Pore-forming protein complexes from *Pleurotus* mushrooms kill western corn rootworm and Colorado potato beetle through targeting membrane ceramide phosphoethanolamine

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Aegerolysins ostreolysin A (OlyA) and pleurotolysin A (PlyA), and pleurotolysin B (PlyB) with the membrane-attack-complex/perforin domain are proteins from the mushroom genus *Pleurotus*. Upon binding to sphingomyelin/cholesterol-enriched membranes, OlyA and PlyA can recruit PlyB to form multimeric bi-component transmembrane pores. Recently, *Pleurotus* aegerolysins OlyA, PlyA2 and erylysin A (EryA) were demonstrated to preferentially bind to artificial lipid membranes containing 50 mol% ceramide phosphoethanolamine (CPE), the main sphingolipid in invertebrate cell membranes. In this study, we demonstrate that OlyA6, PlyA2 and EryA bind to insect cells and to artificial lipid membranes with physiologically relevant CPE concentrations. Moreover, these aegerolysins permeabilize these membranes when combined with PlyB. These aegerolysin/PlyB complexes show selective toxicity toward western corn rootworm larvae and adults and Colorado potato beetle larvae. These data strongly suggest that these aegerolysin/PlyB complexes recognize CPE as their receptor molecule in the insect midgut. This mode of binding is different from those described for similar aegerolysin-based bacterial complexes, or other *Bacillus thuringiensis* Cry toxins, which have protein receptors. Targeting of *Pleurotus* aegerolysins to CPE and formation of transmembrane pores in concert with PlyB suggest the use of aegerolysin/PlyB complexes as novel biopesticides for the control of western corn rootworm and Colorado potato beetle.

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Western corn rootworm (WCR; *Diabrotica v. virgifera* LeConte; Coleoptera, Chrysomelidae) is an important pest on maize in the USA1-2 and Europe3, and it was reported to cause annual economic losses of over 1 billion dollars in the USA4-5. In Europe, the control of WCR by crop rotation3,6, biological control options7,8, and host-plant native resistance and tolerance9 are being evaluated, although these approaches have not achieved widespread success in North America10. In addition to crop rotation, chemical control mainly focuses on the larval stages of WCR6. In 2003, the Environmental Protection Agency approved the first commercial use of transgenic corn hybrids that express proteinaceous crystal toxins (i.e., Cry toxins) from *Bacillus thuringiensis* (e.g., Bt maize) against WCR larvae11-13. This group of toxins comprises approximately 300 proteins that have been divided into 75 subgroups. Cry toxins are species specific, and are toxic due to their binding to species-specific surface proteins
in the microvilli of the larva midgut cells (e.g., cadherins, aminopeptidases, alkaline phosphatases)\(^1,1^2\). Since the discovery of these toxins and their registration as pesticides in the USA in 1961, \(B.\) \textit{thuringiensis} has been the most successful pathogen for the control of WCR, and now commands \(-2\%\) of the total insecticide market\(^1^3\). However, WCR can continuously develop resistance to these toxins through different mechanisms\(^1^4,1^6\).

The Colorado potato beetle (CPB; \textit{Leptinotarsa decemlineata} [Say]; Coleoptera, Chrysomelidae) has been driving the modern insecticide industry since the early days of its spread\(^1^7\). Neonicotinoid insecticides and some endotoxins from \(B.\) \textit{thuringiensis} subsp. \textit{tenetebionis} are generally used for CPB control. However, as for WCR, CPB can continuously develop resistance against these biopesticides through various mechanisms\(^1^8,1^9\).

The search for alternative biopesticides and approaches is therefore of extreme importance, such as the development of efficient attract-and-kill strategies. These efforts very recently resulted in the discovery of novel proteinaceous toxins that are specific for WCR and some other coleoptera. These have been isolated from different bacterial species to \(B.\) \textit{thuringiensis}, and include IPD072a from \textit{Pseudomonas chlororaphis}\(^2^0\), PiP-47a from \textit{Pseudomonas mosselli}\(^2^1\), GNIP1Aa from \textit{Chromobacterium piscinae}\(^2^2\), and AflP-1A/AflP-1b from \textit{Alcaligenes faecalis}\(^2^3\). The \textit{Pseudomonas}-derived toxins IPD072a and PiP-47a do not match any other protein amino-acid sequences currently in databases, while the N-terminal region of GNIP1Aa from \textit{C. piscinae} was assigned to the membrane-attack-complex/perforin (MACPF) protein superfamily. In contrast to the aforementioned proteins that are toxic toward WCR in their monomeric forms, AflP-1A and AflP-1b from \textit{A. faecalis} act as a bi-component toxic complex, in which the AflP-1A partner belongs to the aegerolysin protein family.

The aegerolysins (Pfam 06355; InterPro IPR009413) currently comprise over 350 small (~15–20 kDa), \(\beta\)-structured proteins that are found in several eukaryotic and bacterial taxa\(^2^4,2^5\). The common prominent feature of these proteins is their interactions with specific membrane lipids and lipid domains. Aegerolysins from the fungal genus \textit{Pleurotus} have been shown to interact with sphingomyelin/cholesterol domains in artificial and biological membranes\(^2^7,2^8\). These aegerolysins, namely ostreolysin A (OlyA) from \textit{P. ostreatus} and pleurotolysin A2 (PlyA2) and erylisin A (EryA) from \textit{P. eryngii}, were recently found to interact even more strongly with lipid vesicles that contained equimolar levels of cholesterol and ceramide phosphateethanolamine (CPE)\(^2^9\). CPE is the major sphingolipid in invertebrate cell membranes and is an analog of sphingomyelin, which dominates in vertebrates\(^3^0,3^1,3^2\). The dissociation constants of OlyA and PlyA2 binding to equimolar CPE/cholesterol and sphingomyelin/cholesterol lipid vesicles were reported to be \(10^{-8}\) M and \(>10^{-8}\) M, respectively\(^3^3,3^7\), which suggests that the interactions of these aegerolysins with CPE-containing membrane systems is at least 1000-fold stronger. This discovery led to the use of \textit{Pleurotus} aegerolysins as useful molecular markers of CPE distribution in insect tissues, and for detection of the bloodstream form of \textit{Trypanosoma brucei}\(^3^5\). Furthermore, OlyA, OlyA6, and PlyA were shown to combine with pleurotolysin B (PlyB), a 59-kDa protein partner with a MACPF domain that is produced by \textit{P. ostreatus}\(^2^7,3^0,3^2\), while EryA acts in concert with another MACPF-protein, erlysin B (EryB), which shares 96% amino acid sequence identity with PlyB\(^3^6\). OlyA/PlyB, OlyA6/PlyB and PlyA/PlyB combinations can then act as bi-component pore-forming complexes for artificial and biological cell membranes that contain sphingomyelin/cholesterol domains\(^3^7,3^0,3^1,3^2,4^0\). The membrane-disrupting potential of these aegerolysin/PlyB complexes has, however, not been tested on CPE-containing membranes.

The binary and quaternary cytolytic complexes of bacterial origin where aegerolysin-like proteins are combined with larger, non-aegerolysin protein partner(s) that have been described to date comprise: Cry16Aa/Cry17Aa/Cbm17.1/Cbm17.2 from \textit{Clostridium bifermantans} subsp. \textit{Malaysia}\(^4^1\), Cry34Ab1/Cry35Ab1 from \textit{B. thuringiensis}\(^4^2,4^3\), and AflP-1A/AflP-1b from \textit{A. faecalis}\(^4^4\). These heteromeric aegerolysin-based cytolytic complexes have been exploited as potent biopesticides for specific pests, whereby Cry16Aa/Cry17Aa/Cbm17.1/Cbm17.2 acts against \textit{Aedes} mosquitoes, and Cry34Ab1/Cry35Ab1 and AflP-1A/AflP-1b act against coleoptera species, and mainly WCR. Of the numerous Cry toxins produced by \textit{B. thuringiensis}, Cry34Ab1 is the only aegerolysin-like protein\(^1^1\). Cry34Ab1/Cry35Ab1 were genetically introduced into corn hybrids in 2005 (i.e., Bt maize) as tools for the control of WCR larvae\(^6^4,6^5\). Although no specific Cry34Ab1/Cry35Ab1 receptors have been identified to date, the data obtained so far suggest that they interact with (glyco)protein receptors in the membranes of epithelial cells in the insect midgut, with the resultant damage due to their perforation of these cells\(^6^6,6^7\). WCR larvae were recently reported to evolve resistance to the Bt maize that produces Cry34Ab1/Cry35Ab1\(^1^4\). Recently, it was also shown that laboratory-generated WCR colonies selected for resistance to Cry34Ab1/Cry35Ab1 can feed on AflP-1A/AflP-1b–transfected maize, indicating the similar mode of action of these two protein complexes\(^2^5\), and urging further searches for alternative biopesticides.

Due to their specific interactions with the insect-specific sphingolipid CPE, it can be assumed that \textit{Pleurotus} aegerolysins OlyA, PlyA2, and EryA can target this lipid in cell membranes of insect pests and form pores in the presence of their MACPF-protein partners. The aim of this study was therefore to determine the toxic potential of these aegerolysin/PlyB complexes on major insect pests such as the Coleoptera species WCR (\textit{Diabrotica virgifera virgifera}), CPB (\textit{Leptinotarsa decemlineata}) and mealworm (\textit{Tenebrio molitor}), as well as the grain aphid (\textit{Sitobion arenae}, Hemiptera), the greater wax moth (\textit{Galleria mellonella}, Lepidoptera), and the spotted wing \textit{Drosophila} (\textit{Drosophila sucukii}, Diptera). Furthermore, to elucidate the molecular mechanisms of their toxicities, the binding of \textit{Pleurotus} aegerolysins to biological and artificial membranes that contain biologically relevant CPE concentrations, and the permeabilization of these membranes by the aegerolysin/PlyB complexes, were studied in detail. Since EryA was not applied in combination with its endogenous MACPF-protein partner, EryB, the results obtained with EryA/PlyB complexes were not quantitatively compared with those obtained with OlyA6/PlyB and PlyA2/PlyB combinations.
Results

*Pleurotus* aegerolysins and aegerolysin/PlyB complexes interact with artificial and biological lipid membranes that contain ceramide phosphoethanolamine. Binding to membranes. Using sedimentation assays, we initially confirmed previously reported data by Bhat et al.\(^3\) that OlyA6, PlyA2, and EryA bind to equimolar CPE/cholesterol multilamellar vesicles, while no interactions, or negligible binding, was observed with CPE/1-palmitoyl-2-oleoyl-snglycero-3-phosphocholine (POPC) and POPC/cholesterol vesicles (Fig. 1A, Supplementary Fig. S1). We also confirmed that while OlyA6 and PlyA2 bind to sphingomyelin/cholesterol multilamellar vesicles, EryA was recovered in the supernatant after centrifugation of these vesicles, which suggested no association of EryA with these lipid membranes\(^3\). Cry34Ab1 is an aegerolysin-like protein from *B. thuringiensis*, and it bound nonspecifically to all of the forms of the multilamellar lipid vesicles.

In invertebrate membranes, however, CPE levels are considerably lower, as they represent around 2 to 6 mol% of the total membrane lipids.\(^4\),\(^5\). Here, the *Pleurotus* aegerolysins, and especially OlyA6, bound strongly to the artificial lipid membranes composed of equimolar POPC and cholesterol supplemented with biologically relevant levels (i.e., 5 mol%) of CPE (Fig. 1B, Supplementary Figs S2 and S3). As shown for OlyA6 and EryA, these two aegerolysins differed most for their binding affinities and lipid specificities; their binding was retained even with CPE levels as low as 1 mol%, which was also dependent on the presence of membrane cholesterol (Supplementary Figs S2–S4). In vesicles containing 5 mol% CPE, OlyA6 was partially recovered in the sediments of vesicles containing 30 mol% and higher cholesterol levels, while EryA binding required higher molar proportions of cholesterol (>30 mol%).

Surface plasmon resonance studies with on-chip immobilized large unilamellar vesicles (LUVs; Fig. 2) confirmed the data obtained in the sedimentation assays, and provided deeper insight into the kinetics of the *Pleurotus* aegerolysin interactions with CPE-containing membranes. All of these aegerolysins interacted with LUVs that contained 5 mol% CPE; however, EryA showed the same levels of binding at higher concentrations than those seen for OlyA6 and PlyA2, indicating its weaker interaction with these LUVs. Also, in contrast to OlyA6 and PlyA2, the binding kinetics of EryA during the dissociation phase indicated its reversible interaction with the lipid membranes. In line with the sedimentation assay, the surface plasmon resonance confirmed that the binding of both OlyA6 and EryA to LUVs increased with increased CPE content, and that this was cholesterol dependent (Supplementary Figs S5, S6). Finally, the interactions of all three aegerolysins with the LUVs, including EryA that was not used in combination with its endogenous protein partner (EryB), were considerably stronger in the presence of PlyB (Fig. 2), which suggested the formation of more stable proteolipid complexes.

In agreement with the artificial lipid systems used, the binding of the fluorescent derivatives of OlyA6 and EryA was assayed on the insect Sf9 cells (which contain CPE) and the mammalian MDCK cells (which contain sphingomyelin). These data confirmed the binding of OlyA6-mCherry to both cell types (Fig. 3A,B), while EryA-mCherry bound only to the insect Sf9 cells (Fig. 3C,D). Preincubation of OlyA6-mCherry with equimolar CPE/POPC/cholesterol multilamellar vesicles abolished the binding to MDCK cells, whereas preincubation with the same amount of equimolar SM/cholesterol multilamellar vesicles did not significantly affect this binding (Supplementary Fig. S7), again indicating the preference of the aegerolysin protein for CPE over SM.
Membrane permeabilization. By monitoring the fluorescence of calcein released from the small unilamellar lipid vesicles composed of CPE, POPC, and cholesterol (molar ratio, 5/47.5/47.5). The vesicles were immobilized (Biacore L1 chip) to approximately 7000 RU and the analytes were injected in running buffer (flow rate, 10μL/min). Representative sensorgrams of triplicate analyses are shown. (A–C) Binding to lipid vesicles of 0.25μM OlyA6 alone or in combination with PlyB (A), 0.25μM PlyA2 alone or in combination with PlyB (B), and 5μM EryA alone or in combination with PlyB (C) (aegerolysin/PlyB molar ratio, 12.5/1).

Figure 2. Surface plasmon resonance of the interactions of OlyA6, PlyA2, and EryA with large unilamellar lipid vesicles composed of CPE, POPC, and cholesterol (molar ratio, 5/47.5/47.5). The vesicles were immobilized (Biacore L1 chip) to approximately 7000 RU and the analytes were injected in running buffer (flow rate, 10μL/min). Representative sensorgrams of triplicate analyses are shown. (A–C) Binding to lipid vesicles of OlyA6/PlyB (A), PlyA2/PlyB (B), and EryA/PlyB (C) (aegerolysin/PlyB molar ratio, 12.5/1).
PlyB, and EryA/PlyB indicated their selective affinity against WCR and CPB, with no effects seen on the survival and behaviour of *T. molitor*, *D. suzukii*, *S. avenae*, and *G. melonella* (Supplementary File S1, Fig. S9, Table 1).

Exposure of the WCR larvae to the artificial diet that contained aegerolysin/PlyB or the insecticide (Decis) had significant effects on their survival over these 7-day bioassays ($\chi^2 = 120.8; P < 0.0001$), whereby PlyA2/PlyB, OlyA6/PlyB, and Decis significantly increased the larva mortality (Fig. 6A, Table 2). Various concentrations of the aegerolysin/PlyB complexes had significant detrimental effects on the WCR larvae when evaluated on day 3 (PlyA2/PlyB, $P = 0.002$; OlyA6/PlyB, $P < 0.0001$). PlyA2/PlyB caused significant reduction of the larva length at 11.2 $\mu$g/cm², and OlyA6/PlyB at 2.8, 5.6 and 11.2 $\mu$g/cm² (Fig. 6C,D).

Figure 3. Representative fluorescence microscopy images of cell-surface labeling of insect and mammalian cells with OlyA6 and EryA. Sf9 and MDCK cells were labeled with 5 $\mu$M OlyA6-mCherry or 5 $\mu$M EryA-mCherry, as described in the Methods. Numbers on images indicate fluorescence acquisition times. Scale bars, 20 $\mu$m.

Figure 4. Permeabilization of small unilamellar vesicles composed of CPE, POPC, and cholesterol (molar ratio, 5/47.5/47.5) by OlyA6/PlyB, PlyA2/PlyB, and EryA/PlyB. Fluorescence intensity of calcein released from the lipid vesicles, monitored as described in the Methods. (A) Permeabilization of lipid vesicles by 10 $\mu$g/mL aegerolysin with PlyB (molar ratio, 12.5/1). (B) Concentration dependence of lipid vesicle permeabilization for OlyA6/PlyB, PlyA2/PlyB, and EryA/PlyB. Data are means ± standard error from three repetitions.
Exposure of the WCR beetles to the artificial diet containing the aegerolysin/PlyB complexes or the insecticide (Decis) also had significant effects on their survival over these 7-day bioassays ($\chi^2 = 139.3; P < 0.0001$), whereby OlyA6/PlyB and the insecticide significantly increased beetle mortality (Fig. 6B, Table 2). The individual aegerolysins, PlyB alone, and buffer did not significantly affect the WCR larvae survival or length, or the WCR beetle survival (data not shown).

Exposure of the CPB larvae to the leaf discs that were treated with the aegerolysin/PlyB complexes (9.0 $\mu$g/cm²) or the insecticide (0.1% Actara) had significant effects on their survival ($\chi^2 = 136.7; P < 0.0001$) and weight changes over these 5-day bioassays. The aegerolysin/PlyB complexes and the insecticide significantly increased larva mortality, with the exception of EryA/PlyB with the L3 + L4 CPB larvae (Fig. 7A,B, Table 2). It should be stressed however that the insecticidal effects of EryA/PlyB complexes cannot be directly compared to those obtained with OlyA6/PlyB and PlyA2/PlyB combinations, since EryA was not used in combination with its endogenous protein partner, EryB. The aegerolysin/PlyB complexes and the insecticide also had significant effects on the CPB larva feeding, as indicated by the changes in the larva weights (L1 + L2 and L3 + L4, $P < 0.0001$). Further, all of the aegerolysin/PlyB complexes and the insecticide significantly reduced the larva weight gain for

![Figure 5](https://example.com/figure5.png)

**Figure 5.** Effects of OlyA6/PlyB, PlyA2/PlyB, and EryA/PlyB on insect Sf9 cell and mammalian MDCK cell survival. Survival rates of Sf9 (A) and MDCK (B) cells treated with OlyA6/PlyB, PlyA2/PlyB, and EryA/PlyB (as indicated), expressed as the ratio between the luminescence of the treated and the control cells (x100) after 30 min of exposure. Data are means ± SD from three independent experiments.

| Order     | Species                  | Common name                        | Aegerolysins (μg/mL) | Effects |
|-----------|--------------------------|------------------------------------|----------------------|---------|
| Coleoptera| *Tenebrio molitor*       | Mealworm                           | 500                  | Inactive|
|           | *Diabrotica virgifera virgifera* | Western corn rootworm (adults) | 500                  | Death   |
|           | *Diabrotica virgifera virgifera* | Western corn rootworm (larvae)    | 5.6*                 | Death   |
|           | *Leptinotarsa decemlineata* | Colorado potato beetle (larvae)   | 9.0*                 | Death   |
| Lepidoptera| *Galleria mellonella*    | Greater wax moth (caterpillars)    | 500                  | Inactive|
| Hemiptera | *Sitobion avenae*        | Grain aphid                        | 500                  | Inactive|
| Diptera   | *Drosophila suzukii*     | Spotted wing drosophila            | 500                  | Inactive|

**Table 1.** Insecticidal activities of the *Pleurotus* aegerolysin/PlyB complexes against the various insects. *μg/cm².*
aegerolysin/PlyB. bHazard ratio is defined as the slope of the survival curve, as a measure of how rapidly and PlyA2/PlyB assessed only for WCR larvae at day 7. cInsecticide based on active ingredient: deltamethrin. dInsecticide based on active ingredient: thiametoxam. ND, not determined.

Table 2. Estimates of LT_{50} and LD_{50} of the Pleurotus aegerolysin/PlyB protein complexes against the WCR larvae and beetles and the CPB larvae. aWCR larvae were exposed to 5.6 μg/cm² aegerolysin/PlyB. WCR beetles were exposed to 0.5 mg/mL aegerolysin and 0.04 mg/mL PlyB. CPB larvae were exposed to 9.0 μg/cm² aegerolysin/PlyB. bHazard ratio is defined as the slope of the survival curve, as a measure of how rapidly the subjects died. The hazard ratio was computed for each treatment from the survival curves, compared to the buffer (negative control). cχ² (log-rank test) was computed separately for each pairwise comparison of treatment versus buffer (negative control), and consequently its corresponding P-value compared to Bonferroni-corrected significance thresholds (P/K, where P = 0.05 and K is the number of pairwise comparisons). If the P-value exceeded the Bonferroni-corrected significance threshold, it is reported as >0.05. dLD_{50} of OlyA6/PlyB and PlyA2/PlyB assessed only for WCR larvae at day 7. eInsecticide based on active ingredient: deltamethrin. fInsecticide based on active ingredient: thiametoxam. ND, not determined.

L3 + L4 (Fig. 7C,D). The individual aegerolysins, PlyB alone, and the buffer did not significantly affect the L1 + L2 or L3 + L4 larva survival or weight changes (data not shown).

Discussion

The Pleurotus aegerolysins OlyA6, PlyA2, and EryA target the invertebrate membrane sphingolipid CPE and recruit the MACPF protein, PlyB, to consequently form transmembrane lytic complexes. Here, we have shown these interactions might be useful in the further development of novel insect biopesticides. Such specificity, in turn, renders these aegerolysin/PlyB complexes less toxic to taxa outside their primary specificity range, which is an important requirement in pest control management51.

Previous studies have described the interactions of OlyA6, OlyA, PlyA, PlyA2, and EryA with lipid vesicles composed of equimolar proportions of sphingomyelin/cholesterol or CPE/cholesterol in detail28,30,32,33,52. The potential of the aegerolysins to be used as biomarkers of membrane rafts and for defining the CPE distribution in invertebrates have also been reported31–33,37,53. Here, however, we have initially demonstrated that these interactions might be useful in the further development of novel insect biopesticides. Such specificity, in turn, renders these aegerolysin/PlyB complexes less toxic to taxa outside their primary specificity range, which is an important requirement in pest control management51. Previous studies have described the interactions of OlyA6, OlyA, PlyA, PlyA2, and EryA with lipid vesicles composed of equimolar proportions of sphingomyelin/cholesterol or CPE/cholesterol in detail28,30,32,33,52. The potential of the aegerolysins to be used as biomarkers of membrane rafts and for defining the CPE distribution in invertebrates have also been reported31–33,37,53. Here, however, we have initially demonstrated that these interactions might be useful in the further development of novel insect biopesticides. Such specificity, in turn, renders these aegerolysin/PlyB complexes less toxic to taxa outside their primary specificity range, which is an important requirement in pest control management51.

L3 + L4 (Fig. 7C,D). The individual aegerolysins, PlyB alone, and the buffer did not significantly affect the L1 + L2 or L3 + L4 larva survival or weight changes (data not shown).
insect membrane proteins have been suggested to be the receptors for the recently discovered aegerolysin-like AIP-1A protein from _A. faecalis_.

It has been reported that the combination of OlyA or PlyA with PlyB readily forms 13-meric transmembrane pores in natural and artificial membranes that contain sphingomyelin/cholesterol domains. However, our data in the present study obtained with artificial lipid vesicles and with insect cell cultures also demonstrate the permeabilization activities of the OlyA6/PlyB, PlyA2/PlyB, and EryA/PlyB complexes toward CPE-containing membranes. Importantly, this membrane permeabilization can be observed at ≤ 5 mol% CPE; i.e., at biologically relevant CPE concentrations. The lower membrane-disrupting potential of EryA/PlyB seen here might be explained by the observed lower affinity of EryA toward CPE/cholesterol membranes, and also because EryA was not used in combination with its endogenous pore forming partner protein EryB, which however shares high amino acid sequence identity with PlyB (96%).

Although both Cry34Ab1 and OlyA-A alone have been reported to cause remodeling of biological membranes, in the present study, the individual aegerolysins used and PlyB alone did not have any insecticidal effects on WCR and CPB. Only the combination of these _Pleurotus_ aegerolysins and PlyB significantly increased insect mortality and the nonlethal effects, such as reduced larva growth (for WCR) and decreased larva weight (for CPB).

These data on the insecticidal effects on WCR larvae are directly comparable to some previously published data. Shlotter and Storer reported a LD50 of 11 µg/cm² for WCR larvae treated with Cry34Ab1/Cry35Ab1 complexes, which is in the same range as seen here for these _Pleurotus_ aegerolysin/PlyB complexes. Shlotter and Storer also reported that 35 µg/cm² Cry34Ab1/Cry35Ab1 complex was apparently not high enough to kill 50% of CPB. In the present study, the aegerolysin/PlyB complexes were more potent, as 9.0 µg/cm² killed 50% of CPB larvae in 4.1 to 5.4 days.
It is more difficult to directly compare other studies to the present study due to methodological differences. Moellenbeck et al.\textsuperscript{42} reported a LC\textsubscript{50} of 55 μg (cell biomass)/cm\textsuperscript{2} after 4 days of exposure of WCR larvae to a crude lysate of \textit{B. thuringiensis} strain PS149B1, which produced the Cry34Ab1/Cry35Ab1 toxins. If we approximate the aegerolysin input concentrations in the bioassays of the present study in μg/mL, as would be obtained if they were uniformly incorporated within the assay medium, we would obtain a LD\textsubscript{50} of 33.8 ± 8.5 μg/mL for PlyA/PlyB and 47.0 ± 11.0 μg/mL for OlyA6/PlyB after 5 days of exposure. This is comparable to the findings of Yalpani et al.\textsuperscript{23}, who reported a LD\textsubscript{50} for WCR larvae of 30.4 μg/mL after 3 to 4 days of exposure to AflP-1A/AflP-1B from \textit{Alcaligenes faecalis}. The present data for day 3 (PlyA/PlyB LD\textsubscript{50}, 54.9 ± 23.2 μg/mL; OlyA6/PlyB LD\textsubscript{50}, 58.3 ± 11.7 μg/mL) are similar to the findings of Wei et al.\textsuperscript{21}, who reported a LD\textsubscript{50} for WCR larvae of 52.5 μg/mL after 4 days of exposure to the insecticidal protein PIP-47Aa from \textit{P. mosselii}. Schellenberger et al.\textsuperscript{20} reported a LC\textsubscript{50} of ~120 μg/mL for IPD072Aa from \textit{Pseudomonas chlororaphis} after 8 days of exposure. Here, we report lower LD\textsubscript{50} for 7 days of exposure: PlyA/PlyB, 27.4 ± 12.5 μg/mL; OlyA6/PlyB, 29.0 ± 13.2 μg/mL.

The specificity of these \textit{Pleurotus} aegerolysin/PlyB complexes toward WCR and CPB might be due to physiological differences between the species tested. In contrast to the other insects tested, the midgut pH of these Coleoptera species has been reported to be around 6.66-6.8, which is also the optimal pH for OlyA6 binding to membrane lipids\textsuperscript{61}. Furthermore, aegerolysin-based cytolytic protein complexes might be degraded by proteolytic enzymes present in the midgut of resistant insect species\textsuperscript{62,63}, or they might simply not reach their molecular targets due to the presence of glycocalyx in midgut epithelial cells\textsuperscript{64}. Finally, in addition to CPE, the presence of an additional (glyco)lipid or (glyco)protein aegerolysin receptor site(s) on insect cell membranes cannot be excluded at present.
The loss of larval weight and decrease in their growth, and their death caused by the aegerolysin/PlyB complexes here are similar to the described effects of *B. thuringiensis* Cry34Ab1/Cry35Ab1 toxins. When these *B. thuringiensis* toxins were ingested, they affected the apical surface of the WCR midgut epithelium. Gross changes in the midgut integrity resulted in starvation and dehydration of the larvae. Although the receptors for the Cry34Ab1/Cry35Ab1 toxins and aegerolysin/PlyB complexes are obviously different, we would suggest that the pathophysiological effects of these proteins are similar, and are basically accounted for by permeabilization of the insect midgut epithelium cells.

We can conclude that these CPE-binding aegerolysins from fungal genus *Pleurotus* in complexes with the MACPF-like protein partner, PlyB, are selectively toxic to WCR and CPB larvae, and can be used to develop a novel generation of biopesticides produced by genetically modified plants for the control of these pests. Importantly, due to their interactions with their specific insect membrane lipid receptor, and not with pest proteins that can be prone to mutation, the chances of evolving resistance to these aegerolysin proteins should be extremely small. Finally, the bioinsecticidal effects of protein complexes based on *Pleurotus*-derived aegerolysins and MACPF partners suggest that in their natural environment, these complexes might be involved in the defence of oyster mushrooms against insect larvae.

**Materials and Methods**

**Materials.** All chemicals used in the present study were from Sigma–Aldrich (USA) unless specified otherwise. Porcine brain sphingomyelin, wool grease cholesterol, and POPC were from Avanti Polar Lipids (USA), and CPE was from Matreya (USA). These lipids were stored at −20 °C and dissolved in chloroform prior to use. CPE was dissolved in 1 ml chloroform/methanol (9/1, v/v) with the addition of 3 μL Milli-Q water.

**Cells.** Insect cells derived from the ovarian epithelial cells of the fall army worm (*Spodoptera frugiperda*; Sf9 cells; Thermo Fischer Scientific, USA) were maintained in continuous suspension culture under serum-free conditions at 28 °C in Insect XPRESS protein-free insect cell medium with L-glutamine (Lonza, USA), with agitation at 150 rpm. Mammalian Madin-Darby canine kidney (MDCK) cells (NBL-2; ATCC-CCL-34) derived from canine kidney epithelium were obtained from ATCC (Manassas, USA). The cells were cultured in A-DMEM/F12 (1/1, v/v) with 5% fetal calf serum and antibiotics, and maintained in a controlled atmosphere at 37 °C and 5% CO2.

**Proteins.** The OlyA6, OlyA6-mCherry, and ΔA48PlyB (henceforth PlyB) recombinant proteins were produced as described previously30,31. The EryA, EryA-mCherry, PlyA2, and Cry34Ab1 recombinant proteins were produced following the protocols for OlyA6 and OlyA6-mCherry, with slight modifications, as follows. *Escherichia coli* BL21(DE3) cells containing the respective plasmids were grown in Terrific Broth (1.2% tryptone, 2.4% yeast extract, 0.4% glycerol), and protein expression was induced using 0.5 mM isopropyl 1-thio-β-D-galactopyranoside, overnight at 20 °C. The supernatant of the lysed bacterial cells was centrifuged for 30 min at 100,000 × g and 4 °C, and incubated with Ni2+-NTA beads. Cry34Ab1 was further purified using gel filtration, on Superdex 75 (GE Healthcare, UK) in 0.2 M Na-citrate, pH 4.0. *Pleurotus* proteins were dialyzed against 20 mM Tris-HCl, pH 8.0, overnight at 4 °C, and further purified on a MonoQ anion exchange column (GE Healthcare, UK). The proteins were eluted with a 0 to 250 mM NaCl gradient in the same buffer. Protein concentrations were determined at 280 nm using a microvolume spectrophotometer (Nanodrop2000; Thermo Scientific, USA). Protein size and purity were determined using sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE; Bio-Rad, USA) on homogeneous 12% acrylamide gels. The protein was then stained with SimplyBlue SafeStain (Thermo Fisher Scientific, USA), or detected with anti-His antibodies after Western blotting (Qiagen, Germany).

**Target insects.** *Drosophila suzukii* (Matsumura)) were laboratory reared from a colony initially provided by Julius Kühn (Institute for Biological Control, Darmstadt, Germany). They were reared in 30 × 30 × 30 cm plastic insectoria in a growth chamber at 22 ± 1 °C and 77% relative humidity, and a photoperiod of 14 h/10 h (light/dark). The *Drosophila* were provided with tap water and solid artificial food medium (20 g agar, 20 g sucrose, 10 g wheat flour, 50 g dry baker’s yeast, 500 mL tap water, 400 g grated organic apples, 500 mL organic apple juice, 50 mL apple vinegar, and 4 g nipagin (methyl 4-hydroxybenzoate; Sigma–Aldrich).

Western corn rootworm (*Diabrotica virgifera virgifera* LeConte) beetles were collected from corn fields from central Slovenia a day prior to the experiments. WCR larvae from a US nondiapausing colony were reared in the insectary of the Agricultural Institute of Slovenia. The initial eggs were obtained from CABI Europe (Switzerland) at the Plant Protection Directorate, Hungary.

Individuals of the grain aphid *Sitobion avenae* Fabricius were collected from winter wheat (*Triticum aestivum* L.) fields in central Slovenia. The *S. avenae* were reared on wheat seedlings in a growth chamber at 22 ± 1 °C and 77% relative humidity, and a photoperiod of 14 h/10 h (light/dark). These aphids have been kept in culture for more than 1 year.

The Colorado potato beetle (*Leptinotarsa decemlineata* Say; CPB) larvae were collected from potato fields a day prior to the experiments, and divided into two groups: young larvae (L1 + L2) and old larvae (L3 + L4). The pronotum was entirely black for L1 + L2. For L3, the anterior margin of the pronotum appeared orange-brown, while for L4, about half of the pronotum was light brown, anteriorly.

Mealworms (*Tenebrio molitor* L.) and greater wax moths (*Galleria mellonella* L.) were mass reared in a growth chamber at 30 ± 1 °C and 55% relative humidity, and in the dark, in the insectary of the Agricultural Institute of Slovenia.
Methods. Protein sedimentation assay with multilamellar lipid vesicles. Multilamellar vesicles were prepared with different lipid molar ratios (final concentration, 5 mg/mL) in 140 mM NaCl, 20 mM Tris, 5 μM EDTA, pH 8.0, as described previously. The aegerolysins were added to these vesicles at a 1/10 (w/w) ratio, and incubated for 30 min on a rotary shaker (600 rpm/min) at 25 °C, followed by centrifugation for 60 min at 60,000 × g and 4 °C. The supernatants contained the unbound aegerolysins, and they were transferred to vials, and precipitated with 20% trichloroacetic acid. After a 10-min incubation on ice, the precipitated proteins were sedimented by centrifugation at 14,300 × g for 12 min at 4 °C, and the pellets were washed twice with 300 μL ice-cold acetone. The fractions containing the free and bound aegerolysin proteins were diluted with an equal volume of nonreducing SDS sample buffer (20 mM Tris–HCl, pH 8.0, 5% [w/v] SDS, 2 mM EDTA, 0.1% [w/v] bromophenol blue), heated to 100 °C for 5 min, and applied to homogeneous 12% acrylamide gels. Proteins were stained with SimplyBlue SafeStain (Thermo Fisher Scientific, USA).

The lipid molar ratios and total lipid concentrations in lipid vesicle suspensions were determined colorimetrically using free cholesterol E and phospholipids C kits (Wako Pure Chemical Industries, Japan), and CPE concentrations were determined by quantification of the amine group.

Surface plasmon resonance-based binding studies. To prepare LUVs, suspensions of multilamellar vesicles were subjected to five freeze–thaw cycles and then extruded through 0.1 μm polycarbonate filters (Millipore, Germany) at 40 °C. The size of the LUVs was estimated using dynamic light scattering (Zetasizer Nano ZSP; Malvern Instruments, Malvern, UK). The monomodal distribution of their hydrodynamic radii ranged from 40 nm to 60 nm.

The aegerolysin/LUV interactions were monitored using a surface plasmon resonance-based refractometer (Biacore X; GE Healthcare, USA) and an L1 sensor chip, with 20 mM Tris, 140 mM NaCl, 5 μM EDTA, pH 7.4, as the running buffer. After the initial cleaning of the chip with regeneration solutions of SDS and octyl β-D-glucopyranoside with 1-min injections at a flow rate of 10 μL/min, the LUVs were bound to the second flow cell of the sensor chip to reach responses of ~7,000 RU. The first flow cell was left empty and was used to control for possible nonspecific binding of the aegerolysin proteins to the chip dextran matrix. Loosely bound LUVs were washed from the surface with a 1-min injection of 100 mM NaOH. The nonspecific binding of the aegerolysin proteins was minimized using 1 μM bovine serum albumin at a flow rate of 30 μL/min. The appropriate solutions of the aegerolysins and the aegerolysin/PlyB complexes were injected at a flow rate of 10 μL/min for 1 min, and their dissociation was monitored for an additional 6 min. The interactions with LUVs were tested for OlyA6, PlyA2, and EryA at concentrations from 0.25 μM to 5 μM, as well as for the aegerolysin complexes OlyA6/PlyB, PlyA2/PlyB, and EryA/PlyB (aegerolysin/PlyB molar ratio, 12.5/1). Chip regeneration between injections was achieved with 1-min injections of 0.5% SDS and 40 mM β-D-glucopyranoside, with a flow rate of 10 μL/min. Experiments were performed at 25 °C. The data were processed with the BIAevaluation software (GE Healthcare).

Binding of fluorescently tagged aegerolysins to cell membranes. The Sf9 and MDCK cells were grown on glass coverslips for 1 day and 2 days, respectively. For the labeling with fluorescently tagged aegerolysins, the cells were first fixed using 2% paraformaldehyde, for 15 min at 25 °C, and then washed with phosphate-buffered saline. Then the cells were incubated with 5 μM OlyA6-mCherry or 5 μM EryA-mCherry for 10 min at 25 °C. In some cases, OlyA6-mCherry was preincubated (10 min at 25 °C) with 5 mM multilamellar vesicles composed of equimolar proportions of SM/cholesterol, CPE/POPC/cholesterol or POPC/cholesterol.

After washing with phosphate-buffered saline, the cells were mounted in Vectashield with 4′,6-diamidino-2-phenylindole (DAPI) for nuclei staining, and observed under a fluorescent microscope (Axiolab Z1) using an oil-immersion objective (63 × oil, NA 1.40) and an ApoTome device (Carl Zeiss, Germany), for the generation of optical sections. Images were acquired using the Axio-Vision program (Carl Zeiss, Germany).

Permeabilization of the small unilamellar vesicles. Small unilamellar vesicles loaded with calcine at the self-quenching concentration (80 mM) were prepared as described previously. Vesicle permeabilization was determined using a fluorescence microplate reader (Biotek, UK) with excitation and emission set at 485 nm and 535 nm, respectively. Calcein-loaded vesicles composed of various molar proportions of lipids were exposed to different aegerolysins (concentrations range, 0 to 100 μM) alone or in combination with PlyB at the molar ratio of 12.5/1. The experiments were run for 1 h at 25 °C. The permeabilization induced by the lytic aegerolysin complexes was expressed as the percentage of the maximal permeabilization obtained by addition of the detergent Triton-X 100 to a final concentration of 1 mM.

Cell viability assay. Cell viability assays were performed using the CellTiter-Glo reagent (Promega Corp, USA), following the manufacturer instructions. In brief, Sf9 or MDCK cells were plated in 96-well plates (Costar, USA) at 3.7 × 10^4 cells/cm² and 3 × 10^4 cells/cm², respectively, in their respective growth medium. After 48 h, the cells were exposed to the aegerolysins (0.1, 1, 5 μM) alone or in combination with PlyB at an aegerolysin/PlyB molar ratio of 12.5/1, for 30 min. The luminescence was measured using a microplate reader (Safire 2; Tecan, Switzerland). The data from the viability assays are expressed as percentages of the luminescence of treated to untreated cells, as means ± standard error of three independent experiments, each performed in triplicate.

Insecticidal tests. The insecticidal tests were carried in accordance with the Slovenian law on animal protection (U.I.R.S., 2007). The study was not subject to ethical protocols according to EU legislature (Directive 2010/63/EU).

Western corn rootworm: Western corn rootworm larvae (of a US nonidiapausing strain) were provided with the southern corn rootworm artificial diet (Frontier Agricultural Sciences, USA; product F9800B; 13.8 g), with
some additions. These included agar (1.5 g), lyophilised corn roots (0.63 g), methylene blue solution (product 50484; 100 µL; Sigma–Aldrich, Germany), and demineralized water (88 mL). The pH of the diet was adjusted to 8.0 using 10 M NaOH. Two and a half milliliters of the diet was pipetted into individual wells of sterile six-well plates. Once solidified, 100 µL OlyA6/PlyB, PlyA2/PlyB or EryA/PlyB was added, that was spread evenly over the entire surface of the diet in the wells using a sterile glass stirring rod. The plates were then left to dry for 30 min in a laminar flow chamber.

The mean survival time (LT₅₀) was determined by applying 100 µL 0.5 mg/mL OlyA6, PlyA2 or EryA, and 0.04 mg/mL PlyB onto the artificial diet, which provided the final concentration of 5.6 µg/cm² of the aegerolysin/PlyB complexes on the diet. Five neonate larvae were placed in each well. The wells were sealed using pressure-activated sealing foil (product 4ti-0560; 4titude, UK), which was perforated five times to allow for gas exchange. Each treatment was performed in two wells of four replicate six-well plates, and the experiment was repeated twice independently, thus using a total of 80 larvae for each treatment. The bioassay was carried out over 7 days, with larva survival recorded on days 1, 3, 5, and 7.

Following the same 7-day procedure, to determine the median lethal dose (LD₅₀), six concentrations of the aegerolysin/PlyB complexes were tested, from 0.0 µg/cm² to 11.2 µg/cm². The total sample size tested at each concentration was 15. Additionally, on day 3, the lengths of the living larvae were measured using a motorized, calibrated stereomicroscope (M205 C; Leica, Germany). The bioassays were performed in a growth chamber at 22 ± 1 °C and 77% relative humidity, and with a photoperiod of 14 h/10 h (light/dark). Buffer was used as the negative control (20 mM Tris, 0.5% glycerol, pH 8.0), with the positive control of 0.3% dilution of the insecticide Decis (active ingredient, 2.8% [w/w] deltamethrin; Bayer CropScience, Germany).

The western corn rootworm beetles collected from corn fields the day prior to the experiments received an artificial diet (pH 7.0) that comprised: sucrose (33 g); alphacel (18.6 g); casein (15.9 g); soy flour (12.4 g); wheat germ (12.4 g); brewer’s yeast (5 g); vitamin mix (Vanderzant, 1.2 g; MP Biomedicals, USA); salt mix (Wesson, 1.2 g; Careforde Inc, USA); cholesterol (0.3 g); glycerol (11 mL); and demineralized water (82.5 mL). The recombinant aegerolysin proteins for the bioassays were mixed with this diet (1/1, v/v), to a final concentration of 0.5 mg/mL aegerolysin and 0.04 mg/mL PlyB. Thirty microliters of this mixture was pipetted into each well of a six-well plate. A single WCR adult was placed into each well. Each replicate six-well plates were performed per treatment, and the experiment was repeated twice independently, for a total of 36 beetles for each treatment. The bioassay was performed in a growth chamber at 22 ± 1 °C and 77% relative humidity, and a photoperiod of 14 h/10 h (light/dark). The buffer (20 mM Tris, 0.5% glycerol, pH 8.0) was used as the negative control, and a 0.5% dilution of the artificial food (Decis) was mixed with the artificial food (1/1, v/v) as the positive control. The bioassay was carried out over 7 days, with the survival recorded daily. Additionally, the individual aegerolysins and PlyB were tested for their potential toxicity for the WCR larvae and beetles.

Colorado potato beetle: The Colorado potato beetle larvae collected from potato fields the day prior to the experiments were divided into two groups based on their pronotum coloring: young larvae (L1–L2), and old larvae (L3 + L4). The larvae fed on 14-mm-diameter potato leaf discs that were treated by soaking in OlyA6/PlyB, PlyA2/PlyB and EryA/PlyB (0.5 mg/mL aegerolysin, 0.04 mg/mL PlyB) for 5 min. The calculated concentration of aegerolysin/PlyB on the leaf surface was 9.0 µg/cm². A single treated leaf disc and a CPB larva were placed in each well of a six-well plate. Every second day, the relevant freshly treated potato leaf discs were provided. Two replicate six-well plates per treatment were performed, and the experiment was repeated three times independently, giving a total of 36 larvae of each group for each treatment. The bioassay was performed in an incubator at 22 ± 1 °C and 77% relative humidity, and a photoperiod of 14 h/10 h light/day. The buffer-treated (20 mM Tris, 0.5% glycerol, pH 8.0) leaf discs were used as the negative control, and treatment with a 0.1% dilution of insecticide (Actara 25 WG; active ingredient, 25% [w/w] thiametoxam; Syngenta, Switzerland) was used as the positive control. The survival rate was recorded daily over 5 days. The weights of the larvae were recorded on days 1 and 5. Additionally, the individual aegerolysins, PlyB, and buffer were tested for potential toxicity against the CPB larvae. The insecticidal tests for the other insects tested were carried out as described in Supplementary File S1.

Insecticidal data analysis: The time-based larvae and beetle mortality were analyzed using Kaplan–Meier survival analysis and nonlinear regression. When multiple survival curves were compared, the significance threshold was corrected according to the Bonferroni method. The effects of the aegerolysin proteins on the WCR larva lengths and the CPB larva weight changes were analyzed using Kruskal–Wallis tests, followed by Dunn’s post-hoc tests. The data were analysed using GraphPad Prism (GraphPad Software, Inc., USA).

Data Availability
The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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Author Contributions
A.P., V.H., M.S., M.N., Š.M., I.P., S.P., M.Z., N.R. and J.R. conducted the experiments; A.P., P.M., P.V., J.R. and K.S. contributed laboratory equipment and institutional affiliations.

Competing Interests: Part of the results presented within this paper (bioinsecticidal effects of aegerolysin/MACPF-protein complexes against Colorado potato beetle and Western corn rootworm) are covered by a PCT patent application No. PCT/EP2017/074877, filed on September 29, 2017. The patent applicants are University of Ljubljana and the Agricultural Institute of Slovenia. The inventors are the following employees of patent applicants: Anatasija Panevska, Kristina Sepčić, Peter Maček, Matej Skočaj, Maruša Novak, Matej Butala, Vesna Hodnik, Maja Grundner and Tom Turk (University of Ljubljana), and Jaka Razingar and Spela Modic (Agricultural Institute of Slovenia).

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