Coconut Cadang-Cadang Disease and Its Viroid Agent

In the early 1930s a devastating epidemic of a lethal disease of coconut palm (Cocos nucifera L.) was reported from southern Luzon in the Philippines. It was appropriately named “cadang-cadang” disease from the word gadan-gadan of the local dialect meaning “dying” (13). Anecdotal reports indicate that the disease probably occurred a few years earlier in the Philippines, at about the time a somewhat similar epidemic was reported on the island of Guam. The disease in Guam was called “tinangaja” and was extremely destructive to the island’s coconut population (2). Both diseases have now been shown to be caused by related viroids: coconut cadang-cadang viroid (CCCVd) and coconut tinangaja viroid (CTiVd).

Economic Importance

It has been estimated that over 30 million coconut palms have been killed by cadang-cadang since it was first recognized (22). Based on average yield and copra prices, the loss of production has been valued at about $80–$100 (U.S.) for each planting site occupied by an infected tree (13), an amount exceeding 2 months’ salary for an unskilled worker in the Philippines. According to G. Persley (in a paper presented in 1989 to the Technical Advisory Committee of the Consultative Group on International Agricultural Research), at least 96% of the total world copra production is provided by smallholders. Because coconut is both an important subsistence and a major cash crop in many developing countries, cadang-cadang must be considered a serious economic threat (21).

Tinangaja disease is still prevalent on Guam. Its economic importance is less evident, however, because coconuts are not commercially produced there at present (2).

Symptoms and Host Range

Cadang-cadang develops slowly in palms and cannot be unequivocally identified on the basis of symptoms at a single observation. The disease progresses through three well-defined stages (13) (Fig. 1). In the early stage, nuts become rounded, with characteristic equatorial scarifications, and the first necrotic, translucent, bright yellow leaf spots appear. In the mid stage, inflorescences become necrotic, nut production ceases, new frond production slows down, and leaf spots become larger and more frequent so that fronds begin to appear chlorotic from a distance. In the late stage, preceding death, leaf spots are almost confluent; the whole crown is distinctly yellowish or bronze-colored and very much reduced in size and number of fronds. The early stage lasts 2–4 years, the mid stage approximately 2 years, and the late stage about 5 years. The overall time from first symptoms to death of the tree is about 8 years for 22-year-old palms and about 16 years for 44-year-old palms (22). Usually, palms become naturally infected only after they have reached the age of flowering. In the rare cases where younger palms become infected, they are stunted and fail to produce inflorescences, although they survive well past the age of first flowering. Many cultivars and hybrids have been tested for susceptibility by inoculation in the Philippines, but none have shown any indication of immunity to the viroid. The only known variation in the symptoms of cadang-cadang is a more severe type of lamina reduction that occurs rarely in the field but has been observed in about 3% of mechanically inoculated coconut palms at the Albay Research Center in the Philippines. Parts of the fronds consist only of the midribs, thus showing a “brooming” syndrome (Fig. 2).

Tinangaja disease shows symptoms similar to those of cadang-cadang except that the affected trees bear spindle-shaped nuts with a reduced or absent kernel (2) (Fig. 3). There are a number of diseases of coconut palm with unknown etiology, some of which show abnormalities reminiscent of cadang-cadang or tinangaja, for example, narrow nuts as for Tatipaka disease and a decline as for Kerala wilt, both diseases occurring in India. Some of them have tested negative by gel electrophoresis (3) and solution molecular hybridization assay (12), but they need to be reevaluated by the latest molecular diagnostic methods for the presence of viroid.

Oil palm (Elaeis guineensis Jacq.) naturally infected or inoculated with CCCVd in the Philippines develops bright orange leaf spots that are larger and more numerous on the older fronds (Fig. 4), nut production ceases, and the tree eventually dies (14). Naturally infected buri palm (Corypha elata Roxb.) shows chlorotic leaf spots and stunting (14). Palm species successfully inoculated with CCCVd include oil palm, buri palm, betel nut palm (Areca catechu L.), golden cane palm (Chrysalidocarpus lutescens H. Wendl.), date palm (Phoenix dactylifera L.), royal palm (Roystonea regia (Kunth) Cook), and Manila palm (Veitchia merrillii (Becc.) Moore) (6,13).

Preliminary evidence suggests that several herbaceous monocotyledonous species growing near coconut palms infected with cadang-cadang (Fig. 5) occasionally contain viroidlike molecules with sequence similarity to CCCVd (J. M. B. Rodriguez, D. Hanold, and J. W.
Randles, unpublished). Such plants have no apparent symptoms. Isolation and characterization of these molecules and a comparison with CCCVd isolates is in progress.

Distribution, Epidemiology, and Control

The present distribution of cadang-cadang (Fig. 6) shows the northernmost boundary to be at the latitude of Manila and the southernmost at the latitude of Homonhon Island. The northern and southern boundaries are monitored annually. This is particularly important because of the proximity of the disease to the major coconut and oil palm growing area of Mindanao (16). Rates of spread vary such that some sites show little expansion, while others show active epidemics with outward movement of boundaries at about 0.5 km per year. Diseased palms are not clustered.

Little else is known about the epidemiology of cadang-cadang. The available data suggest that it may not be spread by any one specific route, but that it could be distributed by a variety of means (16). For example, the viroid can be detected in the husk (J. M. B. Rodriguez, unpublished) and embryo (D. Hanold, unpublished) of nuts and is seed-transmitted at the low rate of about one in 300 (18). It has been detected in purified pollen (D. Hanold, unpublished), and trials to determine whether transmission occurs via pollination are in progress at the Albay Research Center. Although many tests have failed to identify any insect vector (22; E. P. Pacumbaba, unpublished), it still seems possible that the viroid could be transmitted unspecifically by certain coleopterous insects through feeding wounds. The possibility that CCCVd can be transmitted by mechanical damage from cultural practices has yet to be adequately tested. Experimental plots have been set up in the Philippines with mechanically inoculated infector plants distributed in different arrays to determine the rate and pattern of spread of the viroid into the healthy population.

No control measures are known. Early attempts at control by eradication failed to prevent spread (22). As mentioned above, no resistant cultivars are available for replanting or as breeding material. No vector is known, so vector control is not an option. At present, the replacement of infected palms is the only practice that allows production to continue in affected areas, since the rate of spread in the new plantings is not influenced by the proximity of infected palms (12). Recently detected CCCVd-related viroids may be mild strains that could be used in mild strain protection, and this is an option for further study.

Molecular Characteristics of the Pathogen

It was reported in 1975 that viroidlike ribonucleic acid (RNA) was associated with cadang-cadang. This RNA was concentrated in a polyethylene glycol 6000 precipitate of leaf extracts, deproteinized, and identified by electrophoresis in polyacrylamide gels (PAGE) (11). The RNA was of low molecular weight, single-stranded (19), predominantly circular (17), and infectious (15). It was also found in naturally infected oil and buri palm (14), and a similar RNA was shown to be associated with tinangaja (3). These results, and the insensitivity of the disease to tetracyclines (15), were the first evidence that the disease had a viroid etiology.

Further analysis of the viroid on gels with a high resolving capacity showed that it comprises two monomeric "slow" and two monomeric "fast" forms (Fig. 7A), as well as dimeric forms of each monomer. Determination of their nucleotide sequences (5) (Fig. 7B) showed that the fast RNAs contain either 246 nucleotides or, by insertion of an additional cytosine residue at a specific site in the molecule, 247 nucleotides. The

Fig. 2. Reduction of lamina in an inoculated palm, giving the unusual "brooming" syndrome.

Fig. 3. Nut abnormalities associated with tinangaja disease: (A) (Left) Scarifications and (right) characteristic spindle shape. (B) Sections of nuts from (left) healthy and (right) diseased palm showing lack of kernel development. (Courtesy G. Boccardo)

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slow RNAs of 296 and 297 nucleotides also differ by the addition of the cytosine, and the slow RNAs are directly derived from the corresponding fast forms by a duplication of the right-hand end of the molecule (Fig. 7).

In coconut palms, frond position is age-related, and an analysis of consecutive fronds in crowns of palms in the early, mid, and late stages of the disease showed that the four distinct molecular forms of the viroid occurred in a progression associated with the stage of the disease (7,13) (Fig. 7A). At the early stage, only the fast forms are found, but on subsequently emerging fronds the slow forms appear and eventually replace the fast forms as the disease proceeds through the mid to late stages. The forms of viroid detected in each frond remain the same as the frond ages and moves downward through the crown. It would be of interest to know whether duplication of the right-hand end of the viroid is involved in the transition of symptoms from the early to the mid and late stages, that is, to inflorescence necrosis and loss of nut production.

CCCVD has the central nucleotide sequence that is conserved among all viroids of the potato spindle tuber viroid group (9). Its physical properties and melting profile are similar to those of other viroids (20), with an intermediate structural form arising from annealing of complementary inverted sequences either side of the central conserved region. It is most closely related by size and nucleotide sequence to CTiVd, which has been shown to be 64% homologous with CCCVD (8). Compared with the other viroids, CCCVD has several unique features. As the 246 nucleotide form, it is the smallest known pathogen; with CTiVd, it is both the only viroid known to affect monocotyledonous plants and the only viroid that is lethal in a host plant; and its pattern of changing molecular forms has not been reported for any other viroid.

The infectivity of purified forms of CCCVD and their ability to induce disease (Fig. 8) has been demonstrated by high-pressure injection into folded leaf tissue (6,10,15), with the highest efficiency being achieved by injection close to the meristem of young sprouts. Increasing the concentration of RNA in the inoculum or increasing the number of injections raises the rate of successful inoculations. The 246/247 nucleotide form apparently has a higher specific infectivity than the others (10).

**Diagnosis**

Because diagnosis of CCCVD by symptoms is unreliable, molecular diag-

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**Fig. 4.** Orange leaf spot development on fronds of oil palm inoculated with coconut cadang-cadang viroid; frond age increases from left to right.

**Fig. 5.** _Alpinia_ sp. (foreground) associated with coconut palms infected with cadang-cadang viroid. RNA related to coconut cadang-cadang viroid has been found occasionally in this and other monocotyledonous weeds.

**Fig. 6.** Distribution of cadang-cadang disease (gray areas) in the Philippines.
nositic methods have been developed that rely on detection of the viroid in test samples. When cellular nucleic acids purified from approximately 1 g of coconut tissue are analyzed on polyacrylamide gels, the molecular forms of CCCVd can be identified by their relative electrophoretic mobility. With use of the highly sensitive silver stain (6), CCCVd can be detected up to about 6 months before the appearance of initial symptoms. Normally, leaf material is used for routine analysis, but the viroid seems to be present in most host tissues. The method of diagnosis by gel electrophoresis has been successfully adapted for work in a mobile laboratory and is now the method of choice for epidemiological field surveys and for the yearly monitoring of disease boundaries in the Philippines.

CCCVd has been cloned (5) and can also be amplified by the polymerase chain reaction (PCR) (J. M. B. Rodriguez, J. W. Randles, and D. Hanold, unpublished). The clones and PCR product can be used as templates for synthesis of radioactively labeled complementary RNA or DNA probes. These are used in hybridization assays to detect nucleotide sequences similar to those of CCCVd (1). Extracts to be tested are applied to a supporting membrane, and the presence of viroid is detected by its specific hybridization with probe (Fig. 9A). The radioactive label causes darkening of exposed x-ray film, while samples without viroid show no signal. This diagnostic method is called “dot blot hybridization”; because it is much more sensitive than electrophoresis, viroid can be detected many months before symptoms appear.

The most conclusive results are obtained by a combination of these two methods in which electrophoretically separated nucleic acids are transferred to a membrane filter and then hybridized with radioactive probe (Fig. 9B). This “gel electrophotoblot hybridization” method tests only for the size and structure, as well as sequence homology, and both one- and two-dimensional (Fig. 9C) gel electrophoresis (13) can be used. It is currently the most reliable and sensitive tool for finding CCCVd-related molecules in any type of host and tissue.

**CCCVd-related Sequences in New Locations**

We have recently found viroid-like molecules related to CCCVd in several areas of the southwest Pacific (4). Electrophores of both one- and two-dimensional gels (Fig. 9C) have identified molecules of a mobility similar to that of CCCVd in oil palms in commercial plantations, in coconut palms, and in several herbaceous monocotyledons. The affected oil palms show symptoms resembling those in naturally infected oil palms (Fig. 10) in the Philippines, but the coconut palms are without the typical cadang-cadang syndrome (Fig. 11). The question as to whether the oil palm and coconut isolates are the same, as well as how closely they are related to CCCVd, must be evaluated by sequencing studies. It must be emphasized that no epidemic of cadang-cadang has been reported in these countries. Factors such as the viroid strain, host reaction, environment, or absence of an efficient putative vector may explain this. Identification of the factors preventing the development of a serious outbreak may eventually help in the development of control measures for CCCVd in the Philippines. For example, certain environmental requirements for viroid transmission may be essential or mild strains of CCCVd may be identified that could be used for “mild strain protection” when present in a host plant before challenge by the severe strain.

**Conclusions**

The viroid etiology of cadang-cadang disease of coconut palm in the Philip-
Fig. 9. Diagnosis of coconut cadang-cadang viroid: (A) Dot blot hybridization assay; 
H = healthy control, D = diseased coconut palm. (B) Electroblot hybridization assay 
showing coconut cadang-cadang viroid in a naturally infected oil palm; s = slow (296/7 
nucleotides) and f = fast (246/7) forms of marker coconut cadang-cadang viroid, 
both showing monomers (lower major bands) and dimers (upper minor bands). (C) 
Two-dimensional polyacrylamide gel electrophoresis in association with an electroblot. 
(Top) Healthy palm; (bottom) coconut cadang-cadang viroid-related sequences in a 
naturally infected oil palm from the South Pacific; arrow indicates viroid-like nucleic 
acid that, because of its structure, moves at a different rate from that of other nucleic 
acids in the sample. Some nonspecific binding of probe in regions other than the 
viroid-like RNA is due to low-stringency washing conditions after hybridization in this 
experiment.

Fig. 10. Orange spotting of oil palm in a commercial plantation: (A) Whole crown. 
(B) Increasing size and density of spots on fronds of (left to right) increasing age.

Fig. 11. A coconut palm from the southwest Pacific containing coconut cadang-
cadang viroid-related sequences. It does not show the typical syndrome, but some 
necrosis of inflorescences and a tendency toward spindle-shaped nuts, as for 
tinangaja, are apparent.

Pines has been established, but no control measures are available and the disease 
continues to spread. Our recent identification of viroid-like sequences related 
to CCCVd in the Pacific outside the known cadang-cadang area raises a 
number of new questions about the origins, epidemiology, and pathogenicity 
of these CCCVd-related sequences. We consider that until more is known about 
the epidemiology of these viroid-like molecules, an embargo should be placed 
on the uncontrolled movement of germ plasm between countries. This is particu-
larly important because the viroid infections appear to be latent and because 
their pathogenicity in new areas is unpredictable.

Current breeding practice is to produce hybrid seednuts of coconut or oil 
palm in seed gardens where the local or introduced female parent is established 
and pollen is either produced nearby or obtained from another country. All germ 
plasm used for this purpose must be assayed by means of the highly sensitive 
diagnostic tests now available, and elite certified planting material must be 
identified.

It is especially important that embryo 
cultures of coconut palm and tissue cul-
tures of oil palm be derived only from 
viroid-tested material. The consequences 
of failure to ensure pathogen freedom 
with such multiplication procedures 
would result in very high levels of infec-
tion of clonally propagated material and 
the introduction of clonally propagated material and 
the introduction of viroid to new areas 
and countries.

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Literature Cited

1. Barker, J. M., McInnes, J. L., Murphy, 
P. J., and Symons, R. H. 1985. Dot blot


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