

Two classes of silencing RNAs move between *Caenorhabditis elegans* tissues

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Organism-wide RNA interference (RNAi) is due to the transport of mobile silencing RNA throughout the organism, but the identities of these mobile RNA species in animals are unknown. Here, we present genetic evidence that both the initial double-stranded RNA (dsRNA), which triggers RNAi, and at least one dsRNA intermediate produced during RNAi can act as or generate mobile silencing RNA in *C. elegans*. This dsRNA intermediate requires the long dsRNA-binding protein RDE-4, the endonuclease DCR-1, which cleaves long dsRNA into double-stranded short-interfering RNA (ds-siRNA), and the putative nucleotidyltransferase MUT-2 (RDE-3). However, single-stranded siRNA and downstream secondary siRNA produced upon amplification by the RNA-dependent RNA polymerase RRF-1 do not generate mobile silencing RNA. Restricting intertissue transport to long dsRNA and directly processed siRNA intermediates rather than amplified siRNA may serve to modulate the extent of systemic silencing in proportion to available dsRNA.

Intercellular transport of RNA has been inferred in plants and animals undergoing gene silencing by RNAi (ref. 1). In plants, siRNA processed from long dsRNA move between cells through intercellular bridges called plasmodesmata and travel long distances through the phloem to convey gene-specific silencing information^{2–4}. Although the nature of mobile silencing signals in animals is unknown, the conserved RNA transporter SID-1 is required for import of these signals in *C. elegans* and has been implicated in RNA transport in other animals^{5–7}. In addition, dsRNA expressed in multiple tissues can generate *sid-1*-dependent mobile silencing RNA through a pathway that is as yet unknown⁸. Because animals transcribe dsRNA from numerous loci⁹, understanding how mobile RNA is produced from dsRNA has broad implications for systemic control of gene expression.

Multiple distinct RNA species are produced during RNAi in *C. elegans*, but it is unclear which of these are mobile (Fig. 1a)^{10–13}. These RNA species include transcribed sense and antisense duplexes (dsRNA), ds-siRNA generated upon cleavage of long dsRNA by the RDE-4–Dicer (DCR-1) complex, primary single-stranded siRNA generated upon cleavage of ds-siRNA by the Argonaute RDE-1 (ref. 13), and the subsequent numerous secondary siRNAs generated by RNA-dependent RNA polymerases (RdRP) that are responsible for potent silencing of the target gene. In addition, enzymes that can modify RNA, such as the putative nucleotidyltransferase MUT-2 (refs. 14–16), which is required for efficient RNAi (Supplementary Fig. 1), may also generate RNA species that act as mobile RNA. Early studies using dsRNA injected into the cytoplasm of gut cells suggested that RNA silencing in gut cells is not required to transport a mobile silencing signal to the germline^{17,18}. However, whether this signal is the injected exogenous dsRNA itself or a dsRNA-derived mobile RNA

or both is unclear, and how endogenously transcribed dsRNA leads to the production of mobile RNA is unknown.

Here, we determine the genetic requirements for silencing due to mobile RNAs using well-characterized promoters to restrict the expression of dsRNA or RNAi pathway genes to specific tissues, and we examine target gene silencing in other tissues. In most experimental systems that use similar approaches, it is difficult to control for low levels of misexpression in the target tissues. Because SID-1 is strictly required for the import of mobile silencing RNAs⁸, the SID-1 dependence of silencing serves to clearly distinguish silencing due to mobile RNA from silencing due to misexpression in the target tissues.

RESULTS

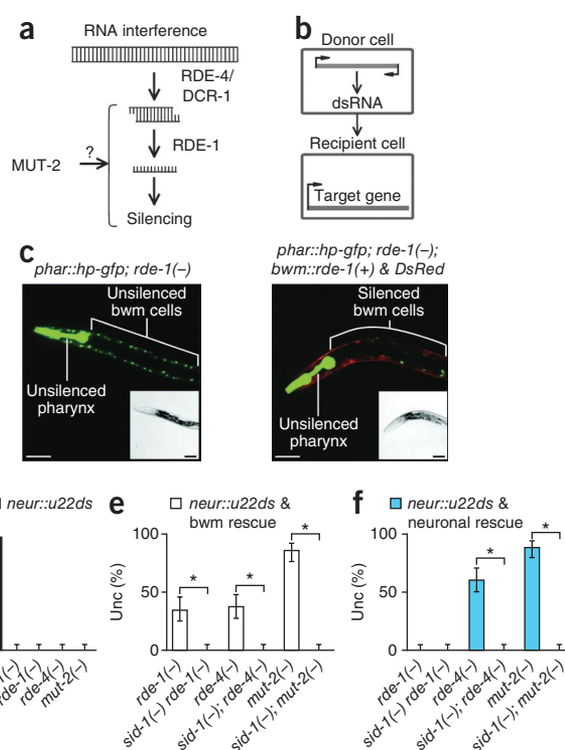
Long dsRNA is mobile in *C. elegans*

We examined how endogenously transcribed dsRNA produces mobile silencing RNA using mosaic animals (animals that have some mutant cells and some wild-type cells) in which a mutant donor tissue expresses dsRNA that targets a gene in a wild-type recipient tissue (Fig. 1b). To determine whether the activity of the primary Argonaute RDE-1 is required to produce a mobile silencing signal, we first expressed dsRNA targeting the green fluorescent protein gene (*gfp*) in the pharynx of *rde-1(-)* animals. We then coexpressed *gfp* and *rde-1(+)* in the body-wall muscle (bwm) cells, making bwm a wild-type recipient tissue (Fig. 1c and Supplementary Fig. 2). A worm or tissue that has a wild-type copy of *rde-1* is indicated as *rde-1(+)*, and a worm or tissue that lacks *rde-1* is indicated as *rde-1(-)*. We observed *gfp* silencing in anterior *rde-1(+)* bwm cells. Thus, RNAi-mediated silencing in the pharynx is not required to produce and transport mobile RNA to the bwm cells. To determine whether RNAi

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Figure 1 RNAi-independent biogenesis of mobile RNA from expressed dsRNA. **(a)** Schematic of RNAi within a cell (see text for details). **(b)** Schematic of assay to measure silencing due to mobile RNA. dsRNA that targets a gene in recipient cells are expressed only in other donor cells. **(c)** A representative *rde-1(-)* L4 animal that expresses *gfp-hairpin* RNA only in the pharynx (*phar::hp-gfp*) but *gfp* in pharynx and bwm cells (left panel) and one that in addition coexpresses *rde-1(+)* and DsRed only in bwm cells (right panel). Silencing was observed in 100% of animals from three independent *bwm::rde-1(+)* & *DsRed* lines, as above. Scale bar, 50 μ m. **(d-f)** A representative transgenic line that expresses *unc-22* dsRNA under the control of the neuronal *rgef-1* promoter (*neur::u22ds*) was generated in wild-type animals and crossed into the genetic backgrounds indicated. The uncoordinated twitching (Unc) due to the silencing of *unc-22* was measured in these animals (black) and in animals that in addition had the corresponding RNAi gene rescued in body-wall muscles (white) or in neurons (blue). $n = 100$ animals, error bars indicate 95% confidence intervals, and asterisks indicate significant differences ($P < 0.05$). Partial silencing in rescued transgenic lines probably indicate that levels of the rescuing genes are inadequate for complete silencing in response to the low levels of neuronal *unc-22* dsRNA. Consistently, feeding *unc-22* dsRNA to animals with the same muscle-rescued transgenic lines above results in complete silencing (**Fig. 3**) (see **Supplementary Fig. 2** for details of constructs used).



pathway genes upstream of RDE-1 are required to produce a mobile silencing signal from expressed dsRNA, we developed a sensitive assay that measures silencing of an endogenous gene due to mobile RNA (**Fig. 1d** and **Supplementary Fig. 2**). Specifically, we introduced a neuronally expressed transgene that produces a dsRNA of ~560 base pairs (bp) that targets the muscle gene *unc-22* (*neur::u22ds*). All *unc-22* silencing detected in animals with the *neur::u22ds* transgene required the RNAi pathway genes and the RNA transporter SID-1, showing that all silencing occurred through RNAi in these animals and was due to mobile RNA-enabled RNAi (**Fig. 1d**). Using this source of mobile RNA, we detected *unc-22* silencing in *rde-4(-)* animals that expressed *rde-4(+)* in bwm cells and in *mut-2(-)* animals that expressed *mut-2(+)* in bwm cells (**Fig. 1e**). Thus, neither dsRNA cleavage through RDE-4 recruitment of Dicer nor modification by the nucleotidyltransferase MUT-2 is required in neurons that express dsRNA for the generation and export of mobile RNA. Together, these results show that an RNA species generated from transcribed long dsRNA, without processing by the canonical RNAi pathway, can act as mobile silencing RNA.

A processed dsRNA also moves between cells

To determine whether products of dsRNA processing by the canonical RNAi pathway are also mobile, we expressed dsRNA in a wild-type RNAi-proficient donor tissue and examined silencing in RNAi-defective recipient tissues. If a processed RNA produced in the wild-type donor tissue can act as or generate a mobile silencing RNA, that RNA may bypass the requirement for the earlier-acting RNAi pathway gene in the recipient tissue and cause silencing. Note that by using this approach, we cannot infer anything about RNAs that move between tissues but fail to cause gene silencing.

To detect silencing triggered by mobile processed RNAs, we rescued RNAi pathway mutants only in neurons of animals that contain the *neur::u22ds* transgene and measured silencing of the target gene *unc-22* in mutant muscle cells. We detected *unc-22* silencing in *rde-4(-)* animals that expressed *rde-4(+)* in neurons. Consistent with silencing due to mobile RNAs, SID-1 was required for the observed silencing (**Fig. 1f**). Because RDE-4 is required for DCR-1 cleavage of long dsRNA into ds-siRNA¹⁹, these mobile RNAs are either ds-siRNA or downstream RNAi products. To distinguish between these two possibilities, we used similar procedures to examine the role of the primary

Argonaute RDE-1 in the production of mobile RNA. In contrast to the analogous experiment with RDE-4, we observed no detectable *unc-22* silencing in *rde-1(-)* animals that express *rde-1(+)* in neurons. This observation suggests that primary siRNA and downstream RNAi products such as RdRP-dependent secondary siRNA are not mobile. Finally, we detected *unc-22* silencing in *mut-2(-)* animals that expressed *mut-2(+)* in neurons (**Fig. 1f**), and this silencing was due to mobile RNA because it required SID-1 (**Fig. 1f**). Therefore, we infer that similarly to RDE-4, MUT-2 functions upstream of RDE-1 to generate a species of mobile RNA that can bypass the need for MUT-2 activity in the recipient tissue.

Mobile RNAs are similarly made from other sources of dsRNA

We next tested whether other sources of silencing RNAs also rely on the same genes to produce mobile RNAs. Multicopy transgenes such as *sur-5::gfp* (which express nuclear-localized GFP in all somatic tissues) can generate mobile RNAs, presumably from trace amounts of dsRNA produced from the transgene^{8,20}. We therefore generated *rde-4(-); sur-5::gfp* animals and moved a representative transgene that expresses *rde-4* in bwm cells (*bwm::rde-4(+)*) into these animals. Significant silencing ($P < 0.05$) was detected in tissues other than muscle in the resultant mosaic animals and was most easily observed in the prominent gut nuclei (**Fig. 2a,b**). Consistent with silencing due to mobile RNAs, SID-1 was required for the observed silencing of gut nuclei (**Fig. 2c,d**). However, when we moved a representative transgene that expressed *rde-1(+)* in bwm cells (*bwm::rde-1(+)*) into *rde-1(-); sur-5::gfp* animals, we observed no detectable silencing of GFP expression in the gut. By contrast, moving a representative transgene that expressed *mut-2(+)* in bwm cells (*bwm::mut-2(+)*) into *mut-2(-); sur-5::gfp* animals resulted in the silencing of GFP expression in the gut (**Fig. 2d**). The observed silencing was dependent on SID-1, showing that mobile RNA triggered the silencing in *mut-2(-)* gut cells (**Fig. 2d**). Therefore, as in the case of expressed dsRNA, multicopy transgenes also generate mobile RNAs that are upstream of RDE-1 and include those that are processed by RDE-4 and MUT-2.

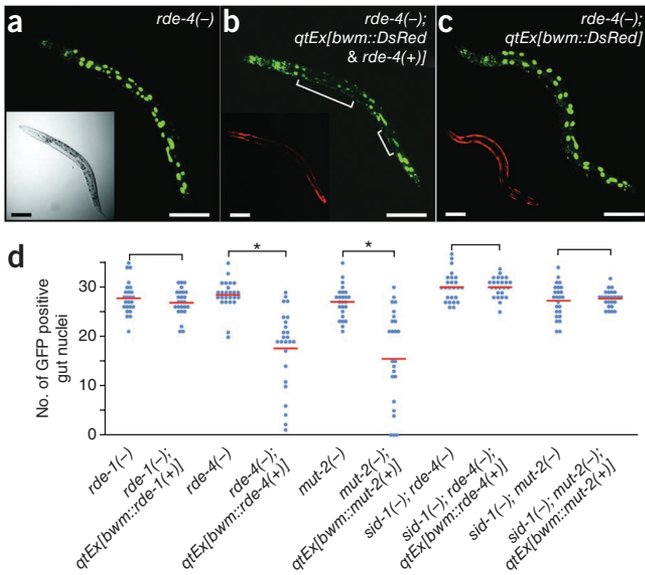


Figure 2 Restricted expression of RDE-4 and MUT-2 but not RDE-1 enables non-cell-autonomous RNA silencing. (a–c) Representative animals that express nuclear-localized GFP in all cells (*sur-5::gfp*). (a) *rde-4(-)*. (b) *rde-4(-)* animals with *rde-4(+)* and *DsRed* expressed in bwm cells (*qtEx[bwm::rde-4(+)] & DsRed*). (c) *rde-4(-)* animals that only express *DsRed* in bwm cells (*qtEx[bwm::DsRed]*). Square brackets indicate silencing in gut nuclei. Insets are wide-field (a) or red channel (b, c) images. Scale bars, 50 μ m. Note that unlike overexpression of *rde-4(+)*, overexpression of the coinjection marker *DsRed* did not result in any silencing of *gfp* expression (compare b and c). When expressed with *rde-4(+)*, *DsRed* expression was lower (enhanced in b (inset) to clearly indicate expression in the bwm), which likely reflects enhanced silencing of the *DsRed* transgene. (d) The number of brightly fluorescent gut nuclei that show *sur-5::gfp* expression were counted in *rde-1(-)*, *rde-4(-)*, and *mut-2(-)* mutant backgrounds as well as in mutant animals with corresponding representative bwm rescue transgenes from **Figure 1e**. Similar experiments done with *rde-4(-)*, *sid-1(-)* and *mut-2(-)*, *sid-1(-)* double mutant backgrounds are also shown. $n = 25$ L4 animals. Averages (red bars), significant differences (square brackets and *, $P < 0.05$) and similar values (square brackets) are indicated. Minor variations in the average number of nuclei (± 2 nuclei) observed between animals were not due to silencing of *gfp* expression but rather due to small changes in the number of intestinal nuclei (see **Supplementary Fig. 4** and the discussion therein).

We next tested whether RDE-4 and MUT-2 but not RDE-1 can similarly process exogenously supplied dsRNA to produce mobile RNA. We fed bacteria that express *gfp*-dsRNA (feeding RNAi)²¹ to the above *rde-1(-)*, *rde-4(-)*, and *mut-2(-)* mutants that contain the *sur-5::gfp* transgene and are rescued in muscle cells, and we examined silencing in the respective mutant gut cells. Consistent with our results with endogenously transcribed dsRNA, we found that *gfp* feeding RNAi increased silencing of GFP in the non-muscle cells of muscle-rescued *rde-4(-)* and *mut-2(-)* animals but not of muscle-rescued *rde-1(-)* animals (**Fig. 3a**). To assay silencing due to feeding RNAi targeting endogenous genes, we removed the *sur-5::gfp* transgenes from the transgenic bwm rescue lines and then fed these muscle-rescued animals bacteria that express dsRNA targeting the muscle gene *unc-22*, or that express dsRNA targeting the skin gene *dpy-7* or that express dsRNA targeting the intestinal gene *act-5*. Results from silencing due to feeding RNAi of these endogenous genes were consistent with our results using *gfp* feeding RNAi and using endogenously transcribed dsRNA. Specifically, although we observed robust silencing of the muscle gene in all three strains of muscle-rescued animals, silencing of the skin and intestinal genes was detectable in muscle-rescued *rde-4(-)* and *mut-2(-)* animals (**Fig. 3b**) but not in muscle-rescued *rde-1(-)* animals (**Fig. 3b**, ref. 22). Thus, the silencing observed in these *rde-4* and *mut-2* mosaic animals by feeding RNAi is probably due to import of ingested long dsRNA into the rescued muscle cells, followed by export of a processed mobile RNA that can silence the target genes in *rde-4(-)* and *mut-2(-)* cells.

Figure 3 RDE-4- and MUT-2- but not RDE-1-processed ingested dsRNA is mobile. (a) Feeding RNAi of *rde-1(-)*, *rde-4(-)* and *mut-2(-)* animals with *sur-5::gfp* and their corresponding representative bwm rescue transgenic lines used in **Figure 2d**. The number of brightly fluorescent gut nuclei that show *sur-5::gfp* expression were counted in L4 animals that were fed either control bacteria (brown, replotted from **Fig. 2d**) or bacteria expressing *gfp*-dsRNA (blue). $n = 25$ animals. Averages (red bars), significant differences (square brackets and *, $P < 0.05$) and similar values (square bracket) are indicated. (b) Feeding RNAi of strains in a after removal of *sur-5::gfp*. L4 animals were fed L4440 (control) or dsRNA targeting the muscle gene *unc-22* (bwm) or the skin gene *dpy-7* (skin) or the gut gene *act-5* (gut), and the percentage of L4 progeny that showed the corresponding defects were determined. $n = 100$ L4 animals; error bars indicate 95% confidence intervals.

Therefore, both multicopy transgenes and ingested dsRNA use the same genetic pathway to produce short mobile silencing RNA.

Two classes of upstream dsRNAs are mobile RNAs

Taken together, our results suggest a model in which upstream dsRNA species such as long dsRNA and ds-siRNA act as or generate mobile RNA, whereas all silencing RNAs produced after cleavage of ds-siRNA by RDE-1 cannot cause silencing in *rde-1(-)* cells (**Fig. 4a**). Because Dicer can cleave long dsRNA in the absence of MUT-2 (ref. 12) and because MUT-2 acts upstream of RDE-1 to generate mobile RNA, one possible role for MUT-2 in RNAi is to modify ds-siRNA. Despite MUT-2 having the required catalytic residues, a systematic test of putative nucleotidyltransferases using *in vitro* assays failed to reveal how MUT-2 might modify RNA²³. Nevertheless, consistent with our model, neither overexpression of *mut-2(+)* nor of *rde-4(+)* in bwm cells of *rde-1(-)*; *sur-5::gfp* animals resulted in detectable silencing (**Supplementary Table 1**). Further, neither overexpression of *mut-2(+)* in bwm cells of *rde-4(-)*; *sur-5::gfp* animals nor overexpression of *rde-4(+)* in the bwm cells of *mut-2(-)*; *sur-5::gfp* animals resulted in detectable silencing (**Supplementary Table 1**), suggesting that RDE-4 and MUT-2 act in the same pathway to generate mobile RNA.

The following results provide additional support for a model in which long and short dsRNAs, but not single-stranded siRNA, act as mobile silencing RNA: (i) The RdRP RRF-1, which makes the numerous downstream secondary siRNAs, was not required for the generation of mobile RNAs (**Fig. 4b**). (ii) Rescuing a partial loss-of-function

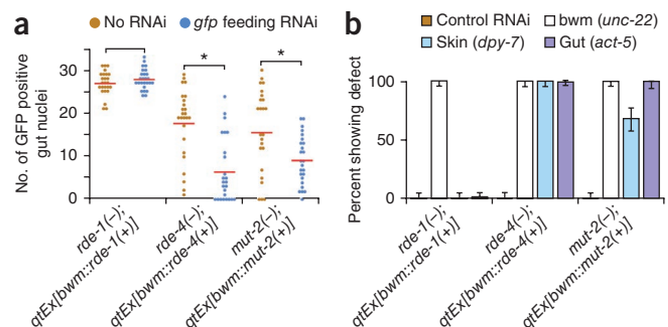
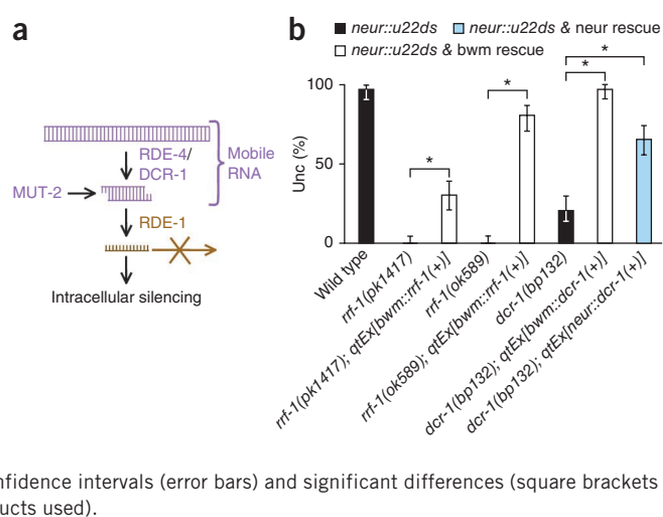


Figure 4 Biogenesis of mobile RNA in *C. elegans*. (a) Schematic of the biogenesis pathway. Double-stranded forms of RNA produced during the early steps of RNA interference act as or generate mobile RNAs. RNAs produced after the Argonaute RDE-1 cleaves ds-siRNA to release single stranded RNA¹³ are restricted to intracellular silencing. MUT-2 expression enables the generation and export of mobile RNA possibly through enzymatic modification of dsRNA. Because expressed as well as ingested dsRNA generate mobile RNA, additional regulation in response to the environment and selection of specific endogenous loci to make mobile RNAs is likely. (b) RdRP activity is not required for mobile RNA production, and dsRNAs both upstream and downstream of Dicer generate mobile RNAs. The representative transgene used in **Figure 1** to express *unc-22*-dsRNA under the control of the neuronal *rgef-1* promoter (*neur::u22ds*) was crossed into the genetic backgrounds indicated. Two deletion alleles of *rrf-1* (*pk1417* and *ok589*) were rescued with *rrf-1(+)* in *bwm*, and the missense allele *dcr-1(bp132)* was rescued with *dcr-1(+)* in *bwm* and in neurons. Silencing of *unc-22* was measured (% Unc) in the mutant animals (black) and in animals with the corresponding RNAi gene rescued in *bwm* (white) and in neurons (blue). $n = 100$ animals. The 95% confidence intervals (error bars) and significant differences (square brackets and *, $P < 0.05$) are indicated (see **Supplementary Fig. 2** for details of constructs used).



dcr-1 mutant²⁴ in the recipient tissue (which increases processing of imported Dicer substrates (long dsRNA)) improved silencing (**Fig. 4b**). (iii) Rescuing the *dcr-1* mutant in donor tissues also increased silencing in recipient cells, presumably by increased transport of ds-siRNA (**Fig. 4b**). (iv) Expression of inhibitors of RNAi in recipient cells, including the conserved exonuclease ERI-1 that can degrade ds-siRNA²⁵, inhibited silencing (**Supplementary Fig. 3**).

DISCUSSION

We provide evidence for the existence of at least two distinct species of mobile RNA in *C. elegans*: one that is produced from long dsRNA independent of RNAi genes in donor tissues but requires all tested RNAi genes in recipient tissues for silencing and one that is produced using RDE-4, DCR-1 and MUT-2 in donor tissues but does not require these proteins in recipient tissues for silencing.

Animal mobile silencing RNAs differ from plant mobile RNAs

In plants, mobile RNAs move between cells through relatively non-selective intercellular bridges called plasmodesmata^{2,3}. In the plant *Arabidopsis thaliana*, grafting experiments between genetically distinct source and target tissues have enabled the molecular identification of mobile RNAs. These studies identified both single-stranded siRNA and ds-siRNA whose movement to distant tissues correlates with mobile RNAs^{2,3}. In addition, accumulating evidence supports the intercellular movement of microRNAs, tasiRNAs and mRNAs²⁶.

Our results indicate that in *C. elegans*, long dsRNA and a form of ds-siRNA can move between cells (**Fig. 4a**). Unexpectedly, and in contrast to what is observed in plants, single-stranded siRNAs produced by RdRP amplification are either not mobile, or if mobile, are incapable of causing detectable silencing in recipient cells. Consistent with mobile silencing signals being restricted to double-stranded forms of RNA, most systemic RNAi silencing observed in *C. elegans* is dependent on SID-1 (refs. 5,8), which is extremely selective for dsRNA²⁷. This restriction couples the extent of RNAi spreading to the amount of primary dsRNA produced within cells or imported from the environment.

A conserved pathway to make animal mobile RNAs

Because a mammalian SID-1 homolog can transport ds-siRNAs into mammalian cells⁷, ds-siRNA, perhaps modified by a nucleotidyltransferase, may move between mammalian cells. Notably, because short dsRNAs can escape the interferon response that results in nonspecific effects in differentiated mammalian cells²⁸, their transport between

differentiated tissues should be tolerated. By contrast, the transport of long dsRNA would result in specific gene silencing only in undifferentiated mammalian cells. Furthermore, the proteins required to make short mobile RNA in worms are found in most animals: dsRNA-binding proteins, such as RDE-4, that act with Dicer (for example, PACT and TRBP with human Dicer²⁹) and β -nucleotidyltransferases, such as MUT-2, that play a role in RNA silencing^{14–16}. Modulation of such conserved biochemical pathways may contribute to the tissue- and environment-dependent differences in silencing due to mobile RNA that are observed in *C. elegans*^{8,30}. Regulated transport of mobile RNA is evident in plants, where mobile RNA produced in metabolic source tissues control gene expression in distant metabolic sink tissues³¹. Similarly, *C. elegans* mobile RNAs are preferentially imported into cells that express SID-1 at high levels^{8,32}, suggesting that SID-1 expression produces a sink for mobile RNA. Therefore, short dsRNAs produced from endogenous loci in a mammalian cell may control gene expression in another cell type that expresses a SID-1 homolog.

METHODS

Methods and any associated references are available in the online version of the paper at <http://www.nature.com/nsmb/>.

Note: Supplementary information is available on the Nature Structural & Molecular Biology website.

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AUTHOR CONTRIBUTIONS

A.M.J. conducted the experiments and G.A.G. generated most of the DNA constructs; A.M.J. and C.P.H. designed the study, analyzed the data and wrote the paper. All authors discussed the results and commented on the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Strains used. N2 wild-type, HC196 *sid-1(qt9)*, WM30 *mut-2* or *rde-3(ne298)*, WM27 *rde-1(ne219)*, WM49 *rde-4(ne301)*, NL2099 *rrf-3(pk1426)*, GR1373 *eri-1(mg366)*, HC70 *rde-1(ne219)*; *mIs11[Pmyo-2::gfp]*; *cIs4251[Pmyo-3::gfp]*; *qtIs3[Pmyo-2::hp-gfp]*, PD4792 *mIs11[myo-2::GFP, gut::GFP, pes-10::GFP]*, HC195 *nrIs20[sur-5::gfp]*, HC731 *sid-1(qt9)*; *eri-1(mg366)*, HC732 *sid-1(qt9)*; *rrf-3(pk1426)*, HC733 *mut-2(ne298)*; *sid-1(qt9)*; *nrIs20*, HC734 *sid-1(qt9)*; *rde-4(ne301)*; *nrIs20*, HC735 *mut-2(ne298)*; *mIs11*, HC736 *qtEx136[Prgef-1(F25B3.3)::unc-22sense; Prgef-1::unc-22antisense; Prgef-1::DsRed line 8]*, HC737 *rde-4(ne301)*; *nrIs20*, HC738 *rde-1(ne219)*; *nrIs20*, HC739 *mut-2(ne298)*; *nrIs20*, HZ202 *dcr-1(bp132)*; *wIs51[scm-1::GFP]²⁴*, RB798 *rrf-1(ok589)*, NL2098 *rrf-1(pk1417)*, HC779 *dcr-1(bp132)* (outcrossed with N2 twice), HC780 *rrf-1(ok589)* (outcrossed with N2 twice), HC781 *rrf-1(pk1417)* (outcrossed with N2 twice), HC782 *sid-1(qt9) rde-1(ne219)*, HC783 *sid-1(qt9)*; *rde-4(ne301)*, HC784 *sid-1(qt9)*; *mut-2(ne298)*.

Strain constructions and analyses of transgenics. Double mutants were made using standard genetic approaches and were verified by genotyping using DNA sequencing or PCR analysis. Additional strains were constructed by crossing representative transgenes into various genetic backgrounds. These include strains generated by (i) crossing HC736 into either single mutants (WM27, HC196, WM27, WM49, NL2099, GR1373, HC779, HC780, HC781) or double mutants (HC731, HC732, HC782, HC783, HC784); (ii) crossing a representative line that coexpresses *rde-4(+)* and DsRed2 in bwm cells of WM49 into HC734 and HC737; (iii) crossing a representative line that coexpresses *rde-1(+)* and DsRed2 in bwm cells of WM27 into HC738; (iv) crossing a representative line that coexpresses *mut-2(+)* and DsRed2 in the bwm cells of WM30 into HC733 and HC739; and (v) crossing a representative line that coexpresses *gfp-dsRNA* and DsRed2 in the pharynx of HC195 and PD4792 into HC739 and HC735, respectively. To avoid bias due to observed phenotypic defects, cross progeny or rehomologized progeny were either selected using the DsRed2 co-injection markers or selected randomly, and the genotype was determined subsequently by PCR.

We analyzed 30–35 animals from three independent transgenic rescue lines ($n = 100$) in all cases except for the rescue of *rde-4(+)* in neurons of *rde-4(-)* animals, where a representative rescue line was crossed into *rde-4(-)* animals with the *neur::u22ds* transgene and 100 double-transgenic animals were analyzed.

Microscopy. Fluorescent images shown are projections of Z-series that were acquired using a Zeiss spinning-disc confocal microscope, except in **Supplementary Figure 1a**, where wide-field fluorescent images taken using a dissecting fluorescent microscope are shown. In all figures, images for strains that are being compared were acquired under the same non-saturating exposure conditions and, with the exception of the DsRed inset in **Figure 2b**, were then adjusted identically using Image J (US National

Institutes of Health) and Adobe Photoshop to allow the images to be viewed under normal printing conditions.

RNAi assays. To measure the extent of GFP silencing, we used a dissecting fluorescent microscope to count the number of brightly fluorescent gut nuclei in animals of the fourth larval stage (L4 stage) that are visible at a fixed magnification. The two nuclei that are located below two other nuclei in the first segment of the intestine (Int 1) are not easily resolved at this level of magnification and were not counted in this assay. Silencing in **Figure 1c** was measured at 25 °C, because some silencing of pharyngeal GFP is observed at lower temperatures, which is consistent with previous reports of RDE-1-independent silencing²⁰. For feeding RNAi, L4-staged animals were fed bacteria that express L4440 control dsRNA or dsRNA matching a target gene on agar plates that contain 1 mM isopropyl β -D-1-thiogalactopyranoside (IPTG). The percentage of the resulting L4 progeny showing the corresponding defects was determined. For *act-5* silencing, the number of animals that survived beyond L4 on day 5 for each genotype is expressed as a percentage of animals of the same genotype that were either L4 or older on day 5 on L4440. To measure *unc-22* silencing in response to expressed *unc-22*-dsRNA or *unc-22* feeding RNAi, we determined the percentage of L4-staged animals that twitched within 3 min in 3 mM levamisole (Sigma Aldrich) or on RNAi feeding plates without levamisole, respectively.

DNA constructs and transgenic animals. PCR fragments for transgenic expression³³ and transgenic animals³⁴ were generated using standard methods as in ref. 8. Briefly, PCR fragments corresponding to the coding sequences and 3' UTRs were amplified and fused to promoter sequences using an overlap extension PCR³². These fragments were then purified using a PCR cleanup column (Qiagen) and injected along with appropriate co-injection markers into *C. elegans* to generate transgenic lines. The specific primers used for PCR (**Supplementary Table 2**) and the specific concentrations and markers used for injections are detailed in **Supplementary Methods**.

Statistical analysis. The statistical significance of differences in average numbers of gut nuclei was calculated using the Student's *t*-test. For all other assays, 95% confidence intervals for single proportions were calculated using Wilson's estimates with continuity correction³⁵, and significant differences were determined using Wilson's pooled estimates.

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