Isolation of Cowpea Mild Mottle Virus from Diseased Soybeans in the Ivory Coast

J. C. THOUVENEL, A. MONSARRAT, and C. FAUQUET, Laboratoire de Virologie, Centre d'Adiopodoumé, Office de la Recherche Scientifique et Technique Outre-Mer, B.P. V 51, Abidjan, Côte d'Ivoire

ABSTRACT

A virus isolated from soybeans showing severe mosaic was identified as an isolate of cowpea mild mottle virus.

Soybean (Glycine max L. Merr.) is a recent introduction in the Ivory Coast. Great development of this crop is planned and numerous cultivars have been tested since 1972. During summer 1978, plants showing mosaic and crinkling of leaves were observed in experimental plots of the Ecole Nationale Supérieure Agronomique near Abidjan (southern Ivory Coast); similar symptoms were observed in fields near Bouake (central Ivory Coast). Characterization of an associated virus that appears to be an isolate of cowpea mild mottle virus (2,3) is reported here.

MATERIALS AND METHODS
Inoculum for mechanical transmission tests was sap from leaves of diseased soybean that were ground in 0.1 M phosphate buffer, pH 7.1, containing 0.02 M cystein hydrochloride. Test plants were grown in insect-proof glasshouses under local climatic conditions (mean temperature of 28°C, average relative humidity 90%).

The aphids Aphis craccivora Koch and A. spiraeola Patch were used in transmission tests. Insects were starved for 2-4 hr, then allowed an acquisition access of about 15 min on diseased leaves of soybean and an inoculation access period of 1-3 days on healthy seedlings of soybean (10 insects per plant).

Thermal inactivation point, dilution end point, and aging in vitro were determined according to the methods of Bos et al. (1) using crude sap from diseased soybeans and soybean seedlings as test plants.

For purification of the virus, leaves of infected soybean were ground in 0.2 M potassium phosphate buffer, pH 7.8, containing 1% thioglycolic acid (1 g/2 ml); chloroform was added during the grinding (1/2 vol). The moisture was centrifuged for 10 min at 10,000 g and the supernatant was centrifuged at 78,000 g for 120 min. Pellets were resuspended in 0.05 M borate buffer, pH 8, containing 0.2% Triton X 100 (1/10 of the initial volume). After removal of the insoluble material by centrifugation for 10 min at 10,000 g, the virus sedimented through a 2-cm-deep layer of 20% sucrose in the SW 27 rotor (95,000 g for 150 min). The pellets were resuspended in the same borate buffer (2% of the initial volume) and further purified by sucrose density gradient centrifugation (10-40% in 0.1 M phosphate buffer, pH 7.5). An opaque virus band was collected, diluted with water, and centrifuged 150 min at 78,000 g before being resuspended in borate buffer, pH 8.

For electron microscopy, virus preparations were stained with 2% aqueous uranyl acetate and observed with a Siemens Elmiskop 102 at the Groupe d'Etude et de Recherches en Microscopie Electronique at Adiopodoumé.

The molecular weight of coat protein was determined by electrophoresis in 7.5, 10, and 12.5% polyacrylamide-sodium dodecyl sulfate gels using the procedure of Weber and Osborne (6). Phosphoglycerate kinase (mol wt 47,000), alcohol dehydrogenase (35,000), carboxy anhydrase (31,000), trypsin inhibitor from soybean (21,000), and lysozyme (14,300) were used as standards.

Antiserum was prepared in rabbits by six weekly intravenous injections of about 1 mg of purified virus. The microprecipitin reaction under paraffin oil in petri dishes was used for serologic tests (5).

RESULTS
Symptomatology and host range. Naturally infected soybeans were stunted and showed a light green mosaic. On inoculated soybean seedlings, veinclaring evolving into yellow mosaic with occasional crinkling developed on youngest leaves 9-12 days after inoculation (Fig. 1); inoculated plants were reduced in size compared with healthy plants.

The virus was readily transmitted by mechanical inoculation as follows: Amaranthaceae: Gomphrena globosa L. (mottle); Chenopodiaceae: Beta vulgaris L., Chenopodium amaranticolor Coste & Reyn., C. quinoa Wild., and C. foetidum Schrad. (chlorotic local lesions); Leguminosae: Arachis hypogaea L. (mild crinkling), Cajanus cajan Lillsp. (chlorosis), Canavalia ensiformis DC. (mosaic and crinkling), Crotalaria usaramoensis Bak. (chlorosis), Glycine max (mosaic), Phaseolus lathyroides L. (mosaic), P. lunatus L. (chlorosis), P. mango L. (chlorosis), P. vulgaris L. (chlorotic spots), Pisum sativum L. (chlorosis), Pseudocordiceps tetragonolobus DC. (mottle), Pigna cylindrica Skeels (mosaic), V. sinensis Sav. (mosaic), V. unguiculata Wild. (mosaic), and Voandzeia subterranea Thouars (light mosaic); Pedaliaceae: Sesamum indicum L. (chlorotic spots); Scrophulariaceae: Penstemon hirsutus Wild. (chlorosis); Solanaceae: Browallia demissa L. (chlorosis), B. spectabilis Hook. (chlorosis), Nicotiana clevelandii Andr. (chlorosis), N. megasplant Ag. (mottle), and Solanum carolinense L. (chlorosis).

Transmission. The virus was not transmitted by A. craccivora (0/100) nor by A. spiraeola (0/100) from soybean to soybean.

The virus was seedborn in G. max cvs. Jupiter (1/200) and Santa Rosa (5/60).

In vitro properties. The dilution end point of the virus was between 10^-4 and 10^-5; the thermal inactivation point was between 65 and 70°C; and infectivity was retained for 2 days at 25°C, 28 days at 4°C, and more than 2 mo at -20°C.

Purification and virus properties. Based on the extinction coefficient E260

![Fig. 1. Symptoms of mosaic on soybean leaf.](image-url)