Detection of *Burkholderia pseudomallei* toxin-mediated inhibition of protein synthesis using a *Caenorhabditis elegans ugt–29* biosensor

Rui-Rui Wong¹, Cin Kong¹, Song-Hua Lee¹,† & Sheila Nathan¹,²

Toxins are believed to play a crucial role in *Burkholderia pseudomallei* pathogenicity, however to date, only a few have been identified. The discovery of additional toxic molecules is limited by the lack of a sensitive indicator of *B. pseudomallei* toxicity. Previously, from a whole genome transcriptome analysis of *B. pseudomallei*-infected *Caenorhabditis elegans*, we noted significant overexpression of a number of worm genes encoding detoxification enzymes, indicating the host’s attempt to clear bacterial toxic molecules. One of these genes, *ugt–29*, a family member of UDP-glucuronosyltransferases, was the most robustly induced phase II detoxification gene. In this study, we show that strong induction of *ugt–29* is restricted to infections by the most virulent species among the pathogens tested. We also noted that *ugt–29* is activated upon disruption of host protein synthesis. Hence, we propose that UGT–29 could be a promising biosensor to detect *B. pseudomallei* toxins that compromise host protein synthesis. The identification of bactobolin, a polyketide-peptide hybrid molecule, as a toxic molecule of *B. pseudomallei* further verifies the utilization of this surveillance system to search for bacterial toxins. Hence, a *ugt–29* based reporter should be useful in screening for other molecules that inhibit host protein synthesis.

*Burkholderia pseudomallei*, a Gram-negative soil bacterium, is the causative agent of melioidosis, an endemic disease which results in high mortality of infected humans¹,². In Northeast Thailand, the number of deaths from melioidosis is the third highest after HIV-acquired immunodeficiency syndrome and tuberculosis³. Despite extensive studies undertaken to unravel the pathogenesis during the establishment of infection, many of the *B. pseudomallei* virulence factors still remain uncharacterized.

In an earlier study, we had analyzed the genome-wide transcriptome profile of *B. pseudomallei* infected *Caenorhabditis elegans* to elucidate host responses to the infection within the context of a whole animal⁴. Approximately 6% of the worm genome was modulated during the infection and amongst the genes that were robustly induced were members of a detoxification enzyme family, proposing the importance of bacterial toxins in the pathogenesis of *B. pseudomallei*. Previously, O’Quinn et al.⁵ suggested that *B. pseudomallei* produces a paralytic endotoxin which leads to the perturbation of Ca²⁺ homeostasis and neuromuscular intoxication in worms. This corresponds to the disease manifestation in higher order animals where paraparesis is reported as a prominent neurological presentation of melioidosis in mice, goats and sheep⁶,⁷. In addition, Ooi et al.⁸ systematically demonstrated that unlike other well-studied pathogens, *B. pseudomallei* does not colonize the worm intestinal lumen during infection. In light of the rapid death of *B. pseudomallei* infected worms, it is very likely that *B. pseudomallei* adopts toxin-mediated killing as the major virulence mechanism in maintaining an active infection and

¹School of Biosciences and Biotechnology, Faculty of Science and Technology, Universiti Kebangsaan Malaysia, 43600 UKM Bangi Selangor, Malaysia. ²Malaysia Genome Institute, Jalan Bangi, 43000 Kajang, Selangor, Malaysia.

†Present address: Centre d’Immunologie de Marseille-Luminy, Aix Marseille Université UM2, Inserm, U1104, CNRS UMR7280, 13288 Marseille, France. Correspondence and requests for materials should be addressed to S. N. (email: sheila@ukm.edu.my)
Results

ugt–29 is expressed ubiquitously upon B. pseudomallei infection. To monitor ugt–29 gene expression patterns in a whole animal, we constructed a transcriptional ugt–29::GFP reporter. The GFP reporter gene was fused to the ugt–29 promoter, microinjected into a pha–1(e2123)ts mutant and maintained extrachromosomally. The pBX plasmid carrying a wild type copy of pha–1 was co-injected to rescue the transgenic worms from embryonic lethality of the pha–1 mutant at 25 °C. Under 100× magnification, ugt–29::GFP transgenic worms exhibited very dim GFP signals when fed on Escherichia coli strain OP50, the standard laboratory food for C. elegans (Fig. 1a). In contrast, the ugt–29::GFP transgene was robustly induced throughout the entire BpR15-infected worm over the period of infection (Supplementary Fig. S1). Next, the tissue distribution of ugt–29 expression was observed using a higher magnification of 400×. As shown in Fig. 1b, constitutive but dim expression of GFP in the uninfected worms was evident specifically at the pharynx, intestine, vulva muscle and tail. In BpR15–infected worms, fluorescence was localized to the same tissues although at a much higher intensity. In addition, intense green fluorescence was also observed at the head and body wall muscle of BpR15-infected worms (Fig. 1c, lower panel).

Robust ugt–29 expression is specific to infections by virulent Burkholderia species and strains.

To determine whether other bacterial pathogens induce ugt–29, we exposed the ugt–29::GFP transgenic worms to four other Burkholderia species (B. thailandensis, B. cepacia, B. vietnamiensis and B. cenocepacia) and three non-Burkholderia pathogens (Pseudomonas aeruginosa, Staphylococcus aureus and Enterococcus faecalis). Aside from B. pseudomallei, B. cepacia was the only pathogen able to induce robust ugt–29::GFP expression (Fig. 2a). Interestingly, data on the mean-time-to-death (TDmean) of nematodes infected by this cohort of bacterial pathogens demonstrated that B. pseudomallei and B. cepacia were significantly more virulent (p < 0.0001, log-rank test) towards the nematodes (Supplementary Fig. S2a). In worms fed with the other bacteria, delayed killing kinetics correlated with mild or negligible ugt–29 expression as indicated by the similar GFP intensity between infected and uninfected worms. The fluorescence profile was consistent with the quantification of endogenous ugt–29 expression by qRT-PCR (Supplementary Fig. S2b). Nevertheless, the observed ugt–29 overexpression might represent changes due to the deadly consequences following infection and may be less apparent at this early time point in worm infections that present with a longer TDmean. Hence, in worms infected by P. aeruginosa, S. aureus and E. faecalis, we observed for fluorescence at a later stage of infection when ~50% of the worm population was killed. Again, we noted no significant increase in GFP intensity in infected worms at this later time point (Supplementary Fig. S3), suggesting that the induction of ugt–29 is a specialized host response to specific infections rather than a generic response towards reduced nematode fitness following infection.

To further delineate the association between ugt–29 expression and bacterial virulence, we challenged ugt–29::GFP transgenic worms with five different B. pseudomallei of varying virulence capacity. When compared with basal fluorescence in uninfected worms, infections with all five B. pseudomallei isolates triggered an observable increase in fluorescence expression, albeit the increase was relatively mild in worms infected by Ovine 3470 (TDmean of 55.44 ± 2.163 hours) and Goat 2124 (TDmean of 69.23 ± 1.097 hours) which were the two least virulent isolates tested (Fig. 2b, see also Supplementary Fig. S1c). Similarly, quantification of endogenous ugt–29 expression by RT-PCR also reinforced the association between ugt–29 overexpression and B. pseudomallei virulence (Supplementary Fig. S1d). Of note, the most virulent strain, Human 3475, triggered markedly enhanced transcription of ugt–29 (134-fold) in worms following infection. In addition, unlike viable BpR15, heat-killed BpR15 which is not pathogenic to worms was not able to trigger robust ugt–29 expression, again implying that induction of ugt–29 is pathogenicity-related (Fig. 2c).

Next, we asked if other abiotic stressors also induce similar robust expression of ugt–29. ugt–29::GFP worms were exposed to three different stressors (heat shock, paraquat and the heavy metal cadmium) and GFP fluorescence intensities were assessed relative to untreated worms. The transgenic worms carrying the hsp–16.2::GFP or pgp–5::GFP transgene were used as controls and exposed to heat shock and cadmium, respectively. As expected, both control worms exhibited an increase in GFP expression relative to untreated worms (Fig. 3a). In contrast, no...
An observable increase in GFP expression was observed in ugt–29::GFP worms under all three stress conditions tested (Fig. 3b). In addition, we eliminated the possibility that the induction of ugt–29 is a consequence of food limitation as a result of pathogen avoidance by *C. elegans* during a *B. pseudomallei* infection. GFP intensities were similar between starved and *E. coli* OP50-fed worms (Fig. 3b, lowest panel). Taken together, these observations suggest that ugt–29 expression is likely a specialized response rather than a general stress response to *B. pseudomallei* infection.

**The conserved zip–2 pathway is required for ugt-29 induction upon *B. pseudomallei* infection.**

When confronted with infecting pathogens, *C. elegans* activates transcriptional responses that involve cooperative
induction of innate immune effectors and detoxification genes to protect itself\(^4,18,19\). The distinct and robust expression of \(ugt–29\) in \(BpR15\)-infected worms suggests a prominent role of this gene in defense against \(B. pseudomallei\) infection. When we reexamined the cohort of \(C. elegans\) genes modulated during the \(B. pseudomallei\) infection, we noted that a number of the genes including \(ugt–29\) are known to be regulated by ZIP-2 in worms infected by \(P. aeruginosa\)\(^4,20\). In \(BpR15\)-infected worms, this cohort of genes was amongst those with the highest magnitude of induction\(^4\). As such, it is likely that ZIP-2 plays an important role in worm defense during a \(B. pseudomallei\) infection.

**Figure 2. Strong \(ugt–29\) expression is specific to infections by virulent \(Burkholderia\) species and strains.** Representative fluorescence micrographs (400× magnification) of \(ugt–29::GFP\) reporter worms exposed to (a) various pathogens at 24 hours post infection, (b) individual \(B. pseudomallei\) isolates at 8 hours post infection and (c) heat-killed \(BpR15\) relative to viable \(BpR15\) at 24 hours post infection. All micrographs were acquired at same exposure time and gain factor. Scale bar, 50\(\mu\)m.
To confirm if ZIP–2 does indeed contribute to increased survival following a *B. pseudomallei* infection, we abolished the **zip–2** signaling pathway in **rrf–3(pk1426);glp–4(bn2)** double mutant worms and evaluated the **zip–2** RNAi worms susceptibility to *B. pseudomallei*. When the survival rate of the infected worms was compared to the control RNAi worms, we noted a significant reduction in **zip–2** RNAi worm survival (Fig. 4a). This verified that **ZIP–2** is required for resistance against the killing effects of *B. pseudomallei*.

Next, we asked if **ugt–29** is a down-stream target of **ZIP–2** in the context of a *B. pseudomallei* infection. We abrogated **zip–2** in **ugt–29::GFP** reporter worms and assessed fluorescence intensity relative to control RNAi worms during a BpR15 infection. As shown in Fig. 4b, when ZIP–2 is compromised, the intense fluorescence in BpR15-infected worms was markedly reduced to an intensity comparable to basal expression of uninfected worms, proposing that **ZIP–2** is required to mediate the induction of **ugt–29**.

Quantification of **ugt–29** transcript levels by RT-PCR confirmed this reduction, further validating that *B. pseudomallei*-induced expression of **ugt–29** requires **ZIP–2** (Fig. 4c).

**Induction of ugt–29 is in response to inhibition of host protein synthesis.** The **zip–2** signaling pathway has recently been identified as a surveillance system utilized by worms to monitor the integrity of core cellular activities. Nematodes interpret any disruption of these activities as a consequence of pathogen attack and subsequently engage various defense responses which include aversion behaviour, detoxification and innate immune responses. Interestingly, many of the critical cellular processes monitored by this cellular surveillance-activated detoxification and defense systems are known targets of bacterial toxins. In light of its transcriptional regulation by **ZIP–2**, we asked if **ugt–29** is induced in response to cellular perturbations caused by *B. pseudomallei* toxins or virulence factors. To answer this, we performed RNAi-mediated gene inactivation on a panel of nematode genes known to encode core cellular components or metabolic enzymes (Table S1). L1 stage sterile **ugt–29::GFP** worms were fed with individual *E. coli* RNAi clones and the level of fluorescence was monitored 48 hours post-treatment. RNAi against four genes encoding translation factors (C37A2.7 or **eef–2**) and tRNA synthetases (**pars–1**, **aars–2**) resulted in significant over-expression of **ugt–29** relative to control RNAi-treated worms (Fig. 5a). Following RNAi knockdown, disruption of protein synthesis most likely triggered the induction of **ugt–29** even in the absence of infection. As translational inhibition is a common mode of toxic action, it is likely that **ugt–29** overexpression following a *B. pseudomallei* infection is a result of toxin-induced translational inhibition.

To validate this, we examined if microbial molecules known to inhibit host translation could cause overexpression of **ugt–29** in worms. **ugt–29::GFP** transgenic worms were exposed to protein synthesis inhibitors, hygromycin B (157 μg/ml) or tetracycline (100 μg/ml), for 24 hours and fluorescence was monitored. Worms treated with either inhibitor exhibited intense fluorescence in comparison to untreated control worms (Fig. 5b). This is in agreement with the observed inducible expression of **ugt–29** in worms whose protein translation machinery was disrupted by RNAi. We further examined if the induction of **ugt–29** by hygromycin and tetracycline is also **ZIP–2**-dependent. To address this, **zip–2**-deficient **ugt–29::GFP** worms were exposed to the inhibitors and

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**Figure 3.** The expression of **ugt–29** was not visibly induced upon challenge by environmental stressors. GFP expression was observed in (a) TJ375 worms exposed to heat stress (37 °C for 1 hour) and 10 mM paraquat; (b) WE5172 worms exposed to 100 μM cadmium and (c) **ugt–29::GFP** worms exposed to heat stress, paraquat, cadmium and starvation. Paraquat-treated worms were examined at 6 hours post treatment whilst for the other three stress conditions, worms were assessed at 24 hours post treatment. The exposure time and gain factor are similar for all the micrographs taken. Scale bar, 200 μm.
data was highly complex, an orthogonal partial least square-discriminant analysis (OPLS-DA) was employed to
replicates of each species and distinct separation among the species tested (Supplementary Fig. S5). As the LC-MS
multivariate analysis, principal component analysis (PCA) indicated close clustering of metabolite data within
time and mass, m/z) derived from XCMS were first clustered using RamclustR to reduce dataset redundancy. For
of the diffusible products of all three
pial targets were shortlisted based on a comparison between all three profiles. Four or five biological replicates
from metabolite profiling was performed on the diffusible products of all three individual species and poten-
trometry) metabolite profiling was conducted. As noted above, B. pseudomallei as
of a size
challenged with the fractions of B. pseudomallei
lar (Fig. 6a). When the pooled diffusible products were size fractionated by membrane ultrafiltration, only worms
fluorescence demonstrating that the substance(s) responsible for inducing
induces nematode
Bactobolin is the toxic molecule that triggers overexpression of ugt–29. As ugt–29 is most likely
induced by the presence of toxic molecules, we determined if the B. pseudomallei BLF-1 toxin triggers the induc-
tion of ugt–29 during a B. pseudomallei active infection. Following worm infection, we noted that the BLF1
mutant retained the capacity to strongly induce ugt–29 (Supplementary Fig. S4). No noticeable reduction in flu-
orescence was observed between worms infected by the BLF–1 mutant and its isogenic wild type strain, K96243.
This suggests that BLF1 does not contribute to or is not solely accountable for the strong induction of ugt–29
observed in C. elegans infected with B. pseudomallei. This was a strong indication that other unidentified toxin(s)
of B. pseudomallei are most likely involved in the activation of the zip–2/ugt–29 surveillance pathway.
As bacterial toxins are usually secreted, the search for the potential toxic molecule(s) of B. pseudomallei that
induces nematode ugt–29 expression was undertaken on the premise that ugt–29::GFP worms would exhibit
increased fluorescence when exposed to BpR15 secreted products. The bacterial secretome was able to promote
fluorescence in untreated worms, suggesting that the target substance(s) is not a protein (Fig. 6c) and is most likely a metabolite.
To enable the identification of B. pseudomallei metabolite(s) responsible for the induction of ugt–29, com-
parative metabolite profiling was conducted. As noted above, B. cepacia but not B. thailandensis shared a similar ability
as B. pseudomallei to strongly induce ugt–29. Hence, this suggested that the target substance(s) is common to both
B. pseudomallei and B. cepacia but not B. thailandensis. Non-targeted LC-MS (liquid chromatography-mass spec-
trometry) metabolite profiling was performed on the diffusible products of all three individual species and potent-
tial targets were shortlisted based on a comparison between all three profiles. Four or five biological replicates of the
diffusible products of all three Burkholderia species were collected and profiled. The features (retention
and mass, m/z) derived from XCMS were first clustered using RamclustR to reduce dataset redundancy. For
multivariate analysis, principal component analysis (PCA) indicated close clustering of metabolite data within
replicates of each species and distinct separation among the species tested (Supplementary Fig. S5). As the LC-MS
data was highly complex, an orthogonal partial least square-discriminant analysis (OPLS-DA) was employed to

Figure 4. ZIP–2 is required to protect C. elegans from B. pseudomallei killing and to regulate the
overexpression of ugt–29 in B. pseudomallei infected worms. (a) Graph denotes the killing kinetics of worms
infected by BpR15 upon zip–2 RNAi knockdown. zip–2 knockdown worms exhibited increased susceptibility
to B. pseudomallei infection (p < 0.0001 log-rank test). Error bars represent mean value ± SD. Shown is the
representative of two independent experiments (n = 120). (b) Representative fluorescence micrographs (100×
magnification) of BpR15-infected ugt–29::GFP reporter worms upon zip–2 RNAi knockdown at 6 hours post
infection. The exposure time and gain factor are similar for all the micrographs taken. Scale bar, 200 μm.
(c) qRT-PCR analysis of ugt–29 in BpR15-infected worms upon zip–2 RNAi knockdown at 6 hours post
infection. Error bars represent mean values ± SD.
identify the compounds that most strongly contributed to the model of comparison and we obtained a shortlist of 36 compounds (Table 1). These compounds were annotated based on spectral matching to five different metabolite databases: an in-house small molecules library of the Proteomics and Metabolomics Core Facility (PMF) at Colorado State University, NISTv12, Golm, Metlin Mass Spectral database and Massbank metabolite databases. However, only five of the compounds had a match in the metabolite databases and were annotated with identities whilst the remaining compounds were resolved as unknowns or peptides. Among the annotated targets, the compounds with the highest fold change between *B. pseudomallei* or *B. cepacia* and *B. thailandensis* were identified as bactobolins i.e. bactobolin A, B and D. Bactobolin is a family of polyketide-peptide hybrid molecules that act as broad-spectrum antibiotics and is cytotoxic to mouse fibroblast cells. Bactobolin targets the prokaryotic 50S ribosome-associated L2 protein and its homologue in eukaryotes, L8e, is presumed to be the conserved target26. To validate whether bactobolin is the bacterial compound that induces nematode *ugt–29* expression, we exposed *ugt–29::GFP* worms to 25 μg/ml or 50 μg/ml synthetic bactobolin A and fluorescence was monitored. As depicted in Fig. 6d, 25 μg/ml of bactobolin was sufficient to strongly induce the expression of *ugt–29* whilst the higher dose intensified the overexpression of this gene in worms. Hence, the toxic action of bactobolin most likely inhibits translation in *C. elegans* through the L8e protein resulting in overexpression of *ugt–29* in worms. To determine if bactobolin A targets L8e, we RNAi inactivated *rpl–2* (the gene encoding L8e) in *ugt–29::GFP* worms and monitored fluorescence intensity following bactobolin treatment relative to control RNAi worms. We propose that abrogation of the bactobolin cellular target would result in decreased *ugt–29* expression. As expected, control RNAi-treated *ugt–29::GFP* worms exhibited strong fluorescence upon supplementation with 25 μg/ml of bactobolin A (Fig. 6e), For *rpl–2* knockdown worms, a marked reduction in fluorescence intensity was noted in comparison to RNAi control worms. This suggests that the inducible expression of *ugt–29* during exposure to bactobolin is essentially dependent on functional RPL–2 protein and that bactobolin affects host protein biosynthesis, which in turn, leads to the overexpression of *ugt–29*.

A partial in-frame deletion of the bactobolin encoding gene in BpR15 (Δbpss1174) was successfully generated, as verified by the amplified truncated sequence of bpss1174 (Supplementary Fig. S6). The gene deletion rendered the mutant *B. pseudomallei* significantly less effective in killing the worms (Fig. 6f) with parallel suppression of *ugt–29* expression (Fig. 6g). The TD$_{\text{mean}}$ of worms infected by the bactobolin mutant was significantly delayed when compared to that of worms infected by wild type BpR15 ($p < 0.0001$, log-rank test). In addition, a nematode lifespan assay demonstrated a dose-dependent reduction in the lifespan of worms supplemented with bactobolin A, further confirming the toxicity of this compound on worms (Supplementary Fig. S7).
together, these data verify that bactobolin is a toxic bacterial compound of *B. pseudomallei* and is responsible for the induction of *ugt–29*.

**Figure 6.** Bactobolin is the toxic molecule that triggers overexpression of *ugt–29*. Representative fluorescence micrographs (400× magnification) of *ugt–29::GFP* worms exposed to (a) *E. coli* OP50 or *B. pseudomallei* diffusible products, (b) <3 kDa or >3 kDa size fractions of BpR15 diffusible products and (c) untreated (control) or Proteinase K-treated BpR15 diffusible products for 24 hours. The *B. pseudomallei* compound that induces *ugt–29* is predicted to be an extracellular non-proteinaceous entity of <3 kDa. (d) Representative fluorescence micrographs of *ugt–29::GFP* worms (100× magnification) treated with 25 μg/ml or 50 μg/ml of bactobolin for 24 hours. (e) Representative fluorescence micrographs of *ugt–29::GFP* worms (100× magnification) fed with RNAi clones of either the empty expression vector (control) or *rpl–2* for 48 hours prior to treatment with 25 μg/ml of bactobolin. (f) Kinetics of *C. elegans* killing by the bactobolin mutant and its isogenic wild type BpR15. Shown is the representative of two independent experiments (n=120). (g) Representative fluorescence micrographs of *ugt–29::GFP* worms infected by the bactobolin mutant and its isogenic wild type BpR15 at 24 hours post-infection. Fluorescence micrographs of the same magnification were acquired at the same exposure time and gain factor. Scale bars, 200 μm (100×) and 50 μm (400×).
Discussion

Lee et al.4 had previously demonstrated that ugt–29 was the most robustly induced C. elegans phase II detoxification gene following B. pseudomallei infection of worms. In parallel with the reports that suggest B. pseudomallei pathogenicity is mediated by bacterial toxins5,8,9, we propose that the robust induction of selected worm detoxification genes is most likely a host response towards the presence of B. pseudomallei toxins. In this study, we constructed a reporter transgenic strain for ugt–29 and demonstrated that ugt–29 expression was specific to infections by B. pseudomallei and B. cepacia, both of which are similar in degree of virulence in the worm infection model. We also demonstrated that induction of ugt–29 is ZIP–2-dependent and the zip–2/ugt–29 activation suggests a role in surveillance of defective mRNA translation. Taken together, we speculate that the over-induced zip–2/ugt–29 in B. pseudomallei-infected worms is indicative of the presence of a B. pseudomallei toxic molecule that interferes with host protein translation.

C. elegans possess a large arsenal of detoxification enzymes that are involved in microbial defense and xenobiotic biotransformation27. Several reports have disclosed evidence that C. elegans elicits different detoxification responses upon challenge by different xenobiotic onslaughts28–31. This implies a specific and distinct response for each detoxification protein-encoding gene which justifies the presence of a large number of these genes in the worm27. Our results support the specific response of detoxification enzymes where a strong induction of ugt–29

Table 1. Identification and annotation of B. pseudomallei (BP), B. cepacia (BC) and B. thailandensis (BT) bacterial compounds by LC-MS. aThe compounds were identified based on mass spectral matching to available databases. bRelative quantity of each compound was determined by calculating the mean area of the chromatographic peak. cFold change in compound abundance of B. pseudomallei or B. cepacia was calculated relative to B. thailandensis.
was observed following infection by only two out of the eight pathogens tested. As ugt–29 is highly induced by \textit{B. pseudomallei} and not by the closely related but less virulent \textit{B. thailandensis}, ugt–29 is most likely required for defense against microbial pathogenic factors as supported by the correlation between strong induction of ugt–29 with strong virulence capacity in killing nematodes.

The zip–2 signaling pathway is an immune pathway that responds specifically to \textit{P. aeruginosa}, in parallel with but independent of, several well-described signaling pathways e.g. the PMK–1 \textit{p38 MAPK} and FSHR–1 pathways\cite{20}. In this study, we demonstrated that zip–2 is also activated during a \textit{B. pseudomallei} infection and that ZIP–2 is essential for complete resistance against killing by \textit{B. pseudomallei}. In fact, ZIP–2 target genes which include the cohort of detoxification genes (CYP–14A5, \textit{ppg–5}, \textit{ppg–7}, \textit{ugt–31} and \textit{ugt–29}) and infection response genes (\textit{irg–1} and \textit{irg–2}) were amongst the most robustly induced transcriptional suite in response to \textit{B. pseudomallei} infection\cite{41}. Interestingly, while zip–2 is comparably active during nematode infection by \textit{P. aeruginosa} or \textit{B. pseudomallei}, ugt–29 is strongly induced by \textit{B. pseudomallei} but not by \textit{P. aeruginosa}. This could be due to different pathogenic mechanisms adopted by these two pathogens. Generally, \textit{P. aeruginosa} relies on bacterial colonization rather than intoxication during slow killing with toxins only playing a minor role, justifying the minimal activation of ugt–29 during a \textit{P. aeruginosa} infection\cite{42}. On the other hand, \textit{B. pseudomallei} does not utilize colonization as the pathogenic strategy in worms\cite{8}.

The zip–2 signaling pathway has been suggested to act as a surveillance mechanism for mRNA translation as well as other essential cellular processes as a means to discriminate pathogens from innocuous microbes. Worms are able to recognize translational inhibition by \textit{P. aeruginosa} ToxA and go on to engage the zip–2 defense\cite{22,23}. Hence, the zip–2/ugt–29 relationship could represent an effector-triggered immune (ETI) response during a \textit{B. pseudomallei} infection. In this study, zip–2/ugt–29 was shown to be responsive to cellular perturbations in mRNA translation, confirming the surveillance role of zip–2. As a highly virulent \textit{B. pseudomallei} isolate could trigger remarkably strong ugt–29 expression, an induction of ugt–29 in \textit{B. pseudomallei}-infected worms implies a lethal toxin action. Together, these observations have paved the way to utilizing ugt–29 expression as a readout of putative \textit{B. pseudomallei} toxic molecules. The finding that bactobolin strongly triggers ugt–29 expression verifies the initial idea of utilizing ugt–29 to sense for microbial compounds that are able to disrupt the host translation apparatus. Bactobolin inhibits the growth of bacterial and mammalian cells by inhibition of protein synthesis\cite{26,33}. The molecular basis of translation inhibition by bactobolin was recently revealed as a consequence of a conformational rearrangement of P-site tRNA that interferes with translation termination\cite{34}. While the toxicity of bactobolin from other bacteria has been extensively studied, our findings propose this small molecule as a new virulence attribute of \textit{B. pseudomallei}. Bactobolin is only the second \textit{B. pseudomallei} bacterial toxin identified to date and more interestingly, both bactobolin and BLF\textsubscript{1}\textsuperscript{9} exploit the host translation machinery to achieve the lethal effect, although targeting different components of the translational apparatus.

While several reports provide evidence for the production of bactobolin by \textit{B. thailandensis}\textsuperscript{26,35,36}, in this study, bactobolin is not secreted in significant amounts by \textit{B. thailandensis}. One possible explanation is that the biosynthesis and secretion of bactobolin are not constitutive and remain cryptic throughout our experimental manipulation. Indeed, it was reported that \textit{B. thailandensis} bactobolin is produced only at 30 °C but not at 37 °C\textsuperscript{26}. The identification of bactobolin as the toxic small molecule also supports the prevalent role of secondary metabolites in bacterial pathogenesis. \textit{P. aeruginosa} strains have been reported to kill worms through secondary metabolites e.g. cyanide and phenazines\textsuperscript{37–39} whilst \textit{B. pseudomallei} siderophore, biofilm ferric and proteasome inhibitor are essential to promote death in the context of a murine infection\textsuperscript{40,41}.

This is the first report on the utilization of a nematode biosensor to successfully identify a potent bacterial toxin. As ugt–29 expression is triggered when the key translation machinery is disrupted, we suggest that the ugt–29:GFP construct is a useful tool for the identification of toxins, particularly those that target the host translational machinery. Our findings have also confirmed that ugt–29 induction is not solely activated by a single xenocompound or bacterial toxin. In addition, hygromycin and tetracycline share the similar ability as bactobolin to trigger strong ugt–29 induction and are also toxic to worms. Worms treated with hygromycin exhibit a significantly reduced lifespan whilst tetracycline negatively affects nematode growth and reproduction\textsuperscript{23,42}.

In summary, we have shown that UGT–29 is part of the ZIP–2-mediated cellular surveillance pathway in response to defects within different factors associated with the host translation machinery. In the context of a \textit{B. pseudomallei} infection, the robust increase in ugt–29 expression suggests significant perturbations in protein synthesis, which is likely the consequence of potent bacterial toxins or virulence factors. Through the utilization of ugt–29:GFP worms and comparative LC/MS, we identified bactobolin as the toxic bacterial molecule that triggers ugt–29 expression. As ugt–29 expression implicates an overall subversion in translation, ugt–29:GFP may serve as a valid nematode biosensor for toxin identification.

**Methods**

**Bacterial and \textit{C. elegans} strains.** The bacterial and worm strains used in this study are listed in Supplementary Table S2. All experiments involving \textit{B. pseudomallei} were approved by Universiti Kebangsaan Malaysia Animal Ethics Committee (UKMAEC) and performed in a BSL2 + level laboratory. Growth and manipulation of \textit{C. elegans} were performed as previously described\textsuperscript{43}.

**Construction of the transgenic ugt–29::GFP expressing strain.** A 964 bp promoter fragment of the \textit{ugt–29} gene (Wormbase ID: WBGene0021709) was amplified from \textit{C. elegans} genomic DNA with primers UGT\textsubscript{F1} (‘\texttt{5’GAGAAGCATTTTGCCAATGGTCTA3’}) and UGT\textsubscript{R1} (‘\texttt{5’AGTGCACTCGCAGCATTGGAAG CTAAAAGATGTGACATTGGTGAAGTT3’}) in which the reverse primer was incorporated with 24–bp of the GFP coding sequence (underlined). Genomic DNA was prepared from individual animals using methods adapted from Barstead \textit{et al.}\textsuperscript{44}. The 1892 bp \textit{Aequorea coeruleascens} \textit{gfp} ORF was amplified from pPD95.75 (a gift
from Prof. Andrew Fire, Stanford University, USA) with the primers GFPF1 (5′ AGCTTGCATGCCTGAGGTCG 3′) and GFPRI (5′ AAGGGCCCGTACGGCAGCTA 3′). Using the PCR fusion technique previously described, both amplicons were fused at the 24-bp overlapping region generated at the 5′ end of the ugt–29 negative strand and gfp ORF positive strand (italicalized sequence), then amplified with a set of nested primers, NUGTF1, NUGTR1 (5′ ATCTTGGCGAATGTCAGACGAG 3′) and NGFPR1 (5′ GGCCGAGTATGAG AACAGTATGAG 3′), to form the ugt–29::GFP construct. High-fidelity Pfu DNA polymerase was used for all DNA amplifications. The resulting fusion product (50 ng/μl) was then microinjected along with 100 ng/μl PBX (pha–1) (a gift from Prof. Andrew Fire, Stanford University, USA) as a coinjection marker into the gonadal syncytium of young adult stage pha-1(e2123)ts animals using a FemtoJet microinjector (Eppendorf AG). Transformed worms were selected by incubating them at 25 °C prior to screening for GFP expression. The transformed worms were identified by observation of GFP expression using a Leica DMRXA2 upright fluorescence microscope equipped with a Leica I3 long-pass GFP filter (Leica Microsystems).

**GFP reporter experiments and microscopy imaging.** Germline proliferation-deficient worms (Glp) were produced as previously described to avoid difficulty in GFP observation within transgenic worms (ugt–29::GFP, hsp–16.2::GFP and pgp–5::GFP) due to the spatial interference by nematode eggs. For every indicated experimental condition, ~ 100 Glp transgenic worms were transferred to the assay plates (pathogen-seeded, RNAi or chemical–supplemented plates) and kept at 25 °C until the completion of the assay. At every indicated time point, 10 to 15 transgenic worms were mounted on a 2% agarose pad prepared on a glass slide for GFP examination. Five microliter of 5 mM levamisole was used as the paralyzing agent. The worms were observed under 100× and 400× magnification using a Leica upright fluorescence microscope equipped with a Leica I3 long pass GFP filter (Leica Microsystems) and all micrographs were captured using the Leica DCF 310 FX digital camera and LAS version 3.8 software (Leica Microsystems).

**Pathogen infection experiments.** To prepare the bacterial lawn, all species of *Burkholderia* including *B. pseudomallei* were cultured overnight at 37 °C in Brain Heart Infusion (BHI) broth (Promadia) and spread on Nematode Growth Media (NGM) plates. For *P. aeruginosa*, the culture was grown in King’s B broth supplemented with 100 μg/ml rifampicin at 37 °C and spread on SK (slow-killing) plates. For *S. aureus* experiments, an overnight culture grown in Trypticase Soy broth (Promadia) at 37 °C was spread on Trypticase Soy agar (Promadia) whilst for *E. faecalis*, bacteria grown overnight in BHI at 37 °C were spread on BHI agar (Promadia). In all cases, 10 μl of the overnight culture was spread on a 3.5 cm assay plate while for the larger 6 cm assay plate, 30 μl of the overnight culture was spread. The assay plates were incubated at 37 °C for 24 hours, cultured at room temperature for 30 minutes to 24 hours before the worms were transferred for infection. Throughout the infection, assays plates were kept at 25 °C until completion of the assay. For heat-killed BpR15, 3 ml overnight culture was pelleted at 3 220 × g for 20 minutes. The pellet was resuspended in 100 μl BHI and incubated at 95 °C for 10 to 15 minutes before seeding on a 6 cm NGM plate.

**RNAi interference.** RNAi interference was performed by feeding the nematodes (Glp ugt–29::GFP transgenic worms for GFP examination or rrf–3(pk1426);gfp–4(bn2) worms for survival assay) with *E. coli* strain HT115 (DE3) expressing double-stranded RNA homologous to the target gene of interest. The *E. coli* RNAi clone was grown in LB medium supplemented with 100 μg/ml carbenicillin at 37 °C overnight. The concentrated culture (25–fold) was seeded onto NGM agar containing 1 mM Isopropyl β-D-1-thiogalactopyranoside (IPTG) (Promega) and was left to dry at room temperature overnight. With the exception of rpl–2, L1 or L2 larval stage Glp worms were transferred to and raised on NGM plates seeded with the individual *E. coli* RNAi clone for 48 hours at 25 °C. zip–2–treated worms were subsequently exposed to *B. pseudomallei* infection or chemical treatments (157 μg/ml hygromycin or 100 μg/ml tetracycline). For rpl–2, a population of older-stage Glp worms (young adult worms) was exposed to the corresponding RNAi clone to avoid developmental arrest. After 48 hours of RNAi treatment at 25 °C, rpl–2–treated ugt–29::GFP worms were later exposed to treatment by 25 μg/ml or 50 μg/ml bactobolin. In all experiments, *E. coli* expressing an empty RNAi expression vector (L4440) served as the control. For chemical treatments, assay plates were prepared by adding the desired concentration of chemicals (157 μg/ml hygromycin, 100 μg/ml tetracycline, 25 μg/ml or 50 μg/ml bactobolin) to the molten NGM during agar preparation. Concentrated *E. coli* OP50 was added as the food source once the chemical-supplemented agars were solidified.

**C. elegans survival and lifespan assay.** *rrf–3(pk1426);gfp–4(bn2)* worms were synchronized and grown at 25 °C to adult stage. For infection assays, thirty age-matched worms were transferred to an assay plate seeded with pathogen (different isolates of *B. pseudomallei*, different *Burkholderia* species, *P. aeruginosa*, *E. faecalis* or *S. aureus*), three plates per strain for each experiment. For lifespan assessment upon treatment with bactobolin, thirty age-matched worms were transferred to NGM plates supplemented with 25 μg/ml or 50 μg/ml of synthetic bactobolin A and the experiment was performed in triplicate. *E. coli* OP50 was seeded on the NGM plates as food source. Survival of infected or treated worms was monitored over time until all the worms died. Worms were considered dead if they failed to respond to probing by a platinum wire picker. Worms that died of desiccation on walls were censored for further analysis. Statistical analysis of worm survival was performed using the Kaplan–Meier nonparametric analysis in the Statview software (version 5.0.1; SAS institute) whereby the statistical significance (p-values) was determined using the log-rank method.

**Environmental stress assays.** Environmental stress assays were performed on transgenic worms as described. Briefly, for the heat shock assay, sterile adult ugt–29::GFP transgenic worms were incubated at 37 °C for 1 hour and transferred to 25 °C over the assessment of GFP profile after heat stress. To establish paraquat and cadmium toxicity assays, sterile adult ugt–29::GFP transgenic worms were transferred and maintained
on NGM agar containing 10 mM paraquat or 100μM cadmium. All the experiments were carried out at 25°C and for every stress condition, 10 to 15 worms were mounted and the GFP profile of stress-challenged worms was compared to untreated control worms at 6 (paraquat) and 24 hours post treatment (heat stress and cadmium). TJ375 (hsp–16.2::GFP) and WE5172 (pgp–3::GFP) transgenic worms were assayed in parallel as the control worms to ensure that the stress conditions were optimally established.

**Total RNA isolation and quantitative RT-PCR (qRT-PCR).** Age-matched sterile adult worms were transferred to pathogen-seeded assay plates for infection, after which, total RNA was isolated using Trizol reagent (Invitrogen) from the population of worms infected by different pathogens at 24 hours post infection and by different isolates of *B. pseudomallei* at 8 hours post infection. Total RNA extracted was further purified using Qiagen RNEasy columns (Qiagen) and DNase-treated using RNase-free DNase (Qiagen). qRT-PCR was performed on purified RNA using iScript™ One-Step RT-PCR kit (BioRad Laboratories) with SYBR green detection on the CFX96 Touch Real-Time PCR Detection System (BioRad Laboratories). The ugt–29 transcript level was measured using the forward primer (5’-TATATGCCAAAGAATGGAGAAC-3’) and the reverse primer (5’-CGAATCTGATAGTGCGGATGTC-3’). Specificity of amplification was confirmed by melt curve analysis whereas amplification efficiency was determined by slope of standard curve. Ct (threshold cycle) values were accurately normalized by means of geometric averaging to three internal control genes (ama-1, F44B95 and pan actin)39. The alteration of ugt–29 transcript level in infected worms was calculated relative to uninfected worms.

**Characterization of BpR15 diffusible products.** To obtain diffusible bacterial products, 30 μl of a 3 ml overnight culture of BpR15 was spread on 0.22 μm cellulose nitrate membrane filters (Whatman) on top of 6 cm NGM agar plates37. The plates were incubated at 37°C for 24 hours after which the filter paper was removed. *E. coli* OP50 was seeded as the food source and ugt–29::GFP transgenic worms were transferred to the plates for GFP examination at 24 hours post exposure. To further determine the size range of the bacterial product(s) that induces ugt–29, three plates of NGM agar with embedded diffusible bacterial products were soaked with 4 ml of sterile distilled water each for 24 hours and pooled together. The pooled diffusible products were size fractionated using Amicon Ultra-15 Centrifugal Filter units (Merck) with a molecular weight cut-off of 3 kDa. The fractionated samples were concentrated in the Eppendorf Concentrator plus™ (Eppendorf) before spreading and drying on NGM agar. Concentrated *E. coli* OP50 was seeded as the food source and ugt–29::GFP transgenic worms were transferred to the plates for 24 hours before the GFP examination. For proteinase K digestion, 166μg/ml proteinase K (Qiagen) was added to the pooled and concentrated sample fraction of <3kDa. The sample was incubated at 55°C for 1.5 hours in parallel with an untreated sample (without proteinase K) that served as a control. The proteinase K-treated and untreated samples were deposited on agar, followed by the seeding of *E. coli* OP50 as food source. ugt–29::GFP transgenic worms were subsequently exposed for 24 hours before GFP examination.

**Metabolite extraction for LC-MS.** NGM plates with embedded diffusible products were prepared as described above. For each biological replicate bacterial sample (*B. pseudomallei*, *B. cepacia* and *B. thailandensis*), three plates of NGM agar were soaked with 4 ml of methanol-water mixture (MW; 1:1, v/v) for 24 hours and the mixture was pooled together. The pooled samples were concentrated in an Eppendorf Concentrator plus™ (Eppendorf) for a complete dry down in order to avoid metabolite degradation, after which, the samples were sent to The Proteomics and Metabolomics Facility (PMF) at Colorado State University for LC-MS analysis. In brief, compounds were annotated based on spectral matching to the PMF NISTv12, Golm, Metlin Mass Spectral database and Massbank metabolite databases. The peak areas for each feature in a spectrum were condensed via the weighted mean of all features in a spectrum into a single value for each compound. Analysis of variance (ANOVA) was conducted on each compound using the aov function in R, and p-values were adjusted for false positives using the p.adjust in R50. PCA was conducted on mean-centered and Pareto-scaled data using the pcaMethods package in R.

**Construction of the bacterobolin mutant (∆bpss1174).** A partial in-frame deletion mutant of bacterobolin in BpR15 was constructed according to the method described by Lopez et al.31 Briefly, two short fragments corresponding to the upstream and downstream sequences of bpss1174 were amplified from BpR15 genomic DNA (Supplementary Fig. S7a). The primer pair used for the amplification by the upstream sequences was US-F (5’-GACCCCGCTATAACAAATCC-3’) and US-R (5’-GAGTCCTCACGACGCAGGAC-3’) whilst the primer pair for the downstream amplification was DS-F (5’-GTTGTGCTGTGGTAAGGACGCCGTTTCTCAAGGG-3’) and DS-R (5’-AAGATTGCTGGTTGTCACGGG-3’). The two amplified fragments were subsequently fused and cloned into the non-replicative plasmid pEXKm5 before transforming into *E. coli* mobilizer strain RH03. The recombinant plasmid harboring the bpss1174 gene fragment was introduced into BpR15 by conjugation and homologous recombination resulting in the formation of either a wild type or mutant strain with the latter counter selected using yeast extract tryptone (YT) agar containing 15% sucrose. The double-crossover clones were further screened by kanamycin selection and colony PCR. Further PCR and DNA sequencing were performed using the V-F (5’-AGTTACTGATCGAAGGCGGCT-3’) and V-R (5’-AGGTTGCACGAGATGTCGGG-3’) primer pair flanking outward from bpss1174 to confirm the successful generation of the bacterobolin mutant.

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**Author Contributions**

R.-R.W., S.-H.L. and S.N. conceived and designed the experiments. R.-R.W., C.K. and S.-H.L. performed the experiments. R.-R.W., S.-H.L. and S.N. analyzed the data and wrote the paper. All authors read and approved the final manuscript.

**Additional Information**

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