Differential modulation of *C. elegans* motor behavior by NALCN and two-pore domain potassium channels

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Abstract

Two-pore domain potassium channels (K2P) are a large family of “background” channels that allow outward “leak” of potassium ions. The NALCN/UNC80/UNC79 complex is a non-selective channel that allows inward flow of sodium and other cations. It is unclear how K2Ps and NALCN differentially modulate animal behavior. Here, we found that loss of function (lf) in the K2P gene *twk-40* suppressed the reduced body curvatures of *C. elegans* NALCN(lf) mutants. *twk-40* (lf) caused a deep body curvature and extended backward locomotion, and these phenotypes appeared to be associated with neuron-specific expression of *twk-40* and distinct *twk-40* transcript isoforms. To survey the functions of other less studied K2P channels, we examined loss-of-function mutants of 13 additional *twk* genes expressed in the motor circuit and detected defective body curvature and/or locomotion in mutants of *twk-2*, *twk-17*, *twk-30*, *twk-48*, *unc-58*, and the previously reported *twk-7*. We generated presumptive gain-of-function (gf) mutations in *twk-40*, *twk-2*, *twk-7*, and *unc-58* and found that they caused paralysis. Further analyses detected variable genetic interactions between *twk-40* and other *twk* genes, an interdependence between *twk-40* and *twk-2*, and opposite behavioral effects between NALCN and *twk-2*, *twk-7*, or *unc-58*. Finally, we found that the hydrophobicity/hydrophilicity property of TWK-40 residue 159 could affect the channel activity. Together, our study identified *twk-40* as a novel modulator of the motor behavior, uncovered potential behavioral effects of five other K2P genes and suggests that NALCN and some K2Ps can oppositely affect *C. elegans* behavior.

Author summary

Two-pore domain potassium channels (K2P) allow outward “leak” of potassium ions. The NALCN channel complex allows inward flow of sodium and other cations. K2Ps and NALCN are implicated in severe human diseases. Using the nematode *Caenorhabditis*...
As a model to study the behavioral effects of K2P TWK-40 and NALCN, we found opposite effects of TWK-40 and NALCN on the body curvature. We identified five other K2P genes, including *twk-2, twk-17, twk-30, twk-48* and *unc-58*, as potential modulators of the motor behavior and confirmed the behavioral effect of the previously studied *twk-7*. These *twk* genes interacted genetically with *twk-40* in a variable manner, and *twk-2* exhibited an interdependent relationship with *twk-40*. In addition, *twk-2* and *unc-58* appeared to affect the body curvature in opposition to NALCN, and *twk-7* appeared to affect the locomotion in opposition to NALCN. Finally, we found that the chemical property of TWK-40 amino acid 159 can affect the channel activity. Together, we suggest that TWK-40 is a novel regulator of *C. elegans* motor behavior, five other K2Ps can potentially modulate the behavior, and some K2Ps and NALCN may oppositely affect the behavior.

**Introduction**

Ion channels are pore-forming membrane proteins that allow specific ions to pass through lipid membranes via the channel pore along the concentration gradient [1]. The major channel types include chloride channels, potassium channels, sodium channels, calcium channels, proton channels and non-selective cation channels. These channels open and close in a highly coordinated manner to regulate the action potentials and set the resting membrane potentials of neurons, muscles, and other cell types. However, much remains to be understood about how these channels differentially affect the behavioral output of a neural circuit.

The two-pore domain K⁺ channels (K2P) are a large group of “leak” channels containing the characteristic four transmembrane domains and two pore-forming domains (4T2P) [2]. K2P is broadly expressed and conserved across species. The human genome encodes 15 K2Ps [3] with diverse functions. The nematode *Caenorhabditis elegans* genome encodes 47 K2Ps named TWKs (TWiK family of potassium channels) [4], most of which have unclear functions. K2Ps are voltage-insensitive in general and can be regulated by pH, temperature, mechanical stimulation, lipids and other factors [2]. K2Ps play important roles in setting the resting membrane potentials of excitable cells [5] and are implicated in the pathophysiology of nociception, neuroprotection, vascular and pulmonary hypertension, cardiac arrhythmias, depression and cancer [2].

Different from K2Ps, NALCN (Na⁺ leak channel, non-selective) is a non-selective cation channel that allows inward passage of Na⁺ and potentially other cations [6–8]. NALCN is conserved across species and regulated by conserved proteins [8]. In mice NALCN affects the tonic firing and excitability of substantia nigra pars reticulata neurons [9], the stable network activity within the respiratory network [7,10], and the amount of rapid eye movement sleep (REMS) and non-REMS [11]. In *Drosophila*, NALCN is required for the proper coupling of locomotion with light and dark [12] and can regulate the circadian locomotion rhythms by affecting the neural output of the circadian pacemaker [13–16]. *C. elegans* has two largely redundant NALCN homologs, NCA-1 and NCA-2, that together regulate the recycling of synaptic vesicles [17], synaptic transmission at neuromuscular junctions [18], neural circuit activity [19], the response to volatile anesthetics [20], locomotory patterns [21], ethanol responses [22] and developmentally timed sleep and arousal [23]. The functional conservation of NALCN is further exemplified by the findings that both *Drosophila* and *C. elegans* NALCN mutants are more sensitive to the volatile anesthetic halothane [20].

NALCN can be regulated by a variety of signals. In mammals, NALCN is affected by substance P via the GPCR TACR1 [24], the Src kinase [25], the M3 muscarinic receptors [26], an
unidentified GPCR [27] and inhibitor GPCRs [28]. Studies in C. elegans suggest that NALCN channels are negatively regulated by dopamine through the DOP-3 receptor in command interneurons [29], are targets of Gq-Rho signaling in head acetylcholine neurons [30], and may also be modulated by the SEK-1 p38 pathway [31]. NALCN can interact with gap junctions [23,32,33].

Three proteins have been identified as specific regulators of NALCN. The ER-associated protein NLF-1 is required for proper axonal localization of C. elegans NALCN channels [34]. NALCN also physically interacts with the conserved proteins UNC79 and UNC80 (orthologs of C. elegans UNC-79 and UNC-80) [27], which are required for the channel function [8]. Loss-of-function mutations in NLF-1, UNC80 or UNC79 cause indistinguishable or highly similar phenotypes like NALCN loss-of-function mutations [15,17,18,20–22,24,25,27,35]. Mutations in NALCN and UNC80 are the cause of a group of human diseases called NALCN channelopathies [36].

We previously found that C. elegans exhibited a strong avoidance response to the odorant methyl salicylate [37]. A screen for new genes affecting this behavior identified multiple loss-of-function mutations in unc-79 and unc-80 [38]. To further understand the function of the NALCN channel, we performed a screen for suppressors of the “fainter” phenotype (slow locomotion and reduced body curvature) of an unc-80 loss-of-function mutant. The screen identified two loss-of-function (lf) mutations in the K2P gene twk-40. twk-40(lf) strongly suppressed the “reduced body curvature” phenotype of unc-80(lf) and other NALCN(lf) mutants. In addition, twk-40 can limit head touch-triggered backward locomotion. We further explored the functions of multiple other twk genes expressed in neurons of the motor circuit and identified five as potential modulators of the motor behavior. These twk genes genetically interacted with twk-40 or NALCN to affect the motor behavior in a variable manner. Together our results suggest that twk-40 is a novel modulator of C. elegans behavior and certain twk genes may affect the motor behavior in opposition to NALCN.

Results

twk-40(lf) mutations suppress the reduced body curvatures of NALCN-related loss-of-function mutants

We recently found that unc-80, unc-79 and NALCN genes were required for C. elegans avoidance to the odorant methyl salicylate [38]. To identify new genes potentially interacting with unc-80, we performed a screen for mutations that can suppress the reduced locomotion and/or reduced body curvature of unc-80(mac379) mutants (Fig 1A, top right panel). mac379 caused a G927R substitution and a W1524stop mutation in the UNC-80a protein and was characterized as a null allele of unc-80 [38] (unc-80(lf) hereafter).

The screen isolated five recessive suppressors (S1 Table, top). mac420, mac422 and mac424 weakly or moderately suppressed the defective forward locomotion of unc-80(lf) mutants (S1A Fig). The other two suppressors, mac425 and mac426, did not obviously suppress the defective basal locomotion of unc-80(lf) mutants or touch-triggered forward locomotion (S1A Fig), but significantly suppressed the reduced body curvature of unc-80(lf) mutants (Fig 1A and 1B). Using genetic analyses and/or SNP mapping, we located these mutations, except mac424, to chromosomes (S1 Table, top). We found that mac425 and mac426 probably affected a same gene on Chr. III (S1 Table, top).

We performed whole-genome sequencing on these mutants. A comparison of genes with deleterious variations on Chr. III found that twk-40 was the only one affected in both mac425 and mac426 mutants (S1 Table, bottom). The analyses of other mutants are ongoing.
twk-40 (Fig 1C and 1D) encodes a *C. elegans* K2P channel homologous to human KCNK9 (also known as TASK-3 or TASK3) and KCNK16 (also known as TALK-1 or TALK1) [3,39–41] (S2 Fig). twk-40 has three annotated isoforms, a, b, and c (Fig 1C) (www.wormbase.org). Exons are shown as boxes and introns as lines. Exonic regions encoding the four transmembrane domains and two pore domains are marked as blue and red, respectively. Positions of loss-of-function mutations are indicated on top. (D) TWK-40a protein domains. TM: transmembrane domain. P: pore domain. The two gain-of-function mutations, mac504 and mac505, affect TM2. (E) twk-40 transgenes rescued the suppression of unc-80(IFS) curvature defect by twk-40(IFS). 40 animals from each of three independent transgenic lines were quantified. (F) twk-40(IFS) suppressed the reduced curvature of unc-79(IFS), nca-2(IFS); nca-1(IFS) and nlf-1(IFS) mutants. 40 animals were quantified for each genotype. Statistics: Bonferroni multiple comparison with one-way ANOVA or two-tailed unpaired Student’s *t*-test. ***, *p* < 0.001.

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caused similarly deep curvature as in the double mutants (Fig 1A, bottom left panel and Fig 1B).

To verify the suppressor activity of \textit{twk-40}, we performed transgene rescue experiments (Fig 1E). A transgene predicted to express all \textit{twk-40} isoforms (\textit{Ptwk-40L::twk-40b\_gDNA}. See S3A Fig for promoter) significantly rescued the deep curvature of \textit{twk-40(mac425); unc-80(lf)} mutants (Fig 1E). In addition, a transgene predicted to express isoforms a and c (\textit{Ptwk-40R::twk-40a/c\_gDNA}. See S3A Fig for promoter) exhibited similar rescuing effects (Fig 1E). Together, these results suggest that \textit{twk-40(lf)} mutations can suppress the reduced curvature of \textit{unc-80(lf)} mutants. To simplify our analyses, we used \textit{twk-40(mac425)} as the reference loss-of-function allele hereafter.

Like \textit{unc-80(lf)}, loss-of-function mutations in \textit{unc-79}, \textit{nca-2; nca-1} (NALCN null) or \textit{nlf-1} all caused reduced curvatures. Similarly, \textit{twk-40(lf)} significantly suppressed the reduced curvatures of these mutants (Fig 1F). Previous studies suggest that \textit{nca-1} and \textit{nca-2} were largely redundant in affecting \textit{C. elegans} behaviors [18,20,21], while differential behavioral effects of these genes were also observed [22,30,38]. We found that \textit{nca-1} and \textit{nca-2} redundantly affected the curvature phenotype (S1B Fig).

twk-40 promoters are active in neurons

To identify the cells in which \textit{twk-40} might be expressed, we generated transgenic animals expressing GFP driven by a 5.8 kb \textit{twk-40} promoter (Figs 2A and S3A, \textit{Ptwk-40F}). The transgene labeled head neurons (Fig 2B, left panel), the ventral nerve cord (Fig 2B, left panel), motor neurons (Fig 2B, central panel) and tail neurons (Fig 2B, right panel).

To investigate whether shorter \textit{twk-40} promoters might exhibit neuronal specificity, we examined an upstream portion (S3A Fig, \textit{Ptwk-40L}, 2.5 kb, upstream of \textit{twk-40b} first exon) and a downstream portion (\textit{Ptwk-40R}, 2.8 kb, upstream of \textit{twk-40a} first exon, including the first exon of \textit{twk-40c}) of the \textit{Ptwk-40F} promoter. A \textit{Ptwk-40L::GFP} reporter and a \textit{Ptwk-40R::mCherry} reporter both labeled \textit{C. elegans} head neurons, the ventral nerve cord, and tail neurons (S3B and S3C Fig). However, it appeared that \textit{Ptwk-40R::mCherry} also labeled motor neurons (S3C Fig) while \textit{Ptwk-40L::GFP} did not (S3B Fig).

We next crossed the two transgenes into same animals (S4 Fig). In these animals, we failed to identify head neurons obviously co-labeled by the two reporters (S4A and S4B Fig). Interestingly, an unidentified tail neuron at the position of DVA/DVB/DVC was co-labeled (S4C Fig). Consistently, multiple motor neurons were labeled by \textit{Ptwk-40R::mCherry} but not by \textit{Ptwk-40L::GFP} (S4D Fig).

twk-40 promoters and an \textit{unc-80} promoter drive co-expression in some neurons

To examine whether \textit{twk-40} and \textit{unc-80} are co-expressed, we crossed a \textit{Ptwk-40L::mCherry} transgene (S5 Fig) or a \textit{Ptwk-40R::mCherry} transgene (S6 Fig) into animals with a \textit{Punc-80::GFP} transgene [38].

In animals co-expressing \textit{Ptwk-40L::mCherry} and \textit{Punc-80::GFP}, the reporters co-labeled some head neurons (Figs 2C and S5) and unidentified tail neuron(s) at the position of DVA/DVB/DVC (S5 Fig). Several co-labeled neurons appeared to be AVA (Figs 2C and S5A), AVB (S5A Fig), AVE (Figs 2C and S5A), AVG (Fig 2C) and RIS (S5B Fig).

In animals co-expressing \textit{Ptwk-40R::mCherry} and \textit{Punc-80::GFP}, the reporters co-labeled some unidentified head neurons (S6A and S6B Fig), unidentified tail neuron(s) at the position of DVA/DVB/DVC (S6C Fig) and a subset of motor neurons (S6D Fig).
We further crossed a Ptwk-40R::GFP or a Ptwk-40L::GFP transgene into animals with a Pnmr-1::mCherry transgene. In these animals, we failed to identify head neurons co-labeled by Ptwk-40R::GFP and Pnmr-1::GFP or a Ptwk-40L::GFP transgene into animals with a Pnmr-1::mCherry transgene. In these animals, we failed to identify head neurons co-labeled by Ptwk-40R::GFP and Pnmr-1::GFP (S7A Fig), including the AVD and RIM neurons known to be labeled by the Pnmr-1 promoter [42,43]. Interestingly, PVC neurons in the tail, which are major command interneurons for forward locomotion [44,45], were co-labeled by Ptwk-40R::GFP and Pnmr-1::mCherry (S7B Fig). Consistent with our earlier results (Fig 2C), Ptwk-40L::GFP and Pnmr-1::mCherry appeared to co-label AVA, AVE and AVG neurons (S7C Fig).
Considering that previous studies suggested the involvement of DVA and DVC neurons in locomotion [46–48], we tried to narrow down the unidentified tail neuron(s) labeled by the Ptwk-40 promoters at the position of DVA/DVB/DVC (S4, S5 and S6 Figs). We generated transgenic animals co-expressing Ptwk-40F::mCherry and PPtrp-4::GFP or Plim-6int::GFP. The PPtrp-4 promoter and the Plim-6int promoter were previously shown to label DVA/DVC neurons [49,50] and DVB neurons [51], respectively. Interestingly, Ptwk-40F::mCherry co-labeled DVA/DVC with PPtrp-4::GFP (S8A Fig) and sometimes DVB with Plim-6int::GFP (S8B Fig).

We next generated transgenic animals co-expressing Plim-6int::GFP with Ptwk-40L::mCherry or Ptwk-40R::mCherry. Here we found that Ptwk-40L::mCherry labeled DVA/DVC but not DVB (S8C Fig), while Ptwk-40R::mCherry labeled DVB/DVC but not DVA (S8D Fig).

Neuronal twk-40 is important for the expression of unc-80(If) curvature phenotype

To examine the neuron-specific activities of twk-40, we used different promoters to drive the expression of a twk-40b cDNA transgene in twk-40(If); unc-80(If) double mutants (Fig 2D). The twk-40b transgene driven by a Punc-119 pan-neuronal promoter [52] and five other promoters, including Ptwk-40L (S3A Fig), Ptwk-40R (S3A Fig), Punc-80 [38], Pnlf-1 [34], and Ptwnmr-1 [42], all exhibited significant rescuing effects (Fig 2D).

To understand whether TWK-40 isoforms might differ in channel activity, we tested a twk-40a cDNA transgene driven by the Ptwk-40F promoter, which is predicted to also express twk-40b and twk-40c, or by the Ptwk-40L promoter. These transgenes exhibited similar rescuing effects compared to the twk-40b cDNA transgenes (Fig 2D). Reasoning that twk-40a encodes the common channel region of all three TWK-40 isoforms, we chose twk-40a cDNA as the representative isoform in the following transgenic experiments.

The effect of neuronal twk-40 on the curvature of unc-80(If) mutants was also confirmed by neuron-specific knockdown experiments. Using a Punc-119:Cas9 transgene and two twk-40-specific sgRNAs to disrupt twk-40 coding sequences, we could suppress the reduced curvature of unc-80(If) mutants to a deeper one (Fig 2E). Cas9 driven by Ptwk-40L or Ptwnmr-1 exhibited similar effects (Fig 2E).

Differential effects of twk-40 promoters on the body curvature

Earlier transgene rescue experiments were primarily performed in twk-40(If); unc-80(If) double mutants (Figs 1E and 2D). To simplify the analyses of twk-40, we performed more rescue experiments in twk-40(If) single mutants using the twk-40a cDNA transgene driven by different promoters.

The first finding was that Ptwk-40F rescued the deep curvature of twk-40(If) mutants to the wildtype level (Fig 3A). This was different from the partial effect of this promoter on twk-40(If); unc-80(If) double mutants, in which the deep curvature was rescued to the wildtype level but not further to the unc-80(If) level (Fig 2D).

Ptwk-40L also rescued the deep curvature of twk-40(If) mutants close to the wildtype level (Fig 3A). However, Ptwk-40R exhibited a weaker rescuing effect, which was like Ptwnmr-1 (Fig 3A).

To examine whether other neurons in which twk-40 promoters were active are involved in the curvature phenotype, we performed transgenic rescue experiments using a Pacr-2 promoter (active in excitatory cholinergic motor neurons) [53], a Pcfi-1 promoter (active in PVC neurons) [54], the PPtrp-4 promoter and the Plim-6int promoter. However, none of these promoters exhibited an obvious rescuing effect on the curvature phenotype of twk-40(If) mutants (Fig 3A).
Differential effects of twk-40 transcript isoforms on the body curvature

To examine how twk-40 isoforms affect the curvature, we used the CRISPR/Cas9 method to generate deleterious mutations only affecting the twk-40b-specific exon, the twk-40c-specific exon, or both (S9 Fig), postulating that such mutations might reveal isoform-specific functions.

We obtained two independent mutants of each category (twk-40b, twk-40c, or both) (S9A Fig). Mutant strains of the same category exhibited indistinguishable phenotypes, and we quantified the curvature of one strain for each category.

\textit{twk-40(mac554)} (1 bp deletion in \textit{twk-40b} exon 1, S9 Fig) mutants exhibited a curvature deeper than wildtype animals but not as deep as \textit{twk-40(lf)} mutants (Fig 3B). However, \textit{twk-40 (mac557)} (61 bp deletion in \textit{twk-40c} exon 1, S9 Fig) mutants exhibited a wildtype-like curvature (Fig 3B). Interestingly, \textit{twk-40(mac558)} mutants (1 bp deletion in \textit{twk-40b} exon 1 and 10 bp deletion in \textit{twk-40c} exon 1, S9 Fig) exhibited a curvature slightly deeper than \textit{twk-40 (mac554)} mutants.

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While generating twk-40(mac558)-like mutations, we isolated twk-40(mac560) (2987 bp deletion spanning the start codons of twk-40b and twk-40c) (S9 Fig). mac560 might also affect twk-40a expression if the deletion disrupted twk-40a endogenous promoter. twk-40(mac560) mutants exhibited a deep curvature like twk-40(lf) mutants (Fig 3B).

So far, these isoform-specific mutant strains were not sufficient for analyzing the effect of twk-40a. To address this question, we used the CRISPR/Cas9 method to mutate the start codon of twk-40a to a codon for isoleucine (S9 Fig, mac564), hoping that this mutation would specifically disrupt TWK-40a expression. The lack of an obvious curvature defect in twk-40 (mac564) mutants (Fig 3B) suggests that this mutation did not significantly disrupt total TWK-40 channel activity, and that twk-40a was not critically involved in the curvature phenotype if the mutation specifically disrupted TWK-40a expression.

**twk-40 affects touch-triggered backward locomotion**

To identify other motor defects caused by twk-40(lf), we quantified touch-triggered forward (body bends/30 sec) or backward locomotion (body bends until stop). We found that tail touch triggered a slightly slower forward locomotion in twk-40(lf) mutants (Table 1, S1 Movie for wild type and S2 Movie for twk-40(lf) mutants). However, head touch triggered

| Genotype | Curvature index | Forward body bends/30 sec | Backward body bends | Notes |
|----------|----------------|---------------------------|---------------------|-------|
| WT       | 0.28 ± 0.07    | 30.0 ± 3.5               | 6.4 ± 1.5           |       |
| twk-40(lf) | 0.47 ± 0.11 *** | 28.2 ± 2.8 *           | 10.1 ± 2.4 ***      | Frequent backward locomotion |
| twk-2(lf) | 0.36 ± 0.08 *** | 29.5 ± 2.7              | 7.5 ± 1.8 *         | Frequent backward locomotion |
| twk-7(lf) | 0.26 ± 0.06    | 36.7 ± 2.8 ***          | 7.4 ± 1.6           |       |
| twk-17(lf) | 0.29 ± 0.07    | 25.2 ± 2.5 ***          | 8.7 ± 1.9 ***       | Longer wavelength |
| twk-30(lf) | 0.27 ± 0.06    | 31.3 ± 2.7              | 10.5 ± 1.7 ***      |       |
| twk-48(lf) | See notes      | 26.8 ± 2.3 ***          | 5.3 ± 1.3 *         | Frequent forward deep bending, brief forward coiling after backward locomotion, irregular curvature |
| unc-58(lf) | 0.46 ± 0.11 *** | 3.4 ± 3.1 ***            | 5.3 ± 8.1           |       |
| twk-2(lf); twk-40(lf) | 0.38 ± 0.10 *** | 29.4 ± 3.0               | 10.8 ± 2.3          |       |
| twk-7(lf) twk-40(lf) | 0.37 ± 0.08 *** | 32.2 ± 2.5 ***          | 10.7 ± 2.1          |       |
| twk-40(lf); twk-17(lf) | See notes | 15.6 ± 2.2 ***          | 10.9 ± 2.1          | Very deep curvature |
| twk-30(lf); twk-40(lf) | 0.45 ± 0.09    | 27.4 ± 2.8              | 10.5 ± 2.4          |       |
| twk-48(lf) twk-40(lf) | See notes | Delta turn/Coiler       | Coiler              | Very deep curvature, frequently coil |
| twk-40(lf); unc-58(lf) | See notes      | 13.6 ± 1.8 ***          | Coiler              | Very deep curvature |
| twk-40(mac505gf) | See notes | 0                      | 0                   | Paralyzed, flaccid |
| twk-2(gf) | See notes      | 0                        | 0                   | Paralyzed, flaccid |
| twk-2(lf); twk-40 (mac505gf) | 0.27 ± 0.07    | 14.9 ± 2.6 ***          | 2.3 ± 0.9 ***       | Slightly flaccid |
| twk-2(gf); twk-40(lf) | See notes      | 1.3 ± 1.1 ***            | 10.3 ± 2.8 ***      | Infrequent forward locomotion with normal curvature, frequent unprovoked backward locomotion with deep curvature |
| twk-40(mac504gf) | 0.15 ± 0.05    | 8.2 ± 4.3               | 3.2 ± 1.8           |       |
| twk-2(lf); twk-40 (mac504gf) | 0.35 ± 0.09 *** | 27.6 ± 2.3 ***          | 9.5 ± 2.0 ***       |       |

For single mutants, comparisons were made with wild type. For double loss-of-function mutants, comparisons were made with twk-40(lf) mutants. For twk-2(lf); twk-40 (mac505gf), twk-2(gf); twk-40(lf) or twk-2(lf); twk-40(mac504gf) double mutants, comparisons were made with twk-40(mac505gf), twk-2(gf) or twk-40(mac504gf) single mutants, respectively. 40 animals were measured for each data. Statistics: Bonferroni multiple comparison with one-way ANOVA or two-tailed unpaired Student’s t-test. *p < 0.05, **p < 0.001. Phenotypes that were difficult to quantify were described.

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significantly extended backward locomotion in twk-40(lf) mutants (6.4 ± 1.5 body bends for wild type, 10.1 ± 2.4 for twk-40(lf)) (Table 1, S3 Movie for wild type and S4 Movie for twk-40 (lf) mutants).

The extended backward locomotion of twk-40(lf) mutants was fully rescued by twk-40a cDNA transgenes driven by Ptwk-40F, and partially by Ptwk-40R or Pnmr-1 (Fig 3C). However, Ptwk-40L did not exhibit an obvious rescuing effect, and neither did Pacr-2, Pcfi-1, Pptrp-4 or Plim-6int (Fig 3C).

Interestingly, twk-40(mac564) mutants, presumptively twk-40a-specific, exhibited obviously extended backward locomotion, though it was weaker than twk-40(lf) mutants (Fig 3D). However, twk-40b-specific mac554 mutants, twk-40c-specific mac557 mutants and twk-40b/c-specific mac558 mutants all exhibited backward locomotion not significantly different from wildtype animals (Fig 3D). Only twk-40(mac560) mutants, which presumably disrupted the expression of all three twk-40 isoforms, exhibited extended backward locomotion like twk-40 (lf) (Fig 3D).

More twk genes affect C. elegans motor behavior

47 twk genes were annotated in the C. elegans genome (S3 Table) [4] (www.wormbase.org), the functions of most of which were unclear. The deep body curvature and extended backward locomotion of twk-40(lf) mutants prompted us to investigate whether other less studied twk genes might also affect C. elegans motor behavior.

To simplify our analyses, we searched the CeNGEN database (www.cengen.org) [55] for twk genes with obvious expression in neurons of the motor circuit. The motor circuit primarily includes backward premotor interneurons AVA/AVD/AVE, forward premotor interneurons AVB/PVC, excitatory backward motor neurons DA/VA, excitatory forward motor neurons DB/VB, and inhibitory motor neurons DD/VD [44,45]. The search identified 14 such twk genes, including twk-40 (S3 Table, genes labeled in red and blue). Though these data probably provided only partial pictures about the in vivo expression of twk genes, we postulated that they would be useful for prioritizing our analyses.

To investigate whether these motor circuit-expressed twk genes might affect the locomotion, we generated two independent presumptive loss-of-function mutations in each gene using the CRISPR/Cas9 method. We also generated a mutation in twk-3, which was included as a negative control because it was not obviously expressed in neurons of the motor circuit (S3 and S4 Tables). Comparison of the two loss-of-function mutants of each twk gene suggested that they exhibited indistinguishable or very similar phenotypes. Hence, we chose one mutant of each gene for further analyses (S4 Table, mutations in bold).

We found that six twk mutants, including twk-2, twk-7, twk-17, twk-30, twk-48, and unc-58, exhibited distinct changes in body curvature and/or touch-triggered locomotion (Table 1). Their phenotypes were briefly described as follows.

twk-2(lf) mutants exhibited a moderately deep curvature (S5 Movie) and a weakly extended backward locomotion. They also frequently switched to backward locomotion without being provoked.

twk-7(lf) mutants exhibited hyperactive forward locomotion (Table 1), consistent with previous findings [56,57].

twk-17(lf) mutants (S6 Movie) exhibited moderately slower forward locomotion, moderately extended backward locomotion, and a longer body wavelength.

twk-30(lf) mutants (S7 Movie) exhibited extended backward locomotion like twk-40(lf) mutants.
twk-48(lf) mutants exhibited an irregular curvature (S8 Movie). These mutants also frequently made deep forward turns, exhibited brief forward coiling upon completing backward locomotion, and exhibited weakly reduced forward and backward locomotion (Table 1).

unc-58(lf) mutants exhibited a deep curvature (S9 Movie) like twk-40(lf) mutants. These mutants also exhibited reduced forward and backward locomotion (Table 1). Previous studies found that gain-of-function mutations in unc-58 caused hyper-activation of body-wall muscles and dramatically perturbed locomotion [58,59].

To confirm that the observed phenotypes of these mutants were caused by loss-of-function mutations, we performed transgene rescue experiments. Indeed, the obvious phenotype of each mutant could be rescued by its respective wildtype transgene under control of the unc-119 promoter (S5 Table).

We failed to observe obvious behavioral changes in eight other twk(lf) mutants, including the twk-3(lf) mutant that was treated as a negative control (S6 Table). These twk mutations did not obviously enhance or suppress the curvature or backward locomotion of twk-40(lf) mutants (S6 Table).

Variable genetic interactions between twk genes

To understand how the twk genes with obvious behavioral effects interact genetically, we generated double mutants between twk-40(lf) and twk-2(lf), twk-7(lf), twk-17(lf), twk-30(lf), twk-48(lf) or unc-58(lf) (Table 1). Their phenotypes were summarized as follows.

twk-2(lf) or twk-7(lf) suppressed the deep curvature of twk-40(lf) mutants (S10 and S11 Movies), twk-30(lf) had no effect, and twk-17(lf) (S12 Movie), twk-48(lf) or unc-58(lf) enhanced.

twk-2(lf) or twk-30(lf) had no effect on the slightly reduced forward locomotion of twk-40(lf) mutants, twk-7(lf) improved it, and twk-17(lf) or unc-58(lf) enhanced it.

However, the extended backward locomotion of twk-40(lf) mutants was not obviously affected by other twk(lf) mutations (Table 1. See below for twk-48 and unc-58), suggesting a dominant effect of twk-40 on this behavior.

Interestingly, twk-48(lf); twk-40(lf) double mutants tended to coil or make delta turns upon head or tail touch (Table 1, S13 and S14 Movies). Coiling was also induced in twk-40(lf); unc-58(lf) double mutants by head touch but not by tail touch (Table 1, S15 and S16 Movies).

Presumptive gain-of-function mutations in twk genes can cause strong paralysis

The variable phenotypes of twk(lf) mutants raised a question as to how gain-of-function mutations in these genes might affect the behavior. To examine this, we used the CRISPR/Cas9 method to generate mutations that changed a residue located in the 2nd transmembrane domain inner helix of the channels (named TM2.6) to asparagine (N). A recent study found that similar substitutions can generate hyperactive K2P channels in vertebrates and C. elegans [60]. We were able to obtain such mutations for TWK-40a (Figs 1D, S2 and S10A, S2 Table), TWK-2, TWK-7, TWK-43, and UNC-58 channels (S4 Table and S10A Fig).

For TWK-40a, the L159N (TM2.6) mutation (mac505gf) (Fig 1D and S2 Table) caused a strong paralysis (Figs 4A and S10B, bottom left panel). The paralysis phenotype was fully phenocopied by twk-40a(mac505gf), cDNA transgenes driven by Ptwk-40F and Ptwk-40L, and partially by Ptwk-40R and Pnmr-1 (Fig 4B). Crossing twk-40(mac505gf) with unc-80(lf) (Fig 4A, top central panel) resulted in a similar paralysis (Fig 4A, bottom central panel and Fig 4C). Interestingly, the coiler phenotype of nca-1(gf) mutants (Fig 4A, top right panel) was suppressed to paralysis by twk-40(mac505gf) (Fig 4A, bottom right panel and Fig 4C).
Among the other four TM2.6>N mutations, TWK-2a(I208N), TWK-7(L309N) and UNC-58a(F332N) all caused strong paralysis (S7 Table and S10B Fig, referred as gf hereafter). The body shapes of twk-2(gf) and twk-7(gf) mutants were flaccid, like twk-40(mac505gf) mutants. The body shape of unc-58(gf) mutants was circled (S10B Fig) or in some cases curved as previously reported [60]. We failed to observe obviously abnormal behaviors in TWK-43a(M208N) mutants (S10A Fig, S4 and S7 Tables). The underlying mechanism was unclear, and it may be worth considering the possibilities that the mutation affected the expression, folding, transportation and/or degradation of TWK-43.

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**twk-40 and twk-2 may be partially interdependent**

To understand whether the paralysis phenotype of twk-40(gf) mutants might be affected by other twk genes, we crossed twk-40(mac505gf) with loss-of-function mutations in twk-2, twk-7, twk-17, and unc-58. Interestingly, twk-2(lf) partially suppressed the paralysis of twk-40(mac505gf) mutants to slow locomotion (Tables 1 and S7) (S17 and S18 Movies), while twk-7(lf), twk-17(lf) or unc-58(lf) failed to do so (S7 Table). The interaction between twk-2 and twk-40 was reciprocal, as twk-40(lf) also suppressed the paralysis of twk-2(gf) mutants to slow locomotion (Tables 1 and S7) (S19 and S20 Movies). However, twk-40(lf) did not obviously affect the paralysis of twk-7(gf) or unc-58(gf) mutants (S7 Table). Therefore, twk-40 and twk-2 might specifically depend on each other for full activities.

To examine whether twk-2 and twk-40 are expressed in same neurons, we generated transgenic animals co-expressing Ptwk-40F::GFP and Ptwk-2::mCherry (Ptwk-2 is a 3.1 kb twk-2 promoter upstream of the start codon). In these animals, we observed co-expression of the reporters in AVA and AVE neurons (S11A Fig), unidentified tail neuron(s) at the position of DVA/DVB/DVC (S11B Fig) and a subset of motor neurons (S11C Fig).

**Differential effects of twk genes on NALCN mutant phenotypes**

To investigate how these newly analyzed twk genes might interact genetically with NALCN-related genes, we crossed twk(lf) mutations with unc-80(lf) and twk(gf) mutations with nca-1(gf). We found that all twk(lf) mutations failed to improve the slow locomotion of unc-80(lf) mutants except twk-7(lf), which did it weakly (Table 2). Interestingly, twk-2(lf) or unc-58(lf), like twk-40(lf), suppressed the reduced curvature of unc-80(lf) mutants to deeper ones, while twk-7(lf) or twk-17(lf) failed to do so (Table 2).

Like twk-40(mac505gf), twk-2(gf), twk-7(gf) and unc-58(gf) all suppressed the coiler phenotype of nca-1(gf) mutants to paralysis phenotypes comparable with those of the individual twk (gf) mutants (Table 2 and S10B Fig).

**Table 2. Genetic interactions between twk and NALCN mutations.**

| Genotype | Phenotype | Curvature index | Forward body bends/30 sec |
|----------|-----------|----------------|---------------------------|
| unc-80(lf) |       | 0.16 ± 0.05 | 2.5 ± 1.4 |
| twk-40(lf); unc-80(lf) |       | 0.46 ± 0.09 *** | 3.5 ± 2.2 |
| twk-2(lf); unc-80(lf) |       | 0.35 ± 0.07 *** | 3.9 ± 2.7 |
| twk-7(lf); unc-80(lf) |       | 0.17 ± 0.05 | 9.9 ± 7.8 *** |
| unc-80(lf); twk-17(lf) |       | 0.19 ± 0.05 | 2.4 ± 1.2 |
| unc-80(lf); unc-58(lf) |       | 0.38 ± 0.07 *** | 3.5 ± 1.6 |
| nca-1(gf) |       |       | Coiler |
| twk-40(mac505gf); nca-1(gf) |       |       | Paralysis |
| twk-2(gf); nca-1(gf) |       |       | Paralysis |
| twk-7(gf); nca-1(gf) |       |       | Paralysis |
| nca-1(gf); unc-58(gf) |       |       | Circled paralysis |

40 animals were measured for each data. Statistics: Bonferroni multiple comparison with one-way ANOVA. *** p < 0.001. Comparisons were made with unc-80(lf).

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A novel form of TWK-40 gain of function

In generating the \textit{twk-40(mac505gf)} mutation, we obtained an in-frame deletion of I158 and L159 in TWK-40a (Fig 1D and S2 Table, mac504). \textit{twk-40(mac504)} mutants were sluggish and moderately flaccid (Fig 4A, middle left panel and Fig 4B). A \textit{twk-40a(mac504)} cDNA transgene driven by \textit{Ptwk-40F} caused a locomotion defect phenocopying that of \textit{twk-40(mac504)} mutants (Fig 4B). Hence, \textit{mac504} might cause a moderate gain of function in TWK-40.

We found that \textit{twk-40(mac504gf)} and \textit{unc-80(\text{lf})} together caused a strong paralysis (Fig 4A, middle central panel, and Fig 4C). Interestingly, \textit{twk-40(mac504gf)} suppressed the coiler phenotype of \textit{nca-1(gf)} mutants to a non-coiler phenotype with very deep curvature (Fig 4A, middle right panel) and these animals exhibited a moderate locomotion (Fig 4C). Like \textit{twk-40(mac505gf)}, the reduced curvature and defective locomotion of \textit{twk-40(mac504gf)} mutants were also suppressed by \textit{twk-2(\text{lf})} (Table 1).

TWK-40a activity might be affected by chemical properties of residue 159 and extracellular K$^+$ concentration

It is intriguing that the deletion of I158 and L159 would confer a hyperactivity in TWK-40a. We examined whether deletion of either residue alone would affect the channel activity using transgene phenocopy experiments.

We found that a \textit{twk-40a} cDNA (I158del) transgene driven by \textit{Ptwk-40F} failed to affect the forward locomotion of wildtype animals, so did the transgene with an L159 deletion (Fig 5).

Realizing that an L159N substitution in wildtype TWK-40a caused hyperactivity (Figs 4 and 5) and T161 in wildtype TWK-40a became T159 in TWK-40a(mac504gf) (I158-L159del) (S10A Fig), we examined whether the hyperactivity of TWK-40a(mac504gf) was related to T161 of the wildtype TWK-40a. In a \textit{twk-40a(I158-L159del)} cDNA transgene, we mutated T161 to asparagine (N) or lysine (K). Both mutations caused paralysis in the transgenic animals (Fig 5). Interestingly, a T161I mutation in the transgene failed to do so (Fig 5).

![Fig 5. Effects of amino acid deletions and/or substitutions on TWK-40a activity.](https://doi.org/10.1371/journal.pgen.1010126.g005)
We next examined whether substituting L159 with amino acids of different chemical properties might affect wildtype TWK-40a channel activity. Indeed, both L159D and L159K substitutions in TWK-40a caused strong paralysis (Fig 5). However, L159I or L159V substitution failed to do so (Fig 5). Together, these results suggest that the hydrophobicity/hydrophilicity of TWK-40a residue 159 may affect channel activity.

Sub-physiological extracellular K\(^+\) concentration is predicted to cause neuronal hyperpolarization by inducing K\(^+\) efflux, which might lead to defective locomotion. We tested this possibility in *C. elegans*. As expected, wildtype animals exhibited reduced locomotion on NGM plates in which K\(^+\) was substituted with Na\(^+\) (S12 Fig). We next explored how *twk-40* might affect this behavior.

In K\(^+\)-deficient plates, *twk-40(lf)* mutants also exhibited reduced locomotion (S12 Fig). Surprisingly, *twk-40(mac504gf)* mutants showed weakly improved locomotion, both forward and backward, in K\(^+\)-deficient plates (S12 Fig). Considering that human K2P TWIK-1 can conduct inward Na\(^+\) leak at sub-physiological extracellular K\(^+\) concentrations or acidic pH [61–63], further analyses of TWK-40(WT) and TWK-40(mac504gf) at low extracellular K\(^+\) concentration might provide new insights into TWK-40 channel activity.

Discussion

In this study, we found that loss of function in the K2P gene *twk-40* suppressed the reduced body curvatures of *C. elegans* NALCN loss-of-function mutants. Neuron-specific *twk-40* expression and *twk-40* transcript isoforms might differentially affect the curvature and backward locomotion. We provide evidence that five other *twk* genes, including *twk-2*, *twk-17*, *twk-30*, *twk-48*, and *unc-58*, may also modulate *C. elegans* behavior. These *twk* genes, together with the previously described *twk-7*, exhibited variable genetic interactions with *twk-40*. Specifically, *twk-2* and *twk-40* might depend on each other for full activities. Like *twk-40*, *twk-2*, *twk-7*, and *unc-58* appeared to affect specific behaviors in opposition to NALCN. Finally, we detected a correlation between the hydrophobicity/hydrophilicity of TWK-40a residue 159 and the channel activity.

K2Ps and NALCN can have opposite effects on animal behavior

K2P channels are key regulators of resting membrane potentials by conducting “leak” outward K\(^+\) current. The activities of K2Ps are affected by a variety of physiological stimuli [2]. NALCN has an opposite effect on resting membrane potentials by conducting primarily inward monovalent cations [6,8], and NALCN activity can be affected by different GPCRs [8,28,29]. Previous studies suggested the existence of a “leak” K\(^+\) channel as a balancing force for NALCN in regulating the spontaneous firing rate of mouse SNr neurons [9] or the rhythmicity of *Drosophila* clock neurons [13]. Alternatively, NALCN could regulate the excitability of pH-sensitive neurons of mouse retrotrapezoid nucleus in opposition to the background “leak” K\(^+\) channels [64]. These findings support the hypothesis that NALCN and certain “leak” K\(^+\) channels might have counterbalancing effects on resting membrane potentials and raise the question whether the “leak” K\(^+\) channels might be K2Ps.

Recently, Kasap et al. [33] found that pharmacological inhibition of different K\(^+\) channels, including K2Ps, can significantly improve the locomotion of NALCN(lf) *C. elegans* mutants, providing an early piece of evidence that NALCN and K2Ps might act as opposite regulators of animal behaviors. Our results are consistent with their findings. The opposite effect was observed between *twk-40(lf)* and NALCN(lf) and was corroborated by the suppression of the coiler phenotype of *nca-1(gf)* mutants by *twk-40(gf)*. Opposite effects on specific behaviors were observed between NALCN and *twk-2*, *twk-7*, or *unc-58* as well. Besides K2Ps, other
channels may also affect the behaviors regulated by NALCN, e.g., increasing Ca$^{2+}$ channel activity pharmacologically improved the locomotion of NALCN(lf) mutants [33]. However, the molecular mechanisms underlying the behavioral effects of these channels are largely unclear and our findings should not be interpreted that TWK-40 or other TWK channels can affect neuronal activities in opposition to NALCN. To address these questions, we need to identify the sites of action of these channels and analyze their effects on neuronal activities by multidisciplinary approaches.

twk-40 activities are associated with its neuron-specific expression

It is intriguing that the Ptwk-40L and Ptwk-40R promoters, which appeared to be active in different neurons, exhibited similar rescuing activities for twk-40(lf) suppression of unc-80(lf) curvature. They also showed rescuing activities for the deep curvature of twk-40(lf) single mutants, though Ptwk-40L appeared to be stronger. However, Ptwk-40L and Ptwk-40R exhibited quite different rescuing activities for the extended backward locomotion of twk-40(lf) mutants: Ptwk-40R was effective while Ptwk-40L was not. We recently made a similar observation about the differential effects of promoters on behaviors in our study of C. elegans avoidance response to the odorant methyl salicylate [38]. In this study, we found that a portion (P$_3$unc-79) of the full-length unc-79 promoter (P$_1$unc-79) exhibited a rescuing activity like P$_1$unc-79 for the avoidance defect of unc-79(lf) mutants. However, P$_3$unc-79 failed to rescue the locomotion defect of unc-79(lf) mutants, which could be strongly rescued by P$_1$unc-79.

Another intriguing finding was that the Pnmr-1 promoter showed rescuing activities for both the curvature and backward locomotion phenotypes of twk-40(lf) mutants. However, Pnmr-1 appeared to be active in a set of neurons that only partially overlapped with those for Ptwk-40L, e.g., in AVA/AVE/AVG, or with Ptwk-40R, e.g., in PVC.

To elucidate the underlying mechanism of these phenomena is beyond the scope of our current study. Nevertheless, it is tempting to speculate a scenario that might accommodate these findings. In this scenario, NALCN might affect an array of behaviors by regulating the activity of a neural network. A component of the network may affect a specific behavior redundantly with other components, or only affect component-specific behaviors, or both. TWK channels, through restricted expression and regulated activities, might be involved in compartmentalizing the expression of some behaviors oppositely regulated by NALCN.

We primarily used genetic and behavioral experiments to analyze the effects of NALCN and twk genes. We found that several tail neurons in which twk-40 promoters were active, including PVC, DVA, DVB and DVC, did not obviously affect the curvature or backward locomotion. However, our analyses were insufficient to reveal whether one or more of the head neurons in which TWK-40 and NALCN were co-expressed, e.g., AVA, AVB and AVE, can affect these behaviors or were the sites where TWK-40 and NALCN exert opposite behavioral effects. Future experiments that can directly measure the activities of these neurons, e.g., calcium imaging or electrophysiology experiments, will provide insights into how TWK-40 and NALCN affect neuronal activities. Also, optogenetic approaches will be important for associating the behaviors affected by these channels with the neurons expressing them.

twk-40 transcript isoforms exhibit differential effects on the behavior

That Ptwk-40L (upstream of the twk-40b isoform) exhibited a stronger rescuing effect on the curvature phenotype of twk-40(lf) mutants while Ptwk-40R (upstream of the twk-40c and twk-40a isoforms) exhibited a stronger rescuing effect on the backward locomotion phenotype implies that twk-40 isoforms might have functional differences. We examined this possibility using presumptive isoform-specific twk-40 mutants.
For the curvature phenotype, twk-40(mac564) (twk-40a-specific) or twk-40(mac557) (twk-40c-specific) had no obvious effect. However, twk-40(mac554) (twk-40b-specific) caused a moderately deep curvature, and with the disruption of more isoforms, e.g., in twk-40(mac558) (disruption of twk-40b and twk-40c) and twk-40(mac560) (disruption of all three isoforms), the mutants exhibited gradually increased deep curvature. Based on these analyses, we speculate that the curvature is more significantly affected by twk-40b and less by twk-40a or twk-40c.

For the backward locomotion phenotype, twk-40(mac554) (twk-40b-specific), twk-40(mac557) (twk-40c-specific) or twk-40(mac558) (disruption of twk-40b and twk-40c) had no obvious effect. However, twk-40(mac564) (twk-40a-specific) caused a moderately extended backward locomotion, and twk-40(mac560) (disruption of all three isoforms) caused a more severe phenotype. Therefore, the backward locomotion appeared to be more significantly affected by twk-40a and less by twk-40b or twk-40c. Together, the isoform-specific mutant phenotypes are consistent with the transgene rescue results, suggesting differential behavioral effects of twk-40 isoforms.

Caution needs to be exercised in interpreting the results of the transgenes and isoform-specific mutants. For example, transgenes may cause non-specific expression in neurons, and the promoters may not be complete, and the fluorescent reporters probably did not provide a full picture about twk-40 expression. In addition, the expression patterns of twk-40 isoforms should not be extracted simply from the reporters, as the promoters may regulate expression of adjacent isoforms as well as distant ones. It is also possible that the presumptive twk-40 isoform-specific mutations may affect the expression of other isoforms, as they were within the full-length promoter region and/or might affect all isoforms. Therefore, new reagents and methods that provide higher-resolution and more accurate pictures about twk-40 expression, e.g., TWK-40 isoform-specific antibodies, isoform-specific knockin of reporters or single-copy transgene experiments, are warranted for validating our findings.

twk genes may fine tune the motor behavior of C. elegans

C. elegans genome contains 47 K2P channel encoding genes [4] (www.wormbase.org). The functions of most TWK channels are largely unclear, with the exception of TWK-18 [65], SUP-9 (TWK-38) [66], TWK-7 [56,57], EGL-23 [59] and UNC-58 [58,59]. TWK-18 can be activated by high temperature [65]. SUP-9 was regulated by the transmembrane proteins UNC-93, SUP-10 and the iodotyrosine deiodinase SUP-18 [66–68]. TWK-7 affected several aspects of adaptive locomotion behavior downstream of the GoS-KIN-1/PKA pathway [56,57]. Gain-of-function mutations in EGL-23 caused egg-laying defects [59,69], while loss-of-function mutations in EGL-23 probably did not cause obviously abnormal behaviors [59]. Gain-of-function mutations in UNC-58 caused a dumpy and spasm phenotype due to hyper-activation of the body wall muscles [58,59]. However, loss-of-function phenotype of unc-58 was unclear.

Prompted by the findings on twk-40 in the early phase of this study, we surveyed the functions of 13 twk genes expressed in neurons of the motor circuit [55] (www.cengen.org) by generating loss-of-function mutations in them. Focusing on the body curvature and touch-triggered forward or backward locomotion, we found that five such twk genes, including twk-2, twk-17, twk-30, twk-48, and unc-58, might be modulators of motor behavior. Our study also confirmed the hyperactive forward locomotion of twk-7(lf) mutants as previously described [56,57].

The behavioral patterns exhibited by these twk(lf) mutants are often displayed by wildtype animals under different circumstances [70]. Therefore, we may speculate that activation or inhibition of these TWK channels might provide dynamic means for modulating C. elegans behavioral patterns.
motor behavior. However, our analyses were mostly descriptive and qualitative. Extensive future studies are needed for understanding the molecular mechanisms underlying the effects of these \textit{twk} genes.

\textbf{An interdependence between \textit{twk-40} and \textit{twk-2} in modulating the locomotion}

We found that loss-of-function mutations in \textit{twk-2} or \textit{twk-40} can partially suppress the paralysis phenotype caused by gain-of-function mutations in the other. However, \textit{twk-40} did not interact with other \textit{twk} genes in such a manner. During the study, we also performed a screen for mutations that can suppress the paralysis phenotype of \textit{twk-40/mac505gf} mutants (Materials and Methods). From the screen, we isolated a potential loss of function mutation in \textit{twk-2} predicted to change TWK-2 G181 to glutamate (E), and G181 is a conserved residue within selectivity filter 1 of K2P channels [60]. Realizing that \textit{Ptwk-40F} and \textit{Ptwk-2} promoters were both active in AVA, AVE, and other neurons, we postulate that \textit{twk-40} and \textit{twk-2} might function in same neurons for their full activities.

\textbf{Gain-of-function mutations in \textit{twk} genes provide new insights into channel activities}

Recently, Ben Soussia et al. found that changing TM2.6 to asparagine can generate hyperactivity in multiple K2Ps [60]. We generated similar mutations in TWK-40, TWK-2, TWK-7, and UNC-58, and found that they caused strong paralysis. The paralysis phenotype of \textit{unc-58(gf)} mutants confirmed that of \textit{unc-58(gf)} mutants generated by Ben Soussia et al. [60]. We also isolated an in-frame mutation \textit{(mac504gf)} that deleted I158 (TM2.5) and L159 (TM2.6) of TWK-40a. \textit{mac504gf} caused a moderately hyperactive channel. Amino acid substitutions in wildtype TWK-40a and TWK-40(mac504gf) channels and transgene phenocopy analyses suggest that the residue at TM2.6 can significantly affect channel activity. Specifically, hydrophilic residues appeared to be related to hyperactivity while hydrophobic residues appeared to be related to inactivity or highly regulated activity. Our results are consistent with recent findings that such a hydrophobic barrier on the mammalian K2P TWIK1 can oppose efficient K$^+$ passage through the channel pore [71,72], and substituting the hydrophobic residue at TM2.6 of K2P channels with different amino acids can modulate the channel activities [60]. In future, detailed structural and electrophysiological analyses are warranted for understanding how these amino acid affect TWK-40 channel activity.

\textbf{A subset of \textit{twk} genes might be recently evolved to modulate the motor behavior}

We illustrated the expression patterns of the \textit{twk} genes with obvious behavioral effects in the motor circuit (Fig 6A). It appeared that there was a correlation between \textit{twk}-expressing neurons and the phenotypes of the \textit{twk(lf)} mutants. For example, among the \textit{twk} genes with expression in interneurons (Fig 6A, \textit{twk-40}, \textit{twk-2}, \textit{twk-7}, and \textit{twk-17}), extended backward locomotion was observed if the expression was in AVA neurons (\textit{twk-40}, \textit{twk-2}, and \textit{twk-17}), while reduced forward locomotion was observed if the expression was in AVB neurons (\textit{twk-40} and \textit{twk-17}). Alternatively, deep curvature was observed for \textit{twk} genes expressed in AVA/AVE neurons (\textit{twk-40} and \textit{twk-2}). However, \textit{twk} expression in interneurons might not be required for the mutants to show these phenotypes: loss-of-function mutations in \textit{twk-30}, \textit{twk-48}, and \textit{unc-58}, genes not obviously expressed in these interneurons (based on CeNGEN), resulted in similar phenotypes (Fig 6A).
To understand the evolutionary relationship of \textit{twk} genes, we built a phylogenetic tree of 47 \textit{C. elegans} TWK proteins (Fig 6B). We found that TWK-40, TWK-2, TWK-7, TWK-17, TWK-30, and UNC-58 were all evolved from the same major branch of the tree (Fig 6B, red box). Among them, TWK-2, TWK-7, and TWK-40 were derived from two most recent evolutionary divergence. Though evolutionary convergence could also lead to such a phylogenetic pattern, it is tempting to speculate that these \textit{twk} genes might have been generated from recent gene duplication events.

Mammalian homologs of TWK-40 are associated with diseases

TWK-40 is homologous to human KCNK16 and KCNK-9. KCNK16 was found to be exclusively expressed in human pancreas [39]. A nucleotide polymorphism in KCNK16 was associated with type 2 diabetes [73]. KCNK16 can negatively modulate human and mouse \(\beta\)-cell excitability and second-phase insulin secretion, probably through altering ER Ca\(^{2+}\) homeostasis and stress [74,75]. In retrospect, NALCN was shown to be required for M3 muscarinic receptor-activated inward current in a mouse MIN6 \(\beta\)-cell line [26]. These findings and our results together imply that NALCN and KCNK16 might counterbalance the excitability and insulin secretion of pancreatic \(\beta\) cells.

KCNK9 is highly expressed in the human brain [76,77]. NALCN is also broadly expressed in the nervous system [7,9,13,64] and can affect various neuronal activities [78]. Mutations in \textit{NALCN} or \textit{UNC80} are related to a group of diseases named “NALCN channelopathies” [36],...
which are characterized by infantile hypotonia, psychomotor retardation, characteristic facies, congenital contractures of the limbs and face, and developmental delays. A loss-of-function mutation in the imprinted \textit{KCNK9} locus was found to be related to the maternally inherited Birk-Barel syndrome \cite{79,80}, which is characterized by congenital hypotonia, variable cleft palate, delayed development, and feeding problems. The similarity between the two types of diseases implies a convergent pathogenetic mechanism involving \textit{KCNK9} and \textit{NALCN}.

**Conclusions**

In short, we identified \textit{twk-40} as a novel modulator of \textit{C. elegans} motor behavior and suggest that the behavioral effects of \textit{twk-40} might be associated with its neuron-specific expression and functional differentiation of transcript isoforms. Two recent studies further analyzed TWK-40 functions in regulating the resting membrane potentials and activities of AVA and DVB neurons \cite{81,82}. We also uncovered potential behavioral effects of five other \textit{K2P} genes and provide genetic evidence that some \textit{K2Ps} and \textit{NALCN} can oppositely affect different aspects of \textit{C. elegans} motor behavior. Understanding the relationship between \textit{K2Ps} and \textit{NALCN} might provide novel insights into the pathogenesis of a spectrum of diseases caused by mutations in \textit{NALCN} and \textit{K2Ps}.

**Materials and methods**

**Screening for \textit{unc-80(lf)} suppressors and mapping of mutations**

Synchronized \textit{unc-80(lf)} L4 animals were mutagenized with EMS (ethyl methanesulfonate) \cite{58}. 400 \(P_0\) animals were picked to 20 NGM plates, with 20 animals per plate. We estimated that each \(P_0\) animal would lay at least 100 eggs within 24 hrs of becoming young adult. After these first-day \(F_1\) eggs (~40,000 in total) became young adults, they were bleached to generate synchronized \(F_2\) progeny. Once growing into young adults, \(F_2\) animals were examined under a dissecting microscope to identify individuals with apparently improved locomotion or body curvature. From the screen, we obtained 5 independent isolates.

All five suppressors were recessive. \textit{mac420} and \textit{mac422} were assigned to Chr. X because \(F_1\) male progeny from the cross between \textit{unc-80(lf)} males and \textit{unc-80(lf)}; \textit{mac420} or \textit{unc-80(lf)}; \textit{mac422} hermaphrodites were suppressed for the defective locomotion, while \(F_1\) hermaphrodite progeny were not. \(F_1\) males from the cross between \textit{unc-80(lf)} males and \textit{unc-80(lf)}; \textit{mac424/mac425/mac426} hermaphrodites were not suppressed for their respective phenotype, suggesting that \textit{mac424}, \textit{mac425} and \textit{mac426} were autosomal. Complementation test suggested that \textit{mac420} and \textit{mac422} might affect different genes. We did not further map \textit{mac420} and \textit{mac422} using SNPs. \textit{mac425} and \textit{mac426} similarly suppressed the reduced body curvature of \textit{unc-80(lf)} mutants and did not complement each other, suggesting that they might affect a same gene. We mapped \textit{mac425} using published single nucleotide polymorphisms (SNP) \cite{83}.

To generate mapping lines, we crossed Hawaiian (CB4856) males with \textit{unc-80(lf)}; \textit{mac425} hermaphrodites to generate \(F_1\) cross progeny (normal curvature), and their \(F_2\) progeny with reduced curvature, which were probably of \textit{unc-80(lf)}; \textit{mac425} or \textit{unc-80(lf)}; +/- genotype, were picked to individual plates. Individual \(F_3\) progeny with suppressed curvature from different \(F_2\) plates were established as mapping lines.

In 10 mapping lines for \textit{mac425}, homozygous CB4856 genotypes were not detected at \(W06F12\) (Chr. III: 21 cM) or \(Y17D7B\) (Chr. V: 18 cM. Note: \textit{unc-80} is at Chr. V: 26 cM) but were detected at other selected SNPs in at least one line. At the same time, genomic sequencing of unoutcrossed \textit{mac425} and \textit{mac426} mutants found that \textit{twk-40} was the only gene on Chr. III affected in both strains. Hence we pursued \textit{twk-40} as the candidate gene for \textit{mac425} and \textit{mac426}.

The mapping of \textit{mac424} is ongoing.
Screening for twk-40(mac505gf) suppressors

We mutagenized twk-40(mac505gf) P₀ animals at the L4 larval stage with EMS. From synchronized F₁ adult progeny of ~40,000 F₁ progeny, we isolated four recessive mutants with improved locomotion. Three isolates exhibited locomotion and body curvature like twk-40(lf) mutants and failed to complement twk-40(lf). We postulated that these isolates probably carried intragenic loss-of-function mutations in twk-40 and did not further analyze them.

In earlier study, we found that twk-2(lf) could suppress the locomotion of twk-40 (mac505gf) mutants (Table 1). We crossed males of the fourth isolate with twk-2(lf); twk-40 (mac505gf) hermaphrodites and found that the male progeny were still suppressed for the locomotion phenotype. Sanger sequencing identified a mutation in twk-2 (named mac537). mac537 caused a G181E change in TWK-2a. TWK-2a G181 is a conserved residue in selectivity filter 1 of K2P channels [60].

Whole-genome sequencing

Each of the original isolates, CSM740 (unc-80(lf); mac420), CSM742 (unc-80(lf); mac422), CSM757 (unc-80(lf); mac424), CSM758 (unc-80(lf); mac425) and CSM759 (unc-80(lf); mac426) before outcrossing, was grown on a single 9 cm NGM plate and washed from the plates before food was completely consumed. Animals were washed twice with M9, resuspended in M9 and starved for several hours. Genomic DNAs were extracted by proteinase K digestion, followed by RNase A treatment and two rounds of phenol–chloroform extraction. Three genomic DNA libraries (380-bp inserts) were constructed by Berry Genomics Co., Ltd (Beijing) using Illumina’s paired-end protocol and paired-end sequencing (100-bp reads) was performed on the Illumina HiSeq 2000. Over 4G clean bases were mapped to the N2 genome (Wormbase release 220) after removal of duplicated reads. SNP calling was performed using Genome Analysis Toolkit (GATK) with the N2 genome as reference. 4449 (mac420), 4623 (mac422), 4633 (mac424), 4712 (mac425), and 4675 (mac426) sequence variants were detected in these mutants. Sequence variants shared among strains were excluded as they were likely derived from common ancestors. To enhance the stringency for mutation identification, we set the depth of reference base (WT) to be < 6 in these mutants. Exon or splice site variants carrying effective mutations with a variant quality greater than 30 were selected. Based on these criteria, we obtained 32 (mac420), 45 (mac422), 46 (mac424), 62 (mac425), and 74 (mac426) variants overall. The potential target genes for each mutant were further narrowed down to the mapped chromosomes.

mac425 and mac426 were mapped to Chr. III and might affect a same gene. Comparing sequence variants on Chr. III identified twk-40 as the only gene that was affected in both mutants.

Quantification of locomotion (body bends) and body curvature

Synchronized L4 animals were transferred to a new NGM plate with thin bacterium lawn and allowed to grow for 12 to 18 hours. A body bend is defined as a forward head turn or backward tail turn. Forward body bends were measured for 30 or 60 seconds after the tail of an animal was gently touched with a worm pick to trigger locomotion. Backward body bends were measured after the head of an animal was gently touched to trigger locomotion until the animal stopped or resumed forward locomotion.

Body curvature was measured based on a previously described method [30] with modification. The curvature index is the ratio of the depth (amplitude) to the period of the body waveform (Fig 1A). Young adults 18–24 hours after the mid L4 larval stage were quantified.
Transgene experiments

Germline transgene experiments were performed as described [84].

For *twk-40* rescue experiments, the transgenic mixtures contained 20 ng/µl pPD95_86 (Pmyo-3::GFP) plasmid as co-injection marker and 20 ng/µl of the transgene of interest.

For transcriptional reporters, a transgene solution containing 20~50 ng/µl of the reporter plasmid was injected to wildtype animals.

For neuron-specific rescue experiments, the transgenic mixtures contained 10~30 ng/µl Promoter::twk-40_cDNA and 20 ng/µl pPD95_86 (Pmyo-3::GFP) plasmid or 2.5 ng/µl pCFJ90 (Pmyo-2::mCherry) plasmid [85] as co-injection marker.

We used a previously described method [86] with modifications to generate neuron-specific knockouts of *twk-40*. The transgenic mixtures contained 25 ng/µl Promoter::Cas9::NLS::3'UTR, 25 ng/µl PU6::twk-40_sgRNA #1 and #2, and 20 ng/µl pPD95_86 (Pmyo-3::GFP) plasmid as co-injection marker.

For phenocopy experiments, transgenic mixtures contained 10~20 ng/µl Promoter::twk-40a (mutations) cDNA and 20 ng/µl pPD95_86 (Pmyo-3::GFP) plasmid as co-injection marker.

Generation of *twk* mutations using the CRISPR/Cas9 method

We followed the method [87] with modifications. Plasmids for microinjection were purified using Omega’s Midi Plasmid Purification kit (Omega Bio-tek). For *twk-40* insertion/deletion mutation, the injected DNA mixture contained 50 ng/µl pPD162 (Peft-3::Cas9-SV40_NLS), 25 ng/µl PU6::sgRNAs (two specific sgRNAs for a *twk* gene), and 20 ng/µl pPD95_86 (Pmyo-3::GFP) plasmid as co-injection marker. To generate *twk-40* knockin mutations, we included a synthesized oligo (500 nM) containing the designed mutation as repair template in the injection mixture.

We normally injected 20 to 40 wildtype *P*₀ animals for knockout or knockin mutations. For *twk-40(mac469)* and *twk-40(mac472)*, we injected unc-80(lf) *P*₀ animals. ~40 F₁ animals with strong GFP signals in body-wall muscles (co-injection marker) were picked to individual plates, and F₂ progeny were sequenced until the expected mutations were detected.

Two approaches were taken to generate mutations in other 14 *twk* genes. For each of *twk-2*, *twk-3*, *twk-7*, *twk-13*, *twk-17*, *twk-43*, *twk-46* and *unc-58*, knockout or knockin mutations were generated using an injection solution containing the aforementioned injection mix together with a repair template. We obtained two or more deletion/insertion mutations for each of the eight genes, one knockin mutation for *twk-2*, *twk-7* or *twk-43*, three knockin mutations for *unc-58*, and no knockin mutation for *twk-3*, *twk-13*, *twk-17* and *twk-46*. For *twk-26*, *twk-29*, *twk-30*, *twk-35*, *twk-48* and *twk-49*, we generated deletion/insertion mutations using an injection solution without repair template (S9 Table). Separate injection and examination were performed for each gene.

Target sequences of sgRNAs are shown in S9 Table and the sequences of repair templates for *twk-40*, *twk-2*, *twk-7*, *unc-58* and *twk-43* are shown in S10 Table. The numbers of lines examined by sequencing and the numbers of knockout lines or knockin lines obtained are shown in S9 Table.

Confocal microscopy and identification of *twk-40*-expressing cells

Confocal pictures of fluorescent transgenic animals were taken with a Zeiss 880 confocal microscope (10X and 63X objective). Multiple animals from each of two transgenic lines were observed for consistency of reporter expression and a subset of animals with strong fluorescence were examined further for neuronal identification. Pictures shown were representative images. We determined the identities of *twk-40*-expressing cells labeled with *Ptwk-40::reporters*
by examining the anatomical positions, morphologies, patterns and neighboring cells and comparing to the descriptions at www.wormatlas.org. The identities of some neurons were further verified by co-labeling experiments with reporters driven by previously described promoters.

**Potassium-deficiency assay**

NGM plates were prepared by replacing K$_2$HPO$_4$/ KH$_2$PO$_4$ with equal molar amount of Na$_2$HPO$_4$/ NaH$_2$PO$_4$. Synchronized L1 animals were allowed to grow on the K$^+$-deficient plates until reaching the L4 larval stage. The animals were then transferred to fresh K$^+$-deficient NGM plates with thin bacterium lawn. Touch-triggered body bends were measured in 18 to 24 hours.

**Neuronal expression pattern based on CeNGEN**

We followed the instruction at www.cangen.org to search for the graphed expression of each twk gene in *C. elegans* neurons. The top 20 neuronal classes in which a twk gene was expressed were listed in S4 Table.

**Phylogenetic tree**

Maximum likelihood phylogenetic tree was constructed with the Molecular Evolutionary Genetics Analysis software MEGA X [88].

**Statistics**

P values were determined by two-tailed unpaired Student’s t-test for pairwise comparison or Bonferroni multiple comparison with one-way ANOVA for multiple comparison using GraphPad Prism 7.0 software.

Strains used in this study and plasmid constructions are shown in S1 Text. Raw data are shown in S1 Data.

**Supporting information**

**S1 Fig. Behavioral quantification of unc-80(lf) suppressors and nca(lf) mutants.** (A) Touch-triggered forward locomotion of unc-80(lf); sup mutants. mac420, mac422 and mac424 could significantly improve the defective locomotion of unc-80(lf) mutants. (B) Curvature indices of nca-1(lf) single, nca-2(lf) single and nca-2(lf); nca-1(lf) double mutants. 40 animals were quantified for each genotype. Statistics: Bonferroni multiple comparison with one-way ANOVA. ***, p < 0.001.

(TIF)

**S2 Fig. Protein sequence alignment of TWK-40 and homologs.** The four transmembrane domains are enclosed in blue boxes, and the two pore domains are in red boxes. The amino acid at TM2.6 is indicated with a red arrowhead. The first 50 amino acids of the *Drosophila* CG43155 were omitted. C-terminal regions of TWK-40 (aa 338–393), *Drosophila* CG43155 (aa 364–411), *M. musculus* KCNK3 (aa 285–409), *H. sapiens* KCNK9 (aa 287–401) and *H. sapiens* KCNK16 (aa 260–262) were not included in the alignment.

(TIF)

**S3 Fig. twk-40 promoters were active in the nervous system.** (A) The genomic positions of Ptwk-40F, Ptwk-40L and Ptwk-40R promoters. (B) GFP driven by the Ptwk-40L promoter labeled some head neurons, ventral nerve cord and tail neurons. Ventral cord motor neurons
were not obviously labeled. (C) mCherry driven by the Ptwk-40R promoter labeled multiple head neurons, ventral nerve cord, ventral motor neurons and tail neurons. a: anterior; v: ventral.

(TIF)

**S4 Fig. Ptwk-40R and Ptwk-40L appeared to be active in mostly non-overlapping neurons.**

(A) Confocal pictures of head neurons labeled by Ptwk-40R::mCherry (left panel) and Ptwk-40L::GFP (middle panel). The merged picture was shown on the right. (B) Confocal pictures of the same animal as in (A) at a different focal plane. (C) A tail neuron(s) at the position of DVA/DVB/DVC neurons was co-labeled by mCherry and GFP. (D) Ventral nerve cord in the middle region of an animal showing the mCherry-labeled motor neurons and GFP-labeled cord. For all pictures, a: anterior; v: ventral.

(TIF)

**S5 Fig. Ptwk-40L::mCherry and Punc-80::GFP appeared to co-label AVA, AVE, AVB and RIS neurons.**

(A) Confocal pictures of adult head neurons expressing the Ptwk-40L::mCherry transgene (left panel) and the Punc-80::GFP transgene (middle panel). (B) Confocal picture of the same animal as in (A) on a different focal plane. (C) A tail neuron(s) at the position of DVA/DVB/DVC was co-labeled by mCherry (left panel) and GFP (middle panel). The merged pictures were shown on the right. For all pictures, a: anterior; v: ventral.

(TIF)

**S6 Fig. Ptwk-40R::mCherry and Punc-80::GFP appeared to co-label a subset of unidentified interneurons.**

(A) Confocal pictures of adult head neurons expressing the Ptwk-40R::mCherry transgene (left panel) and the Punc-80::GFP transgene (middle panel). The merged picture was shown on the right. (B) Confocal picture of the same animal as in (A) at a different focal plane. (C) A tail neuron(s) at the position of DVA/DVB/DVC appeared to be co-labeled by mCherry (left panel) and GFP (middle panel). (D) Ventral cord in the middle region of an animal showing several motor neurons co-labeled by mCherry and GFP. For all pictures, a: anterior; v: ventral.

(TIF)

**S7 Fig. twk-40 promoters and the nmr-1 promoter appeared to co-label some neurons.**

(A) Confocal pictures of adult head neurons expressing the Ptwk-40R::GFP transgene (left panel) and the Pnmr-1::mCherry transgene (middle panel). The merged picture was shown on the right. (B) Confocal pictures of adult tail neurons expressing the Ptwk-40R::GFP transgene (left panel) and the Pnmr-1::mCherry transgene (middle panel). (C) Confocal pictures of adult head neurons expressing the Ptwk-40L::GFP transgene (left panel) and the Pnmr-1::mCherry transgene (middle panel). For all pictures, a: anterior; v: ventral.

(TIF)

**S8 Fig. twk-40 promoters were active in DVA, DVB and DVC nervous.**

(A) Confocal pictures of adult tail neurons expressing the Ptwk-40F::mCherry transgene (left panel) and the Pprp-4::GFP transgene (middle panel). (B) Confocal pictures of adult tail neurons expressing the Ptwk-40F::mCherry transgene (left panel) and the Plim-6::GFP transgene (middle panel). (C) Confocal pictures of adult tail neurons expressing the Ptwk-40L::mCherry transgene (left panel) and the Plim-6::GFP transgene (middle panel). (D) Confocal pictures of adult tail neurons expressing the Ptwk-40R::mCherry transgene (left panel) and the Plim-6::GFP transgene (middle panel). The merged pictures were shown on the right. For all pictures, a: anterior; v: ventral.

(TIF)
S9 Fig. Mutations affecting \textit{twk-40} transcript isoforms. (A) List of presumptive \textit{twk-40} isoform-specific mutations. (B) Genomic positions of \textit{twk-40} isoform-specific mutations. (TIF)

S10 Fig. TWK 2\textsuperscript{nd} transmembrane domain sequence alignment and phenotypes of TWK gain-of-function mutants. (A) Alignment of the 2\textsuperscript{nd} transmembrane domains of indicated TWK channels. TM2.6 was substituted with asparagine (N). (B) Representative pictures of \textit{twk (gf)} mutants. (TIF)

S11 Fig. The \textit{Ptwk-40F} and \textit{Ptwk-2} promoters appeared to be active in same neurons. (A) Confocal pictures of adult head neurons expressing the \textit{Ptwk-40F}:GFP transgene (left panel) and a \textit{Ptwk-2}:mCherry transgene (middle panel). (B) An unidentified tail neuron(s) at the position of DVA/DVB/DVC was co-labeled by GFP (left panel) and mCherry (middle panel). The merged pictures were shown on the right. (C) Ventral cord in the middle region of a transgenic animal showing two motor neurons co-labeled by GFP and mCherry. For all pictures, a: anterior; v: ventral. (TIF)

S12 Fig. TWK-40(mac504gf) mutants exhibited improved locomotion in K\textsuperscript{+}-deficient environment. 40 animals were quantified for each assay condition. Statistics: two-tailed unpaired Student’s \textit{t}-test. ***, \textit{p} < 0.001. (TIF)

S1 Table. List of \textit{unc-80(lf)} suppressors and deleterious genetic variations identified in \textit{unc-80(lf); mac425/mac426} mutants. Chromosomal locations of the suppressors are shown at the top. Genetic variations identified from genomic sequencing of unoutcrossed \textit{unc-80(lf); mac425} and \textit{unc-80(lf); mac426} mutants are shown at the bottom. (XLSX)

S2 Table. List of \textit{twk-40a} mutations. (XLSX)

S3 Table. List of \textit{C. elegans twk} genes and their expression in the motor circuit neurons based on the CeNGEN project (www.cengen.org). Genes in red were expressed in the motor circuit and exhibited detectable effects on the motor behavior. Genes in blue were expressed in the motor circuit and did not obviously affect the motor behavior. Genes in bold black were previously studied but not analyzed in this study. Behaviors of loss-of-function mutants were considered here. \textit{twk-40} was not annotated to be obviously expressed in PVC neurons at CeNGEN. \textit{twk-3} was chosen as a presumptive negative control due to its lack of obvious expression in the motor circuit. The relative expression levels of \textit{twk} genes (based on CeNGEN) in each neuronal type are qualitatively indicated with the numbers of asterisks: * , weak; ** , moderate; *** , strong. (XLSX)

S4 Table. List of mutations and neuronal expression patterns of 14 \textit{twk} genes. We used the CRISPR/Cas9 method to generate mutations in these \textit{twk} genes. For each gene, top 20 neuron classes with high to low transcript signals (www.cengen.org) were listed. Mutations in bold were used as reference alleles. Key neurons in the motor circuit are shown in red. (XLSX)

S5 Table. Transgene rescue of \textit{twk(lf)} mutant phenotype. Wildtype transgenes under control of the \textit{unc-119} promoter were used. 20 animals from each of two transgenic lines were
quantified and the data were used together for comparison. Obvious phenotypes of twk-17(lf) and twk-48(lf) mutants other than the body curvature or locomotion were not quantified due to strong rescuing effects by the transgenes. Statistics: two-tailed unpaired Student’s t-test. ***, p < 0.001. NA: not applicable.

S6 Table. Curvature indices and backward body bends of twk loss-of-function mutants without obviously abnormal behavioral phenotypes. 40 animals were measured in each dataset. Statistics: Bonferroni multiple comparison with one-way ANOVA, ***, p < 0.001.

S7 Table. Genetic interactions between twk loss-of-function and presumptive gain-of-function mutations.

S8 Table. List of PCR primers for generating promoters, transgenes, and the indicated mutations.

S9 Table. twk target sequences used in CRISPR/Cas9-based mutagenesis. For all twk genes, two sgRNAs were used simultaneously. The use of twk-40 sgRNAs was as follows: twk-40a sgRNA # a1 and # a2 together with a repair template (S10 Table) for mac564; twk-40a sgRNA # a3 and # a4 for mac469, mac472 and neuron-specific knockouts; twk-40a sgRNA # a5 and # a6 together with a repair template for mac504 and mac505; twk-40b sgRNA # b1 and # b2 for mac554 and mac555; twk-40c sgRNA # c1 and # c2 for mac556 and mac557; twk-40b sgRNA # b2 and twk-40c sgRNA # c2 for mac558, mac559 and mac560. For each gene or twk-40 isoform, the number of transgenic lines examined, and the number of lines carrying the indicated mutations are shown.

S10 Table. Repair template sequences for introducing twk-40a isoform-specific mutations and TWK TM2.6>N mutations. The twk-40a #1 repair template was used for generating mac564 and twk-40a #2 for mac505. Nucleotides in red are for introducing the missense mutation and nucleotides in blue are silent mismatch mutations of the target sequences.
S7 Movie. *twk-30(lf)* Backward. (MP4)

S8 Movie. *twk-48(lf)* Forward. (MP4)

S9 Movie. *unc-58(lf)* Forward. (MP4)

S10 Movie. *twk-2(lf); twk-40(lf)* Forward. (MP4)

S11 Movie. *twk-7(lf) twk-40(lf)* Forward. (MP4)

S12 Movie. *twk-40(lf); twk-17(lf)* Forward. (MP4)

S13 Movie. *twk-48(lf) twk-40(lf)* Forward. (MP4)

S14 Movie. *twk-48(lf) twk-40(lf)* Backward. (MP4)

S15 Movie. *twk-40(lf); unc-58(lf)* Forward. (MP4)

S16 Movie. *twk-40(lf); unc-58(lf)* Backward. (MP4)

S17 Movie. *twk-2(lf); twk-40(gf)* Forward. (MP4)

S18 Movie. *twk-2(lf); twk-40(gf)* Backward. (MP4)

S19 Movie. *twk-2(gf); twk-40(lf)* Forward. (MP4)

S20 Movie. *twk-2(gf); twk-40(lf)* Backward. (MP4)

S1 Text. Strains and plasmid constructions. (DOCX)

S1 Data. Raw data for curvature indices and body bends. (XLSX)

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