Nephrocytes Remove Microbiota-Derived Peptidoglycan from Systemic Circulation to Maintain Immune Homeostasis

Graphical Abstract

Highlights

- *Klf15* flies display increased resistance to infection and a shorter lifespan
- *Klf15* flies show systemic PGN accumulation and Toll pathway activation
- Germ-free conditions rescue the pathology of *Klf15* flies
- Nephrocytes uptake microbiota-derived PGN and degrade it inside lysosomes

Authors

Katia Troha, Peter Nagy, Andrew Pivovar, Brian P. Lazzaro, Paul S. Hartley, Nicolas Buchon

Correspondence

nicolas.buchon@cornell.edu

In Brief

Troha et al. reveal that renal filtration of microbiota-derived peptidoglycan at steady state prevents aberrant immune activation, thus maintaining immune homeostasis in *Drosophila*. This function is likely conserved in mammals, with relevance to the chronic immune activation seen in settings of impaired blood filtration.
Nephrocytes Remove Microbiota-Derived Peptidoglycan from Systemic Circulation to Maintain Immune Homeostasis

Katia Troha,1 Peter Nagy,1 Andrew Pivovar,1 Brian P. Lazzaro,1 Paul S. Hartley,2 and Nicolas Buchon1,3,*
1Cornell Institute of Host-Microbe Interactions and Disease, Department of Entomology, Cornell University, Ithaca, NY, USA
2Department of Life and Environmental Science, University of Bournemouth, Talbot Campus, Poole, Dorset BH12 5BB, UK
3Lead Contact
*Correspondence: nicolas.buchon@cornell.edu
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SUMMARY

Preventing aberrant immune responses against the microbiota is essential for the health of the host. Microbiota-shed pathogen-associated molecular patterns translocate from the gut lumen into systemic circulation. Here, we examined the role of hemolymph (insect blood) filtration in regulating systemic responses to microbiota-derived peptidoglycan. Drosophila deficient for the transcription factor Klf15 (Klf15NN) are viable but lack nephrocytes—cells structurally and functionally homologous to the glomerular podocytes of the kidney. We found that Klf15NN flies were more resistant to infection than wild-type (WT) counterparts but exhibited a shortened lifespan. This was associated with constitutive Toll pathway activation triggered by excess peptidoglycan circulating in Klf15NN flies. In WT flies, peptidoglycan was removed from systemic circulation by nephrocytes through endocytosis and subsequent lysosomal degradation. Thus, renal filtration of microbiota-derived peptidoglycan maintains immune homeostasis in Drosophila, a function likely conserved in mammals and potentially relevant to the chronic immune activation seen in settings of impaired blood filtration.

INTRODUCTION

As the first line of defense against invading microorganisms, the innate immune system senses and responds to both pathogen-associated molecular patterns (PAMPs) and damage-associated molecular patterns (DAMPs) (Newton and Dixit, 2012). Peptidoglycan (PGN), a major constituent of the microbial cell wall, is an immune-stimulatory PAMP found in all bacteria. Following infection, detection of PAMPs in systemic circulation triggers a cascade of immune reactions that, if uncontrolled, can ultimately lead to sepsis (Cecconi et al., 2018). Even in the absence of infection, organisms are not a sterile environment—the animal gut and other mucosal tissues harbor numerous microbial species, which constitute the microbiota. A consequence of the presence of microbiota is the translocation of microbiota-shed PAMPs from the gut lumen into systemic circulation. This is a phenomenon that has been documented in a variety of organisms, including Drosophila and mammals (Capo et al., 2017; Clarke et al., 2010; Corbitt et al., 2013; Gendrin et al., 2009; Paredes et al., 2011; Zaidman-Rémy et al., 2006).

Drosophila is a powerful system to study innate immune responses to both pathogens and gut microbes (Buchon et al., 2014; Liu et al., 2017). To resist infection, Drosophila relies on both cellular and humoral innate immune responses. The cellular response consists of encapsulation and phagocytosis, while the humoral response involves the melanization cascade and the synthesis of antimicrobial peptides (AMPs) by the fat body, an organ analogous to the liver and adipose tissue of mammals. Production of AMPs is controlled by two principal signaling cascades: the Toll and Imd pathways. Both pathways are activated in response to PGN: Lys-type PGN from Gram-positive bacteria triggers the Toll pathway, while DAP-type PGN from Gram-negative bacteria and certain Gram-positiv bacilli induces the Imd pathway (Buchon et al., 2014). In the fly, PGN is detected by peptidoglycan recognition proteins (PGRPs). PGRP-LC and PGRP-LE sense DAP-type PGN, and PGRP-SA recognizes Lys-type PGN (Kaneko et al., 2006; Michel et al., 2001). Drosophila possesses several immune mechanisms to both shape the microbiota and prevent excessive immune responses upon detection of microbial stimuli (Basmous et al., 2011; Cao et al., 2013; De Gregorio et al., 2002; Gordon et al., 2008; Levashina et al., 1999; Maillet et al., 2008; Scherfer et al., 2008; Thevenon et al., 2009). In the case of the Imd pathway, these mechanisms include negative regulators that avert excessive immune activation in response to the microbiota. For instance, the transcription factor Caudal suppresses Imd-dependent expression of AMPs in the gut, thereby shaping microbiota composition (Ryu et al., 2008). Another regulator, Pirk, sequesters PGRP-LC to prevent its exposure to microbiota-derived PGN and subsequent activation of the Imd pathway (Agarwal et al., 2008; Klein et al., 2008; Lhocine et al., 2008). Finally, secreted PGRPs with amidase activity scavenge and degrade immunostimulatory DAP-type PGN in order to block Imd activation (Paredes et al., 2011). A notable example of this is PGRP-LB, which is released into the hemolymph (extracellular fluid analogous to blood) to degrade translocated, microbiota-derived PGN, thus preventing systemic immune activation (Paredes et al., 2011; Zaidman-Rémy et al., 2006). Although the Toll...
pathway also responds to the presence of PGN, little is known about the mechanisms that suppress Toll activation in response to microbiota and whether these have a role in the regulation of systemic responses.

The excretory system of *Drosophila* is composed of nephrocytes (which regulate hemolymph composition by filtration followed by filtrate endocytosis) and Malpighian tubules (which modify and secrete urine) (Denholm and Skaer, 2009; Hartley et al., 2016). *Drosophila* nephrocytes can be divided into two distinct groups: the garland cells, which appear as a necklace-like structure surrounding the esophagus, and the pericardial cells that form two rows of cells flanking the heart (Aggarwal and King, 1967; Crossley, 1972; Na and Cagan, 2013). In the adult stage, pericardial nephrocytes serve as the primary filtration units (Zhang et al., 2013). Hemolymph filtration occurs in a stepwise manner. First, hemolymph is filtered across the nephrocytes’ negatively charged basement membrane and a specialized filter known as the nephrocyte diaphragm. The filtrate then enters the lacunae, also known as the labyrinthine channels, which extend several microns into the nephrocyte’s cortical region (Kosaka and Ikeda, 1983). It is in these chambers where the filtrate is finally endocytosed by nephrocytes (Denholm and Skaer, 2009). Nephrocytes possess significant molecular, anatomical, and functional similarities to the glomerular podocyte, a cell type of the mammalian kidney important for the kidney’s filtration function (Weavers et al., 2009; Zhuang et al., 2016). These flies thus enable study of the impact of hemolymph filtration in the maintenance of immune homeostasis.

Here, we found that flies devoid of nephrocytes (*Klf15<sup>NN</sup>* null allele), or with diminished nephrocyte function, were more resistant to a variety of microbial infections. *Klf15<sup>NN</sup>* flies exhibited improved survival upon infection but also a shorter lifespan stemming from abnormal Toll pathway activation. Aberrant Toll signaling in *Klf15<sup>NN</sup>* flies was dependent on the presence of Lys-type PGN microbiota. Microbiota-derived PGN accumulated in the hemolymph of these flies, triggering chronic stimulation of the Toll pathway. In wild-type (WT) flies, microbiota-derived Lys-type PGN found in systemic circulation was taken up by nephrocytes via endocytosis and degraded within lysosomes. Thus, renal filtration of microbiota-derived PGN maintains immune homeostasis in *Drosophila*, a function likely conserved in mammals.

RESULTS

**Klf15<sup>NN</sup>** Flies Are Less Susceptible to Microbial Infection

In order to evaluate the role of hemolymph filtration in immune function and homeostasis, we turned to flies that are mutant for the transcription factor *Klf15* (*Klf15<sup>NN</sup>* null allele), which lack nephrocytes. First, we confirmed that *Klf15<sup>NN</sup>* flies fail to develop nephrocytes (Figure 1A; Ivy et al., 2015). *Klf15<sup>NN</sup>* flies exhibited a significantly shorter basal lifespan compared to WT controls of the same genetic background (Figure 1B). Despite having a curtailed life expectancy, *Klf15<sup>NN</sup>* flies survived sterile wounding comparably to WT animals (Figures 1C–1G). To determine whether immune competence was affected by the loss of hemolymph filtration, we conducted survival assays with *Klf15<sup>NN</sup>* flies following systemic infection with the bacterial pathogens *Serratia marcescens* type strain, *Salmonella typhimurium*, *Listeria innocua*, *Enterococcus faecalis*, *Staphylococcus aureus*, and *Providencia rettgeri*. *Klf15<sup>NN</sup>* flies displayed significantly increased survival against five of these infections (Figures 1C–1G). The only exception was infection with *P. rettgeri*, to which the mutant proved more sensitive (Figure S1A). We did not observe a survival phenotype following challenge with two fungal agents, *Metarhizium anisopliae* and *Beauveria bassiana* (Figures S1B and S1C). Overall, these results suggest that *Klf15<sup>NN</sup>* flies are broadly protected against systemic infection by bacterial pathogens.

To verify that the enhanced survival observed in *Klf15<sup>NN</sup>* flies was a direct consequence of the loss of nephrocytes, we generated nephrocyte-deficient flies through complementary means. Using the nephrocyte-specific driver *Dot-Gal4*, we decreased *Klf15* expression by *in vivo* RNAi throughout development (*Dot-Gal4* > *UAS-Klf15-IR*), which results in adult flies lacking nephrocytes (Ivy et al., 2015). Upon infection with *S. aureus* and *E. faecalis*, these flies displayed increased survival relative to the WT controls (Figures 1H and S1D). Next, we set out to determine whether the survival phenotype of *Klf15<sup>NN</sup>* flies resulted from loss of hemolymph filtration in the mutant or from a developmental defect associated with the loss of nephrocytes. To distinguish between the two possibilities, we took advantage of the fact that adult-specific loss of *Klf15* halts the endocytic function of mature nephrocytes (Ivy et al., 2015). We decreased *Klf15* expression by *in vivo* RNAi specifically during the adult stage using the conditional, nephrocyte- and heart-specific driver *Hand-Gal4<sup>ts</sup>2* (*Hand-Gal4<sup>ts</sup>2* > *UAS-Klf15-IR*) and infected these flies separately with *S. aureus* and *E. faecalis*. Diminishing the endocytic competence of adult nephrocytes by decreasing *Klf15* expression was sufficient to increase survival to infection with both bacteria (Figures 1I and S1E). Altogether, our results support the conclusion that loss of nephrocyte function generally increases survival against microbial infection.

**Klf15<sup>NN</sup>** Flies Are More Resistant to Infection, Independent of Phagocytosis and Melanization

Multicellular organisms employ two complementary strategies to combat infection: resistance, to eliminate microbes; and disease tolerance, to allow them to withstand the infection and/or its deleterious consequences (Ayres and Schneider, 2012). To determine whether the improved survival of *Klf15<sup>NN</sup>* flies was due to an increase in disease tolerance, we compared the bacterial load upon death (BLUD) of WT and *Klf15<sup>NN</sup>* flies following infection with *S. aureus* and *E. faecalis*. BLUD represents the maximal quantity of bacteria that an infected fly can sustain before it dies (Duneau et al., 2017) and is therefore one measure of disease tolerance. We found that control flies and *Klf15<sup>NN</sup>* flies die carrying similar numbers of each bacterium tested.
indicating that this marker of disease tolerance is not altered in the mutant. Next, we tested whether the survival advantage of the mutant stemmed from improved resistance to infection. We monitored bacterial load during the course of *S. aureus*, *E. faecalis*, *S. marcescens*, *L. innocua*, and *S. typhimurium* infections (Figures 2A, 2B, and S2B–S2D). *Klf15NN* flies carried significantly lower bacterial burdens than WT flies as soon as 3 h post-infection with *S. aureus*, 4.5 h after challenge with *E. faecalis*, and 6 h post-inoculation with *S. marcescens*, *L. innocua*, and *S. typhimurium*, demonstrating that flies without nephrocytes are more resistant to pathogens in the early stages of infection.

*Figure S2A*, indicating that this marker of disease tolerance is not altered in the mutant. Next, we tested whether the survival advantage of the mutant stemmed from improved resistance to infection. We monitored bacterial load during the course of *S. aureus*, *E. faecalis*, *S. marcescens*, *L. innocua*, and *S. typhimurium* infections (Figures 2A, 2B, and S2B–S2D). *Klf15NN* flies carried significantly lower bacterial burdens than WT flies as soon as 3 h post-infection with *S. aureus*, 4.5 h after challenge with *E. faecalis*, and 6 h post-inoculation with *S. marcescens*, *L. innocua*, and *S. typhimurium*, demonstrating that flies without nephrocytes are more resistant to pathogens in the early stages of infection.

*Figure 1. Loss of Nephrocyte Function Increases Survival against Infection*

(A) Adult pericardial nephrocytes stained with WGA Alexa Fluor 594 conjugate (red). Phalloidin-FITC (green) marks the heart tube. Staining is shown for both wild-type (WT) and *Klf15NN* flies.

(B) Lifespan curve comparing WT to *Klf15NN* flies.

(C–G) Survival curves over 7 days following infection of WT and *Klf15NN* flies with the bacterial pathogens: *S. marcescens* Type strain (C), *S. typhimurium* (D), *L. innocua* (E), *E. faecalis* (F), and *S. aureus* (G).

(H and I) Survival of flies expressing nephrocyte-specific RNAi against *Klf15* throughout development (*Dot-Gal4 > UAS-Klf15-IR*) (H) or only during the adult stage (*Hand-Gal4ts > UAS-Klf15-IR*) (I) after infection with *S. aureus*. The curves represent the average percent survival ± SE of three or more biological replicates. **p < 0.01 ***p < 0.001 ****p < 0.0001 in a Log-rank test.

*Drosophila* relies primarily on three effector mechanisms to control bacterial growth: phagocytosis, melanization, and the production of AMPs. First, we evaluated a role for phagocytosis in the resistance phenotype of *Klf15NN* flies. We injected nephrocyte-deficient and WT flies with pH-sensitive pHrodo bacteria, which become fluorescent only after being engulfed into a fully mature, acidified phagosome (Guillou et al., 2016). After quantification, we observed close to 50% less fluorescence in *Klf15NN* flies relative to controls (Figure S2E). Moreover, injection of flies with latex beads prior to systemic infection with both *S. aureus* and *E. faecalis*, a treatment that blocks phagocytosis (Eirod-Erickson et al., 2000), did not alter
the survival phenotype of Klf15NN flies (Figures S2F and S2G). These results demonstrate that phagocytic activity does not contribute meaningfully to the increased resistance of Klf15NN flies. Assessment of phenoloxidase (PO) activity, the terminal enzymatic step driving melanization, revealed that while Klf15NN flies had significantly higher PO activity in basal conditions, they also displayed significantly lower PO activity relative to controls 3 h post infection with S. aureus and E. faecalis (Figure S2H). To clarify whether melanization played any role in the survival phenotype of Klf15NN flies, we generated a mutant deficient for both Klf15 and key genes required for the melanization response (PPO1-D, D,31) (Binggeli et al., 2014; Dudzic et al., 2015). Upon infection with S. aureus, the quadruple mutant (Klf15NN; PPO1-D, D,31) exhibited improved survival relative to the triple mutant (PPO1-D, D,31) (Figure S2I), suggesting that melanization is not required for the protection observed in nephrocyte-deficient flies. In sum, our data indicate that loss of nephrocytes confers increased resistance to hosts independent of phagocytosis and melanization.

The Toll Pathway Is Constitutively Active in Klf15NN Flies

Nephrocytes are major regulators of hemolymph content via filtration followed by filtrate endocytosis (Hartley et al., 2016; Soukup et al., 2009). Therefore, we considered whether changes in circulating proteins in the mutant could account for the increased resistance observed in Klf15NN flies. Previously, we performed a proteomic analysis of hemolymph composition in both WT and Klf15NN unchallenged flies (Hartley et al., 2016). An in-depth analysis of this dataset revealed that among 130 proteins enriched (≥1.5-fold) or detected only in the hemolymph of nephrocyte-deficient mutants, 65 proteins were annotated as having an immune-related function (Figure 2C). All 65 proteins were predicted to have a signal sequence (SignalP 4.1), which is expected for secreted hemolymph proteins. Of these 65 proteins, 19 were encoded by core genes of the Drosophila immune response (i.e., genes with increased transcription in response to most bacterial infections [Troha et al., 2018]), 30 were the products of genes that are induced only by a subset of microbial infections, and 16 were coded by genes that, while not regulated in response to infection themselves,
have been ascribed an immune function. We also noted that a majority (33 of 65) of these proteins are known targets of the Toll pathway (e.g., the antimicrobial peptide genes IM2, IM23, and CG15067). Thus, the hemolymph of Klf15NN flies is enriched in proteins of immune function primarily encoded by target genes of the Toll pathway, suggesting that changes in Toll pathway activity may explain the increase in resistance to pathogens observed in Klf15NN flies.

We developed two competing hypotheses to explain the accumulation of Toll pathway targets in the hemolymph of Klf15NN flies. The first hypothesis posited that because nephrocytes are critical regulators of protein turnover in the hemolymph, the rise in immune effectors could be the result of a decrease in protein turnover in these flies. Alternatively, the accumulation of immune gene products could be due to aberrant activation of the Toll and/or Imd pathways in nephrocyte-deficient flies. In agreement with the latter hypothesis, our proteomic analysis also identified proteins that were depleted (≥ 1.5-fold) in the hemolymph of Klf15NN flies relative to controls (Figure S2J). Six of these proteins are encoded by genes that typically show decreased transcription in response to bacterial infection in a Toll-dependent manner (e.g., Lsp1β and CG2233) (Troha et al., 2018), arguing that changes in hemolymph protein content are due to Toll pathway activation rather than protein turnover. To test this idea directly, we surveyed the activation of the Toll and Imd pathways by measuring the mRNA expression of 5 Toll target genes and 4 Imd target genes in WT and Klf15NN flies under basal conditions. qRT-PCR data from whole fly showed that the mRNA expression of all 5 Toll target genes—IM2, CG15067, Drs, CG18067, and CG15293—was significantly increased in Klf15NN flies compared to controls (Figures 2D and S2K). In contrast, we did not find any appreciable differences in gene expression between WT and Klf15NN flies for the Imd target genes Dpt, AttC, and TotA; the exception was AttD, for which the mutant had significantly lower mRNA expression relative to WT (Figures 2E and S2K). These data indicate that the Toll pathway, but not the Imd pathway, is constitutively activated in Klf15NN flies in unchallenged conditions. In agreement with these data, we also detected abnormal Toll activation in flies in which Klf15 expression was decreased by in vivo RNAi, specifically during the adult stage (Hand-Gal4> UAS-Klf15-IR), demonstrating that the loss of nephrocyte scavenging function is solely responsible for Toll activation in these flies (Figure 2F). In contrast to our results in unchallenged conditions, infection of Klf15NN flies with either S. aureus or E. faecalis revealed no significant differences between WT and mutant in terms of Toll or Imd target gene expression at any of the time points surveyed (3, 8, and 12 h post challenge) (Figures S2L and S2M). In conclusion, our data establish that Klf15NN flies present elevated basal mRNA expression of Toll target genes in conjunction with increased immune resistance to pathogens.

Increased Pathogen Resistance in Klf15NN Flies Is Contingent on Higher Baseline Toll Activity

Next, we asked whether Toll pathway activity could be responsible for the increased resistance observed in Klf15NN flies. We began by verifying that the increase in baseline Toll target gene expression was dependent on the Toll pathway itself. qRT-PCR of the Toll target genes IM2 (Figure 3A), CG15067, and Drs (Figures S3A and S3B) showed that a null mutation in the gene coding for the Toll cytokine Spz completely abolished the increase in Toll target gene expression found in Klf15NN flies (Klf15NN; spzmut double mutants). This was also true for a null mutation in the gene coding for SPE (Klf15NN; SPEmut), a key enzyme involved in the maturation of Spz and subsequent activation of the Toll pathway (Figures S3C–S3E), demonstrating that the increase in Toll target gene expression in Klf15NN flies is due to elevated Toll pathway activity. Notably, suppression of the Toll pathway by either spzmut or SPEmut completely abrogated the survival advantage of Klf15NN flies against pathogenic infection (Figures 3B, 3C, 3F, and 3G). These results indicate that a surge in Toll pathway signaling is directly accountable for the augmented resistance of Klf15NN flies to infection.

The Toll pathway can be induced by endogenous DAMPs, which trigger the maturation of the circulating serine protease Persephone (Psh), or by the recognition of PAMPs, leading to the activation of the serine protease ModSP (Buchon et al., 2009; Gottar et al., 2006; Ming et al., 2014). Consequently, we set out to investigate whether aberrant Toll signaling in Klf15NN flies was dependent on the detection of DAMPs or PAMPs by the host. While a null allele of psh was unable to rescue the elevated basal expression of the Toll target genes IM2 (Figure 3D), CG15067, and Drs (Figures S3H and S3I) in Klf15NN flies (Klf15NN; pshh), a null mutation in modSP fully reverted this increase in Klf15NN; modSP flies (Figures 3G, 3J, and 3K). Accordingly, while the improved survival phenotype of Klf15NN flies was still present in Klf15NN; pshh flies (Figures 3E and 3F), Klf15NN; modSP flies no longer exhibited it (Figures 3H and 3I). Thus, our data support the conclusion that elevated Toll signaling in response to PAMPs is responsible for the Klf15NN phenotype.

ModSP activity, and therefore Toll pathway signaling, can be induced by the binding of pattern recognition receptors (PRRs) to two types of PAMPs: β-(1,3)-glucan derived from the fungal cell wall is recognized by GNBP3 (Gottar et al., 2006), while bacterial PGN is detected by PGRP-SA (Michel et al., 2001). Thus, we moved to resolve whether the increase in Toll pathway activity observed in Klf15NN flies was due to sensing of PGN by PGRP-SA or β-(1,3)-glucan by GNBP3. qRT-PCR of the Toll target genes IM2, CG15067, and Drs demonstrated that the increase in basal Toll pathway signaling present in Klf15NN flies was downstream of PGRP-SA (Klf15NN; PGRP-SAsem) (Figures 3J, 3L, and 3M) but not GNBP3 (Klf15NN; GNBP3sem) (Figures S3N–S3P). The enhanced survival phenotype of Klf15NN flies was also lost in the double mutant Klf15NN , PGRP-SAsem (Figures 3K and 3L), but not in the double mutant Klf15NN , GNBP3sem (Figures S3Q and S3R), indicating that the surge in Toll signaling observed in nephrocyte-deficient flies is likely downstream of PGN recognition. Finally, bacterial load data from the double mutants Klf15NN; modSP1 and Klf15NN , PGRP-SAsem infected with S. aureus or E. faecalis showed that in absence of a functional Toll pathway, Klf15NN flies no longer display reduced pathogen load compared to WT (Figures S3S and S3T), confirming that the increase in resistance observed in Klf15NN flies is dependent on the Toll pathway.

Microbiota-Derived PAMPs Trigger Aberrant Toll Pathway Activation in Klf15NN Flies

Gut microbes are a source of PAMPs, such as PGN, and therefore can act as elicitors of the immune system (Clarke et al.,
Figure 3. Increased Resistance to Infection in Klf15NN Flies Is PGRP-SA-Dependent

(A–C) Comparison of Klf15NN; spzm7 double mutants to WT, Klf15NN, and spzm7 single mutants in experiments measuring IM2 (Toll target) gene expression via qRT-PCR (A), survival against S. aureus (B), and survival against E. faecalis (C).

(legend continued on next page)
In Drosophila, multiple mechanisms are in place to prevent undue activation of the Imd pathway in response to microbiota. These include the expression of a plethora of negative regulators (e.g., Caudal and Pirk) and enzymes that degrade DAP-type PGN (e.g., PGRP-LB and PGRP-SC). However, no similar mechanism has been described for the Toll pathway despite the fact that it also senses PGN (Lys-type) (Bischoff et al., 2004; Park et al., 2007). Because the increase in Toll pathway activity in Klf15NN flies depends on PGRP-SA, we hypothesized that the phenotype could stem from an errant immune response against the microbiota. To test this idea, we used qRT-PCR to measure the mRNA expression of the Toll target genes IM2, CG15067, and Drs (Figure S4A) in WT and Klf15NN flies raised in both conventionally reared (CR) and germ-free (GF) conditions. We found that the increase in Toll signaling in Klf15NN flies was fully dependent on the presence of microbiota, as GF WT and GF Klf15NN flies displayed similar mRNA expression for all measured Toll target genes.

Since Klf15NN flies did not have a higher microbiota load or show any alteration in gut-barrier integrity—as determined by the SMURF assay and measurements of both circulating bacteria in the hemolymph and whole fly microbiota (Figures S4B–S4D)—our results indicate that Klf15NN flies display aberrant Toll pathway activation in response to microbiota.

Because the increase in Toll pathway activity found in nephrocyte-deficient flies is both downstream of PGRP-SA and microbiota-dependent, we postulated that this phenotype could arise from an abnormal response to microbiota-derived PAMPs. In agreement with this hypothesis, mono-colonization of GF Klf15NN flies with the Gram-positive, Lys-type PGN-carrying microbes E. faecalis and Lactobacillus brevis (L. brevis, like many other Lactobacilli spp., carries Lys-Asp-type PGN as previously described [Salvetti et al., 2012; Schleifer and Kandler, 1972]) triggered aberrant Toll pathway activation, while recolonization with the Gram-negative, DAP-type PGN-containing Acetobacter pomorum did not (Figures 4A and S4A). Of note, E. faecalis,

For qRT-PCR experiments, mean values of three or more repeats are presented ± SE (*p < 0.05 **p < 0.01 ***p < 0.001 in a Student’s t test). Survival curves show the average percent survival ± SE of three biological replicates (**p < 0.01 ****p < 0.0001 in a Log-rank test).

Figure 4. Nephrocytes Prevent Excessive Immune Activation Against Gram-Positive Microbiota

(A) Quantification of mRNA transcripts in conventional (CR), germ-free (GF), and germ-free flies recolonized with either live A. pomorum (Gram-negative), live L. brevis (Gram-positive), or live E. faecalis (Gram-positive), qRT-PCR measurements of Toll target genes IM2 and CG15067 are shown. (B) Quantification of mRNA transcripts in CR, GF, and germ-free flies fed either heat-killed A. pomorum (Gram-negative), heat-killed L. brevis (Gram-positive), or heat-killed E. faecalis (Gram-positive), qRT-PCR measurements of Toll target genes IM2 and CG15067 are presented. For qRT-PCR experiments, mean values of three or more repeats are given ± SE. *p < 0.05 **p < 0.01 ***p < 0.001 in a Student’s t test. (C and D) Survival curve over 7 days following infection of WT and Klf15NN flies with S. aureus (C) and E. faecalis (D) in both CR and GF conditions. Survival curves give the average percent survival ± SE of three biological replicates (**p < 0.01 ****p < 0.0001 in a Log-rank test).

2010; Kaneko et al., 2004). In Drosophila, multiple mechanisms are in place to prevent undue activation of the Imd pathway in response to microbiota. These include the expression of a plethora of negative regulators (e.g., Caudal and Pirk) and enzymes that degrade DAP-type PGN (e.g., PGRP-LB and PGRP-SC). However, no similar mechanism has been described for the Toll pathway despite the fact that it also senses PGN (Lys-type) (Bischoff et al., 2004; Park et al., 2007). Because the increase in Toll pathway activity in Klf15NN flies depends on PGRP-SA, we hypothesized that the phenotype could stem from an errant immune response against the microbiota. To test this idea, we used qRT-PCR to measure the mRNA expression of the Toll target genes IM2, CG15067 (Figure 4A), and Drs (Figure S4A) in WT and Klf15NN flies raised in both conventionally reared (CR) and germ-free (GF) conditions. We found that the increase in Toll signaling in Klf15NN flies was fully dependent on the presence of microbiota, as GF WT and GF Klf15NN flies displayed similar mRNA expression for all measured Toll target genes.
A. pomorum, and L. brevis are normal constituents of the Drosophila gut microbiota (Broderick et al., 2014). This result suggested that the microbiota could act to elevate Toll pathway signaling in nephrocyte-deficient flies by providing a source of Lys-type PGN, thus stimulating PGRP-SA in the absence of infection. Additional experiments confirmed that gut microbiota-derived PAMPs were sufficient to trigger the Toll pathway in GF Klf15NN flies. Feeding GF Klf15NN hosts with heat-killed L. brevis or E. faecalis, but not A. pomorum, was enough to elicit abnormal Toll pathway activity as measured by IM2, CG15067, and Drs expression (Figures 4B and S4E). Altogether, these results established that in Klf15NN flies, gut-microbiota-derived Lys-type PGN induces an errant, Toll pathway-mediated immune response.

Next, we explored whether this abnormal response to the microbiota could be responsible for the increase in resistance to infection observed in Klf15NN flies. Unlike flies raised in CR conditions, GF Klf15NN flies infected with the bacterial pathogens S. aureus and E. faecalis did not exhibit increased survival to infection relative to GF WT controls (Figures 4C and 4D). These results suggest that microbiota-derived Lys-type PGN primes the Toll pathway in Klf15NN flies, leading to enhanced resistance. Chronic immune activation is costly and harmful to hosts (Charroux et al., 2018; Guo et al., 2014; Paredes et al., 2011). As we noted that flies devoid of nephrocytes had a shorter lifespan (Figure 1B), we asked whether this could also be due to chronic immune activation in response to the microbiota. Klf15NN flies reared in GF conditions significantly outlived their CR counterparts, and no difference in lifespan was found between Klf15NN and WT flies reared in GF conditions (Figure 4E). Our results demonstrate that nephrocytes are part of a program that prevents microbiota-dependent Toll pathway activation, thus avoiding its deleterious effect on lifespan.

Nephrocytes Endocytose PGN from the Hemolymph to Avert Excessive Immune Activation in Response to Microbiota

We find that the Toll pathway is significantly activated in response to microbiota-derived PAMPs in the absence of nephrocytes. This could be the result of either the presence of elevated amounts of microbiota-shed PGN in the hemolymph (as microbiota-shed PAMPs are commonly translocated from the gut lumen to systemic circulation) or the hyper-reactivity of these flies to PAMPs. SPE, a signaling component of the Toll pathway, accumulates in the hemolymph of WT and Klf15NN flies (Figure 2C) despite its mRNA not being transcriptionally increased (Figure S4I). Overexpression of SPE is also sufficient to trigger Toll pathway activation (Jang et al., 2006). Consequently, we hypothesized that accumulation of SPE could result in an aberrant response to the microbiota in Klf15NN flies. While overexpression of SPE alone resulted in increased expression of three target genes of the Toll pathway, IM2, CG15067, and Drs (Figures S4F–S4H), the mRNA expression of these genes was identical between CR and GF conditions, suggesting that this effect was not dependent on the presence of microbiota. It is therefore unlikely that the microbiota-dependent induction of Toll in Klf15NN flies is due to SPE accumulation. In light of this result, we moved on to the next hypothesis. As nephrocytes regulate hemolymph composition by filtration followed by filtrate endocytosis, we reasoned that in the absence of nephrocytes, microbiota-derived Lys-type PGN could accumulate in the hemolymph. We therefore measured the amount of PGN circulating in the hemolymph of WT and Klf15NN flies in both CR and GF conditions. Using a colorimetric assay (SLP), we detected three times more circulating PGN in Klf15NN flies than in WT controls under CR conditions, with no difference found between the two genotypes under GF conditions (Figure 5A). These data establish that nephrocytes participate in the removal of microbiota-shed PGN from systemic circulation.

Subsequently, we focused on determining which mechanisms underlie nephrocyte-mediated PGN removal from hemolymph. Nephrocytes are filtration devices. Their surface is covered by extensive membrane invaginations, which are sealed at the top by slit diaphragms. These chambers, known as lacunae or labyrinthine channels, are where most of their endocytic activity takes place. Once endocytosed, internalized cargo is either trafficked to lysosomes for degradation, metabolized and released back into circulation, or stored in vacuoles for the lifespan of the fly (Denholm and Skaer, 2009; Psathaki et al., 2018). To assess whether nephrocytes internalize circulating PGN, we immunostained nephrocytes with an anti-PGN antibody (raised against PGN from a Gram-positive Streptococcus sp.). Confocal sectioning of nephrocytes revealed a strong punctate signal pattern, indicating that nephrocytes do indeed internalize PGN (Figure 5B, specifically SB1). PGN staining disappeared in flies reared in GF conditions, suggesting that nephrocytes take up microbiota-derived PGN in order to remove it from the hemolymph (Figure 5B2, see Figure S5A for quantification). Immunostaining against PGN in nephrocytes expressing either a reporter for the early endosomal marker Rab5 (Hand-Gal4>UAS-Rab5-YFP) or a reporter for the late endosomal marker Rab7 (Hand-Gal4>UAS-Rab7-YFP) showed co-localization of PGN with both markers (Figure 5C, see Figure S5B for Pearson correlation coefficients). We also detected co-localization of PGN with the lysosomal markers cathepsin L and Lamp1 (Lysosomal associated membrane protein 1, Figures 5C, S5B, and S5C), implying that PGN is internalized by endocytosis and routed to the lysosomal compartment.

We then moved to evaluate the role of endocytosis in the uptake of PGN by nephrocytes. The dynamin Shibire is involved in the early steps of endocytosis. Blocking Shibire triggers elongation of the lacunae (also referred to as labyrinthine channels) within nephrocytes (Figure 5D, Duf labels the lacunae) (Kosaka and Ikeda, 1983; Psathaki et al., 2018). Because the filtration and endocytic functions of nephrocytes are separate, blocking endocytosis—but not filtration via expression of the thermosensitive shibireT allele results in the accumulation of filtrate in the lacunae, a phenomenon previously observed with the circulating serpin Necrotic (Soukop et al., 2009). When we blocked the endocytic pathway using this same allele (Hand-Gal4>UAS-shiT), we observed the accumulation of PGN in the lacunae of nephrocytes (Figures 5B3, 5D, S5D1, and S5E), signifying that nephrocytes endocytose PGN by a Shibire-dependent mechanism. In addition, nephrocyte-specific expression of a dominant negative form of Rab5 (Hand-Gal4>UAS-Rab5DN, Rab5 is a key regulator of early endosomal trafficking) and Rab7 (Hand-Gal4>UAS-Rab77DN, Rab77DN reroutes all traffic to clear vacuoles, thereby blocking access to the lysosome [Fu et al., 2017]) led to cyttoplasmic accumulation of endocytosed PGN relative
Since our data indicated that PGN is routed to the lysosomal compartment inside nephrocytes, we next assessed whether acidification of the lysosome is important for PGN degradation. Using in vivo RNAi, we decreased nephrocyte-specific expression of two key components of the vacuolar proton pump V-ATPase (Dot-Gal4 > UAS-Vha16-1-IR and Dot-Gal4 > UAS-Vha44-IR), which functions to acidify the endolysosomal compartment (Mauvezin et al., 2015). Blocking acidification of the endolysosomal compartment led to substantial PGN accumulation (Figures 5B6 and S5D3, quantification in S5A). Taken together, these results demonstrate that PGN is endocytosed and degraded by nephrocytes in a Shibire-, Rab5-, Rab7-, and V-ATPase-dependent manner.

Finally, we examined the consequences of arresting the endocytic function of nephrocytes on Toll pathway activity. Blocking endocytosis in nephrocytes via expression of the same shibirets1 and Rab5DN alleles was sufficient to induce abnormally high transcription of Toll target genes (Figures 5E and 5F). These results are in agreement with the increase in Toll pathway signaling we detected in flies in which Klf15 expression was decreased by in vivo RNAi in adult nephrocytes (Figure 2F), a condition known to block their endocytic capability (Ivy et al., 2015). Altogether, our results establish that nephrocytes remove microbiota-derived PGN from systemic circulation, thus preventing deviant immune activation in response to gut microbes.

DISCUSSION

Toll and Imd, the two primary immune recognition pathways in the fly, detect the presence of invading bacteria through sensing of specific forms of PGN: the Toll pathway recognizes Lys-type PGN from Gram-positive bacteria, while the Imd pathway detects DAP-type PGN from Gram-negative bacteria and certain Gram-positive bacilli (Lemaitre and Hoffmann, 2007). Because the gut is constantly exposed to microbes and their PAMPs, it relies on specialized mechanisms to prevent local immune activation against the microbiota. Given
that microbiota-shed PGN translocates from the gut lumen into general circulation (Capo et al., 2017; Clarke et al., 2010; Corbitt et al., 2013; Gendrin et al., 2009; Paredes et al., 2011; Zaidman-Rémy et al., 2006), mechanisms are required to prevent systemic immune activation in response to these microbiota products. Without such processes, chronic immune induction can lead to abnormal development (Bischoff et al., 2006) and/or a shortened lifespan, indicating that uncontrolled immune activity can be costly to the host’s health (Charroux et al., 2018; Guillou et al., 2016; Paredes et al., 2011). Mechanisms that prevent the systemic activation of Imd in response to the microbiota include the secretion of amidase PGRPs into the hemolymph, which act to degrade DAP-type PGN (Bischoff et al., 2006; Charroux et al., 2018; Guo et al., 2014; Paredes et al., 2011; Zaidman-Rémy et al., 2006). Here, we found that filtration of hemolymph by nephrocytes prevents chronic activation of the Toll pathway. Mechanistically, nephrocytes endocytose Lys-type PGN from systemic circulation and route it to lysosomes for degradation, thus maintaining immune homeostasis.

Why would an organism evolve distinct mechanisms to eliminate 2 types of PGN? One possibility is that efficient degradation of Lys-type PGN requires specialized enzymes, such as lysozymes, that work best in the acidified environment of a mature lysosome than in circulation. The optimal pH for Drosophila lysozyme activity is ~5 (Regel et al., 1998). By contrast, hemolymph pH is considerably higher, with pH values ranging from 7.3 to 7.4 (Ghosh and O’Connor, 2014). As nephrocytes are professional endocytic cells, they are well suited to rapidly and proficiently uptake Lys-type PGN from the hemolymph and route it for degradation to lysosomes. In support of this idea, it is worth noting that nephrocytes express at least 6 lysozyme-like genes (Chintapalli et al., 2007). Due to redundancy and a lack of genetic tools for all 6 lysozyme genes, we were not able to functionally test the role of these lysozymes in the degradation of PGN. Therefore, a role for lysozyme remains speculative. Interestingly, our data also established that the Imd pathway is not activated in the absence of nephrocytes. This may be a result of the efficient degradation of DAP-type PGN by amidase PGRPs, such that there is no remaining intact PGN of this class that needs to be filtered and endocytosed. Alternatively, the intrinsic negative charge of the nephrocyte basement membrane, which is known to act as a charge-selective filter (Denholm and Sker, 2009), may act to exclude passage of DAP-type but not Lys-type PGN.

Nephrocytes uptake Nec, a secreted serpin and negative regulator of the Toll pathway, and target it for lysosomal degradation (Soukup et al., 2009). Our work not only confirmed this finding, as Nec protein concentration was higher in the hemolymph of Klf15NN flies compared to WT, but also found that other signaling components of the Toll pathway, such as SPE, accumulated in the hemolymph of Klf15NN flies in the absence of transcriptional changes. These results suggest that hemolymph filtration by nephrocytes may serve to regulate Toll pathway homeostasis on multiple levels—regulating the concentration of both Lys-type PGN and Toll pathway components in the hemolymph. We were unable to determine whether the accumulation of signaling components of the Toll pathway was also important for the loss of immune homeostasis in flies lacking nephrocytes. However, the fact that GF Klf15NN flies did not show an increase in Toll activity suggests that PGN filtration, rather than accumulation of Toll pathway signaling components, is the critical mechanism at work. In addition, we note that the phenotype associated with a lack of nephrocytes is not easily predicted. At first glance, accumulation of Nec in flies devoid of nephrocytes would suggest a possible decrease in immune reactivity. However, we found that the loss of PGN filtration primes the immune system and increases resistance to infection.

Chronic kidney disease (CKD), characterized by a gradual loss of glomerular filtration rate, leads to alterations in plasma protein content similar to those observed in Klf15NN flies. Specifically, proteomic analyses show that patients with CKD progressively accumulate in their plasma high quantities of 24 proteins involved in the complement system, as well as 62 proteins associated with the acute phase response (Glorieux et al., 2015). Given the remarkable functional, structural, and molecular similarities between nephrocytes and the glomerular podocytes of the mammalian kidney, we propose that renal filtration by the kidneys could also act to regulate the amount of microbiota-derived PAMPs, such as PGN, in the blood, thus maintaining immune homeostasis. In support of this idea, we highlight that the alternative complement pathway, several components of which are enriched in the plasma of CKD patients, is activated by PGN, including Lys-type PGN (Kawasaki et al., 1987). In both Klf15NN flies and CKD patients, proteomic studies also showed accumulation of lysozymes in circulation, with lysozyme C increasing in the plasma and the lysozyme encoded by CG6426 rising in hemolymph. It is possible that lysozyme accumulation may result, in both cases, from induction of the immune system in response to PGN, especially as CG6426 is a target of the Toll pathway (De Gregorio et al., 2001; Troha et al., 2018). Finally, it has been proposed that nephrocytes are functionally analogous to endocytic scavenger cells of the mammalian reticuloendothelial system (Serensen et al., 2012; Wigglesworth, 1970). Therefore, it is possible that additional cells with scavenging function, such as hepatocytes, may also be involved in the regulation of circulating microbiota-shed PAMPs.

Altogether, our results reveal a role for podocyte filtration in the maintenance of insect immune homeostasis. The results suggest that renal clearance may be a major and conserved mechanism to remove PGN from circulation, thus preventing aberrant immune activation in response to the gut microbiota. Because of the parallels between the filtration systems of flies and mammals, as well as the similar consequences of altering renal function in both species, we propose that at least part of the immune activation observed in patients suffering from glomerular diseases may stem from the accumulation of PGN in plasma.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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- **EXPERIMENTAL MODEL AND SUBJECT DETAILS**
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we thank Gábor Juhasz for providing the

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The authors declare no competing interests.

DECLARATION OF INTERESTS

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AUTHOR CONTRIBUTIONS


DATA AND CODE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.immuni.2019.08.020.

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REFERENCES


## STAR METHODS

### KEY RESOURCES TABLE

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### CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Nicolas Buchon (nicolas.buchon@cornell.edu).

This study did not generate new unique reagents.

### EXPERIMENTAL MODEL AND SUBJECT DETAILS

#### Rearing of Drosophila melanogaster

Flies were maintained on standard sucrose-cornmeal-yeast medium: 50 g baker’s yeast, 60 g commeal, 40 g sucrose, 7 g agar, 26.5 mL Moldex (10%), and 12 mL Acid Mix solution (4.2% phosphoric acid, 41.8% propionic acid) per 1L of deionized H2O. Wild-type and mutant flies were raised at 24°C. Flies originating from crosses that employ the UAS-Gal4-Gal80\[^{\text{ts}}\] gene expression system were raised at 18°C (Gal80\[^{\text{ts}}\] On, Gal4 Off) and transferred to 29°C (Gal80\[^{\text{ts}}\] Off, Gal4 On) 5 days after eclosion. Males were used for all experiments, with the exception of immunostaining (larger female size is preferred for dissection and visualization of cells). For experiments with mutants, 5- to 8-day-old adult flies were used. For experiments with UAS transgenes, 10- to 14-day-old flies were used (to allow for the expression of the pertinent construct).
**Drosophila melanogaster strains**

Kif15\textsuperscript{SR}, spz\textsuperscript{rm7}, SPE\textsuperscript{Skr6}, psh\textsuperscript{1}, modSp\textsuperscript{1}, GNPB2\textsuperscript{Hades}, PGRP-SA\textsuperscript{sent}, and PPO1\textsuperscript{d}, 2\textsuperscript{d}, 3\textsuperscript{d} mutants and the dLamp-3xmCherry lysosomal marker have been previously described (Buchon et al., 2009; Dudzic et al., 2015; Gobert, 2003; Gottar et al., 2006; Hegedüs et al., 2016; Ivy et al., 2015; Jang et al., 2006; Michel et al., 2001; Ming et al., 2014). The nephrocyte-specific drivers, Dot-Gal4 and Hand-Gal4\textsuperscript{HS}, are detailed in (Ivy et al., 2015). The following lines were purchased from the Bloomington Drosophila Stock Center: UAS-shits1\textsuperscript{HS} (44222), UAS-Rab5\textsuperscript{HS} (9771), UAS-Rab7\textsuperscript{HS} (9778), UAS-Rab5-YFP (24616), UAS-Rab7-YFP (23270). The following lines were purchased from the Vienna Drosophila Resource Center: UAS-Vha16-IR (49290) and UAS-Vha44-IR (46563).

**Culturing of microbes**

The following bacteria were cultured overnight in LB broth and adjusted to the specified density: 

- *Salmonella typhimurium* (OD\textsubscript{600} = 1),
- *Listeria innocua* (OD\textsubscript{600} = 1),
- *Enterococcus faecalis* (OD\textsubscript{600} = 1),
- *Staphylococcus aureus* (OD\textsubscript{600} = 1),
- *Providencia rettgeri* (OD\textsubscript{600} = 1),
- *S. typhimurium* and *L. innocua* were grown at 37°C. All other bacteria were grown at 29°C. The fungi *Beauveria bassiana* and *Metarhizium anisopliae* were grown at 29°C on YPG-agar plates.

**METHOD DETAILS**

**Infection, survival, and lifespan experiments**

Flies were systemically infected with bacteria via septic pinprick to the thorax. Pinprick infection with an OD\textsubscript{600} = 1 for the bacteria aforementioned results in inoculation with ~3,000 CFU/fly. For natural infections with fungi, CO\textsubscript{2}-anaesthetized flies were placed directly on the sporulating lawn of a fungal culture plate and the plate was shaken for ~15 s to coat the flies in spores. Flies were then transferred to a new, clean food vial to recover. All flies, regardless of infection method, were maintained at 29°C for 24 hours. For survival experiments, death was recorded daily following inoculation, with flies transferred to fresh vials every 2 to 3 days. For lifespan measurements, adults were transferred to 29°C 5 days post-eclosion and remained at that temperature for the duration of the experiment. All experiments were performed at least 3 times.

**Quantification of bacterial CFUs**

At specified time points following inoculation, flies were individually homogenized by bead beating in 500 μL of sterile PBS using a tissue homogenizer (OPS Diagnostics). Dilutions of the homogenate were plated onto LB agar plates using a WASP II autoplate spiral plater (Microbiology International), incubated overnight at 29°C, and CFUs were counted. All experiments were performed at least 3 times.

**RT-qPCR**

For all experiments involving RT-qPCR, total RNA was extracted from pools of 20 flies using the standard TRIzol (Invitrogen) extraction. RNA samples were treated with DNase (Promega), and cDNA was generated using murine leukemia virus reverse transcriptase (MLV-RT-Promega). qPCR was performed using the SSO Advanced SYBR green kit (Bio-Rad) in a Bio-Rad CFX-Connect instrument. Data represent the relative ratio of the target gene and that of the reference gene Rpl32. Mean values of at least three biological replicates are represented ± SE. The oligonucleotide sequences used can be found in Table S1.

**Phagocytosis assays**

To assay phagocytosis, flies were injected in the thorax with 69 nL of pHrodo Red Bioparticles (Invitrogen) using a Nanoject (Drummond Scientific). The fluorescence within the abdomen of the flies was then imaged at 3 h post-injection with a Leica MZFLIII fluorescence microscope and quantified using ImageJ (NIH) as previously described (Guillou et al., 2016). To block phagocytosis, adult flies were pre-injected with a solution containing latex beads as previously described in (Elrod-Erickson et al., 2000). Twenty-four h post-injection, the flies were subjected to systemic infection as described above.

**Hemolymph extraction**

Hemolymph was collected using a centrifugation or capillary method. In the first method, ~100 anesthetized flies are loaded into a modified spin column (QIAGEN), in which the filter was removed and thoroughly washed with water before use, and 2 metal beads are placed on top of the flies. Flies are then centrifuged twice at 5,000 g for 5 min at 4°C. For more details, see (Troha and Buchon, 2019). For the capillary method, a pulled glass needle is used to prick flies in the thorax. Hemolymph is extracted into the needle by capillary action.

**DOPA assay**

Extracted hemolymph was immediately diluted in a 1:10 ratio using a protease inhibitor cocktail (Sigma: 11697498001) and kept on ice. Briefly, 50 μL of diluted hemolymph was combined with 150 μL of a 5 mM CaCl\textsubscript{2} solution and 800 μL of L-DOPA (Sigma: D9628) reagent. Following thorough mixing, 200 μL of sample/well was loaded into a 96-well plate. Using a spectrophotometer set to 29°C, a kinetic assay was performed at OD\textsubscript{492}.
**Generation of GF and mono-colonized flies**

Collected eggs were surface sterilized by immersion in 70% ethanol for 2 min. Eggs were then dechorionated via treatment with a 10% bleach solution for 10 min. This was followed by rinsing the eggs in sterile water 3 times to remove any leftover bleach. The eggs were then transferred to pre-autoclaved media vials, where they were allowed to develop. The entire procedure was performed using sterile technique in a laminar flow hood. For mono-colonized flies, pre-autoclaved media vials were seeded with 200 µL of the desired individual bacterial culture (OD$_{600}$ = 200). After the bacterial solution was absorbed into the media, adult germ-free flies were flipped into the mono-colonized media vial. Experiments with mono-colonized flies were carried out 5 days after the flies were first exposed to the bacteria.

**PGN detection by SLP assay**

The Silkworm Larvae Plasma (SLP) assay was used. After diluting extracted hemolymph (1:10 ratio), 50 µL of hemolymph sample/condition was used for the SLP assay (Fujifilm Wako Pure Chemical Corporation: 297-51501) following the manufacturer’s instructions.

**Gut barrier integrity (Smurf) assay**

Adult flies were fed standard medium supplemented with Blue Dye No. 1 (2.5%). A fly was counted as a Smurf when the blue dye could be observed outside of the digestive tract.

**UAS/GAL4/GAL80ts gene expression system**

For RNAi and overexpression experiments, we used the UAS/GAL4 gene expression system in combination with GAL80ts to restrict the expression of the constructs specifically to the adult stage. Flies were collected 5 to 8 days after eclosion from the pupal case and shifted to 29°C for an additional 8 days prior to any experiments. See our Rearing of Drosophila melanogaster section for additional details.

**Immunohistochemistry and fluorescence imaging**

Dissected nephrocytes were fixed in a 4% paraformaldehyde solution in PBST (PBS with 0.5% Triton X-100) for 1 h. After repeated washes in PBST, samples were blocked in 3% BSA PBST for 3 h and incubated overnight with primary antibodies in 1% BSA PBST at 4°C. Samples were labeled with secondary antibodies in 1% BSA PBST for 2 h. Samples were washed after each antibody labeling step with PBST containing 4% NaCl to reduce non-specific background labeling. The primary antibodies used in this study were: mouse anti-peptidoglycan (GeneTex: GTX39437) diluted 1:200, chicken anti-GFP (Invitrogen: A10262) diluted 1:1500, rabbit anti-Cathepsin L (Abcam: ab58991) diluted 1:1000, rabbit anti-Dumbfounded diluted 1:100 (Psathaki et al., 2018), and rabbit anti-RFP (Invitrogen: R10367) diluted 1:2000. The secondary antibodies were: Alexa Fluor 488 anti-chicken (A11039), Alexa Fluor 488 anti-rabbit (A21206), Alexa Fluor 488 anti-mouse (A21202), Alexa Fluor 555 anti-rabbit (A31572), and Alexa Fluor 555 anti-mouse (A31570), all diluted 1:1500 and from Invitrogen. Imaging was performed on a Zeiss LSM 700 confocal inverted microscope. Pearson correlation coefficients were calculated using ImageJ (FIJI, version: 2.0.0-rc-69/1.52n).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Aside from one exception, all analyses were performed in Prism (GraphPad Prism V7.0a, GraphPad Software, La Jolla, CA, USA). For survival assays, the curves represent the average percent survival ± SE of three or more biological replicates (n = 20 flies for each biological replicate). A Log-rank test was used to determine significance (*p < 0.05 **p < 0.01 ***p < 0.001 ****p < 0.0001). In bacterial load quantification assays, the horizontal lines represent median values for each time point. Three biological replicates were included. Following normalization, results were analyzed using a two-way ANOVA followed by Sidak’s post-test for specific comparisons (*p < 0.05 **p < 0.01 ***p < 0.001 ****p < 0.0001). For all other experiments, mean values of three or more biological repeats are presented ± SE. Significance was calculated by a Student’s t test following normalization (*p < 0.05 **p < 0.01 ***p < 0.001 ****p < 0.0001). Whenever survival curves crossed, a Cox’s proportional-hazards model was used instead of a Log-rank test to assay significance. In this case, SPSS (IBM Corp. Released 2017. IBM SPSS Statistics for Mac OS X, Armonk, NY: IBM Corp.) was used for the analysis.

**DATA AND CODE AVAILABILITY**

This study did not generate datasets/code.