Independent Recruitment of a Flavin-Dependent Monooxygenase for Safe Accumulation of Sequestered Pyrrolizidine Alkaloids in Grasshoppers and Moths

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Abstract

Several insect lineages have developed diverse strategies to sequester toxic pyrrolizidine alkaloids from food-plants for their own defense. Here, we show that in two highly divergent insect taxa, the hemimetabolous grasshoppers and the holometabolous butterflies, an almost identical strategy evolved independently for safe accumulation of pyrrolizidine alkaloids. This strategy involves a pyrrolizidine alkaloid N-oxidease that transfers the pyrrolizidine alkaloids to their respective N-oxide, enabling the insects to avoid high concentrations of toxic pyrrolizidine alkaloids in the hemolymph. We have identified a pyrrolizidine alkaloid N-oxidease, which is a flavin-dependent monooxygenase, of the grasshopper Zonocerus variegatus. After heterologous expression in E. coli, this enzyme shows high specificity for pyrrolizidine alkaloids of various structural types and for the tropane alkaloid atropine as substrates, a property that has been described previously for a pyrrolizidine alkaloid N-oxidease of the arctiid moth Grammia geneura. Phylogenetic analyses of insect flavin-dependent monooxygenase sequences suggest that independent gene duplication events preceded the establishment of this specific enzyme in the lineages of the grasshoppers and of arctiid moths. Two further flavin-dependent monooxygenase sequences have been identified from Z. variegatus sharing amino acid identities of approximately 78% to the pyrrolizidine alkaloid N-oxidease. After heterologous expression, both enzymes are also able to catalyze the N-oxidation of pyrrolizidine alkaloids, albeit with a 400-fold lower specific activity. With respect to the high sequence identity between the three Z. variegatus sequences this ability to N-oxidize pyrrolizidine alkaloids is interpreted as a relict of a former bifunctional ancestor gene of which one of the gene copies optimized this activity for the specific adaptation to pyrrolizidine alkaldoid containing food plants.

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Introduction

Chemical defense against herbivory is essential for plants to be able to survive in their natural habitat. During evolution, many insect herbivores have developed counterstrategies to cope with these toxic compounds. In some cases, they have even acquired these chemicals for their own benefit. One of the best studied examples of plant toxins sequestered by adapted insects are the pyrrolizidine alkaloids. After ingestion by a vertebrate or insect herbivore, the N-oxides are easily reduced to the protoxic free base, the substrate for cytochrome P450-mediated bioactivation [2,3].

Strategies for PA sequestration in adapted insects have evolved in various insect lineages under the identical selection pressure to avoid higher concentrations of PAs in the form of their free base in the hemolymph [recent reviews see [1,3,4,5]]. Leaf beetles of the genus Platypatra (Chrysomelidae, Coleoptera) have developed a strategy to transfer the free base from the hemolymph into defense secretions with such an efficiency that all other tissues outside the secretory glands are almost devoid of PAs [6]. In the related leaf beetle genus Oreina (Chrysomelidae, Coleoptera), the reduction of ingested PAs is suppressed. Instead, the N-oxides are directly absorbed and accumulated in their hemolymph and defense glands [7,8]. A third strategy to handle sequestered PAs is the stabilization of the PAs by enzyme-catalyzed N-oxidation of the alkaloids within the insect. This mechanism is realized in larvae of the tiger moth family (Arctiidae, Lepidoptera) [9–13], in several Longitarsus flea beetle species [14,15], and in the grasshopper genus Zonocerus [13,16]. Recently, we have been able to show that the enzyme responsible for the N-oxidation of PAs in arctiids belongs to the family of flavin-dependent monooxygenases (FMOs) [12,13] and was recruited by gene duplication early in this lineage during the adaptation to PA-containing plants [17]. To date, these PA N-oxigenases are the only functional characterized FMOs of insects.

FMOs are well characterized in vertebrates, where they are involved in the detoxification of nucleophilic nitrogen- and sulfur-
containing xenobiotics [18]. Although insects have to cope with a wide variety of xenobiotics, none of these so-called microsomal multi-substrate FMOs has been detected in insects, suggesting that the xenobiotic metabolism is guaranteed by cytochrome P450 monooxygenases (CYPs). Despite the different mechanism by which these two classes of monooxygenases operate, both convert lipophilic compounds into more hydrophilic metabolites that are readily excreted and have reduced bioactivity [3,18]. Both, FMOs and CYPs have been shown to catalyze the N-oxygenation of PAs in vertebrates [19,20].

Here, we report that PA N-oxygenation as one strategy for insect adaptation to PA-containing food plants evolved independently in the grasshopper *Z. variegatus* and the arctiid moths. We show that convergent evolution resulted in two almost identical systems. This identity refers to the findings that, in both cases, an FMO has been recruited, and that the respective enzymes show almost identical substrate specificity.

**Materials and Methods**

**Collection and rearing of Zonocerus**

Larvae and adults of *Zonocerus variegatus* were collected in the vicinity of Accra, Ghana, and reared in the laboratory under a light/dark regime of 16/8 h at 20 to 25°C. Egg pots were incubated in moist sand at a constant temperature of 30°C. Nymphs and adults were fed with leaves of *Rubus fruticosus* unless otherwise stated. Leaves of *R. fruticosus* are regarded as non-toxic as they do not contain PAs.

RNA isolation and cDNA synthesis

Adult insects were dissected, and fat body tissue was frozen in liquid nitrogen before total RNA was isolated by using TRIZOL® reagent (Invitrogen); 2 µg RNA was used for cDNA synthesis with Superscript III reverse transcriptase (Invitrogen) and an oligo(dT)17 primer (P1, 0.1 µM; Table S1) at 55°C in a total volume of 20 µl.

Identification and expression of FMO-like cDNA sequences

Degenerate primers P2 and P3 (for primer sequences see Table S1) were designed based on an alignment of FMO sequences of the lepidopteran species *Arctia caja, A. villica, Tyria jacobaeae*, and *Grammia geneura* (Syn.: *G. incorrupta*) in combination with two FMO-like sequences identified in each of the genomes of *Anopheles gambiae* and *Drosophila melanogaster*. In a semi-nested PCR approach, 3 µl of *Z. variegatus* cDNA was amplified with AccuTaq LA polymerase (Sigma) and primer pair P2/P1 by using a touch-down protocol with decreasing annealing temperatures from 60°C to 52°C (−0.5°C per cycle for 16 cycles and 52°C constant for 20 further cycles). The reaction was diluted 1:100, and 3 µl was used as template for the second PCR with primer pair P2/P3 and a decreasing annealing temperature from 55°C to 47°C in a total volume of 25 µl. The resulting fragment of 473 bp was subcloned by using the pGEM-T Easy vector (Promega). Sequencing revealed similarity to the FMO-encoding sequences of insects. Gene-specific primers were designed and used for 3'- and 5'-rapid amplification of cDNA ends (RACE, P4–P7) as described previously [21]. 3'-RACE resulted in the identification of two sequences covering the 3'-cDNA end and sharing 86% sequence identity. One of these sequences overlapped with the fragment identified with the degenerate primers and with the 5'-RACE fragment and was assembled to the full-length sequence (ZvFMOa). According to the second 3'-end cDNA, gene-specific
For LC-MS analysis, samples were diluted by using 50% and identities of senecionine were used, respectively, at an annealing temperature of 55°C with Platinum HPLC DNA polymerase (Invitrogen), which possesses proof-reading activity. The resulting fragments were digested with NdeI/Xhol and cloned into an NdeI/Xhol-linearized pET22b vector for heterologous expression with a C-terminal hexahistidine tag in E. coli BL21(DE3). Restriction of the fragment resulting from amplification with primer pair P13/P14 indicated that the PCR product was not homogeneous, as a fragment was detected that contained an internal, non-predicted restriction site for XhoI. Cloning of this fragment into the pGEM-T easy vector, screening with Xhol, and sequencing resulted in the identification of another FMO-encoding sequence, termed FMO2. For this sequence, the 3’- and 5’-cDNA ends were identified by RACE with the primers P15–P19. The full ORF of FMO2 was amplified with primer pair P20/P21 at 55°C with Platinum HPLC DNA polymerase, digested with BsaI and NotI, and cloned into an Neo/NotI-linearized pET28a vector for expression with a C-terminal hexahistidine tag in E. coli BL21(DE3). Expression of the recombinant proteins FMO3, FMO4, and FMO2 was induced with 0.1 mM isopropyl β-D-thiogalactoside at 30°C, before the proteins were purified by metal chelate affinity chromatography by using Ni2+-nitrilotriacetic acid agarose (Qagen). For determination of the native enzyme was applied to a Superdex200HighLoad column (GE Healthcare) using 20 mM glycine/NaOH buffer pH 9.0 containing 200 mM NaCl. As reference proteins thyroglobulin (669 kDa), ferritin (440 kDa), bovine serum albumin (67 kDa), ovalbumin (43 kDa), and chymotrypsinogen A (25 kDa) have been used.

Sequence analysis

Comparison of amino acid sequences was performed with the Bestfit software of the Wisconsin Sequence Analysis Package (version10, Genetics Computer Group, Madison, WI). An alignment of amino acid sequences [Figure S1] was generated with ClustalX [22] and used to estimate phylogenies with the PHYLIP program package [23]. The maximum likelihood tree was calculated with PROML by using the Jones-Taylor-Thornton model for amino acid changes [24]. Bootstrap values were estimated with the programs SEQBOOT and CONSENSE. The bootstrap estimates are the result of 1000 replicates. The sequences reported in this paper have been submitted to the EMBL Nucleotide Sequence Database with the accession numbers FR696371 (FMO3), FR696372 (FMO4), and FR696373 (FMO2).

Assay for PA N-oxygenase activity

Enzyme activity was assayed in a 100 mM glycine/NaOH buffer pH 9 containing 0.2 mM NADPH at 37°C photometrically or in a radio assay as described previously [12]. All substrates were assayed at a concentration of 0.2 mM by using the photometric assay or the radio assay (0.023 μCi assay), except for retronecine, seneconine, and isoseneconine, which were tested as 3H-labeled substrates, and phalaenopsine as a 14C-labeled substrate in the radio assay (0.023 and 0.045 μCi assay, respectively). For enzyme kinetics, initial rates were determined by varying the concentration of alkaloid (1–40 μM) for seneconine, 4–60 μM for monocrotaline and atropine, and 0.1–1.5 mM for heliotrine and phalaenopsine) while maintaining the concentration of NADPH at 200 μM. The identity of seneconine N-oxide, seneconine N-oxide, monocrotaline N-oxide, phalaenopsine N-oxide, and atropine N-oxide was confirmed by liquid chromatography mass spectrometry (LC-MS). For LC-MS analysis, samples were diluted by using 50% acetonitrile containing 1% formic acid. Solutions were infused directly into a 3200 Q Trap MS instrument (Applied Biosystems/MD Sciex). The mass spectrometer was equipped with an electrospray ionization interface (ESI, Turbo V) and was operated in positive-ion mode and enhanced product ion (EPI) scan mode. Ionization and EPI conditions were optimized individually for each compound in its non-oxidized form, and the obtained settings were used to analyze extracts of the corresponding enzymatic assays. Nitrogen was used as a curtain and auxiliary gas.

Results

Identification and Heterologous Expression of PA N-Oxygenase of Z. variegatus

In contrast to the arctic moths in which the PA N-oxygenase is a soluble protein in the hemolymph, the arctic N-oxygenating activity in the grasshopper Z. variegatus is associated with the fat body [13]. Separate incubation of a particulate fraction and the supernatant obtained from the insects fat body with 14C-seneconylline showed that N-oxygenation activity is detectable in the soluble fraction, but not in the pellet. These results indicate that PNO is expressed as a soluble protein, excluding the involvement of a membrane-bound CYP. Therefore, we hypothesized that the enzyme might belong to the FMOs in this species. Using cDNA preparations of fat body tissue of Z. variegatus, we applied a polymerase chain reaction (PCR)-based approach, in order to identify FMO-encoding cDNA sequences. Degenerate primers were designed according to an alignment of FMO sequences of Drosophila melanogaster and Anopheles gambiae present in the database and of four arctic species identified recently in our group, viz., Actia caja, A. sillica, Tyria jacobaeae, and Grammia geneura [17]. Three cDNA sequences were identified, FMOa, FMOc, and FMO2, with open reading frames of 1242 bp, 1245 bp, and 1242 bp in length, encoding proteins with a subunit size of 47,726 Da, 47,939 Da, and 47,793 Da, respectively. The isoelectric point of the three proteins is predicted to be 6.6, 6.2, and 6.0, respectively. The sequence motifs most characteristic for FMO, viz., the nucleotide-binding sites (Rossman folds, consensus GxGxxG) for binding of FAD as a prosthetic group and of the NADPH cofactor, respectively, and the FMO-identifying sequence (consensus FxxGxxxF) are highly conserved in sequence position in comparison with the FMOs of other eukaryotes [25–28] (Figure S1). For one of the three sequences an N-terminal signal peptide was predicted as it was shown to be present in the SNO of Tyria jacobaeae and the FMOs of other arctic species [12,17]. No transmembrane helices were detected as they are present in mammalian FMOs close to the C-terminus for membrane attachment [29]. The FMO-like sequences of Zv-FMOa, Zv-FMOc, and Zv-FMO2 showed, in assays for PA N-oxygenase of T. jacobaeae (Diptera), and to the human FMO1 (Table 1).

Biochemical Characterization of PA N-Oxygenase of Z. variegatus

After the expression and purification by metal chelate affinity chromatography, FMOa, FMOc, and FMO2 showed, in assays for PA N-oxygenation, specific activities of 123 pkat/mg, 146 pkat/mg, and 59 pkat/mg, respectively (100 mM glycine/NaOH buffer pH 9.0 with 0.2 mM seneconine as substrate). The identity of the reaction product as seneconine N-oxide was confirmed by LC-MS (liquid chromatography with mass spectrometry). Therefore, Zv-FMO2 was designated as PA N-oxygenase protein.
activity with most of the substrates that show activity with atropine. The apparent K selected substrates representing the major classes of PAs and of cysteamine is accepted by the PNO of Zonocerus Zv substrates. Cysteine, dimethylaniline, hydroxylamine, or glutathione are not mammalian FMOs and for the FMO from senecionine and seneciphylline. Of the substrates described for almost the same activity as the best substrates tested, e.g., certain solaneceous plants, is converted to its substrate (Figure 1). Atropine, a tropane alkaloid common to carries a methyl group at the ring-bound nitrogen, is not a With regard to structures of PAs, other alkaloids and compounds known to be types are 11.9, 9.8, and encountering a huge diversity of plant allelochemicals [30].

Table 1. Degree of identity between amino acid sequences encoding FMOs of Z. variegatus, other insects, and the FMO1 of human.

| (%)   | ZvFMOa | ZvFMOc | ZvPNO | TjSNO | DmFMO3006 | DmFMO3174 | HsFMO1 |
|-------|--------|--------|-------|-------|-----------|-----------|-------|
| ZvFMOa| 100    | 77.0   | 77.8  | 35.2  | 36.4      | 36.3      | 39.8  |
| ZvFMOc|        | 77.0   | 35.3  | 38.2  | 36.6      | 38.8      |       |
| ZvPNO |        |        | 36.3  | 38.2  | 37.1      | 37.7      |       |

For comparison, the full coding regions were used. TjSNO, Senecine N-oxygenase of Tyria jacobaeae; DmFMO3006 and DmFMO3174, FMO sequences of unknown function present in the genome of Drosophila melanogaster; HsFMO1, FMO1 of human.

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Table 2. Substrate specificity of recombinant flavin-dependent monoxygenases of Z. variegatus.

| Substrate     | Specific activity (nkat/mg) | G3PNO | ZvPNO | ZvFMOa | ZvFMOc |
|---------------|----------------------------|-------|-------|--------|--------|
| Pyrrolizidine alkaloids |                           |       |       |        |        |
| Senecine type |                           |       |       |        |        |
| Senecionine   | 55.0                      | 32.0  | 0.1   | 0.1    |        |
| Seneciphylline| 53.2                      | 29.4  | 0.6   | 0.1    |        |
| Senecivernine | -                         | 24.0  | 1.3   | 0.6    |        |
| Retrosine     | -                         | 18.6  | 1.5   | 0.5    |        |
| Senkirkine    | n.d.                      | n.d.  | n.d.  | n.d.   |        |
| Monocrotaline type |                         |       |       |        |        |
| Monocrotaline | 90.2                      | 23.4  | 1.5   | 0.9    |        |
| Axillarline   | 62.2                      | 18.6  | n.d.  | n.d.   |        |
| Axillaridine  | -                         | 11.8  | n.d.  | n.d.   |        |
| Lycopasine type |                         |       |       |        |        |
| Heliotrine    | 44.2                      | 19.5  | n.d.  | n.d.   |        |
| Rinderine     | 38.8                      | 11.5  | 1.3   | 0.4    |        |
| Indicine      | -                         | 4.8   | n.d.  | n.d.   |        |
| Triangularine type |                     |       |       |        |        |
| Sarracine     | 16.0                      | 0.4   | n.d.  | n.d.   |        |
| Phalenopsine type |                     |       |       |        |        |
| Phalenopsine  | 17.1                      | 9.6   | n.d.  | n.d.   |        |
| Necine bases  |                           |       |       |        |        |
| Retronecine   | n.d.                      | 10.2  | n.d.  | 0.1    |        |
| Heliotidine   | -                         | n.d.  | n.d.  | n.d.   |        |
| Supinidine    | n.d.                      | n.d.  | n.d.  | n.d.   |        |
| Other alkaloids |                         |       |       |        |        |
| Ephedrine     | -                         | n.d.  | n.d.  | n.d.   |        |
| Nicotine      | n.d.                      | 2.3   | 0.3   | 0.4    |        |
| Atropine      | 18.9                      | 29.1  | 0.3   | n.d.   |        |
| Other substrates |                         |       |       |        |        |
| Dimethylaniline | n.d.                     | n.d.  | n.d.  | n.d.   |        |
| Cysteamine    | n.d.                      | 7.0   | 0.5   | 0.1    |        |
| L-Cysteine    | n.d.                      | n.d.  | n.d.  | n.d.   |        |
| Hydroxylamine | -                         | n.d.  | n.d.  | n.d.   |        |
| Glutathione   | 37.0                      | n.d.  | n.d.  | n.d.   |        |

n.d., not detectable; -, not tested.

For comparison, the full coding regions were used. TjSNO, Senecine N-oxygenase of Tyria jacobaeae; DmFMO3006 and DmFMO3174, FMO sequences of unknown function present in the genome of Drosophila melanogaster; HsFMO1, FMO1 of human.

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(PNO). Recombinant ZvPNO shows a pH optimum of 9.0 in potassium phosphate buffer (10 mM), Tris/HCl buffer (10 mM), and in glycine/NaOH buffer (100 mM), respectively (Figure S2). Using the potassium phosphate-based buffer system (pH9.0) in the standard assay, linear rates of product formation were maximal at a temperature of 42°C. Using a temperature range of 20–42°C, the activation energy was estimated to be 70.7 kJ/mol. The molecular mass of the heterologously expressed protein has been estimated by gel filtration chromatography to be 91,200 Da, indicating that the active enzyme is a homodimer.

Substrate Specificity of Recombinant FMO of Z. variegatus

Z. variegatus is a generalist feeding on a wide array of food plants and encountering a huge diversity of plant allelochemicals [30]. Therefore, in testing the substrate specificity of recombinant ZvFMOa, ZvFMOc, and ZvPNO, we used, in addition to various structures of PAs, other alkaloids and compounds known to be accepted as substrates of mammalian and yeast FMO (Table 2). With regard to ZvPNO, PA structures of the five main structural types are N-oxygenized. The otonecin derivative senkirkine, which carries a methyl group at the ring-bound nitrogen, is not a substrate (Figure 1). Atropine, a tropane alkaloid common to certain solaneceous plants, is converted to its N-oxide form with almost the same activity as the best substrates tested, e.g., senecione and seneciphylline. Of the substrates described for mammalian FMOs and for the FMO from Scaurachyges ceresiter, cysteamine is accepted by the PNO of Zonocerus, whereas L-cysteine, dimethylaniline, hydroxylamine, or glutathione are not substrates. ZvFMOa and ZvFMOc show low but unequivocal activity with most of the substrates that show activity with ZvPNO.

For ZvPNO, enzyme kinetics have been established by using selected substrates representing the major classes of PAs and of atropine. The apparent K values of 1.1 μM (senecione), 11.9 μM (monocrotaline), 263.9 (heliotrine), 837.6 μM (phalaenopsine), and 9.8 μM (atropine) show that the PNO has a higher affinity to the macrocyclic diesters senecione and monocrotaline than to the monesters heliotrine and phalaenopsine. The affinity to the tropane alkaloid atropine is in the same range as that for the macrocyclic diester type. The values for the catalytic efficiency (kcat/Km) support the interpretation that senecione is the most efficient substrate for the PNO followed by monocrotaline and atropine (Table 3).

Comparative Sequence Analysis of Insect FMOs

The amino acid sequences of the three FMOs of Z. variegatus and of various FMOs of insects have been used for phylogenetic analysis (Figure 2). The sequences form well-supported individual clusters with regard to the sequences of the Diptera (D. melanogaster and ZvPNO, enzyme kinetics have been established by using selected substrates representing the major classes of PAs and of atropine. The apparent K values of 1.1 μM (senecione), 11.9 μM (monocrotaline), 263.9 (heliotrine), 837.6 μM (phalaenopsine), and 9.8 μM (atropine) show that the PNO has a higher affinity to the macrocyclic diesters senecione and monocrotaline than to the monesters heliotrine and phalaenopsine. The affinity to the tropane alkaloid atropine is in the same range as that for the macrocyclic diester type. The values for the catalytic efficiency (kcat/Km) support the interpretation that senecione is the most efficient substrate for the PNO followed by monocrotaline and atropine (Table 3).

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and *A. gambiae* and of *Z. variegatus*, two clusters for the sequences of *Tribolium castaneum*, and three separate clusters for lepidopteran sequences, recently termed FMO1 to FMO3 [17]. Within the Lepidoptera, gene duplication events have been shown to be responsible for the separation of the related FMO1 and FMO2 cluster and, within the Arctiids, for two separated clusters, of which one encodes the PA \( \text{N} \)-oxygenases [17]. The tree topology clearly supports two independent origins of PA \( \text{N} \)-oxygenizing enzymes in insects, one early in the arctiid lineage and one in grasshoppers.

### Discussion

Our studies show that, in Arctiids and in the grasshopper *Z. variegatus*, FMOs were recruited independently as \( \text{N} \)-oxygenases for the safe handling of plant-derived PAs in these insects. Therefore, PA \( \text{N} \)-oxygenation in insects seems to be a unique feature of FMOs. Of note, CYPs in insects form a far larger gene family than FMOs (the genomes of *D. melanogaster* and *B. mori* encode 83 and 86 putative CYPs, respectively, in contrast to only 2 and 3 putative FMOs, respectively [12,17,31,32]) and are well-known for their importance in the metabolism of natural and artificial xenobiotics, including insecticides [33]. Specific and inducible CYP-encoding genes in *Papilio* butterflies (Papilionidae, Lepidoptera) for the detoxification of furanocoumarins represent only one of several fascinating examples of gene evolution during adaptation of insects to plant allelochemicals [34,35]. The array of substrates of the respective enzymes has been shown to be correlated with the feeding habits of the insects, *i.e.*, a broad substrate specificity in generalists that feed on a wide diversity of host plants containing a broad spectrum of allelochemicals, and a narrow substrate specificity in specialist insects that feed on only one or a few plant species [36]. The same has been shown for FMOs involved in PA \( \text{N} \)-oxygenation of a specialist and a generalist arctiid species [17]. Concretely, the respective enzyme of the specialist *T. jacobaeae* is highly specific for PAs of the toxic senecionine type, the

### Table 3. Enzyme kinetics of recombinant ZvPNO.

| Substrate          | \( K_m \) (\( \mu \)M) | \( V_{\text{max}} \) (nkat/mg) | \( k_{\text{cat}} \) (s\(^{-1}\)) | \( k_{\text{cat}}/K_m \) (s\(^{-1}\) mol\(^{-1}\) l) |
|--------------------|-------------------------|-----------------------------|----------------------|----------------------------------|
| Senecionine        | 1.08 ± 0.07             | 20.73 ± 0.24                | 0.99                 | 916.666                          |
| Monocrotaline      | 11.93 ± 0.76            | 19.99 ± 0.40                | 0.96                 | 80.469                           |
| Heliotrine         | 263.89 ± 10.06          | 2.49 ± 0.03                 | 0.12                 | 454                              |
| Phalaenopsine      | 837.61 ± 77.37          | 2.66 ± 0.11                 | 0.13                 | 155                              |
| Atropine           | 9.83 ± 1.04             | 7.68 ± 0.23                 | 0.37                 | 37.640                           |

For calculations a molecular mass of ZvPNO of 47793 g/mol was used. doi:10.1371/journal.pone.0031796.t003

### Figure 2. Unrooted maximum-likelihood tree of amino acid sequences derived from cDNA encoding FMOs of various insect species.

Framed sequences were heterologously expressed and functionally analyzed. The other sequences should be regarded as putative FMO-coding cDNA. Branch lengths are proportional to the number of amino acid substitutions per site (scale: 0.1 substitutions per site). Bootstrap proportions resulted from 1000 replicates and are given for values >50. Ac, *Arctia caja*; Ag, *Anopheles gambiae*; Bm, *Bombyx mori*; Dm, *Drosophila melanogaster*; Gg, *Grammia geneura*; Ha, *Helicoverpa armigera*; Tc, *Tribolium castaneum*; Tj, *Tyria jacobaeae*; Zv, *Zonocerus variegatus*. Accession numbers for all sequences are listed in Figure S1. doi:10.1371/journal.pone.0031796.g002
predominant PA type of its almost exclusive food plant Jacobaea vulgaris (syn. Senecio jacobanus). In contrast, the N-oxygenase of the generalist G. geneura (Syn.: G. incornuta) accepts a wider range of substrates, including the non-toxic 1,2-saturated PA phalaenoepine and the tropaeoline alkaloid atropine [17]. Most interestingly, the substrate specificity of the PA N-oxygenase of G. geneura is almost identical to that of the PNO of Z. variegatus that also accepts atropine as a substrate. We postulate that the almost identical substrate specificity of the PA N-oxygenases of the two unrelated insects is the result of convergent evolution under identical selection pressure of generalist feeding. The wide substrate specificity of the PA N-oxygenase of Z. variegatus correlates with the extremely polyphagous behavior of this grasshopper, which is ranked as one of the most important economically pests of agriculture in west and central Africa [37,38].

Of note, the N-oxygenation of PAs in specialized insects is not a detoxification mechanism that converts xenobiotics into more polar metabolites for efficient excretion from the body. Instead, this enzymatic conversion allows the sequestration of these plant toxins in a deactivated, metabolically safe form [5]. The recruitment of an enzyme that catalyzes PA N-oxygenation can be regarded as a “key innovation” during the insect’s adaptation to PA-containing food plants [17]. In arctiids, this enzyme was the prerequisite for the evolution of several further, highly specific adaptations, such as the positive feeding response triggered by traces of PAs sensed by specialized taste receptors, the ability to transfer the alkaloids from the larvae to the adult stage, and the insect’s behavior that ensures optimal egg protection by PAs of both parents governed by PA-derived pheromones [5,39].

The high degree of sequence identity of the PA N-oxygenase to two further FMOs identified from Z. variegatus, and the fact that the latter two enzymes are also able to catalyze the N-oxygenation of alkaloids, albeit with a much lower specific activity, indicate the close relationship of these sequences. The data suggest a duplication of an ancestor gene encoding an FMO of unknown function that already possessed the ability to N-oxygenize PAs, most probably as a side activity. One of the gene copies was recruited and optimized for N-oxygenation of plant-derived PAs. The finding that PAs are strong phagostimulants for Zonocerus variegatus [40] suggests a central role of PAs in the insect’s ecology. Further research has to show whether this is also the case for the N-oxygenation of further plant-derived toxins, such as atropine or nicotine, that are accepted as a substrate by the PNO of Z. variegatus.

Supporting Information

Figure S1 Amino acid alignment of flavin-dependent monoxygenases of various insect species. Only the central part of the alignment that was used for the estimation of the phylogenies is shown, spanning the region from amino acid position 5 to 402 with respect to ZPNO. The sequence motifs for binding of FAD and NADPH and the FMO-identifying sequence are boxed. The accession numbers of the three FMO sequences of Zonocerus variegatus and of all sequences taken from the databases are given at the end of the alignment. (PDF)

Figure S2 PNO activity in three buffer systems with pH variation. Enzyme activities have been determined with senecionine as substrate at 37°C. (PDF)

Table S1 Sequences of primers used for the identification and cloning of cDNAs of flavin-dependent monoxygenases of Zonocerus variegatus. Recognition sites of restriction endonucleases used for cloning are underlined. (DOC)

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Author Contributions

Conceived and designed the experiments: LW DO JT. Performed the experiments: LW TB. Analyzed the data: LW TB DO. Contributed reagents/materials/analysis tools: JT TB. Wrote the paper: DO. LC-MS experiments: TB.

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