Bombardier Enables Delivery of Short-Form Bomanins in the Drosophila Toll Response

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Toll mediates a robust and effective innate immune response across vertebrates and invertebrates. In Drosophila melanogaster, activation of Toll by systemic infection drives the accumulation of a rich repertoire of immune effectors in hemolymph, including the recently characterized Bomanins, as well as the classical antimicrobial peptides (AMPs). Here we report the functional characterization of a Toll-induced hemolymph protein encoded by the bombardier (CG18067) gene. Using the CRISPR/Cas9 system to generate a precise deletion of the bombardier transcriptional unit, we found that Bombardier is required for Toll-mediated defense against fungi and Gram-positive bacteria. Assaying cell-free hemolymph, we found that the Bomanin-dependent candidacidal activity is also dependent on Bombardier, but is independent of the antifungal AMPs Drosomycin and Metchnikowin. Using mass spectrometry, we demonstrated that deletion of bombardier results in the specific absence of short-form Bomanins from hemolymph. In addition, flies lacking Bombardier exhibited a defect in pathogen tolerance that we trace to an aberrant condition triggered by Toll activation. These results lead us to a model in which the presence of Bombardier in wild-type flies enables the proper folding, secretion, or intermolecular associations of short-form Bomanins, and the absence of Bombardier disrupts one or more of these steps, resulting in defects in both immune resistance and tolerance.

Keywords: Drosophila melanogaster, immunity, Toll, Bomanins, humoral

INTRODUCTION

Innate immune pathways are found in plants, fungi, and animals and provide a rapid defense against a broad range of pathogens (1–3). In the fruit fly Drosophila melanogaster, the two major innate immune pathways are Toll and Imd (4–6). The Toll pathway is activated by Gram-positive bacteria with Lys-type peptidoglycan and by fungi, and is required for defense against these microbes (7–10). Conversely, the Imd pathway is activated by and plays a major role in survival against Gram-negative bacteria and Gram-positive bacteria with DAP-type peptidoglycan (11, 12). These pathways, which are both mediated by NF-κB transcription factors, are broadly conserved as initiators of innate immune responses. Activation of either pathway induces robust production of an array of immune molecules, including antimicrobial peptides (AMPs) (13–17).
AMPS are found in all kingdoms of life (18–22). These peptides have long been thought to play the principal effector role in innate immune defense due to their demonstrated in vitro antimicrobial activity and their marked upregulation after infection. However, recent research in *D. melanogaster* suggests that AMPS play a major role in Imd-mediated defense, but a relatively minor role in Toll-mediated immunity (23).

In contrast to the AMPS, the *Drosophila*-specific Bomanin peptides (Boms), which are highly induced after infection, are indispensable for resistance against pathogens controlled by the Toll pathway (24). BomΔ55C flies, which lack 10 of the 12 Bom genes, succumb to fungal and Gram-positive bacterial infections at rates indistinguishable from Toll-deficient flies (23, 24), suggesting that Boms rather than AMPS are the primary Toll effectors.

Bom peptides, like AMPS, are secreted from the fat body, the *Drosophila* immune organ, into the hemolymph, the *Drosophila* circulatory fluid. The family is comprised of three groups. The short-form peptides are 16–17 residues long and contain only the C-terminal tail. Finally, the bicipital forms consist of two Bom peptides, like AMPs, are secreted from the fat body, the *Drosophila* immune organ, into the hemolymph, the *Drosophila* circulatory fluid. The family is comprised of three groups. The short-form peptides are 16–17 residues long and contain only the C-terminal tail. Finally, the bicipital forms consist of two Bom peptides, like AMPs, are secreted from the fat body, the *Drosophila* immune organ, into the hemolymph, the *Drosophila* circulatory fluid. The family is comprised of three groups. The short-form peptides are 16–17 residues long and contain only the C-terminal tail. Finally, the bicipital forms consist of two Bom peptides, like AMPs, are secreted from the fat body, the *Drosophila* immune organ, into the hemolymph, the *Drosophila* circulatory fluid. The family is comprised of three groups. The short-form peptides are 16–17 residues long and contain only the C-terminal tail. Finally, the bicipital forms consist of two Bom peptides, like AMPs, are secreted from the fat body, the *Drosophila* immune organ, into the hemolymph,

**MATERIALS AND METHODS**

**CRISPR/Cas9 Deletion of *bombardier* Locus**

The *bombardier* gene (CG18067) was deleted using CRISPR/Cas9 technology according to established protocols (26). Briefly, a pair of gRNAs designed to delete the region 2R: 20,534,248–20,536,154 were cloned into pU6-Bbs1-chiRNA (Addgene plasmid #45946). Homology arms (1,017 bp left and 1,022 bp right) were cloned into pDsRed-atTP (Addgene plasmid #51019). The plasmid pBS-Hsp70-Cas9 (Addgene plasmid #46294) was used as the Cas9 source. Constructs were injected into *w*1118 embryos. F1 progeny were screened for DsRed eyes and homozygous lines were established. See **Supplemental Table 1** for gRNA and homology arm primer sequences.

**Toll Activation, *Drosophila* Infection, and Survival Analysis**

Flies were raised at 25°C on cornmeal molasses agar media1. The *w*1118 strain was used as the wild type. Microbial isolates, culture conditions, and conditions for infection for *Enterococcus faecalis*, *Enterobacter cloacae*, *Fusarium oxysporum*, and *Candida glabrata* were as described previously (24), except that *C. glabrata* was concentrated to OD$_{600}$ = 100. Flies were incubated at 25°C after live bacterial infection and at 29°C after fungal infection. For heat-killed challenge, bacterial cultures were autoclaved and resuspended in 20% glycerol to OD$_{600}$ = 10 for *E. faecalis* and OD$_{600}$ = 300 for *M. luteus*. For both survival assays and hemolymph preparation, flies challenged with heat-killed bacteria were incubated at 29°C.

**Hemolymph Antimicrobial Assays**

Candidalcidal activity of hemolymph was assayed as described previously (25), except that hemolymph was prepared from groups of 30 flies and all activity assays were carried out for 1 h at room temperature. The number of colonies representing zero percent killing was set as the value obtained by assaying uninduced hemolymph.

**MALDI-TOF Analysis of Hemolymph**

The Toll pathway was activated in flies using heat-killed *M. luteus*, then incubated at 29°C for 24 h. Hemolymph was extracted as in Lindsay et al. (25), with slight modifications. Hemolymph extracted with glass capillaries from five male flies was pooled and transferred into 0.1% trifluoroacetic acid (TFA)/50% acetonitrile (ACN). One µl of each mixture was spotted on a Bruker MSP 96 ground steel plate, mixed 1:1 with a saturated solution of Universal MALDI matrix (Sigma-Aldrich) in 0.1% TFA/78% ACN, and air-dried. MALDI-TOF spectra were acquired using a Bruker Autoflex mass spectrometer. Data were collected from 1,500 to 10,000 m/z in positive linear mode, and 1,000–5,000 m/z in positive reflectron mode. Peptide calibration standard II (Bruker) was mixed with Universal MALDI matrix and used as an external calibration standard. At least ten independent samples were collected for each genotype. For peptide identification, peaks were matched to those of corresponding peaks in prior studies (13, 25). Representative spectra were visualized using R 3.3.2 and ggplot2 2.2.1 (27, 28).

**Gene Expression Quantitation**

The Toll pathway was activated with heat-killed *M. luteus*. Using TRIzol (Ambion), total RNA was extracted 18 h after Toll activation from four to six adult flies (2–5 days old). Next, cDNA was synthesized from 500 ng total RNA using the SuperScript II Reverse Transcriptase kit (Invitrogen). Quantitative RT-PCR was performed on an iQ5 cycler (BioRad) using iQ SYBR Green Supermix (BioRad). Quantification of mRNA levels was calculated relative to levels of the ribosomal protein gene *rp49* using the Pfaffl method (29). Three independent replicates were completed. See **Supplemental Table 1** for qPCR primer sequences.

**Hemolymph LC-MS**

Flies were challenged with heat-killed *M. luteus* to activate the Toll pathway. Hemolymph was extracted from 100 to 110 each of *w*1118, *Δbbd*, and BomΔ55C flies using the same method as in the hemolymph antimicrobial assays, with 50–60 flies processed per Zymo-Spin IC column (Zymo Research).

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1http://blogs.cornell.edu/drosophila/dssc-cornmeal-recipe/
and yielding a total of ~10 μl hemolymph per genotype. Three independent biological replicates were processed for ∆bbd and BomΔ55C, and two independent biological replicates were processed for w1118. Extracted hemolymph was mixed 1:1 (vol/vol) with denaturing buffer (8 M Urea, 50 mM Tris, pH 7.8, 150 mM NaCl, protease and phosphatase inhibitors) and protein concentration was determined using a BCA assay. For each sample, 40 μg of hemolymph was diluted to 1 M urea using 50 mM ammonium bicarbonate and digested overnight with trypsin (Promega, V511A) at a 1:100 (trypsin:protein) ratio. After digestion, peptides were reduced with 1 mM dithiothreitol at room temperature for 30 min and then alkylated with 5 mM iodoacetamide at room temperature in the dark for 30 min. Formic acid was added to a 0.1% final concentration and peptides were desalted using the C18-Stage-Tip method and then vacuum dried. The dried peptides were reconstituted in 5% formic acid/5% acetonitrile and 1 μg of total peptide for each sample was loaded for MS analysis. Samples were run in technical triplicates on a Q-Exactive mass spectrometer with instrument and chromatography settings as described previously (30), except for the following modifications: the RAW files were analyzed using Andromeda/MaxQuant (version 1.6.7.0) (31) with default settings (32) except the match between the run and LFQ quantitation settings was enabled for label free quantification. Data were searched against a concatenated target-decoy database comprised of forward and reversed sequences from the unreviewed UniprotKB/Swiss-Prot FASTA Drosophila database (2019). A mass accuracy of 20 ppm was assigned for the first search and 4.5 ppm for the main search. The statistical analysis was calculated using the DEP analysis R-package (33).

**Bacterial Load Quantification**

Bacterial load upon death (BLUD) was obtained as in Duneau et al. (34), with slight modifications. Briefly, flies were infected with E. faecalis and vials were monitored every 30 min for newly dead flies. These flies were then individually homogenized with a pestle in 400 μl LB media. Homogenates were also prepared from individual live w1118 flies 120 h post-infection (hpi). Homogenates were diluted serially in LB and spread on LB agar plates for incubation at 37°C overnight. Colonies were counted manually and the number of viable bacteria per fly was calculated. Data were obtained from three independent experiments.

**Data Analysis**

GraphPad Prism 5 was used for statistical tests. Survival data were plotted as Kaplan-Meier curves and were analyzed using the Gehan-Breslow-Wilcoxon test to determine statistical significance. Statistical differences in candidacidal activity were calculated using one-way ANOVA followed by Tukey’s test. Multiple Mann-Whitney U tests were used to calculate differences between BLUD samples (p = 0.0085 after Sidak correction for multiple comparisons, α = 0.05, k = 6). Spearman rank correlation was used to assess the relationship between BLUD and time of death.

**RESULTS**

**The bombardier Gene Is Specifically Required for Toll-Mediated Defense**

The bombardier (bbd) gene contains a consensus Toll-responsive NF-κB binding site within its promoter region and is strongly expressed upon Toll activation by Gram-positive bacterial infection or other inducers (14, 17, 35, 36). The encoded protein is predicted to be secreted and to generate a mature protein of 222 amino acids with a coiled coil near its C-terminus (37, 38). Orthologs of Bombardier are found across the Drosophila genus, but in no other genera (39).

We began our analysis of the bombardier gene by generating a null mutant, using CRISPR/Cas9 to delete 1,906 bp encompassing the annotated transcriptional unit. Flies homozygous for this deletion (hereafter ∆bbd) were viable and morphologically wild-type. Given that bombardier is Toll-inducible, we assayed ∆bbd flies for a potential loss-of-function phenotype in Toll-mediated immunity. Specifically, we infected adult ∆bbd flies with various pathogens and then monitored survival. Two additional genotypes were used as controls: w1118 flies, which served as the wild type, and BomΔ55C flies, which lack Toll-mediated humoral defenses due to deletion of the 12 Bom genes (24).

As shown in Figure 1, we observed a marked immunodeficiency when ∆bbd flies were challenged with representative species for the three classes of microbes against which Toll provides defense. With the yeast Candida glabrata, more than 90% of w1118, but no ∆bbd flies, survived 5 days after infection (Figure 1A). In the case of the filamentous fungus Fusarium oxysporum, 70% of w1118 adults, but fewer than 20% of ∆bbd adults, were alive 5 days post-infection (Figure 1B). Finally, with Enterococcus faecalis, a Gram-positive bacterium, 50% of wild-type flies, but no ∆bbd flies, were alive 5 days after infection (Figure 1C).

The impairment of Toll-mediated defenses by deletion of bombardier was significant for all three pathogens (p < 0.0001). In the case of C. glabrata, the immunodeficiency of ∆bbd phenocopied that observed for BomΔ55C flies (n.s., p > 0.05). In contrast, with either F. oxysporum or E. faecalis, the rate of death was greater for BomΔ55C than for ∆bbd (∗∗∗p < 0.0001 for both infections). The ∆bbd mutant thus displays a substantial, but not complete, loss of Toll-mediated defense.

The expression of bombardier is strongly induced by Toll, but not Imd activation (14). We therefore hypothesized that Imd-mediated defenses would not require bombardier function. To test this prediction, we infected ∆bbd flies with Enterobacter cloacae, a Gram-negative bacterium. In this experiment, ∆bbd flies were as immunocompetent as w1118 flies: more than 90% of both genotypes survived at least 5 days post-infection (Figure 1D). In contrast, 100% of Rel20 flies, which are deficient in Imd signaling (40), succumbed to infection within 1 day. Thus, bombardier functions in defense against a range of pathogens for which Toll mediates defense—yeast, filamentous fungi, and Lys-type Gram-positive bacteria—but not against Gram-negative bacteria, against which the Imd pathway is active.
The Candidacidal Activity of Hemolymph Requires Bombardier, but Neither Drosomycin Nor Metchnikowin

Next, we investigated the potential humoral role of Bombardier by preparing and assaying cell-free hemolymph. We have previously shown that hemolymph from wild-type flies exhibits a Toll-dependent and Bomanin-dependent candidacidal activity (25). However, we were also curious as to the identity of the active antifungal component. In particular, we considered the potential role of Metchnikowin (Mtk) and Drosomycin (Drs), two antimicrobial peptides (AMPs) that have documented antifungal activity in vitro and are strongly Toll-induced in vivo (14, 41, 42). We therefore took advantage of the recently described ΔAMPs strain, which is deficient for Mtk and Drs, as well as all other induced AMPs other than the Cecropins (23). Extracting and assaying Toll-induced hemolymph, we found that hemolymph from ΔAMPs flies had a killing activity against C. glabrata comparable to that of wild-type hemolymph (Figure 2). In contrast, we failed to detect any killing of C. glabrata by Δbbd hemolymph. We conclude that Boms and Bombardier, but neither Mtk nor Drs, are required for humoral defense against C. glabrata.

Short-Form Bom Peptides Are Specifically Absent From Δbbd Hemolymph

MALDI-TOF provides a robust tool for characterizing small (<5,000 MW) peptides present in hemolymph after Toll activation. As shown in Figures 3A,B, such a readout includes the aforementioned AMPs (Mtk and Drs), several short-form Boms (BomS1, S2, S3, and S6; see Supplemental Table 2 for updated Bomanin nomenclature), and other induced peptides (e.g., IM4). We have previously shown that deleting the 55C Bom gene cluster removes the peaks attributable to the short-form Boms, while leaving the remaining signals unaffected (25). Remarkably, analysis of Δbbd hemolymph yielded a similar pattern. As shown in Figures 3C,D, the short-form Boms that were readily detectable in the wild type—S1, S2, S3, and S6—were absent in Δbbd hemolymph, whereas the remaining peptides, including Mtk, Drs, and IM4, displayed a wild-type profile.

Although Δbbd disrupts the accumulation of short-form Bom peptides in hemolymph, this effect does not reflect a disruption in transcription or stability of the corresponding Bom mRNAs: robust induction of Toll-regulated genes, including genes of short-form Boms, was readily detectable with qRT-PCR (Supplemental Figure 1).
Because proteins such as Bombardier and bicipital Boms are too large to be detected by our MALDI-TOF protocol, we used LC-MS to further characterize the relationship between Bombardier and the Boms in hemolymph. For these studies, we prepared Toll-induced hemolymph from three genotypes: \( w^{118} \), \( \Delta bbd \), and \( Bom^{\Delta 55C} \). In wild-type hemolymph, we readily detected Bombardier protein (Figure 4), consistent with the presence of a canonical secretion signal sequence in the Bombardier coding sequence. Bombardier, like the Boms, is thus secreted into hemolymph upon Toll induction. We also detected all three bicipital Boms—BomBc1, BomBc2, and BomBc3. The LC-MS studies thus complemented the MALDI-TOF studies, with bicipital Boms detected by the former and short-form Boms by the latter (tailed Boms are not detected by either protocol). Next, we assayed \( \Delta bbd \) hemolymph. As expected, Bombardier was not detected. However, the three bicipital Boms were present at comparable levels in wild-type and \( \Delta bbd \) hemolymph (see Figure 4). Combined with the MALDI-TOF studies, these results demonstrate that \( \Delta bbd \) blocks accumulation in hemolymph of short-form, but not bicipital, Boms. Lastly, we analyzed hemolymph from \( Bom^{\Delta 55C} \) flies, which lack 10 of the 12 Bom genes. As expected, the products of the two deleted bicipital genes (BomBc1 and BomBc2) were absent, whereas the product of the remaining bicipital gene (BomBc3) was present at wild-type levels (see Figure 4). Turning our attention to Bombardier, we observed no effect of the 55C Bom deletion. Thus, Bombardier is required for the presence of short-form Boms in hemolymph, but the 55C Boms are not required for the presence of Bombardier.

**Bombardier Mediates Both Infection Resistance and Tolerance**

The \( \Delta bbd \) survival phenotype could be due to an inability to control pathogen growth—a defect in resistance—or an inability to endure infection—a defect in tolerance. Because flies lacking Bombardier demonstrate an increased susceptibility to infection and decreased levels of known resistance factors, the short-form Boms, it seemed likely that \( \Delta bbd \) flies, like \( Bom^{\Delta 55C} \) flies, have a defect in infection resistance. In exploring this hypothesis, we found that the model recently developed by Duneau et al. provided a useful framework (34). Following infection of an individual fly, there are two stereotyped outcomes: either the pathogen replicates, reaches a lethal burden, and the fly dies; or the pathogen is controlled at a level below the lethal burden and the fly survives with a persistent infection. Variation in survival curves for different pathogens and fly genotypes reflects variation in both the time required to reach lethal burden and in the fraction of flies that are able to control the infection before it reaches such a threshold. In cases where a fraction of flies control infection, group survival typically drops after infection and then reaches a plateau (23).

The survival curve for \( \Delta bbd \) flies infected with *E. faecalis* does not plateau (see Figure 1C). Instead, it exhibits a profile that we hypothesize reflects two phases of death. In the first phase, extending roughly 2 days post-infection, some \( \Delta bbd \) flies reach a lethal burden of *E. faecalis* and die, as reflected in a sharp decline in survival; the remainder control the infection. In the second phase, from 2.5 days onward, those flies with a persistent infection die at a reduced but steady rate, due to a defect in tolerance. If this hypothesis is correct, flies dying in the first phase should have a bacterial load upon death (BLUD) comparable to that of wild-type flies dying from infection. Furthermore, those dying in the second phase should have a much lower pathogen burden, comparable to that of wild-type survivors with a persistent infection.

To test our predictions regarding pathogen burden, we measured the BLUD of individual flies after infection with live *E. faecalis* and divided the data into two time intervals (Figure 5). For the earlier interval (dead flies obtained between 17 and 51.5 hpi), both \( Bom^{\Delta 55C} \) and \( \Delta bbd \) bacterial loads upon death were not significantly different from \( w^{118} \) (Figure 5, red, \( p > 0.05 \)). For the later time interval (flies obtained between 68 and 120.5 hpi), \( \Delta bbd \) flies perished at significantly lower bacterial loads compared to that of \( \Delta bbd \) flies which died earlier (Figure 5, \( \Delta bbd \) early compared to \( \Delta bbd \) late, \( p < 0.0001 \)), indicating that these two groups die from distinct causes. Importantly, late-death \( \Delta bbd \) flies perished at significantly lower bacterial loads than those of \( w^{118} \) suffering early deaths (\( p < 0.0001 \)), demonstrating that \( \Delta bbd \) flies have a defect in tolerance.

Together, the survival curve and BLUD data offer strong support for our two-phase-model: \( \Delta bbd \) flies died early in infection with high bacterial loads, due to a defect in resistance, and died later with lower bacterial loads, reflecting a deficiency in tolerance. However, we note that the bacterial loads of
Δbbd flies dying in the later phase were still significantly greater than those of w^1118 flies alive 120 hpi (Figure 5, Δbbd late compared to live w^1118, p < 0.0001). This indicates that the later-death Δbbd group has not completely controlled infection compared to the live w^1118 flies, and suggests that both resistance and tolerance contribute to the later Δbbd fly deaths. Although we cannot rule out a minor resurgence in bacterial proliferation preceding late death of bbd flies, we note that BLUD and time of death were not significantly correlated for these flies (Supplemental Figure 2, Spearman correlation test, r = −0.2654, p = 0.1564).

**Immune Activation, Specifically Bom Expression, Is Deleterious in the Absence of Bombardier**

What is the nature of the tolerance defect we observed in Δbbd flies? More specifically, is their health impaired by an excessive or toxic immune response, or is death due to another class of impaired tolerance (43)? To distinguish between these explanations, we assayed the effect of activating the immune response in Δbbd flies in the absence of infection.

When Δbbd flies were challenged with heat-killed E. faecalis, we observed a decrease in survival that first was apparent 3 days post-challenge followed by a steady decline in the number of live flies in the following days (Figure 6A), consistent with the timing of the late-phase deaths (see Figure 5). Overall, the death rate was slower than that of live infection, but the extent of killing was similar between heat-killed and live E. faecalis: fewer than 20% of flies survived (compare Figures 1C, 6A). In contrast, no effect on survival was observed upon challenge of either w^1118 or Bom^Δ55C flies with heat-killed E. faecalis: >95% flies survived seven or more days post-challenge.

The effect of immune stimulation on Δbbd survival was not specific to E. faecalis. When we repeated the challenge experiments with heat-killed Micrococcus luteus, which activates the Toll response [see Supplemental Figure 1, as well as (25, 44)], the effect on Δbbd survival was again marked: 5 days after challenge, fewer than 5% of Δbbd flies were alive, compared to

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**FIGURE 3** | Short-form Bom peptides are specifically absent in hemolymph from Δbbd flies. MALDI-TOF mass spectra of w^1118 (A,B) and Δbbd (C,D) hemolymph samples were collected in linear (A,C) and reflectron mode (B,D). For peptide identification, peaks were matched to those of corresponding peaks in prior studies (13, 25). Spectra were obtained from at least ten independent biological replicates and representative spectra are shown (a.u., arbitrary units; m/z, mass/charge).

**FIGURE 4** | The presence of bicipital Bomarins in hemolymph is unaffected by loss of Bombardier. MS/MS counts for the indicated proteins as determined by Andromeda/MaxQuant were normalized to total MS/MS counts in each run. Error bars represent standard deviation for biological replicates (n = 3 for Δbbd and Bom^Δ55C, n = 2 for w^1118). Supplemental Table 3 shows the full dataset.
survival of >95% of w^{118} and 85% of Bom^{Δ55C} flies over the same period of time (Figure 6B).

As both M. luteus and E. faecalis induce the Toll pathway, Toll activation could be the key factor in Δbbd mortality. To address this hypothesis, Δbbd flies were crossed with MyD88^{ken1} (Toll-deficient) flies to generate the MyD88^{ken1} Δbbd double mutant, and the resulting flies were challenged with heat-killed E. faecalis and M. luteus. Unlike Δbbd flies, MyD88^{ken1} Δbbd flies survived challenge with Toll activators (Figures 7A,B). Because blocking the Toll pathway with MyD88^{ken1} rescues the Δbbd phenotype triggered by heat-killed bacteria (p < 0.0001 compared to Δbbd, p > 0.05 compared to MyD88^{ken1} for both heat-killed bacteria), we conclude that Toll activation underlies the death of Δbbd flies in the absence of infection.

As described above, Bom genes are transcribed in Δbbd flies (Supplemental Figure 1), but short-form Bom peptides do not appear in hemolymph (Figure 3). This suggests a mislocalization of these peptides, perhaps in an unprocessed or misfolded state. Given that short-form Bom genes are among the most abundantly transcribed genes after infection (17, 36), such mislocalized or misfolded Boms could rapidly accumulate to high levels in Δbbd flies. Could this explain the death of Δbbd flies upon immune stimulation? To address this question, we generated Bom^{Δ55C} Δbbd double mutants and assayed the effect of immune induction alongside both Bom^{Δ55C} and Δbbd flies (Figures 7A,B). The result was unequivocal: introducing Bom^{Δ55C}, which deletes all of the short-form Boms, eliminated the effect of Δbbd on survival following immune stimulation (p < 0.0001 compared to Δbbd, p > 0.05 compared to Bom^{Δ55C} for both heat-killed bacteria). The fact that Bom^{Δ55C} is epistatic to Δbbd demonstrates that Toll-driven expression of Bom genes is specifically responsible for the death of immune-stimulated Δbbd flies.

**DISCUSSION**

The results presented in this study identify a key factor that regulates humoral and Bom-mediated defense in Drosophila. We demonstrate that Δbbd flies are defective in resistance to pathogens controlled by the Toll pathway. The results support the hypothesis that this defect results from the absence of short-form Boms in Δbbd hemolymph. Absence of Boms is sufficient to cause a defect in resistance (24) and Δbbd hemolymph appears to be lacking the short-form Boms but no other component, save Bombardier itself. Furthermore, Δbbd phenocopies Bom^{Δ55C} with regard to survival after C. glabrata infection, and resistance to C. glabrata can be restored in Bom^{Δ55C} flies by expression of short-form Boms (25). Finally, Δbbd hemolymph lacks candidacidal activity, which is dependent on short-form Bom peptides (25) and which we show here does not require Drs or Mtk.

For pathogens other than C. glabrata, the effect of deleting Bombardier is less severe than that of deleting the ten Bom genes clustered at 55C. Our mass spectrometry data suggest an explanation. Whereas, short-form Boms are absent from Δbbd hemolymph, bicipital Boms are present. (Tailed Boms were not detected with either mass spectrometry method.) Therefore, we postulate that the bicipital Boms, which are not required for resistance to C. glabrata (25), are functional against other pathogens. This would explain why Δbbd flies are more resistant than Bom^{Δ55C} flies upon infection with E. faecalis or F. oxysporum (Figure 1). In this regard, we note that Bombardier and all three forms of Bom proteins—short, tailed, and bicipital—are found across the Drosophila genus, supporting the notion that all three classes of Boms are immunoprotective and therefore maintained across the Drosophila genus.

It might appear that our discovery of Bombardier was serendipitous, given our role in defining the Bomanin gene family (24, 25). In hindsight, however, the link was forged in our approach. We selected CG18067 from the most strongly inducible Toll-regulated loci, a group that also includes eight of the Bomanin genes. Next, we engineered a CRISPR/Cas9 deletion of CG18067 and assayed this knockout with the identical set of pathogens that we had used for the Bom^{Δ55C} deletion, screening for loss of survival upon infection. Having examined a gene that is as strongly induced as the Bomanins, present in the same range of species as the Bomanins, and with a spectrum of loss-of-function phenotypes similar to that of the Bomanins, it is not particularly surprising that we would find ourselves studying a gene that affects the Bomanins.

**Bombardier Function and Structure**

What is the function of Bombardier? Deleting the gene results in the absence of short-form Boms from hemolymph, an effect
we find is at the level of protein. Other mature immune peptides are present at normal levels in the hemolymph, and there is thus no general defect in translation, secretion, or processing. Based on these findings, we propose that Bombardier normally functions either to chaperone short Boms as they are secreted from the fat body into the hemolymph or, alternatively, to protect the Boms from misfolding or aggregation while in the hemolymph. We further hypothesize that it is the ectopic localization or aberrant form of short-form Boms in \( \Delta bbd \) flies that generates morbidity upon Toll pathway activation. In support of this idea, we showed that Bom expression underlies the lethality observed in \( \Delta bbd \) flies (Figure 7). Whether the short-form Boms physically interact with Bombardier, perhaps in the context of a larger antimicrobial complex, is currently unknown.

Activation of Toll-like receptor (TLR) signaling is important for innate immunity, but induction of the pathway can lead to autoimmune disorders and chronic inflammatory disease (45–48). Here we report an autoimmune activity driven by Toll-induced Bom expression in flies lacking a downstream pathway component, Bombardier. To what extent this parallel can be exploited in the context of understanding autoimmune disorders promises to be a significant focus for future investigation.

DATA AVAILABILITY STATEMENT

The LC-MS datasets generated for this study can be accessed on the MassIVE data repository using the accession identifier: MSV000084509.
AUTHOR CONTRIBUTIONS

SL and SW: conceptualization, writing—original draft, and visualization. SL, AF, and EB: methodology. SL and AF: formal analysis. SL, AF, and LC: investigation. SL, AF, LC, EB, and SW: writing—review and editing. EB and SW: supervision and funding acquisition.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2019.03040/full#supplementary-material
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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