Reactive Oxygen Species-Dependent Innate Immune Mechanisms Control Methicillin-Resistant *Staphylococcus aureus* Virulence in the *Drosophila* Larval Model

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**ABSTRACT** Antibiotic-resistant *Staphylococcus aureus* strains constitute a major public health concern worldwide and are responsible for both health care- and community-associated infections. Here, we establish a robust and easy-to-implement model of oral *S. aureus* infection using *Drosophila melanogaster* larvae that allowed us to follow the fate of *S. aureus* at the whole-organism level as well as the host immune responses. Our study demonstrates that *S. aureus* infection triggers H$_2$O$_2$ production by the host via the Duox enzyme, thereby promoting antimicrobial peptide production through activation of the Toll pathway. Staphylococcal catalase mediates H$_2$O$_2$ neutralization, which not only promotes *S. aureus* survival but also minimizes the host antimicrobial response, hence reducing bacterial clearance in vivo. We show that while catalase expression is regulated *in vitro* by the accessory gene regulatory system (Agr) and the general stress response regulator sigma B (SigB), it no longer depends on these two master regulators *in vivo*. Finally, we confirm the versatility of this model by demonstrating the colonization and host stimulation capabilities of *S. aureus* strains belonging to different sequence types (CC8 and CC5) as well as of two other bacterial pathogens, *Salmonella enterica* serovar Typhimurium and *Shigella flexneri*. Thus, the *Drosophila* larva can be a general model to follow *in vivo* the innate host immune responses triggered during infection by human pathogens.

**IMPORTANCE** The pathogenicity of methicillin-resistant *S. aureus* (MRSA) strains relies on their ability to produce a wide variety of tightly regulated virulence factors. Current *in vivo* models to analyze host-pathogen interactions are limited and difficult to manipulate. Here, we have established a robust and reliable model of oral *S. aureus* infection using *Drosophila melanogaster* larvae. We show that *S. aureus* stimulates host immunity through the production of reactive oxygen species (ROS) and antimicrobial peptide (AMP) and that ROS potentialize AMP gene expression. *S. aureus* catalase plays a key role in this complex environment and acts *in vivo* independently from SigB and Agr control. We propose that fly larvae can provide a general model for studying the colonization capabilities of human pathogens.

**KEYWORDS** *Staphylococcus aureus*, *Drosophila melanogaster*, intestinal infection, virulence, catalase, Duox, gastrointestinal infection

*S. aureus* is a facultative aerobic Gram-positive bacterium that behaves as a commensal microorganism (up to 30% of the healthy human population carries *S. aureus* through nasal, skin, and intestinal colonization) or as a pathogen causing...
a wide range of infections in humans and in wild and companion animals (1–3). The emergence and diffusion of methicillin-resistant *S. aureus* (MRSA) clones that express numerous virulence factors, including toxins and adhesins increasing their toxicity and colonization capacities, are a major public health issue. Expression of these numerous virulence factors are correlated with severe symptoms among previously healthy colonized individuals (4–6). During infection, *S. aureus* must face host innate immunity, i.e., phagocyte-mediated elimination via oxidative stress (by macrophages and neutrophils) and antimicrobial peptides secretion (7). *S. aureus* undergoes both endogenous oxidative stress (notably caused by incomplete aerobic respiration) and exogenous host-induced oxidative stress aimed at killing the bacteria (8, 9). Host reactive oxygen species (ROS) are secreted by the Nox/Duox NADPH (NAD phosphate) oxidases. In mammals, Nox and Duox families are composed, respectively, of five Noxs (Nox1 to Nox5) and two Duoxs (Duox1 and Duox2) members. Nox1 and Duox2 are found in the gastrointestinal tract, and Nox2 was identified in phagocytic cells (10–12), while other enzymes are expressed in other tissues, such as airway epithelium, kidneys, endothelial cells, etc. (13, 14). Duox enzymes generate hydrogen peroxide (H₂O₂), whereas Nox enzymes catalyze superoxide production (O₂⁻). Of note, mitochondria from phagocytic cells also contribute in ROS production to counteract infection (15). To neutralize the deleterious effects of ROS, *S. aureus* USA300 expresses multiple direct detoxifying enzymes, including (i) the two superoxide dismutases SodA and SodM, which convert the superoxide anion O₂⁻ to H₂O₂ and O₂; (ii) the H₂O₂-detoxifying catalase KatA and the alkyl hydroperoxide reductase AhpC, which, respectively, quench high and low H₂O₂ concentrations; and (iii) the two glutathione peroxidases GpxA1 and GpxA2, which also catalyze H₂O₂ reduction into water through glutathione (GSH) oxidation (16). It is known that *sodA* transcription depends on two sigma factors, the first being sigma A (σA)-type promoters and the second being the alternative stress-activated sigma B factor (σB), which represses *sodA; sodM* is repressed solely by σB (17). More recently, it was shown that *sodA* and *sodM* expression is repressed by the *msaABCR* operon (18). In contrast, the *katA* gene is upregulated through the ferric uptake regulator Fur under iron-rich conditions (19). Under low-iron or manganese-rich conditions, the metal-dependent transcription factor PerR inhibits Fur, resulting in the downregulation of *katA* (20).

Although most in vivo studies rely on the mouse model, mechanistic and genetic analyses can be performed with powerful alternative animal models, such as *Drosophila melanogaster* (fly) or *Danio rerio* (zebrafish). Of note, flies share many similarities with humans with respect to gastrointestinal anatomy and physiology (21), while the zebrafish model displays several disadvantages for intestinal infection research, mainly due to limited pH variations (the pH remains around 7.5) (22, 23). Furthermore, the *Drosophila gut microbiota**, which includes only a few bacterial species (mainly from the two bacterial families *Lactobacillaceae* and *Acetobacteraceae*) (24, 25), is closer to the human microbiota than it is to that of zebrafish, which is colonized mainly by the class gammaproteobacteria and more specifically by the *Aeromonas* genus (26). The *D. melanogaster* intestine consists of a simple ciliated epithelium layer surrounded by a muscle layer (27) that has the ability to develop a proper innate immune response to intestinal bacteria, including tolerating mechanisms for beneficial microbiota (28), similarly to mammals (29). The peritrophic matrix establishes a physical barrier that isolates pathogenic bacteria and their toxins from the epithelium layer (30).

When infected, the *Drosophila* intestinal epithelium, at all stages, can generate a robust antimicrobial response. On one hand, it involves the secretion of antimicrobial peptides (AMPs). They are produced (i) either upon Toll pathway activation, similarly to the MyD88 Toll-like receptor pathway in mammals, reacting to Gram-positive bacteria and fungi, (ii) or upon immune deficiency (IMD) pathway activation, which shares many similarities with the tumor necrosis factor (TNF) cascade, reacting to Gram-negative bacteria (31, 32). In addition to activating these two pathways, the fly can clear pathogenic bacteria by activating the production of microbicidal reactive oxygen
species (ROS) via the Duox pathway (33). Several studies showed that the *Drosophila* model recapitulates many aspects of the human intestinal pathologies (34) and has already allowed the successful evaluation of the harmfulness of human pathogens such as *Mycobacterium tuberculosis* (35), *Listeria monocytogenes* (36), *Vibrio cholerae* (37), *Francisella tularensis* (38), *Pseudomonas aeruginosa* (39), and *Yersinia pestis* (40).

The lack of a satisfactory *in vivo* model to study *S. aureus* virulence prompted us to develop an alternative *D. melanogaster* model that mimics mammalian immune responses to bacterial infections. To date, several *S. aureus* infection models have been assessed with adult flies. Systemic infections (via pricking in the thorax) result in different outcomes that depend on the dose used and the strain tested, while oral infections showed a limited, or no, infection cost for the host (41–46). Specifically, of the previously published research papers presenting models of intestinal infection in *Drosophila*, none used the epidemic methicillin-resistant *S. aureus* USA300 strain. Thus far, *S. aureus* USA300 virulence has been assessed only by septic injury in flies, leading to animal death in a more severe way than with poorly virulent strains, i.e., *S. aureus* NCTC8325 RN1 and CMRSA6 or the colonization strain M92 (43–46).

In this work, we took advantage of the *Drosophila* larval stage, where animals feed continuously and massively, to establish a new infection model based on the virulence of *S. aureus* USA300. We also demonstrate the colonization capabilities of *Salmonella enterica* serovar Typhimurium and *Shigella flexneri*, suggesting that *Drosophila* larvae can serve as a general model for studying multiple human pathogens.

**RESULTS**

*D. melanogaster* larvae as a new model to study *S. aureus* USA300 virulence.

We established a 24-h infection course (Fig. 1A), after a 30-min period in which mid-L3 larvae were fed a mixture of mashed banana and bacteria (see Materials and Methods). We observed that after 24 h of infection, 93% of the larvae were killed by a bacterium-enriched medium containing $10^8$ bacteria, while lower bacterial doses ($5 \times 10^7$, $2.5 \times 10^7$, or $1 \times 10^7$ bacteria) killed only 62, 51, and 20% of the larvae, respectively (Fig. 1B). We then monitored the kinetics of larval killing using wild-type *S. aureus* USA300 (USA300 WT), compared with the Gram-positive opportunistic entomopathogen *Micrococcus luteus*, which is known to be nonpathogenic to *D. melanogaster* (47).

Larvae were infected with medium enriched with $10^8$ bacteria for 30 min, and killing was followed over a 24-h period. Under these conditions, *S. aureus* USA300 WT was able to kill larvae, with a drop in animal survival occurring between 12 h and 18 h (Fig. 1C). In contrast, infection with *M. luteus* did not affect animal survival. We hypothesized that the death of the animals is related to the bacterial load in the gut. To avoid quantifying intestinal microbiota, we generated an *S. aureus* USA300 WT strain carrying the pRN11 plasmid, which expresses a chloramphenicol resistance (Cmr) gene (48). In support of the mortality data, the numbers of bacteria (CFU) in the larval gut were found to be 10-fold lower at 6 h and 20-fold lower at 24 h with an initial infective dose of $1 \times 10^7$ bacteria than with an initial infective dose of $10^8$ bacteria (Fig. 1D). These results suggest that larval death is related to the number of bacteria in the gut. We then confirmed the absence of effective tracheal colonization. As shown in Fig. S1A in the supplemental material, numbers of CFU remained low in the tracheal system throughout the experiment, reaching the highest count at 6 h, with an average of 447 CFU for an infecting dose of $10^8$ bacteria. In support of the mortality data, the numbers of bacteria (CFU) in the larval gut were found to be 10-fold lower at 6 h and 20-fold lower at 24 h with an initial infective dose of $1 \times 10^7$ bacteria than with an initial infective dose of $10^8$ bacteria (Fig. 1D). These results suggest that larval death is related to the number of bacteria in the gut. We then confirmed the absence of effective tracheal colonization. As shown in Fig. S1A in the supplemental material, numbers of CFU remained low in the tracheal system throughout the experiment, reaching the highest count at 6 h, with an average of 447 CFU for an infecting dose of $10^8$ bacteria. Furthermore, we showed that bacteria were not able to diffuse in the systemic compartment. As shown in Fig. S1B, *S. aureus* USA300 WT was almost undetectable in the hemolymph, as it reached only 8 and 6 CFU for 10 larvae, respectively, at 6 h and 18 h. Similar values were obtained with the nonpathogenic strain *M. luteus* (Fig. S1B). All together, these data indicate that *S. aureus* USA300 WT, in this model of oral infection, is pathogenic to *D. melanogaster* larva in a dose-dependent manner and that infection is constrained to the gut, where it persists for at least 24 h.

Since oral infection of adult flies with different *S. aureus* strains, including *S. aureus*...
**FIG 1** The *D. melanogaster* larva, a model to study *S. aureus* USA300 virulence. (A) Mid-L3 larvae were placed in a microcentrifuge tube with 100 μl of crushed banana and 100 μl of bacteria for 30 min. Then larvae were briefly washed with 30% ethanol and transferred to a petri dish with fresh fly medium until further processing. (B) Survival of *w1118 D. melanogaster* larvae following 30 min of oral infection with wild-type *S. aureus* USA300 at the indicated infectious doses. Animals were checked 24 h after infection. Data are means ± SEM (n = 3) with 20 animals/point. One-way ANOVA and multiple-comparison tests were performed between infected animals and BHI agar-treated (noninfected) animals (*, P < 0.05; **, P < 0.001). (C) Survival of *w1118 D. melanogaster* larvae upon 30 min of oral infection with 10^8 *S. aureus* USA300 WT bacteria and the nonpathogenic entomopathogen *Micrococcus luteus*. Animals were monitored at 0, 6, 12, 18, and 24 h after infection. Seventy animals from 3 independent experiments were used. The Kaplan-Meier test was applied to the whole group (***, P < 0.001). NI, noninfected larvae. (D) *w1118 D. melanogaster* larvae were orally infected for 30 min with 1 × 10^8 and 10 × 10^8 chloramphenicol-resistant *S. aureus* USA300 WT bacteria (carrying the pRN11 plasmid)/larva. Bacterial counts (CFU) in the gut were determined at 0.5, 6, 12, 18, and 24 h p.i. in live larvae. Tissues were homogenized in DPBS, serially diluted, and plated on BHI agar supplemented with chloramphenicol (10 μg/ml). Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests were performed between 1 × 10^8 and 10 × 10^8 bacteria-infected larva groups (**, P < 0.01). (E) Representative images of guts from noninfected larvae and larvae infected with mCherry-*S. aureus* USA300 WT (carrying pRN11) (Continued on next page)
USA300, does not interfere with animal survival (41, 42, 49), we assumed that larval death was related to a high number of ingested bacteria, due to their hyperphagic behavior. To confirm this, adult flies were first starved for 2 h and then placed on filters soaked with $10 \times 10^6$ bacteria for 1 h. Neither S. aureus USA300 WT nor M. luteus oral infection affected the survival of adult flies (Fig. S2A). Indeed, after 1 h of feeding, numbers of CFU recorded were only about 2% of those recorded with larvae (i.e., $6.9 \times 10^5$ bacteria/10 adult flies compared to $3.4 \times 10^7$ bacteria/10 larvae) (Fig. S2B), suggesting that animal killing, when flies are orally infected by S. aureus USA300 WT, may depend on the bacterial load ingested. The number of bacteria counted at day 1 in adult flies is consistent with the results of the study from Hori et al. (42), where the authors retrieved $8 \times 10^4$ bacteria per fly gut (compared to an average of $1.8 \times 10^5$ bacteria per fly in our study). Of note, we found that at day 1, adult flies had bacteria in the middle midgut (Fig. S2C).

We next analyzed S. aureus localization in the larval gut using fluorescence microscopy (Fig. 1E) and Lightsheet three-dimensional (3D) imaging (Fig. 1F and Movie S1). For this, we used the S. aureus USA300 WT strain carrying the pRN11 plasmid, which expresses the mCherry gene (48) (red fluorescence). Imaging from 6-h-infected larvae with mCherry-expressing S. aureus USA300 WT revealed that bacteria were clustered in the posterior midgut (Fig. 1E and F and Movie S1). This specific localization in larvae might be explained by the local pHs of the gut, as the first half of the hind midgut is at neutral to acidic pH, compared with the middle midgut, which corresponds to a very acidic region ($\text{pH} < 3$), and the second half of the posterior midgut, which corresponds to an alkaline region ($\text{pH} > 10$) (50). To test this hypothesis, we tested S. aureus USA300 susceptibility to a wide range of different pHs (from 3 to 11) (Fig. S3). We observed that S. aureus USA300 was highly susceptible to the highly acidic pH 3, as well as to basic pH 11. In contrast, bacteria were able to survive at pH 5 and 9 for at least 2 h and multiplied at pH 7. This sensitivity to environmental pH may explain the specific localization of S. aureus USA300 WT in the neutral region of the Drosophila larval gut. Interestingly, the site of infection was associated with apparent inflammation (a swelling of the gut in this area of about 1.3 times that of uninfected larvae) (Fig. S4). Overall, these data show that S. aureus USA300 successfully colonizes D. melanogaster larvae after a 24-h infection and preferentially localizes to the anterior half of the midgut. This prolonged infection results in tissue inflammation and correlates with animal death.

**Drosophila larvae are a suitable model for human pathogen studies.** To assess the versatility of the larval model and confirm its ease of implementation, we tested the infection of larvae with three other clinical S. aureus strains belonging to different sequence types (STs) that are predominant among clinical S. aureus isolates sampled in human infections. These strains have been designated as follows: S. aureus P1 (ST1), S. aureus P2 (ST5), and S. aureus P3 (ST30) (see Materials and Methods for details). As shown in Fig. S5A, we observed that animals infected with S. aureus P1 and P2 followed survival curves identical to those of larvae infected with S. aureus USA300 WT (with, respectively, 24, 23, and 31% of animals surviving at 24 h postinfection [p.i.]), while S. aureus P3 showed a slightly higher virulence (with 9% of larvae surviving at 24 h p.i.). No difference in bacterial survival was observed (the CFU counts recorded for the three isolates at 6 h and 24 h p.i. were identical to those of the WT strain) (Fig. S5B). Interestingly, the three isolates differentially activated the Drosophila Toll pathway. At 6 h p.i., both S. aureus P1 and P2 triggered a 44% decrease in the expres-
sion of the antimicrobial peptide drosomycin (Drs; one of the main readouts for the Toll pathway in *D. melanogaster*) compared with that of *S. aureus* USA300 WT, whereas *S. aureus* P3 induced a 66% increase in the expression of *Drs* (Fig. S5C). These data suggest that the *Drosophila* larvae can serve as a model to evaluate the virulence of *S. aureus* clinical isolates as well as their ability to activate the host immune response.

We have also tested the value of this model with two human enteric pathogens: *Salmonella enterica* serovar Typhimurium and *Shigella flexneri*. Notably we observed significant larval death when organisms were fed 10 × 10⁸ bacteria/larva, under conditions similar to those of *S. aureus* infection (see Materials and Methods). Indeed, after 24 h of infection, 53.8% and 46.6% of the larvae were killed, respectively, when they were fed *S. Typhimurium*- and *S. flexneri*-enriched medium (Fig. S6A). After 30 min of feeding, the larvae were infected with 8.2 × 10⁶ *S. Typhimurium* or 7.9 × 10⁶ *S. flexneri* bacteria per 10 larvae. At 6 h p.i., 5 × 10⁸ and 2 × 10⁹ bacteria per 10 larvae were recorded, respectively (Fig. S6B). We confirmed that these two pathogens also triggered immune responses in *Drosophila*, as we observed a significant production of intestinal H₂O₂ at 2 h when larvae were infected with *S. Typhimurium* (43% increase) and *S. flexneri* (57% increase) (Fig. S6C). This correlated with a significant increase in the expression of the gene for the antimicrobial peptide diptericin (*Dpt*), which is dependent on the Gram-negative, sensitive immune deficiency pathway (51). We observed 42.9-fold (*S. Typhimurium*) and 37.9-fold (*S. flexneri*) increases in *Dpt* expression at 6 h postinfection (Fig. S6D). Interestingly, by using DsRed-expressing strains, we observed that at 6 h postinfection, each strain preferentially localized to the foregut and midgut (Fig. S6E, white arrowheads), where pH values ranged from neutral to acidic (50). This trait may be explained by their ability to survive in an acidic environment (52). We confirmed that the sensitivity of *S. Typhimurium* and *S. flexneri* to pH 5 after 2 h of treatment (Fig. S6F) was lower than that of *S. aureus*. After 2 h in brain heart infusion (BHI) broth adjusted to pH 5, the number of CFU of *S. aureus* was 25 times lower than after treatment at pH 7. In contrast, for *S. Typhimurium* and *S. flexneri*, the values recorded at pH 5 were only 5- to 3-fold lower than those recorded at pH 7, respectively.

**S. aureus USA300 triggers host intestinal hydrogen peroxide production.** It has previously been shown that adult intestinal infection triggers the production of reactive oxygen species (ROS) by the Duox enzyme to eliminate invading pathogens, complementing AMP actions (53). Therefore, we monitored the transcripts level of the *Duox* gene in intestines from infected larvae (10 × 10⁸ bacteria). As shown in Fig. 2A, we observed an 86.8-fold increase in *Duox* transcription at 2 h p.i. over that of noninfected animals. Interestingly, the use of heat-killed (HK) bacteria also induced *Duox* transcription but to a lesser extent (22.2-fold increase), suggesting that the *S. aureus* cell wall, as well as secreted factors, modulates *Duox* transcription. To confirm this induction of intestinal ROS in this context, we used the H₂O₂-specific detector 2′,7′-dichlorodihydrofluorescein diacetate (H₂DCFDA). We observed a 20% increase in signal detection in infected intestines 2 h p.i. compared to that in noninfected intestines (Fig. 2B). This was confirmed by live imaging (Fig. 2C). Interestingly, H₂DCFDA fluorescence (green) often colocalized with bacteria (mCherry *S. aureus* USA300, red) in the intestine. We also noticed strong H₂DCFDA fluorescence in Malpighian tubules when animals were infected (Fig. 2C, white arrowheads). Of note, it was recently shown that Malpighian tubules also play an active role during oral infection by sequestering excessive ROS and oxidized lipids (54). All together, these results demonstrate that *S. aureus* USA300 oral infection rapidly triggers H₂O₂ production at the intestinal epithelium, through Duox activation.

**Catalase is a key enzyme in the *S. aureus* antioxidant defense.** Since H₂O₂ generation through Duox enzyme is a key mechanism for controlling pathogen load (55), we then tested which *S. aureus* antioxidant enzymes, catalase (*katA* encoded), superoxide dismutases (*sodA* and *sodM* encoded), or glutathione peroxidases (*gpxA1* and *gpxA2* encoded), might contribute to intestinal persistence during host infection. For this, larvae were orally infected with *katA*, *sodA*, *sodM*, *gpxA1*, or *gpxA2* transposon insertion mutants (kindly obtained from BEI Resources [see Materials and Methods]), and we
FIG 2  ROS quenching in vivo is a key mechanism for successful colonization. (A) w^{1118} mid-L3 larvae were fed for 30 min with $10^8$ live or heat-killed (HK) S. aureus USA300 WT bacteria/larva. A quantitative real-time PCR analysis of Duox transcripts was done with total RNA extracts from guts (15 animals, n = 3) recovered at 2 h p.i. Bar graph data are relative to RP49. Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests between infected groups and noninfected (NI) larvae were applied (*, P < 0.05; **, P < 0.01). (B) w^{1118} mid-L3 larvae were fed for 30 min with $10^8$ S. aureus USA300 WT bacteria/larva. Generation of intestinal H$_2$O$_2$ was measured with the H$_2$DCFDA dye (10 µM) on noninfected samples and at 2 h p.i. Data (Continued on next page)
evaluated bacterial persistence at 6 h p.i. We used pRN11-transformed bacteria to allow chloramphenicol resistance gene expression and specific clone selection on chloramphenicol-supplemented BHI agar.

The $\Delta sodA$, $\Delta gpxA1$, and $\Delta gpxA2$ mutants behaved like the wild-type strain. In contrast, the growth of the $\Delta sodM$ mutant was 60-fold less than that of the WT strain (Fig. 2D), supporting an earlier report of a mouse abscess model (17). The $\Delta katA$ mutant was the most attenuated strain compared to the WT strain (92-fold), prompting us to evaluate its role in $S. aureus$ virulence and persistence. Genome-wide sequencing and analysis of the $\Delta katA$ mutant confirmed the transposon insertion site in the $katA$ gene and the absence of unintended secondary mutations. We showed that the $\Delta katA$ mutant strain was more sensitive to H$_2$O$_2$ than the WT strain (15 mM H$_2$O$_2$ in Dulbecco’s phosphate-buffered saline (DPBS)) (Fig. S7A). As shown in Fig. 2E, we observed that $S. aureus$ USA300 $\Delta katA$ killed larvae to a much lesser extent than the WT strain. However, expression of the catalase gene in the mutant strain restored its virulence phenotype. This difference in larval survival was correlated with 17.5-, 190-, and 122-fold decreases in the number of intestinal $\Delta katA$ mutant CFU compared to the number of WT CFU, respectively, at 2, 4, and 6 h p.i. (Fig. 2F). Restoration of catalase expression in the $\Delta katA$ strain restored $S. aureus$ survival in the Drosophila gut to WT $S. aureus$ levels at 6 h p.i. (Fig. S7B). Supporting the idea that this greater bacterial clearance may be related to a defect in H$_2$O$_2$ quenching, we observed a significant increase in the amount of ROS, by H$_2$DCFDA measurement, in the intestines of $\Delta katA$ mutant-infected larvae compared to that in WT-infected larvae (with respective increases in fluorescence intensity of 43% and 34% at 2 and 6 h p.i.) (Fig. 2G). These results suggest that $S. aureus$ USA300 $\Delta katA$ is defective for H$_2$O$_2$ quenching. Second, we confirmed that bacterial persistence in the larval gut and the bacterium’s ability to kill larvae are closely related to ROS content. For this, we evaluated numbers of bacterial CFU of the WT and $\Delta katA$ strains in flies fed N-acetyl-L-cysteine (NAC; 1 mM), an antioxidant drug that was shown to quench H$_2$O$_2$ molecules (56). We observed that NAC counteracted the deleterious intestinal environment for the $\Delta katA$ strain, as NAC abolished the $\Delta katA$ mutant’s defect compared to the WT at 2 h p.i. and promoted a 5-fold increase in the CFU count of the $\Delta katA$ mutant at 6 h p.i. (Fig. 2H). In parallel, we tested the survival of the WT and $\Delta katA$ strain in NP3084-GAL4 $\rightarrow$ Duox-RNAi larvae that are defective for Duox expression specifically in the intestine. NP3084-GAL4 (or MyoD1-GAL4) primarily drives gene expression in the larval midgut in enterocytes (57). Notably, larvae with abolished Duox

FIG 2 Legend (Continued)

are means ± SEM (n = 4). The Mann-Whitney test was applied (*, P < 0.05). A.U., arbitrary units. (C) Representative live imaging of posterior midguts from noninfected larvae (NI) and orally infected larvae (2 h p.i.; mCherry-S aureus USA300 WT, red). Intestines were dissected, treated with H$_2$DCFDA (10$\mu$m green) for 15 min, and imaged with an epifluorescence microscope. TL, transmitted light. White arrowheads indicate Malpighian tubules. Scale bar, 10 $\mu$m. (n = 3, 10 guts/experiment, for each condition.) (D) $w^{1118}$ D. melanogaster larvae were orally infected for 30 min at 10$^6$ bacteria/larva with chloramphenicol-resistant S aureus USA300 WT or the $\Delta katA$, $\Delta sodA$, $\Delta sodM$, $\Delta gpxA1$, or $\Delta gpxA2$ (carrying the pRN11 plasmid) strain. Bacterial counts (CFU) were determined at 0 6 h p.i. After homogenization and serial dilution, samples were plated on BHI agar supplemented with chloramphenicol (10$\mu$g ml$^{-1}$). Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests between mutant-infected groups and the WT-infected group were applied (*, P < 0.05). (E) Survival of $w^{1118}$ D. melanogaster larvae following 30 min of oral infection with S aureus USA300 WT $\Delta katA$ $\Delta katA$-cp-katA with 10 $\times$ 10$^6$ bacteria/larva. The experiment was monitored until 24 h after infection. Sixty-two animals were pooled from 3 independent experiments. The Kaplan-Meier test was applied for the whole group (***, P < 0.001). (F) $w^{1118}$ D. melanogaster larvae were orally infected for 30 min at 10$^6$ bacteria/larva with chloramphenicol-resistant S aureus USA300 WT or the $\Delta katA$ mutant (carrying the pRN11 plasmid). Bacterial counts (CFU) were determined at 0.5, 2, 4, and 6 h p.i. After homogenization and serial dilution, samples were plated on BHI agar supplemented with chloramphenicol (10$\mu$g ml$^{-1}$). Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests were performed between WT and $\Delta katA$-infected larval groups (**, P < 0.01; ***, P < 0.001). (G) $w^{1118}$ mid-L3 larvae were fed for 30 min with chloramphenicol-resistant S aureus USA300 WT or the $\Delta katA$ mutant at the infectious dose of 10 $\times$ 10$^6$ bacteria/larva. The intestinal ROS titer was measured at 0.5, 2, and 6 h p.i. After dissection, intestines were homogenized in 400 $\mu$l DPBS and treated with H$_2$DCFDA (10$\mu$m) for 30 min. Fluorescence was measured at 490 nm. Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests were performed between WT and $\Delta katA$-infected larval groups (*, P < 0.05; **, P < 0.01). (H) $w^{1118}$ D. melanogaster mid-L3 larvae were orally infected for 30 min with chloramphenicol-resistant S aureus USA300 WT or the $\Delta katA$ mutant (carrying the pRN11 plasmid) at the infectious dose of 10 $\times$ 10$^6$ bacteria/larva. Then animals were transferred to fresh fly medium supplemented, or not, with NAC (1 mM). Bacterial counts (CFU) were determined at 0.5, 2, and 6 h p.i. After homogenization and serial dilution, samples were plated on BHI agar supplemented with chloramphenicol (10$\mu$g ml$^{-1}$). Data are means ± SEM (n = 4). One-way ANOVA and then multiple-comparison tests were performed (*, P < 0.05; **, P < 0.001). (i) NP3084-GAL4 $\rightarrow$ Duox-RNAi larvae were orally infected for 30 min with chloramphenicol-resistant S aureus USA300 WT or the $\Delta katA$ mutant (carrying the pRN11 plasmid) at the infectious dose of 10 $\times$ 10$^6$ bacteria/larva. Bacterial counts (CFU) were determined at 0.5, 2, and 6 h p.i. After homogenization and serial dilution, samples were plated on BHI agar supplemented with chloramphenicol (10$\mu$g ml$^{-1}$). Data are means ± SEM (n = 4). One-way ANOVA and then multiple-comparison tests were performed (*, P < 0.05; **, P < 0.01; ***, P < 0.001).
expression in the midgut (NP3084-GAL4 > Duox-RNAi) showed significant 9.8- and 11.7-fold increases in CFU counts for the ΔkatA strain, respectively, at 2 h and 6 h p.i. in NP3084-GAL4 > Duox-RNAi compared to counts in NP3084-GAL4 > w^{1118} larvae. In contrast, the S. aureus USA300 WT strain showed nonsignificant 0.8- and 1.7-fold changes in CFU counts in NP308-GAL4 > S. aureus USA300 ΔkatA strain at the infectious dose of 10 × 10^8 bacteria/larva. At 6 h p.i., guts were dissected for quantitative real-time PCR analysis of *Drosomycin* transcripts. Data were normalized to the corresponding RP49 levels. Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests were performed (*, P < 0.05). (C) w^{1118} mid-L3 larvae were fed for 2 h with fly medium supplemented with stabilized H_2O_2 (0.5%). Guts were dissected for quantitative real-time PCR analysis of *Drosomycin* transcripts. Transcripts levels were normalized to the corresponding RP49 levels. Data are means ± SEM (n = 3). The Mann-Whitney test was applied to compare the nontreated (NT) group and H_2O_2-treated group (*, P < 0.05). (D) w^{1118} mid-L3 larvae were orally infected for 30 min with S. aureus USA300 WT or S. aureus USA300 ΔkatA at the infectious dose of 10 × 10^8 bacteria/larva. Then animals were transferred to fresh fly medium supplemented, or not, with NAC (1 mM). At 6 h p.i., guts were dissected for quantitative real-time PCR analysis of *Drosomycin* transcripts. Data were normalized to the corresponding RP49 levels. Results were compared to those for noninfected larvae transferred to supplemented NAC medium (NI+NAC) or not (NI). Data are means ± SEM (n = 4). One-way ANOVA and then multiple-comparison tests were performed (*, P < 0.05; **, P < 0.01).

**Catalase-mediated ROS quenching limits Toll pathway activation in the host.**

Like other Gram-positive bacteria, *S. aureus* is known to induce the Toll pathway, a key innate immune signaling pathway in *D. melanogaster*, through its lysine-type peptidoglycan (58). This prompted us to test the expression of the Drs gene in wild-type *yw* larvae and in the derivative *spz*^{mut7} mutated line (larvae lacking the expression of the Toll ligand spätzle) when larvae were infected with the *S. aureus* USA300 WT strain. As shown in Fig. 3A, we observed a significant 126-fold increase in intestinal Drs expression in the midgut (NP3084-GAL4 > Duox-RNAi) showed significant 9.8- and 11.7-fold increases in CFU counts for the ΔkatA strain, respectively, at 2 h and 6 h p.i. in NP3084-GAL4 > Duox-RNAi compared to counts in NP3084-GAL4 > w^{1118} larvae. In contrast, the *S. aureus* USA300 WT strain showed nonsignificant 0.8- and 1.7-fold changes in CFU counts in NP308-GAL4 > S. aureus USA300 ΔkatA strain at the infectious dose of 10 × 10^8 bacteria/larva. At 6 h p.i., guts were dissected for quantitative real-time PCR analysis of *Drosomycin* transcripts. Data were normalized to the corresponding RP49 levels. Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests were performed (*, P < 0.05). (B) w^{1118} mid-L3 larvae were orally infected for 30 min with *S. aureus* USA300 WT or the ΔkatA or ΔkatA-cp-ΔkatA strain at the infectious dose of 10 × 10^8 bacteria/larva. At 6 h p.i., guts were dissected for quantitative real-time PCR analysis of *Drosomycin* transcripts. Data were normalized to the corresponding RP49 levels. Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests were performed (*, P < 0.05). (C) w^{1118} mid-L3 larvae were fed for 2 h with fly medium supplemented with stabilized H_2O_2 (0.5%). Guts were dissected for quantitative real-time PCR analysis of *Drosomycin* transcripts. Transcripts levels were normalized to the corresponding RP49 levels. Data are means ± SEM (n = 3). The Mann-Whitney test was applied to compare the nontreated (NT) group and H_2O_2-treated group (*, P < 0.05). (D) w^{1118} mid-L3 larvae were orally infected for 30 min with *S. aureus* USA300 WT or *S. aureus* USA300 ΔkatA at the infectious dose of 10 × 10^8 bacteria/larva. Then animals were transferred to fresh fly medium supplemented, or not, with NAC (1 mM). At 6 h p.i., guts were dissected for quantitative real-time PCR analysis of *Drosomycin* transcripts. Data were normalized to the corresponding RP49 levels. Results were compared to those for noninfected larvae transferred to supplemented NAC medium (NI+NAC) or not (NI). Data are means ± SEM (n = 4). One-way ANOVA and then multiple-comparison tests were performed (*, P < 0.05; **, P < 0.01).
expression, in comparison to levels under noninfected conditions (using a $10 \times 10^8$ bacteria/larva setup), in yw flies. This activation was proportional to the initial bacterial load, as a 10-fold-lower infectious dose ($1 \times 10^9$) induced only a 16-fold increase in Drs gene transcription. Notably, using $spz^{m7}$ larvae considerably reduced the Drs transcript amount, even using medium enriched with $10 \times 10^8$ bacteria, suggesting that Drs activation is almost exclusively controlled by the Toll pathway. In Drosophila, links between ROS and the Toll/NF-κB pathway have already been established. Under wasp infestation (at the larval stage), the lymph gland (the main hematopoietic organ) undergoes a burst of ROS in the posterior signaling center, resulting in Toll pathway activation, whose purpose is to redirect hemocyte progenitor differentiation into the lamellocyte subtype (59). This led us to wonder if H$_2$O$_2$ generated during the infection plays a role in Toll pathway activation in the intestine. We first evaluated Drs expression in animals infected with S. aureus USA300 WT or the ΔkatA or ΔkatA cp katA strain (S. aureus USA300 ΔkatA complemented with the pCN57-cp-katA plasmid [see Materials and Methods]). Interestingly, the ΔkatA strain induced a 65% increase in Drs transcription at 6 h p.i. in Drosophila intestine compared to that in WT-infected animals. Infection with the complemented ΔkatA strain allowed restoration of Drs expression to levels under WT-infected conditions (Fig. 3B). We then tested the direct effect of H$_2$O$_2$ on intestinal Drs expression. Animals treated with H$_2$O$_2$ (0.5% in fly medium) for 2 h showed a 15-fold induction of Drs expression in the gut (Fig. 3C). We then evaluated the expression of the Drs gene in flies infected with the WT and ΔkatA strains, fed with NAC or not. At 6 h p.i., the WT and ΔkatA strains induced 91- and 169-fold increases in Drs expression, respectively, relative to levels under noninfected condition. Of note, at this time point (Fig. 2F), the number of ΔkatA bacteria was 122-fold lower than that of the WT strain, suggesting that factors other than the bacteria themselves modulate Drs expression. Feeding animals with NAC induced relative 1.8- and 3.5-fold decreases in Drs expression. Animals treated with H$_2$O$_2$ underwent a higher transcription in the WT strain (16.5-fold increase)

**The katA gene is differentially regulated by SigB and Agr in vitro and in vivo.**

We then tested the contribution of the master regulators SigB and Agr in the virulence of S. aureus in fly larvae and more particularly their influence on the expression of catalase encoded by katA. The expression of the alternative factor sigma B is linked to environmental stress and plays a key role in resistance to oxidative stress, heat, and antibiotics, while the two-component quorum-sensing system encoded by the accessory gene regulator (the Agr loci, composed of the AgrA, AgrB, AgrC, and AgrD loci) locus regulates multiple virulence components (60). To address their role, we used sigB and agrC (agrC codes for the receptor histidine kinase AgrC) transposon insertion mutants. As for the katA transposon insertion mutant, the whole-genome sequence analysis of the agrC and sigB transposon insertion mutants confirmed the correct transposon insertion site and the absence of unintended mutations in these mutant strains.

We first tested the virulence of ΔsigB and ΔagrC mutant strains in our model. We observed that 72% and 69% of larvae survived upon infection with the sigB and agrC mutants, respectively, at 24 h (compared to 16% of larvae surviving with the WT strain). The virulence of the ΔsigB and ΔagrC mutants was partially or totally restored when the strains were complemented with pCN57-cp-sigB and pCN57-cp-agrC, respectively (see Materials and Methods) (Fig. 4A). The attenuated virulence of the two mutants was accompanied by a reduced number of bacteria in intestines, which was also restored partially or totally when the strain was complemented with the corresponding WT allele of sigB and agrC (Fig. 4B). Partial phenotype restoration of the ΔsigB strain might be explained by differences in genomic and plasmid-carried sigB expression.

We then first assessed the specific role of SigB and Agr in katA transcription upon oxidative stress. For this, exponentially growing bacteria were subjected to 15 mM H$_2$O$_2$ in DPBS for 30 min. We observed that under nontreated conditions, katA expression remained unchanged in the WT, ΔsigB, and ΔagrC strains. Under H$_2$O$_2$ treatment, the katA gene underwent a higher transcription in the WT strain (16.5-fold increase).
**FIG 4** *katA* expression is differently mediated *in vitro* and *in vivo*. (A) Survival of *w*1118 *D. melanogaster* larvae following 30 min of oral infection (10 × 10^8 bacteria/larva) with *S. aureus* USA300 WT or the ΔsigB, ΔsigB-cp-sigB, ΔagrC, or ΔagrC-cp-agrC strain against noninfected animals (NI). The experiment was monitored until 24 h after infection. At least 60 animals were pooled under each condition from 3 independent experiments. The Kaplan-Meier test was used to compare the whole group (***, P < 0.001). (B) *w*1118 *D. melanogaster* larvae were orally infected for 30 min at 10 × 10^8 bacteria/larva with chloramphenicol-resistant *S. aureus* USA300 WT (carrying the pRN11 plasmid, plated on BHI agar with chloramphenicol) or the ΔsigB, ΔsigB-cp-sigB, ΔagrC, or ΔagrC-cp-agrC strain (plated on BHI agar with erythromycin). Bacterial counts (CFU) were determined at 6 h p.i. Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests were performed (*, P < 0.05). (C, D) Exponentially grown *S. aureus* WT (C), the ΔsigB mutant (D, left panel), or the ΔagrC mutant (D, right panel) were incubated for 30 min in DPBS with or without (NT) H₂O₂ 15 mM. Quantitative real-time PCR analyses of *katA* (C), *sigB* (D, left panel), or *agrC* (D, right panel) transcripts were performed. Transcripts levels were normalized to the corresponding *gyrB* levels. Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests were performed (**, P < 0.01; ***, P < 0.001). (E) NP3084-GAL4 > *w*1118 and NP3084-GAL4 > Duox RNAi larvae were orally infected for 30 min with *S. aureus* USA300 WT (10 × 10^8 bacteria/larva). At 2 h p.i., quantitative real-time PCR analyses of the *katA* strain were performed. Transcripts levels were normalized to the corresponding *gyrB* levels and expressed relative to WT results for panel F. Data are means ± SEM (n = 3). One-way ANOVA and then multiple-comparison tests were performed, and results were compared to those for the WT-infected group (*, P < 0.05; **, P < 0.01).
than in the \(\Delta\text{sigB}\) and \(\Delta\text{agrC}\) strains (4.1- and 3-fold increases) (Fig. 4C). Unexpectedly, transcriptional analyses revealed that \(\text{H}_2\text{O}_2\) downregulated \(\text{sigB}\) expression independently from Agr (Fig. 4D, left), whereas repressive control from SigB on \(\text{agr}\) expression, under nontreated conditions, was lost upon \(\text{H}_2\text{O}_2\) treatment (Fig. 4D, right).

Finally, we monitored \(\text{katA}\) gene regulation \textit{in vivo} during larval intestinal infection. We found that \(\text{S. aureus}\) \(\text{katA}\) expression was dependent on intestinal \(\text{H}_2\text{O}_2\) generation, as we observed a 14.7-fold decrease in expression in \(\text{Duox RNAi}\)-expressing larvae (\(\text{NP3084-GAL4} > \text{Duox RNAi}\)) compared to that in WT larvae (\(\text{NP3084-GAL4} > \text{w}\)) at 2 h.p.i. (Fig. 4E). Unexpectedly, the \(\text{katA}\) gene was upregulated in \(\Delta\text{sigB}\) and \(\Delta\text{agrC}\) strains compared to the WT during intestinal infection (respectively, 25.5- and 4.7-fold increases) (Fig. 4F). Experiments performed in animals also showed that Agr repressed \(\text{sigB}\) expression at 6 h.p.i. (Fig. 4G, left), whereas \(\text{sigB}\) mutation still resulted in a higher expression of \(\text{agr}\) (Fig. 4G, right). All together, these results support a role for Agr as a new regulator of \(\text{katA}\) transcription upon \(\text{H}_2\text{O}_2\) challenge, in addition to SigB and other contributors of the \(\text{S. aureus}\) oxidative-stress response.

**DISCUSSION**

Here, we present an \textit{in vivo} \textit{Drosophila} larval model that allows for easy and rapid monitoring of both bacterial infection and innate host immune responses simultaneously. The use of this invertebrate model offers great potential to dissect complex host-pathogen interactions (61–63) because it has remarkable homology to mammals in innate immunity, in addition to available genetic tools and husbandry facilities.

\textbf{Intestinal infection with MRSA and MSSA clinical isolates induces larval death.}

During oral infection of \textit{Drosophila} larvae by \(\text{S. aureus}\), the bacteria reach and establish themselves in the first half of the posterior part of the larval gut, probably due to the neutral pH specifically encountered there. This localized colonization is associated with intestinal enlargement, as previously observed with the invertebrate model \textit{Caenorhabditis elegans} (64). Notably, we found that infection with \(10 \times 10^8\) bacteria significantly killed larvae after 24 h when they were infected with the MRSA USA300 strain (ST8) or clinical isolates of methicillin-susceptible \(\text{S. aureus}\) (MSSA) with different sequence types (ST1, -5, and -30). In contrast to a previous study with non-antibiotic-resistant \(\text{S. aureus}\) strains that did not identify a bacterial killing effect in the adult stage (42), we suggest here that the killing phenotype observed in larvae is due primarily to the amount of bacteria ingested. The epidemic strain \(\text{S. aureus}\) USA300 carries an hypervirulent phenotype characterized by the expression of multiple toxins (such as the enterotoxins K and Q and the Panton-Valentine leukocidin [PVL] pore-forming toxin) and the arginine catabolic mobile element (ACME), which displays adhesive properties and improves bacterial colonization (5, 65). All these specificities may play an important role in successful intestinal establishment.

It was recently shown that larvae orally infected with \textit{Erwinia carotovora} subsp. \textit{carotovora} 15 (\(\text{Ecc15}\)), \textit{Pseudomonas aeruginosa}, or \textit{Pseudomonas entomophila} are more susceptible to pathogens than adult flies infected with similar doses (66). The adult intestine undergoes basal turnover characterized by proliferation of intestinal stem cells (ISCs), which differentiate into intermediate progenitor cells named enteroblasts (EBs) and then into enterocytes (ECs) or enteroendocrine cells (EEs) (67). Upon infection with the Gram-negative pathogen \(\text{Ecc15}\) (68, 69) or \(P. \text{entomophila}\) (70), compensatory mechanisms, respectively, activated by the epidermal growth factor receptor (EGFR) and the JAK/STAT pathways initiate a strong mitotic response in the midgut, without modifying ISC number. This phenomenon is complementary to the intestinal antimicrobial response and essential to resist infection. Interestingly, it was also shown that the IMD pathway plays a key role in ECs shedding during infection, also favoring epithelial turnover (71). In contrast, \textit{Drosophila} larvae do not possess ISCs and, upon \(\text{Ecc15}\) intestinal infection, rely on adult midgut progenitors (66). These progenitor cells differentiate into ECs; however, the authors raise the hypothesis that these cells are
insufficient in number to meet the need of both intestinal repair and antimicrobial response. This may explain the particular sensitivity of the larvae to intestinal infections.

**Intestinal infection with *S. aureus* triggers host ROS production.** We found that *S. aureus* infection triggered ROS production in the gut of *Drosophila* larvae early in the infection and in a transient way. The *Drosophila* genome encodes one Duox enzyme, whereas two Duox homologs are identified in mammals (72). In flies, it was shown that the Duox enzyme could be activated by pathogen-derived uracil, unlike with commensal bacteria, which do not secrete this molecule (73). Of note, *S. aureus* USA300 is capable of generating uracil through pyrimidine metabolism (Kyoto Encyclopedia of Genes and Genomes pathway). We herein showed that live bacteria, as well as heat-killed bacteria, were able to increase Duox gene transcription, suggesting that both the *S. aureus* cell wall and secreted factors play a role in this mechanism. We confirmed Duox activation by measuring an increase in ROS generation, occurring in the first hours of infection. These data somewhat contradict an earlier work done by Hori et al. (42) reporting that *S. aureus* *Drosophila* feeding did not induce ROS production. This apparent discrepancy is likely due to the ROS quantitation method used in both studies. To quantify the ROS amount, Hori and colleagues used hydro-Cy3, a compound for which measurement may be influenced by mitochondrial membrane potential (74), which was shown to be modified during bacterial infection (75). This discrepancy may also be due to the method used to dissect the larval intestine. In insects, Malpighian tubules play a key role in detoxification and hemolymph filtering (as with the kidneys and liver in mammals). They are intimately linked to the stress status of the fly (76), and it was recently shown that Malpighian tubules play an active role during oral infection by sequestering excessive ROS and oxidized lipids (54). Including them during dissection might greatly affect results by hiding the specific intestinal ROS signal. In another model of orally infected black soldier flies, *S. aureus* was shown to be able to induce Duox gene expression, as well as to increase the H$_2$O$_2$ concentration, in a short time frame (77). Overall, this work confirms the importance of generating intestinal oxidative stress to clear colonizing pathogens as well as the necessity for the bacterium to acquire efficient oxidative-stress-resistant systems. Our results demonstrate that the *S. aureus* USA300 katA gene is necessary to increase bacterial virulence in *vivo* and to assess its colonization capacities. Of note, the importance of the *S. aureus* catalase gene has previously been shown in *vitro* during intracellular infection in murine macrophages or in *vivo* through intraperitoneal injection with a ΔkatA clinical bovine strain (78).

Our work has demonstrated the link between ROS production and activation of Toll signaling in the guts of *Drosophila* larvae after exposure to *S. aureus* USA300 (Fig. 5). Presumably, this mechanism potentiates the host immune response against harmful pathogens, such as *S. aureus*. The established link between ROS and Toll pathway initiation in *Drosophila* was established previously, and it was also shown that *Wolbachia*-infected mosquitoes exhibited an increase in Duox2 transcription that was sufficient to induce transcription of the Toll pathway-sensitive AMPs cecropins and defensins (79). Interestingly, research in mammals suggests that ROS may alter the activity of the ikB kinase (IKK) complex in the cytoplasm or the DNA-binding capacity of NF-κB in the nucleus. (80). These observations highlight the need for bacteria to consistently control the host clearance strategy by simultaneously acting on the immune response and the ROS pool.

**katA undergoes different regulatory networks in *vitro* and in *vivo***. The present work also shed light on the role of SigB and Agr in the expression of *S. aureus* katA in *vitro* or in *vivo* (Fig. 5). We first observed *in vitro* that SigB and Agr did not influence katA transcription under the nontreated condition and showed that, upon H$_2$O$_2$ treatment, SigB upregulated katA expression. These data are in agreement with previous observations reporting the increased susceptibility of a ΔsigB mutant (Newman and MSSA backgrounds) to H$_2$O$_2$ treatment (81, 82). However, we have also shown that SigB played a minor contribution to the regulation of katA because, in the presence of H$_2$O$_2$, sigB expression was reduced and its negative influence on Agr expression (83).
was repressed. Hence, we propose here that Agr is an important new intermediate in katA upregulation.

In vivo, S. aureus can undergo multiple intestinal stresses, such as pH changes, iron sequestration (84), interactions with microbiota (85), antibiotic treatments (86), and oxygen or nutrient limitations (87). Somewhat unexpectedly, we found that the katA gene was upregulated in sigB and agrC mutants, suggesting a minor role for these master regulators in its in vivo regulation and implying the contribution of multiple additional cues. Notably, several infection models suggest that SigB was not important for bacterial virulence in the mouse abscess model (88), osteomyelitis, pyelonephritis (89), or nematode digestive infection (90). Thus, one can speculate that in this complex environment, resistance to oxidative stress, despite a clear role in limiting bacterial clearance, may also allow the acquisition of a fitness advantage.

Finally, we reported that intestinal infections with S. Typhimurium and S. flexneri induced death in Drosophila larvae and that this was associated with an antimicrobial response consisting of the IMD pathway and ROS production. We therefore propose that this infection model can serve as an easily manipulated alternative to mammalian models to study innate host immune responses triggered during infection with human bacterial pathogens.

MATERIALS AND METHODS

Bacterial strains. The epidemic clone S. aureus USA300-LAC (designated S. aureus USA300 WT) as well as its isogenic derivatives were provided by Biodfense and Emerging Infections Research Resources (BEI Resources). S. aureus growth was performed in brain heart infusion (BHI) broth or agar or chromID MRSA agar at 37°C. Micrococcus luteus was grown in Luria-Bertani (LB) broth at 30°C. Salmonella Typhimurium SL1344 was grown in LB broth or agar at 37°C, and Shigella flexneri was grown in tryptic soy broth or agar (TSA) supplemented with Congo red dye (final concentration, 0.01%) to induce a type 3 secretion system (T3SS)-dependent secretion of virulence factors (91). Only pigmented colonies from TSA plates were used to prepare liquid cultures. pRN11 (48)- and pCNS7 (92)-carrying strains were transformed by electroporation, with the following settings: 2,450 V, 100 μF, 25 μF, and a time constant of 2.3 to 2.5 ms. For heat-killed bacteria, overnight-grown cultures were centrifuged, the supernatant was removed, and bacteria were suspended in DPBS. Bacteria were heated at 80°C for 20 min and further used for infection.
For complemented strains, we used the pCN57 shuttle vector. For the pCN57-<i>cp-kata</i> plasmid, the PCR product of the pCN57 vector treated with restriction enzyme DpnI to eliminate the template plasmid was fused to the catalase gene amplicon, followed by a kanamycin resistance cassette (kanamycin resistance gene <i>aphII</i> fused with the <i>pGro</i> promoter; 1,111 bp in total). All PCR products were assembled with the HiFi DNA assembly master mix kit. Similarly, for the pCN57-<i>cp-agrC</i> plasmid, the three PCR products (pCN57, <i>agrC</i>, and <i>kan</i> cassette) were purified and fused with the HiFi DNA assembly master mix. For the pCN57-<i>cp-sigB</i> plasmid, we used pCN57-<i>cp-agrC</i> as the template to amplify the pCN57 vector with the kanamycin cassette. This PCR product (after treatment with the restriction enzyme DpnI) was fused to the <i>sigB</i> PCR product using the HiFi DNA assembly master mix kit.

All strains and plasmids are defined in Table 1.

### Whole-genome sequencing and analysis

Genomic DNA was extracted using a DNeasy blood and tissue kit. Genomic libraries were prepared using a Nextera XT kit and then multiplexed and sequenced on an Illumina MiniSeq instrument (2 × 150 paired-end reads).

Single-nucleotide polymorphisms (SNPs) and small indels were assessed using Snippy v3.1 (https://github.com/tseemann/snippy). Briefly, Snippy was used to map the raw short reads against the annotated assembly of the parental strain (<i>Staphylococcus aureus</i> strain JE2; GenBank accession no. CP020619.1).

Assemblies were collected with a Unicycler assembler, available through PATRIC version 3.6.8 (93, 94). Prokka version 1.14.0 (95) was used to annotate the assemblies by allowing the verification of the transposon insertion site.

### Drosophila stocks and rearing

<i>Drosophila melanogaster</i> was maintained on a fresh medium prepared with the Nutri-Bloomington formulation (Genesee Scientific, San Diego, CA, USA), supplemented with 64 mM propionic acid. N-Acetyl-cysteine (NAC)-supplemented medium was prepared at the final concentration of 1 mM (A9165; Sigma) (96).

All <i>Drosophila</i> stocks are defined in Table 2.

### Infection experiments

Oral infections were performed on mid-L3 larvae (3.5 days after egg laying). For each test, animals were placed in a 2-ml microcentrifuge tube filled with 200 μl of crushed banana and 200 μl of overnight bacterial culture for 30 min. Bacterial infectious doses were adjusted by measuring culture turbidity at an optical density at 600 nm (OD<sub>600</sub> considering that an OD<sub>600</sub> of 1 is 5 × 10<sup>8</sup> bacteria/ml). Animals were blocked by a foam plug to be sure that they remained in the bottom of the tube for the whole infection time. After 30 min, they are washed briefly in 30% ethanol and placed in a petri dish with fresh fly medium without yeast. Infections and waiting times were performed at 29°C. Larvae were dissected at the indicated time points for reverse transcription-quantitative PCR (RT-qPCR) analyses, bacterial counts, and ROS quantification.

For oral infection of adult flies, 5- to 7-day-old adults were starved for 2 h in empty vials at 25°C. After starvation, flies were flipped into an infection vial with medium and completely covered with a Whatman paper disk. The disk was soaked with 100 μl of a 5% sucrose solution supplemented or not with antibiotics at the indicated infectious doses. After 30 min of oral infection, flies were flipped to fresh fly medium without yeast (changed every day).

**CFU counts.** At the indicated time points, larvae were dissected (at least 10 animals per point) and guts homogenized in 400 μl of DPBS (Gibco, ThermoFisher Scientific, MA, USA) by following the manufacturer's instructions. For CFU counts from hemolymph, animals were briefly washed in 70% ethanol, rinsed in sterile DPBS, and bled into a 200-μl DPBS drop on the slide. Samples were directly plated on BHI agar plates for <i>S. aureus</i> counts or LB agar for <i>M. luteus</i>.

**pH survival assay.** Assays were performed in BHI agar with an adapted pH. pH was adjusted with sodium hydroxide or hydrochloric acid solutions at the selected conditions: pHs of 3, 5, 7, 9, and 11. Media adjusted at pHs 3 and 5 were buffered with 2-(N-morpholino)-ethane sulfonic acid (MES; 30 mM; Euromexed, Safflewysersheim, France), at pH 7 with Tris-(hydroxymethyl) ammonium (Tris; 30 mM; Merck), and at pHs 9 and 11 with 2-(N-cyclohexylamino)ethane-sulfonic acid (CHES; 30 mM; Sigma-Aldrich). Fresh bacterial cultures that reached an OD<sub>600</sub> of 0.3 to 0.6 were washed once with PBS and then diluted in the different buffers to reach the concentration of 2 × 10<sup>7</sup> bacteria/ml. At the indicated time points, 50 μl from each culture was sampled, serially diluted, and plated on BHI agar.

**ROS quantification and visualization.**

(i) **ROS quantification.** The amount of ROS in dissected guts (from 10 animals) was estimated using 2′,7′-dichlorodihydrofluorescein diacetate (H<sub>2</sub>DCFDA; C6827; ThermoFisher Scientific, MA, USA) by following the manufacturer's instructions. For larval gut dissection, we carefully removed the Malpighian tubules, as they can strongly influence ROS level, and then tissues were homogenized in H<sub>2</sub>DCFDA mix. Fluorescence was measured 30 min after the mix preparation in a Berthold TriStar LB941 plate reader (Berthold France SAS, Thoiry, France). Results were normalized to those for total protein for each sample. The protein concentration was quantified using the Pierce bicinchoninic acid (BCA) colorimetric assay (Life Technologies, CA, USA) by following the manufacturer's instructions.

(ii) **ROS visualization.** Guts were dissected at the indicated times on glass slides, incubated in H<sub>2</sub>DCFDA (10 μM) for 15 min, and live imaged with a Zeiss Axio Imager Z2 Apotome microscope.

### Larval imaging

(i) **Whole-gut stainings.** Guts were dissected in PBS, fixed for at least 1 h at room temperature in 4% paraformaldehyde in PBS, and permeabilized in PBS plus 0.1% Triton X-100 for 30 min. They were stained with BODIPY 493/503 at a 1/100 dilution (D3922; ThermoFisher Scientific, MA, USA) for 1 h, stained with DAPI (4′,6-diamidino-2-phenylindole) at the dilution 1.43 μM for 10 min, washed with PBS, and mounted in Mowiol 4-88 (17951-500; BioValley, France).

(ii) **Peroxidase staining.** After fixation in 4% paraformaldehyde in PBS, guts were incubated in a peroxidase staining solution for 30–60 min at room temperature. They were washed 3× for 30 min in PBS, and washed again in 5% sucrose in PBS to permeabilize the gut lining. The gut was incubated for 1 h in a paraformaldehyde solution containing 10% sucrose in PBS, washed 3× in 5% sucrose in PBS, and then incubated overnight in an immersion solution containing 5 mg/ml diaminobenzidine (DAB; Sigma-Aldrich), 0.05% hydrogen peroxide, and 0.001% sodium azide. After rinsing in PBS, guts were incubated in a clearing solution containing 5% sucrose in PBS and 5% glycerol for 1 h.

(iii) **Hemolymph microbiota.** Larvae were killed by freezing and were washed 3× with PBS to remove bacteria from the outer cuticle. Larvae were homogenized in PBS, serially diluted, and plated on BHI agar.
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<th>Strains and plasmids used in this study</th>
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<td><strong>Plasmids</strong></td>
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**TABLE 1** Strains and plasmids used in this study. | Relevant characteristics |
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For sample preparation, animals were first fixed in ScaleCUBIC-1 (reagent 1) for at least 4 days and cleared in ScaleCUBIC-2 (reagent 2) for at least 2 days according to the method of Susaki et al. (97). Briefly, to prepare 500 g of reagent 1 solution, 125 g of urea and 156 g of 80% (by weight) Quadrol were dissolved in 144 g of distilled water (dH2O). After complete dissolution under agitation, we added 75 g of Triton X-100 and then degassed the reagent with a vacuum desiccator (0.1 MPa, 30 min) (97). Then, samples were cleared with ScaleCUBIC-2 (reagent 2). To perform the Lightsheet fluorescence microscopy (LSFM) imaging, samples were embedded in 4% low-melting-point agarose (ThermoFisher Scientific, France) dissolved in R2 medium by using a glass cylindrical capillary, and we allowed embedding overnight. Images were acquired with a Lightsheet Z.1 microscope (Carl Zeiss, Germany) equipped with a Plan-Apochromat 20/R2-immersion lens objective with left and right illumination.

RT-qPCR. For mRNA extraction, dissected guts were collected at the indicated time points and homogenized with a Mikro-Dismembrator S (Sartorius Stedim, Aubagne, France). Total RNA was isolated using Trizol reagent and dissolved in RNase-free water. Five hundred nanograms of total RNA was then reverse transcribed in a 20-μl reaction volume using the LunaScript RT supermix kit (E3010; New England Biolabs, MA, USA). Quantitative PCR was performed by transferring 2 μl of the RT mix to the qPCR mix prepared with Luna Universal qPCR master mix (M3003; New England Biolabs, MA, USA) according to the manufacturer’s recommendations. All the primers used for this experiment are defined in Table 3, and their amplification efficiency was checked before any further analysis. Reactions were performed on a 7900HT Fast real-time PCR system (Applied Biosystems) according to the standard settings of the system software. The thermal cycling conditions were as follows: an initial denaturation at

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<td>ywDD1; (control line)</td>
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<tr>
<td>yw drs-GFP dipt-LacZ;spz+/-/TM6c (ywDD1 + spz+/-/TM6c)</td>
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<td>w NP3084-GAL4</td>
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<td>w UAS-Duox RNAI/CYO</td>
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Acknowledgments

We thank D. Ferrandon and S. Liégeois (IBMC, Strasbourg, France) and W. J. Lee (SNU, Seoul, South Korea) for sharing fly stocks and bacteria. We are grateful to the core imaging facility of the Structure Fédérative de Recherche Necker, INSERM US24 CNRS UMS 3633, for its technical support. We thank Thierry Capiod (INEM, Paris, France) for technical help in measuring H2-DCFDA fluorescence. The following reagents were provided by the Network on Antimicrobial Resistance in Staphylococcus aureus (NARSA) for distribution by BEI Resources, NIAID, NIH: USA300 WT and USA300 ΔkatA, USA300 ΔsodA, USA300 ΔsodM, USA300 ΔgpxA1, USA300 ΔgpxA2, USA300 ΔsigB, and USA300 ΔagrC.

This work was supported by grant ANR-15-CE15-0017 from StopBugEntry, INSERM, CNRS, and by grant ANR-18-IDEX-0001 from the IdEx Université de Paris.

We declare no conflict of interest.

Supplemental Material

Supplemental material is available online only.

Fig S1, TIF file, 0.1 MB.
Fig S2, TIF file, 0.3 MB.
Fig S3, TIF file, 0.1 MB.
Fig S4, TIF file, 0.05 MB.
Fig S5, TIF file, 0.1 MB.
Fig S6, TIF file, 11.7 MB.
Fig S7, TIF file, 0.1 MB.
Movie S1, MOV file, 3.8 MB.

References

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