrodidae) and infect dicotyledonous plants. Diseases which were caused by whitefly-transmitted geminiviruses (WTG) have become a serious constraints to crops in tropical and subtropical areas throughout the world (Idris & Brown 1998; Samretwanich et al. 2000). In the last five years, geminivirus has been reported to cause significant yield loss in chilli pepper and tomato in vegetable growing areas in Java (Hidayat, unpublished data). However, tobacco leaf curl disease caused by geminivirus in Indonesia was reported earlier by Thung in 1932 (Trisusilowati et al. 1990). That was probably the first report on geminivirus infection in Indonesia. In 1984, Tobacco leaf curl virus (TLCV) caused serious damage in Bojonegoro, East Java, with up to 30% disease incidence (Poerbokoesoemo 1984 in Trisusilowati et al. 1990). The symptoms of TLCV infection include leaf curling, vein banding, uneven leaf surface, and rigid leaves. The disease may effect not only the yield but also the quality of tobacco leaf, especially when the leaves were targeted for cigar wrapping.

Observation conducted by Trisusilowati *et al.* (1990) using electron microscope demonstrated that a unique twin isometric particle was associated with leaf curl disease in tobacco. They

*Corresponding author. Phone: +62-251-629363, Fax: +62-251-629362, E-mail: srihendrastutihidayat@gmail.com were also proved that the virus was transmitted through whitefly, *B. tabaci*. Furthermore, Aidawati *et al.* (2002) carried out a transmission study to elucidate the characteristics of TLCV transmission by its vector, *B. tabaci*. A single whitefly was able to transmit the virus and the efficiency of transmission were increased when the number of adult whiteflies was increased up to 20 per plant. Inoculation access period of 1 h could cause transmission up to 20% and the optimum inoculation access period was 12 h. Acquisition access period of 30 min resulted in 70% transmission while 100% transmission occurred with a 24 h-acquisition access period. The virus was proven to be persistently but not transovarially transmitted.

The evidence above has confirmed the association of geminivirus in tobacco leaf curl disease in East Java. Later on, polymerase chain reaction (PCR) using specific degenerate primers for geminivirus, PAL1v1978, and PAR1c715, was successfully amplified a 1.6 kb DNA fragment from infected tobacco plants as well as viruliferous whiteflies (Aidawati *et al.* 2002).

In this paper we reported our attempt to clone and sequence the geminivirus causing tobacco leaf curl in Jember, East Java. Its relationship with other geminiviruses, especially those reported from Indonesia and those causing leaf curl disease on tobacco from other part of the world, was analysed based on nucleotide sequence homology.

MATERIALS AND METHODS

Collection and Maintenance of Virus Isolate. Leaves showing TLCV symptom were collected from tobacco field at Arjasa and Tegal Gede villages, Jember, East Java. The virus was maintained on tobacco plants (*Nicotiana tabacum*) H382 by insect transmission (Aidawati *et al.* 2002). Adults *B. tabaci*

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were obtained from broccoli plants in Bogor and identified using identification key of Martin (1987). The insect were reared on tobacco and broccoli (*Brassica oleareceae* var. *italica*) plants in whitefly-proof cages. Tobacco plants for maintaining virus isolates were grown in a whitefly-proof screenhouse.

PCR-Based Detection Using Geminivirus Degenerate Primers. DNA template for PCR was prepared from infected tobacco plants following method developed by Dellaporta et al. (1983). The DNA pellet was resuspended in 50 μl of sterile distilled water. Amplification of geminivirus genome was proceeded using a pair of degenerate primers designed for the amplification of the DNA A genomic component, pAL1v 1978 (5'GCATCTGCAGGCCCACATYGTCTTYCCNGT3') and pAR1c 715 (5'GATTTCTGCAGTTDATRTTYTCRTCCAT CCA3') (Rojas et al. 1993). Amplification with PCR technique was carried out in a 25 µl reaction mixture containing 1 µl of sample DNA solution and 0.2 µM of each primer using Ready To Go PCR kit (Amersham Life Science). PCR was performed in thermalcycler Gen Amp PCR System 9700 (Perkin Elmer) with 30 cycles of melting, annealing and DNA extension at 94 °C for 1 min, 55 °C for 2 min, and 72 °C for 2 min, respectively. PCR products were then analysed by electrophoresis in 1% agarose gels in Tris-buffer EDTA.

Cloning and Sequencing of DNA from TLCV Isolate. DNA fragments of approximately 1.6 kbp, as a product of PCR amplification, was cloned into pGEM- T Easy vector (Biorad) with *Pst*I cloning site. Selected DNA clone was then sequenced by the dideoxy nucleotide chain termination method (Maniatis *et al.* 1982). A 1.6 kbp viral DNA fragment was completely sequenced on both strands using internal primers that was designed based on the prior nucleotide sequences.

Once the sequence was completed, it was compared with those of other whitefly-transmitted geminiviruses available in Genbank (Table 1) using Clustal W program version 1.82 European Bioinformatics Institute (EMBL-EBI: www.ebi.ac.uk/serve/clustalW). Phylogenetic analyses was conducted with PAUP program version 4.0 b4a using Maximum-Parsimony method with heuristic searches using the TBR branch swapping option and 10,000 random addition sequences.

RESULTS

Symptoms of TLCV-Infected Tobacco Plants. Whitefly transmission of TLCV to tobacco plants was conducted following a procedure established by Aidawati *et al.* (2002). Whiteflies were given a 24 h acquisition feeding period on virus source before transferred to healthy tobacco for a 48 h inoculation feeding period. In general, the symptoms of TLCV on tobacco plants (*N. tabacum*) H382 was developed 7-10 days after inoculation feeding period. Infected plants will show leaf malformation involving upward curling and vein banding with crinkle-like symptom (Figure 1).

Identification of TLCV by PCR and Sequence Characterization. Specific DNA fragments of 1,600 bp was successfully amplified from infected tobacco plants using geminivirus-specific degenerate primers pAL1v 1978 and pAR1c 715. The amplified DNA fragment, denoted as "top

region", include part of replicase region, full common region, and part of coat proteion region. The PCR product was then cloned into the PGEM-T easy vector. Following screening of recombinant DNA, a clone identified as pTobT8 was selected for viral sequence analyses. Nucleotide sequence of TLCV (pTobT8) from base 1 to 1474 was determined and submitted to Genbank (Accession No. AB246171). The nucleotide sequence of 33 base stem-loop region was found in the sequence of pTobT8 as well as the conserved nonanucleotide sequence TAATATTAC which is known as TATA-box region (Figure 2). Both stem-loop and TATA-box regions has been found in all geminiviruses sequenced so far (Ikegami et al. 1988), as it was observed in other sequences of begomoviruses from Indonesia. The isolate of Begomovirus associated with leaf curl disease of tobacco in Jember, East Java was tentatively denoted as Tobacco leaf curl Indonesia virus-Jember (TLCIV- Jbr).

Phylogenetic Analyses of TLCIV-Jbr. Relationship between TLCIV-Jbr and other selected begomovorises was evaluated based on "top region" sequences. Although conserved sequence of the geminivirus's common region was observed, it was found that TLCIV-Jbr was different with the other geminiviruses that has been reported from Indonesia, i.e those infecting chillipepper, tomato, and weed (*Ageratum conyzoides*). Analyses of sequence identity using Maximum-Parsimony method furthermore revealed that TLCIV-Jbr was distinct from any other tobacco leaf curl viruses (Figure 3).

DISCUSSION

A geminivirus was first demonstrated to be the causative agent of tobacco leaf curl disease in Indonesia in 1990. However, the disease has been reported earlier by Thung in 1932 and caused serious problem in tobacco plantation in East Java (Trisusilowati *et al.* 1990). Infection of TLCV may cause various symptoms including leaf curling, vein banding, uneven leaf surface, and rigid leaves (Aidawati *et al.* 2002). Therefore, in Indonesia the disease is commonly called *penyakit kerupuk*. Variation on disease symptom developed from TLCV infection has also been documented. A perplexing feature of tobacco leaf curl since its earliest reports has been the observed variation in symptom severity. For instance in Southern Africa, at least three symptom phenotypes in tobacco have been identified that could be classified as

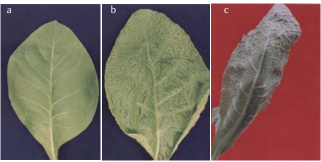


Figure 1. Symptom of *Tobacco leaf curl virus* on tobacco plants (N. tabacum) H382. a. Healthy leaf, b. Leaf showing mild symptom, c. Leaf showing severe symptom.

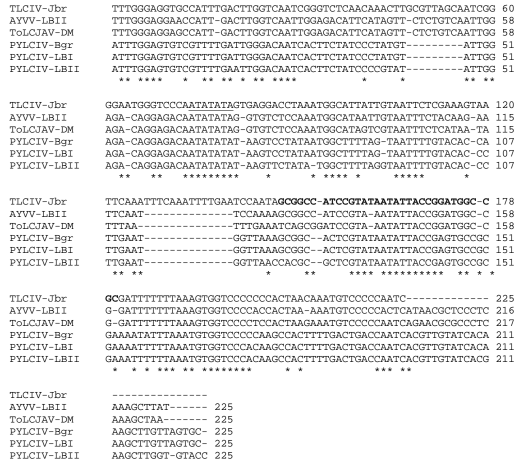


Figure 2. Alignment of nucleotide sequences of the common region of Tobacco leaf curl Indonesia virus-Jember (TLCIV-Jbr) with other reported Indonesia begomoviruses as listed in Table 1. The alignment showing TATA sequences (underlined sequences) and the stemloop region (bold letters for TLCIV-Jbr).

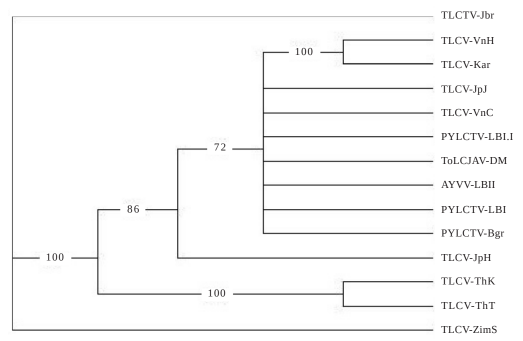


Figure 3. Phylogenetic tree based on the alignments of nucleotide sequences of "top region" of Tobacco leaf curl Indonesia virus-Jember (TLCIV-Jbr) with other begomoviruses as listed in Table 1.

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Table 1. List of Begomoviruses used for viral sequence analysis

Genbank accession number	Organism	Nucleotide length (bp)	Geography origin	Host plant	Acronim
AB189845	Pepper yellow leaf curl Indonesia virus	1563	Indonesia: Lembang, West Java	Lycopersicon esculentum	PYLCIV-LBI
AB246170	Pepper yellow leaf curl Indonesia virus	1491	Indonesia: Segunung, West Java	Capsicum frutescens	PYLCIV-Bgr
AB189913	Ageratum yellow vein virus-Indonesia	1557	Indonesia: Lembang, West Java	Ageratum conyzoides	AYVV-LBII
AB189848	Tomato leaf curl java virus-Magelang	1562	Indonesia: Magelang, Central Java	L. esculentum	TolCJAV-DM
IB189850	Pepper yellow leaf curl Indonesia virus	1555	Indonesia: Lembang, West Java	C. annuum	PYLCIV-LBII
AM051086	Tobacco leaf curl virus-[Vietnam]	526	Vietnam: Cao Bang Province	N. tabacum	TLCV-VnC
AM051085	Tobacco leaf curl virus-[Vietnam]	526	Vietnam: Hang Tay Province	N. tabacum	TLCV-VnH
AY007616	Tobacco leaf curl virus-[India]	771	India: Karnataka	Tobacco	TLCV-Kar
AY633751	Tobacco leaf curl virus-[Thailand]	771	Thailand: Tak Province	Tobacco	TLCV-ThT
AY633750	Tobacco leaf curl virus-[Thailand]	771	Thailand: Kamphaensaen	Tobacco	TLCV-ThK
AB108838	Tobacco leaf curl virus-[Japan]	1570	Japan: Kagoshima	Honeysuckle	TLCV-JpH
E15418	Tobacco leaf curl virus-[Japan]	2766	Japan	Eupatorium japonicum	TLCV-JpJ
AF077749	Tobacco leaf curl virus-[Japan]	777	Zimbabwe	Ageratum conyzoides	TLCV-ZimS

tobacco leaf curl. Based on sequence analyses, different virus strains are unlikely to be found in the infected plants (Paximadis & Rey 1997). The discovery of defective DNA molecules in leaf curl-affected tobacco plants may possibly explain the variable symptom severity. The environmental factors and type of tobacco varieties may also contribute to symptom severity. Agroinoculation study using a full-length genomic DNA -A derived from a field tobacco plant exhibiting mild symptoms in N. tabacum, but N. benthamiana did exhibit severe leaf curl symptoms (Paximadis & Rey 2001). The possibility of satellite virus similar to DNA β associated with Ageratum yellow vein virus from Singapore (Saunders et al. 2000), Cotton leaf curl virus from Pakistan (Briddon et al. 2001), or Tomato leaf curl Java virus from Indonesia (Kon et al. 2006) might contribute to symptom severity. Therefore it is also interesting to determine whether TLCIV-Jbr infection is associated with the presence of its DNAβ.

Begomoviruses has been reported to infect tomato, chilli pepper, and *A. conyzoides* in Indonesia and it was reported that tobacco plant, *N. tabacum*, is one of the host plants of all those begomoviruses (Rusli *et al.* 1999; Aidawati *et al.* 2002; Hidayat *et al.* 2002; Haerani & Hidayat 2003; Sulandari *et al.* 2006). Conserved sequence in the common region was observed for all Indonesia begomoviruses (Figure 2); however, the phylogenetic analysis revealed that TLCIV-Jbr was grouped in different cluster from those of other *Begomovirus* isolates reported from Indonesia (Figure 3). We understand from previous sequence analysis that *Begomovirus* species from Indonesia, *Tomato leaf curl Indonesia virus* (TolCIDV), *Tomato leaf curl Java virus* (TolCJAV), *Ageratum yellow vein virus-Java* (AYVV-[Java]), PYLCIV-Bgr, had close relationship one to each other (Sukamto *et al.* 2005; Kon *et al.* 2006; Hidayat

et al. 2006). The fact that TLCIV-Jbr was not clustered with the other Indonesian begomoviruses might indicated that TLCIV-Jbr was possibly different Begomovirus species. Furthermore, when analysis was conducted using nucleotide sequences of other geminiviruses causing leaf curl disease on tobacco, TLCIV-Jbr was clustered only with TLCV-Zims (Figure 3). This facts strengthened the hypotheses that there was a high degree of genetic diversity among Begomovirus group, which might be emerged due to a high degree of pre-exisiting genetic diversity among begomoviruses or possible recombination between different virus species during infection (Ribeiro et al. 2003). The knowledge regarding genetic diversity among members of geminivirus infecting different crops and weeds in Indonesia should be considered in developing disease control strategies.

REFERENCES

Aidawati N, Hidayat SH, Suseno R, Sosromarsono S. 2002. Transmission of an Indonesian isolate of tobacco leaf curl virus (Geminivirus) by *Bemisia tabaci* Genn. (Hemiptera: Aleyrodidae). *Plant Pathol J* 18:231-236.

Bock KR. 1982. Geminivirus diseases. Plant Dis 66:266-270.

Briddon RW et al. 2001. Identification of DNA components required for induction of cotton leaf curl disease. Virology 285:234-243.

Dellaporta SL, Wood J, Hicks JB. 1983. A plant DNA minipreparation: Version II. *Plant Mol Biol Rep* 1:19-21.

Haerani, Hidayat SH. 2003. Geminivirus causing yellow disease on *Ageratum conyzoides*: molecular detection and transmission study on some solanaceae plants. *J Fitopatol Indones* 7:65-70.

Hidayat SH, Chatchawankanpanich O, Rusli E, Aidawati. 2006. Begomovirus associated with pepper yellow leaf curl disease in West Java, Indonesia. J Mikrobiol Indones 11:87-90.

Hidayat SH, Mulyati S, Kadir M, Sulandari S, Aidawati N. 2002. Tobacco leaf curl virus from Indonesia: its detection and host range study. J Agrik 13:157-161.

- Ribeiro SG et al. 2003. Distribution and genetic diversity of tomatoinfecting begomoviruses in Brazil. Arch Virol 148:281-295.
- Phytopathology 88:648-657. Rojas MR, Gilbertson RL, Russell DR, Maxwell DP. 1993. Use of Ikegami M, Morinaga T, Miura K. 1988. Potential gene products of whitefly-transmitted geminiviruses. Plant Dis 77:340-347.
- bean golden mosaic virus have higher sequence homologies to those of tomato golden mosaic virus than those of cassava latent virus. Virus Genes 1:191-203.

Idris AM, Brown JK. 1998. Sinaloa tomato leaf curl geminivirus: biological and molecular evidence for a new subgroup III virus.

- Kon T, Hidayat SH, Hase S, Takahashi H, Ikegami M. 2006. The natural occurrence of two distinct Begomovirus associated with DNAB and a recombinant DNA in a tomato plant. Phytopathology 96:517-525.
- Maniatis T, Fritsch EF, Sambrook J. 1982. Molecular Cloning: a Laboratory Manual. New York: Cold Spring Harbor Laboratory
- Martin JH. 1987. An identification guide to common whitefly pest species of the world (Homoptera: Aleyrodidae). Trop Pest Manag 33:298-322.
- Paximadis M, Rey MEC. 1997. Aetiology of tobacco leaf curl in Southern Africa. Ann Appl Biol 131:449-457.
- Paximadis M, Rey MEC. 2001. Genome organization of tobacco leaf curl Zimbabwe virus, a new, distinct monopartite Begomovirus associated with sub genomic defective DNA molecules. J Gen Virol 82:3091-3097.

- degenerate primers in the polymerase chain reaction to detect
- Rusli ES, Hidayat SH, Suseno R, Tjahjono B. 1999. Virus Gemini asal cabai: kisaran inang dan cara penularan. Bulletin HPT 11:26-31.
- Samretwanich K, Chiemsombat P, Kittipakorn K, Ikegami M. 2000. A new geminivirus associated with a yellow leaf curl disease of pepper in Thailand. Plant Dis 84:1047.
- Saunders K et al. 2000. A unique virus complex causes Ageratum yellow vein disease. Proc Natl Acad Sci USA 97:6890-6895.
- Sulandari S, Suseno R, Hidayat SH, Harjosudarmo J, Sosromarsono S. 2006. Deteksi dan kajian kisaran inang virus penyebab daun keriting kuning cabai. Hayati 13:1-6.
- Sukamto et al. 2005. Begomovirus associated with leaf curl disease of tomato in Java, Indonesia. J Phytopathol 153:562-566.
- Trisusilowati EB et al. 1990. Transmission, serological aspects and morphology of the tobacco krupuk virus. Indones J Agric 1:75-
- Van Rogenmortel MHV et al. 2000. Virus Taxonomy. Seventh Report of the International Committee on Taxonomy of Viruses. San Diego: Academic Pr.