**A Stenotrophomonas maltophilia Strain Evades a Major Caenorhabditis elegans Defense Pathway**

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*Stenotrophomonas maltophilia* is a ubiquitous bacterium and an emerging nosocomial pathogen. This bacterium is resistant to many antibiotics, associated with a number of infections, and a significant health risk, especially for immunocompromised patients. Given that *Caenorhabditis elegans* shares many conserved genetic pathways and pathway components with higher organisms, the study of its interaction with bacterial pathogens has biomedical implications. *S. maltophilia* has been isolated in association with nematodes from grassland soils, and it is likely that *C. elegans* encounters this bacterium in nature. We found that a local *S. maltophilia* isolate, JCMS, is more virulent than the other *S. maltophilia* isolates (R551-3 and K279a) tested. JCMS virulence correlates with intestinal distension and bacterial accumulation and requires the bacteria to be alive. Many of the conserved innate immune pathways that serve to protect *C. elegans* from various pathogenic bacteria also play a role in combating *S. maltophilia* JCMS. However, *S. maltophilia* JCMS is virulent to normally pathogen-resistant DAF-2/16 insulin-like signaling pathway mutants. Furthermore, several insulin-like signaling effector genes were not significantly differentially expressed between *S. maltophilia* JCMS and avirulent bacteria (*Escherichia coli* OP50). Taken together, these findings suggest that *S. maltophilia* JCMS evades the pathogen resistance conferred by the loss of DAF-2/16 pathway components. In summary, we have discovered a novel host-pathogen interaction between *C. elegans* and *S. maltophilia* and established a new animal model with which to study the mode of action of this emerging nosocomial pathogen.

*S. maltophilia*, a Gram-negative bacillus previously classified as both *Pseudomonas* and *Xanthomonas* (1), is an emerging nosocomial pathogen. From 1993 to 2004, *S. maltophilia* was found to be among the 11 organisms most frequently recovered from intensive care unit (ICU) patients in U.S. hospitals (2). A more recent study of patients with *S. maltophilia* bacteremia revealed that most cases were related to hospital admission, with some being associated with health care, such as outpatient intravenous antibiotics, treatment in a long-term-care facility, or chemotherapy (3). That study also found that intubation and ICU stay at the time of *S. maltophilia* bacteremia were associated with mortality (3). This is likely correlated with the propensity of *S. maltophilia* to adhere to plastics and form biofilms (4) and the opportunistic infection of patients with preexisting illnesses. In fact, *S. maltophilia* is the predominant bacterium recovered from the respiratory tract of patients with cystic fibrosis (5) and enhances the pathogenesis of Crohn’s disease and ulcerative colitis (6, 7). Additionally, *S. maltophilia* can cause nosocomial pneumonia (8) and is associated with a number of infections, such as meningitis and endocarditis (reviewed in reference 9). Thus, *S. maltophilia* is a significant and medically important human pathogen. Furthermore, characterization of various *S. maltophilia* strains is imperative, as the genome sequences of environmental (including R551-3) and clinical (including K279a) *S. maltophilia* isolates contain heterogeneity that might help mediate adaptations to different environments (10).

Despite its medical importance, there are few model systems to investigate the mode of action of *S. maltophilia* (10–13). Furthermore, little is known about what is required for *S. maltophilia* virulence and if these requirements are bacterium and/or strain specific. Here, we develop *Caenorhabditis elegans* as a model for the study of *S. maltophilia* strain-specific responses. *C. elegans* has proven to be an excellent model for understanding development, neurobiology, behavior, and, more recently, innate immunity (14). Like other metazoans, *C. elegans* has evolved in the presence of microbes. The interaction between *C. elegans* and its associated microbes is multifaceted, as these nematodes feed on potentially pathogenic bacteria. Consequently, the *C. elegans* innate immune response has evolved both conserved and specific aspects to cope with the microbial world. Whereas the conserved Toll-like receptor (TLR) pathway plays a central role in innate immunity in other animals, the single *C. elegans* TLR gene, tol-1, is not involved in the response to several pathogens (15) and does not appear to control the expression of antimicrobial genes (16). Uniquely, tol-1 plays a role in the protection of pharyngeal tissue upon challenge with *Salmonella enterica* (17), suggesting a specific response that differs from the usual TLR signaling. Conversely, the functions of other innate immune pathways are conserved and studying the nematode immune response can be informative in understanding how higher organisms mount pathogen defenses (18, 19). For example, the highly conserved p38 mitogen-activated protein kinase (MAPK) pathway plays a major role in the response to human bacterial pathogens (20) such as *Pseudomonas aeruginosa* (21) and...
Staphylococcus aureus (22). The unfolded-protein response (UPR) ire-1–xbp-1 module is a downstream target of the p38 MAPK pathway in response to pore-forming toxin (PFT) (23), a virulence factor for a number of bacterial pathogens, including Bacillus thuringiensis (24). Other conserved pathways, such as the DBL-1/transforming growth factor β (TGF-β) pathway, play a role in the nematode response to a number of pathogens, including Drechmeria coniospora (25) and Serratia marcescens (26). The conserved insulin-like receptor DAF-2 negatively regulates the transcription factor DAF-16/forkhead box O (FOXO), which induces the expression of numerous downstream effector genes. As a result, daf-2 mutants are long-lived on most bacteria tested to date, including Escherichia coli, Enterococcus faecalis, as well as several other human pathogens (15, 27). The DAF-2/16 pathway also regulates longevity, aging, and diapause in addition to its role in innate immunity (28–30). However, the regulation of longevity and innate immunity is distinct, involving pathway components that either play a role in both processes or have an exclusive role in longevity (31).

Each of the conserved innate immune pathways specifically regulates downstream effectors such as lysozymes, lectins, neuropeptide-like downstreamers (NDLPs), and antimicrobial factors (reviewed in reference 20). For example, dbi-1 regulates caenacins but not the structurally related NDLPs (25). Innate immune effector genes such as members of the lysozyme family and the C-type lectin family are also pathogen specific (32), and innate immune pathway components can exhibit effector regulation independent of other pathway constituents (33). Finally, the p38 MAPK and DAF-2/16 pathways appear to function in parallel and positively regulate unique downstream genes (34). Thus, in addition to sharing conserved pathways with human innate immunity, C. elegans seemingly shares a conserved genetic architecture employing multiple pathways and corresponding effectors that function in parallel to combat various microbial assaults. Therefore, C. elegans is a valuable model system to study innate immune pathway function and specificity.

Previous work demonstrated that S. maltophilia K279a can kill C. elegans (13), and another strain, S. maltophilia G2, was shown to have nematicidal activity (35). S. maltophilia has also been detected in association with the nematode Pristionchus pacificus (36) as well as in the natural environment of C. elegans (B. Samuel, personal communication). Thus, it is likely that C. elegans encounters S. maltophilia in natural settings, suggesting that their interaction is evolutionarily significant. In this study, we report our discovery of an S. maltophilia strain, JCMS, that also kills C. elegans. Our results indicate that JCMS accumulates within C. elegans, does not appear to employ a toxin to confer virulence, and needs to be alive to be pathogenic. Our data also showed that the UPR, p38 MAPK, and DBL-1/TGF-β pathways are involved in a nonspecific innate immune response to bacteria. Furthermore, normally pathogen-resistant C. elegans mutants, such as daf-2, akt-1, and ins-7 mutants, were susceptible to S. maltophilia JCMS, suggesting that JCMS evades the downstream effects of these DAF-2/16 pathway components. These findings correlate with our observation that daf-2–regulated effector genes were not differentially expressed when nematodes were fed S. maltophilia JCMS compared to E. coli OP50. In summary, these results demonstrate the value of evaluating several conserved genetic pathways as a whole and provide evidence for specificity in the C. elegans innate immune response mounted against S. maltophilia.

**MATERIALS AND METHODS**

**Nematode strains.** C. elegans strains containing the following alleles were obtained from the Caenorhabditis Genetics Center: LG (linkage group) I: daf-16(mud6), tol-1(mz2033), LG II: age-1(lx546), ire-1(x33), rss-1(aq3), sma-6(wk7), LG III: daf-2(e1370), daf-2(e1368), sma-4(e729), sma-3(e941), tir-1(qd4), xbp-1(zc12), LG IV: hta-1 ins-7(ok1573), pmk-1(km25), sma-2(c502), LG V: akt-1(ok525), dbi-1(nk3), LG X: akt-2(ok393), pdk-1(sa680), sek-1(km4). Strain ZD350 aqls219 [T248R5:: GP::unc-54-3′ UTR + ttx-3::GP::unc-45-3′ UTR] af-7(qd137)III was provided by D. Pagano (MIT). N2 was used as the wild-type strain and was thawed yearly from a frozen stock for experimentation.

**Bacterial strains and growth.** S. maltophilia JCMS was isolated by our laboratory from a culture of Mesorhabditis species nematodes found in soil samples from the Konza Prairie Biological Station in Manhattan, KS. Briefly, nematodes were isolated from soil cores, washed in sterile M9 buffer, and allowed to crawl on nematode growth medium (NGM) plates without any bacteria for 1 h. Nematodes were then moved to a plate seeded with E. coli OP50 for rearing. Bacteria that grew on the initial NGM plate were considered to be “nematode-associated bacteria.” Despite our efforts to ensure that JCMS was indeed associated with native soil nematodes, it is possible that this strain could have been present in the soil from which the nematodes were isolated or a laboratory contaminant. E. coli OP50 and OP50-GFP were obtained from the Caenorhabditis Genetics Center. S. maltophilia K279a was obtained from R. Ryan (University College Cork), S. maltophilia R551-3 was obtained from D. van der Lelie (Brookhaven National Laboratory), E. faecalis V583 was obtained from L. Hancock (Kansas State University), and P. aeruginosa PA14 was obtained from F. M. Ausubel (Harvard Medical School). Transformation of S. maltophilia strains was completed via the insertion of a mini-Tn7 expression cassette that expresses green fluorescent protein (GFP) (37), miniTn7KSGFP(pUR25), obtained from T. Ciche (Michigan State University), into the genome of each S. maltophilia strain. All bacterial strains were frozen at −80°C upon retrieval and were thawed regularly for use in experiments. S. maltophilia strains are naturally amplicillin (AMP) resistant and were streaked for colony isolation from a frozen stock onto Luria broth (LB) agar containing 100 µg/ml ampicillin to selectively prevent the growth of other bacterial contaminants. E. coli OP50 was streaked onto LB agar for colony isolation. For each bacterial strain, liquid LB was inoculated and shaken overnight at 32°C. Bacterial lawns used for survival were seeded onto NGM with the bacterial culture at log/lag phase and grown overnight at room temperature.

**Nematode survival assays.** Nematodes were reared and synchronized as larval stage 4 (L4s) at 20°C on E. coli OP50 lawns. For survival analysis, 10 to 15 L4s were picked onto replicate lawns of bacteria and maintained at 25°C. The number of surviving nematodes was recorded daily, and death was determined by the lack of motion in response to prodding with a platinum wire pick. Nematodes were picked to new bacterial lawns for the first 5 to 6 days after the start of the experiment to separate them from their progeny. Dead nematodes were removed upon discovery. Sample sizes (number of nematodes) varied due to contamination and the removal of specimens that died via means other than the specified bacterial treatment, such as desiccation that occurs when nematodes leave the bacterial lawn and die at the plate edge. The infrequent presence of contamination was determined by observing bacterial lawn morphology, and contaminated replicates were discarded. Kaplan-Meier estimates of survival over time and determination of survival curve statistics using Cox proportional-hazard models were performed with R (R Foundation for Statistical Computing, Vienna, Austria). Survival curves can be statistically compared by using the log-rank or Cox proportional-hazard tests. Cox proportional-hazard models were used to test the effects of independent variables such as genotype and bacterium on the hazard, a dependent variable defined as the probability of dying at a given time (38). The model used for analysis is indicated in a footnote for each table, and the effect of the designated independent variable was considered significant if the P value was <0.05. Some models included a categorical variable that speci-
fied the date on which the experiment was completed. This categorical variable and the interaction between this variable and genotype or bacterial treatment were included in the model if they were found to be significant. Models were evaluated by testing for a nonzero slope and visualizing the Schoenfeld residuals (39). A nonzero slope is an indication of the violation of the proportional-hazard assumption, and models were fit to the data aiming to meet this assumption.

**Bacterial load assays.** Bacterial load was determined by measuring CFU using methods modified from those reported in previous studies (40, 41). Synchronized L4s were fed non-GFP-expressing strains (except for *E. coli* due to the presence of ampicillin resistance) for 6 days on NGM plates. For each time point (30 min; 1 h; 6 h; 12 h; and days 1, 2, 4, and 6), three replicates of 7 to 10 nematodes were picked after exposure to *E. coli* OP50-GFP or *S. maltophilia* JCMS, R551-3, and K279a and fed non-GFP-expressing *E. coli* OP50 for 1 h to clear the intestinal lumen of nonadhering bacteria that we reasoned would be swept away during this feeding period. Only living nematodes were used, and the sample size varied due to the number of living individuals available at each time point. Nematodes were then placed onto unseeded plates of NGM containing 120 μg/ml doxycycline (NGM-doxycycline plates) for washing once with 25 mM levamisole-M9 (LM) buffer, twice with LM buffer with doxycycline (120 μg/ml), and twice with M9 buffer. Washed nematodes were then placed into a 1.7-ml microcentrifuge tube containing 50 μl of M9 buffer plus 1% Triton X-100 (Sigma-Aldrich) and homogenized by using a pestle motor. Crushed nematodes were diluted and plated onto LB agar containing 100 μg/ml ampicillin to select for the growth of adherent bacteria. Bacterial load was quantified by plating five replicates of OP50, JCMS, R551-3, and K279a bacterial lawns onto LB agar containing 100 μg/ml ampicillin. For each replicate, 5 ml of M9 buffer was used to gather the bacterial lawn from an NGM plate (see “Bacterial strains and growth,” above).

**Bacterial accumulation, distension, and pharyngeal pumping.** Synchronized L4s were fed GFP-containing bacteria for 11 days and maintained at 25°C. Prior to observation, nematodes were fed non-GFP-containing bacteria for 1 h of clearing. Nematodes were anesthetized (10 mM sodium azide) for observation daily at a ×1,000 magnification using a Zeiss Axiosplan II microscope equipped with epifluorescence and differential interference contrast (DIC) optics. The GFP accumulation pattern (punctate or diffuse) was scored, and the degree of intestinal distension was quantified by using a micrometer. Only living nematodes were scored. The sample sizes varied depending upon the length of time that nematodes survived on the different bacteria. The GFP accumulation pattern throughout the nematode was observed, and distension was recorded for the anterior section of the intestine (most proximal to the pharynx) for a maximum of 30 worms per day for each bacterial treatment. To measure pharyngeal pumping rates, synchronized L4s were picked onto each treatment bacterium. Each treatment included three replicates of 10 to 15 nematodes. During the survival analysis, six nematodes from each treatment were randomly selected and observed daily. The number of pumps of the posterior bulb of the pharynx was counted for 30 s and extrapolated to 60-s intervals. The mean pumping rate is the average of the number of nematodes. During the survival analysis, six nematodes from each treatment strain were removed, and the plate was seeded with *E. coli* OP50 bacteria.

**Germ line removal.** To assess whether the extended survival of age-1 (hx546) mutants compared to that of wild type was dependent on the presence of the germ line, we used RNA interference (RNAi) to knock down *daf-2* (c25.1 as previously reported (44)). Briefly, adult nematodes were placed onto RNAi plates (1 mM isopropyl-β-D-thiogalactopyranoside [IPTG] and 50 μg/ml ampicillin) seeded with *E. coli* HT115(DE3) bacteria expressing either double-strand RNA (dsRNA) or the empty vector (L4440) and allowed to lay eggs. Prior to seeding, each bacterial strain was shaken overnight at 32°C in LB containing ampicillin (50 μg/ml), and dsRNA expression was induced via shaking in 2 ml of fresh LB containing AMP in the presence of IPTG (1 mM) for 3 h. Treated adult nematodes were removed, and their eggs were allowed to develop into adults. Those without a proliferating germ line (Glp) were then picked onto NGM plates seeded with *E. coli* OP50 or *S. maltophilia JCMS* for survival analysis.

**Reverse transcription-quantitative PCR.** Synchronized wild type and *daf-2(e1368) mutant* L4s were grown with *E. coli* OP50 or *S. maltophilia* JCMS at 25°C for 24 h, collected in M9 buffer, and lysed in TRizol reagent (Life Technologies). RNA extraction and DNase treatment were completed by using the PureLink RNA minikit (Invitrogen) and on-column PureLink DNase treatment (Invitrogen). RNA quality was checked by visualizing 28S and 18S rRNA bands using gel electrophoresis and checking the ratios of absorbance at 260 nm/280 nm and at 260 nm/230 nm using a NanoDrop 2000 spectrophotometer. Intact RNA was used for cDNA synthesis using a SuperScript Vilo cDNA synthesis kit (Invitrogen). Reverse transcription-quantitative PCR (RT-qPCR) was completed by using 96-well plates and the CFX96 Touch real-time PCR detection system (Bio-Rad). Each amplification reaction was performed in triplicate, and three biological replicates were completed for each bacterium-nematode combination. Primer sequences for *dcr-85* (5′-CCTGTGACTCTCATTTCGCCG-3′) and 5′-CTGAGGAAGCTGCCCTCAA-3′ and 5′-GCAAATCCAGCCTAACCCACT-3′ and 5′-AACGGCAACAGCATA GTCCA-3′ were designed using NCBI Primer3. Primer sequences for *csq-1* (5′-AACCTGAGGTTCGCAAGGAG-3′ and 5′-TACTGGTCTC GTCTGAGTCTGC-3′) were designed using Geneious software (Biomatters Ltd.). All designed primer sequences were checked for specificity by using NCBI BLAST. Previously reported primer sequences for *dod-2*, K08D8.5, lyc-7, and lyc-1 were used (32). The efficiency of each primer pair was determined by using a standard curve on a pooled sample of cDNA. The efficiencies of the target and the reference gene *csq-1* were determined to be approximately equal (45) and were assumed to be 100% during normalization and ΔCq quantification. The reference gene *csq-1* was chosen due to its low level of variance between the bacterial treatments used. Differential expression was determined by comparing the 2−ΔΔCq values for biological replicates of the target gene in a *daf-2* mutant background versus wild type on JCMS or OP50 and in wild-type nematodes grown on JCMS versus OP50 (46). Statistical significance (*P < 0.05) was determined with Student’s t test assuming equal variance.

**Nucleotide sequence accession number.** The full-length *S. maltophilia* JCMS 16S rRNA gene sequence was deposited in GenBank with the accession number KF724885.

**RESULTS**

*S. maltophilia* JCMS kills *C. elegans*. *S. maltophilia* strain JCMS was isolated in our laboratory (see Materials and Methods). We amplified the complete 16S rRNA gene, and sequence comparisons indicated that JCMS is more closely related to the clinical strain K279a than to the environmental isolate R551-3 (see Fig. S1 in the supplemental material). *S. maltophilia* strains R551-3, K279a, and JCMS are differentially virulent to *C. elegans*, with JCMS being the most severe (Fig. 1; see also Table S1 in the supplemental material), as seen by comparing the hazard ratios determined by the corresponding Cox proportional-hazard model.
Survival of nematodes fed PA14, K279a, and R551-3 was significantly different (see Materials and Methods). Briefly, a hazard is the probability that an individual nematode dies at a given time. Therefore, in this case, the hazard ratio compared the relative hazards of two bacterial strains. Nematodes fed JCMS were ~10 times more likely to die than those fed K279a and 4 times more likely to die than those fed R551-3 (see Table S1 in the supplemental material). Survival of nematodes fed PA14, K279a, and R551-3 was significantly different (P < 0.05) from survival of nematodes fed JCMS. JCMS was more virulent than OP50 and K279a, and PA14 was more virulent than JCMS. Survival of nematodes fed JCMS was not significantly different from that of nematodes fed V583 (P = 0.0663).

**S. maltophilia** JCMS accumulates in the intestine. Given the diverse levels of virulence displayed by the different *S. maltophilia* strains, we sought to determine whether the whole-nematode bacterial load was related to pathogenicity. In order to quantify the bacterial load, we measured CFU from crushed nematodes exposed to *E. coli* OP50-GFP (used instead of OP50 because it is ampicillin resistant) (see Materials and Methods) or *S. maltophilia* JCMS, K279a, or R551-3. Nematodes exposed to JCMS had 10-fold more CFU than any other strain tested at 30 min (Fig. 2A). The bacterial load in nematodes grown on JCMS increased exponentially after 12 h and leveled off after 4 days. Thus, the whole-nematode bacterial load correlates with the degree of pathogenic effect for all the bacterial strains tested. It was possible that the differences in bacterial loads could be caused by differences in the densities of bacteria in the lawns of the different strains; however, the number of CFU in each lawn was not correlated with *S. maltophilia* virulence (Fig. 2B). *C. elegans* feeds by the pharynx pumping bacteria from the mouth into the intestine. It is possible that the observed difference in the bacterial load was related to feeding behavior, as a rapid uptake of bacterial cells could cause an increase in the number of CFU. To address this question, we measured the pharyngeal pumping rates for nematodes fed each bacterial strain daily until death. Since pumping rates are known to decline with age (48), we included only data from the first 4 days. The pumping rate for nematodes fed JCMS was not substantially higher on any of the days observed (Fig. 2C). Thus, bacterial lawn density and nematode bacterial uptake do not correlate with pathogenicity or contribute to the observed strain-specific differences in bacterial loads.

In order to further investigate the differences in the bacterial loads and pathogenicities of the *S. maltophilia* strains, we sought to determine where bacteria were localized within the nematode and whether localization was also correlated with pathogenicity. To better visualize and track bacteria within the *C. elegans* intestine, we introduced GFP into each *S. maltophilia* strain (JCMS, K279a, and R551-3). Integration of GFP did not cause a significant difference in nematode survival on these *S. maltophilia* strains (see Table S3 in the supplemental material). Bacterial pathogenicity was previously correlated with bacterial accumulation within the *C. elegans* intestine and distension of the intestinal lumen (40, 49). Thus, the extent of intestinal lumen distension (Fig. 2D) was measured when the pattern of GFP accumulation was observed (Fig. 3). We focused on the anterior portion of the intestine since it has been shown to be sensitive to the effects of pathogenic bacteria (50, 51). Intestinal distention was observed in nematodes fed each *S. maltophilia* strain but occurred earlier and to a greater extent for those fed the more pathogenic strains JCMS and R551-3 (Fig. 2D).

Nematodes grown on *E. coli* OP50-GFP accumulated GFP-expressing bacteria within the intestinal lumen that had primarily a “punctate” pattern (Fig. 3B and C; see also Fig. S2A in the supplemental material). This pattern of accumulation appears to illuminate intact bacterial cells. Worms fed JCMS-GFP accumulated bacteria primarily in a “diffuse” pattern that appeared to involve higher numbers of tightly packed bacteria, some of which might have been lysed, resulting in the release of GFP into the intestinal lumen (Fig. 3D to F; see also Fig. S2B in the supplemental material). Efforts to characterize the exact cellular mechanism responsible for the diffuse GFP accumulation pattern have been inconclusive. However, the diffuse pattern of bacterial GFP accumulation was characteristic of all *S. maltophilia* strains and
roughly correlated with the degree of virulence (see Fig. S2B in the supplemental material). Few GFP-labeled *S. maltophilia* cells were observed in tissues outside the intestinal lumen and not until nematodes were older and deteriorating, suggesting that *S. maltophilia* pathogenesis initially involved intestinal colonization, which became systemic after prolonged exposure. The extent of this effect, and the degree of intestinal distension, appeared to be generally consistent for all sections of the nematode (data not shown), signifying that the accumulation and distention patterns were representative.

Intriguingly, whole-nematode bacterial loads increased more quickly than the GFP-labeled bacterial loads in the intestine and did not coordinate with intestinal distension. When the accumulation of GFP-expressing bacteria in all parts of the intestine was measured, between 7 and 17% of nematodes had a GFP signal on day 1, depending upon the strain (see Fig. S2C in the supplemental material). These data correlate better with whole-nematode CFU (Fig. 2A), but differences remain. One possible explanation for these differences is that nematodes in the accumulation experiments were cleared of nonadherent bacteria via feeding on the same non-GFP-bearing strain, while in the bacterial load assay, nematodes were cleared of nonadherent bacteria on *E. coli*. To investigate this experimental difference, we compared accumulations of JCMS-GFP within nematodes after clearing on non-GFP-bearing OP50 or JCMS. JCMS-GFP initially accumulated to a greater extent when cleared on *E. coli* OP50 (see Fig. S2D in the supplemental material). Thus, JCMS is better at dislodging itself than OP50. This bacterium-specific difference in the ability to clear nonadherent bacteria may have contributed to the discrepancy between the GFP accumulation and bacterial load experiments. However, it should be noted that the measurements of nematode CFU and GFP accumulation within the nematode quantify two distinct aspects of *S. maltophilia* pathogenicity. A recent report demonstrated that the presence of a GFP signal and/or intestinal distension does not indicate that an infection has been established, suggesting that bacterial load and intestinal accumulation are not necessarily coupled (52). Thus, although GFP accumulation, distension, and bacterial load should intuitively be correlated, these phenomena likely represent different aspects of *S. maltophilia* pathogenicity.

**Virulence of *S. maltophilia* JCMS is not mediated by a toxin and requires live bacteria.** The ability of *S. maltophilia* JCMS to colonize and accumulate within *C. elegans* (Fig. 2A) suggested that living bacteria were involved in *S. maltophilia* virulence. To test the requirement of living bacteria for virulence, we used heat to kill *E. coli* OP50 and each *S. maltophilia* strain. Nematodes fed heat-killed JCMS survived longer than did nematodes fed living bacteria, while survival of nematodes fed heat-killed R551-3 and K279a bacteria was not statistically different from survival of nematodes fed living bacteria of the same strains (Fig. 4A; see also FIG 2
Both antibiotic treatments were optimized to remove all prolifer-maltophilia data and determine if a heat-labile toxin could be involved in antibiotic treatment should denature proteins, we conclude that a protein toxin does not play a significant role in JCMS and likely required for either antibiotic can induce a toxin that is detrimental to the nematode. However, taken together, these data indicate that proliferating bacteria and/or a factor associated with bacterial growth is not the cause of the decreased survival of nematodes fed the respective living bacteria (Fig. 4B and C; see also Table S4 in the supplemental material). Surprisingly, heat-killed OP50 bacteria were more hazardous to nematodes than living bacteria. This effect is not likely due to a lack of nutrition, as these bacteria were concentrated in the same manner as the S. maltophilia strains. While the curves shown in Fig. 4A appear to indicate that heat-killed R551-3 and K279a bacteria were also more hazardous to nematodes, these differences were not statistically significant. These data suggest that a heat-stable compound might contribute to the virulence of the mildly pathogenic S. maltophilia strain R551-3. Additionally, JCMS-mediated killing appeared to be distinct from that of other S. maltophilia isolates in that virulence clearly required these bacteria to be alive. To confirm these data and determine if a heat-labile toxin could be involved in S. maltophilia JCMS virulence, we used the antibiotics doxycycline and ciprofloxacin to treat E. coli and each S. maltophilia strain. Both antibiotic treatments were optimized to remove all proliferating cells. As expected with antibiotic treatment (53), nematodes fed doxycycline-treated OP50 bacteria survived significantly longer than did nematodes fed living bacteria (Fig. 4B; see also Table S4 in the supplemental material). Nematodes fed JCMS and K279a bacteria treated with either antibiotic and R551-3 bacteria treated with ciprofloxacin also survived longer than did nema-todes fed the respective living bacteria (Fig. 4B and C; see also Table S4 in the supplemental material). Surprisingly, survival of nematodes fed OP50 and R551-3 showed a treatment-specific interaction with each antibiotic, as survival of nematodes fed doxycycline-treated R551-3 or ciprofloxacin-treated OP50 bacteria was significantly shorter than survival of nematodes fed the respective living bacteria. This treatment-specific effect suggests that either antibiotic can induce a toxin that is detrimental to the nematode. However, taken together, these data indicate that proliferating bacteria and/or a factor associated with bacterial growth is required for S. maltophilia virulence. Moreover, since neither antibiotic treatment should denature proteins, we conclude that a protein toxin does not play a significant role in JCMS and likely also R551-3 pathogenicity. We also attempted to use UV treatment as another means to kill bacteria without denaturing proteins. However, we were unable to find a UV dose that completely killed all cells (data not shown).

We next used a filter assay to test for the contribution of S. maltophilia secretions to pathogenicity. We found that treatment of OP50 with secretions from any S. maltophilia strain did not cause a decrease in survival compared to treatment with OP50 secretions (Fig. 4D; see also Table S5 in the supplemental material). In fact, we observed a significant extension in survival time for nematodes fed OP50 bacteria treated with K279a secretions. However, it is possible that the filter bound the toxic secretion and/or that the molecule was too large to pass through the filter. While this issue could be addressed through the use of a nonfiltered bacterium-free supernatant obtained by centrifugation of bacterial cultures, we were unable to remove all viable S. maltophilia cells from the supernatant (data not shown). Therefore, we cannot rule out the role of a large and/or filter-bound toxin. However, taken together, these data strongly suggest that living bacteria, rather than a toxin, play a larger role in S. maltophilia nematode pathogenesis.

**DFD-16 insulin-like signaling pathway component mutants are not resistant to S. maltophilia JCMS.** C. elegans daf-2 mutants display increased survival and increased pathogen resistance to practically all bacteria tested to date (14, 27, 31). To determine whether this was the case for S. maltophilia JCMS, we analyzed the survival of various daf-2 pathway mutants on JCMS and E. coli OP50 as a control. Compared to wild type, daf-2(e1368) and daf-2(e1370) mutants were long-lived on OP50, and neither mutant allele conferred extended survival on S. maltophilia JCMS (Fig. 5A and Table 1). Specifically, daf-2(e1370) mutants were more susceptible to S. maltophilia JCMS, while survival of daf-2(e1368) mutants was not significantly different from that of wild type. While we observed increased susceptibility for daf-2(e1370) mutants, these mutants also displayed pleiotropic effects (28) such
as reduced brood size and abnormal development, which might have contributed to the observed phenotype for nematodes fed JCMS. Mutants of other DAF-2/16 pathway components, including the insulin-like ligand \textit{ins-7} and a serine/threonine kinase ortholog, \textit{akt-1}, had extended survival times compared to that of wild type grown on OP50 but not on JCMS, with \textit{ins-7} mutants being more susceptible (Fig. 5Band Table 1). Intriguingly, mutants of another serine/threonine kinase ortholog, \textit{akt-2}, were significantly longer-lived than wild type grown on JCMS, suggesting a role that is similar to that for growth on OP50. The differential roles of \textit{akt-1} and \textit{akt-2} in growth on JCMS may be linked to their differential regulation of antimicrobial gene expression (31).

Contrary to data from previous reports (27,31), we observed that \textit{age-1} mutants were not significantly longer-lived than wild type on OP50. This anomaly led us to ask whether previously reported effects of sterility on life span and pathogen resistance (54, 55) might be responsible. We used RNAi of \textit{cdc-25.1} to remove the germ line (44) in wild-type nematodes and \textit{age-1} mutants and found that \textit{age-1;cdc-25.1}(RNAi) nematodes survived significantly longer than did \textit{cdc-25.1}(RNAi) nematodes grown on OP50 (Fig. 5C and Table 1). There was not a significant difference between wild type and \textit{age-1} mutants fed the RNAi vector only, confirming that the knockdown of \textit{cdc-25.1} was the cause of the increased life span. Furthermore, there was not a significant difference in survival between \textit{age-1;cdc-25.1}(RNAi) and \textit{cdc-25.1}(RNAi) nematodes on JCMS. When the germ line-dependent effects of \textit{age-1} loss of function are taken into account, like most other DAF-2/16 pathway components, \textit{age-1} mutants were not resistant to JCMS. Finally, survival of forkhead box O (FOXO) homolog \textit{daf-16} mutants was not significantly different from survival of wild type fed JCMS or OP50, which is consistent with previously reported results (27,34, 56). In summary, all tested components of the DAF-2/16 pathway except \textit{daf-16} were similarly involved in the response to OP50. We observed that most of the pathway mutants were not significantly different from wild type fed \textit{S. maltophilia} JCMS, suggesting that DAF-2/16 signaling plays little or no role in the \textit{C. elegans} defense response against these bacteria. Furthermore, the usually pathogen-resistant \textit{daf-2(e1370)} and \textit{ins-7} mutants were slightly more susceptible to JCMS than were wild-type nematodes, indicating that their role in the response to JCMS is different than that for other bacterial pathogens.

The degree to which mutations of DAF-2/16 pathway components affect survival was inferred from the value of the mutant-to-wild-type hazard ratio (from the corresponding Cox proportional-hazard model) (see Materials and Methods). Hazard ratio values near 1 reveal that the mutant hazard is not different from that of wild type and suggest that the mutated gene is not involved in the response to the given bacterium, while values that deviate from 1 suggest involvement. Values of >1 indicate that the mu-
Survival of representative DAF-2/16 insulin-like signaling pathway mutants grown on OP50 (Fig. 6 and Table 1). These results suggested an involvement of these C. elegans defense pathways and were bolstered by our analysis of mutants affecting multiple genes within each pathway (Fig. 6 and Table 1; see also Fig. S3 and S4 in the supplemental material). Mutants of the p38 MAPK pathway signaling components sek-1, nsy-1, tir-1, and ayt-7 were each susceptible to OP50 and JCMS, but pmk-1 mutants were susceptible to JCMS only (Fig. 6A and Table 1; see also Fig. S3 in the supplemental material). These results agreed with previously reported data in that pmk-1 mutants were susceptible to pathogenic P. aeruginosa and that the absence of pmk-1 in nematodes fed E. coli is irrelevant (34). The hazard ratios for p38 MAPK mutants were higher than those for other pathway mutants for growth on both JCMS and OP50 (Table 1). In addition, the hazard ratio for a given mutant fed JCMS was higher than that for OP50 for these genes. For example, the hazard ratios for sek-1 and nsy-1 for growth on JCMS were 228 and 31.3, while for growth on OP50, they were 7.18 and 13.9 (Table 1). Thus, the loss of the p38 MAPK pathway is the most detrimental for survival on both bacteria, with the loss being more severe for growth on JCMS.

Mutations in the DBL-1/TGF-β pathway components that form the SMAD complex (sma-2, sma-3, and sma-4) also increased susceptibility to both OP50 and JCMS (Table 1; see also Fig. S4 in the supplemental material). Thus, the SMAD complex components have similar roles in growth on JCMS and OP50. However, mutants of the upstream components dbl-1 and sma-6 were specifically susceptible to JCMS and long-lived when grown on OP50 (Fig. 6B and Table 1). The hazard ratios of these mutants were closer to 1 for growth on OP50 than on JCMS. Thus, it appears that these components are necessary for the response to JCMS and, perhaps, less important for the response to OP50. In fact, with the exception of sma-2, the hazard ratios for DBL-1/TGF-β pathway mutants deviated more from 1 for growth on JCMS. Comparatively, the deviations from 1 for this pathway are not as great as those for the p38 MAPK pathway. Thus, the loss of the DBL-1/TGFβ pathway as a whole is only mildly detrimental to nematodes fed JCMS.

UPR pathway components were also found to play similar roles in the response to JCMS and OP50. Both ire-1 and xbp-1 mutants had significantly shorter life spans than those of wild-type nematodes during growth on both JCMS and OP50 (Fig. 6C and Table 1). Similar to the p38 MAPK and DBL-1/TGFβ pathways, the hazard ratios for UPR pathway mutants were higher for growth on JCMS. In addition, the hazard ratio of ire-1 mutants was higher than that of xbp-1 mutants grown on JCMS and OP50, suggesting a greater role for this serine/threonine protein kinase in both bacteria. Finally, survival of the Toll-like receptor (TLR) tol-1 mutant was not significantly different from that of wild type grown on OP50 but was slightly increased with growth on JCMS.
significantly differentially expressed between p38 MAPK 2/16-regulated genes chosen, only each effector gene on DAF-2/16 signaling by comparing expres-

sion. Accordingly, we sought to determine the dependence of clec-85 focus on nematodes grown on S. maltophilia, sites of pathogen contact (32). Our analysis of DAF-2/16 expression of innate immune effector genes in C. elegans and wild type 4.69 0.088 150 NA a NA 8.53 0.187 244 NA NA

dafl-2/16 pathogenic bacterium like OP50. However, since clec-85, dod-22, and lys-7 are regulated by DAF-2/16 signaling and DAF-2/16 sig-

differential regulation of immune effector genes in nema-

todes grown on S. maltophilia JCMS. We used RT-qPCR to in-
vestigate the expression of several innate immune effector genes in nematodes fed S. maltophilia JCMS and E. coli OP50. We chose to focus on clec-85, lys-1, lys-7, dod-22, K08D8.5, and spp-1 due to their demonstrated regulation by the p38 MAPK, DBL-1/TGF-β, and/or DAF-2/16 signaling pathway (32, 57). These genes were also of interest because they are expressed in the intestine and pharynx, sites of pathogen contact (32). Our analysis of DAF-2/16 pathway mutant survival suggested that these genes do not play a role in the C. elegans innate immune response to S. maltophilia JCMS. Accordingly, we sought to determine the dependence of each effector gene on DAF-2/16 signaling by comparing expression levels in wild type and daf-2 mutants. Of the putative DAF-

2/16-regulated genes chosen, only clec-85, lys-7, and dod-22 were significantly differentially expressed between daf-2 and wild type for growth on JCMS (Fig. 7A). Consistent with data from previous work (32, 57), clec-85 and lys-7 were upregulated and dod-22 was downregulated in daf-2 mutants grown on OP50. However, the expression of spp-1 and K08D8.5 in the daf-2 background was not significantly different from that of wild type fed either bacterium, and these genes were not evaluated further since their expression trends did not agree with data from previous studies (32, 57). Additionally, as expected for a daf-2-independent gene, the expression of lys-1 in the daf-2 background was not significantly different from that for wild type with either bacterium. The expressions of lys-1, lys-7, dod-22, and clec-85 are known to be up-

regulated when C. elegans comes into contact with pathogenic bacteria (26, 32). Thus, the expression of these immune effector genes is typically expected to be upregulated during growth on pathogenic bacteria such as JCMS versus a less pathogenic or non-

pathogenic bacterium like OP50. However, since clec-85, dod-22, and lys-7 are regulated by DAF-2/16 signaling and DAF-2/16 sig-

naling was not primarily involved in the JCMS response, we hy-

pothesized that these effectors would not be differentially ex-

pressed. As expected, only lys-1 was significantly upregulated during growth on JCMS versus OP50 (Fig. 7B). These results support a role for lys-1 and the upstream pathways that regulate lys-1 in the C. elegans defense response to JCMS, while DAF-2/16-de-

TABLE 1 C. elegans defense pathway mutant responses

1. The resistance of tol-1 mutants to JCMS was only marginally significant, with the hazard ratio indicating little involve-

ment.

**DIFFERENTIAL REGULATION OF IMMUNE EFFECTOR GENES IN NEMATODES GROWN ON S. MALTOPHILIA JCMS.** We used RT-qPCR to in-

vestigate the expression of several innate immune effector genes in nematodes fed S. maltophilia JCMS and E. coli OP50. We chose to focus on clec-85, lys-1, lys-7, dod-22, K08D8.5, and spp-1 due to their demonstrated regulation by the p38 MAPK, DBL-1/TGF-β, and/or DAF-2/16 signaling pathway (32, 57). These genes were also of interest because they are expressed in the intestine and pharynx, sites of pathogen contact (32). Our analysis of DAF-2/16 pathway mutant survival suggested that these genes do not play a role in the C. elegans innate immune response to S. maltophilia JCMS. Accordingly, we sought to determine the dependence of each effector gene on DAF-2/16 signaling by comparing expression levels in wild type and daf-2 mutants. Of the putative DAF-2/16-regulated genes chosen, only clec-85, lys-7, and dod-22 were significantly differentially expressed between daf-2 and wild type for growth on JCMS (Fig. 7A). Consistent with data from previous work (32, 57), clec-85 and lys-7 were upregulated and dod-22 was downregulated in daf-2 mutants grown on OP50. However, the expression of spp-1 and K08D8.5 in the daf-2 background was not significantly different from that of wild type fed either bacterium, and these genes were not evaluated further since their expression trends did not agree with data from previous studies (32, 57). Additionally, as expected for a daf-2-independent gene, the expression of lys-1 in the daf-2 background was not significantly different from that for wild type with either bacterium. The expressions of lys-1, lys-7, dod-22, and clec-85 are known to be up-regulated when C. elegans comes into contact with pathogenic bacteria (26, 32). Thus, the expression of these immune effector genes is typically expected to be upregulated during growth on pathogenic bacteria such as JCMS versus a less pathogenic or non-pathogenic bacterium like OP50. However, since clec-85, dod-22, and lys-7 are regulated by DAF-2/16 signaling and DAF-2/16 sig-

naling was not primarily involved in the JCMS response, we hy-

pothesized that these effectors would not be differentially ex-

pressed. As expected, only lys-1 was significantly upregulated during growth on JCMS versus OP50 (Fig. 7B). These results support a role for lys-1 and the upstream pathways that regulate lys-1 in the C. elegans defense response to JCMS, while DAF-2/16-de-

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Dependent effector genes have little or no role in the response to these bacteria.

**DISCUSSION**

*S. maltophilia* JCMS is a *C. elegans* bacterial pathogen. We isolated a strain of the emerging nosocomial pathogen *S. maltophilia* that kills *C. elegans*. Like *E. faecalis*, *P. aeruginosa*, and *S. marcescens* (20, 40), *S. maltophilia* JCMS accumulates in the gut and causes intestinal distention. In contrast to *P. aeruginosa* (58) and/or *B. thuringiensis* (24), JCMS virulence requires the presence of proliferating bacteria and does not involve a toxin. Furthermore, JCMS causes a diffuse GFP accumulation pattern that may represent a bacterial lysate and/or the presence of an intestinal biofilm. Biofilm formation is thought to be a survival mechanism of clinically relevant organisms such as *E. faecalis* (reviewed in reference 59). *E. faecalis* also establishes a proliferating infection within the *C. elegans* intestine (40), and survival on *S. maltophilia* JCMS was indistinguishable from survival on *E. faecalis* V583 (Fig. 1 and Table 1). Also, like *S. maltophilia* JCMS, *E. faecalis* does not kill nematodes when treated with antibiotics (40). These data suggest that the modes of action utilized by *S. maltophilia* JCMS and *E. faecalis* could be similar, possibly involving opportunistic proliferation and biofilm formation.

Comparison of the effects of *S. maltophilia* JCMS, R551-3, and K279a on *C. elegans* revealed several characteristics that likely underlie their differences in virulence. For example, differences in the onset of bacterial accumulation suggest distinct modes of action. Additionally, unlike JCMS, R551-3 virulence did not always require that these bacteria be alive, as heat killing does not change nematode survival, while antibiotic treatment produces differential effects (Fig. 4; see also Table S5 in

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**FIG 6** p38 MAPK, DBL-1/TGF-β, and UPR defense pathway mutants. (A) Survival of wild-type nematodes and *pmk-1*(*km25*), *sek-1*(*km4*), and *nsy-1*(*ag3*) mutants grown on *E. coli* OP50 and *S. maltophilia* JCMS. (B) Survival of wild-type nematodes and *dbl-1*(*nk3*) and *sma-6*(*wk7*) mutants grown on *E. coli* OP50 and *S. maltophilia* JCMS. (C) Survival of wild-type nematodes and *xbp-1*(*zc12*) and *ire-1*(*v33*) mutants grown on *E. coli* OP50 and *S. maltophilia* JCMS. Results plotted are the proportions of surviving nematodes determined by using Kaplan-Meier estimates generated in R for at least three replicate samples of 10 to 15 nematodes. Sample sizes and P values from the application of Cox proportional-hazard models are included in Table 1. All pathway mutants had significantly decreased survival during growth on JCMS. Survival for *pmk-1* mutants was not significantly different from that of wild type, and the *dbl-1* and *sma-6* mutants had significantly extended survival during growth on OP50. All other immune pathway mutants were significantly susceptible to OP50. Values of <0.05 were considered significant.

**FIG 7** *daf-2*-regulated genes are not regulated in nematodes grown on *S. maltophilia* JCMS. RNA was extracted from synchronized wild-type nematodes and *daf-2*(*e1368*) mutants that were grown on *E. coli* OP50 or *S. maltophilia* JCMS for 24 h. Differential expression was determined by comparing biological replicates of the target gene in *daf-2*(*e1368*) mutants versus wild-type (*WT*) nematodes (control) grown on OP50 or JCMS (A) and wild-type nematodes grown on JCMS versus OP50 (control) (B). Fold changes are shown in reference to expression levels in the control sample (wild type grown on JCMS and/or OP50). Statistical significance (*P* < 0.05) was determined with Student’s t test, assuming equal variance. The effector genes *dod-2*, *lys-7*, and *clec-85* are significantly regulated by *daf-2* on JCMS. Only *clec-85* was significantly regulated by *daf-2* when nematodes were grown on OP50. *lys-1* was marginally significantly (*P* = 0.058) upregulated during growth on JCMS versus OP50 and was not significantly regulated by *daf-2* during growth on either bacterium.
the supplement material). Curiously, our data suggest that unlike the other S. maltophilia strains that we tested, R551-3 produces a heat-resistant, ciprofloxacin-sensitive toxin that is induced by doxycycline. If true, the mode of action of R551-3 might be similar to that of P. aeruginosa, which, when heat treated, is as virulent as living bacteria on fast-killing medium (47) and employs heat-stable diffusible toxins called phenazines (58). However, unlike for R551-3, antibiotic treatment of P. aeruginosa renders these bacteria less pathogenic to nematodes (47). Thus, it appears that P. aeruginosa and S. maltophilia R551-3 do not act similarly in all respects.

The DAF-2/16 pathway plays a relatively small role in defense against S. maltophilia JCMS. Binding of the insulin/insulin growth factor (IGF) receptor ortholog DAF-2 negatively regulates the transcription factor DAF-16/FOXO. When the functions of DAF-2 or other members of the DAF-2/16 pathway are disrupted, DAF-16 is free to enter the nucleus to promote the expression of DAF-2 or other members of the DAF-2/16 pathway are disrupted, growth factor (IGF) receptor ortholog DAF-2 negatively regulates similarly in all respects.

Thus, our data suggest that the activation of these genes is insufficient protection against JCMS and that JCMS causes cellular stress that is distinct from that caused by other bacteria that require the action of the DAF-2/16 pathway. Our observation that daf-2-regulated effector genes were not differentially expressed between JCMS and avirulent E. coli strain OP50 (Fig. 7) also supports this notion. Taken together, these results support a response to JCMS that is independent of the DAF-2/16 signaling pathway.

However, survival of ins-7 and akt-2 mutants on S. maltophilia JCMS revealed the possibility of a small, but perhaps significant, role for these genes. Specifically, our data suggest that ins-7 functions to protect nematodes from JCMS. However, the role of ins-7 appears to be relatively minor, as the hazard ratio was the lowest among those for the significantly susceptible innate immune pathway mutants fed JCMS. On the other hand, the removal of akt-2 in nematodes fed JCMS appears to be beneficial. This result may be attributed to the dual role of akt-2 in pathogen resistance and longevity (31). As age-1 also has a dual role (31), it is possible that this resistance is dependent on the germ line, as we demonstrated for age-1 mutants. In fact, akt-2 and age-1 mutants with an intact germ line had similar survival phenotypes and hazard ratios (Table 1). However, like ins-7 mutants, the mutant-to-wild-type hazard ratio is fairly low compared to the hazard ratios of other mutants (Table 1). Therefore, if C. elegans akt-2 has a role in growth on S. maltophilia JCMS, this role is nearly negligible. Still, the resistance of akt-1 mutants to JCMS is curious and suggests that akt-1 and akt-2 might have different functions in the response to these bacteria. This is not unprecedented, as akt-1 and akt-2 have been shown to differentially regulate innate immune effectors such as thr-2 on E. coli and spp-1 on E. coli and P. aeruginosa (31). Thus, akt-1 and akt-2 might play different roles in the response to JCMS by regulating different effector genes.

Conserved defense pathways are involved in the C. elegans response to S. maltophilia JCMS. We performed a survey of the known C. elegans bacterial defense pathways and found that several conserved pathways were involved in the response to S. maltophilia JCMS and E. coli OP50, while the Toll-like receptor gene tol-1 was not. Specifically, we found that tol-1 was minimally involved in the response to JCMS and, as previously shown for other nematode-bacterium interactions (15), was not involved in the response to OP50. On the other hand, the role of the UPR, p38 MAPK, and TGF-β-like pathways is largely conserved, as these pathways are involved in the response to other pathogenic bacteria (21–23, 25, 26). JCMS and OP50. The differential expression of the p38 MAPK- and TGF-β-regulated gene lys-1 (32) in nematodes fed JCMS and OP50 further highlights the necessity of these pathways in C. elegans defense. Furthermore, lys-1 was up-regulated in nematodes fed JCMS (Fig. 7B), as observed previously for other bacterial pathogens (26, 32), suggesting that it functions as an antimicrobial gene. We observed that the loss of the more universally acting p38 MAPK pathway (60; reviewed in reference 14) was the most detrimental, followed by the UPR and TGF-β pathways. Our data also revealed that the loss of genes encoding signaling proteins, such as nsy-1, was very detrimental. This suggests that signaling through these gene products plays a central role in multiple pathways and/or a number of biological processes.

However, a closer inspection of the hazard ratios of S. maltophilia JCMS and E. coli OP50 suggests that individual components within the p38 MAPK and TGF-β pathways might function in a bacterium-specific manner. For example, PMK-1 is the terminal kinase in the p38 MAPK pathway and is important for the switch of the leucine zipper transcription factor ATF-7 from a repressor to an activator in response to P. aeruginosa (61). Our data suggest that PMK-1 mediates this switch in response to JCMS, and thus, nematode defense against these bacteria requires both atf-7- and pmk-1-dependent genes. On the other hand, we have evidence that the response to OP50 is atf-7 dependent and pmk-1 independent. For the TGF-β pathway, it appears that sma-6 and dbi-1 activities during growth on OP50 are somehow disadvantageous for wild-type nematodes but are required for the response to JCMS. Furthermore, the role for TGF-β-like pathway components during growth on JCMS seems similar to their role in S. marcescens infection (26).

Therefore, in wild-type nematodes, all the p38 MAPK and TGF-β pathway components likely play a defense role in the response to JCMS, while only some components are required for the response to OP50, suggesting a bacterium- or pathogen-specific action of individual pathway components. Intriguingly, we did not observe a bacterium-specific action for the tested UPR pathway components, which is likely due to a more generalized role in nematode stress responses (62). Together, these data support an interaction between C. elegans and bacteria involving conserved mechanisms that include genes that are specific to the bacterial environment. While the significance of bacterium-specific action for components within individual pathways is not clear, it can be revealed only by evaluating the functions of multiple pathway components and suggests that further investigation of the roles of individual components in the response to different bacteria is warranted.
S. maltophilia Evades Major C. elegans Defenses

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