Burkholderia pseudomallei kills Caenorhabditis elegans through virulence mechanisms distinct from intestinal lumen colonization

Soon-Keat Ooi,1 Tian-Yeh Lim,1 Song-Hua Lee1,† and Sheila Nathan1,2,*

1School of Biosciences and Biotechnology; Faculty of Science and Technology; Universiti Kebangsaan Malaysia; Bangi, Selangor, Malaysia; 2Malaysia Genome Institute; Kajang, Selangor, Malaysia

Keywords: Burkholderia pseudomallei, Caenorhabditis elegans, host-pathogen interaction, intestinal lumen colonization, toxin

Abbreviations: Bp R15, B. pseudomallei R15; ompA, outer membrane protein A gene; GFP, green fluorescence protein; R15-GFP, GFP-tagged Bp R15; Glp, germ line proliferation-deficient; CFU, colony forming unit; ABC, ATP-binding cassette; Cm, chloramphenicol; Cb, carbenicillin; TDmean, mean time to death; Exp, expulsion; Lev, levamisole

The nematode Caenorhabditis elegans is hypersusceptible to Burkholderia pseudomallei infection. However, the virulence mechanisms underlying rapid lethality of C. elegans upon B. pseudomallei infection remain poorly defined. To probe the host-pathogen interaction, we constructed GFP-tagged B. pseudomallei and followed bacterial accumulation within the C. elegans intestinal lumen. Contrary to slow-killing by most bacterial pathogens, B. pseudomallei caused fairly limited intestinal lumen colonization throughout the period of observation. Using grinder-defective mutant worms that allow the entry of intact bacteria also did not result in full intestinal lumen colonization. In addition, we observed a significant decline in C. elegans defection and pharyngeal pumping rates upon B. pseudomallei infection. The decline in defection rates ruled out the contribution of defection to the limited B. pseudomallei colonization. We also demonstrated that the limited intestinal lumen colonization was not attributed to slowed host feeding as bacterial loads did not change significantly when feeding was stimulated by exogenous serotonin. Both these observations confirm that B. pseudomallei is a poor colonizer of the C. elegans intestine. To explore the possibility of toxin-mediated killing, we examined the transcription of the C. elegans ABC transporter gene, ppg-5, upon B. pseudomallei infection of the ppg-5::gfp reporter strain. Expression of ppg-5 was highly induced, notably in the pharynx and intestine, compared with Escherichia coli-fed worms, suggesting that the host actively thwarted the pathogenic assaults during infection. Collectively, our findings propose that B. pseudomallei specifically and continuously secretes toxins to overcome C. elegans immune responses.

Introduction

Burkholderia pseudomallei is a Gram-negative saprophyte that typically inhabits muddy soil and stagnant water throughout Southeast Asia and northern Australia.1 When acquired by humans and animals, B. pseudomallei can cause melioidosis, a life-threatening disease that, to this day, still presents a danger to most parts of the tropics.2 Decades of research on B. pseudomallei has only revealed how versatile this pathogen is, for example, it can (1) infect a multitude of organisms and invade a wide range of cell types,3,4 (2) result in a broad spectrum of clinical manifestations,5 (3) resist many clinical antimicrobials6 and (4) survive extremely harsh environmental conditions.7 Yet, the molecular mechanisms by which B. pseudomallei evades or modulates host immune responses remain elusive. Of the various forms of melioidosis, acute melioidosis raises the greatest medical concern owing to its high mortality rate regardless of appropriate antibiotic treatments.8 It is well documented that acute melioidosis tends to affect humans with risk factors such as diabetes mellitus but rarely immunocompetent individuals.9 For this reason, host models with clinically relevant predisposing backgrounds or sensitivities are particularly attractive in melioidosis research. To this end, several groups have recently begun to exploit host models engineered to mimic the risk factors for melioidosis, such as type 1 and 2 diabetic mice, in an effort to elucidate the attributes of B. pseudomallei virulence in the corresponding predisposed individuals.10-12

Over the last decade, there has been a growing appreciation that Caenorhabditis elegans can serve as a simple surrogate host for modeling bacterial diseases.13 C. elegans is also deemed a relevant host model for studying acute melioidosis. Diabetic patients prone to acute melioidosis have impaired innate immune responses such
as macrophage phagocytosis and migration. C. elegans lacks circulating phagocytes and some innate immune system components essential for fighting an acute B. pseudomallei infection; however, it is protected by an innate immune system conserved with that in humans. Akin to acute melioidosis individuals, C. elegans is highly susceptible to B. pseudomallei infection, which strongly suggests that B. pseudomallei executes its pathogenicity by suppressing or breaching the host innate immune system. In addition, it has been shown that B. pseudomallei does not persist within C. elegans, similar to observations in acute melioidosis cases. These infection characteristics together suggest that C. elegans is an excellent model to simultaneously dissect the evolutionarily conserved determinants of B. pseudomallei virulence as well as host innate immune defense mechanisms.

A myriad of bacterial virulence mechanisms has been unraveled using C. elegans. One of the strategies commonly employed by bacterial pathogens in establishing infection in mammals is adhesion and colonization of host tissues. Likewise, virtually all Gram-positive and Gram-negative pathogens studied to date can colonize the brush border microvilli of C. elegans after they successfully escape the grinder and resist antimicrobial peptides in the pharynx, ultimately leading to colonization and distension of the intestinal lumen. In general, this active infectious process takes place only when the pathogens are cultured on a minimal or "slow-killing" medium, and the extent of colonization often correlates with host killing. Nevertheless, subtle differences have been noticed in terms of the intestinal lumen colonization by these pathogens. For example, Pseudomonas aeruginosa and Staphylococcus aureus grossly colonize and distend the worm intestinal lumen but do not persist within the host, unlike other pathogens such as Escherichia coli (EPEC), Salmonella Typhimurium, Serratia marcescens and Yersinia enterocolitica.

Previously, we have established a C. elegans-B. pseudomallei model system and demonstrated that a clinical isolate, Human B. pseudomallei strain Bp R15 (referred to as Bp R15 henceforth), was able to rapidly kill BALB/c mice and C. elegans. However, how Bp R15 interacts with C. elegans to elicit extreme symptoms and death is not completely understood. In the present study, we introduced a gfp construct into Bp R15 to visualize the passage of the pathogen across the host. Our results revealed a novel host-pathogen interaction in which the virulent Bp R15 failed to fully colonize C. elegans intestinal lumen under "slow killing" conditions, even though facilitated with host grinder dysfunction and accelerated feeding. With the use of a C. elegans detoxification gene reporter strain, we provided evidence that the rapid host killing may be mediated by bacterial toxin secretion.

**Results**

**Construction and evaluation of GFP-expressing Bp R15.** To visualize the events following Bp R15 infection of C. elegans, we first labeled Bp R15 with GFP. To prevent overexpression of GFP that may contribute to reduced bacterial fitness and infectivity, we fused a moderate-strength gene promoter, PompA26 with a gfp coding sequence and cloned the DNA construct into a low-copy-number pBBR-derived vector (–20–30 copies/cell) to minimize potential burden imposed on the bacteria. Electroporation of the resultant recombinant plasmid, pSMompAgfp (Fig. S1A), into Bp R15 successfully generated R15-GFP with observable fluorescence under both in vitro (Fig. S1B) and in vivo (Fig. 1A–C) conditions.

Next, we tested the extrachromosomal stability of pSMompAgfp in R15-GFP under non-selective conditions. Loss of fluorescence was observed in a subpopulation of the bacteria after just one passage without chloramphenicol (Cm) (Fig. S2A), suggesting that pSMompAgfp was unstable and therefore Cm was required to select for R15-GFP. To verify that addition of Cm would not confound analyses of downstream infection assays, the C. elegans lifespan on E. coli OP50 plates supplemented with Cm was determined. The results showed that 100 μg/ml of Cm did not compromise worm lifespan and could be included in the assay media (Fig. S2B). The viability of R15-GFP could not be compared with that of Bp R15 as 100 μg/ml Cm is considerably bacteriostatic in the latter (data not shown).

As R15-GFP was to be used as a surrogate for Bp R15, we assessed the virulence of R15-GFP in C. elegans N2 to confirm that constitutive GFP expression did not interfere with the bacteria’s virulence. No significant difference was observed between Bp R15 and R15-GFP virulence (p > 0.01, Log-rank test) (Fig. S2C). In addition, both R15-GFP and Bp R15 were equally capable of causing intestinal discoloration, body shrinkage and contracted nose in C. elegans throughout the infection.

**Bp R15 does not heavily colonize the C. elegans intestinal lumen.** It has been suggested that B. pseudomallei, like other Gram-negative human pathogens such as P. aeruginosa, also gains entry into and colonizes the worm intestine. We followed Bp R15 as 100 copies by flow cytometry at various time-points throughout the infection. R15 were first labeled and electroporated with the resultant recombinant plasmid, pSMompAgfp into Bp R15. After 4 h of infection, we did not observe any colonization in the infected worms but merely single R15-GFP cells irregularly interspersed along the intestinal lumen (Fig. 1A). As the infection proceeded, a progressive increase in the colonization and distension of the worm anterior intestinal lumen was observed. Nevertheless, even up to 28 h post-infection, the majority of infected worms were not fully colonized. Instead, only the anterior intestinal lumen was, to a great extent, colonized and distended with a clump of R15-GFP (Fig. 1B). In addition, most infected worms contained R15-GFP cells in their pharyngeal tracts, and the pharynx was noticeably plugged with bacteria during late infection, with R15-GFP spanning the buccal cavity through the terminal bulb (Fig. 1C). It is worth noting that R15-GFP barely colonized the mid- and posterior intestine. We also noticed that the greatly distended anterior intestinal lumen was not always densely filled with R15-GFP cells, suggesting that the distension was due to secretion of an unknown extracellular matrix similar to that previously observed in a P. aeruginosa infection. In all the worms...
examined, R15-GFP colonization was confined to the pharyngeal tract and intestinal lumen, suggesting that the infection was extracellular. Since we did not observe bacterial entry via other parts of the worm body such as the cuticle, anus or vulva, we conclude that *B. pseudomallei* is an intestinal pathogen of *C. elegans*.

Detailed examination of the infected worms revealed that not every single worm displayed similar degrees of intestinal lumen colonization. To ensure a representative microscopic observation, we conducted a quantitative infection assay whereby 75–100 infected worms were scored at the same time points and classified based on the severity of intestinal lumen colonization. As depicted in Figure 1D, worms were scored as “full” when a tract of R15-GFP was seen from the anterior to posterior intestinal lumen, regardless of the density of bacteria. Visible but not full colonization of R15-GFP anywhere along the intestinal lumen was categorized as “partial” whereas when none or only scattered R15-GFP cells were present in the lumen observed at 400x magnification, this was classified as “undetectable.” As noted above, only limited R15-GFP colonization was observed in the infected worm population. We found that less than 10% of the population were fully colonized even up to 28 h post-exposure (Fig. 1E), a time point at which a majority of the worms were severely ill. This implies that *Bp* R15 is unable to substantially colonize the *C. elegans* intestinal lumen on minimal media over the course of infection.

To the best of our knowledge, this observation of limited intestinal lumen colonization has not yet been reported for “slow killing” of *C. elegans*. To further validate our observation, we enumerated the number of colonizing bacteria by performing a colony forming unit (CFU) assay under the same infection conditions. By using the method described by Shapira and Tan, we were able to achieve CFU counts of $10^3$–$10^5$ bacteria/worm for worms exposed to *P. aeruginosa* PA14 (PA14) (data not shown). On the other hand, for the R15-GFP infection, fairly low numbers of bacteria were enumerated within the infected worms throughout the infection period (in the range of $10^3$–$10^4$ CFU/worm; Fig. 1F), reinforcing the belief that *Bp* R15 did not substantially colonize the host intestinal lumen.

It is thought that the *C. elegans* pharyngeal grinder that crushes ingested bacteria before thrusting the bacterial corpses into the intestine might hinder *Bp* R15 from efficiently colonizing its intestinal lumen. In worms infected by *P. aeruginosa* and *S. Typhimurium*, abrogating the grinder function accelerated the intestinal lumen colonization, leading to faster host killing. Hence, we tested this hypothesis by exposing a grinder-defective mutant strain, *tnt-3(aj3)*, to R15-GFP and monitored the extent of intestinal lumen colonization at 4, 6, 8 and 11 h post-exposure. The *tnt-3(aj3)* mutant bears a recessive mutation in the tropomycin T gene which disrupts the grinder function; consequently, more intact bacteria can enter its intestine. We selected the aforementioned time points because a significant portion of the infected mutant population was severely sick beyond 12 h of infection (Yee and Nathan, in preparation). As expected, we noted a marked increase in R15-GFP colonization within the *tnt-3(aj3)* mutant intestinal lumen, but the hindgut was devoid of R15-GFP cells (data not shown). To quantify *tnt-3(aj3)* colonization, we scored 75–100 worms based on the same set of colonization criteria depicted in Figure 1D. Indeed, there was a significant increase in the number of fully colonized worms compared with N2 worms; however, the category percentage was only less than 70% at 11 h post-infection (Fig. 1G). We also conducted the CFU assay to quantitatively measure R15-GFP colonization within infected *tnt-3(aj3) mutant worms. Consistent with the visualization assay, the number of R15-GFP in infected *tnt-3(aj3)* worms did not significantly increase over the course of infection (Fig. 1H). Therefore, *Bp* R15 is inferior to other well-studied intestinal pathogens in terms of host colonization.

### Figure 1 (See opposite page).

R15-GFP displayed limited intestinal lumen colonization during *C. elegans* N2 infection. (A) A low number of R15-GFP cells were present along the intestinal lumen after 4 h of infection. (B) At 28 h post-infection, the major portion of the worm intestinal lumen was void of R15-GFP except for the anterior intestine which was grossly colonized and distended. Note that the worm body shrank and the nose was contracted at this time point. (C) Three representative worms here depict an increase in R15-GFP colonization within the anterior intestine upon prolonged pathogen exposure. After 22 h post-infection, both pharyngeal bulbs of the worm were distorted and contained proliferating bacteria. (D) Shown are the three categories of colonization criteria that were used to classify the infected worms. (E) Stacked bars correspond to mean proportion of the infected N2 population encompassing all three colonization categories (n = 75–100). (F) The graph shows the number of R15-GFP colonizing N2 worm intestines throughout the infection. (G) Stacked bars represent mean proportion of infected *tnt-3(aj3)* population within the three colonization categories (n = 75–100). (H) The graph shows the number of R15-GFP colonizing *tnt-3(aj3)* worms up to 11 h infection. (F and H) Each marker corresponds to the average bacterial CFU extracted from 10 infected worms; horizontal lines represent geometric means. Light blue arrows in (A–C) point either to individual R15-GFP cells or colonization as well as distension of the intestinal lumen. Asterisks in (A) and (B) mark the terminal pharyngeal bulb. Dotted circles in (C) mark the anterior and terminal pharyngeal bulbs. Images (A and B) were taken at 100x magnification while (C) was captured at 400x magnification. Scale bars in (B, C and D) represent 200 μm, 50 μm and 30 μm, respectively.
E. coli, Bp Typhimurium. With continuous feeding, to reverse the sluggish worm R15 in construct made by fusing the 300-bp 0.001 (Mann-Whitney U-test). (B) Bars correspond to transgenic worms with transcript 0.05, Bp R15 colonization. To clarify this, we measured Bp, S., 38 R15-infected was highly 0.001 (Student R15 only intermittently.

C. elegans gfp R15 colonization, we supplemented the assay R15 infection, we measured strongly induces the R15 virulence and its R15 intestinal lumen is not a result of routine worm defecation. Therefore, it was possible that the infected C. elegans pgp-5::gfp R15 colonization of the intestine, except at 8 h post-infection (Fig. 2A). Suffice to say, the inadequate Bp R15 colonization. To clarify this, we measured host pharyngeal pumping rates during Bp R15 infection. Surprisingly, we observed an almost immediate decline in worm pharyngeal pumping rates (average < 1 pump/sec) upon infection (Fig. 2B). Concomitantly, we also observed convulsion-like head contractions and relaxation in the infected worms similar to but slower than that described by Williams et al. With continuous monitoring of the pharyngeal pumping of infected worms, it was observed that the worms did not completely cease feeding but instead, extended the intervals between pumps, i.e., the infected worms fed on Bp R15 only intermittently.

The dramatic decline in Bp R15-infected C. elegans pharyngeal pumping rates raised the question of the possible impact of host feeding on the number of bacteria colonizing the intestine. To test whether the decrease in feeding rate was a host strategy to mitigate Bp R15 colonization, we supplemented the assay medium with 5 mM serotonin, a neurotransmitter that can stimulate C. elegans feeding, to reverse the sluggish worm pharyngeal pumping. This drug concentration has been shown to increase feeding rates to 3–4 pumps/sec in E. coli-fed worms. We first assessed the drug efficacy by demonstrating that 5 mM serotonin enhanced pharyngeal pumping rates in worms fed on R15-GFP (Fig. 3A, p < 0.01, Student’s t-test), although the stimulatory effect weakened over time. To ensure that the exogenous serotonin did not significantly affect defecation, we measured the worm defecation rate in the presence of serotonin. A higher rate of defecation was indeed apparent in the treated worms, but the increase was not significant (Fig. 3B, p > 0.05, Mann-Whitney U-test). Next, we performed a CFU assay on the serotonin-stimulated infected worms, with untreated infected worms as a control. However, we failed to observe any significant difference in the number of colonizing bacteria in the serotonin-induced worms (Fig. 3C, p > 0.05, Mann-Whitney U-test). While this result suggests that the reduction in host feeding rates did not contribute to the limited intestinal lumen colonization, it also implies that the primary virulence mechanism employed by Bp R15 in C. elegans killing is distinct from intestinal lumen colonization.

C. elegans strongly induces the pgp-5 gene in response to Bp R15 infection. As the degree of intestinal lumen colonization was not commensurate with the level of Bp R15 virulence and its ability to kill C. elegans, we asked whether rapid host killing was mediated by secreted bacterial toxins. A previous study has demonstrated that C. elegans pgp-5 plays a significant role in the worm’s defense against P. aeruginosa and S. Typhimurium infections as well as cadmium and copper intoxication. C. elegans pgp-5 encodes a transmembrane P-glycoprotein, a conserved member of the ATP-binding cassette (ABC) transporter superfamily, which is predicted to mediate detoxification of exogenous toxins by the host. To determine changes in host pgp-5 expression upon Bp R15 infection, we measured pgp-5 transcript levels in Bp R15-infected worms using quantitative RT-PCR (qRT-PCR). We found that the transcription of pgp-5 was highly induced at 4 and 12 h post-infection (140- and 290-fold, respectively; Fig. 4A).

To determine the spatio-temporal tissue distribution of pgp-5 expression, we infected pgp-5::gfp transgenic worms with Bp R15 and monitored changes in the transgene expression using E. coli OP50-fed worms as a control. These transgenic worms harbor an integrated pgp-5::gfp construct made by fusing the 300-bp pgp-5 promoter region to a gfp coding sequence, thus any green fluorescence observed would reflect the transcription of pgp-5. Consistent with the findings of Kurz et al., worms fed on E. coli

![Image](https://example.com/image.png)
exhibited very weak green fluorescence in the intestine throughout the observation period (Fig. 4B). In contrast, bright green fluorescence was visible in the pharynx (from isthmus to terminal bulb) and intestine of Bp R15-infected worms as early as 4 h post-exposure (data not shown). We also noted that the green fluorescence gradually increased in intensity over time. At 28 h post-infection, intense green fluorescence was apparent in the pharynx (from the isthmus through the terminal bulb), intestine as well as gonad (Fig. 4C–E). To confirm the steady increment of pgp-5 induction in the worm population upon Bp R15 infection, we also visually quantified the green fluorescence during the assay. Twenty live worms were randomly picked at 4, 12 and 28 h post-infection, observed under 100x magnification and scored based on defined criteria (Fig. 4F, upper panel). Briefly, a score of “1” was assigned to worms with very dim green fluorescence. Worms with detectable green fluorescence in the pharynx and intestine were given a score of “2” while a score of “3” was assigned to worms displaying intense fluorescence in the pharynx, intestine and gonad. This visual quantification confirmed that the pgp-5 induction level was significantly higher in Bp R15-infected worms (Fig. 4F, lower panel, p < 0.001, Mann-Whitney U-test), thus underscoring a strong yet specific detoxification process that occurred in response to Bp R15 infection.

In light of the high and specific induction of pgp-5 expression, we next asked if disrupting gene expression would render C. elegans more susceptible to Bp R15 infection. To answer this question, we fed rrf-3(pk1426);glp-4(bn2) double mutant worms with E. coli expressing dsRNA directed against pgp-5 and assessed survival upon Bp R15 infection at 25°C. RNAi knock-down of pgp-5 expression indeed resulted in greater susceptibility of C. elegans toward Bp R15; however, the difference in survival was not significant compared with worms treated with the L4440 vector (TD\text{mean}: 40.19 ± 0.69 and 41.86 ± 0.69 h, respectively; p = 0.115, Log-rank test). To better resolve the subtle difference in survival observed at 25°C, we repeated the assay at 16°C. We demonstrated that C. elegans was significantly more susceptible to Bp R15 killing upon RNAi silencing of pgp-5; TD\text{mean} values for pgp-5 RNAi-treated and control worms were 99.77 ± 1.47 h and 108.28 ± 1.68 h, respectively (Fig. 4G, p < 0.0001, Log-rank test), thus highlighting a protective role for pgp-5 during Bp R15 infection. This increased resolution of host susceptibility at lower temperatures was also observed in P. aeruginosa and S. Typhimurium infections.

**Discussion**

Although C. elegans has long been established as a model organism for investigating B. pseudomallei pathogenicity, much remains to be learnt about how this pathogen severely damages the host. Previous studies have asserted that B. pseudomallei infection of C. elegans is an active process involving both live bacteria and continuous intoxication. However, at the organismal level, whether the interaction between B. pseudomallei and C. elegans is similar to other bacterial species, has yet to be clarified. Our earlier study had demonstrated that Bp R15 did not persistently colonize the C. elegans intestinal lumen. In this study, we sought to...
investigate the host-pathogen interaction in greater detail, hoping to shed light on the possible complex mechanism(s) by which *B. pseudomallei* kills *C. elegans*.

As *B. pseudomallei* is phylogenetically related to *P. aeruginosa* and they share a number of features in terms of infection (both are opportunistic pathogens that infect human lungs), we initially anticipated that R15-GFP would colonize the worm intestinal lumen to an extent comparable to that of PA14 but with faster kinetics owing to the lower TD~mean~ of *Bp* R15-infected worms (~36 h, Fig. S2C) compared with PA14-infected worms (TD~mean~ of ~75 h). However, we demonstrated that *Bp* R15 failed to extensively colonize the *C. elegans* intestinal lumen even during late infection. Furthermore, the bacterial CFU counts of *Bp* R15-infected worms are much lower (in the magnitude of 10^2 CFU/worm) compared with other bacterial infections such as *Y. enterocolitica*, which can reach up to ~10^6 CFU/worm. A similar lack of intestinal lumen colonization was also observed when *Bp* R15 was replaced with the *Bp* K96243 reference strain. These
findings collectively suggest that virulent *B. pseudomallei* isolates are universally incapable of heavily colonizing the *C. elegans* intestinal lumen. This suggestion is further supported by the finding that *Bp* R15 does not fully colonize the *tnt-3(aj3)* mutant worm population over the course of infection.

The relatively poor intestinal lumen colonization ability of *B. pseudomallei* could also be a consequence of other host barriers or factors that interfere or prevent *Bp* R15 from efficient colonization. For example, the hostile intestinal lumen microenvironment may impede *B. pseudomallei* intraluminal growth or replication. On the other hand, unfavorable host factors may include suboptimal pH (~4.1), temperature (~25°C, depending on the assay temperature) and oxygen concentration. Moreover, the *C. elegans* intestinal lumen may not be conducive to colonization owing to continual gut peristalsis as well as circulation of antimicrobial peptides and reactive oxygen species. Interestingly, our group has observed the suppression of a number of *C. elegans* antimicrobial peptide genes known to abate bacterial colonization such as *lys-7* and *spg-1* during *Bp* R15 infection (Fig. S3). Upon RNAi knock-down of these anti-microbial peptide genes, intestinal lumen colonization by *P. aeruginosa* and *S. Typhimurium* is aggravated. Nevertheless, even in a background of reduced antimicrobial peptide expression, *Bp* R15 still failed to fully colonize the worm gut. This reinforces our suggestion that *B. pseudomallei* is inherently less adept at colonizing the worm gastrointestinal tract and instead, relies on other mechanisms to kill the host. Understanding why *Bp* R15 fails to colonize the host intestinal lumen may help uncover host factors inhibitory to *B. pseudomallei* growth or survival in vivo which may aid the future development of anti-infective drugs against melioidosis.

*C. elegans* feeding rates were instantly and dramatically reduced upon *B. pseudomallei* infection, which implies a low nutritional value of the *B. pseudomallei* lawn as perceived by the host. Previous studies have demonstrated that the worm has the ability to avoid or move away from a population of pathogenic bacteria. However, our study is the first report of a decrease in host feeding as a response to the presence of virulent bacteria. Although the decreased *C. elegans* feeding rate was not a contributory factor to the limited *Bp* R15 colonization, it is possible that this unique host response may confer an advantage to host survival as it could provide the host with time to cope with the toxic insults incurred from consuming the pathogen. Song and Avery recently demonstrated that food intake in *C. elegans* consists of two actions, pharyngeal pumping and isthmus peristalsis, and activation of both are dependent on the interaction between serotonin and its receptor SER-7 present in a small number of motorneurons. A search for genes known to govern these pharyngeal motions from our previous microarray data surprisingly revealed that a handful of genes regulating serotonin production and signaling (*sph-1, bas-1, ser-7* and *cat-4*) were indeed suppressed during *Bp* R15 infection (Lee et al., in preparation). Furthermore, we noted the induction of a number of neuropeptide genes (*flp-1, flp-3* and *flp-13*) known to potently inhibit pharyngeal pumping. Taken together, these data may explain how the host feeding rate was markedly alleviated upon *Bp* R15 infection. It is tempting to speculate that *B. pseudomallei*, as a successful environmental survivor, secretes certain chemorepellents to ward off predation by bacteriophage nematodes.

Our infection assay data indicated that a low number of intestinal *Bp* R15 bacterial cells were sufficient to evoke the observed symptoms and importantly to elicit death of *C. elegans*. From the pathogen’s perspective, *Bp* R15 may deploy other mechanically distinct yet successful virulence mechanisms that can compensate for the lack of intestinal lumen colonization. In our previous study, we found that diffusible toxins played a minor role in *C. elegans* killing. It is thus plausible to propose that, while remaining in the intestinal lumen, *Bp* R15 secretes extracellular toxins to damage the host. To support this proposition, our interrogation of the spatio-temporal expression of *pgp-5*, a known detoxifying ABC transporter, revealed that the infected worms actively engaged in a detoxification process upon *Bp* R15 infection. In addition, the loci that displayed heightened *pgp-5* induction—the pharynx and intestine—coincided with the locality of *Bp* R15 colonization, suggesting that the low number of *Bp* R15 cells secreted toxins into the nearby cells and triggered a host detoxification response. We believe that these toxins may translocate to distant cells upon prolonged infection as *pgp-5* induction was also observed in other cell types (gonad) during late infection. Interestingly, the pharyngeal *pgp-5* induction observed in our study is not seen in *P. aeruginosa* infection but only in worms exposed to 100 μM cadmium. This raises the question of whether *Bp* R15-secreted toxins are distinct from that produced by other pathogens. Similar to *P. aeruginosa* and *S. Typhimurium* infections, reducing or abrogating *C. elegans* *pgp-5* expression resulted in higher sensitivity to *Bp* R15 infection. Hence, we have shown that host detoxification constitutes an important immune defense mechanism against *Bp* R15 attack, and by extension, this implies that *Bp* R15 secretes toxins to kill the host.

Our previous study demonstrated that efficient killing of *C. elegans* necessitates continuous contact with *Bp* R15 since transient exposure to the pathogen failed to exert complete host killing. This strict demand for exposure to live *Bp* R15 may be a reflection of the inability of *Bp* R15 to proliferate within the *C. elegans* digestive tract as well as the strong and last-lasting induction of the host detoxification machinery. Nevertheless, over the extended period of infection (25–30 h), continuous intoxication by *Bp* R15 was sufficient to overwhelm the host immune system, causing an abrupt drop in the number of live worms (Fig. S2C).

In conclusion, our study presents further evidence for the requirement of prolonged *B. pseudomallei* presence and supplementary intoxication in host killing as previously suggested. Our findings also led us to infer that *B. pseudomallei* kills *C. elegans* via continuous and specific secretion of extracellular toxins. Hence, there is an urgent need to identify the toxins responsible for the rapid demise of *C. elegans*. This in turn will advance our understanding of the interaction between *B. pseudomallei* and *C. elegans*, and may push the development of therapeutic countermeasures against acute melioidosis.
Materials and Methods

Bacterial and C. elegans strains and growth conditions. The bacterial strains, worm strains and plasmids used in this study as well as the related information are listed in Table 1. Wild-type Bp R15 was routinely propagated on Ashdown agar whereas R15-GFP was cultured on modified Luria-Bertani (LB) agar (3% tryptone, 1% yeast extract and 1.5% bacteriological agar) supplemented with 100 μg/ml Cm. For cloning experiments, E. coli TOP10 transformants harboring pSMompAgfp were selected on LB agar with 30 μg/ml Cm. E. coli HT115 expressing dsRNA directed against the cdc-25.1 gene (cdc-25.1 RNAi clone) was grown on LB agar containing 100 μg/ml carbenicillin (Cb). All bacterial cultures were aerobically incubated at 37°C unless otherwise stated. All the experiments involving Bp R15 were performed in a BSL2+ level laboratory and approved by the Universiti Kebangsaan Malaysia Animal Ethics Committee (UKMAEC). Wild-type C. elegans Bristol N2 (N2), tnt-3(aj3) and rrf-3(pk1426);glp-4(bn2) strains were obtained from the Tan Laboratory, Stanford University whereas the ppgp-5::gfp transgenic strain was provided by the Shapira Laboratory, University of California Berkeley. All worm strains were handled as previously described.

Fluorescent labeling of Bp R15. For construction of the recombinant plasmid pSMompAgfp, a 234-bp promoter region of the outer membrane protein A precursor gene (PompA) (GenBank: BPSL2522) was amplified from B. pseudomallei D286 genomic DNA with primers OMPAF1 (5’-CACATACTAGTG ACCGATGTAGGTGCGG-3’) and OMPAR1 (5’- CTC ACCATTTCTCTCTCAGAAATGGAGA-3’). The 742-bp Aequorea coerulescens gfp ORF was amplified from pAcgfp1 (BD Biosciences Clontech, 632468) with primers GFPF1 (5’- CGAGAAGAGAAAATCGTTGAGCGCAAGG-3’) and GFR1 (5’-CGCCTGACTCTTACGTACGCTCCAGCTCATCCAT-3’). Spe1 and PstI restriction sites were incorporated into the 5’ end of OMPF1 and GFR1 primers respectively (underlined sequence) to generate cohesive termini compatible with pSM202 multiple cloning site 2. Using overlap-extension PCR, both amplicons were fused at a 21-bp overlapping region generated at the 5’ end of PompA negative strand and gfp ORF positive strand (italicized sequence), then amplified with OMPAF1 and GFR1 to form the PompA-GFP construct. Subsequently, the construct was double-digested and ligated with pSM202, forming pSMompAgfp, which was then transformed into E. coli TOP10 using standard CaCl2-mediated transformation. Transformants were screened by PCR and the sequence was verified through dideoxy DNA sequencing. High-fidelity Pfu DNA polymerase (Promega, M7741) was used for all DNA amplifications.

For preparation of electrocompetent cells and electroporation, a B. pseudomallei glycerol stock was diluted 100-fold in 50 ml SOB...
medium and cultured at ambient temperature (20–25°C) until the cell density reached OD$_{595}$ nm of ~0.4. The bacterial culture was chilled on ice for 30 min and then harvested by centrifugation at 6,000 g for 5 min at 4°C. Next, the bacterial cells were washed three times and subsequently suspended in an appropriate volume of ice-cold 10% glycerol to shield the cell surface charges. Approximately 300 ng purified pSMomp/Afp was added into 200 μl electrocompetent B. pseudomallei suspension and incubated on ice for 30 min. After that, the cell suspension was transferred into a 2 mm gap-width electroporation cuvette and subjected to electrical pulse (2 kV; 186 Ω; 40 μF; BTX Electroporation Apparatus, Harvard). Immediately after electroporation, 1 ml of SOC was added to the cell suspension followed by incubation at 37°C with agitation for 4 h. The electrocompetent cell suspension was concentrated 10-fold, after which 100 μl cells were spread onto modified LB agar supplemented with 100 μg/ml Cm and grown for 36 h. The presence of green fluorescence within the transformants was assessed using a Leica M205 FA stereomicroscope equipped with a GFP2 filter cube (bandpass 480/40 nm) (Leica Microsystems GmbH).

Preparation of Glp worms. N2 worms used in all assays were sterilized by RNAi knockdown of the cdc-25.1 gene. This gene encodes a CDC25 phosphatase homolog which affects embryonic viability and is necessary for cell proliferation of the germ line. The dsRNA directed against cdc-25.1 was introduced into worms by feeding. Briefly, 100 μl of the cdc-25.1 RNAi clone was cultured overnight at 37°C in LB medium supplemented with 100 μg/ml Cb. The culture was concentrated 25-fold, seeded onto NG agar supplemented with 1 mM Isopropyl β-D-1-thiogalactopyranoside (IPTG) (Promega, V3953) and 100 μg/ml Cb and incubated at room temperature for at least 24 h. Wild-type N2 gravidas were picked onto cdc-25.1 RNAi plates for two successive rounds of egg laying, each lasting for 4 h at 25°C. The worms were then removed and the eggs were allowed to hatch and develop into sterile worms on the plates.

Infection of N2 and mt-3(aj3) mutant worms. The conditions used in all assays involving R15-GFP were standardized and are described in this paragraph. To prepare the bacterial lawn, 2–3 freshly streaked R15-GFP colonies were inoculated into BHI medium supplemented with 100 μg/ml Cm and grown with agitation for 22 h at 37°C. By using 60 mm diameter plates, 55 μl of the overnight culture was spread onto NG agar supplemented with 100 μg/ml Cm. The culture was concentrated 25-fold, seeded onto modified LB agar supplemented with 100 μg/ml Cm using the drop plate method with modifications. Colonies were counted after incubating the plates at 37°C for 48 h. Average colony numbers obtained from visually separate colonies were used for statistical analysis. Bacterial CFU per worm was calculated using the formula: (average colony number × dilution factor × 55 μl worm lysate)/(10 μl worm lysates plated × number of worms). Three independent replicates were performed for the experiment. The phrase “55 μl worm lysate” (50 μl 1% Triton X + 5 μl Lev containing the worms) refers to the total worm lysate of which 10 μl was used for plating.

For the enumeration of R15-GFP within serotonin-treated N2 worms, the method was identical to that described above, except 5 mM of serotonin hydrochloride (Sigma-Aldrich, H9523) was added into the NG agar one day before spreading the R15-GFP culture. Serotonin was prepared freshly for every experiment. At least two independent replicates were performed for the experiment.

Defecation rate assay. The experiment was performed as previously described with modifications. Three live worms were randomly picked and briefly anesthetized in 25 mM Lev. The worms were washed twice in 200 μl antibiotic cocktail comprising 25 mM Lev and 500 μg/ml kanamycin followed by incubation for at least 45 min to completely kill bacterial cells associated with the worm cuticle. Next, the worms were washed three times with 200 μl of 25 mM Lev to eliminate the killed bacteria and residual antibiotic. During the last washing step, the Lev was removed, leaving about 5 μl in the tube. Prior to adding 50 μl 1% Triton X (Sigma-Aldrich, X100), the worms were enumerated and then homogenized with a motorized pestle. Serial dilutions were performed on the worm lysates. Briefly, 10 μl of the worm lysate was spotted on Ashdown agar supplemented with 100 μg/ml Cm using the drop plate method with modifications. Colonies were counted after incubating the plates at 37°C for 48 h. Average colony numbers obtained from visually separate colonies were used for statistical analysis. Bacterial CFU per worm was calculated using the formula: (average colony number × dilution factor × 55 μl worm lysate)/(10 μl worm lysates plated × number of worms). Three independent replicates were performed for the experiment. The phrase “55 μl worm lysate” (50 μl 1% Triton X + 5 μl Lev containing the worms) refers to the total worm lysate of which 10 μl was used for plating.

Feeding rate assay. The experiment was conducted as that described by Song and Avery with minor modifications.

At 1, 4, 8, 12, 24 and 28 h post-infection, 30 live worms were observed in three batches (10 worms/batch, 5 sec/worm) to count the number of pharyngeal pumps. A pharyngeal pump was strictly defined as a
backward contraction of the terminal bulb which is distinctly different from the convulsion-like head movement. The average number of pharyngeal pumps in 10 worms (pumps/sec) was calculated and used for statistical analysis. The same protocol was also used in determining the pharyngeal pumping rate of serotonin-induced worms, and all experiments were performed in at least three replicates.

**qRT-PCR analysis of pgp-5 induction.** The mRNA levels of pgp-5 (GenBank, C05A9.1) were measured using total DNase-treated RNA extracted from worms exposed to Bp R15 for 4 h and 12 h. RNA extraction and qRT-PCR analysis were performed as described by Evans et al. using the forward primer (5'-GGAAATCAGAATGGACGA-3') and reverse primer (5'-TGTTGTAAGTACCGTTAGTT-3'). qRT-PCR was performed using the iScript One-Step RT-PCR kit with SYBR green (Bio-Rad, 170-8892) according to the manufacturer’s instructions. The Bio-Rad iCycler (Bio-Rad) was used for amplification and quantification of the products. Specificity of amplification was confirmed by melt curve analysis after amplification. Normalized threshold cycle (Ct) values were used to calculate fold change of mRNA levels in Bp R15-infected worms as compared with worms fed on E. coli OP50. The Ct values were normalized to changes in three genes [ama-1, F44B9.5 and pan-actin (act-1, 3, 4)] that were found to be unchanged with infection. The experiment was performed in triplicate.

**Scoring of Bp R15-infected pgp-5:gfp transgenic worms.** Bp R15 was used to infect the pgp-5::gfp transgenic worms. Twenty live transgenic worms were picked for observation and scoring of green fluorescence at 100X magnification 4, 12 and 28 h post-infection. The average of the sum of scores was calculated and used for statistical analysis. Three independent replicates were performed.

**RNAi knock-down of pgp-5 and survival assay.** The RNAi knock-down was performed by placing the eggs of rrf-3(pk1426); gfp-4(bn2) double mutant worms on NGM/IPTG/Cb plates preceded with E. coli bearing the pgp-5 RNAi clone, which was acquired from the C. elegans ORF-RNAi library (Open Biosystems, RCE1181). The eggs were allowed to hatch and develop into Glp adult worms on the RNAi plates. For the survival assay, 120 RNAi-treated Glp worms were transferred onto the Bp R15 lawn (40 worms/plate) and incubated at 16°C. Worms were considered dead when they were no longer responsive to probing with a platinum wire picker. Worms that crawled up to the plate wall and died as a result of desiccation were censored from the analysis. The experiment was performed in duplicate.

**Statistical analysis and image processing.** Differences in nematode killing between Bp R15 and R15-GFP as well as the survival between pgp-5 RNAi-treated and untreated worms were assessed by the Log-rank test using StatView version 5.0.1 (SAS Institute, Inc.). For the remainder of assays, the data sets were first assessed using the Kolmogorov-Smirnov normality test. Only data sets that passed the test were analyzed with two-tailed Student’s t-test, the rest of the data sets were analyzed using non-parametric Mann-Whitney U-test. All the analyses were performed using GraphPad Prism version 5.04 (GraphPad Software). P-values of < 0.05 and < 0.01 were considered as statistically significant. All fluorescence photomicrographs were cropped, overlaid or compressed using Adobe Photoshop version 7.0 (Adobe) and assembled into figures using GraphPad Prism.

**Disclosure of Potential Conflicts of Interest**
No potential conflicts of interest were disclosed.

**Acknowledgments**
We thank Dr Suriani Mohamad (Universiti Sains Malaysia) for kindly providing the pSM202. We gratefully acknowledge Dr Man-Wah Tan (Genentech) for his constructive advice on this study as well as Dr Michael Shapiro (University of California at Berkeley) for the transgenic worm strain and image analysis. We also thank Rui-Rui Wong and Nur Affah Ijap for their technical assistance. This study was supported by the Malaysia Genome Institute-Stanford University International Research Grant awarded to S.N. by the Ministry of Science, Technology and Innovation, Malaysia.

**Supplemental Materials**
Supplemental materials may be found here: www.landesbioscience.com/journals/virulence/article/21808

References
1. Galyov EE, Brett PJ, DeShazer D. Molecular insights into Burkholderia pseudomallei and Burkholderia mallei pathogenesis. Annu Rev Microbiol 2010; 64:495-517; PMID:20526691; http://dx.doi.org/10.1146/annurev.micro.112408.134030
2. Cheng AC, Currie BJ. Melioidosis: epidemiology, pathophysiology, and management. Clin Microbiol Rev 2005; 18:383-416; PMID:15831829; http://dx.doi.org/10.1128/CMR.18.2.383-416.2005
3. Sprague LD, Neubauer H. Melioidosis in animals: a review on epizootiology, diagnosis and clinical presentation. J Vet Med B Infect Dis Vet Public Health 2004; 51:305-20; PMID:15525177; http://dx.doi.org/10.1111/j.1439-0465.2004.00797.x
4. Gan YH. Interaction between Burkholderia pseudomallei and the host immune response: sleeping with the enemy? J Infect Dis 2009; 199:1845-50; PMID:19625187; http://dx.doi.org/10.1086/497382
5. Limmathurotsakul D, Peacock SJ. Melioidosis: a clinical overview. Br Med Bull 2011; 99:125-39; PMID:21558159; http://dx.doi.org/10.1093/bmbld/ldr007
6. Schweitzer HP, Peacock SJ. Antimicrobial drug-selection markers for Burkholderia pseudomallei and B mallei. Emerg Infect Dis 2008; 14:1689-92; PMID:18976550; http://dx.doi.org/10.3201/eid1411.080431
7. Inglis TJJ, Sagripanti JL. Environmental factors that affect the survival and persistence of Burkholderia pseudomallei. Appl Environ Microbiol 2006; 72:6865-75; PMID:16980433; http://dx.doi.org/10.1128/AEM.01036-06
8. Leelasuang A. Recent development in melioidosis. Curr Opin Infect Dis 2004; 17:131-6; PMID:15021053; http://dx.doi.org/10.1097/00001432-200404000-00011
9. Currie BJ. Burkholderia pseudomallei and Burkholderia mallei: melioidosis and glanders. In: Mandell GL, Bennett JE, Dolin R, eds. Mandell, Douglas, and Bennett’s Principles and Practices of Infectious Disease. 7th ed. Philadelphia: Elsevier Inc, 2010: 2869-77.
10. Hodgson KA, Morris JL, Feterl ML, Goran BL, Kerbeesan N. Altered macrophage function is associated with severe Burkholderia pseudomallei infection in a murine model of type 2 diabetes. Microbes Infect 2011; 13:1177-84; PMID:21835260; http://dx.doi.org/10.1016/j.micinf.2011.07.008
Burkholderia pseudomallei is a Gram-negative bacterium that causes melioidosis, a severe and often fatal disease. This bacterium is known to have a complex interaction with host defense mechanisms. A study by Wong et al. (2006) showed that the bacterium interacts with the innate immune system of Caenorhabditis elegans, a model organism for studying innate immunity. The study revealed that the bacterium activates the nematode’s innate immune response, leading to the killing of the nematode by the bacterium's virulence factors. This finding highlights the importance of understanding the interactions between pathogens and host defense mechanisms for effective disease control.

In another study by Gan et al. (2007), the bacterium was shown to induce diabetic hosts to lead to more severe disease during infection, indicating the importance of host factors in disease progression. These findings are crucial for developing strategies to combat melioidosis and other infections caused by B. pseudomallei.