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NilD CRISPR RNA contributes to *Xenorhabdus nematophila* colonization of symbiotic host nematodes

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Summary

The bacterium *Xenorhabdus nematophila* is a mutualist of entomopathogenic *Steinernema carpocapsae* nematodes and facilitates infection of insect hosts. *X. nematophila* colonizes the intestine of *S. carpocapsae* which carries it between insects. In the *X. nematophila* colonization-defective mutant *nilD6::Tn5*, the transposon is inserted in a region lacking obvious coding potential. We demonstrate that the transposon disrupts expression of a single CRISPR RNA, NilD RNA. A variant NilD RNA also is expressed by *X. nematophila* strains from *S. anatoliense* and *S. websteri* nematodes. Only *nilD* from the *S. carpocapsae* strain of *X. nematophila* rescued the colonization defect of the *nilD6::Tn5* mutant, and this mutant was defective in colonizing all three nematode host species. NilD expression depends on the presence of the associated Cas6e but not Cas3, components of the Type I-E CRISPR-associated machinery. While *cas6e* deletion in the complemented strain abolished nematode colonization, its disruption in the wild-type parent did not. Likewise, *nilD* deletion in the parental strain did not impact colonization of the nematode, revealing that the requirement for NilD is evident only in certain genetic backgrounds. Our data demonstrate that NilD RNA is conditionally necessary for mutualistic host colonization and suggest that it functions to regulate endogenous gene expression.

Keywords

small non-coding RNA; mutualism; symbiosis; entomopathogen

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Introduction

Entomopathogenic *Steinernema spp.* nematodes are mutualistically associated with bacteria of the genus *Xenorhabdus* (Stock and Goodrich-Blair, 2008). Together, these symbiont pairs infect, kill, and reproduce within insect hosts. A specialized infective juvenile (IJ) stage of the *Steinernema* nematode transmits bacterial symbionts between insects, ensuring maintenance of the symbiosis through generations. The association between *S. carpocapsae* and its symbiont *X. nematophila* has been well studied with regard to cellular and molecular aspects of symbiosis, particularly with respect to the mechanisms by which the IJ is colonized (Goodrich-Blair, 2007; Herbert and Goodrich-Blair, 2007; Chaston *et al.*, 2013). The bacteria occupy a region known as the receptacle in the anterior portion of the IJ intestine (Poinar, 1966; Wouts, 1980; Bird and Akhurst, 1983; Martens *et al.*, 2003; Flores-Lara *et al.*, 2007; Snyder *et al.*, 2007). Although the processes by which *X. nematophila* bacteria are selected and gain entry to the receptacle remain obscure, only one or two individual clones are founders for the final population (~30–200 CFU/IJ) that ultimately fills the space (Martens *et al.*, 2003; Chaston *et al.*, 2013).

To better understand the bacterial molecular factors necessary during colonization of the IJ nematode, a signature tagged mutagenesis screen to identify *X. nematophila* mutants defective in this process was conducted in the *S. carpocapsae* nematode-associated strain *X. nematophila* HGB081 AN6/1 (hereafter referred to as XnSc 081) (Heungens *et al.*, 2002). In one of the mutants identified in this screen, *nilD6::Tn5*, the transposon had inserted into a region of the genome lacking obvious coding potential. Complementation studies then confirmed the *nilD* region was necessary for nematode colonization but was dispensable for virulence in an insect model of infection (Heungens *et al.*, 2002). Bioinformatic analyses have since indicated that the *nilD* locus encodes a single, free-standing CRISPR (clustered regularly interspaced short palindromic repeats) element comprising one spacer and two repeats, which was disrupted by the transposon insertion (Fig. 1).

CRISPRs are genetic elements broadly distributed among bacteria and archaea and can provide resistance to foreign nucleotide sequences (Barrangou and Marraffini, 2014). CRISPRs comprise a series of short repeat sequences that are separated by variable regions, called spacers. Many CRISPR spacers exhibit identity to sequences, termed proto-spacers, within bacteriophage genomes or other mobile DNA elements (Bolotin *et al.*, 2005; Mojica *et al.*, 2005; Pourcel *et al.*, 2005; Deveau *et al.*, 2008). This observation led to the discovery that CRISPR elements encode a rapidly evolving acquired immune defense system against incoming phages and plasmids (reviewed in (Sorek *et al.*, 2013; Barrangou and Marraffini, 2014)). CRISPR arrays are transcribed as single RNA molecules that are then processed by components of the Cas (CRISPR associated sequences) machinery into individual CRISPR RNA (crRNA) molecules (Barrangou *et al.*, 2007; Brouns *et al.*, 2008; Carte *et al.*, 2008). There are three major classes of CRISPR systems (Types I–III). These three types are further subdivided into several subtypes, differentiated by criteria including the phylogeny of the *cas* genes and the sequence of the CRISPR repeats (Makarova *et al.*, 2011a; Makarova *et al.*, 2011b). In *E. coli*, a type I-E system, five proteins, Cse1 (Cas subtype E. coli), Cse2, Cas7, Cas5, and Cas6e (previously named CasA, B, C, D, and E, respectively) are associated in a complex termed “Cascade” and Cas6e, a putative RNA binding protein,

is the subunit responsible for RNA processing (Brouns *et al.*, 2008; Carte *et al.*, 2008; Jore *et al.*, 2011). Processed crRNAs target and interact with proto-spacer encoding DNA or RNA molecules, resulting in gene silencing and/or degradation (Gasiunas *et al.*, 2014). A 6–8 nt seed sequence within the crRNA exhibits 100% identity to the target and is predicted to guide the interaction between the crRNA and the proto-spacer (Semenova *et al.*, 2011; Wiedenheft *et al.*, 2011). CRISPR targeting and silencing also require the presence of a short, proto-spacer adjacent motif (PAM) within the exogenous target sequence, located directly upstream of the seed sequence. The PAM provides a mechanism by which the system differentiates between target and non-target (e.g. endogenous) sequences, thereby preventing potentially lethal auto-immunity due to targeting of chromosomal “self” sequences (Mojica *et al.*, 2009; Westra *et al.*, 2013). The hybridization of the crRNA molecule to the PAM-encoding DNA sequence results in the formation of an R-Loop within the crRNA that acts as a binding site for another member of the Cas protein family, Cas3. Cas3 contains helicase and nuclease activities that are responsible for degradation of the target molecule (Sinkunas *et al.*, 2011; Westra *et al.*, 2012; Sinkunas *et al.*, 2013). Evolution of resistance to new challenges occurs by the addition of spacers to the promoter-proximal end of the CRISPR repeat array, in a Cas1 and Cas2-dependent process (Barrangou and Marraffini, 2014).

In addition to providing resistance to exogenous sequences, *E. coli* CRISPRs have activity against lysogeny and induction of temperate bacteriophages (Edgar and Qimron, 2010). Induction of the CRISPR-Cas system results in *E. coli* cell death if crRNA targets are present on the chromosome, but the system also selects for bacterial populations that have lost prophages. Other functions attributed to CRISPR-Cas systems include modulation of bacterial community behaviors, gene expression, and DNA repair (Methe *et al.*, 2005; Viswanathan *et al.*, 2007; Zegans *et al.*, 2009; Aklujkar and Lovley, 2010; Babu *et al.*, 2010). Lastly, recent studies have implicated or established a role for CRISPR-Cas systems in promoting virulence of several pathogens including *Legionella pneumophila*, *Campylobacter jejuni*, and *Francisella novicida* (Gunderson and Cianciotto, 2013; Louwen *et al.*, 2013; Sampson *et al.*, 2013; Louwen *et al.*, 2014). Thus, the repertoire of cellular activities impacted by CRISPR-Cas appears to be diverse and much remains to be learned about these versatile elements particularly with regard to their influence on host interactions.

The work presented here was undertaken to determine the relationship of *nilD* to the CRISPR-Cas system and its function in mutualistic colonization of host nematodes. We demonstrate that the *nilD* locus expresses a CRISPR RNA molecule that contributes, in a Cas6e-dependent manner, to colonization of three distinct nematode species. Our data are consistent with a model that NilD functions to regulate endogenous bacterial sequences in a way that requires neither Cas3 nor perfect sequence identity to the target.

Results

The *nilD* locus is encoded within a CRISPR-Cas region

Heungens *et al.* (2002) previously reported a 3185-bp sequence (AY077466) of XnSc 081 containing the *nilD* locus required for association with *S. carpocapsae* nematodes. Further sequence analysis revealed this locus encodes a CRISPR repeat element, comprising

one spacer and two repeats, which is disrupted by the transposon insertion in the colonization deficient strain *nilD6::Tn5* (Fig. 1) (Heungens *et al.*, 2002). Additional CRISPR repeat sequences were noted upstream of *nilD* (bracketed region in Fig. 1A). These data indicate that *X. nematophila* encodes multiple CRISPR loci and that disruption of one of these, *nilD*, can inhibit nematode colonization.

To gain further information on the chromosomal context of *nilD* and to identify other CRISPR loci, the genome of XnSc 081 was sequenced and compared to that of the published sequence of *X. nematophila* strain HGB800 (NC_014228; ATCC 19061, hereafter referred to as XnSc 800) (Table S1). In both genomes there are six CRISPR elements (Table S2), labeled alphabetically in order of their occurrence in the chromosome (A–E and G) in addition to *nilD* (CRISPR-F). In both genomes, the *nilD* locus is located approximately 250-nt downstream of CRISPR-E. The *nilD* CRISPR is most similar to loci C and E: Each of these three loci (*nilD*, CRISPR-C, and CRISPR-E) encodes 32-nt spacer sequences and 29-nt repeats that are similar in sequence to those of *E. coli* K12 (Heungens *et al.*, 2002) (Fig. 1C). In turn, the *nilD* repeat sequences are similar, but not identical, to those of the loci C and E. In the 29 nt comprising each repeat, 6 and 4 differences occurred in the left and right repeats of *nilD* respectively, relative to the CRISPR-E repeats (Fig. 1C), indicating the *nilD* locus has diverged from the other CRISPR loci in the genome.

Encoded downstream of *nilD* is the previously sequenced *gloA* gene, as well as *cas* genes and CRISPR locus G (Fig. 1A). The sequences of each of these regions are identical between XnSc 081 and XnSc 800. CRISPR loci are preceded by A/T rich leader sequences containing promoters driving CRISPR transcription (Pul *et al.*, 2010). These leaders can be necessary for CRISPR function (Marraffini and Sontheimer, 2008; Sorek *et al.*, 2008) and their sequence tends to be conserved within, but not across, species (Jansen *et al.*, 1999; Lillestol *et al.*, 2006). In both XnSc 800 and XnSc 081, a 99-nt sequence adjacent to CRISPR-E (Fig. 1A) is 93% identical to that found upstream of CRISPR-C, and is likely the leader sequence. This sequence is not found adjacent to any other CRISPR locus (A, B, D, or G), nor is it found adjacent to *nilD*, suggesting that these loci may be regulated in a manner distinct from CRISPR-C and -E.

The *X. nematophila cas* genes are of the Type I-E subset and include the broadly conserved *cas1*, *cas2*, and *cas3* genes, as well as *cse1* (*casA*), *cse2* (*casB*), *cas7* (*casC*), *cas5* (*casD*), and *cas6e* (*casE*) (Fig. 1A) (Haft *et al.*, 2005; Makarova *et al.*, 2006; Chakraborty *et al.*, 2010; Makarova *et al.*, 2011b). Based on comparisons to *E. coli* and other systems (Brouns *et al.*, 2008; Sinkunas *et al.*, 2011; Westra *et al.*, 2012; Sinkunas *et al.*, 2013), we predict *cas3* encodes a protein with nuclease and helicase activity necessary for mediating degradation of crRNA-DNA hybrids, while the other five genes encode components of the ribonucleoprotein Cascade complex necessary for CRISPR RNA processing and target DNA degradation. In other systems *cas1* and *cas2* genes are not necessary for CRISPR RNA processing or activity, but rather encode nucleases that form a complex necessary for acquisition of new spacers (Fineran and Charpentier, 2012; Nuñez *et al.*, 2014). In addition, *cas1* is involved in DNA repair and chromosome segregation (Babu *et al.*, 2010), while *cas3* promotes plasmid replication in *E. coli* (Ivancic-Bace *et al.*, 2013).

Genomic analyses indicate that the spacer composition of the *E. coli* Type I-E system diversifies more slowly than would be expected if the CRISPRs were primarily involved in immunity (Touchon *et al.*, 2011). To address if *X. nematophila* spacer content diversifies during association with nematode and insect hosts we isolated DNA from 10 individual colonies of *X. nematophila* from our laboratory stock population of *S. carpocapsae* IJ nematodes that had been maintained for ~15 years by repeated (~monthly) passage through *Galleria mellonella* insect larvae. These “evolved” *X. nematophila* are the result of >1500 rounds of the natural life cycle, comprising persistence in non-feeding IJ nematodes during storage in water, infection and growth within insect larvae (including exposure to insect-associated microbiota), and colonization of nematode IJs (Richards and Goodrich-Blair, 2009). In contrast, XnSc 800 and XnSc 081 stocks have been stored for a similar period frozen in glycerol without propagation. There were no spacer sequence differences in CRISPR loci C, E or *niID* in the 10 isolated colonies relative to each other or to the frozen stock strains (data not shown), indicating that these loci are not evolving during laboratory passage through nematodes and insects. While we did not examine spacer content of the other 4 CRISPR loci in the evolved strain, the absence of spacer content changes in CRISPR loci C and E after more than 1500 passages through insects supports the idea that the CRISPR-Cas machinery in *X. nematophila* may function in a role outside of the canonical immunity against exogenous nucleic acids (Takeuchi *et al.*, 2012). Overall, the genomic analyses described above indicate that the *niID* locus, which is necessary for *X. nematophila* to colonize *S. carpocapsae* nematodes, encodes a non-canonical CRISPR element.

The *niID* CRISPR sequence is sufficient to promote nematode colonization

We previously reported that the colonization defect of the *niID6::Tn5* mutation was partially rescued by introduction of a plasmid (pSR2-312, Table 1) carrying a 312-bp fragment of wild type *niID*-containing DNA, confirming this region is necessary for colonization (Heungens *et al.*, 2002). Likewise, insertion of a 387-bp fragment encoding the *niID* locus (pEVS107-*niID*, Table 1) in single copy at the *attTn7* insertion site of the XnSc 081 *niID6::Tn5* genome (referred to hereafter as *niID6::Tn5 + niID*) was sufficient to restore nematode colonization, in this case to wild type levels (Fig. 2B). The DNA surrounding the transposon insertion encodes two putative divergent and overlapping small open reading frames, *orf1* and *orf2* that encompass the CRISPR element (Heungens *et al.*, 2002) (Fig. 1B) and may encode small peptides that could be involved in colonization. A plasmid carrying the *niID* sequence with a mutation at the common “T” of the start codons of these two ORFs did not rescue the colonization defect of the *niID6::Tn5* mutant (Heungens *et al.*, 2002). However, since this nucleotide is also the first within the 32-nt spacer (see underlined nucleotide in Fig. 1C), these previously reported data did not clarify if *orf1*, *orf2*, or the CRISPR-like element is involved in colonization. To help address this question, we transformed the *niID6::Tn5* mutant with derivatives of plasmid pSR2-312 (Heungens *et al.*, 2002), containing the 312-bp fragment sufficient to rescue colonization. Deletions were made in the pSR2-312 backbone such that the 5' (L) and/or the 3' (R) ends of the 312-bp *niID* region were truncated (Table 1, Fig. S1). These constructs were transformed into either the *niID6::Tn5* mutant or its wild-type parent XnSc 081 and transformants were tested for the ability to colonize IJ nematodes (Table S3). Constructs lacking substantial regions of *orf1* or *orf2* were able to rescue the colonization defect of the *niID6::Tn5* mutant, indicating

that neither ORF is required in its entirety to encode the colonization determinant. In contrast, the deletion constructs in which portions of the CRISPR repeat-spacer sequence are truncated were unable to rescue the colonization defect of the *nilD6::Tn5* mutant. Furthermore, a plasmid (pSR2- P90/ L84, Table 1) carrying a 137-bp central fragment containing the 90-nt *nilD* CRISPR-like region did rescue the colonization defect of the *nilD6::Tn5* mutant. These data support the hypothesis that within the *nilD* locus, the CRISPR-like sequences, not the short coding regions are necessary for colonization.

The role of the predicted NilD RNA in colonization was investigated further by introducing base substitutions that would alter the putative NilD RNA sequence but not the Orf1 or Orf2 peptide coding sequences (Fig. 2A). This construct (pEVS107-*nilD*-SDM, Table 1) was introduced in single copy in the *attTn7* site on the XnSc 081 *nilD6::Tn5* genome and the colonization phenotypes of the resulting strains were examined. Complementation with this construct failed to restore nematode colonization, indicating that NilD RNA rather than either Orf peptide is essential for nematode colonization and further supporting the predicted function of NilD as a crRNA (Fig. 2B). Consistent with this conclusion, attempts to detect expression of the Orf1 and Orf2 peptides by immunoblot and assaying *lacZ* translational fusions were unsuccessful (data not shown), indicating that these factors may not be stably expressed.

***nilD* encodes a small RNA transcript expressed during growth in lab culture and colonization of nematodes**

E. coli CRISPR repeats are transcribed and processed into small RNAs of ~57 nt (Brouns *et al.*, 2008). We sought to determine if the *nilD* locus similarly expresses a small RNA. Northern blots were insufficiently sensitive to detect an RNA transcript from the *nilD* region (data not shown). We therefore performed an RNase protection assay (RPA) using probes containing *nilD* sequence specific for either sense or anti-sense RNAs (relative to the transcript orientation of the CRISPR-like element predicted by comparison to *E. coli* (Brouns *et al.*, 2008)). No protected signal was observed in any reactions specific for anti-sense RNA (data not shown). In contrast, RNA harvested from wild type cultures, but not from the *nilD6::Tn5* mutant, protected a fragment of approximately 58-nt when probes specific for the sense strand were used (Fig. 3A), indicating the *nilD* region encodes a 58-nt RNA that is expressed under *in vitro* growth conditions. Similar results were observed when RNA was extracted from wild type bacteria harvested from IJ stage *S. carpocapsae* nematodes (Fig. 3B), demonstrating that the NilD RNA is also expressed during mutualistic interactions with its nematode host. Additionally, RPA analysis of RNA isolated from *X. nematophila* wild type cells grown under various *in vitro* conditions indicate that NilD RNA levels are elevated in nutrient-limited or aged cells (Fig. S2A). Higher NilD RNA levels were detected after growth in LB supplemented with 2, 2-dipyridil (an Fe(II) chelator) relative to LB alone or with additional supplementation with exogenous Fe₂SO₃, indicating NilD RNA levels may increase after iron limitation (Fig. S2B).

Primer extension was used to determine the 5' end of the ~58-nt NilD RNA (Fig. 3C). The run-off fragment indicates that the 5' end of NilD RNA is an adenine (designated +1 and indicated by an asterisk in Fig. 3C) seven nucleotides upstream of the spacer region. In

addition to this 5' end, we occasionally observed run off fragments consistent with the +2U as the 5' end (data not shown), which may indicate flexibility in processing or transcription initiation. The mapped 5' end of NiID RNA and the predicted 5' end match the 5' and 3' handles identified for *E. coli* crRNAs (Brouns *et al.*, 2008) (Fig. 1C). These results confirm that the *niID* locus encodes a small CRISPR RNA (genome coordinates: 3579434...3579491), hereafter referred to as NiID RNA (Fig. 1C).

NiID RNA expression requires Cas6e

To investigate the role of the Cas machinery in NiID RNA processing and nematode colonization, we used allelic exchange to generate mutations in *cas6e*, a gene predicted to encode an endoribonuclease responsible for processing of CRISPR transcripts into small RNAs, and in *cas3*, which is predicted to encode a helicase/nuclease required for CRISPR-mediated resistance to infection (Brouns *et al.*, 2008; Sinkunas *et al.*, 2011). RPA was used to detect *X. nematophila* NiID RNA and northern hybridization, using a general crRNA probe, was used to detect total CRISPR RNA in wild type and *cas* mutants (Fig. 4). In the XnSc 081 background, CRISPR small RNAs were absent in the *cas6e* mutant, but were present in the *cas6e* mutant complemented with the *cas6e* gene on a plasmid (compare Fig. 4B lanes 4 and 5 with lane 6). CRISPR RNAs were also apparent in the *cas3* mutant (Fig. 4B, lane 3). These data indicate that, as in *E. coli*, *cas6e* but not *cas3* is necessary for normal processing of crRNAs (Brouns *et al.*, 2008). Furthermore, crRNAs were expressed in the *niID6::Tn5* mutant (Fig. 4B, lane 2), suggesting that the nematode colonization defect of this mutant is not due to general disruption of crRNA expression. RPA analysis revealed NiID RNA, like other crRNAs, is apparent in the *cas3*-deficient (Fig. 4A, lanes 3 and 8) but not the *cas6e*-deficient strains (Fig. 4A, lanes 4, 5, 9, and 10), when expressed from its native locus (in the HGB081 background) or from the *attTn7* locus (in the *niID6::Tn5 + niID* background). Furthermore, providing a wild-type copy of the *cas6e* gene on a plasmid restored NiID processing (Fig. 4A, lanes 6 and 11). These results establish that the expression of the 58-nt NiID RNA depends on a component of the Cas machinery, further supporting its identity as a CRISPR RNA.

niID and *casE* are only necessary for colonization in a specific genetic background

Our data indicate that NiID RNA is not processed in the absence of *cas6e*. Furthermore, in *E. coli*, Cas3 is predicted to be required for processed crRNA function. We therefore predicted that in *X. nematophila* *cas6e* and *cas3*, like *niID*, would be required for nematode colonization. We first tested this by replacing the *cas6e* and *cas3* genes with a streptomycin resistance cassette in XnSc 081 and the *niID6::Tn5* mutant with (Tn7-*niID*) or without (eTn7) *niID*, generating a panel of *cas6e::strep* and *cas3::strep* strains. As predicted, disruption of *cas6e* in the *niID6::Tn5 + niID* strain resulted in a significant colonization defect, which was rescued by providing a wild type copy of *cas6e* on a plasmid (Fig. 5B). These data are consistent with the model that the function of Cas6e in colonization is to process NiID RNA into a crRNA, although we have not ruled out the possibility that it has a NiID-independent function in colonization. Contrary to our prediction, deletion of *cas3* resulted in no significant effects on nematode colonization. Therefore, in contrast to crRNAs in other systems, NiID activity is independent of Cas3.

Surprisingly, neither the *cas6e::strep* nor the *cas3::strep* mutations caused a colonization defect in the XnSc 081 strain background (the parent of the *nilD6::Tn5* mutant) (Fig. 5A), raising the possibility that the requirement for NilD RNA is specific to the *nilD6::Tn5* background. To further explore this hypothesis, *nilD* was replaced by allelic exchange with a kanamycin resistance cassette in the XnSc 081 and XnSc 800 wild type backgrounds. Like XnSc 081 *cas6E::strep*, the resulting strains, XnSc 081 *nilD56::kan* and XnSc 800 *nilD56::kan* mutants colonized *S. carpocapsae* to wild type levels (Table 2), despite a lack of NilD RNA detected by RPA (Fig. 3E and Fig. 4A).

These data verify that NilD and Cas6e are required for mutualism, but only within a specific genetic background of *X. nematophila*, suggesting a synthetic allele arose during the transposon mutagenesis process that gave rise to *nilD6::Tn5*. To examine possible synthetic mutations present in this background, we sequenced and compared the *nilD6::Tn5* draft genome to that of its parent XnSc 081 and found only one difference, a single nucleotide variation (SNV) located within the intergenic region of genes XNC1_3346 and XNC1_3347 (T-3271541-C). Gene XNC1_3346 is predicted to encode a P4-like DNA primase while XNC1_3347 is a small hypothetical gene in the DUF1795 superfamily that is followed immediately by XNC1_3348, predicted to encode an Rhs-like, YD-repeat-containing protein of unknown function (Fig. S3). Overlapping XNC1_3346 is an 1147-bp repeat sequence that occurs with 80–87% identity in two other locations of the genome (one full length copy and one truncated copy), also overlapping with genes with homology to those encoding P4-like DNA primases (Fig. S3). The SNV occurs within this repetitive region and an alignment of the three repeat regions shows variability in the nucleotides around the SNV (Fig. S3D). That the SNV associated with the *nilD6::Tn5* strain background is located within a phage-encoding region is consistent with it being involved in the activity of the CRISPR-Cas system, potentially as a target for NilD RNA. However, no obvious regions of sequence identity or complementarity were observed between NilD RNA (repeats or spacer) and the region surrounding the SNV.

A suppressor of the *nilD6::Tn5* colonization phenotype

The low level of colonization observed for the *nilD6::Tn5* mutant (~0.1 CFU/IJ; see for example Table 3) could reflect a majority of nematodes colonized by few bacterial cells, or full colonization (~50 bacteria) in one of many nematodes. The former phenotype might indicate the *nilD6::Tn5* mutant has a defect in outgrowth (Martens *et al.*, 2003), whereas the latter phenotype might indicate the *nilD6::Tn5* mutant has a defect in initiation of colonization which can be occasionally suppressed. To distinguish between these possibilities we examined by epifluorescence microscopy the frequency of colonized nematodes after cultivation on a GFP-expressing *nilD6::Tn5* strain (XnSc 829; Table 1). This analysis showed that the majority of individual nematodes were uncolonized and that approximately 1 in 600 nematodes were fully colonized (data not shown). To determine if rare colonization events were due to genetic or epi-genetic suppression of the *nilD6::Tn5* mutation we examined the colonization phenotype of a colony isolate derived from *nilD6::Tn5*-colonized nematodes. Upon re-cultivation with nematodes, this colony isolate, XnSc 1186 (*sup-1*) exhibited significantly higher levels of colonization than its *nilD6::Tn5* parent (5.48 ± 0.57 vs. 0.15 ± 0.47 Avg. CFU/IJ respectively, $n = 6$, $p < 0.001$ by unpaired

Student's t-test), despite the absence of detectable NilD RNA by RPA (Fig. 3E). These data indicate that the phenotype caused by the *nilD6::Tn5* mutation can be suppressed, presumably by nucleotide or epigenetic changes elsewhere on the chromosome. The latter seems most likely, as sequencing of the XnSc 1186 *sup-1* genome did not reveal mutations that could explain the suppression phenotype (data not shown).

The *nilD* locus contributes to XnSc 081 association with different nematode species

In addition to *S. carpocapsae*, *X. nematophila* associates with two other nematode species, *S. websteri* and *S. anatoliense* (Lee and Stock, 2010). To examine if NilD RNA is required for colonization of these other nematode hosts, the colonization phenotypes of the *nilD6::Tn5* mutant and XnSc 081 in *S. anatoliense* and *S. websteri* were assessed. Similar to the phenotypes observed in *S. carpocapsae* colonization assays, XnSc 081 colonized *S. anatoliense* and *S. websteri* while the *nilD6::Tn5* mutant displayed a marked defect in its ability to associate with these host nematode species (Table 3). In all three nematode hosts the colonization defect of the *nilD6::Tn5* mutant was rescued by insertion of the XnSc 081 *nilD* sequence at the *attTn7* insertion site (Table 3).

Strain specific NilD RNA variants are expressed in the *S. anatoliense* and *S. websteri* symbionts but do not rescue the *nilD6::Tn5* colonization defect

To determine if the *nilD* locus is present in all *X. nematophila* strains regardless of the identity of their natural nematode host, we searched for it in *X. nematophila* strains from *S. anatoliense* and *X. websteri* (XnSa 1418 and XnSw 1419 respectively). Oligonucleotides (NilD 5' ApaI and NilD 3' KpnI, Table S4) complementary to flanking regions around the *nilD* region of XnSc 081 successfully amplified products from XnSa 1418 and XnSw 1419 genomic DNA. In each case a product was obtained of similar size to that amplified from XnSc 081 and XnSc 800 (data not shown). The products amplified from XnSa 1418 and XnSw 1419 genomic DNA were cloned and sequenced. Relative to the *S. carpocapsae*-derived *X. nematophila* strains, the XnSa 1418 and XnSw 1419 *nilD* regions encode an identical left repeat, 4-nt differences within the 32-nt spacer, and an identical right repeat except for the last 3 nt (Fig. 1D). Also, unlike the XnSc 081 *nilD* sequence, the *nilD* regions of XnSa 1418 and XnSw 1419 are not predicted to encode small overlapping open reading frames (data not shown).

To assess if the *nilD* loci of XnSa 1418 and XnSw 1419 encode a NilD RNA molecule, we conducted RNase protection assays (RPA) using probes that match the XnSa 1418 *nilD* sequence. RPA using this probe detected protected fragments in both the XnSa 1418 and XnSw 1419 samples that were similar in size to those that were detected (using XnSc specific probe) in the XnSc 081 and XnSc 800 samples (Fig. 3D). These data indicate that NilD RNA is expressed and processed similarly in all four Xn strains.

Given the divergence of the XnSa 1418 and XnSw 1419 *nilD* spacers from those of Sc *X. nematophila* strains, we examined if the former could rescue the colonization defect of the *nilD6::Tn5* mutant. The XnSa 1418 and XnSw 1419 *nilD* sequences were cloned and inserted at the *attTn7* site of the XnSc 081 *nilD6::Tn5* mutant, generating strains *nilD6::Tn5* + *nilD*-XnSa (*nilD* + Sa) and *nilD6::Tn5* + *nilD*-XnSw (*nilD* + Sw). RPA analysis was used

to verify expression of the NilD RNA (Fig. 3D) while the colonization proficiency of the complemented strains was assessed using *Sc* nematodes (Table 3). As expected, insertion of the XnSc *nilD* sequence (strain *nilD6::Tn5 + nilD*) was sufficient to restore wild type levels of colonization. In contrast, providing the XnSa 1418 and XnSw 1419 sequences failed to restore levels of colonization above the level of the *nilD6::Tn5* mutant strain (Table 3). These results are consistent with our findings described above that alteration of the NilD RNA sequence was sufficient to disrupt the activity of this molecule. *nilD6::Tn5 + nilD-XnSa* and *nilD6::Tn5 + nilD-XnSw* were also deficient for colonization of *S. anatoliense* and *S. websteri* nematodes (Table 3), indicating that distinct *nilD* sequences do not confer specificity for these nematode species.

To further explore the possible role of *nilD* in host range specificity, we introduced it into *X. bovienii* (the symbiont of the nematode *S. jolietii*) that naturally lacks *nilD* (Chaston *et al.*, 2011; Sugar *et al.*, 2012). *X. bovienii* is unable to colonize *S. carpocapsae* unless it expresses the host-range specificity determinants *nilA*, *B*, and *C* (Cowles and Goodrich-Blair, 2008; Chaston *et al.*, 2013). We therefore expressed *nilD* in *X. bovienii* with and without the *nilA*, *B*, and *C* genes. The presence of *nilD* did not impact the colonization levels of *X. bovienii*: colonization of *S. carpocapsae* nematodes was below the level of detection (0.005 CFU/IJ) when *nilD* was present without *nilA*, *B*, and *C*, and colonization levels of *X. bovienii* carrying *nilA*, *B*, and *C* were not increased by the presence of *nilD* (data not shown).

Discussion

The goal of this study was to elucidate the mechanistic role of the *X. nematophila nilD* locus during colonization of *S. carpocapsae* host nematodes. Our work demonstrates that *nilD* encodes a small RNA and that expression of this molecule is sufficient to rescue the colonization defect of the *nilD6::Tn5* strain. Bioinformatic predictions indicated that NilD RNA is a member of the CRISPR RNA family. Consistent with this, we present experimental evidence that NilD RNA processing and colonization function requires *cas6e*, encoding a homolog of the *E. coli* CRISPR RNA processing Cascade complex endonuclease (Cas6e) (Westra *et al.*, 2012). However, unlike the CRISPR-Cas systems of other bacteria, NilD RNA function does not appear to require the helicase-nuclease Cas3 (Sinkunas *et al.*, 2011), since a *cas3* mutant does not display a colonization defect (Fig. 5). This may indicate that the function of NilD RNA diverges from that of other crRNAs, and that its requirement in colonization does not include Cas3-mediated target degradation.

Several lines of evidence argue against the idea that the colonization function of NilD RNA is to restrict entry of exogenous DNA (plasmids and lytic bacteriophages), the initial primary function ascribed to crRNAs (Brouns *et al.*, 2008; Marraffini and Sontheimer, 2008). Instead, our data support the model that NilD RNA regulates endogenous sequences, as has been observed or suggested in other CRISPR-Cas systems (Zegans *et al.*, 2009; Aklujkar and Lovley, 2010; Cady and O'Toole, 2011; Sampson *et al.*, 2013). First, the *nilD6::Tn5* colonization defect is apparent in a closed system consisting only of a bacterial clonal population and the nematode host. Therefore, the only source of potentially toxic foreign DNA is the nematode. However, nematode lysates do not inhibit growth of

Xenorhabdus strains in liquid culture and do not form plaques on bacterial lawns (data not shown). Further, BLASTn analyses (Altschul *et al.*, 1997) against the NCBI GenBank sequence database and to 13 other *Xenorhabdus* bacterial genomes (H. Goodrich-Blair, unpublished) failed to identify putative proto-spacers with identity to NilD (data not shown). While not conclusive, this fact is contrary to the idea that the NilD RNA targets a mobile genetic element present in other *Xenorhabdus* spp. Second, differences in the endogenous chromosome can bypass or cause the need for NilD RNA. The *nilD6::Tn5* colonization defect is only apparent in a specific genomic background (Fig. 5) in which suppressor alleles (e.g. *sup-1*) can arise that are able to colonize despite the absence of *nilD* expression (Fig. 3). Our genomic analysis indicates the strain background associated with the requirement for *nilD* in colonization has a single distinguishing SNV, a T to C change at nt 3271541 in the intergenic region between a phage P4 primase and a region predicted to encode Rhs-like and associated elements (Fig. S3), while sequencing of the *sup-1* strain failed to identify any genetic alterations that could account for the suppression phenotype (data not shown). These findings suggest that a single nucleotide change in the bacterial genome may confer dependence upon *nilD* for colonization, whereas suppression may result from epigenetic or phase variability.

An alternative explanation for the role of NilD RNA in colonization is that it is required to control expression of an endogenous genetic element that is detrimental for host interactions, with the NilD RNA 32-nt spacer region conferring specificity for this element. Similar to the *E. coli* CRISPR system, NilD may control expression of its targets (Edgar and Qimron, 2010). Two models of CRISPR-Cas-mediated gene regulation are that the Cascade complex, including the crRNA binds to target mRNA to block translation or to promote Cas-3-mediated cleavage, or the Cascade complex binds to the DNA target and prevents transcription. Since a *cas-3* mutant does not display the same colonization defects as the *nilD6::Tn5* mutant (Fig. 5), our data are most consistent with the idea that the NilD RNA-Cascade complex blocks either transcription or translation, rather than triggering target degradation. Further insights into the mechanism of NilD RNA function await identification of its target(s). No proto-spacer with 100% identity is apparent in the genomes of XnSc 800 or XnSc 081, suggesting that low levels of similarity may be sufficient for NilD targeting. Conversely, complementation experiments using mutagenized XnSc 081 *nilD* and the divergent *nilD* loci of XnSa 1418 and XnSw 1419 failed to restore the colonization defect of *nilD6::Tn5* (Table 3), revealing that some sequence integrity is essential. Furthermore, these data may indicate that the NilD RNA targets in strains XnSa 1418 and XnSw 1419 have diverged in sequence, and that the *nilD* loci in those strains have co-evolved to maintain sequence identity. If true, further comparative sequence analysis of these strains could yield putative targets.

This report extends the limited number of studies demonstrating an impact of CRISPR-Cas systems on host-microbe interactions. The *cas2* gene of *Legionella pneumophila* is required for intracellular growth within host amoebae (Gunderson and Cianciotto, 2013). Similar to our work, these experiments were conducted in the absence of exogenous DNA or phage, suggesting that the requirement for *cas2* in *L. pneumophila* was independent of any interference-related functions. Likewise, a *cas2* mutant was not more sensitive upon

exposure to UV light or to treatment with mitomycin C or nalidixic acid, indicating that the virulence defect was not due to the a potential role for Cas2 in DNA repair (data not shown). A recent study demonstrated *F. novicida* Cas9-mediated negative regulation of an endogenous gene encoding a lipoprotein. In the absence of *cas9*, aberrant expression of the lipoprotein triggered host immunity and reduced virulence of the pathogen (Sampson *et al.*, 2013). Together these and other studies have established a role for CRISPR-Cas machinery in facilitating pathogen virulence (Louwen *et al.*, 2014). The work presented here demonstrates these systems can also function in mutualistic associations. Though still enigmatic, the role of NilD RNA and its synthetic allele in nematode colonization should be clarified by identifying its molecular target, and in turn the function of this target in either promoting or inhibiting the colonization process.

Experimental Procedures

Organisms and growth conditions

Strains used in this study are listed in Table 1. Unless otherwise noted, cultures were grown at 30°C in LB broth (Miller, 1972). *X. nematophila* growth media were stored in the dark or supplemented with 0.1% pyruvate (Xu and Hurlbert, 1990). Media were supplemented with kanamycin (Km, 50 µg/ml), rifampicin (Rif, 100 µg/ml), ampicillin (Amp, 150 µg/ml), streptomycin (Sm, 25 µg/ml), or chloramphenicol (Cm, 30 µg/ml) where appropriate. For indicated RNA isolations, cultures were supplemented with 500 µM Fe₂SO₃, 100 µM 2,2-dipyridyl or 1 µM deferoxamine (Sigma-Aldrich, St. Louis, MO). The nematodes *Steinernema carpocapsae* (Weiser) All, obtained from Harry Kaya, and *S. anatoliense* (Al-Jubiha Jordan) and *S. websteri* (Peru), obtained from S. Patricia Stock, were reared in *Galleria mellonella* wax moth larvae (Kaya and Stock, 1997). For *in vitro* co-cultures nematodes were grown at room temperature (20–26°C) on lipid agar (LA) plates with lawns of test *X. nematophila* strains as previously described (Vivas and Goodrich-Blair, 2001). Defined medium was as previously described (Orchard and Goodrich-Blair, 2004) except that glutamate was added at 100 mg l⁻¹ and Bacto agar (Sigma-Aldrich, St. Louis, MO) was used instead of noble agar. *X. nematophila* strains from *S. anatoliense* and *S. websteri* were isolated by surface sterilization of 1000–10000 IJ nematodes for 3 min in 0.5% NaOCl as described previously (Heungens *et al.*, 2002). Surface-sterilized nematode were sonicated for 1 min (Cowles and Goodrich-Blair, 2004) and dilution plated onto LB + 0.1% pyruvate agar. Individual colonies were streaked for isolation, cultured overnight at 30°C, and frozen in glycerol stocks. Bacterial identity was verified by Sanger sequencing of the 16S gene using primers 27F and 1492R (Table S4) (Lane, 1991).

DNA manipulations and biochemical assays

Plasmids used in this study are listed in Table 1. To create pCR2.1-TOPOmini, which lacks the Kan^R cassette, primers TOPO2.1mini_Fwd_NcoI and TOPO2.1mini_Rev_NcoI (Table S4) were used to amplify the backbone of the plasmid pCR2.1-TOPO. The amplified product was cut with NcoI and self-ligated. Standard protocols were used for the isolation of chromosomal DNA, DNA digestion, electrophoresis, and electroporation (Sambrook *et al.*, 1989). Enzymes for DNA manipulations were obtained from Promega (Fitchburg, WI), New England Biosciences (Ipswich, MA) or Fermentas (Pittsburg, PA). Plasmid isolations and

gel purifications were performed using appropriate kits (Qiagen, Germantown, MD). PCR amplification was performed using ExTaq polymerase, Primestart polymerase (Takara Shuzo, Kyoto, Japan) or Pfu Ultra (Agilent Technologies, Madison, WI) and appropriate buffers on 100 ng *Xenorhabdus* chromosomal template-DNA, 0.2 μ M each primer, 0.4 mM dNTPs, and 2.5 U of polymerase. After 2 min incubation at 95°C, 30 cycles of 20 s at 95°C, 30 s at an annealing temperature appropriate for each primer pair, and 60-s kb^{-1} at 72°C, were conducted, followed by 7 min incubation at 72°C.

pECMXb-Empty and pECMXb-SR2 construction, conjugation into *X. bovienii*

The pECMXb-Empty vector was constructed from pECM20 (Martens *et al.*, 2003) by replacing the *X. nematophila* insertion sequence with a 588-bp fragment of *X. bovienii* intergenic genomic DNA (coordinates 390649-391236 of *X. bovienii* SS-2004; NC_013892.1) to facilitate homologous recombination of the plasmid into the *X. bovienii* genome. The pECM20 plasmid was divergently amplified on either side of the *X. nematophila* insert site to replace the insertion with the restriction sites for ApaI and KpnI. Primers used were pECM20_Xb_F and pECM20_Xb_R (Table S4). A predicted intergenic region of *X. bovienii* was amplified using primers pECMXb_insert_F and pECMXb_insert_R, and the sequence was inserted into pECMXb using ApaI and KpnI. The insert was confirmed by sequencing using primers pECMXb_seq_F and pECMXb_seq_R (Table S4). For construction of pECMXb-SR2, the SR2 region was sub-cloned from pTopoSR2-2 into the pECMXb XbaI site using XbaI and SpeI.

The pECMXb-Empty and pECMXb-SR2 plasmids were conjugated in *X. bovienii* 1166 and 1167 using previously described methods (Murfin *et al.*, 2012). Ex-conjugants were grown on LB media supplemented with 15 μ g/ml of chloramphenicol to select for insertion of the plasmid. Integration of pECMXb-Empty and pECMXb-SR2 into the genome was confirmed by positive PCR results using primers pECMXb_integratation_F and pECMXb_integratation_R (Table S4).

Isolation of *X. nematophila cas* and *niID* mutants

The 4,857 bp DNA fragment containing 2,751 bp *cas3* gene and its up-stream (1,213 bp) and down-stream (893 bp) sequences were amplified from HGB800 chromosomal DNA using Pfu DNA polymerase (Stratagene, Santa Clara, CA) and primers cas3UpFwd_SpeI and cas3DownRev_XbaI. Likewise, the 2,553 bp DNA fragment containing 678-bp *cas6e* (termed *casE* in strain designations) and its up-stream (1,233 bp) and down-stream (642 bp) sequences were amplified using primers casEUpF_SpeI and casEDownR_XbaI, respectively (Table S4). The resulting fragments were digested with XbaI and SpeI and cloned into plasmid pCR2.1-TOPOmini between XbaI and SpeI sites. The *ahp* kanamycin resistance cassette and its promoter region were amplified from plasmid pEVS107 using primers Kan-CleanRev_EcoRV_NEW and Kan-FullFwd_NheI_NEW (Table S4) digested with NheI and EcoRV, and used to replace the 2,362 bp NheI-EcoRV region (89-2451 bp) within the *cas3* gene and the 26-321 bp region of the *cas6e* gene, generating pCR2.1 mini *cas3::kan* and pCR2.1 mini *casE4::kan*.

To create *cas6e::strep* and *cas3::strep* insertion constructs used in this study, the Kan^R cassettes in pCR2.1 mini *casE4::kan* (HGB1692) and pCR2.1 mini *cas3-3::kan* were removed by cutting with EcoRV and NheI. The remaining backbone of the plasmid was ligated to EcoRV/SpeI fragment, containing the Sm^R cassette from pKNG101, to form pCR2.1 mini *casE6::strep* and pCR2.1 mini *cas3-5::strep*. The *casE6::strep* and *cas3-5::strep* fragments were cut from the pCR2.1 mini backbone using SpeI and XbaI and cloned into the SpeI site of the mobilizable suicide plasmid pKNG101, generating pKNGD*casE6::strep* and pKNGD*cas3-5::strep*. The resulting constructs were maintained by electroporating into *E. coli* SM10 (*λpir*) cells and then introduced into HGB081, HGB800 and HGB1940 (Table 1) by conjugation, as described previously (Forst and Tabatabai, 1997). Ex-conjugants were grown on LB agar containing 25 µg/ml streptomycin overnight and subsequently grown on LB agar plus 5% sucrose to select for sucrose-resistant ex-conjugants that had excised the vector. The Sm^R phenotype was verified, and deletion of the *cas6e* or *cas3* gene fragments was confirmed by PCR amplification.

For deletion of the *nilD* region, a 1,542 bp fragment upstream of *nilD* was amplified from the XnSc 081 chromosome using PFU Ultra polymerase and the dNilD Up 5' SalI and dNilD Up 3' ApaI primers. Likewise, a 1,082 bp fragment downstream of *nilD* was amplified using the dNilD Dwn 5' ApaI and dNilD 3' SacI primers while the Kan^R cassette was amplified from plasmid pEVS107 using primers Kan 5' ApaI and Kan 3' ApaI (Table S4). PCR fragments were digested using appropriate restriction enzymes and then ligated into pKR100 plasmid, linearized with Sall and SacI enzymes, generating pKR100-*nilD56::kan*. The resulting construct was maintained by electroporation of *E. coli* S17.1 (*λpir*) and introduced into HGB081 and HGB800 by conjugation. Ex-conjugants were grown on LB agar containing 50 µg/ml kanamycin overnight and then screened for loss of resistance to chloramphenicol. The deletion of the *nilD* locus was confirmed by PCR and RPA was utilized to confirm NilD RNA was not being produced.

Complementation studies

To generate *nilD* truncation constructs, portions of the *nilD* locus were amplified (using primers indicated in parentheses) and cloned into pCR2.1[®]-TOPO to create pTopoSR2-R90 (KHP62 and KHP58), - R126 (KHP62 and KHP64), - L84 (KHP57 and KHP63), - L132 (KHP55 and KHP63), - L161 (KHP36N and KHP63), and - R90/ L84 (KHP57 and KHP58) (Table S4). Once constructed, all fragments were subcloned from pCR2.1[®]-TOPO into pBCSK+ using ApaI and SacI to create pSR2- R90, pSR2- R126, etc. To generate a stable *nilD* complemented strain, a 387-nt fragment encoding the *nilD* crRNA region and approximately 175-nt upstream, was amplified using Primestart polymerase (Takara Shuzo, Kyoto, Japan) and primers nilD 5' ApaI and nilD 3' KpnI (Table S4). The *nilD* PCR fragment and the Tn7-insertion vector, pEVS107 (Table 1), were digested with ApaI and KpnI restriction enzymes and ligated using T4 DNA ligase (New England Biosciences). The resulting vector, pEVS107-*nilD*, was maintained by electroporating into competent S17.1 *λpir E. coli* cells followed by selection on LB plates supplemented with kanamycin.

To complement the *nilD*-deficient strain (*nilD6::Tn5*) with a *nilD* region encoding synonymous base mutations within the spacer sequence, pEVS107-*nilD* was subjected to a series of site directed mutagenesis reactions. The pEVS107-*nilD* vector was first amplified using the NilD SDM set 1F and 1R primers (Table S4) to generate base substitutions within codons 2 and 3 in the *nilD* spacer region. The resulting construct was then further mutated using NilD SDM sets 2 through 5 (Table S4) to generate synonymous base substitutions within codons 4–11, forming pEVS107-*nilD*-SDM (Table 1). For site directed mutagenesis reactions, approximately 3 µg of pEVS107-*nilD* DNA were mixed with 15 pmol of each primer, 4 µl of DMSO, 50 µmol dNTP mix, 5 µl PFU Ultra buffer and 1 µl PFU ultra polymerase (2.5 units/µl) (Agilent Technologies) in 50 µl of total volume. After 1 minute at 95°C, the resulting mixtures were incubated at 95°C for 1 min, 56°C for 50 s, and 72°C for 10 min for 25 cycles, followed by 10 additional min at 72°C. Following amplification, template DNA was digested by incubation with 10 units of ApaI restriction enzyme at 37°C for 1 hr and the resulting PCR product was maintained by electroporation into S17.1 λ pir *E. coli* cells and selection on LB supplemented with kan. All plasmids were sequenced to ensure that the appropriate mutations were present and that no additional mutations had occurred within the *nilD* region.

To generate the *casE* complementation construct, the *cas6e* gene was amplified using the CasE 5' XbaI and CasE 3' EcoRV primers (Table S4) and Primestar polymerase (Takara Shuzo, Kyoto, Japan). The PCR product and pBluescript (pBCSK+) vector (Stratagene, La Jolla, CA) were subjected to restriction digestion with XbaI and EcoRV enzymes and ligated using T4 ligase. The resulting vector, pBC-*casE* (Table 1), was introduced into Top 10 *E. coli* cells (Invitrogen, Carlsbad, CA) and maintained by selection on LB plates supplemented with chloramphenicol.

Complementation of *Xenorhabdus* strains was performed as previously described (Bao *et al.*, 1991; Forst and Tabatabai, 1997). Briefly, for complementation of the *nilD* mutation, overnight cultures of *nilD6::Tn5*, S17.1 + pEVS107-*Tn7/nilD* and the transposition helper strain S17.1 + pUX-BF13 (Bao *et al.*, 1991), were diluted 1:100 in LB and incubated for 3 h at 30°C. After incubation, 900 µl of *X. nematophila* culture was mixed with 600 µl of S17.1 + pEVS107-*Tn7/nilD*, and 500 µl S17.1 + pUX-BF13. The mixture was then pelleted by centrifugation, resuspended in 30 µl of LB and spotted onto LB plates supplemented with 0.1% pyruvate. Approximately 18 h after plating, cells were scraped into 1 ml of LB and 50 µl were plated onto LB supplemented with 0.1% pyruvate, and containing ampicillin and erythromycin for selection. Resistant colonies were screened for proper insertion at the Tn7 site by PCR analysis. For complementation with *nilD* truncation or *cas6e* constructs, chemically competent *X. nematophila* strains were generated as previously described (Xu *et al.*, 1989) and transformed with 200 ng of individual vector constructs. Transformants were plated on LB supplemented with 0.1% pyruvate and chloramphenicol.

RNA isolations and analyses

For RNA isolation from cultured bacteria, *X. nematophila* strains were grown for 18 h in either liquid LB or LB containing 100 µM dipyrindyl (to promote optimal NilD expression) and 0.1% pyruvate. Cultures were then diluted and cells were harvested during stationary

phase or during late log phase (OD₆₀₀ of 1.0). Alternatively, for analysis of gene expression on solid media, cells were harvested from LA agar plates after 1 or 8 d of incubation, or from defined medium plates after 1 d. All strains were grown at either 20° or 30°C. RNA for RPA was isolated from individual strains using Trizol Reagent (Life Technologies, Madison, WI), as previously described (Wassarman and Storz, 2000). The presence of approximately equal amounts of RNA between treatments was confirmed by agarose gel-electrophoresis (data not shown). Small RNA for northern analysis of CRISPR RNAs was isolated using mirVana kits (Applied Biosystems, Life Technologies, Madison, WI) according to manufacturer methods except for modifications to facilitate cell lysis as previously described (Cavanagh *et al.*, 2012).

For isolation of RNA from symbiotic bacteria, approximately 100,000 nematodes were harvested from lipid agar plates, suspended in LB, and lysed by sonication for 1 minute in a water bath sonicator. *Xenorhabdus* bacteria were harvested by centrifugation and RNA was isolated using Trizol Reagent (Life Technologies, Madison, WI), as described above.

Primer extension experiments

For primer extension, PAGE purified primers AAP1 and AAP2 (Table S4) were radioactively labeled using T4 Polynucleotide Kinase and $\gamma^{32}\text{P}$ -ATP (Perkin Elmer, Waltham, MA) for 30 min at 37°C. Excess nucleotides were removed using a QIAquick® (Nucleotide Removal kit, Qiagen, Valencia, CA). Labeled primer was hybridized to 5 μg of total cell RNA in Avian Myeloblastosis Virus Reverse Transcriptase (AMV-RT) buffer (Promega, Madison, WI) by heating to 80°C for 10 min and slow cooling to 37°C. The primer was extended by AMV-RT enzyme at 37°C, precipitated, and resuspended in gel loading buffer (Promega, Madison, WI). A plasmid-based copy of *X. nematophila* SR2-2 region was cycle sequenced using *fmol*® DNA Cycle Sequencing System kit (Promega, Madison, WI) and the labeled primer. The sequencing reaction was stopped using *fmol*® sequencing stop solution (Promega, Madison, WI). Samples were electrophoresed on a 12% polyacrylamide gel (National Diagnostics, Charlotte, NC) that was then dried and imaged on a Storm860 phosphorimager (Molecular Dynamics, Sunnyvale, CA).

RNase protection assays

RPA analyses were performed using the Ambion RPA III kit™ method (Ambion Inc., Austin, TX), following manufacturer recommendations. Probes for XnSc *nilD* were generated by transcription from pAWA1 (Table 1) template containing a 137 bp fragment containing the *nilD* locus of XnSc. For generation of a probe complementary to the *nilD* region of XnSa and XnSw, site directed mutagenesis was utilized to alter the spacer region of *nilD* within TOPOSR2-2. SDM was performed as described above using PFU Ultra polymerase and primers RPA SDM 5' and 3' (Table S4) to generate pTOPO-*nilD*-Sa (Table 1). Complementary transcripts were amplified and radioactively labeled with $\alpha^{32}\text{P}$ -UTP (PerkinElmer, Waltham, MA) using the MAXIscript®-T7 reaction (Ambion, Austin, TX). Labeled transcripts were separated on a 10% polyacrylamide gel (National Diagnostics Inc, Charlotte, NC) and probes were gel purified. 10 μg of sample RNA was then hybridized to 1–2 x 10⁵ cpm of purified probe and RNase treated using a 1:100 dilution of RNase A/T1 enzyme (Ambion, Austin, TX). The RNase treated samples were electrophoresed on 10%

polyacrylamide gels and visualized using XAR ECL- film (Kodak, Rochester, NY). A no-RNase control was also run to confirm probe integrity.

Northern

Total small RNA was separated on 12% denaturing polyacrylamide MOPS gels, transferred to uncharged nylon membrane, and probed first for crRNAs with pAAP1 oligonucleotide (which anneals to the conserved 3' repeat of all crRNAs; Table S4) using methods previously described for LNA probes except that 2.5 µg small RNA was loaded per lane and hybridization was done at 50°C (Cavanagh *et al.*, 2012). Membranes were then reprobbed with a full length RNA probe to *E. coli* 5S RNA (generated from pBS-5S) as previously described (Wassarman and Storz, 2000).

Nematode colonization assays

For colonization assays, lawns of individual bacterial strains were grown on lipid agar for 48 hours. Aposymbiotic infective juvenile-stage nematodes were generated by cultivation on a non-colonizing *rpoS* mutant, then surface sterilized, using a diluted bleach solution, and applied to the bacterial lawns (Vivas and Goodrich-Blair, 2001). In each experiment, two independent cultures of each strain were used as replicates, and 3 plates per replicate were seeded. Approximately one week post-inoculation, plates were water trapped for harvesting of IJs. Roughly 10⁴ progeny IJs were then harvested from each plate (Vivas and Goodrich-Blair, 2001), surface sterilized and disrupted by sonication. The macerates were dilution plated on selective media to quantify CFU/IJ as previously described (Heungens *et al.*, 2002). For *nilD* deletion analyses, XnSc 081 and *nilD6::Tn5* were transformed (Xu *et al.*, 1989) with the plasmids indicated for each experiment and transformants were co-cultivated with nematodes on LA plates containing rifampicin and chloramphenicol. For *casE* complementation analyses, strains containing pBC-*casE* were grown on lipid agar plates supplemented with 10 µg/ml Cm and 1 mM IPTG.

Sequencing of *X. nematophila* strains

The genomes of *X. nematophila* strains XnSc 081, *nilD6::Tn5*, and *nilD6::Tn5 sup-1* were sequenced using Illumina paired-end libraries (mean insert length = 300bp) yielding approximately 20–30 million 75 base-pair, paired-end reads for each strain. The resulting reads were trimmed for quality and then used in a reference alignment with respect to XnSc 800 using CLC Genomics Workbench 5.1 (CLC Bio). Assembled genomes were then analyzed for deletions, insertions and single nucleotide variations using CLC Genomics Workbench 5.1. SNVs were manually inspected for verification. Regions where low sequence coverage was obtained (fewer than 8 reads of coverage) were amplified and cloned from individual genomes and then sequenced. Because of significant genomic differences between XnSc 800, XnSa 1418 and XnSw 1419, performing reference alignments was not feasible. As a result, CLC Genomics Workbench 5.1 software was utilized to generate *de novo* genome assemblies using the reads from XnSa 1418 and XnSw 1419.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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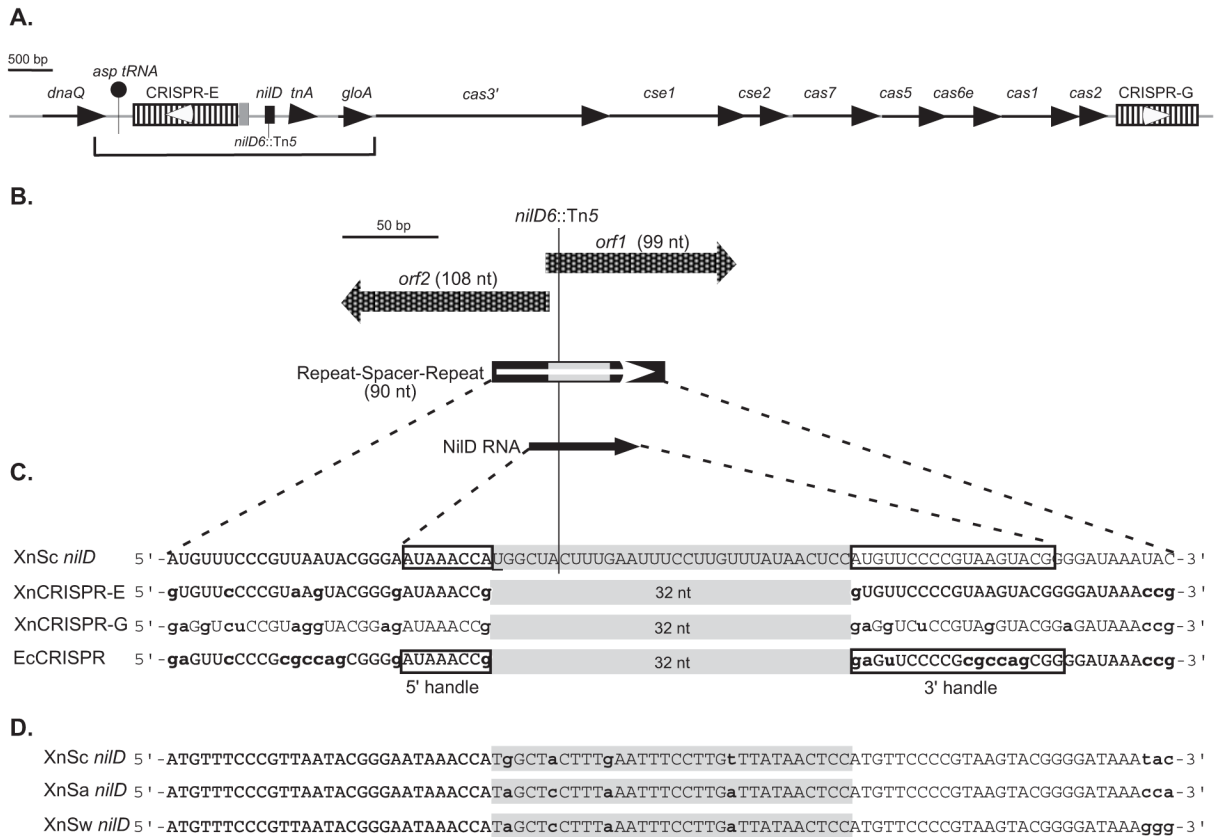


Figure 1.

Schematic representation of the *nilD* CRISPR locus.

A. Schematic diagram of the *X. nematophila* genomic regions containing CRISPR loci, *cas/cse* genes, and *nilD*. The bracket indicates the 3240-bp region previously sequenced in the HGB081 (XnSc 081) strain background (AY077466) (Heungens *et al.*, 2002), which is identical in the sequenced ATCC 19061 (HGB800/XnSc 800) genome (NC_014228.1). Line arrows represent open reading frames, with gene names indicated above each. CRISPR loci are represented by hatched rectangles and are named CRISPR-E and CRISPR-G according to their position on the *X. nematophila* genome, with *nilD*, shown as a black rectangle, considered CRISPR-F. The location of the *nilD6::Tn5* transposon insertion within *nilD* is indicated. The gray shaded box represents the 135-bp leader sequence of CRISPR-E. White arrowheads indicate the predicted orientations of CRISPR-E and -G transcription, based on comparison to *E. coli* CRISPR transcription.

B. Detail of the *nilD* locus, showing the two small overlapping open reading frames (*orf1* and *orf2*) represented by checkered block arrows. The positions of *nilD* locus repeats and spacer are indicated by black and gray rectangles, respectively. The white arrow indicates the predicted orientation of transcription based on comparison to *E. coli* CRISPR transcription. The black arrow represents the position of the small RNA transcript encoded by the *nilD* locus. The position of the *nilD6::Tn5* insertion site is indicated by a line.

C. Sequence of NiID RNA aligned with CRISPR small RNAs predicted to be encoded by *X. nematophila* (XnCRISPR-E and -G) and CRISPR RNAs expressed in *E. coli* (EcCRISPR).

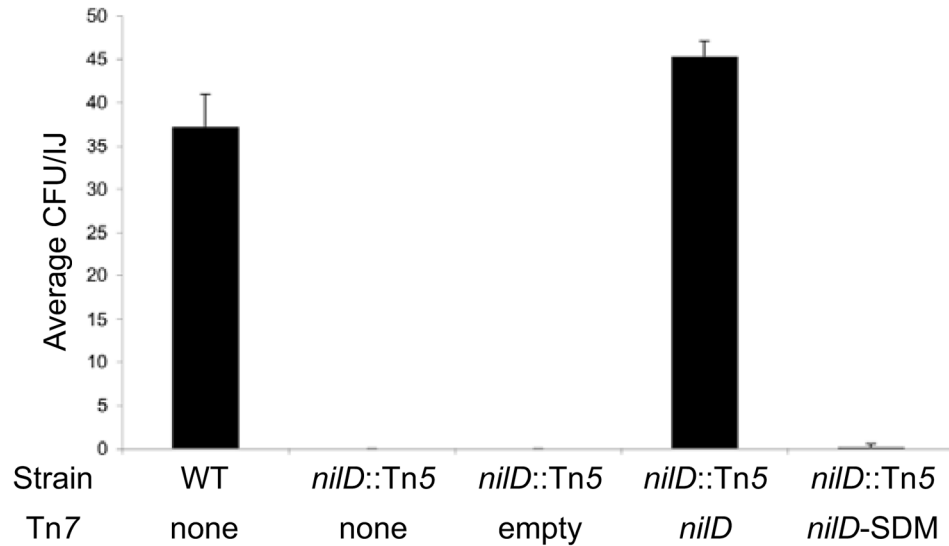
Spacer regions are shaded in gray. Lower case, bold nucleotides indicate those that differ from NiID RNA. The 5' and 3' handles as described by Brouns *et al.* (2009), and experimentally determined for NiID RNA are boxed. The underlined “U” in the niID spacer sequence is necessary for colonization (Heungens *et al.*, 2002).

D. Alignment of *niID* locus repeat-spacer-repeat sequences of *X. nematophila* (carpocapsae) (XnSc *niID*), *X. nematophila* (anatoliense) (XnSa *niID*), and *X. nematophila* (websteri) (XnSw *niID*). Lower case bold letters indicate nucleotides that differ among the three sequences.

A.

nilD-XnSc CCA-TGGCtACTtTgAAATTCcTtGTtTATAAcTCc-ATG
nilD-SDM CCA-TGGCaACaTTaAAaTTaCTaGTaTAAAAaTCa-ATG

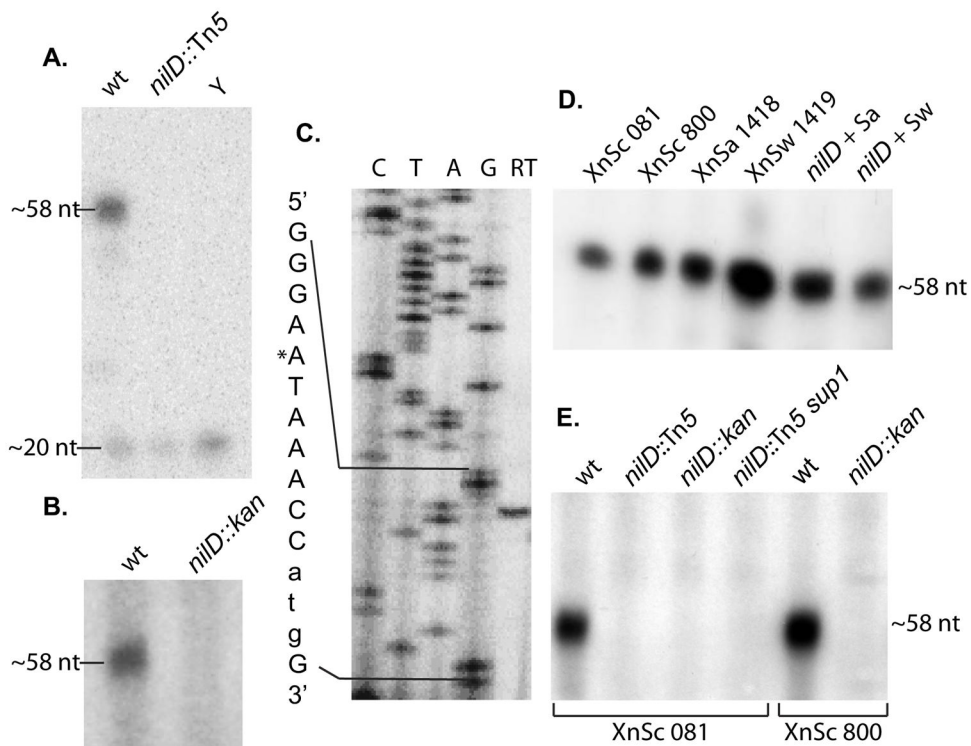
B.

**Figure 2.**

Wild type, but not mutant *nilD* provided in trans complements the colonization defect of the *nilD6*::Tn5 mutant

A. Alignment of the spacer sequences of *X. nematophila* (from *S. carpocapsae*) wild type *nilD* (*nilD*-XnSc) and the mutated allele (*nilD*-SDM).

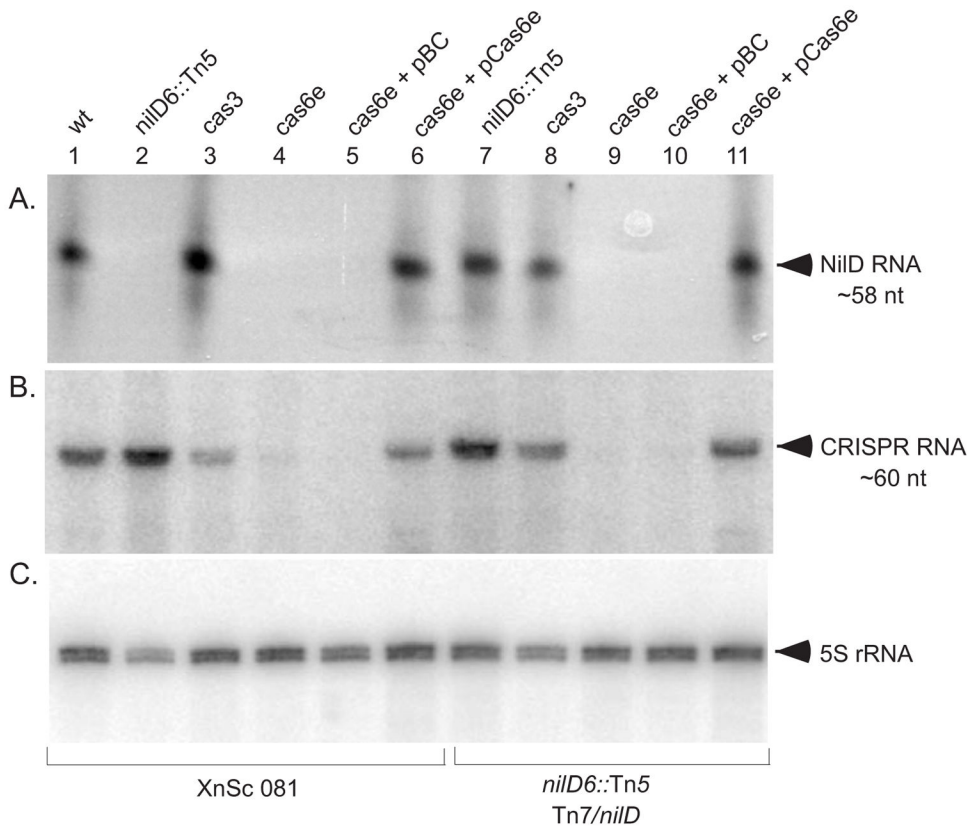
B. XnSc 081, XnSc 081 *nilD*::Tn5, and complemented strains were co-cultivated with axenic *S. carpocapsae* nematodes. The average colony forming units (CFU) colonizing the resulting progeny infective juveniles was measured by sonication and dilution plating.

**Figure 3.**

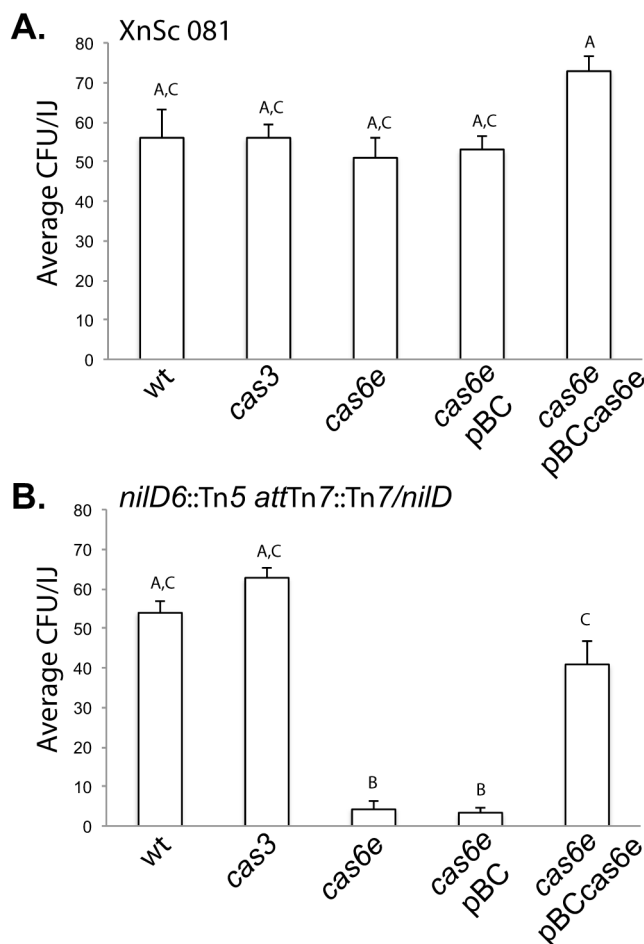
The *nilD* locus of *X. nematophila* encodes a small RNA

RNase protection analysis, using a radiolabeled NilD RNA-specific probe, was used to detect expression of the NilD RNA in RNA harvested from *X. nematophila* strains during both *in vitro* growth (panels **A**, **C**, and **E**) and mutualistic interactions with the nematode host (panel **B**). Primer extension analysis was performed on RNA isolated from laboratory grown *X. nematophila* cultures to map the 5' end of NilD RNA (panel **C**). **A**. RNA was harvested from stationary phase cultures of *X. nematophila* Sc 081, and *X. nematophila* Sc 081 *nilD6::Tn5*. The probe was also incubated with yeast RNA (Y) as a negative control. Radioactive markers (not shown) were used to estimate the sizes of the labeled fragments as indicated on the left. The 20-nt band represents the smallest nuclease-resistant single stranded RNA product of RNase cleavage. **B**. To assay for *in vivo* expression of NilD, RNA was extracted from Sc 081 and Sc 081 *nilD56::kan* harvested from infective-juvenile stage *S. carpocapsae* nematodes after co-cultivation. **C**. RNA isolated from wild type *X. nematophila* HGB800 (ATCC19061) after overnight growth in LB was used as a template for reverse transcriptase extension (RT) from the radioactively labeled primer AAP2. The resulting products were loaded on each gel adjacent to a sequencing ladder (C, T, A, and G lanes) derived from the same primer on pAWA1 template DNA. The relevant sequence is indicated to the left of the panel. The asterisk represents the starting nucleotide of the observed product. The lower case atg in the sequence to the left of the panel indicates the predicted start codon of *orf1*. **D**. To determine if the NilD RNA is expressed in distinct *X. nematophila* strains, RNA was harvested from the indicated strains: XnSc 081, XnSc 800 and strains harvested from *S. anatoliense* and *S. websteri* nematodes (XnSa 1418 and XnSw 1419, respectively). Likewise, RNA was harvested from *nilD6::Tn5* strains complemented

with the *nilD* locus derived from XnSa 1418 and XnSw 1419 (*nilD* + Sa and *nilD* + Sw, respectively). **E.** RNA was isolated from wild type *X. nematophila* strains (XnSc 081 and XnSc 800), *nilD* mutant strains (*nilD*::Tn5 or *nilD*::*kan*), and the *nilD6*::Tn5 suppressor strain (XnSc 081 *nilD*::Tn5 *sup1*). For each panel, each image was processed in its entirety with Adobe Photoshop CS3 to optimize visibility of bands by adjusting brightness and contrast.

**Figure 4.**

cas6e is necessary for presence of CRISPR RNA, but not for NiID RNA or colonization of *S. carpocapsae* nematodes. RNA was isolated from XnSc 081 wild type (wt), *nilD6::Tn5*, *cas-3::strep* (*cas3*), and *casE::strep* (*cas6e*) with or without the empty vector control (pBC) or the *cas6e* complement plasmid (pCas6e). All cells were grown to stationary phase in LB at 30°C. RNase protection assays using radiolabeled AAP2 primer (A) were used to detect NiID RNA while radiolabeled primers AAP1 (B) or Xn 5S RNA (C) were used in northern blots to detect CRISPR RNA and 5S RNA respectively. CRISPR RNA is detected as a band of ~60 nt. 5S RNA was detected as a band of ~113 nt.

**Figure 5.**

cas6e is necessary for colonization in the *nilD6::Tn5 attTn7::Tn7/nilD* background. Colonization ability was measured for (A) XnSc 081 and (B) *nilD6::Tn5 attTn7::Tn7/nilD* (the *nilD* mutant with wild type *nilD* expressed in trans at the *attTn7* site). Each strain carried wild type *cas* genes (wt) or *cas3* and *cas6e* mutations. In each background, the colonization phenotypes of the *cas6e* mutant carrying the control vector pBC or wild type *cas6e* (pBC*cas6e*) was also assessed. Each strain was co-cultivated with axenic *S. carpocapsae* nematodes and colony forming units (CFU) within the resulting progeny infective juveniles (IJ) was measured by sonication and dilution plating. The average CFU of strain/IJ \pm standard error ($n = 5$) is shown. In this experiment the uncomplemented *nilD6::Tn5* strain colonized at 0.22 ± 0.11 CFU/IJ. Different letters indicate significant differences in colonization levels between bacterial strains: $p < 0.0001$ One-way ANOVA with Tukey's post-test.

Table 1

Strains and plasmids used in this study

Strain or plasmid	Relevant Characteristics	Source or Reference
<i>E. coli</i>		
DH5α	General cloning host	(Sambrook <i>et al.</i> , 1989)
DH5α (λ pir)	General cloning strain for maintenance of oriR6K plasmids	
S17-1 (λ pir)	<i>E. coli</i> donor strain for conjugations	
<i>X. nematophila</i>		
HGB081 (XnSc 081)	Rifampicin resistant derivative of wild-type <i>X. nematophila</i> AN6/1 (carpocapsae)	S. Forst
HGB151	<i>X. nematophila</i> ATCC 19061 <i>rpoS1::kan</i>	(Vivas and Goodrich- Blair, 2001)
HGB315	HGB081 <i>nilD6::Tn5</i>	(Heungens <i>et al.</i> , 2002)
HGB829	HGB315 <i>nilD6::Tn5</i> pECM20- <i>gfp</i>	(Martens <i>et al.</i> , 2003)
HGB1186	HGB315 <i>nilD6::Tn5</i> pECM20- <i>gfp sup-1</i>	This study
HGB1578	HGB081 <i>cas3-3::kan</i>	This study
HGB1695	HGB081 <i>casE4::kan</i> (<i>cas6e</i> mutant)	This study
HGB1418 (XnSa 1418)	<i>X. nematophila</i> (anatoliense) isolated from <i>S. anatoliense</i> nematodes	This study
HGB1419 (XnSw 1419)	<i>X. nematophila</i> (websteri) isolated from <i>S. websteri</i> nematodes	This study
HGB1421	<i>X. nematophila</i> strain of unknown origin	S. P. Stock
HGB007 (XnSc 007)	Wild-type <i>X. nematophila</i> (carpocapsae) ATCC 19061, acquired in 1995	ATCC
HGB800 (XnSc 800)	Wild-type <i>X. nematophila</i> (carpocapsae) ATCC 19061, acquired in 2003	ATCC
HGB1764	XnSc 081 <i>nilD56::kan</i>	This study
HGB1756	XnSc 800 <i>nilD56::kan</i>	This study
HGB1940	HGB315 <i>nilD6::Tn5 attTn7::Tn7/nilD</i>	This study
HGB1986	HGB315 <i>nilD6::Tn5 attTn7::Tn7/nilD-SDM</i>	This study
HGB1901	XnSc 081 <i>cas3-5::strep</i>	This study
HGB1909	XnSc 081 <i>casE6::strep</i> (<i>cas6e</i> mutant)	This study
HGB1907	HGB1940 <i>cas3-5::strep</i>	This study
HGB1915	HGB1940 <i>casE6::strep</i> (<i>cas6e</i> mutant)	This study
<i>X. bovienii</i>		
HGB1166	ATCC 35271 <i>X. bovienii attTn7::miniTn7</i>	
HGB1167	ATCC 35271 <i>X. bovienii attTn7::miniTn7/SR1</i> (containing <i>nilA</i> , <i>nilB</i> , and <i>nilC</i>)	(Cowles and Goodrich- Blair, 2004)
HGB1649	HGB1166 pECMXb-Empty	This study
HGB1651	HGB1166 pECMXb-SR2; <i>nilD</i> +	This study
HGB1653	HGB1167 pECMXb-Empty; <i>nilABC</i> +	This study
HGB1655	HGB1167 pECMXb-SR2; <i>nilD</i> + <i>nilABC</i> +	This study
Plasmids		
pBCSK+	General cloning vector, Cm ^R ,	Stratagene
pCR2.1 [®] -TOPO	General cloning vector, Amp ^R , Kan ^R	Invitrogen, Carlsbad, CA
pCR2.1-TOPOmini	General cloning vector, Amp ^R	This study
pTopoSR2-2	312-bp XnSc 007 <i>nilD</i> region amplified with KPH62 and KPH63 primers and cloned into pCR2.1 [®] -TOPO	(Heungens <i>et al.</i> , 2002)

Strain or plasmid	Relevant Characteristics	Source or Reference
pSR2-312	ApaI-SacI fragment from pTopoSR2-2 subcloned into pBCSK+, formerly named pBCSR2-2	(Heungens <i>et al.</i> , 2002)
pAWA1	137-bp XnSc 007 <i>nilD</i> region PCR amplified from HGB007 chromosomal DNA with primers KPH57 and KPH58 and cloned into pCR®II-TOPO	This study
pCR2.1-TOPO- <i>nilD</i> - <i>XnSa</i>	pCR2.1-TOPO- <i>nilD</i> modified by site directed mutagenesis to match the <i>nilD</i> spacer sequence of XnSa and XnSw. Use for RPA analysis.	This study
pEVS107	Source of Kan ^r cassette for <i>cas3</i> and <i>cas6e</i> mutations	(McCann <i>et al.</i> , 2003)
pEVS107- <i>nilD</i>	Kan ^R ; vector for insertion of 387-bp <i>nilD</i> fragment at <i>attTn7</i> site of XnSc 081	This study
pEVS107- <i>nilD</i> -SDM	Kan ^R ; pEVS107- <i>nilD</i> altered by site directed mutagenesis to change NilD RNA spacer sequence	This study
pKNG101	Sm ^R ; <i>oriR6K</i> suicide vector	(Kaniga <i>et al.</i> , 1991)
pECM20	pECM2 containing a 614-bp chromosomal insert from XnSc 007	(Martens <i>et al.</i> , 2003)
pECMXb-Empty	pECM20 with <i>X. bovienii</i> sequence replacing the <i>X. nematophila</i> insertion sequence	This study
pECMXb-SR2	pECMXb-Empty with 312bp <i>nilD</i> region from pTopoSR2-2 in the XbaI site	This study
pKNG <i>cas3-5::strep</i>	Sm ^R ; pKNG101 with <i>cas3::strep</i> for replacing <i>cas3</i> gene with Sm ^R cassette	This study
pKNG <i>casE6::strep</i>	Sm ^R ; pKNG101 with <i>casE6::strep</i> for replacing <i>cas6e</i> gene with Sm ^R cassette	This study
pKR100	Cm ^R ; <i>oriR6K</i> suicide vector	
pKR100 <i>nilD56::kan</i>	Kan ^R , Cm ^R ; pKNG101 with <i>cas-3::kan</i> for replacing <i>nilD</i> encoding region with Kan ^R cassette.	This study
pBS-5S	Amp ^R	(Trotochaud and Wassarman, 2005)

Table 2

A *nilD::kan* mutation does not confer a colonization defect in the HGB081 and HGB800 backgrounds.

Relevant Alleles	Average CFU of strain/IJ \pm standard error (n = 3)^a	
	XnSc 081	HGB800
none	66.36 \pm 2.57	50.03 \pm 4.02
<i>nilD::kan</i>	60.03 \pm 3.86	61.73 \pm 0.96
<i>nilD::kan cas3::str</i>	60.52 \pm 5.97	60.05 \pm 3.17
<i>nilD::kan cas6e::str</i>	58.83 \pm 2.76	58.72 \pm 2.80

^aIn this experiment the *nilD6::Tn5* strain colonized at 0.10 ± 0.01 CFU/IJ. None of the values shown in the table were significantly different from each other, but all were significantly different from the *nilD6::Tn5* strain ($p < 0.001$ using one-way ANOVA with Tukey's post-test).

Table 3

nilD is required for *X. nematophila* colonization of *S. anatoliense* and *S. websteri* nematodes.

Bacterial Strain	<i>attTn7</i> locus insertion ^a	Average CFU/IJ ± standard deviation (n=3)		
		<i>S. carpocapsae</i>	<i>S. anatoliense</i>	<i>S. websteri</i>
XnSc 081	none	42.07 ± 11.01 ^A	34.40 ± 4.12 ^A	26.31 ± 6.20 ^A
XnSc 081 <i>nilD6::Tn5</i>	none	0.10 ± 0.04 ^B	0.36 ± 0.06 ^B	0.12 ± 0.15 ^B
XnSc 081 <i>nilD6::Tn5</i>	<i>nilD-Sc</i>	54.40 ± 7.95 ^A	38.60 ± 4.77 ^{A,C}	44.10 ± 7.03 ^{A,C}
XnSc 081 <i>nilD6::Tn5</i>	<i>nilD-Sa</i>	1.09 ± 0.24 ^B	1.08 ± 0.40 ^B	0.45 ± 0.23 ^B
XnSc 081 <i>nilD6::Tn5</i>	<i>nilD-Sw</i>	0.03 ± 0.02 ^B	0.25 ± 0.17 ^B	0.02 ± 0.14 ^B
XnSa	none	77.90 ± 16.58 ^{A,C}	59.40 ± 4.13 ^C	55.50 ± 16.61 ^{C,D}
XnSw	none	63.55 ± 12.97 ^{A,C}	28.45 ± 6.66 ^A	74.57 ± 4.74 ^D

^a A Tn7 transposon carrying *nilD* loci from XnSc (*nilD-Sc*), XnSa (*nilD-Sa*), or XnSw (*nilD-Sw*) was integrated at the *attTn7* locus. Different letters indicate significant differences between bacterial strains for colonization within each nematode species: p<0.05 using one-way repeated measures ANOVA (Prism v2.0, GraphPad, La Jolla, CA) with Tukey's post-test. Colonization levels achieved by each bacterial strain in the three nematode species were not significantly different except XnSw for which colonization of *S. anatoliense* nematodes was significantly lower than those of the other two nematode species (not shown).