Stigmasterol-3-O-glucoside, an allelopathic molecule responsible for pest resistance of Thenkaali (AAB), a Musa cultivar against Odoiporus longicollis [Olivier]

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INDIA is one of the major biodiversity centres of Musa cultivars and stands first in the world in the production of bananas and plantains¹. Musa cultivars are sterile plants which are unable to set viable seeds and are propagated vegetatively by suckering. These plants are originated through natural hybridization of two plants under a single genus, Musa acuminata (A genome) and M. balbisiana (B genome)². The cultivars are mostly triploid 2n = 3x = 33 and those with short, robust pseudostem and broad leaves come under the genome constitution of AAA, while plantains AAB or AB and starchy cooking banana ABB, all are equally diverse with major influence of ecological factors on their distribution and survival². A few diploid cultivars with genome constitution AA are also cultivated, but yield comparatively small fruit bunches and have chromosome number 2n = 2x = 22, which is also common in the agro-ecosystem³. The cultivars of Musa have diversified into complex germplasm pool, mostly through somatic mutation⁴ and somaclonal variation⁵. The Cavendish AAA group of Musa cultivars is increasingly susceptible to a range of fungal and viral diseases and also to insect pests, which necessitate chemical control measures.

The diversity of Musa cultivars in India is high and the commercially viable cultivars (CVCs) usually have desirable characters such as large fruit bunch, palatable ripe fruits, short duration to set flowers, etc. However, these CVCs are highly susceptible to pseudostem borer, Odoiporus longicollis [Olivier]⁷,⁸. They possess very low quantities of total phenols (TP), total flavonoids (TF) and very low activities of phenylalanine ammonia lyase (PAL), polyphenol oxidase (PPO) and peroxidase (PO)⁹,¹⁰. O. longicollis is the most destructive pest in the commercial cultivation of Musa¹¹. The Musa cultivars have been categorized into highly susceptible (HS), susceptible (S), nonpreferred (NP) and resistant (R) based on the preference of O. longicollis⁷,⁸. Rearing of larvae in live pseudostem of HS, S and NP cultivars has resulted in completion of life cycle, but in R cultivars has resulted in mortality of larvae in one week⁷,⁸. If HS or S cultivars were available in the agro-ecosystem, mother weevil never visited NP cultivars. They showed sporadic attack by the pest when away from the vicinity of HS or S cultivars⁷,⁸. Rearing of O. longicollis in live R cultivars has resulted in mortality of larvae in seven days. Analysis of 85 cultivars of Musa in Kerala, India showed that 28 of them are under R category and all cultivars possessed...
very high content of TP, TF and elevated activity of PAL, PPO and PO\textsuperscript{9,10}.

Thenkaali (Figure 1\textit{a}), a \textit{Musa} cultivar of genome constitution AAB is among the R category cultivars, identified in Pathanamthitta district of Kerala. It is a commercially non-viable cultivar identified through detailed field study supplemented with GIS Arc software\textsuperscript{7}. Thenkaali showed strong allelopathic interactions against \textit{O. longicollis}, when reared in the live pseudostem. Bioassay-guided extraction of dry pseudostem powder of Thenkaali using organic solvents and subsequent fractionation of the active larvicidal extract resulted in the isolation and characterization of the allelopathic molecule, which forms the subject matter of this study.

Materials and methods

Experimental organism

Larvae of \textit{O. longicollis} dissected out from the infested pseudostem of susceptible cultivars were used. Larvae up to second instar are small and highly sensitive to handle. Fourth-instar larvae are preferably large and voracious feeders, and hence these were used for the study.

Rearing of larvae in the live pseudostem

Four-month-old \textit{Musa} cultivars with trunk circumference of 25–30 cm were used for the study. The crown of the plant was cut in such a way that the live stump of 1 m height remained in viable state. Larvae of \textit{O. longicollis} (six numbers) were carefully placed at the free end so that they could bore into the pseudostem. The free end was covered with a piece of mosquito net and kept undisturbed for eight days. On the eighth day, the pseudostem was cut 15 cm below the first cut, and the larvae were carefully taken out and compared with those in susceptible cultivars. If all the larvae were dead on the eighth day the cultivar was considered as resistant to \textit{O. longicollis} infestation\textsuperscript{7,8}. Palayankodan (Figure 1\textit{b}), a CVC and HS cultivar with genome constitution AAB was used as control\textsuperscript{8}.

Host plant enzymes

Tender leaf at unfurl cigar state of 20–30 cm length was cut from the free terminal (5 cm) and kept in ice-cold conditions till further processing. The content of TP\textsuperscript{12}, TF\textsuperscript{13}, and activities of enzymes such as PAL\textsuperscript{14}, PO\textsuperscript{15} and PPO\textsuperscript{16} were assayed according to standard protocols.

Phytochemical isolation of compounds

Pseudostem of \textit{Musa} cultivars which are highly susceptible or resistant to infestation were cut vertically into small chips and dried under shade (at ambient temperature 28°–33°C) for three weeks, after which they were ground to a fine powder. Thereafter, 400 g sample was kept overnight in organic solvents and first extracted by petroleum ether, followed by acetone and finally with methanol. The extracts were filtered and concentrated in rotary vacuum evaporator and larvicidal activity was observed as described in the following section. As the larvicidal activity was found only in the acetone extract, it was subjected to column chromatography using a suitable solvent system. Elution was carried out by gradient polarity system starting from 100% petroleum ether to 100% methanol. Each fraction was monitored by TLC and they were combined, based on their TLC profiles, which resulted in nine fractions. Larvicidal activity of each fraction was noted. As the eighth fraction showed larvicidal activity, it was again subjected to column chromatography to isolate the specific compound in pure form.

Spectral analysis of active molecule

NMR (\textsuperscript{1}H, \textsuperscript{13}C, HMBC and HSQC) and LC-Q-T-MS analysis of the active compound were done at IIRBS and School of Environmental Science, Mahatma Gandhi University, Kottayam, Kerala.

Testing of larvicidal activity

Freshly cut pseudostem of 100 g pieces of circular rods from Palayankodan cultivar was used for the study. Active compound or extract was dissolved in water containing 0.5% Tween. The dissolved extract (2 ml) from susceptible or resistant pseudostem or active molecule in known concentration was injected into the 100 g piece of pseudostem evenly at ten sites using insulin syringe. Each larva was allowed to bore into the pseudostem. Healthy
fourth-instar larvae were used for the study. Percentage mortality was calculated as follows:

\[
\text{Percentage mortality} = \frac{\left( \frac{\text{Percentage of test mortality}}{\text{percentage of control mortality}} \right) \times 100}{100 - \text{Percentage of control mortality}}.
\]

Every day each piece containing a single larva inside was kept close to the ear of the observer six times (every 30 min continuously for 3 h) to hear the feeding sound, which is an indication of the health of the larva inside. Wriggling movements of the apodous larvae and creamy-white colour of the body are other conspicuous characters which indicate health of the larvae. They became flaccid and gradually acquire a bluish colouration at death. Each piece with a larva was placed in a plastic container covered with a cheese cloth and kept in a dark, cool place. Fresh piece of pseudostem, administered with the test sample was provided once in two days. On the eighth day, the pseudostem pieces were carefully dissected to observe the larvae. Minimum quantity of active compound required for mortality within seven days was studied. As 100% mortality of the larvae in live pseudostem of Thenkaali cultivar occurred between the seventh and eighth day, seven days was taken as the observation limit to assess toxicity.

**Haematological studies**

The larvae on the fourth day of observation (before death) were used for study. They were kept on a glass plate with ice cubes placed above and a small cut was given on the ventral side with great care, not to puncture the gut, to collect haemolymph. Total haemocyte count was measured using by haemocytometer. The number of haemocytes per cubic millimetre was calculated using the formula suggested by Jones. A thin film of smear was prepared using haemolymph to study the differential hemocyte count. The smear was air-dried and stained using Giemsa stain for 20 min. The slides were then washed in running water to remove excess stain. The slides were then air-dried and the number of different types of haemocytes of treated and control samples was counted. The percentage of different types of haemocytes in each of the samples was also calculated.

**Assay of larval enzymes**

Cell-free haemolymph of fourth-instar larvae was used for the study. Quantitative estimation of protein, total free amino acids, transaminases (aspartate amino transferase [E.C. 2.6.1.1] and alanine amino transferase [E.C. 2.6.1.2])

leucine aminopeptidase [E.C. 3.4.11.1] and cathepsin D [E.C. 3.4.23.5] was done according to standard protocols. Enzyme immunoassay of 20-hydroxyecdysone was done using an enzyme kit (Bertin Pharma, France).

Uric acid content was estimated by uricase-peroxidase end-product assay kit (Span Diagnostics).

Electrophoresis of cell-free haemolymph of larvae was carried out using the method devised by Laemmeli; 12% resolving gel and 4% stacking gel were used for SDS-PAGE. The electropherogram was subjected to Gel-Doc analysis.

**Statistical analysis**

The results obtained are represented as mean ± standard deviation. Statistical comparison of results was performed by one-way analysis of variance [ANOVA] using SPSS 21 software. The results are considered statistically significant if \( P \leq 0.05 \) (ref. 28).

**Results**

An indigenous *Musa* cultivar of Kerala, locally called as Thenkaali (AAB), did not get infested by *O. longicollis* under field conditions. Rearing of *O. longicollis* larvae in the live pseudostem of Thenkaali cultivar occurred between the seventh and eighth day, seven days was taken as the observation limit to assess toxicity.

See Figure 2. Total phenols and total flavonoids content of resistant and susceptible *Musa* cultivars. All values are mean ± SD, \( n = 6 \). Value of each parameter is significantly different in Duncan’s multiple range test at 0.05 level.

[Figure 2](#)
NMR (H¹ and C¹³) and LC-Q-T-MS revealed that the compound was a glycoside and was named stigmasterol-3-O-glucoside (Figure 4). ESI-MS: m/z 575.2375 [M + H]+. ¹H NMR (CDCl₃–CD₃OD, 400 MHz): δ (ppm): 5.42 (1H, br.s, H-6), 5.09–5.21 (2H, m, H-22 and H-23), 4.47 (1H, br.s, H-1′), 3.28–3.93 (6H, m, H-2′–H-6′). ¹³C-NMR (CDCl₃–CD₃OD, 100 MHz): δ (ppm) 141.2, 139.2, 130.1, 122.7, 102.0, 79.7, 77.4, 77.0, 74.5, 71.1, 62.4, 57.7, 56.8, 51.1, 50.7, 43.1, 39.4, 38.1, 37.5, 36.9, 32.7, 32.6, 30.4, 30.1, 25.0, 23.4, 21.8, 21.5, 21.3, 19.8, 19.4, 12.7, 12.4.

The ¹³C NMR spectrum showed a total of 33 peaks which included four olefinic carbon signals (δ 141.2, 139.2, 130.1 and 122.7), similar to those of stigmasterol. However, there was a chemical shift value of 77.4 ppm observed instead of 71 ppm at C-3, indicating a different attachment at C-3 instead of the hydroxyl group. The attachment at C-3 was found as glucose moiety from the peaks between δ 3.28–3.93 and 4.47 in ¹H NMR, and carbon signals at δ 102.0, 62.4, 71.1, 74.5, 77.0 and 79.7 in the ¹³C NMR spectrum. The mass data (m/z 575.2375 [M + H]+) further confirmed the presence of sugar attachment to the stigmasterol skeleton. The ¹³C NMR values of the compound were once again matched with the literature values and the compound was confirmed as stigmasterol-3-O-glucoside.

The larvae in the pseudostem (100 g) which was administered with 1 mg (10 ppm) of active compound stopped feeding within 24 h; thereafter, no feeding sounds were heard. Table 1 shows the percentage mortality induced by stigmasterol-3-O-glucoside at different concentrations, viz. from 1 to 10 ppm. At higher concentration of 5 and 10 ppm, the larvae became immobile from the fourth day onwards, and died between the sixth and eighth day. Even though the larvae in pseudostem at 1 ppm concentration of stigmasterol-3-O-glucoside did not die even after one week, they died after 15 days without undergoing pupation. No feeding sounds were heard after seven days, which is an indication that the larvae stopped feeding on the seventh day at 1 ppm. They were weak and flaccid, and unable to feed or enter into the pseudostem.

Toxicity of stigmasterol-3-O-glucoside in O. longicollis larvae resulted in dose-dependent haemocytopenia in the haemolymph (Figure 5). The differential count also

### Table 1. Percentage of mortality exhibited by stigmasterol-3-O-glucoside

| Stigmasterol-3-O-glucoside (ppm) | Four days | Seven days | 15 days |
|----------------------------------|-----------|------------|--------|
| Control                          | –         | –          | 100    |
| 1                                | –         | –          | 100    |
| 2                                | –         | 20         | 100    |
| 5                                | 30        | 50         | 100    |
| 10                               | 70        | 100        | 100    |

*Fifteen larvae were used for each concentration.*

Figure 3. Activity of host-plant enzymes of resistant and susceptible Musa cultivars. All values are mean ± SD, n = 6. Value of each parameter is significantly different in Duncan’s multiple range test at 0.05 level.

Figure 4. Structure of stigmasterol-3-O-glucoside.

Figure 5. Effect of stigmasterol-3-O-glucoside on total haemocyte count. All values are mean ± SD, n = 6. Value of each parameter is significantly different in Duncan’s multiple range test at 0.05 level.
showed significant change during toxicity, indicating imbalance on the proportionate distribution of haemocytes. Among the different populations of haemocytes, the number of granulocytes showed a sharp increase but the number of plasmatocytes showed a sharp decrease (Figure 6).

Stigmasterol-3-O-glucoside intoxication at doses of 5 and 10 ppm resulted in sharp elevation of haemolymph protein in *O. longicollis* larvae (Figure 7). Along with hyperproteinemia, sharp qualitative changes in protein profile, indicating appearance of new protein bands and disappearance of existing bands were observed (Figure 8). Sharp elevation of the content of haemolymph protein was clearly visible in the electropherogram. The changes in protein profile specifically at bands of molecular weight 64.53, 49.73, 36, 36.52, 30.48 and 21.23 kDa are indicated as arrows in the electropherogram. Significant decrease in the content of free amino acids was also observed at 5 and 10 ppm concentration of the larvicide (Figure 7). Significant elevation of haemolymph uric acid (Figure 9) was observed in intoxicated larvae. The activity of cathepsin D showed a sharp elevation (Figure 10), but another protein-degrading enzyme, viz. leucine amino peptidase (LAP) showed sharp inhibition of activity (Figure 11). Significant elevation in the activities of aspartate amino transferase and alanine amino transferase were also observed in the larvae intoxicated with stigmasterol-3-O-glucoside (Figure 12).
β-Ecdysone is a steroid hormone which plays a major role in insect metamorphosis. Larvae reared in the pseudostem containing different concentrations of stigmastrol-3-\textit{O}-glucoside caused a dose-dependent elevation in the activity of the enzyme 20-hydroxy ecdysone monooxygenase responsible for converting \textit{α}-ecdysone to \textit{β}-ecdysone. Figure 13 shows the effect of stigmastrol-3-\textit{O}-glucoside on the activity of 20-hydroxy ecdysone monooxygenase.

**Discussion**

Host-plant resistance is the most viable and sustainable strategy for the management of agriculturally important insect pests. Field study and laboratory experiments on 85 \textit{Musa} cultivars from different agro-ecosystems in Kerala, showed that the cultivars with high content of TP, TF and high activity of PAL, PPO and PO can cause allelopathy in \textit{O. longicollis} larvae\textsuperscript{8-10}. Isolation of a steroid glycoside which is highly toxic to the larvae which are completing their life cycle within the \textit{Musa} cultivars is new to science, but stigmasterol has been isolated from many plants and the compound isolated from the bark of \textit{Butea monosperma} with hypoglycaemic activity in experimental animals\textsuperscript{31}. Crushed juice of fresh pseudostem of \textit{Musa} cultivar Thenkaali is used as a remedy for hyperacidity and urolithiasis among villagers of Chittar in Kerala\textsuperscript{8}. Antiulcerogenic activity of two acyl steryl glycosides, sitoindoside I and sitoindoside II, has been reported from peeled fruits of another \textit{Musa} cultivar from India\textsuperscript{32}.

Allelopathy induced by the host plant (Thenkaali) through stigmasterol-3-\textit{O}-glucoside on the larvae within...
the pseudostem was not directed against any particular metabolism, but it affected transamination, protein degradation, protein profile of haemolymph, endocrine disruption and haemocytes. Attack on plasmatocytes and granulocytes will result in a defective immune system. Haemocytes of the intoxicated larvae showed a sharp decrease in number. Widespread cytopathological changes were induced by the live cultivar Thenkali, on larvae of O. longicollis during laboratory rearing. This type of haemocytopenia was also observed in Dyscercus sirgentatus and Papilo demoleus on intoxication by azadiractin. In the present study, stigmasterol-3-O-glucoside caused sharp increase in the number of granulocytes and decrease in the number of plasmatocytes. Plasmatocytes and granulocytes are major cell types responsible for immune defence in insects. Plasmatocytes are more involved in engulfing non-self or invading microorganisms, while granulocytes are involved in removing cell debris. Cytopathological studies revealed that most of the cells affected during toxicity were plasmatocytes, and this may be the reason for the low number of plasmatocytes observed during differential count. The debris of the lysed plasmatocytes in the haemolymph may be the reason for the low number of plasmatocytes. Plasmatocytes affected during toxicity were plasmotocytes, and immunochemical staining of feeding was one of the reasons for the death of larvae in the pseudostem containing different concentrations of the active compound. It is well established that elevation of 20-hydroxyecdysone in the haemolymph caused cessation of feeding activity in almost all final instar larvae and pupae of holometabolous insects.

Conclusion

Presence of a highly toxic larvicidal molecule in an indigenous Musa cultivar against the chief pest of Musa cultivars paves the way for its use in pest management. Identification of the genetic mechanism involved in the synthesis of this larvicide and its subsequent modulation in susceptible cultivars need further studies. Development of host-plant resistance is the sustainable strategy for pest management.
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