Research Article

Characterisation of adult green lacewing (Chrysoperla carnea) digestive physiology: impact of a cysteine protease inhibitor and a synthetic pyrethroid

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Abstract

BACKGROUND: In spite of concern regarding potential non-target effects of GM crops, few studies have compared GM pest control with conventional methods. The impacts of cypermethrin and oilseed rape expressing oryzacystatin-1 (OC-1) were compared in this study on the predator Chrysoperla carnea (Stephens).

RESULTS: Adults fed purified rOC-1 showed a subtle shift in digestive protease profile, with an increasing reliance on serine proteases (chymotrypsin), increase in aspartic proteases and a slight reduction in elastase activity. Although there were no effects on mortality, onset of oviposition was delayed; however, once egg production commenced, egg laying and hatching success rates were comparable with those of controls. Oryzacystatin-1 expressed in pollen showed no detrimental effects. Cypermethrin had no effect on mortality owing to high levels of non-specific esterase activity resulting in partial breakdown of the insecticide. In spite of this, there was a significant delay in onset of oviposition and a significant reduction in egg production and viability.

CONCLUSION: This study demonstrates the potential for pest management to impact on predators, but importantly it highlights the ability of the predator to detoxify/respond to treatments with different modes of action. In this case, exposure to an insecticide carried a greater fitness cost than exposure to a protease inhibitor expressed in transgenic crops.

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Keywords: lacewing; Chrysoperla carnea; cypermethrin; transgenic crops; cysteine protease inhibitors

1 INTRODUCTION

Chrysoperla carnea (Stephens) is an important predator in agriculture. The value of this predator as a biological control agent arises not only from its widespread occurrence and broad range of prey but also from the fact that each of its three larval stadia are voracious polyphagous feeders, capable of consuming their own body mass in prey each day. 1 It is thus important that the effects of insect pest control treatments on adults of C. carnea be minimised to ensure the production of successive generations of this biological control agent. 2 The adult stages of C. carnea have a palyno-glycophagous diet, and are noted to feed on plant exudates, sap, honeydew, nectar and pollen. 1 A recent study has shown that their ability to digest and utilise pollen is very high. 3 With C. carnea larvae and pupae remaining in constant contact with the plant, and adults feeding directly on plant products, C. carnea has a potentially high exposure to plant protection chemicals.

Deleterious effects caused by herbicides, acaricides and insecticides on the survival and fecundity of beneficial insects have been well documented, 4–7 as have the effects of insecticides on C. carnea. 8–10 Pyrethroids are the most widely used insecticide in many countries and in many different crops. Plapp and Bull 11 reported that C. carnea larvae were relatively tolerant to synthetic pyrethroids compared with other classes of insecticide. Larvae of the lacewing have been shown to have a contact LD 50 of 17 000 µg g −1 for permethrin and can thus tolerate doses in excess of the recommended application rates; 12 this value is also above that reported for many other resistant pests and beneficial insects. Tolerance may be due to a slow rate of pyrethroid binding and rapid clearance of these binding sites in neuronal sodium channels 7 or to high levels and activity of esterases. 12 Such increased tolerance was also linked to the ability of esterases rapidly to detoxify the cis-isomer of the common pyrethroids
permethrin and cypermethrin. cis-Isomers are the more toxic of the two common isomeric forms (cis and trans), and in many insect species esterases are only capable of detoxification of cis-isomers at a slow rate, and are thus unable effectively to protect the insect from pyrethroid toxicity.13 To the authors’ knowledge, no work has been carried out to date on the specific mechanisms by which adult *C. carnea* tolerate pyrethroids.

The use of transgenic crops expressing *Bt* toxins is an ever-growing market,14 and today the impact of *Bt* maize and cotton varieties on natural enemies is relatively well understood. This includes *C. Carnea*,15,16 either when fed directly using artificial diets or when fed via the plant/pest/predator interaction.17–23 All stages of the lacewing may be exposed to transgenic products, either directly or via trophic feeding. However, in spite of the fact that adults potentially have greater exposure levels to these products in cases where they are expressed in the pollen, there is only one study addressing the risks of *Bt* maize on the adult stage.3 Feeding adult *C. carnea* with *Bt* maize pollen expressing either Cry1Ab or Cry3Bb1 or elevated doses of the two toxins dissolved in an artificial diet does not cause any effect on a range of important life-table parameters.

Very little documented evidence exists on the impact of GM crops expressing non-*Bt* genes on *C. carnea*, although one study by Lawo and Romeo24 reported no detrimental effects of direct feeding on the protease inhibitor SBTI.

The potential for pest control measures based on the use of plant protease inhibitors (PIs) has long been noted.25 Biotechnology offers the opportunity both to increase expression of endogenous protease inhibitors and/or to express foreign inhibitors in crop plants.26 Protease inhibitors have been shown to have the potential to control insect pests such as *Lacanobia oleracea* (L.),27 *Plutella xylostella* (L.), *Mamestra brassicae* (L.)28 and *Chrysomela tremulae* (F.).29 The efficacy of these protease inhibitors on insects is linked to a number of factors including their expression levels in plant tissues, the binding affinity between pest proteases and the protease inhibitor and the ability of the pest to adapt to inhibition.

The level of protease inhibitor expression is very important in providing effective control of pests, with low expression levels only achieving low levels of inhibition of digestive proteases, resulting in little effect either on insect development or on insect feeding patterns.28,30 High expression levels, on the other hand, can cause compensatory feeding effects in insects, thus causing an increase in crop damage.31 Insects can respond to protease inhibitors by altering their digestive proteases by increasing production of native enzymes31 or by synthesising insensitive forms of enzymes of the same or different mechanistic classes.32–39 The most successful examples of insect control via transgene expression of protease inhibitors result from use of protease inhibitors with a high affinity for the digestive protease in a targeted insect, as exemplified by Edmonds et al.40 Many PIs not only are effective against insect pests but also have the potential adversely to affect beneficial insects such as predators, either as a consequence of altered prey quality41,42 or by direct inhibition of the predator’s digestive proteases.

As little information is currently available on the digestive proteases present in *C. carnea*, one of the aims of this study was to further characterise protein digestion in this important predatory species. The most recent work carried out on neuropteran insects has revealed the presence of an acidic gut,43 suggesting a predominance of cysteine forms of digestive proteases. The protease inhibitor selected for use in this trial was oryzacystatin (OC-1), a cysteine protease inhibitor that is thermodynamically stable and has been shown to have strong affinity for cysteine proteases in many different insects.36,40,44

The major aim of the study, however, was to compare the effects of two pest control strategies on life history parameters of the beneficial insect *C. carnea*, as well as effects on the digestive proteases and xenobiotic detoxifying enzymes [P450s and esterases (ESTs)]. The potential effects of cypermethrin, one of the most widely utilised pyrethroid pest control measures, were compared with pest control based on transgenic oilseed rape (OSR) expressing the cysteine protease inhibitor OC-1.

## 2 MATERIALS AND METHODS

### 2.1 Insect cultures

Stocks of *C. carnea* were raised from cultures maintained at the Agricultural Research Station ART, Zurich, Switzerland. Larval cultures were maintained on UV sterilised eggs of *Ephestia kuehniella* (Zeller) stored at 2–4 °C. Adults were kept on an artificial diet (honey + yeast + water, 7 + 4 + 4) in cages within controlled environment rooms at 22 ± 3 °C and 70 ± 5% RH under a 16:8 h light:dark photoperiod.

### 2.2 Plant material

Homozygous transgenic spring oilseed rape seed (*Brassica napus* (L.) cv. Drakkar, line OC-1 Drakkar 4B expressing the cysteine protease inhibitor oryzacystatin-1) was grown from seed generated as previously described by Bonadé-Bottino.45 Control *B. napus* cv. Drakkar was grown under identical conditions of a 16:8 h light:dark photoperiod at a temperature of 20 ± 2 °C. Pollen was harvested and stored at −80 °C until required.

### 2.3 Reagents

EnzChek Protease assay kits (Molecular Probes) were obtained from GE Healthcare, Amersham. Transepoxysuccinyl-L-leucylamido (4-guanidio)butane (E-64), phenylmethylsulfonyl fluoride (PMSF), pepstatin A, ethylenediaminetetra-acetic acid disodium salt (EDTA), soybean Bowman–Birk inhibitor (SBBI) and papain were purchased from Bachem (UK) Ltd. Polyclonal antibodies were raised in rabbits by standard protocols against OC-1. Cypermethrin 100 g L−1 EC (Pyrimet) was supplied by United Agricultural Products Ltd, Greets Farm, Welburn, North Yorkshire. Ethyl acetate used in GC detection of cypermethrin was obtained from Fisher and was HPLC grade. All other chemicals were from Sigma (Poole, UK) and were of analytical grade unless otherwise stated.

### 2.4 Production of recombinant oryzacystatin (OC-1)

Recombinant oryzacystatin (rOC-1) was purified from *E. coli* by ion-exchange chromatography and purified according to the methods of Edmonds et al.40

### 2.5 Insect bioassays

*Chrysoperla carnea* were separated from stock culture at the pupal stage and stored individually in 10 mL sealed glass vials at 22 ± 3 °C and 70 ± 5% RH under a 16:8 h light:dark photoperiod. Immediately after emergence, gender was determined and males and females separated. Groups of five insects were each placed into one of fourteen 250 mL sealed pots, with access to damp cotton and artificial diet treated for the appropriate group (seven
pots of five males and seven pots of five females, n = 35 per sex per treatment). After 5 days, individual male/female pairs were separated into 100 mL pots and provided with damp cotton, and were fed 5 µg artificial diet per day for the remaining 17 days of the trial. Mortality and egg production was recorded daily throughout the trial. Egg hatching success was recorded for eggs laid on day 17 of the trial (day 17 was selected as it was only possible to assess hatching success once in the trial, and, on this day, high levels of oviposition occurred in all treatment groups). Hatching was classified as successful at the emergence of an L1 (first larval stadium). All adults in the trial were maintained in controlled environment rooms at 25 ± 3 °C and 70 ± 5% RH under a 16:8 h light:dark photoperiod regime. The four treatment groups were as follows (basic artificial diet: honey + yeast + water, 7 + 4 + 4):

- **control**: artificial diet containing 1% w/w control OSR pollen;
- **OC-1 control**: artificial diet containing 1% w/w control OSR pollen + 1% w/w rOC-1;
- **cypermethrin**: artificial diet containing 1% w/w control OSR pollen and 19 µM cypermethrin (1/100 dilution of recommended field application);
- **OC-1 pollen**: artificial diet containing 1% w/w OSR pollen expressing OC-1.

### 2.6 Determination of transgene expression levels in OSR leaves and pollen

Leaf samples were taken at random from transgenic and non-transformed control plants, flash frozen in liquid nitrogen, ground to a fine powder and extracted in 50 mM Tris-HCl buffer, pH 8.0 (containing 1% PMSF), as described previously. Extracts were centrifuged at 10,000 × g for 15 min, and total soluble protein of the supernatants was estimated by Bradford assay. For immunoassay by western blotting, samples (containing 50 µg total protein) were separated on an SDS-PAGE 15% minigel, and proteins were transferred electrophoretically to a 0.2 µm nitrocellulose filter. Oryzacystatin-1 was detected by enhanced chemiluminescence (ECL), as previously described, using polyclonal antibodies raised in rabbits against recombinant OC-1 as the primary antibody, with HRP-conjugated goat antirabbit IgG (Bio-Rad laboratories, Hertfordshire, UK) as the secondary antibody; rOC-1 (produced in E. coli by L Ceci, Institute for Biomembranes and Bioenergetic-CNR, Trian, Italy) was used as a positive standard. Following positive confirmation of expression of OC-1 in OSR plants, pollen was collected and levels of expression within the pollen grains were determined via the addition of OC-1 in OSR plants, pollen was collected and levels of expression positive standard. Following positive confirmation of expression of Biomembranes and Bioenergetic-CNR, Trani, Italy) was used as a light:dark photoperiod regime.

### 2.7 Determination of cypermethrin levels in Chrysoperla carnea guts by gas chromatography

Cypermethrin levels in C. carnea were analysed by gas chromatography with electron capture detection using a Hewlett Packard 5890 series II operated at 50 °C, and a Hewlett Packard 6890 series injector with ECD radioactive source for detection, as described by Mulligan et al. Extracts were prepared from C. carnea guts. Known weights of digestive tract samples were extracted in ethyl acetate (1 mL) for approximately 1 min by vortexing, then allowed to stand for a further 5 min. The extract (0.5 mL) was dried under nitrogen and then reconstituted in ethyl acetate (50 µL, using a microsyringe), thus equating to a tenfold concentration. The reconstituted extract was mixed using a vortex mixer, sonicated to ensure that the extract was fully redissolved, transferred to a 2 mL clear vial containing a 250 µL tapered insert and analysed by GC-ECD.

### 2.8 Demonstration of proteolytic activity in guts of Chrysoperla carnea

Digestive tracts from the different developmental stages of C. carnea were dissected into chilled distilled water (1 gut section to 20 µL), homogenised on ice and centrifuged at 14,000 × g for 10 min at 4 °C, the supernatants were precipitated with ammonium sulfate (to 65% saturation) and finally dialysed overnight at 4 °C against distilled water. Extracts from the different larval stadia and from male and female adults were kept separately so that subsequent activity could be related to the different developmental stages. General proteolytic activity was determined using the fluorescent protein substrate BODIPY-FL casein. Chrysoperla carnea extract (2 µL) from L1/L2/L3/adult female/adult male was incubated with 50 mM buffer (188 µL) at 25 °C. The reaction was initiated by the addition of 10 mM substrate (10 µL) to give a final substrate concentration of 0.5 mM. Fluorescence was monitored in a Fluoroskan Ascent FL fluorescence microtitre plate reader from Thermo LabSystems at 25 °C, excitation/emission maxima 485/538 nm, every 2 min over a 60 min period against appropriate controls. All assays were performed in triplicate. To determine the pH optimum, a range of overlapping buffer systems was used: acetate (pH 4 – 5.5), MES (2-morpholinoethanesulfonic acid; pH 5 – 7), HEPES [2-(4-(2-hydroxyethyl)piperazin-1-yl)ethanesulfonic acid; pH 6 – 7.5] and bis-Tris propane (pH 7 – 11). All buffers contained DTT and Brij35 at final assay concentrations of 5 mM and 0.1% (v/v) respectively.

### 2.9 Determination of digestive proteases by SDS-PAGE

General gut proteolysis was also determined using gelatin PAGE, where total soluble protein (10 µg) was run on a 12.5% SDS-PAGE minigel copolymerised with 0.1% gelatin. Sample-loading buffer did not contain 2-mercaptoethanol, and samples were not boiled prior to loading. Subsequently, SDS was eluted from the gel following electrophoresis with 1% (v/v) Triton X-100 in distilled water for 30 min at 4 °C. The gel was cut into strips and incubated in 100 mM borate/NaOH, pH 8.0, for 2 h at 37 °C prior to staining in Kenacid Blue.
2.10 Enzyme inhibition studies

Enzyme activity in the digestive tract of all *C. carnea* life stages was partially characterised using the class-specific protease inhibitors E-64, pepstatin A, EDTA and PMSF (final assay concentrations 10 μM, 1 mM, 10 mM and 1 mM respectively) and a range of synthetic and natural serine/cysteine protease inhibitors: TLCK, TPCK, chymostatin, elastatinal, CptP, SBBi, rOC-1 and leupeptin (final assay concentrations 100 μM, 100 μM, 100 μM, 100 μM, 1.2 μM, 1.25 μM, 1.2 μM, 2 μM and 100 μM respectively). Inhibitors were pre-incubated with crude enzyme preparations at 25 °C for 10 min, prior to addition of the substrate. Using pH papers, the digestive tract of *C. carnea* was determined to be approximately pH 8; all assays were carried out at pH 8, using bis-Tris propane buffer, and performed in triplicate with appropriate controls. All buffers contained DTT and Brij35 at final assay concentrations of 5 mM and 0.1% (v/v) respectively.

2.11 Enzyme preparations for analysis of detoxifying enzyme activity

Enzyme extracts were prepared from adult *C. carnea* guts. Guts were removed, washed with distilled water, homogenised (10 guts mL⁻¹) in ice-cold 0.1 M potassium phosphate buffer (pH 7.2, containing 1 mM EDTA, 1 mM DTT and 1 mM PMSF) and centrifuged at 4 °C, 10 000 × g for 20 min. The supernatant was removed and quantified, and appropriate dilutions (for the tests below) were made using potassium phosphate buffer; preparations were assayed immediately for monooxygenase, glutathione S-transferase (GST) and esterase activity (EST), using a range of different substrates, as detailed below. All enzyme assays throughout were carried out in triplicate with appropriate controls. Assays for *O*-demethylation of p-nitroanisole (PNOD) and the *O*-deethylation of 7-ethoxycoumarin (ECOD) by cytochrome P450 monooxygenases were conducted using the procedures of Yang et al.⁴⁹ Rose et al.⁵⁰ and Ullrich and Weber.⁵¹ Non-specific esterase activity was measured with the substrate α-naphthyl acetate (α-NA) at a final assay concentration of 1.9 mM, following the method of Yang et al.⁴⁹

The ability of *C. carnea* to hydrolyse the insecticide cypermethrin in vitro was determined by incubating cypermethrin in 5 μg mL⁻¹ enzyme extract in a total volume of 250 μL, 100 mM potassium phosphate buffer pH 7.2 (final substrate concentration 1 mM). The reaction was terminated at time points of 15 and 45 min by the addition of 250 μL of acetonitrile; the samples were centrifuged at 13 000 × g to pellet the precipitated protein, and 80 μL of the supernatant was analysed by HPLC. A Kontron Instrument HPLC 430 detector and Kontron 460 autosampler were used with a mobile phase of (A) 1% H₃PO₄ and (B) acetonitrile, stationary phase Phenomenex 250 × 4.60 mm and a 5 μm C18 100A column. The column was equilibrated in 65% B, samples were injected and a gradient to 100% B was carried out over a 2 min period, with a flow rate of 1 mL min⁻¹. Eluate from the column was analysed for UV absorbance at 264 nm.

2.12 Statistical analysis

Statistical analysis was carried out using one-way analysis of variance (ANOVA) (followed by mean separation via Tukey–Kramer tests) and two-tailed Student’s t-tests. All tests were performed using Minitab v.14 software on an iMac computer. Differences between treatments were considered significant at the *P* < 0.05 level. When data were not normally distributed, the non-parametric Kruskal–Wallis test replaced one-way ANOVA and the Mann–Whitney U-analysis replaced Student’s t-test.

### Table 1

Detection of OC-1 in *Chrysoperla carnea* frass collected after 17 days of feeding on artificial diet containing 1% w/w control OSR pollen (control), 1% w/w control OSR pollen + 1% rOC-1 w/w (OC-1 control), 1% w/w control OSR pollen and cypermethrin (cypermethrin) and 1% w/w pollen from OC-1-expressing OSR (OC-1 pollen). Figures represent the mean (± standard error) of three independent replicates from pooled insect samples.

<table>
<thead>
<tr>
<th>Feeding treatment group</th>
<th>Total OC-1 detected via ELISA (± SE) (ng)</th>
<th>OC-1 detected (% of total soluble protein)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0 (±0.02)</td>
<td>0</td>
</tr>
<tr>
<td>OC-1 control</td>
<td>3.8 (±0.04)</td>
<td>0.0019</td>
</tr>
<tr>
<td>Cypermethrin</td>
<td>0 (±0.01)</td>
<td>0</td>
</tr>
<tr>
<td>OC-1 pollen</td>
<td>1.1 (±0.02)</td>
<td>0.0005</td>
</tr>
</tbody>
</table>

3 RESULTS

3.1 Production and purification of recombinant oryzacystatin-1 (rOC-1)

Recombinant oryzacystatin was produced using *Escherichia coli* as the microbial expression system. Following purification by ion-exchange and reverse-phase chromatography, the protein was visualised as a single band of 14 kDa on SDS PAGE. Biological activity was confirmed by activity assay against papain, following the methods of Edmonds et al.⁴⁰ (data not presented).

3.2 Expression of OC-1 in pollen of transgenic OSR plants and detection in *Chrysoperla carnea*

A standard curve was constructed from rOC-1 standards (0–10 ng) and the concentration of OC-1 present in pollen extracts was calculated from the regression equation. An average of 0.4 ng OC-1 protein was detected from the 2 μg of total soluble protein in pollen extracts, equating to an expression level of 0.02% OC-1 in the transgenic pollen grains. Exposure of the predator to OC-1, either via OC-1-expressing pollen or artificial diet, was demonstrated in the frass by immunoassay (ELISA), where it was detected at a level of 0.001% in the experimental group fed purified rOC-1 in artificial diet, while in the group fed transgenic OC-1-expressing pollen it was only present at 0.0005% of total soluble protein (Table 1).

Unfortunately, insufficient material was available to quantify the levels present in the digestive tract.

3.3 Detection of cypermethrin in *Chrysoperla carnea* digestive tract tissue

Cypermethrin was quantified in the digestive tracts of *C. carnea* that had been exposed to the insecticide via a diet containing pollen treated with the recommended field dose. The results demonstrated that cypermethrin could be readily detected, with a mean value of 0.16 μg μL⁻¹, although these levels were highly variable (range 0.05–0.2 μg μL⁻¹). No insecticide was detected in control samples (limit of detection 0.02 μg μL⁻¹).

3.4 Characterisation of gut digestive proteases of *Chrysoperla carnea*

Analysis of gelatin PAGE gels, visualising the major gut proteases of the different life stages of *C. carnea*, revealed differences in proteolytic enzymes between the different larval instars and adults. First-instar larvae revealed an unresolved area of digestive enzymes between 50 and 70 kDa, and one further major band of activity at approximately 37 kDa (Fig. 1, L1), with trace levels of activity associated with two further lower-molecular-mass bands.
Impact of a cysteine protease inhibitor and a synthetic pyrethroid on *C. carnea* 

**Figure 1.** Digestive proteases from three different life stages of *C. carnea*. L1, first instar; L2, second instar; L3, third instar; Male, adult male; Female, adult female. Clear regions on the gel indicate gelatin hydrolysis at pH 8.

Figure 2. Determination of pH optima of digestive tract proteolysis in larval instars (a) and adults (b) of *C. carnea*, using the protein substrate BODIPY-FL casein. L1, first instar; L2, second instar; L3, third instar; Male, adult male; Female, adult female. Points and bars represent mean ± SE for triplicate independent determinations.

**Table 2a.** Endogenous gut proteolytic activity of male *Chrysoperla carnea* following ingestion of an artificial diet containing: 1% w/w control OSR pollen (control), 1% w/w control OSR pollen + 1% wOC-1 w/w (OC-1 control), 1% w/w control OSR pollen and cypermethrin (cypermethrin) and 1% w/w pollen from OC-1-expressing OSR (OC-1 pollen). Protease activity was measured with BODIPY-FL casein as substrate at pH 8. Figures represent the mean of three independent replicates carried out from pooled gut samples.

<table>
<thead>
<tr>
<th>Inhibitor (µM)</th>
<th>Control</th>
<th>OC-1 control</th>
<th>Cypermethrin</th>
<th>OC-1 pollen</th>
</tr>
</thead>
<tbody>
<tr>
<td>PMSF (1000)</td>
<td>71.9</td>
<td>58.3</td>
<td>75.0</td>
<td>76.7</td>
</tr>
<tr>
<td>Chymostatin (10)</td>
<td>68.7</td>
<td>55.6</td>
<td>66.7</td>
<td>70.0</td>
</tr>
<tr>
<td>SBBI (2.5)</td>
<td>68.7</td>
<td>83.3</td>
<td>66.7</td>
<td>63.3</td>
</tr>
<tr>
<td>E-64 (10)</td>
<td>6.2</td>
<td>33.4</td>
<td>6.2</td>
<td>10.7</td>
</tr>
<tr>
<td>Pepstatin A (1)</td>
<td>21.9</td>
<td>25.0</td>
<td>19.8</td>
<td>20.8</td>
</tr>
<tr>
<td>Elastatinal (10)</td>
<td>31.2</td>
<td>55.6</td>
<td>29.2</td>
<td>36.0</td>
</tr>
<tr>
<td>Leupeptin (10)</td>
<td>21.9</td>
<td>8.3</td>
<td>25.0</td>
<td>26.7</td>
</tr>
<tr>
<td>EDTA(1000)</td>
<td>25.0</td>
<td>22.2</td>
<td>22.9</td>
<td>26.7</td>
</tr>
</tbody>
</table>

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(approximately 30 and 24 kDa). Second-instar larvae again demonstrated the presence of unresolved protease activity, although this area included proteases as low as 40 kDa in size. Similar banding patterns were also present in the later larval instars, although the level of activity appeared to increase with increasing instar, with the exception of the band of approximately 24 kDa (Fig. 1).

The adult life stages showed no detectable differences in banding patterns between the two genders (Fig. 1), although metamorphosis did markedly alter the protease profile of *C. carnea*. Proteases capable of degrading gelatin were present, though unresolved, in the size range 100–55 kDa, with three major bands of activity detectable below this molecular mass. A band at approximately 39 kDa appeared to be unique to the adult life stage, whereas the band at 37 kDa present in the larval forms was absent. Bands present at 30 kDa and 24 kDa were present at significantly higher levels in adults than in the larvae, and appeared to be more prevalent in male *C. carnea* (Fig. 1).

Alteration of digestive proteases in the life cycle of the insect was reflected via changes in the pH profile of enzymes from the digestive tract. All three instars exhibited an optimal pH of 9.5 for the proteolysis of the protein substrate BODIPY-FL casein; these results suggest a predominance of serine proteases in the digestive tract. A second peak of activity was also observed at lower pH (5.5–7.5), which is more indicative of cysteine proteolytic digestion (Fig. 2a). All larval instars showed very low levels of proteolytic activity compared with adults (Fig. 2b). While possessing different protease profiles, a broadly similar pH profile was observed for larvae and adults. Male lacewings, in particular, exhibited a pronounced pH optimum of between 8 and 9.5, with significant proteolytic activity present in both sexes at pH 6–11 (Fig. 2b).

The use of diagnostic classes of inhibitor confirmed that the digestive profile of adult *C. carnea* was dominated by serine proteases (Tables 2a and b). Both the serine protease inhibitor PMSF and the proteinase inhibitor SBBI caused approximately 70% inhibition of proteolysis in both sexes; similarly, chymostatin was shown to cause a 64% and 69% inhibition of proteolysis in females and males respectively.

Inhibition by elastatinal (31% in males, 18% in females) suggested the presence of elastase-like enzymes, in addition to the dominant serine groups. The low levels of inhibition caused by the cysteine inhibitor E-64 suggested a low level of cysteine protease activity. This was confirmed with the use of leupeptin, although male *C. carnea* showed greater inhibition with leupeptin than females. Male *C. carnea* also showed substantial inhibition with EDTA compared with females, suggesting a role for metallo proteases in the adult male digestive tract (Table 2). Pepstatin A showed a slight difference between the sexes. The increased effect of this inhibitor (14% inhibition in females, 22% in males) in males may be linked to the greater intensity observed for gelatin digestion at 24 kDa (Fig. 1).

All three larval instars showed a strong susceptibility to inhibition by serine protease inhibitors. PMSF caused almost 100% inhibition of proteolysis in first- (L1) and second-instar (L2) larvae, but its effects were reduced in third-instar (L3) insects (with a 1 mM concentration only causing 35% inhibition) (Fig. 3). Chymostatin, a specific inhibitor of chymotrypsin forms of serine proteases, caused >80% inhibition in all larval instars; the dual trypsin/chymotrypsin
inhibitor, SBBI, caused very similar effects. These results strongly suggest that both trypsin and chymotrypsin enzyme forms are prevalent in the serine-dominated digestive profile of *C. carnea* (Fig. 3). As indicated by the somewhat elevated activity around pH 7 (Fig. 2a), cysteine protease activity was expected to increase slightly in later instars, and, in line with this, E-64 caused 5% inhibition in L1, 15% in L2 and 25% in L3, indicating relatively more cysteine protease activity in older instars. The inhibitor leupeptin, an inhibitor of both serine and cysteine proteases, on the other hand, caused a 51% decrease in activity in first-instar larvae, a 29% decrease in activity in second-instar larvae, but negligible effects upon third-instar larvae (Fig. 3). Similarly, EDTA, an inhibitor of metallo proteases, caused negligible inhibition in the final larval instar, but 62% in L1 and 36% in L2; this suggests the presence of metallo proteases early in the life cycle of *C. carnea*. Pepstatin A had no discernable effect upon proteolysis in any instar (data not shown).

### 3.5 Effects of rOC-1 in artificial diet on *Chrysoperla carnea*

#### 3.5.1 Effects on life history parameters

When fed at 1% w/w on artificial diet, the cysteine protease inhibitor OC-1 had no significant impact on the mortality of adult *C. carnea*, with 98% survival during the 17 day trial. Evidence that adults not only were exposed to the inhibitor but consumed it was confirmed by immunoassay, which clearly demonstrated the presence of rOC-1 in the frass (Table 1). Treatment with the purified cysteine protease inhibitor did, however, significantly reduce the number of eggs laid throughout the trial (df = 3, 88, F = 29.25, P = 0.002) (Table 3). Although six females fed rOC-1 did not produce eggs during this period, a figure comparable with that of the control group, females that did produce eggs laid an average of ca 20 fewer eggs female⁻¹ throughout the 17 day period (Tukey–Kramer test *P* = 0.002) (Table 3). This reduction in egg production was linked to a significant delay (*W* = 790, df = 1, *P* = 0.035) in the onset of egg production by female *C. carnea* by comparison with control females (Fig. 4). Once egg production had commenced, however, females feeding on rOC-1 laid at a rate comparable with that of the control females. Fecundity was also assessed, in terms of egg viability (Table 3). Eggs produced from females fed on a 1% w/w rOC-1 diet showed a level of hatching success (79.7%) equal to that of control *C. carnea* eggs, thus demonstrating that the inhibitor did not directly affect hatching success; however, over the limited trial period, fecundity was reduced as a consequence of a delay in the onset of oviposition.

#### 3.5.2 Effects on endogenous gut proteases

The effects of OC-1 on digestive proteases of instars and adult *C. carnea* were investigated in vitro (Fig. 5). The cysteine protease inhibitor had a strong effect on proteases of first-instar (L1) *C. carnea*, causing almost 50% inhibition of protease activity (at a concentration of 5 µM). However, the later larval instars were less susceptible to the effects of the inhibitor, with inhibition of protease activity decreasing slightly in later instars, and, in line with this, E-64 caused 51% decrease in activity in first-instar larvae, a 29% decrease in activity in second-instar larvae, but negligible effects upon third-instar larvae (Fig. 3).

### Table 2a. Endogenous gut proteolytic activity of female *Chrysoperla carnea* following ingestion of an artificial diet containing: 1% w/w control OSR pollen (control), 1% w/w control OSR pollen + 1% rOC-1 w/w (OC-1 control), 1% w/w control OSR pollen and cypermethrin (cypermethrin) and 1% w/w pollen from OC-1-expressing OSR (OC-1 pollen). Protease activity was measured with BODIPY-FL casein as substrate at pH 8. Figures represent the mean of three independent replicates carried out from pooled gut samples

<table>
<thead>
<tr>
<th>Inhibitor (µM)</th>
<th>Control</th>
<th>OC-1 control</th>
<th>Cypermethrin</th>
<th>OC-1 pollen</th>
</tr>
</thead>
<tbody>
<tr>
<td>PMSF (1000)</td>
<td>74.4</td>
<td>66.0</td>
<td>74.4</td>
<td>70.4</td>
</tr>
<tr>
<td>Chymostatin (10)</td>
<td>64.1</td>
<td>62.0</td>
<td>65.8</td>
<td>70.4</td>
</tr>
<tr>
<td>SBBI (2.5)</td>
<td>74.4</td>
<td>88.0</td>
<td>74.4</td>
<td>77.1</td>
</tr>
<tr>
<td>E-64 (10)</td>
<td>0</td>
<td>16.0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Pepstatin A (1)</td>
<td>14</td>
<td>7.7</td>
<td>4.3</td>
<td>14.8</td>
</tr>
<tr>
<td>Elastatinal (10)</td>
<td>17.9</td>
<td>32.0</td>
<td>17.9</td>
<td>29.6</td>
</tr>
<tr>
<td>Leupeptin (10)</td>
<td>7.7</td>
<td>22.0</td>
<td>9.4</td>
<td>11.1</td>
</tr>
<tr>
<td>EDTA (1000)</td>
<td>15.4</td>
<td>18.0</td>
<td>12.8</td>
<td>11.1</td>
</tr>
</tbody>
</table>

### Table 3. Egg production and egg hatching success of *Chrysoperla carnea* following ingestion of artificial diet containing: 1% w/w control OSR pollen (control), 1% w/w control OSR pollen + 1% rOC-1 w/w (OC-1 control), 1% w/w control OSR pollen and cypermethrin (cypermethrin) and 1% w/w pollen from OC-1-expressing OSR (OC-1 pollen). The numbers of eggs produced per female from the different treatments were submitted to one-way ANOVA, while pre-oviposition periods and hatching success were analysed by Mann–Whitney U-tests. In all cases, values followed by different letters are significantly different (*P* < 0.05)

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>Number of fertile females</th>
<th>Number of non-laying females</th>
<th>Median day females commenced oviposition</th>
<th>Total number of eggs laid by day 17</th>
<th>Mean number of eggs laid per female (± SE)</th>
<th>Overall % hatching success (± SE) from surviving insects on day 17 of the trial</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>30</td>
<td>5</td>
<td>8.5 a</td>
<td>1912</td>
<td>63.7 (±19) a</td>
<td>80.0 (±2.1) b</td>
</tr>
<tr>
<td>OC-1 control</td>
<td>29</td>
<td>6</td>
<td>11.0 b</td>
<td>1276</td>
<td>44.0 (±15) b</td>
<td>79.7 (±2.2) b</td>
</tr>
<tr>
<td>Cypermethrin</td>
<td>27</td>
<td>8</td>
<td>12.0 c</td>
<td>750</td>
<td>27.7 (±14) c</td>
<td>75.6 (±3.6) a</td>
</tr>
<tr>
<td>OC-1 pollen</td>
<td>31</td>
<td>4</td>
<td>8.0 a</td>
<td>2073</td>
<td>66.8 (±19) a</td>
<td>80.9 (±2.6) b</td>
</tr>
</tbody>
</table>
Susceptible to inhibition, decreasing to 40 and 30% in second (L2) and third-instar (L3) larvae respectively. These results are in contrast to predicted results, as cysteine proteolytic activity was seen to increase slightly at a lower pH with larval age, but, more importantly, the cysteine inhibitor E-64 was a more potent inhibitor in the older larvae (Fig. 3), indicating relatively greater cysteine protease activity. Both sexes of adult C. carnea showed only 10% inhibition (in vitro) to OC-1; this is consistent with results demonstrating a limited effect of the cysteine inhibitor E-64 on adult digestive proteases (Tables 2a and b).

Although consumption of OC-1 by C. carnea adults did not appear to alter the protease profile as visualised by gelatin PAGE (data not shown), the presence of the inhibitor caused a change in the relative activities of the proteases present in the digestive tract (Tables 2a and b). Male C. carnea showed a decrease in sensitivity to inhibition by PMSF and chymostatin, indicating an increase in serine and particularly chymotrypsin proteases in response to feeding on rOC-1, while SBBI caused a greater inhibitory effect, indicating a switch away from trypsin proteases. Feeding on rOC-1 at 1% w/w also caused increased sensitivity to inhibition by both E-64 and elastatinal (by approximately 30% and 24%, respectively, compared with the controls), but pepstatin A, an inhibitor of aspartic proteases, showed almost identical levels of inhibition to that in control male adult lacewings (Table 2a).

Results from inhibition studies demonstrated that OC-1 was a weak inhibitor of adult lacewing digestive proteases in vitro (Fig. 5), and that, when the purified inhibitor was ingested, the predator responded by an alteration in levels of different protease classes (Tables 2a and b; Section 3.5.2). However, when adults were fed on a diet containing 1% transgenic pollen, very few changes were observed in the digestive protease profile (Tables 2a and b). Female C. carnea showed a slight increase in inhibition by elastatinal, a response clearly discernable when they were fed the purified protein; males also showed a slight increase in susceptibility to this inhibitor.
3.7 Effects of cypermethrin on *Chrysoperla carnea*

3.7.1 Effects on life history parameters

Cypermethrin fed at a 100-fold dilution of the recommended field preparations was high (average) than either control-fed females (Tukey–Kramer = 0.001), with eight females laying no eggs (Table 3), the lowest recorded number of eggs laid for any treatment group. Females that did produce eggs laid significantly fewer eggs (just 35.7 on average) than either control-fed females (Tukey–Kramer P = 0.001) or any other experimental group (Tukey–Kramer P = 0.022 for OC-1 control and Tukey–Kramer P = 0.0002 for OC-1 pollen) (Table 3). The insecticide-treated group also exhibited the greatest delay in the onset of oviposition (Fig. 4), this being significantly greater compared with either the control-fed group or the group fed transgenic pollen (W = 662, df = 1, P = 0.02 and W = 263, df = 1, P = 0.03 respectively). Egg viability was also significantly reduced in the insecticide-fed group compared with the control, the OC-1 control and the OC-1 pollen (W = 616, 321 and 744 respectively, df = 1 and P ≤ 0.05 in all cases) (Table 3).

3.7.2 Effects on endogenous gut proteases and enzymes involved in detoxification

Feeding on cypermethrin caused no significant changes in the digestive protease profile of adult *C. carnea*. Gelatin PAGE did not reveal the presence of any novel forms of protease (data not shown), and analysis with diagnostic inhibitors showed no alterations in the levels of protease forms present in adult *C. carnea* of either sex (Tables 2a and b). HPLC analysis of cypermethrin digests (Fig. 6) exposed for different time periods to *C. carnea* digestive tract enzyme preparations *in vitro* clearly demonstrated that insects with no previous contact with the insecticide were capable of rapidly degrading (within 15 min) 1 mM cypermethrin. Exposure of adults to the insecticide when fed caused no alteration in this response, as the enzymes remained capable of rapidly degrading the insecticide *in vitro*, so producing a variety of digestion products (Fig. 6). The addition of NADPH caused no alteration in either the pattern of breakdown products or the rate of breakdown, suggesting a limited role of P450 activity in insecticide degradation.

### Table 4. Activity of *Chrysoperla carnea* digestive tract crude enzyme extract against substrates specific for P450, GST and esterase (EST) activity following ingestion of artificial diet containing: 1% w/w control OSR pollen (control), 1% w/w control OSR pollen + 1% rOC-1 w/w (OC-1 control), 1% w/w control OSR pollen and cypermethrin. Insecticide and 1% w/w pollen from OC-1-expressing OSR (OC-1 pollen). Assays measured the O-demethoxylation of p-nitroanisole (PNOD), the O-deethoxylation of 7-ethoxycoumarin (ECOD) and the breakdown of 1-chloro-2,4-dinitrobenzene (CBND) and α-naphthyl acetate (α-NA). Figures represent the mean of three independent replicates carried out from three pooled insect samples per treatment.

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>P450 assay</th>
<th>GST assay</th>
<th>EST assay</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PNOD V&lt;sub&gt;max&lt;/sub&gt;</td>
<td>ECOD V&lt;sub&gt;max&lt;/sub&gt;</td>
<td>CDNB V&lt;sub&gt;max&lt;/sub&gt;</td>
</tr>
<tr>
<td>Control</td>
<td>0.03</td>
<td>0.05</td>
<td>0.26</td>
</tr>
<tr>
<td>OC-1 control</td>
<td>0.01</td>
<td>0.03</td>
<td>0.42</td>
</tr>
<tr>
<td>Cypermethrin</td>
<td>0.04</td>
<td>0.04</td>
<td>0.51</td>
</tr>
<tr>
<td>OC-1 pollen</td>
<td>0.04</td>
<td>0.02</td>
<td>0.32</td>
</tr>
</tbody>
</table>
Preliminary enzyme assays were carried out with synthetic substrates for P450s, GSTs and ESTs. While PNODE, ECOD (P450 substrate) and CDND (GST substrate) were hydrolysed at a negligible rate (Table 4), α-NA, the substrate for non-specific EST activity, was degraded rapidly (V_{max} = 21.45 ηOD min⁻¹ 100 μg protein⁻¹) in both control samples (no exposure to insecticide) and all experimental groups (the rate was highest, though not significantly so, in the insecticide fed group). This may suggest that ESTs play a key role in the insect’s ability to degrade insecticides.

4 DISCUSSION

4.1 Characterisation of major digestive proteases in the gut of Chrysoperla carnea

The use of synthetic substrates and diagnostic inhibitors suggests that serine proteolytic enzymes dominate the digestive tract of C. carnea, with both trypsin and chymotrypsin being present at high levels in the larvae and adults. In adults, approximately 80% of the protease activity was shown to be susceptible to inhibition by serine protease inhibitors, with the remaining activity being attributed to metallo or cysteine proteases. Furthermore, gelatin PAGE analysis revealed the presence of a protease (37 kDa) that increased in activity with each subsequent instar; the evidence suggests that this is most likely a further serine protease. Across both adult and larval instars the digestive tract of C. carnea is dominated by serine proteases, with large changes seen in the levels of the different protease forms throughout the life cycle.

First-instar larvae show higher levels of sensitivity to inhibition by metallo and leupeptin inhibitors compared with the later stages, with the converse being the case for cysteine protease inhibitors. Cysteine proteases, on the other hand, were shown to play a limited role in protein digestion, with levels of proteases sensitive to inhibition by the cysteine protease inhibitor E-64 being negligible in first-instar larvae, increasing to approximately 30% of the total proteolytic activity in third instars and declining to just 10% in the adult. To date, relatively few studies have been carried out to characterise the digestive enzymes present in the order Neuroptera, and in particular C. carnea. Data provided in the present study support earlier findings by Ferran et al. and Yazlovskey, who demonstrated the dominance of trypsin and chymotrypsin forms of proteases present in the larvae of this predator. More recently, Mochizuki et al. demonstrated that the beneficial predator Harmonia axyridis (Pallas), when exposed to OC-1-expressing OSR via a tritrophic interaction, was able to respond by the upregulation of native proteases, while the carabid Pterostichus madidus (Fabricius) responded to MTI-2-expressing plants by switching from predominantly trypsin to predominantly chymotrypsin.

4.2 Effects of the cysteine protease inhibitor OC-1 on Chrysoperla carnea

While in vitro studies would suggest a limited impact of OC-1 on C. carnea, and the feeding of rOC-1 in the diet at 1% w/w caused no negative impact on survival, the fecundity of the beneficial insect was reduced. There was both a significant delay (three days) in oviposition and in the total numbers of eggs laid over the 17 day trial. A reduction in fecundity has previously been noted with protease inhibitors, and in particular with OC-1 for other arthropods. However, in these cases a much greater impact on digestive proteases, nutrient availability and consequently development was linked to the decrease in fecundity. Furthermore, Rahbe et al. also demonstrated a decrease in aphid survival in the presence of the Bowman–Birk inhibitor which targets both trypsin and chymotrypsin.

Analysis of the levels of OC-1 detected in the frass of adult C. carnea showed that predators exposed to the transgenic pollen had only 25% of the level of inhibitor compared with those consuming the diet containing the rOC-1. Feeding OC-1-expressing pollen had only 25% of the level of inhibitor compared with those consuming the diet containing the rOC-1. Nevertheless, the purified recombinant OC-1 did cause limited effects on protein digestion and fecundity, demonstrating a potential for toxicity.

To date, a wide variety of protease inhibitors have been expressed in crop plants. Thus, given the potential for rOC-1 to affect lacewing fecundity, it is important that other protease inhibitors, particularly those targeted to the serine-dominated C. carnea digestive tract, be evaluated for potential negative effects. Interestingly, a study by Lawo and Romeis revealed no detrimental effect on important life-table parameters of C. carnea larvae when directly fed with SBTI at a dose of 1% w/v in a sucrose solution. As control strategies based on the expression of a combination of different protease inhibitors are being investigated, it is important that the effects of such combinations be evaluated more thoroughly on beneficial insects.
4.3 Effects of the pyrethroid cypermethrin on Chrysoperla carnea

One of the major objectives of the present study was to compare the effects of recombinant DNA technology with the use of synthetic insecticides for insect pest control on beneficial insects. The insecticide utilised in this study was the pyrethroid cypermethrin. Results obtained clearly demonstrate that this insecticide exhibited more deleterious effects on C. carnea than either the transgenic OC-1-expressing pollen or the purified rOC-1. While the insecticide exhibited little effect on mortality, fecundity (rate of egg production) and fertility (hatching success) were significantly reduced. Chrysoperla carnea adults were able to tolerate cypermethrin in the digestive tract at 0.16 µg L⁻¹, presumably as a consequence of rapid degradation of the compound owing to the increased levels of non-specific esterase (EST) activity that were observed. Similar findings were reported by Ishaaya and Casida for C. carnea larvae in response to exposure to the insecticide permethrin. While this increase in activity appeared to protect mature lacewings from mortality, it did not prevent a significant reduction in fecundity. In the present study, insecticide-fed females laid the lowest number of eggs throughout the 17 day trial, with subsequent hatching success in this group being significantly lower than either the control or transgenic pollen-fed groups; however, fertility, and in particular fecundity, remained high (74%), even in cypermethrin-treated insects. Determination of the significance of such a reduction in fecundity on active field populations of the chrysopid would be essential for a thorough evaluation of the environmental impact of synthetic pyrethroids. Although the trial was terminated before the completion of the reproductive cycle of C. carnea, feeding on a diet containing purified OC-1 (1% w/w) did delay the onset of egg production; however, subsequent fecundity rates and fertility were the same as for the control. It would be interesting to determine whether ingestion of a serine protease inhibitor, which more effectively targets the digestive proteases of the predator, would have caused more deleterious effects. At day 17 of the trial, insecticide-fed C. carnea produced eggs at a significantly lower rate than in all three other feeding treatments; further trials are needed to establish the effects on fecundity throughout the adult female life stage.

Few studies have investigated the effects of feeding cypermethrin-contaminated food to adult C. carnea. Studies by Ishaaya and Casida estimate the LD₅₀ of this insecticide in larval C. carnea to be 17 µg L⁻¹ when applied topically, although Rumpf et al. demonstrated that topical doses of 2.4 µg L⁻¹ caused no impact on survival of larval C. carnea. In contrast to results presented here, where continuous exposure to cypermethrin at 19 µL in artificial diet caused sublethal effects, work by Grafton-Cardwell and Hoy reported no significant reduction in either survival or fecundity when the two pyrethroids permethrin and fenvalerate were applied at recommended field rates to C. carnea. Work by Hassan et al. is the only study to show a direct effect of pyrethroids (permethrin and fenvalerate) on C. carnea, although effects on larval mortality were analysed in conjunction with adult fecundity and it is thus difficult to determine which stage was affected.

While this study has demonstrated the potential for pest management to impact on predators, more importantly it highlights the ability of the predator to detoxify/respond to two diverse treatments. In this case, exposure to the insecticide carried a greater fitness cost (in terms of impact on fecundity) than exposure to a PI expressed in transgenic crops.

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REFERENCES
Impact of a cysteine protease inhibitor and a synthetic pyrethroid on C. carnea


