Expression of *Pseudomonas* phosphotriesterase activity in the fall armyworm confers resistance to insecticides

D. P. Dumas, J. R. Wild and F. M. Raushel

Departments of Chemistry and Biochemistry and Biophysics, Texas A & M University, College Station (Texas 77843, USA)

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Summary. The gene encoding for the phosphotriesterase (opd) from *Pseudomonas diminuta* has been subcloned into a baculovirus expression system. Functional enzyme is produced when the recombinant baculovirus is used to infect either cultured *Spodoptera frugiperda* sF9 cells or the larval stage of the fall armyworm. The LD$_{50}$ for paraaxon toxicity was found to increase 280-fold in the larvae after infection with the recombinant baculovirus and expression of the functional phosphotriesterase.

Key words. Phosphotriesterase; fall armyworm; pesticide resistance; paraaxon.

The resistance of bacteria to antibiotics is due in many cases to the existence of a gene which encodes for an enzyme that catalyzes the chemical modification of the particular antibiotic to a nontoxic molecule. The hydrolytic action of the various β-lactamases with the penicillin-type antibiotics is a classic and well-characterized example. A *Pseudomonas diminuta* gene (opd, organophosphorus degrading) encoding an enzyme that catalyzes the rapid hydrolysis of many of the phosphotriesters currently used as commercial insecticides has been successfully cloned. These insecticides are highly toxic due to their ability to rapidly inactivate the enzyme acetylcholinesterase from a variety of sources. The problem to be addressed in this report is whether the expression of this gene in insects could confer reduced toxicity to these phosphotriester-type insecticides due to the chemical detoxification of the pesticide within the insect itself. The baculovirus expression system developed by Summers et al. appeared to be a well-suited model for this hypothesis since the cloned opd gene could be inserted into the baculovirus and functionally expressed either in cultured *Spodoptera frugiperda* sF9 cells or, more appropriately, into the larval caterpillar stage of the fall armyworm.

Materials and methods

All kinetic measurements were performed at 25 °C using a Gilford 260 spectrophotometer. Enzymatic activity was measured by monitoring the absorbance of p-nitrophenol at 400 nm generated upon the hydrolysis of diethyl-p-nitrophenylphosphate (paraaxon) at pH 9.0 in 150 mM 2-cyclohexylamino]-ethanesulfonic acid (CHES). A unit of activity is defined as the amount of enzyme needed to hydrolyze 1 μmol of paraaxon per min.

A recombinant baculovirus containing the opd gene was constructed as in Harper et al. using standard recombinant DNA techniques as described by Summers and Smith. The construction of this baculovirus utilized the insertion of the opd gene into the baculovirus transfer vector pVL941 to give the recombinant transfer vector pLH1170opd. pVL941 is a derivative of pAC311 in which the ATG start has been changed to an ATT triplet by site directed mutagenesis. The translation of a foreign gene is thus expressed from the polyhedron protein promoter and initiated at the first codon in the open reading frame of that gene. Recombinant virus containing the opd gene was obtained by the co-transfection of *Spodoptera frugiperda* sF9 cells with pH1170opd and wild-type *Autographa californica* nuclear polyhedrosis virus (AcNPV) viral DNA followed by the isolation of those cells which were able to hydrolyze paraaxon to diethylphosphate and p-nitrophenol. Among the viral lines capable of expressing the phosphotriesterase, isolate A5B-opd was shown to produce the greatest amount of phosphotriesterase activity.

The fall armyworms used in this study were cultured from eggs provided by Dr James Fuwa of Louisiana University. These eggs were from a strain originally isolated in Hammond, Louisiana. The larvae of the fall armyworms were raised in 1-oz plastic creamer cups with cardboard lids, and fed a commercially prepared diet (BioServ, Inc. Frenchtown, NJ). The temperature was maintained at 27 °C with a 16-h photoperiod. After pupation, the insects were transferred to oviposition cages made from 3-gallon ice cream cylinders covered with cheese cloth. Upon emergence, the moths fed on a solution of beer and honey (12 oz beer, 30 ml honey, and 10 g ascorbic acid diluted to 1 l). Eggs were harvested daily from the cheese cloth and placed in a sealed container with food until hatching when they were separated into plastic creamer cups.

In order to assess whether the expression of a functional phosphotriesterase by a baculovirus within living insects would protect against the lethal effects of an insecticide, the recombinant baculovirus was injected directly into the larval stage of the fall armyworm after sterilization of the insect in carbon dioxide and disinfection by dipping in 2% bleach followed by dipping in 70% ethanol and finally submerged for 5–10 min in 0.9% NaCl. The caterpillars were injected late in the third instar by taking
up 5 μl of either 2 × 10⁶/ml A5B-opd or 2 × 10⁸/ml AcNPV virus into a fine capillary needle and injecting s.c. into the hemolymph through one of the prolegs. The expression of phosphotriesterase activity was assessed by the sacrifice of ten caterpillars each day for several consecutive days by forcing the insects individually through the small orifice of a disposable 10-ml syringe. The crushed caterpillars were centrifuged and the supernatant fluid was decanted and recentrifuged to further separate the body fluids from the fat and body parts that remained after the first centrifugation. After preparing the body fluids, enzymatic activity was determined on an aliquot of the solution without further dilution and the values for the ten caterpillars were averaged. The larvae were exposed to different concentrations of paraoxon by placing 1-μl droplets of paraoxon/acetone solutions on the top of the last five segments of the insect. Above 0.5 mg, paraoxon was placed directly on the back of the caterpillar. Mortality was defined as a moribund state 24 h after exposure when the caterpillar could no longer roll onto its legs when placed on its back. A total population of 222 A5B-opd infected caterpillars were used to construct the mortality curve of infected insects, while 331 caterpillars were used to determine LD₅₀ of the control group. This control group included 90 caterpillars that had been infected with the wild-type AcNPV baculovirus. All larvae were tested in the fifth instar of development, four days after infection. The median lethal dose was calculated using the computational method of Finney on the corrected mortality as determined by Abbott’s formula.

**Results and discussion**

The production of active phosphotriesterase in A5B-opd infected s9 cells increased logarithmically after a short initial lag with maximum enzymatic activity of more than 10 units/ml per 10⁶ cells being achieved 5–6 days after infection when the cells begin to lyse. This protein production mimics the expression of polyhedrin protein in the AcNPV wild-type baculovirus system, and represents an overproduction of the phosphotriesterase 40 times greater than the native expression system in *Pseudomonas diminuta*. While this is a substantial increase over the native system, the amount of phosphotriesterase produced (about 7 μg/ml per 10⁶ cells) is on the low end of the 1–600 μg/l per 10⁶ cells range reported for the baculovirus expression systems.

Shown in figure 1 is the time course for the appearance of phosphotriesterase activity in the caterpillars following injection. After a relatively short lag phase, enzymatic activity could be detected in the caterpillar when paraoxon was utilized as a substrate. For the first four days after injection the phosphotriesterase activity increased about one order of magnitude every 24 h until it reached a maximum of approximately 11 units of paraoxon hydrolyzing activity per caterpillar (based upon an average of 250 μl fluids recovered from each caterpillar). No phosphotriesterase activity (< 5 × 10⁻⁵ units/caterpillar) could be detected in uninfected caterpillars or caterpillars infected with the wild-type baculovirus. Therefore, the phosphotriesterase activity expressed within the A5B-opd infected caterpillars is due solely to the recombinant baculovirus containing the opd gene. The effect of the insecticide paraoxon was determined on both the infected and uninfected caterpillars. Shown in figure 2 is a plot of probit mortality versus the amount of paraoxon that could be applied directly to the caterpillars. The LD₅₀ for the A5B-opd infected caterpillars was calculated to be 1800 ± 800 μg while the control group had an LD₅₀ of 6.5 ± 3 μg.

This graph clearly demonstrates that those caterpillars containing a functional phosphotriesterase are resistant to all but the highest concentration of paraoxon that could be applied. The slopes of the two mortality curves, being roughly the same, demonstrate the death of caterpillars in the A5B-opd infected group occurs through a similar mode of action as for the control group. The phosphotriesterase in the caterpillars infected with the recombinant virus is therefore hydrolyzing the paraoxon.
faster than the acetylcholinesterase is inactivated. Thus, the lethal dose increased by a factor of at least 280 due to the presence of the phosphotriesterase. This pesticide resistance, however, was short-lived since all caterpillars exposed to recombinant or wild type baculovirus died after approximately six days. In the moth stage, the LD₅₀ was found to be 1.2 ± 0.3 μg of paraoxon.

The results presented above demonstrate that reduced toxicity to paraoxon and perhaps other pesticides can be induced in insects by expression of an enzyme that is known to efficiently hydrolyze these molecules to nontoxic products. In this manner, a symbiotic relationship is created between the host insect and the baculovirus. Similarly, the use of a nonpathogenic bacterium as a symbiont vector may be feasible for the incorporation of the phosphotriesterase into other insects.

* To whom correspondence should be addressed.
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