Inheritance of Resistance to Passionfruit Woodiness Virus in Common Bean (*Phaseolus vulgaris* L.)

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**Abstract.** Passionfruit woodiness virus (PWV) can infect bean (*Phaseolus vulgaris* L.), causing a light and dark green foliar mosaic, veinbanding, downward curling, and plant stunting. The intensity of these symptoms can vary with the strain of the virus and cultivar, but they resemble those caused by bean common mosaic virus. In genetic populations derived from crosses and backcrosses involving cultivars that are resistant ('Black Turtle 1', 'Clipper', and 'RedKote') or susceptible ('Black Turtle 2', 'California Light Red Kidney', and 'Pioneer'), a single dominant gene conferred resistance to an Australian strain PWV-K. To this gene, the symbol *Pw* (Passionfruit woodiness virus) is tentatively assigned. In plants derived from rooted cuttings of backcross populations, the same factor also conditioned resistance to three other Australian strains, PWV-Mild, PWV-51, and PWV-Tip Blight.

Passionfruit woodiness virus (PWV) is considered to be one of the major Potyviridae affecting passionfruit (*Passiflora edulis* Sims.) and related species. It was originally described by McKnight (1953) and has since been found in other tropical and subtropical regions of the world (Provvidenti and Niblett, 1994; Taylor and Greber, 1973). The host range of this virus also involves members of the Leguminosae, including the common bean (*Phaseolus vulgaris* L.) Taylor and Greber, 1973). Very little is known about the presence of PWV in common bean crops grown in warm areas of the world. However, a few years ago, it was found to occur in the Federal District and Cristalina, Goias State, Brazil, where it caused severe symptoms on local bean cultivars (Inoue et al., 1995). Experimentally, they found that a few cultivars were resistant. In our work on biological properties of PWV from Australia (PWV-K), Puerto Rico (PWV-PR), and Thailand (PWV-T), American common bean cultivars were useful in differentiating these strains. The Australian strain systematically infected more bean lines (16/32) than did the other two strains (Provvidenti, 1992), and was one of the few strains capable of infecting pea (*Pisum sativum* L.). However, among the pea lines tested, 23 of 30 were found to be highly resistant. Inheritance studies revealed that in peas, the resistance is conferred by a single recessive gene (Provvidenti and Niblett, 1994) Conversely, our preliminary tests indicated that the resistance to PWV-K in common beans was dominant (Provvidenti, 1993). The data presented herein confirm the dominance of resistance in common beans to PWV-K and three other Australian strains.

**Materials and Methods**

A few cultivars reported to be resistant (R) or susceptible (S) to PWV-K (Provvidenti, 1992), were selected for inheritance studies. The following crosses were made: ‘Black Turtle 1’ (BT-1) (R) x ‘Black Turtle 2’ (BT-2) (S); ‘RedKote’ (R) x ‘California Light Red Kidney’ (CLRK) (S); and ‘Clipper’ (R) x ‘Pioneer’ (S). Both BT-1 and BT-2 have black seedcoats and belong to the Andean Group, whereas the Middle American Group includes CLRK and ‘RedKote’ (red seedcoats) and ‘Clipper’ and ‘Pioneer’ (white seedcoats). A culture of the PWV-K strain was available from previous studies (Provvidenti, 1992, 1993; Provvidenti and Niblett, 1994), whereas the Australian PWV-Mild (M), PWV-51, and PWV-Tip Blight (TB) (Shukla et al., 1988), were kindly supplied by Dr. R.S. Greber (Queensland, Australia). Cultures of these strains were maintained in CLRK plants, and inocula were prepared by macerating young infected leaves with a phosphate buffer (0.05 x K,HPO4, at pH 8.5. For inheritance studies involving PWV-K, plants of the parental lines and populations of F1, F2, and backcross F1 to both parents were mechanically inoculated on the expanded primary leaves and reinoculated a week later on the first trifoliate. This dual inoculation minimized the number of escapes. To confirm infectivity, bean plants were assayed by enzyme-linked immunosorbent assay (ELISA) using an antiserum to PWV previously obtained from Dr. Greber (Provvidenti and Niblett, 1994). When ELISA was inconclusive, recovery tests were made using CLRK as a systemic host. To determine whether the same resistance factor(s) conferred resistance to different Australian strains of PWV, four cuttings were made from each plant of three backcross populations: (BT-1 x BT-2) x BT-2, (‘Clipper’ x ‘Pioneer’) x ‘Pioneer’, and (‘RedKote’ x CLRK) x CLRK. After rooting, the resulting plants were inoculated individually with one of the four strains (TWV-K, -M, -51, or -TB). All of the plants were grown in sterilized clay pots (500-mL volume) containing the Cornell artificial mix (sphagnum peatmoss, Whittomevermiculite, Baker’s dolomitic limestone, and CaNO3). Every week, plants were fertilized with the water-soluble ‘Start-N-Gro’ (14N-28P-14K) (Agway, Syracuse, N.Y.), and treated with ENSTAR II [S-Kinoprene [2-propynyl (2E,4E)3-7S,11-trimethyl-2,4,5-dodecadienoate]] (Sandoz Agro, Des Plains, Ill.) This insecticide was used for the control of whiteflies, aphids, armored scales and mealybugs, and kept plants completely free of viral vectors. During the winter months, natural light was supplemented with fluorescent lights for 16 h daily. Plants were kept in a greenhouse, where temperature ranged from 25 to 30°C, with a relative humidity (RH) of 40% to 60%.

**Results and Discussion**

Plants of BT-1 and ‘RedKote’ remained free of local and systemic symptoms after two inoculations with PWV-K, but ‘Clipper’ developed small necrotic lesions confined to the inoculated primary leaves. ELISA confirmed that the virus had infected inoculated leaves of plants of these three cultivars, but had failed to move systemically (systemic resistance). The virus infected locally and systemically plants of BT-2, CLRK, and ‘Pioneer’. Initially, necrotic local lesions were visible only on primary leaves of ‘Pioneer’, but eventually, veinal browning and interveinal chlorosis also developed on those of BT-2 and CLRK. Systemic symptoms initially consisted of a light and dark green mottle, followed by a prominent mosaic, green veinbanding, downward leaf curling, and plant stunting. The F1 plants of the crosses BT-1 x BT-2, ‘Clipper’ x ‘Pioneer’, and ‘RedKote’ x CLRK responded to inoculation with local infection, but, in the resistant parents, the virus failed to move systemically. The F1 plants of these crosses segregated in a ratio of 3 systemically resistant: 1 susceptible. Systemically infected plants exhibited the same type of symptoms displayed by those of the susceptible parents. Backcross populations involving the susceptible parents (BT-1 x BT-2) x BT-2, (‘Clipper’ x ‘Pioneer’) x ‘Pioneer’, and (‘RedKote’ x CLRK) x CLRK segregated in a ratio of 1 systemically resistant: 1 susceptible. The backcross populations involving resistant parents (BT-1 x BT-2) x BT-1, (‘Clipper’ x ‘Pioneer’) x ‘Clipper’, and (‘RedKote’ x CLRK) x ‘RedKote’, were all systemically resistant. Thus, based upon the data presented in Table 1, the resistance to PWV-K in BT-1, ‘Clipper’, and ‘RedKote’ is conferred by a single dominant factor. There-
In common beans, the type of symptoms incited by the four Australian strains of PWV were essentially similar, but they ranged from mild (PWV-M), to moderate (PWV-51), to severe (PWV-TB). The prominence of symptoms on susceptible genotypes incited by PWV-K helped in analyzing genetic populations during inheritance studies. Gough and Shukla (1992) reported that the coat protein sequence of PWV-K differed in the N-terminus from those of other strains, but serologically they were essentially similar those caused by bean common mosaic virus (BCMV), also of common occurrence in warm regions, very few efforts have been made to determine the presence, distribution, and economical importance of PWV. BCMV and PWV are not serologically related and the latter has a wider host range than the former (Taylor and Breber, 1973). Resistance to BCMV in common beans can be strain-specific and conditioned by: 1) recessive genes (bc-1, bc-1', bc-2, bc-2', and bc-3); 2) a dominant gene (I); and 3) a combination of a recessive and the dominant factor (Drijfhout et al., 1978). In ‘Clipper’, the resistance to BCMV is recessive, and in BT-1 and ‘Redkote’, it is dominant. Our work has established that in these three cultivars, the resistance to PWV is conferred by a non-strain-specific dominant factor (Pwv). However, a number of common bean cultivars possess resistance to BCMV and PWV (e.g., ‘Alliance’, BT-1, ‘Benton’, ‘Bonanza Wax’, ‘Bounty’, ‘Bush Blue Lake 47’, ‘King Horn Wax’, ‘Labrador’, ‘Midnight’, ‘RedKloud’, ‘RedKote’, ‘RoyalRed’, ‘Vitagreen’, and others) (Provvidenti, 1992).

### Table 1. Inheritance of resistance in common bean (Phaseolus vulgaris L.) to an Australian strain of passionfruit woodiness virus (PWV-K).

| Parents and cross progeny | No. of plants | Expected ratio | Goodness of fit |
|---------------------------|---------------|----------------|-----------------|
|                           | Resistant     | Susceptible    | (R : S) | $\chi^2$ | (P) |
| Black Turtle 1 (BT-1)     | 54            | 0              |         |         |     |
| Black Turtle 2 (BT-2)     | 0             | 51             |         |         |     |
| Clipper                   | 43            | 0              |         |         |     |
| ‘Pioneer’                 | 0             | 37             |         |         |     |
| Redkote                   | 35            | 0              |         |         |     |
| CA Light Red Kidney (CLRK)| 0             | 48             |         |         |     |
| **BT-1 x BT-2**           |               |                |         |         |     |
| F₁                         | 35            | 0              |         |         |     |
| F₂                         | 84            | 24             | 3:1    | 0.444  | 0.51|
| BC (F₁ x BT-2)            | 42            | 0              |         |         |     |
| BC (F₂ x BT-2)            | 35            | 31             | 1:1    | 0.242  | 0.64|
| **Clipper x Pioneer**     |               |                |         |         |     |
| F₁                         | 47            | 0              |         |         |     |
| F₂                         | 97            | 29             | 3:1    | 0.264  | 0.62|
| BC (F₁ x Clipper)         | 77            | 0              |         |         |     |
| BC (F₂ x Pioneer)         | 43            | 38             | 1:1    | 0.604  | 0.48|
| **Redkote x CLRK**        |               |                |         |         |     |
| F₁                         | 28            | 0              |         |         |     |
| F₂                         | 76            | 22             | 3:1    | 0.264  | 0.57|
| BC (F₁ x Redkote)         | 39            | 0              |         |         |     |
| BC (F₂ x CLRK)            | 27            | 24             | 1:1    | 0.176  | 0.68|

*Resistant.

'Susceptible.