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To cite this version:
Michalis Barkoulas, Amhed Vargas Velazquez, Alexandre Peluffo, Marie-Anne Félix. Evolution of New cis-Regulatory Motifs Required for Cell-Specific Gene Expression in Caenorhabditis. PLoS Genetics, Public Library of Science, 2016, 12 (9), pp.e1006278. 10.1371/journal.pgen.1006278. inserm-02159894

HAL Id: inserm-02159894
https://www.hal.inserm.fr/inserm-02159894
Submitted on 19 Jun 2019

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Evolution of New \textit{cis}-Regulatory Motifs Required for Cell-Specific Gene Expression in \textit{Caenorhabditis}

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Abstract

Patterning of \textit{C. elegans} vulval cell fates relies on inductive signaling. In this induction event, a single cell, the gonadal anchor cell, secretes LIN-3/EGF and induces three out of six competent precursor cells to acquire a vulval fate. We previously showed that this developmental system is robust to a four-fold variation in \textit{lin-3/EGF} genetic dose. Here using single-molecule FISH, we find that the mean level of expression of \textit{lin-3} in the anchor cell is remarkably conserved. No change in \textit{lin-3} expression level could be detected among \textit{C. elegans} wild isolates and only a low level of change—less than 30%—in the \textit{Caenorhabditis} genus and in \textit{Oscheius tipulae}. In \textit{C. elegans}, \textit{lin-3} expression in the anchor cell is known to require three transcription factor binding sites, specifically two E-boxes and a nuclear-hormone-receptor (NHR) binding site. Mutation of any of these three elements in \textit{C. elegans} results in a dramatic decrease in \textit{lin-3} expression. Yet only a single E-box is found in the \textit{Drosophilae} supergroup of \textit{Caenorhabditis} species, including \textit{C. angaria}, while the NHR-binding site likely only evolved at the base of the \textit{Elegans} group. We find that a transgene from \textit{C. angaria} bearing a single E-box is sufficient for normal expression in \textit{C. elegans}. Even a short 58 bp \textit{cis}-regulatory fragment from \textit{C. angaria} with this single E-box is able to replace the three transcription factor binding sites at the endogenous \textit{C. elegans} \textit{lin-3} locus, resulting in the wild-type expression level. Thus, regulatory evolution occurring in \textit{cis} within a 58 bp \textit{lin-3} fragment, results in a strict requirement for the NHR binding site and a second E-box in \textit{C. elegans}. This single-cell, single-molecule, quantitative and functional evo-devo study demonstrates that conserved expression levels can hide extensive change in \textit{cis}-regulatory site requirements and highlights the evolution of new \textit{cis}-regulatory elements required for cell-specific gene expression.
Author Summary

Diversification of mechanisms regulating gene expression of key developmental factors is a major force in the evolution of development. However, in the past, comparisons of gene expression across different species have often been qualitative (i.e. ‘expression is on versus off’ in a certain cell) without precise quantification. New experimental methods now allow us to quantitatively compare the expression of gene homologs across species, with single cell resolution. Moreover, the development of genome editing tools enables the dissection of regulatory DNA sequences that drive gene expression. We use here a well-established “textbook” example of animal organogenesis in the microscopic nematode, *Caenorhabditis elegans*, focusing on the expression of *lin-3*, coding for the main inducer of the vulva, in a single cell called the anchor cell. We find that the *lin-3* expression level is remarkably conserved, with 20–25 messenger RNAs per anchor cell, in species that are molecularly as distant as fish and mammals. This conservation occurs despite substantial changes and compensation in the regulatory elements required for cell-specific gene expression.

Introduction

Developmental systems operate in the presence of stochastic, environmental and genetic perturbations. While the output of a developmental system may be under stabilizing selection and remain mostly invariant, many internal variables such as the expression of a key gene or the activity of signalling pathways can be sensitive to perturbations. To reach a quantitative understanding of developmental systems, a key approach is to measure the sensitivity of the developmental system output to induced variation in an intermediate developmental phenotype. Whether and how this intermediate developmental phenotype varies within and among species then becomes a relevant evolutionary question [1]. The present work addresses the evolution of the expression level of the inducer of vulval development, *lin-3*, on which we previously performed a sensitivity analysis by manipulating its genetic dosage and addressing the phenotypic consequences for the developmental system [2].

The site and level of transcription of a gene can be modulated both in cis to the gene through cis-regulatory DNA sites directly influencing its transcription, or in trans due to evolution of trans-factors modifying the cellular context in which the gene is acting [3]. cis-regulatory sites containing binding sites for transcription factors often occur upstream of the coding region or in introns. These binding sites are often organized in modules, hence the designation as cis-regulatory modules (CRMs), acting in concert to enhance or repress gene expression in a given tissue at a given time. Changes in the number, relative order, orientation and spacing of transcription factor binding sites can affect transcription, often in a tissue-specific manner [4–6]. Tissue-specificity of CRMs is important for organismal evolution as it is thought to contribute to evolutionary novelty by minimizing pleiotropy [7–12]. Comparative studies in closely related species have revealed that transcriptional regulation can evolve through either extensive rewiring, or quantitative variation in the molecular components of a conserved network [11,13–17]. In particular, changes in cis-regulatory elements directly influencing the expression of critical developmental regulators have been shown to be a driving force for evolutionary innovation and phenotypic novelty in a variety of organisms. One example in *Caenorhabditis* concerns evolution between *C. elegans* and *C. briggsae* in the expression pattern of the transcription factor *lin-48* in the excretory system, resulting in a morphological change in excretory cell position. In this case, *lin-48* expression was gained in the excretory duct cell of *C. elegans* due to the acquisition of upstream binding sites for the transcription factor CES-2 [18,19].
Several features now make nematodes excellent experimental systems to understand gene expression evolution. First, rhabditid nematode species present a great advantage because homologous cells are easy to identify [20] so gene expression can be measured in a given cell. Second, the model organism *Caenorhabditis elegans* and other congeneric nematodes are amenable to functional genetics, transgenesis and now genome editing [21–26]. While transgenesis in *C. elegans* has long relied on formation of extra-chromosomal arrays containing many copies of the injected DNA that rearrange in an uncontrolled manner [27], the integration of a single copy at a defined locus is now possible, either at the endogenous locus using CRISPR/Cas9-mediated replacements [24–26,28] or at a controlled insertion locus using Mos1-mediated single-copy insertions (MosSCI) [29]. Third, *Caenorhabditis* species are highly divergent at the molecular level [30,31]. For example, *C. elegans* is as molecularly distant to *C. briggsae* as human is to mouse, and *C. angaria* as far as zebrafish to mouse [31], providing an opportunity to study the turnover of regulatory sequences at a large evolutionary scale where the nucleotide turnover is many times saturated yet the cellular context unchanged [32]. Many new *Caenorhabditis* species have recently been found and fully sequenced genomes are now available [33,34] (M. Blaxter, pers. comm.). Finally, the recent advent of quantitative methods, such as single-molecule fluorescent *in situ* hybridisation (smFISH) [35,36], allows to compare gene expression across species. The intensity of the conventional *in situ* hybridization signal cannot be meaningfully compared among species (regardless of whether the same probes or different probes targeting orthologs are used), while in the smFISH method the number of spots reflecting individual mRNA molecules can be counted, allowing a quantitative study of gene expression evolution.

Here, we take advantage of these recent developments to study the expression and requirement of *lin-3*, a model developmental gene involved in *C. elegans* vulval induction. The vulva is the egg-laying and copulatory organ of nematodes, and the *C. elegans* vulva is now a ‘textbook’ example of animal organogenesis [37]. *C. elegans* vulval development involves induction of three ventral epidermal cells (P5.p-P7.p) in response to the secretion of the LIN-3 signal from the anchor cell of the somatic gonad. LIN-3 activates the EGF receptor in the vulval precursor cells closest to the anchor cell and thereby acts as the upstream major inducer of vulval fates, in three precursor cells out of the six competent cells (Fig 1A). Induction of vulval fates involves interactions between EGF-Ras-MAPK, Notch and Wnt signalling, including some established pathway crosstalks [38]. We previously showed by modulating *lin-3* expression via single-copy transgenesis that the genomic level of *lin-3* expression is limited within a four-fold range for the vulva to develop normally in the *C. elegans* N2 background [2].

The *C. elegans lin-3* gene has two alternative promoter regions, each including transcriptional and translational start sites. The *lin-3* anchor cell isoform is driven by a specific *cis*-regulatory module lying immediately 5' of the second promoter, which is located in the first intron of the mRNA driven by the upstream promoter. Within this region, a 59 bp element was shown to be sufficient to drive expression in the anchor cell, acting as a transcriptional enhancer if placed upstream of a minimal promoter [39]. Anchor cell expression was shown to rely on two types of transcription factor binding sites in this 59 bp element, conserved in *C. briggsae* [39] (Fig 2): an NHR-binding site and two E-boxes. The *lin-3*(e1417) mutation substitutes a single nucleotide within the NHR-binding site and results in a strong reduction of *lin-3* expression in the anchor cell [2,39]. This site can be bound *in vitro* by nuclear hormone receptors such as *C. elegans* NHR-25. The two E-boxes surround the NHR-binding site (E-boxL for left to the NHR and E-boxR for right), each consisting of the conserved sequence "CACCTG" but on opposite DNA strands to each other. When either of them is mutated in a *lin-3::GFP* transgene context, GFP expression in the anchor cell is strongly reduced [39]. We refer for simplicity to the ensemble of these three regulatory elements as the “regulatory triplet”.

cis-Regulatory Evolution at the *lin-3* Locus

A

![Diagram](Image)

B

C. *elegans*

D

C. *angaria*

C

C. *briggsae*

E

O. *tipulae*

F

![Bar chart](Image)

lin-3 mRNAs
We show here that a relative stability in *lin-3* mRNA expression in the anchor cell and conservation of *LIN-3* vulval induction activity contrasts with the turnover of *cis*-regulatory binding sites at the *lin-3* locus. We show that the difference in requirement of regulatory elements for anchor cell expression is due to evolution in *cis* to the *lin-3* locus without a need to infer evolution in *trans*. This evolution in *cis* occurs in a very short 58bp region upstream of the *lin-3* vulval specific isoform. This study uncovers the evolution of new *cis*-regulatory motifs required for cell-specific gene expression.

**Fig 1.** *lin-3* expression level in the anchor cell is overall conserved in different *Caenorhabditis* species. (A) Cartoon depicting the position of the anchor cell (AC) and Pn.p cells at the time of induction. Three Pn.p cells (P5.p – P7.p) are induced upon LIN-3 secretion. (B-E) smFISH using a *lin-3* probe in *C. elegans* N2 (data from [2]), *C. briggsae* AF16 (C), *C. angaria* RGD1 (D) and *O. tipulae* CEW1 (E). Red arrow marks the position of the anchor cell. (F) Quantification of the number of spots detected in the anchor cell of these species at the time of induction (n = 32* animals for N2, n = 24 for AF16, n = 26 for RGD1 and n = 22 for CEW1). *: these include 20 animals from [2] (see Fig 6 for an independent dataset with a similar result). The difference between *C. elegans* and *C. briggsae* is not statistically significant with a Tukey’s multiple comparison test (P value = 0.99), whereas the difference between *C. elegans* and *C. angaria*, or *C. elegans* and *O. tipulae* is significant (P values < 0.0002).

doi:10.1371/journal.pgen.1006278.g001

**Fig 2.** *lin-3* activity in vulval induction is conserved in *Caenorhabditis* species. (A) Comparative *lin-3* RNAi effect on vulva induction in *C. elegans*, *C. briggsae* and *C. remanei*. Tables show graphically the observed defects in vulval cell fate pattern after scoring at least 100 nematodes. Every column is a distinct Pn.p cell (3 to 8) and 1° fate is depicted in blue, 2° fate is depicted in red and 3° fate in yellow. Half fates represent cases where the Pn.p daughter cells adopt different cell fates after the first cell division. The defects are ordered based on their consequence on vulval induction index, from high index to low. (B) Treatment with the MEK inhibitor U0126 decreases vulval induction in *C. elegans* (n = 15 for DMSO control and 10 μM U0126 treatment), *C. angaria* (n = 32 for control, n = 27 for 150 μM U0126 treatment) and *C. atra* (n = 100 for control, n = 30 for 150 μM treatment), as measured by the vulval induction index (average number of induced Pn.p cells at the population, wild-type index = 3). In all cases P<0.0001 with a Mann Whitney test.

doi:10.1371/journal.pgen.1006278.g002
Results

Evolutionary conservation of lin-3 mRNA expression in the anchor cell of Caenorhabditis and Oscheius

To determine the level of intraspecific variation in lin-3 expression, we quantified lin-3 expression in different C. elegans wild isolates. In the reference strain N2, a mean level of 25.4 lin-3 mRNA spots was detected using smFISH [2,40] (Fig 1B; S1A Table). We found that the mean and range of lin-3 expression in the anchor cell at the time of vulval induction are comparable between the C. elegans reference strain N2 and the most genetically divergent C. elegans isolates such as DL238 and QX1211 (S1A Fig; S1A Table).

We further explored lin-3 expression in different rhabditid species. First, we searched for the lin-3 ortholog in other available genomes (S2 Fig). The LIN-3 proteins can be aligned along their whole length, with a conserved signal peptide, EGF and trans-membrane domains. Interestingly, the most conserved parts of the proteins are the N-terminal part following the signal peptide and the intracellular domain [41].

We designed smFISH probes for the lin-3 gene of C. briggsae, C. avara, C. angaria and Oscheius tipulae and found that lin-3 is expressed in a single cell within the somatic gonad, immediately dorsal to P6.p, which we identified by DAPI staining as the anchor cell (Fig 1C–1E; S1B Fig; S1B Table). Similar to C. elegans, we also detected lin-3 expression at a lower level in the gonad outside the anchor cell and in the pharynx. We quantified fluorescent spots in the anchor cell and found no significant difference between C. elegans and C. briggsae (mean of 26.5±1 standard error in C. elegans vs. 25±1 in C. briggsae) (Fig 1F). In C. angaria and O. tipulae, we only found a small decrease compared to C. elegans (Fig 1F). Although lin-3 was clearly detected in the anchor cell of C. avara (S1B Fig), the inferior quality of the hybridisation signal compared to the background did not allow us to quantify fluorescent spots in this species. We conclude that despite the great genetic distance between these nematodes [31], the mean number of lin-3 mRNAs is remarkably conserved at least in C. briggsae and may only vary within a narrow range in C. angaria and O. tipulae.

Conserved role of LIN-3 in inducing vulval cell fates

The vulval cell fate pattern is conserved throughout the Rhabditidae family, to which the Caenorhabditis and Oscheius genera belong [42], nevertheless molecular underpinnings of vulval induction in species other than C. elegans remain mostly unknown. lin-3 RNAi experiments in C. briggsae so far produced a weak effect [43]. In Pristionchus pacificus, an outgroup and the only nematode species for which we currently have substantial molecular information related to vulval induction, vulval formation relies on Wnt signalling and is thought to be independent of the EGF pathway [44,45].

To address whether the lin-3 homolog plays a functional role in vulval induction in different Caenorhabditis species, we used a combination of RNAi and pharmacological inhibition. First, we used recently established strains of C. remanei and C. briggsae that are rendered sensitive to RNAi administered by feeding due to the expression of the C. elegans intestinal transporter sid-2 [21,46]. lin-3 RNAi treatment in these C. briggsae and C. remanei strains resulted in substantial reduction in vulval induction (Fig 2A; S3A–S3D Fig). We observed vulval cell fate phenotypes upon lin-3 RNAi that are not found in C. elegans, but are in keeping with published results revealing cryptic variation in vulval fate patterning following anchor cell laser ablations. Specifically, we found that P(5–7).p adopted a 2°-3°-2° cell fate pattern in C. remanei and a 2°-2°-2° pattern in C. briggsae [17,43]. Second, we used the MAP kinase (MEK) inhibitor U0126 that inhibits the downstream signalling events following EGF
receptor activation. Application of this inhibitor has been previously shown to decrease vulval induction in *O. tipulae* [47]. Consistent with this result, we also obtained evidence for loss of overall vulval induction both in *C. angaria* and *C. afru* (Fig 1B; S3D Fig). Thus, we conclude that *lin-3* is expressed in the anchor cell and plays a conserved role in inducing vulval fates in the *Caenorhabditis* genus.

The *lin-3* regulatory triplet evolved at the base of the *Elegans* species group

Three transcription-factor binding sites, an NHR-binding site and two E-boxes, are required for *lin-3* expression in the anchor cell of *C. elegans* [39]. In light of the conserved expression pattern and level, we wondered whether these regulatory elements required for AC expression of *lin-3* are also conserved. The regulatory triplet was found to be present in different species of the *Elegans* group of *Caenorhabditis* including *C. briggsae* (Figs 3, S4 and S5). However, in the sister clade, called the *japonica* group, we were able to find the two E-boxes, but no putative NHR-binding site within a window of 2.5 kb upstream of the translational start site of the vulval isoform of *lin-3*. In further outgroup species, such as *C. angaria*, we only found a single E-box, and no NHR-binding site in this region. One E-box within the *lin-3* CRM was also detected in the outgroup *Oscheius tipulae* (Fig 3). In *C. sp. 1*, we were able to detect a single ATG and the first E-box was only found 2 kb upstream. Overall, these observations suggest that the NHR-binding site was acquired in the branch leading to the *Elegans* group of the *Caenorhabditis* genus. The evolution of the second E-box at the base of the *Caenorhabditis* genus remains unclear: the second E-box may have been acquired in the branch leading to the *Elegans* supergroup or else be lost in the *Drosophilae* supergroup. No other sequence similarity could be found in the region upstream of the ATG of the vulva-expressed isoform of *lin-3* (S4 Fig).

The above results raised an interesting conundrum. How is it possible that some elements that are required for *lin-3* anchor cell expression in *C. elegans* are completely missing in related species, without any significant consequence for *lin-3* spatial and quantitative expression?

A single *C. elegans* E-box cannot drive *lin-3* expression in the anchor cell

We first aimed to confirm that one E-box is not sufficient for *lin-3* expression in the anchor cell in *C. elegans*. We used CRISPR-mediated genome editing [48] to select deletions of *cis*-regulatory elements of the *C. elegans* *lin-3* gene. We generated a variety of alleles, in which either all three elements are deleted (mf90), or NHR and E-boxR are deleted leaving E-boxL intact (mf72-mf74) or only E-boxR is left intact (mf75), the latter recapitulating the *cis*-regulatory context of the *C. angaria* *lin-3* upstream module (Fig 4A). All these alleles result in fully penetrant vulvaless phenotypes with no cell induced to a vulval fate, thus a stronger phenotype than the *lin-3*(*e1417*) allele with one-nucleotide substitution in the NHR binding-site (Fig 4B). We used smFISH to detect *lin-3* transcripts and found no *lin-3* expression in the anchor cell, which was visualised by the unperturbed expression of *lag-2*. Interestingly, we still detected *lin-3* expression in the gonad of these mutant animals (Fig 4C and 4D). We conclude that these new *lin-3* alleles are anchor cell-specific null alleles.

These results confirmed that one E-box in the upstream *cis*-regulatory module of *lin-3* is not sufficient for *lin-3* expression in the anchor cell of *C. elegans*—whereas it appears sufficient in species of the *Drosophilae* group such as *C. angaria*. 

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*PLOS Genetics* **DOI:10.1371/journal.pgen.1006278 September 2, 2016** 7 / 23
Compensatory evolution occurs in cis to the lin-3 locus

The evolution in the requirement of transcription-factor binding sites for lin-3 expression in the anchor cell could be due to changes in cis or in trans to the lin-3 locus or both. We reasoned that if differences in trans were important, we would expect lin-3 genomic fragments derived from species missing one or two cis-regulatory elements from the regulatory triplet to be unable to be expressed in the anchor cell of C. elegans. We tested this hypothesis and obtained multiple lines of evidence suggesting no role for changes in trans to the lin-3 locus in explaining the differential binding site requirement.

First, we overexpressed in C. elegans a C. angaria lin-3 genomic fragment containing 200 bp of upstream sequence, the coding region and the 3′ UTR. This fragment drove anchor cell expression of Can-lin-3 and triggered vulval hyperinduction in C. elegans, further showing that the Can-LIN-3 protein could activate the C. elegans LET-23/EGF receptor (S6A and S6B Fig). Vulval hyperinduction was also observed when an equivalent genomic fragment from C. elegans was expressed in C. angaria or a fragment from C. afr a was expressed in C. elegans (S6C and S6D Fig). These results indicate that the injected lin-3 fragments from different Caenorhabditis species contain the necessary information for anchor cell-specific expression, despite the fact that a superficially equivalent C. elegans fragment with only one E-box, as in the new lin-3 alleles described above, cannot be expressed in this cell.

Since the regulatory triplet for C. elegans anchor cell expression is missing in these transgenes, we tested whether sequences in the introns, exons or 3′UTR sequences were required for expression of the C. angaria transgene in the anchor cell. To this end, we fused the Can-lin-3 upstream sequences to a fragment containing the C. briggsae lin-3 coding sequence and 3′ UTR. We expressed this fragment in C. elegans N2 and again observed clear expression in the anchor cell. As expected, in control injections containing only the promoterless C. briggsae fragment, the transgene was not expressed anywhere in the body (S7 Fig). To further strengthen these results, we fused the lin-3 cis-regulatory modules amplified from C. elegans, C.
briggsae, C. afra and C. angaria to sequences encoding an unrelated protein, the fluorescent protein Cherry, and the unrelated unc-54 3'UTR. In all cases, we observed clear expression in the anchor cell (Fig 5A), indicating again that these short cis-regulatory modules alone contain the necessary information for anchor cell-specific expression in C. elegans. We conclude that evolution within the 200 bp upstream cis-regulatory module of lin-3 is sufficient to explain the difference in requirement of regulatory elements for anchor cell expression within Caenorhabditis. The C. angaria lin-3 transgene quantitatively mimics a C. elegans lin-3 transgene

Above, we used multicopy transgenesis, which may cause sufficient expression and hyperinduction due to summing of weak transcriptional activity of many copies. We thus next asked whether the C. angaria lin-3 fragment had quantitatively a similar activity to that of its C. elegans counterpart when introduced in single copy at a targeted genomic location outside the lin-3 locus (using MosSCI transgenesis, see Methods). We found that a single-copy Can-lin-3 insertion in C. elegans N2 is expressed in the anchor cell (Fig 5B) and does not cause hyperinduction, like an equivalent Cel-lin-3 transgene copy [2]. Most interestingly, this single copy transgene could completely rescue the induction and brood size of lin-3(e1417) mutants, both
in homozygous and hemizygous states (Figs 5C, S8). This quantitative behavior of the Can-lin-3 transgene (rescue in the hemizygous and homozygous state, no effect when added to the endogenous locus) recapitulates the activity of a C. elegans copy inserted at the same genomic location [2]. This experiment shows that the C. angaria lin-3 gene driven by its cis-regulatory element acts in a similar quantitative manner to the C. elegans fragment, even in the absence of the regulatory triplet.

A 58 bp cis-regulatory fragment from C. angaria with a single E-box can replace the entire C. elegans regulatory triplet

To pin down the regulatory elements in the C. angaria transgene that are required for anchor cell expression, we mutated the E-box, which is the only distinguishable regulatory element in this short upstream region. We found that Can-lin-3 genomic fragments with a mutated E-box lose their ability to be expressed in the anchor cell of C. elegans and to trigger vulval

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doi:10.1371/journal.pgen.1006278.g005

doi:10.1371/journal.pgen.1006278.S005
hyperinduction when expressed as multi-copy transgenes (Fig 6B, 6D and 6E). This shows that the single E-box$_R$ of \textit{C. angaria} is necessary for \textit{lin-3} expression in the anchor cell of \textit{C. elegans}.

Changes in the flanking sequences to core binding sites have been shown to contribute to binding efficiency of transcription factors, so we reasoned that perhaps the difference in requirement of regulatory elements for \textit{lin-3} expression in the anchor cell may rely on nucleotides adjacent to the single E-box. To this end, we synthesised a chimeric CRM, where a 58 bp central portion harbouring the regulatory triplet in \textit{C. elegans} was replaced with 58 bp from \textit{C. angaria} containing E-box$_R$ (Fig 6E). We first showed that this chimeric fragment can be expressed in the \textit{C. elegans} anchor cell when used in multiple-copy extra-chromosomal array transgenesis (Fig 6C). Furthermore, we used genome editing at the \textit{Cel-lin-3} locus to replace the endogenous \textit{lin-3} CRM with this chimeric CRM. We found that the genome-edited animals expressed \textit{lin-3} in the anchor cell at a normal level and produced a phenotypically wild-type vulva (Fig 6F; S2 Table).

These results demonstrate that the difference in requirement of \textit{cis}-regulatory elements between \textit{C. elegans} and \textit{C. angaria} is explained by compensatory evolution within a very short \textit{cis}-regulatory fragment (58 bp), rendering the presence of a second E-box and the NHR binding site unnecessary in \textit{C. angaria}. Despite this loss of transcription factor binding sites, the activity of the \textit{cis}-regulatory module in driving transcription in the anchor cell remains at the same quantitative level.

The compensation could be explained by the gain of new transcription factor binding sites in the \textit{C. angaria} 58 bp regulatory region. To identify putative transcription factor binding sites, we performed a motif discovery approach in the anchor cell \textit{cis}-regulatory \textit{lin-3} regions of \textit{Caenorhabditis} species close to \textit{C. angaria} and an exhaustive search of transcription factors that could bind the 58 bp sequence (see Methods). We found the GTTTATG sequence, a possible Forkhead-binding site, to be significantly over-represented. This sequence is only one bp to the right of the \textit{C. angaria} E-box. We tested whether modifying this sequence in the 58 bp \textit{C. angaria} replacement would change the \textit{lin-3} expression level. Indeed, when scrambling these 7 bp (see Methods; S2 Fig), \textit{lin-3} expression was reduced significantly to about 60% of the wild-type level (\textit{mf95} allele in Fig 6F; t-test, p-value < 6 $\times$ 10$^{-8}$). However, as expected from a less than two-fold decrease [2], this new replacement, like the intact \textit{C. angaria} CRM, produced phenotypically wild-type vulva cell fate induction (Fig 6F). Thus, we could affect the expression of the \textit{C. angaria} CRM by modifying a motif adjacent to the E-box. This motif contributes to the compensation in \textit{cis} in the 58 bp, but does not explain all of it, as \textit{lin-3} expression in the \textit{mf95} mutated replacement allele was still much higher than with a single \textit{C. elegans} E-box.

**Discussion**

**A quantitative account of gene expression evolution**

This study addressed the level of expression of a critical developmental regulator in a single cell. We showed that both \textit{lin-3} expression level in the anchor cell and its requirement for the induction of vulval cell fates are conserved in \textit{Caenorhabditis} and \textit{Oscheius} nematode species. We found that the mean \textit{lin-3} mRNA level in the anchor cell only varies within 30%, despite the vast genetic divergence in this group—corresponding to that found among the most diverged vertebrates. We previously showed using quantitative perturbations that the mean level of \textit{lin-3} expression in \textit{C. elegans} needs to stay within a four-fold range for a correct vulva pattern to arise and that the mean \textit{C. elegans} N2 level is in the very middle (on a log scale) of this permissible zone. Therefore, it is likely that stabilizing selection acting on vulva formation [49] leads to stasis both in \textit{lin-3} expression level and in its effect on vulval induction.
Fig 6. A 58 bp C. angaria fragment with a single E-box is able to replace the C. elegans regulatory triplet. (A) Expression of a Can-lin-3 fragment in C. elegans containing the Can-lin-3 CRM, coding sequences and 3' UTR leads to Can-lin-3 expression in the anchor cell, as detected by FISH. (B) Expression of the same fragment with a mutated E-box in the CRM results in loss of anchor cell expression. (C) A chimeric CRM that is mostly C. elegans apart from a 58 bp region around the regulatory triplet that is taken from C. angaria is also expressed in the anchor cell of C. elegans. (A) and (C) are using classical transgenesis in multicopy arrays experiments. (D) Over-expression of the Can-lin-3(+) fragment in C. elegans causes an increase in vulval induction, but not if the Can-lin-3 fragment with a mutated E-box is used. (E) Summary of the compensatory changes in cis to the C. angaria lin-3 locus allowing lin-3 expression in the anchor cell. C. elegans sequences are depicted in green and C. angaria sequences in blue. Orange box corresponds to the E-box. (F) Modification at the Cel-lin-3 endogenous locus, replacing the regulatory triplet with 58 bp from C. angaria containing a single canonical E-box with its genomic context (mf91 and mf92) or with 7 bp modified to its right (mf95 and mf112). The violin plots show the number of lin-3 mRNAs spots quantified in the anchor cell in the CRISPR replacements compared to N2 and lin-3(e1417) which are used as control strains. The whisker plot is superimposed in red. The average number of induced VPCs is shown below with the number of scored animals being 130, 159, 111, 42, and 35 from left to right.

doi:10.1371/journal.pgen.1006278.g006
By contrast with this evolutionary stasis in vulval pattern and in the lin-3 mRNA level, we showed that this cell-specific level of lin-3 expression involves substantial turnover of key cis-regulatory elements, namely the appearance of a novel binding site (NHR) and the turnover of a second copy of an existing binding site (E-box). Each of these elements is required for anchor cell expression in C. elegans yet is absent in some Caenorhabditis species. We further focused on the difference in requirement of cis-regulatory elements for lin-3 expression between C. elegans and C. angaria. A 58 bp fragment from C. angaria with a single E-box can replace the three C. elegans binding sites, demonstrating that compensatory evolution within this short cis-regulatory fragment at the lin-3 locus is sufficient to explain this difference in transcriptional regulation.

Among evo-devo studies that center on comparisons of gene expression patterns and the evolution of cis-regulatory sequences, this is to our knowledge the first study taking advantage of the latest available capabilities to edit genomes and to quantify the level of mRNA expression at the single-cell level in a multicellular eukaryote.

Turnover of transcription-factor binding sites

Gene expression may evolve due to changes in cis or in trans to a given locus, two possibilities that are not mutually exclusive. Cis-regulation may occur from sites quite distant to the transcriptional unit due to long-range chromatin interactions. Our data provided strong support for compensatory cis-changes, and this in a DNA fragment directly upstream of the translational start site of the vulva specific isoform of lin-3. We cannot exclude that some further trans-changes facilitate the difference in requirement of regulatory elements between the two species. However, the cis-regulatory changes that we uncovered in this work are at least sufficient to explain the difference in requirement of regulatory elements for anchor-cell-specific gene expression in Caenorhabditis.

We have narrowed down the compensatory changes that allow the C. angaria lin-3 to be expressed in the anchor cell in a very short region of 58 bp. To explain the compensatory changes, we performed an exhaustive search of transcription factor binding sites and found a putative Forkhead binding site immediately adjacent to the E-box in C. angaria and absent from the replaced 58 bp region of C. elegans. Mutation of this site significantly lowered lin-3 expression, but insufficiently to affect the vulval induction level and it thus only partially explained the compensatory evolution in cis (Fig 6E). We further note that, because this putative Forkhead binding site is immediately adjacent to the E-box, we cannot distinguish between two scenarios: a role for another specific transcription factor binding site versus an alteration of the affinity of the E-box itself. An alternative model would indeed be that compensation occurs through a stronger affinity of the E-box in the C. angaria regulatory region, while the C. elegans E-box is insufficient to drive expression. Such differences in affinity may arise from changes in the sequences flanking the core binding sites as it has been shown for bHLH factors binding to E-boxes [50,51]. Variation in the flanking sequences next to core transcription factor binding sites has also recently been shown to influence both the levels and sites of gene expression for another developmentally important gene [52]. We conclude that the GTTTATG sequence contributes to the compensation, but does not explain it entirely.

Evolution of transcriptional regulation without change in gene expression

Here we described some evolution in cis-regulatory elements that occurs without consequences at the level of gene expression, as observed in many other genes and various groups of organisms [53–56]. This cis-regulatory element turnover in the absence of phenotypic consequence can be viewed as an extension to the notion of developmental systems drift, which posits that
distinct molecular mechanisms may underlie the emergence of similar developmental pheno-
types [57]. In a similar way, the conservation of gene expression pattern and level may depend
on distinct molecular mechanisms due to the loss and gain of binding sites. Indeed, if the
invariant output phenotype that we consider is *lin-3* expression level in the anchor cell, the
molecular events leading to it, such as transcription factor binding, do vary in evolution.

The best-studied example for conservation of gene expression pattern despite turnover of
cis-regulatory elements is the stripe 2 enhancer of the *Drosophila* pair-rule gene *even-skipped*.
The minimal stripe 2 enhancer (*eve2*) in *D. melanogaster* is a DNA region of approximately
500 bp that consists of multiple binding sites for activators such as Bicoid and Hunchback and
for repressors such as Giant and Krüppel: their combination allows a confined expression in
the second stripe along the antero-posterior axis of the early *Drosophila* embryo [58]. Com-
pared to the described *lin-3 cis*-regulatory module, the *eve2* stripe element involves more tran-
scription-factor binding sites and results in expression in a group of cells (nuclei) rather than
in a single cell. Similar to the *lin-3 CRM*, the transcription-factor binding sites change in Dro-
sophila species in a way that binding sites required for correct expression in *D. melanogaster*
are absent in the stripe 2 element of other species, though without leading to alteration in the
expression domain, due to compensatory cis-changes [53,59]. Here we went further in replac-
ing the endogenous cis-regulatory sequences at the locus by those of a distant species, and
show a quantitative rescue of gene expression and vulval induction.

One previous example in *C. elegans* of turnover of binding sites involves *lin-48* expression
in hindgut cells, which is conserved between *C. elegans* and *C. briggsae* despite turnover of
EGL-38 upstream response elements [60]. This turnover shows both similarities and differ-
ences to the described evolution of *lin-3 cis*-regulatory elements. The similarity is that there is
an increase in the number of EGL-38 response elements in *C. elegans*. However, in the *lin-48*
case, there is evolution towards redundancy because the gain in one EGL-38 response element
decreases the reliance on the existing element for correct gene expression.

More recently, evolution of cis-regulatory elements between *C. elegans* and *C. briggsae* has
been studied by placing exogenous cis-regulatory elements from *C. briggsae* into *C. elegans*. A
main result over several genes whose expression is conserved between the two species is the
appearance of ectopic gene expression domains in these transgenic experiments, implying evo-
lution both in cis and in trans [61,62]. In one case, the ability of the *unc-47* proximal promoter
from *C. briggsae* to drive ectopic expression in some *C. elegans* neurons was mapped next to a
conserved cis-regulatory motif [61].

We note that the *C. angaria* fragment conveys the same level of transcriptional activity yet
that a few vulval cell fate patterning "errors" occur in the replacement lines (Fig 6F). We
observed both hypoinduced and hyperinduced variants in each of the two replacement lines
(S2 Table), but the very low frequency of these variants make them difficult to study quantita-
tively. In the case of the *eve2* enhancer, the minimal stripe element is embedded within a larger
region of approximately 800 bp, and these flanking sites contribute to robustness to some
genetic and environmental perturbations [63]. In *Caenorhabditis*, the distal promoter of *unc-
47*, although largely not conserved, is also important for robust gene expression, acting perhaps
in a sequence-independent manner [64]. It remains unclear whether any regions within and/or
outside the *lin-3 CRM* can play a similar role in stabilizing expression of *lin-3* in *Caenorhabdi-
tis* to different perturbations.

**An evolutionary gain in binding site requirement**

The distribution of *lin-3 cis*-regulatory elements in different *Caenorhabditis* nematodes and the
mapping of changes on the phylogeny suggests as the most likely evolutionary scenario a gain
of regulatory sites: the likely acquisition of an E-box before the common ancestor between the *Elegans* and *Japanica* groups and a gain of an NHR-binding site before the origin of the *Elegans* group. In addition, these sites not only appeared, but also became indispensable for *lin-3* anchor cell expression at least in *C. elegans*.

The acquisition of such new short regulatory motifs (6 bp) is easy and gains of regulatory motifs have been observed in other systems as well [65]. Given the high robustness of vulval development to several perturbations, the evolution towards a dependence on a higher number of sites for anchor cell expression is counter-intuitive and suggestive of evolution towards fragility. It is currently unclear what drove the evolution of these novel motifs with a conserved gene expression, whether selection or drift. Gains in interconnectedness between components of transcriptional networks may often occur non-adaptively, for example if they do not disrupt the underlying regulation [66]. Such gains can also be reshaped in equivalent network configurations and eventually become necessary depending on the evolution of the transcriptional network [67].

**Materials and Methods**

**Nematode culture, genetics and pharmacology**

A complete list of strains used in this study is presented in the supplement (S3 Table). All strains were maintained at 20°C and handled according to standard procedures [68]. We used the Bristol N2 strain as a reference *C. elegans* strain on standard NGM plates with OP50 as a food source. The U0126 treatments were performed by supplying the DMSO-dissolved inhibitor to NGM plates at a concentration between 10–150 μM and letting synchronised L2 stage nematodes develop into L4 larvae. Control treatments in this case were performed by growing nematodes on plates supplemented with DMSO only.

For the *Can-lin-3* rescue of the *C. elegans lin-3(e1417)* mutant, JU2495 hermaphrodites were crossed to JU2498 males and the F1 or F2 progeny were analysed for hemizygous or homozygous insertion phenotypic rescue, respectively.

**Identification of *lin-3* orthologs in *Caenorhabditis* genomes**

The *lin-3* genomic sequences of the different species were accessed in WormBase (www.wormbase.org; version WS252) or from the Caenorhabditis Genomes Project by Mark Blaxter’s laboratory (http://bang.bio.ed.ac.uk:4567) or from Matt Rockman’s laboratory. The *Oscheius tipulae* genome was sequenced and assembled as a collaborative effort between M. Blaxter’s and our lab (Besnard, Kotsouvolos et al., in preparation) and is available (http://oscheius.bio.ed.ac.uk/). We first used the TBLASTN algorithm conditioning only to the most identical hits, favouring those with high similarity in the N-terminal part and signal peptide, and lower e-value. Afterwards, we proceeded to predict gene bodies in these contigs using FGNESH (http://www.softberry.com) with a hidden Markov model specific to *C. elegans*. Finally, manual curation and annotation of the *lin-3* sequences were performed using as a reference the amino-acid sequence of the closest available *lin-3* ortholog.

**Transcription-factor motif recognition in *lin-3* promoter sequences**

To study the evolution of the regulatory triplet in the *Caenorhabditis* clade, we analysed the promoter regions upstream of the downstream ATG corresponding to the N-terminal exon homologous to that known to be expressed in the AC of *C. elegans* (S2 Fig). First, to address whether the cis-regulatory *C. elegans* NHR-binding sites and E-boxes were present in the other species, we performed a scan in the promoters with the position weight matrices of HLH-2 and
NHR-proteins available in JASPAR [69] using matrix-scan [70] and a n = 2 Hidden Markov Model specific to C. elegans (Fig 3). Similarly, we looked in these regions for DNA patterns known to be binding sites of bHLH proteins [51] using the dna-pattern tool present in the RSAT suite [71]. Once we had the position of these sites across the promoter regions, we proceeded to plot their location using RSAT feature-map tool (S5 Fig).

Additionally, we looked for DNA motifs different from the cis-regulatory C. elegans NHR and E-boxes binding sites by performing a motif-discovery approach in Caenorhabditis lin-3 promoters using the RSAT tool oligo-analysis [71]. The top over-represented words of length 6, 7 and 8 base pairs were compared to known motifs available in JASPAR. We thus identify the GTTTATG to the right of the E-box. Finally, to identify possible transcription factors acting on the AC lin-3 expression in the 58 bp C. angaria fragment, we performed an exhaustive search of the full JASPAR motif repertoire in the 58 bp replaced sequence using RSAT matrix-scan. This search found the putative Forkhead-binding motif and a putative overlapping bZIP-binding motif (Fos/Jun repressors). The 7 bp modification in the mf95 replacement also affected this predicted binding site of bZip transcription factors.

Cloning

All lin-3 CRMs reside directly upstream of the ATG of the vulval isoform of lin-3. To create the lin-3 CRM::Cherry::unc-54 constructs, we used a three-fragment Gateway approach merging the lin-3 CRMs cloned in pDONOR P4-P1R, the Cherry ORF cloned in pDONOR 221 and the unc-54 3’UTR cloned in pDONOR P2R-P3. All primer sequences containing attB4 forward and attB1 reverse recombination sites used to amplify the CRMs from gDNA from different species are shown in S4 Table.

unc-54 3’ UTR was amplified from N2 genomic DNA using primers unc-54attB2 and unc-54attB3. Worm-optimised Cherry was amplified from pAA64 using primers containing the attB1 and attB2 sites. All constructs were injected at 10 ng/μl with myo-2::GFP as co-injection marker and pBluescript as carrier DNA.

To create the Can-lin-3 insertion by MosSCI, we amplified a 2.9 kb lin-3 fragment from C. angaria genomic DNA using primers Canlin-3AvrII and Canlin-3XhoI. The amplicon was cloned into pCFJ151 (chromosome II targeting vector) [29] as an AvrII/XhoI fragment. Injections and recovery of insertions were performed using the direct insertion protocol, as previously described.

To overexpress lin-3 fragments in C. elegans or C. angaria, we amplified genomic fragments amplified from C. elegans (5.2 kb), C. angaria (3.2 kb) and C. afr (5.1 kb) using primer pairs RH9for/RH9rev, Canlin-3F2/Canlin-3R1 and Caflin-3oxF2/Caflin-3oxR1, respectively. The PCR products were injected directly (30 ng/μl) together with pBluescript as carrier and myo-2::GFP as co-injection marker.

To mutagenize the E-box in the C. angaria lin-3 CRM, the above 3.2 kb fragment was cloned into pGEM-Teasy and the 5’-CAGGTG-3’ sequence was modified to 5’-CAGGAA-3’ using primers t211a_g212a/t211a_g212a_anti and standard in vitro site directed mutagenesis.

The chimeric construct replacing a 58 bp region containing the C. elegans regulatory triplet (5’-cacctgtgtattttatgctggttttttcttgtgaccctgaaaactgtacacacaggtg-3’) with a similar in length sequence from C. angaria containing only one E-box (5’-attttttgcaagattttttgctggcggctggttttattatatgtgctggtctggttctgctgac-3’) was synthesised by Genewiz. This construct was used as PCR template to permute 7 bp to the right of the C. angaria E-box (5’-CAGGTGTGGTTTATG-3’ to 5’-CAGG TGTGGGATT-3’).

The chimeric construct to drive Cbr-lin-3 under the C. angaria CRM was built using fusion PCR. Briefly, the Can-lin-3 CRM was amplified from C. angaria genomic DNA with primers
Canlin-3 F2 and CaACFusion and the Cbr-lin-3 region coding region and 3' UTR from C. briggsae genomic DNA with primers Cbrlin-3F1 and Cbrlin-3R1. The two amplicons were then fused together using a third PCR reaction with primers Canlin-3F2 and Cbrlin-3R1. The final product was injected as a PCR fragment at 20 ng/μl concentration.

**Single molecule fluorescence in situ hybridization**

smFISH was performed in synchronized populations of L3 stage animals using short fluorescently labelled oligos as probes, as previously described [2]. The animals were age-synchronized by bleaching, followed by hatching of embryos in M9 buffer. The L1 larvae were then placed onto culture plates with food until the L3 stage, as determined by Nomarski microscopy, and then fixed.

The C. elegans lin-3 and lag-2 probes have been previously described [2]. The low level of genetic divergence within C. elegans allowed us to detect fluorescent spots while using the same FISH probe as in the N2 strain. For all other species we followed the same protocol as with C. elegans with the following two modifications to decrease the more pronounced background fluorescence. We used 20% formamide in the hybridisation and wash solutions and performed three washes post-hybridisation instead of two in C. elegans. Given that we are using different probes consisting of fewer oligos for the detection of lin-3 transcripts in these species together with slightly more stringent hybridisation conditions, the observed difference in the number of fluorescence spots may thus even be due to technical rather than biological reasons. The sequences of the new lin-3 probes can be found in **S5 Table**. The probes were labelled with Quasar 670 (Biosearch Technologies) and diluted to 100–200 nM for the overnight hybridisation.

**RNAi**

RNAi was performed by feeding the animals with dsRNA-expressing bacteria, as previously described [2]. The C. elegans lin-3 RNAi feeding clone used in this study is from the Ahringer RNAi library (Source Bioscience). A Cre-lin-3 fragment was amplified using oligos Crlin-3RNAiF1 and Crlin-3RNAiR1 that contain an XhoI restriction site. The PCR product was cloned into L4440 as an XhoI fragment. To create the C. briggsae lin-3 RNAi clone, a fragment was amplified using primers Cbrlin-3RNAiF1 and Cbrlin-3RNAiR1 and then cloned into pDONR 221 (Invitrogen) using attB1F and attB2R universal oligonucleotides. The lin-3 fragment was sequence verified and transferred to a Gateway compatible L4440 plasmid. Both constructs were transformed into E. coli HT115 for use in C. elegans feeding.

**Phenotypic characterisation**

To score the vulval cell fate pattern, nematodes were mounted with M9 on 3% agar pads containing 10 mM sodium azide and analysed under Nomarski optics. Standard criteria were used to infer cell fates based on the topology and number of cells at the L4 stage [43,72]. Half fates were assigned when two daughters of the Pn.p cells acquired distinct fates after the first cell division.

**Genome editing**

We followed the CRISPR/Cas9 target design and used reagents as previously described [48]. We targeted the following region at the C. elegans lin-3 CRM 5’-acctgaaactgtaaacAGG-3’ with AGG representing the PAM motif. We replaced the unc-119 target site under the pU6 promoter [48] with the lin-3 target site using fusion PCR first with primers E-box2A gRNA-F/
U6prom HindIII and E-box2A gRNA-R/ oligos U6prom EcoRI F followed by amplification of the full sgRNA fragment with U6prom EcoRI F/ U6prom HindIII R. The only modification was that we did not clone the lin-3 sgRNA in a vector but injected it directly as a PCR product (40 ng/μl, together with 40 ng/μl eft-3::Cas-9 and myo-2::GFP as co-injection marker).

To replace the endogenous lin-3 cis-regulatory element of C. elegans by a 58 bp lin-3 element from C. angaria, we first obtained a chimeric double-stranded DNA as homologous recombination template, using Gibson assembly of C. elegans lin-3 promoter extremities with 58 bp of the C. angaria lin-3 upstream sequence. In a similar fashion, we obtained a homologous recombination template identical to the previous but with modified bases next to the E-box. Oligonucleotide sequences are found in S4 Table.

Supporting Information

S1 Fig. Single-molecule FISH of lin-3 in C. elegans and C. afra. (A) smFISH quantification of Cel-lin-3. The level of lin-3 expression in other C. elegans isolates is similar to that in the N2 reference strain (n≥14 animals; S1A Table. (B) smFISH localising lin-3 transcripts in the anchor cell of C. afra. Serial optical sectioning through the anchor cell of a single animal showing lin-3 fluorescent spots.

S2 Fig. (Text file): lin-3 sequences from the different Caenorhabditis species, Oscheius tipulae and the Cel:Can-lin-3(mf91&mf92) and Cel:Can-lin-3(mf95) replacement, with annotations of cis-regulatory binding sites. The sequences of the enhancers used in Fig 5A are in bold. The endogenous 3′ UTR used for the overexpression experiments in Figs 5B, 6A–6D, S6 and S7 is underlined.

S3 Fig. lin-3 RNAi and MEK inhibitor treatment in different Caenorhabditis species. (A-D) Nomarski images of L4 stage animals upon lin-3 RNAi (A-C) or MEK inhibitor (U0126) treatment (D). (A-C) lin-3 RNAi by feeding in C. remanei strain JU1184 results in 2°-3°-2° (B) or 3°-3°-3° (C) vulval cell fates for P(5–7);p compared to the 2°-1°-2° of the wild-type (A). (D) Treatment with U0126 decreases vulval induction in C. angaria and C. afra. Note uninduced cells in both cases. The vulva in control L4 animals shows the typical "Christmas tree" morphology in all species.

S4 Fig. lin-3 cis-regulatory sequence alignments in different Caenorhabditis species. (A-C) Alignment of the 300 bp region upstream of the lin-3 ATG shows no other similarity in different species outside the E-box (B) and NHR (C) binding sites. Cbr = C. briggsae, Csi = C. sinica, Cre = C. remanei, Cwa = C. wallacei, Ctr = C. tropicalis, Cbn = C. brenneri, Cel = C. elegans, Cja = C. japonica, Caf = C. afra, Can = C. angaria. (D) Comparison of the Drosophila Fushitarazu/F1 (FTZ-F1) binding site, the NHR-binding site in wild-type C. elegans and lin-3 (e1417) mutant. At least two nucleotide changes are required to align putative NHR binding
sites from the Japonica group of the Caenorhabditis genus to the sequence in C. elegans and multiple changes are required for C. angaria.

(TIF)

S5 Fig. Distribution of bHLH cis-regulatory binding sites upstream the AC's specific lin-3 TSS in Caenorhabditis species. Location of transcription factor binding sites belonging to the bHLH protein family (as described in [51]) across DNA sequences upstream the TSS of the vulval form of lin-3 mRNA. The location of the NHR-binding site belonging to the lin-3 regulatory triplet is also depicted. Only the first 500 bp before the ATG are displayed for most of the species, except for C. virilis (only 300 bp) and C. sp. 1 (up to 1.9 Kb).

(TIF)

S6 Fig. Cross-species lin-3 transgenesis. (A) Wild-type C. elegans vulval invagination in the L4 stage as seen by Nomarski optics. (B) Over-expression of Can-lin-3(+) in C. elegans via transgenesis with repeated extra-chromosomal arrays results in vulval hyper-induction, with several additional invaginations in the L4 stage (arrowheads). (C) Injection of a Cel-lin-3 fragment in C. angaria leads to vulval hyperinduction. (D) Over-expression of a Caf-lin-3 fragment in C. elegans with repeated extra-chromosomal arrays leads to vulval hyperinduction.

(TIF)

S7 Fig. The Can-lin-3 CRM is able to drive specific expression in the anchor cell in C. elegans. (A) A promoterless C. briggsae fragment introduced into C. elegans is not expressed in N2. (B) The same fragment under the Can-lin-3 CRM drives expression in the anchor cell of N2, as monitored using Cbr-lin-3 FISH. Green corresponds to lin-3 expression and blue is DAPI staining of nuclei.

(TIF)

S8 Fig. Rescue of brood size defects by a single-copy insertion of Can-lin-3, in the homozygous or hemizygous state. A single-copy insertion of Can-lin-3(+) rescues brood size defects of lin-3(e1417) mutants (n>15). Note that the presence of a myo-2::GFP transgene linked to lin-3(e1417) in the background enhances the lin-3(e1417) brood size defects and does not allow rescue to wild-type brood size. Vulval induction in this experiment is presented in Fig 5C.

(TIF)

S1 Table. lin-3 mRNA single-molecule FISH quantification in the anchor cell in different C. elegans wild isolates (sheet A) and different Caenorhabditis species (sheet B). Each entry is an individual animal. Mean and standard deviation (SD) are indicated at the bottom.

(XLSX)

S2 Table. lin-3 mRNA single-molecule FISH quantification in the anchor cell of different C. elegans genotypes differing in their lin-3 cis-regulatory region, and corresponding vulval indexes (last sheet). The genotype is indicated in the name of the sheet. Gonad size of each individual is given in pixels. Experimental batch is indicated in the "Batch" column.

(XLSX)

S3 Table. Nematode strains used in this study.

(XLSX)

S4 Table. Sequences of oligonucleotides used in this study (except those for smFISH).

(XLSX)

S5 Table. Sequences of the oligonucleotides used as pools for smFISH experiments.

(XLSX)
Acknowledgments

We thank Lewis Stevens, Mark Blaxter and the Caenorhabditis Genomes Project, as well as Matt Rockman and Luke Noble for early access to unpublished genome assemblies. We are grateful to Jean-Louis Bessereau and Aurélien Richaud for advice and technical help on CRISPR replacements. We gratefully acknowledge John Reece-Hoyes and Marian Walhout for one-hybrid screens with lin-3 fragments from different species. Some strains were provided by the CGC, which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440). We also thank WormBase.

Author Contributions

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Writing – original draft: MB AMVV MAF.

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