Antifeedant Activity of *Ginkgo biloba* Secondary Metabolites against *Hyphantria cunea* Larvae: Mechanisms and Applications

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Abstract

*Ginkgo biloba* is a typical relic plant that rarely suffers from pest hazards. This study analyzed the pattern of *G. biloba* pest hazards in Beijing; tested the antifeedant activity of *G. biloba* extracts, including ginkgo flavonoids, ginkgolide, and bilobalide, against *Hyphantria cunea* larvae; determined the activities of glutathione transferase (GSTs), acetylcholinesterase (AChE), carboxylesterase (CarE) and mixed-functional oxidase (MFO), in larvae after feeding on these *G. biloba* secondary metabolites; and screened for effective botanical anti-feedants in the field. In this study, no indicators of insect infestation were found for any of the examined leaves of *G. biloba*; all tested secondary metabolites showed significant antifeedant activity and affected the activity of the four larval detoxifying enzymes. Ginkgolide had the highest antifeedant activity and the most significant effect on the detoxifying enzymes (P < 0.05). Spraying leaves with *G. biloba* extracts or ginkgolide both significantly repelled *H. cunea* larvae in the field (P < 0.05), although the former is more economical and practical. This study investigated the antifeedant activity of *G. biloba* secondary metabolites against *H. cunea* larvae, and the results provide new insights into the mechanism of *G. biloba* pest resistance. This study also developed new applications of *G. biloba* secondary metabolites for effective pest control.

Introduction

During the long-term co-evolution with insects, plants gained the ability to produce a plethora of secondary metabolites through accidental genetic mutation and genetic recombination. Secondary metabolites are irrelevant for normal plant growth, but have key roles for chemical defense against insects by killing, repelling, inhibiting the feeding, or hindering the growth of insects [1]. These secondary metabolites also have important effects on insects’ selection of host plants [2]. During co-evolution, plants developed a chemical defense system of secondary metabolites to prevent insect feeding, and insects adaptively developed detoxifying enzymes to defeat this plant chemical defense system [3].
Ginkgo biloba is a rare and endangered tree species unique to China, and the only living member of the G. biloba order of plants, which historically included at least seven genera as indicated by the fossil record. G. biloba has been widely cultured in China for at least 1,000 years, and all G. biloba cultured in other countries are derived from China, making it a typical cash crop [4]. G. biloba produces many important secondary metabolites, among which ginkgo flavonoids and ginkgolide are unique to this plant. These secondary metabolites accumulate to relatively high contents in G. biloba, and the levels may differ among different plant parts and in different seasons. Ginkgo flavonoids accumulate to the highest levels in tender leaves, and ginkgolides have the highest levels during July to September [5–6].

Hyphantria cunea (Lepidoptera, Arctiidae) originated in North America. It is a major invasive species on the international quarantine list, and was first identified in China in 1979. These insects feed on the leaves of many plants, rapidly spread, and cause serious ecological disaster [7–8]. The detoxifying system of H. cunea includes glutathione transferase, carboxylesterase and mixed-functional oxidase, which have important roles in degrading plant secondary metabolites. Acetylcholinesterase is a key enzyme involved in synaptic conduction of insects, and is an important target for insecticides [9–10].

Botanical antifeedants are a class of compounds that inhibit insect feeding, although they do not directly kill insects. Most plant secondary metabolites that show antifeedant activity can be classed into the following four categories: sesquiterpene lactones, heterogeneous flavonoids, quassins, and limonoids. Some of these agents show only relative antifeedant activity within a certain period, but they do affect insect host selection. Botanical antifeedants are rapidly degraded after application, thereby causing little environmental impact [11].

Ginkgo biloba is a typical relic plant with strong resistance to various insects. Field observations of G. biloba rarely reveal any signs of pest damage. Sometimes, mild and sparsely distributed insect damage can be observed, but G. biloba rarely exhibits disastrous pest damage [12–14]. So, this study investigated the antifeedant activity of secondary metabolites that are unique to G. biloba. The results provide new insights into the mechanism underlying the strong resistance to insects, and show that application of G. biloba secondary metabolites provides effective antifeedant activity.

Materials and Methods
Antifeedant Agents and Source of Insects
Ginkgo biloba extracts contain approximately 25% ginkgo flavonoids and 5% ginkgolide. The ginkgo flavonoids used in this study contained quercetin, kaempferol, and isorhamnetin (Sigma-Aldrich, USA) at a ratio of 2:2:1 (w/v). The ginkgolide used in this study contained bilobalide and ginkgolide A/B/C (Sigma-Aldrich, USA) at a ratio of 5:2:2:1 (w/v).

Normal artificial diet and H. cunea egg mass were purchased from the Chinese Academy of Forestry. The H. cunea larvae grew, developed, and reproduced normally after feeding the normal artificial diet [15]. H. cunea larvae for indoor studies were all hatched from the same egg mass, and were starved for 8 hours before using in experiments.

Analysis of Pest Feeding
This study was conducted in areas populated by G. biloba in Beijing City in June 2014. A total of 30 G. biloba trees were selected in each of 10 sample plots. Permission for each sample investigation was authorized by Beijing Municipal Bureau of Landscape and Forestry, and activities at each location were supervised by relevant staff from the department responsible. Three distinct classes rating tree health were established to analyze pest infestation (Table 1) [16]. Tree height, diameter, health status, and site and type of insect infestation were measured and
recorded. The pest resistance percentage was calculated as the number of pest-free trees among the total number of trees in the plot.

\[
\text{Pest resistance \%} = \frac{\text{Number of pest free trees}}{\text{Total number of trees in the plot}} \times 100
\]

Verification of Antifeedant Activity

To make the normal diet, feed powder (20 g), distilled water (80 ml), and agar power (1.6g) were mixed and heated to completely dissolve the agar. To make the test diet, G. biloba secondary metabolites were dissolved in 10 ml ethanol before mixing them with the normal diet at different concentrations. All the diets were poured into a rearing box (100ml, d = 5cm) and allowed to solidify. The resulting circular pie (δ = 2mm) was then cut into two equal semicircles for future use. Six other rearing boxes of the same shape was prepared, and a semicircular pie of the normal diet and six semicircular pies containing 4%, 8%, or 16% G. biloba extract, 2% flavonoids, 0.4% ginkgolides, or 0.2% bilobalide were placed in the boxes. Then 30 third-instar H. cunea larvae were put into each box, and the number of larvae choosing to feed on each of the semicircular pies in the boxes was recorded after 24h. These experiments were repeated five times [17–20].

Detoxifying Enzyme Activity Assay

Circular diet pies containing 8% G. biloba extracts, 2% ginkgo flavonoids, 0.4% ginkgolide, 0.2% bilobalide, or no extra additive were prepared as described above, to feed 120 third-instar H. cunea larvae in each group. The activities of glutathione transferase, acetylcholinesterase, carboxylesterase and mixed function oxidase were measured at 12, 24, 36, and 48 h for each group of 30 larvae. Third-, fourth-, and fifth-instar H. cunea larvae were fed on a diet containing the above four G. biloba secondary metabolites, and the activities of the above detoxifying were measured at 24 h in each group of enzymes in each group of 30 larvae. These experiments were repeated five times.

Glutathione transferase (GSTs) activity was measured using Lee's method. Briefly, the tissue sample was weighed, and 9 volumes of normal saline were added to produce a 10% tissue homogenate, which was then centrifuged at 12,000 rpm for 30 min at 4°C. The supernatant was considered as the enzyme solution. The enzyme reaction was measured in a cuvette containing 2.4 mL phosphate buffer (66 mmol/L, pH 7.0), 0.1 mL enzyme solution, 0.3 mL of 50 mmol/L glutathione (GSH), and 0.1 mL of 30 mmol/L 1-chloro-2,4-dinitrobenzene (CDNB), and changes in absorption at 340 nm were recorded. The measurement was repeated five times [21].

Acetylcholinesterase (AChE) activity was measured using a modified Ellman method. Briefly, the tissue sample was weighed, and 9 volumes of normal saline were added to produce a 10% tissue homogenate, which was then centrifuged at 3,500 rpm for 10 min at 4°C. The

| Classification | Number of emerged pests or defecation holes | Proportion of leaves with symptoms of pest feeding |
|---------------|--------------------------------------------|--------------------------------------------------|
| Class 1       | 0                                          | 0%                                               |
| Class 2       | 1–5                                        | 1–50%                                            |
| Class 3       | >5                                         | >50%                                             |

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supernatant was collected for enzyme activity measurements. Supernatant (0.1 mL) and ATCh (0.1 mL) (a substrate of AChE) were mixed together and incubated at 30°C for 15min, and then 3.6 mL DTNB was added to terminate the reaction. Absorbance was measured at 412nm. The experiment was repeated five times [22–23].

Carboxylesterase (CarE) activity was measured using Asperen’s method. Briefly, the tissue sample was weighed, and 9 volumes of normal saline were added to produce a 10% tissue homogenate, which was then centrifuged at 12,000 rpm for 30 min at 4°C. The supernatant was collected and diluted 50× with phosphate buffer before using for enzyme activity measurement. The enzyme reactions contained 0.3 mL of the diluted supernatant, 0.7 mL of 0.04 mol/L phosphate buffer, and 5 mL substrate, which were mixed and incubated on a shaker for 30 min at 37°C. Then, 1 mL of chromogenic reagent was added, and the reactions were incubated for a further 30 min at room temperature, followed by measurement of absorbance at 600 nm. The experiment was repeated five times. Known concentrations of α-naphthol were used to obtain a standard curve [24].

Mixed function oxidase (MFO) activity was measured using Yu and Nguyen’s method. Tissue samples were weighed and homogenized in 200μl ice-cold homogenization buffer (10% glycerol in 0.1 mol/L phosphate buffer, pH7.5). Then the homogenate was centrifuged at 4°C, 12000r/min for 10min, and the supernatant was centrifuged again. The final supernatant was collected for enzyme activity measurements. Paranitroanisole was added to the supernatant as a substrate for MFO, and the para-nitrophenolate produced was used to calculate MFO activity. The activity of each sample was measured five times [25].

Protein content was measured according to the Bradford’s method using bovine serum albumin as the standard.

Leaf-Disc Assay

*Ginkgo biloba* extracts, ginkgo flavonoids, ginkgolide, and bilobalide were dissolved in 20 mL absolute alcohol:water [1:7 (v/v)] solution in a concentration series (Table 2). Fresh *Fraxinus americana* leaf discs (1.5 cm diameter) were prepared with a drilling machine from healthy and pesticide-free leaves, and were soaked in the secondary metabolite solutions or alcohol: water [1:7 (v/v)] (as a control) for 10 sec. Four of the treated leaf discs were placed into each empty rearing box (100 mL, d = 5cm) to feed 30 third-instar *H. cunea* larvae for 24 h. Then, the leaves were removed from the boxes and placed on 1×1 mm graph paper to measure the eaten area by counting the grids. The experiment was repeated five times, and the antifeedant rate was calculated [26].

\[
\text{Antifeedant rate} = \frac{\text{Eaten area of the control group} - \text{Eaten area of the test group}}{\text{Eaten area of the control group}} \times 100
\]

**Field Antifeedant Test**

For field antifeedant tests, 2 g of *G. biloba* extracts, 0.5g of ginkgo flavonoids, 0.1g ginkgolide and 0.05g of bilobalide were respectively dissolved in 10 mL absolute alcohol, mixed with seven volumes of water, and stored in a 100 mL plastic watering can that was properly labeled. Alcohol: water [1:7 (v/v)] was used as the vehicle control. The tests included 183 branches of 24 *Salix* trees with a diameter of 12–21 cm in the suburban area of Tianjin, China, an area of 0.08 km² where Salix trees suffered severe damage from *H. cunea* larvae infestation in August 2015 (Table 3). Photographs were taken to record the insect feeding situation. The five solutions (four secondary metabolites and one control) were sprayed on infested leaf surfaces 3–5 times,
and the larval feeding situation was recorded by photographing again from the same perspective 5 h later. The repelling rate was calculated by comparing the number of larvae on the photographs taken before and after spraying [27].

\[
\text{Repelling rate} = \frac{\text{Number of larvae before}}{\text{Number of larvae after}} \times 100
\]

Data Analysis

Variance analysis and data processing were performed using SPSS 17.0 or Prism 6.0. One-way ANOVA followed by least significant difference test (LSD, at a level of statistical significance of \( P = 0.05 \)) were used to analyze the percentage of larva feeding on a given diet or food source, the detoxifying enzyme activities, antifeedant rate, and repelling rate. All of the data expressed as percentages were arcsine transformed and analyzed using SPSS 17.0.

Results

Pest Damage to \( G. \) biloba Trees in Beijing City

The average height of \( G. \) biloba trees in the 10 sample plots was 6.1–15.1 m, and the average diameter was 13.3–49.9 cm. Symptoms of pest feeding on the trees are summarized in Table 4. All pest feeding symptoms were confined to the trunk, and no symptoms were observed on leaves. Approximately 93.3% or more trees in the selected plots were free of insect damage in the same year. Only six \( G. \) biloba trees in Plots A, C, F, and G were damaged by \( Holcocerus \) insularis Staudinger (Lepidoptera, Cossidae) larvae, and only two trees were rated as Class 3 damage. No damage by any insect was found in the other six plots.

Table 2. Concentration series for four \( Ginkgo \) biloba secondary metabolites.

| Secondary metabolite | High(g/L) | Medium(g/L) | Low(g/L) | Control group(g/L) |
|----------------------|-----------|-------------|----------|---------------------|
| GBE                  | 50        | 25          | 12.5     | 0                   |
| GF                   | 12.5      | 6.25        | 3.125    | 0                   |
| GL                   | 2.5       | 1.25        | 0.625    | 0                   |
| BB                   | 1.25      | 0.625       | 0.3125   | 0                   |

GBE, \( Ginkgo \) biloba extract; GF, gingko flavonoids; GL, ginkgolide; BB, bilobalide.

Table 3. Instars of \( Hyphantria \) cunea larvae infesting branches treated with \( Ginkgo \) biloba secondary metabolites.

| Instar | 1st | 2nd | 3rd | 4th | 5th | Total |
|--------|-----|-----|-----|-----|-----|-------|
| GBE    | 10  | 11  | 6   | 6   | 6   | 39    |
| GF     | 13  | 9   | 7   | 5   | 5   | 39    |
| GL     | 15  | 10  | 8   | 5   | 6   | 44    |
| BB     | 9   | 10  | 7   | 6   | 5   | 36    |
| CG     | 5   | 5   | 5   | 5   | 5   | 25    |

GBE, \( Ginkgo \) biloba extract; GF, gingko flavonoids; GL, ginkgolide; BB, bilobalide; CG, control group.

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Antifeedant Activity of Different *G. biloba* Secondary Metabolites against *H. cunea* Larvae

The two curves (Fig 1) show that the proportion of *H. cunea* larva feeding on the diet containing *G. biloba* secondary metabolites was significantly lower than the number feeding on the normal diet (F = 36.579; df = 5, 48; P < 0.0001). *Ginkgo biloba* extracts significantly repelled *H. cunea* larvae (F = 6.492; df = 2, 12; P = 0.023), and the effect was stronger at higher extracts concentrations (Fig 1A–1C). The concentration of ginkgo flavonoids, ginkgolide, and bilobalide were defined according to 8% *G. biloba* extracts. Comparing these results with those of Fig 1D–1F and 1B, it can be concluded that ginkgo flavonoids, ginkgolide, and bilobalide significantly repel *H. cunea* larvae (F = 13.585; df = 3, 16; P = 0.009), and ginkgolide and bilobalide show stronger repelling effects than that of the 8% *G. biloba* extracts.

Effects of *G. biloba* Secondary Metabolites on Detoxifying Enzyme Activities

The activities of four detoxifying enzymes of *H. cunea* larvae increased after feeding on diet containing *G. biloba* secondary metabolites, but the extent to which the activities varied differed because of differences in the detoxification mechanisms of the enzymes. GSTs activity was affected most strongly by *G. biloba* extracts in the Fig 2A (F = 37.965; df = 4, 80; P < 0.0001) and B (F = 40.861; df = 4, 60; P < 0.0001), CarE activity responded more rapidly to ginkgolide and bilobalide in the Fig 2C (F = 67.249; df = 4, 60; P < 0.0001) and D (F = 30.852; df = 4, 80; P = 0.001), and AChE activity was affected most strongly by ginkgolide in the Fig 2E (F = 30.327; df = 4, 80; P = 0.001) and F (F = 33.985; df = 4, 60; P < 0.0001), MFO activity responded more rapidly to ginkgo flavonoids and ginkgolide in Fig 2G (F = 33.416; df = 4, 80; P < 0.0001) and H (F = 32.589; df = 4, 60; P < 0.0001). GSTs (F = 37.965; df = 4, 80; P < 0.0001), CarE (F = 40.861; df = 4, 60; P < 0.0001), and MFO (F = 33.416; df = 4, 80; P < 0.0001) activities were relatively higher in the first 24 h (Fig 2A and 2E), CarE activity was relatively higher at 24 and 48 h (Fig 2C) (F = 30.852; df = 4, 80; P = 0.001) and MFO activity was relatively higher at 24 h (Fig 2G) (F = 33.416; df = 4, 80; P < 0.0001). GSTs (F = 30.852; df = 4, 60; P < 0.0001), CarE (F = 67.249; df = 4, 60; P < 0.0001), and MFO (F = 42.589; df = 4, 60; P < 0.0001) activities were relatively lower in the first 24 h (Fig 2A and 2E).
Antifeedant Activity of G. biloba Secondary Metabolites against H. cunea Larvae as Assessed by the Leaf-Disc Assay

The leaf-disc assay results show that all tested G. biloba secondary metabolites have antifeedant activity against H. cunea larvae (Fig 3). The antifeedant activity of all four secondary metabolites showed concentration-dependent larval responses ($F = 22.986; \text{df} = 3, 48; \ P = 0.002$). At low concentrations, the antifeedant activities of all four secondary metabolites were not prominent. At medium and high concentrations, all four secondary metabolites showed strong antifeedant activity. The antifeedant efficiency in descending order was ginkgolide $>$ G. biloba extracts $>$ bilobalide $>$ ginkgo flavonoids ($F = 18.773; \text{df} = 3, 32; \ P = 0.003$).

Antifeedant Activity of G. biloba Secondary Metabolites against H. cunea Larvae in the Field

All four tested secondary metabolites significantly repelled H. cunea larvae compared with that of the vehicle control (Fig 4) ($F = 58.175; \text{df} = 4, 100; \ P < 0.0001$). In all larval instars, ginkgolide and G. biloba extracts showed the strongest repelling effects, whereas ginkgo flavonoids had the weakest. Third-instar larvae were the most sensitive to G. biloba secondary metabolites, whereas newly hatched and aged larvae showed relatively lower sensitivities to these agents ($F = 28.276; \text{df} = 4, 100; \ P = 0.001$).
Fig 2. Analysis of the enzyme activities of four detoxifying enzymes of Hyphantria cunea larvae fed artificial diets with Ginkgo biloba secondary metabolites. (A) The enzymatic activity of GSTs were measured in different feeding time. (B) The enzymatic activity of GSTs were measured in different larvae instar. (C) The enzymatic activity of CarE were measured in different feeding time. (D) The enzymatic activity of CarE were measured in different larvae instar. (E) The enzymatic activity of AChE were measured in different feeding time. (F) The enzymatic activity of AChE were measured in different larvae instar. (G) The enzymatic activity of MFO were measured in different feeding time. (H) The enzymatic activity of MFO were measured in different larvae instar. EGB: extract of ginkgo biloba; GF: gingko flavonoids; GL: ginkgolide; BB: bilobalide; CG: control group. Different letters above bars indicate significant differences (P < 0.05).

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Ginkgo biloba is a typical relic plant that rarely exhibits severe pest infestations, and reports about pest hazards to G. biloba leaves are rare. We tested the antifeedant activity of G. biloba extracts and isolated secondary metabolites, including ginkgo flavonoids, ginkgolide, and bilobalide, against H. cunea larvae. The results showed that all four preparations conferred significant antifeedant activity in leaf-disc assays and in artificial diets, and the activities of larval

**Discussion and Conclusions**

*Ginkgo biloba* is a typical relic plant that rarely exhibits severe pest infestations, and reports about pest hazards to *G. biloba* leaves are rare. We tested the antifeedant activity of *G. biloba* extracts and isolated secondary metabolites, including ginkgo flavonoids, ginkgolide, and bilobalide, against *H. cunea* larvae. The results showed that all four preparations conferred significant antifeedant activity in leaf-disc assays and in artificial diets, and the activities of larval
detoxifying enzymes significantly increased after feeding on diet containing these substances. These preparations also significantly repelled H. cunea larvae in field tests. Ginkgolide, which is unique to G. biloba, showed the strongest antifeedant activity, the strongest effect on detoxifying enzyme activity, and the strongest repelling effect in field tests. Studies have shown that ginkgolides can be detected in various tissues of G. biloba. Their contents ranged from high to low in the order leaf > root > stem [28]. No sign of insect feeding was observed on leaves of G. biloba, but fresh and past-year emergence holes of H. insularis Staudinger were observed on the stem, which may be due to differences in ginkgolide concentrations in these tissues. The ginkgolide used in this study was a mixture of four monomers, and the efficacy of the major lactone, bilobalide, alone also showed significant antifeedant efficacy toward H. cunea larvae. All of these monomers are secondary metabolites unique to G. biloba, which highlights the antifeedant activity of secondary metabolites produced by relic plants that co-evolved with insects [29].

MFO is one kind of oxidase system, and can carry on the oxidation of foreign substances in the insect body, GSTs can inactivate exogenous and endogenous toxin molecules and convert them into water-soluble compounds, CarE can convert compounds containing ester compounds into alcohol and acid [30–34]. Ginkgo biloba secondary metabolites affected the activities of glutathione transferase, carboxylesterase, acetylcholinesterase and mixed-functional oxidase activity, which is involved in the nervous system. These results suggest that the antifeedant activity of G. biloba secondary metabolites against H. cunea larvae may be multifactorial.

Under field conditions, G. biloba secondary metabolites do not kill the insects; rather, the strong antifeedant activity affects plant host selection by the insects. Larvae rapidly move away after spraying preparations containing G. biloba secondary metabolites on Salix branches damaged by H. cunea larvae. This result indicates the strong repelling effect conferred by these agents [35]. Although ginkgolide showed similar repelling effect as that of G. biloba extracts, it is more practical and economical to use preparations containing G. biloba extracts.

Plants can be classified into different families, genera, and species. After long-term evolution and specialization, insects feeding on certain categories of plants developed detoxification mechanisms to infest more effectively plants that produce these metabolites. Secondary substances produced by non-host plants show antifeedant effects on insects. G. biloba is a relic plant, with no parallel of the same family or genus, and plays host to a few insects. Its unique secondary metabolites show significant antifeedant and repelling effects, but they are not toxic to H. cunea larvae, so insects will not normally develop resistance to these metabolites after genetic selection. The unique secondary metabolites of G. biloba undergo rapid degradation after spraying in the wild, and thus have little impact on the environment. Beside G. biloba, some other plant species are unique to the family and genus; thus, the unique secondary metabolites of G. biloba could actually represent a large group of potential plant protection agents. Secondary metabolites of different relic plants can be mixed to widen the anti-insect spectrum, or alternatively, the genes encoding these secondary metabolites could be introduced into other plant to generate new resistant variants [36].

The long-term co-evolution of insects and plants promoted the development of plant secondary metabolites, which emerged to have key roles in chemical defense and insect resistance. G. biloba survived from the quaternary glaciation period at least partly owing to its strong pest resistance conferred by its unique secondary metabolites. G. biloba has a rich array of secondary metabolites that may accumulate and decline with changing seasons, so that the peak content of chemical agents largely overlaps with seasonal peaks in insect activity [28]. This study presented evidence for the antifeedant and repelling effects of G. biloba secondary metabolites. The results provide new insights into the function of unique G. biloba secondary metabolites, and a new understanding of why G. biloba is largely free of insect pests [37].
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Author Contributions
Conceived and designed the experiments: YL LP. Performed the experiments: LP. Analyzed the data: LP LR. Contributed reagents/materials/analysis tools: LP YF. Wrote the paper: LP LR FC.

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