

1 The ketone body β -hydroxybutyrate ameliorates neurodevelopmental deficits
2 in the GABAergic system of *daf-18/PTEN* *Caenorhabditis elegans* mutants

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Highlights

**daf-18/PTEN* deficiency in *C. elegans* results in a specific impairment of inhibitory GABAergic signaling, while the excitatory cholinergic signaling remains unaffected.

*The dysfunction of GABAergic neurons in these mutants arises from the inactivity of the transcription factor DAF-16/FOXO during their development, resulting in conspicuous morphological and functional alterations.

*A diet enriched with the ketone body β -hydroxybutyrate, which induces DAF-16/FOXO activity, mitigates the functional and morphological defects in the development of GABAergic neurons

* β -hydroxybutyrate supplementation during the early stages of development is both necessary and sufficient to achieve these rescuing effects on GABAergic signaling in *daf-18/PTEN* mutants.

4 **Abstract**

5 A finely tuned balance between excitation and inhibition (E/I) is essential for proper brain function.
6 Disruptions in the GABAergic system, which alter this equilibrium, are a common feature in
7 various types of neurological disorders, including Autism Spectrum Disorders (ASDs).
8 Mutations in PTEN, the main negative regulator of the PI3K/Akt pathway, are strongly associated
9 with ASD. However, it is unclear whether PTEN deficiencies can differentially affect inhibitory and
10 excitatory signaling. Using the *C. elegans* neuromuscular system, where both excitatory
11 (cholinergic) and inhibitory (GABAergic) inputs regulate muscle activity, we found that *daf-*
12 *18/PTEN* mutations specifically impact GABAergic (but not cholinergic) neurodevelopment and
13 function. This selective impact results in a deficiency in inhibitory signaling. The specific defects
14 observed in the GABAergic system in *daf-18/PTEN* mutants are due to reduced activity of DAF-
15 16/FOXO during development. Ketogenic diets (KGDs) have proven effective for disorders
16 associated with E/I imbalances. However, the mechanisms underlying their action remain largely
17 elusive. We found that a diet enriched with the ketone body β -hydroxybutyrate during early
18 development induces DAF-16/FOXO activity, therefore improving GABAergic neurodevelopment
19 and function in *daf-18/PTEN* mutants. Our study provides valuable insights into the link between
20 PTEN mutations and neurodevelopmental defects and delves into the mechanisms underlying
21 the potential therapeutic effects of KGDs.

22 **INTRODUCTION**

23 Maintaining a delicate balance between excitatory and inhibitory (E/I) neurotransmission is critical
24 for optimal brain function¹. Disruptions in this balance are commonly observed in
25 neurodevelopmental disorders²⁻⁴. In particular, deficits in inhibitory (GABAergic) signaling have
26 been reported in Autism Spectrum Disorders (ASD) and other related physiopathological
27 conditions^{4,5}.

28 *PTEN* is a classical tumor suppressor gene that antagonizes the highly conserved
29 phosphatidylinositol 3-phosphate kinase (PI3K)/protein kinase B (PKB/Akt) pathway. Several
30 reports using animal models have highlighted the importance of *PTEN* in neurodevelopment⁶⁻¹².
31 Moreover, mutations in *PTEN* were frequently found in human patients presenting ASD¹³. The
32 molecular events underlying the neurodevelopmental deficits in *PTEN* mutants remain poorly
33 understood.

34 The *C. elegans* neuromuscular system, where both excitatory (cholinergic) and inhibitory
35 (GABAergic) motor neurons regulate muscle contraction and relaxation, provides an excellent
36 platform for studying the function, balance, and coordination between excitatory and inhibitory
37 signals¹⁴⁻²⁰. This system has yielded valuable insights into fundamental synaptic transmission
38 mechanisms^{21,22}. Over the last decade, numerous studies focused on this simple yet highly
39 informative system have significantly contributed to our understanding of the functioning and
40 dysregulation of human genes associated with neurodevelopmental disorders, epilepsy, and
41 familial hemiplegic migraine^{14,19,20,23}. Furthermore, the substantial conservation of the main
42 components of the PI3K/Akt pathway in *C. elegans*^{24,25}, enhances the applicability of this model
43 system for investigating the role of this pathway in neurodevelopment.

44 We here found that mutations in *daf-18* (the ortholog for *PTEN* in *C. elegans*) result in specific
45 impairments in GABAergic inhibitory signaling due to decreased activity of the transcription factor
46 DAF-16 (the ortholog for FOXO in *C. elegans*) during neurodevelopment. Interestingly, cholinergic
47 excitatory motor neurons remain unaffected. This targeted impairment of inhibitory signals causes

48 an imbalance between excitatory and inhibitory (E/I) neurotransmission in the animal's
49 neuromuscular system.

50 In humans, ketogenic diets (KGDs), which force fatty acids beta-oxidation into ketone bodies,
51 have been utilized for decades to treat pathologies associated with E/I imbalances, such as
52 refractory epilepsies²⁶⁻²⁸. More recently, KGDs have also demonstrated effectiveness in
53 alleviating autistic symptoms in humans²⁹ and rodent models of ASD^{30,31}. The mechanisms
54 underlying these beneficial effects remain largely unknown.

55 We demonstrated that exposing *daf-18/PTEN* mutants to a diet enriched with the ketone body β -
56 hydroxybutyrate (β BHB) early in development enhances DAF-16/FOXO activity, mitigates
57 morphological and functional defects in GABAergic neurons, and improves behavioral
58 phenotypes. This study not only provides a straightforward system for studying the role of the
59 conserved PI3K/Akt/FOXO pathway in neurodevelopment but also contributes to our
60 understanding of the mechanisms underlying the effects of ketone bodies in neurodevelopment.

61

62 **RESULTS**

63 *Mutants in daf-18/PTEN and daf-16/FOXO are Hypersensitive to Cholinergic Drugs*

64 Disturbances in *C. elegans* cholinergic or GABAergic activity can be detected by analyzing the
65 sensitivity to the paralyzing effects of drugs that exacerbate cholinergic transmission^{14,15}. We
66 analyzed the sensitivity of *daf-18/PTEN* deficient animals to the acetylcholinesterase inhibitor
67 aldicarb and the cholinergic agonist levamisole. Exposure to aldicarb leads to an increase in ACh
68 levels at cholinergic motor synapses, resulting in massive activation of muscular cholinergic
69 receptors and subsequent paralysis³² (Figure 1A). Levamisole also induces paralysis by directly
70 activating muscular cholinergic receptors³² (Figure 1A). We found that *daf-18/PTEN* mutants are
71 hypersensitive to the paralyzing effects of both drugs (Figures 1B and 1C). Hypersensitivity to
72 cholinergic drugs is typical of animals with an increased E/I ratio in the neuromuscular system,
73 such as mutants in *unc-25* (the *C. elegans* orthologue for glutamic acid decarboxylase, an

74 essential enzyme for synthesizing GABA)^{14,15}. While *daf-18/PTEN* mutants become paralyzed
75 earlier than wild-type animals, their hypersensitivity to cholinergic drugs is not as severe as that
76 observed in animals completely deficient in GABA synthesis, such *unc-25* null mutants (Figures
77 1B and 1C) indicating a less pronounced imbalance between excitatory and inhibitory signals.
78 Reduced activity of DAF-18/PTEN has been largely shown to exacerbate the PI3K pathway,
79 precluding the activation of DAF-16/FOXO, the *C. elegans* ortholog of the FOXO transcription
80 factors family²⁵ (Figure S1A). We analyzed aldicarb and levamisole sensitivity of mutants in this
81 transcription factor. Similar to *daf-18/PTEN* mutants, we found that *daf-16/FOXO* null mutants are
82 hypersensitive to the paralyzing effects of aldicarb and levamisole (Figure 1C). Furthermore, we
83 did not observe significant differences in aldicarb and levamisole sensitivity between *daf-18*; *daf-*
84 *16* double null mutants and the respective single mutants, suggesting that both genes affect
85 neuromuscular signaling by acting in the same pathway (Figure 1C). In addition to *daf-16/FOXO*
86 and *daf-18/PTEN*, we assessed the sensitivity to the paralyzing effects of aldicarb and levamisole
87 in loss of function mutants of other components of the PI3K pathway, such as *age-1/PI3K*, *pdk-1*,
88 *akt-1*, and *akt-2* (Figure 1D). Unlike mutations in *daf-18/PTEN*, in these mutants the PI3K pathway
89 is downregulated (Figure S1A). We did not observe significant differences compared to the wild-
90 type. Due to the complete dauer arrest observed in double mutants of *akt-1* and *akt-2*^{33,34}, we
91 were unable to explore the potential redundancy of these two genes. Interestingly, a gain-of-
92 function mutant in *pdk-1*, *pdk-1(mg142)*²⁴, is hypersensitive to aldicarb and levamisole, similar to
93 *daf-16/FOXO* and *daf-18/PTEN* mutants (Figure 1D). Given that increased *pdk-1* activity is linked
94 to hyperphosphorylation and inactivation of DAF-16/FOXO (Figure S1A), these results support
95 the hypothesis that low activity of DAF-16/FOXO leads to hypersensitivity to these drugs.
96 In vertebrates, alterations in PTEN activity have been largely shown to impact neuronal
97 development and function by affecting the mTOR pathway³⁵. Consequently, we analyzed whether
98 mutations in components of the *C. elegans* TOR complexes (TORC) would lead to significant
99 changes in sensitivity to aldicarb and levamisole. Our findings indicate that neither animals with

100 loss of the essential TORC1 component *raga-1*/RagA nor animals with a loss of function in the
101 essential TORC2 component *rict-1*/Rictor exhibited significant alterations in sensitivity to
102 cholinergic drugs compared to wild-type animals (Figure 1E). This suggests that the mTOR
103 pathway is not involved in *daf-18/PTEN* pharmacological phenotypes.

104

105 *daf-18/PTEN and daf-16/FOXO Mutants Show Phenotypes Indicative of GABAergic Deficiency*

106 In *C. elegans*, the body wall muscles receive cholinergic innervation, which induces contraction,
107 and GABAergic innervation, which leads to relaxation. The contralateral activity of cholinergic and
108 GABAergic neurons facilitates the characteristic undulatory movement of the animal (Figure 2A).
109 Hypersensitivity to cholinergic drugs has long been observed in worms where GABAergic
110 signaling is deficient^{14,15} (Figure 1). In mutants with severe deficits in GABA transmission,
111 prodding induces a bilateral contraction of the body wall muscles that shortens the body (shrinker
112 phenotype)³⁶. When *daf-18/PTEN* mutants are touched, there is a slight but significant shortening
113 in body length (Figure 2B). As expected, this shortening is not as noticeable as in animals with a
114 complete deficit in GABAergic signaling, such as mutants in *unc-25*. Similar to *daf-18/PTEN*
115 mutants, *daf-16/FOXO* animals also exhibit a mild decrease in body length after prodding.
116 Consistent with our aldicarb and levamisole results, there are no significant differences in body
117 shortening between *daf-18;daf-16* double mutants and the corresponding single mutants (Figure
118 2B), further supporting the notion that both genes act in the same pathway to impact
119 neuromuscular signaling.

120 We also analyzed other behaviors that require a concerted activity of GABAergic and cholinergic
121 systems, such as the omega turns during the escape response³⁷. In *C. elegans* the escape
122 response can be induced by a gentle touch on the head and involves a backward movement that
123 is usually followed by a sharp omega turn and a 180° change in its direction of locomotion ³⁷
124 (Movie 1). The execution of the omega turn involves a hypercontraction of the ventral muscles
125 and relaxation of the dorsal muscles, allowing the animal to make a sharp turn, where the animal's

126 head slides on the ventral side of the body (closed omega turn), and resumes locomotion in the
127 opposite direction (Movie 1). In response to anterior touch, the vast majority of wild-type worms
128 make a closed omega turn^{37,38}(Figure 2C). Ventral muscle contraction is triggered by cholinergic
129 motor neurons (VA and VB neurons) that synapse onto ventral muscles, while dorsal muscle
130 relaxation is induced by GABAergic motor neurons (DD neurons) that synapse onto dorsal
131 muscles (Figure 2A)^{37,38}. Ablation of DD GABAergic neurons reduces dorsal muscle relaxation,
132 therefore preventing the head from touching the ventral side of the body during the escape
133 response (open omega turn)³⁷. In agreement with previous reports, we found that 91% of wild-
134 type animals exert a closed omega turn within the escape response (Figure 2C). We observed
135 that similar to wild-type animals, gentle anterior touch with an eyelash induces *daf-18* mutants to
136 move backward and initiate an omega turn (Movie 2). However, only 53% of *daf-18* mutants
137 exhibit the typical head-to-tail contact during the omega turn (Figure 2C and Movie 2). Akin to *daf-*
138 *18* mutants, *daf-16/FOXO* mutants exhibited a decrease in the proportion of closed omega turns
139 (Figure 2C). No additive effects were observed in the *daf-18*; *daf-16* double mutant, suggesting
140 that the increased inactivation of *daf-16/FOXO* is primarily responsible for the defects observed
141 in the escape response of *daf-18* mutants.

142 Given that our results suggest a deficit in GABAergic functionality in *daf-18/PTEN* mutants, we
143 used optogenetics to specifically activate these neurons in mutant worms. The expression of
144 Channelrhodopsin (ChR2) in GABAergic motor neurons (using the *unc-47* promoter, orthologue
145 for the vesicular GABA transporter SLC32A1) elicits a flaccid paralysis of the worms upon
146 exposure to blue light. This obvious and robust response results in an increase in body length
147 that can be used as a clear readout³⁹⁻⁴¹(Figure 2D and Movie 3). Interestingly, we found that the
148 elongation of the animal after the specific activation of GABAergic neurons is significantly
149 decreased in *daf-18/PTEN* and *daf-16/FOXO* mutants compared to wild-type worms (Figure 2D).
150 While these results suggest a defect in GABAergic transmission, it could also be possible that
151 general neuronal transmission is affected. Consequently, we reciprocally activated the cholinergic

152 motor neurons in animals expressing ChR2 under the *unc-17* promoter, a gene encodes the
153 vesicular acetylcholine transporter (VACHT), which leads to muscle contraction and shortened
154 body length^{39,41}(Figure 2E and Movie 4). Rather than observing reduced shortening in *daf-*
155 *16/FOXO* and *daf-18/PTEN* mutants, we found that cholinergic activation caused
156 hypercontraction of these mutant animals (Figure 2E). Since the activation of cholinergic motor
157 neurons not only activates muscles but also stimulates GABAergic neurons to produce
158 counteractive muscle relaxation in the other side of the animal (Figure 2A), it is expected that a
159 GABAergic deficit would lead to increased muscle contraction and body shortening upon
160 cholinergic activation. In summary, these results strongly suggest that in *daf-18/PTEN* and *daf-*
161 *16/FOXO* mutants, there is a specific functional defect in GABAergic neurons, while excitatory
162 neurons do not appear to be affected.

163

164 *Disruption of daf-18/PTEN Alters Commissural Trajectories in GABAergic Motor Neurons*
165 Since our previous results imply perturbations of neuromuscular transmission, we explored the
166 morphology of *C. elegans* motor neurons. The cell bodies of both cholinergic (A and B-type) and
167 GABAergic (D-type) motor neurons that innervate body wall muscles are located in the ventral
168 nerve cord (VNC), and a subset extends single process commissures to the dorsal nerve cord
169 (DNC)⁴² (Figures 3A and S2). The commissures have proved useful for studying defects in motor
170 neuron development or maintenance^{43,44}. We analyzed the morphology of GABAergic motor
171 neurons in L4 animals expressing *mCherry* under the control of the *unc-47* promoter⁴⁵. We found
172 that *daf-18/PTEN* mutants exhibit a higher frequency of commissure flaws, including guidance
173 defects, ectopic branching, and commissures that fail to reach the dorsal cord (Figures 3B and
174 3C). In contrast to our findings in GABAergic neurons, we observed no obvious differences in the
175 frequency of commissure defects when we compared cholinergic motor neurons in control and
176 *daf-18/PTEN* animals (Figure S2).

177 GABAergic motor neurons can be classified based on the muscles they innervate: those that
178 innervate the dorsal muscles are called DDs, while those that innervate the ventral muscles are
179 called VDs. Both types of D neurons send commissures to the DNC^{46,47}. We found defects in
180 various commissures, some of which correspond to VD and others to DD neurons (Figure S3).
181 We also analyzed GABAergic commissures at the beginning (1 hour post-hatching) of the first
182 larval stage (L1), when only the six DD neurons are formed^{46,48,49}. We found that in this early larval
183 stage, *daf-18/PTEN* mutants exhibit defects in the GABAergic commissures (Figures 3D, 3E and
184 S4). While the DD neurons are born and mostly develop embryonically, limited postembryonic
185 axonal outgrowth has been observed in these neurons⁴⁹. We did not find an increase in the
186 number of errors in larvae 5-6 hours post-hatching compared to recently hatched larvae (Figure
187 S4B), indicating that deficiencies in DD neurons in *daf-18/PTEN* mutants mainly occur during their
188 embryonic development. In contrast, we observed that the prevalence of errors increases
189 significantly at the L4 stage (Figure S4B and S4C). The greater number of defects in L4s likely
190 arises from defects in the VDs, which are born post-embryonically between the mid-L1 larval
191 stage and the L2 stage, and add to the defects already present in the DDs. Taken together, these
192 observations suggest that reduced DAF-18/PTEN activity affects the neurodevelopment of the
193 GABAergic motor system. Since the transcription factor DAF-16/FOXO is one of the main targets
194 of DAF-18/PTEN signaling, we analyzed the morphology of GABAergic motor neurons in *daf-*
195 *16/FOXO* null mutants. These animals also exhibit an increased number of defects in GABAergic
196 commissures compared to the wild type (Figure 3C).

197 Given that *daf-18* is ubiquitously expressed in all tissues²⁵, we asked whether DAF-18/PTEN acts
198 autonomously in GABAergic neurons to ensure proper development. We found that specific *daf-*
199 *18/PTEN* expression in GABAergic neurons increased the proportion of closed omega turns in
200 *daf-18/PTEN* null mutants (Figure 3F). In addition, the morphological defects in GABAergic
201 commissures were significantly reduced (Figure 3G), suggesting that DAF-18/PTEN acts
202 autonomously in GABAergic motor neurons to regulate their development.

203 Collectively, our findings demonstrate that mutations in *daf-18/PTEN* and *daf-16/FOXO* result in
204 developmental defects in GABAergic neurons, leading to both altered morphology and function,
205 while leaving cholinergic motor neurons unaffected. Our experiments strongly suggest that these
206 specific defects in the inhibitory transmission arise from the hyperactivation of the PI3K pathway,
207 along with subsequent DAF-16/FOXO inhibition, in GABAergic neurons of *daf-18/PTEN* mutants.

208

209 *A diet enriched with the ketone body β-hydroxybutyrate (βHB) Ameliorates Defects in daf18/PTEN*
210 *Mutants but not in daf-16/FOXO Mutants*

211 Mutations in *PTEN* are linked with Autism Spectrum Disorders (ASD)⁶. Ketogenic diets, which
212 force the endogenous production of KBs, have proved to be effective for the treatment of
213 neurological disorders associated with E/I imbalances, such as epilepsy and, more recently, ASD
214²⁷⁻²⁹. It has been shown that the KB βHB induces DAF-16/FOXO activity⁵⁰. Therefore, we asked
215 whether it is possible to improve the observed phenotypes by modulating the activity of DAF-
216 16/FOXO with βHB. We first evaluated the expression of *sod-3*, which codes for a superoxide
217 dismutase and is a DAF-16/FOXO transcriptional target gene⁵¹. We used a strain expressing a
218 GFP transcriptional reporter for *sod-3* and determined fluorescence intensity upon dietary
219 supplementation of βHB. Consistent with previous reports, the levels of SOD-3::GFP are reduced
220 in *daf-18/PTEN* and *daf-16/FOXO* mutant strains. Furthermore, we observed that βHB (20 mM)
221 induces the expression of *sod-3* in *daf-18/PTEN* but not in *daf-16/FOXO* mutants (Figure S5).
222 Importantly, we did not detect increased *sod-3* expression in *daf-18; daf-16* double deficient
223 animals, strongly suggesting that βHB induces *sod-3* expression in *daf-18/PTEN* mutants through
224 the transcription factor *daf-16/FOXO* (Figure S5).

225 Next, we evaluated behavioral phenotypes and GABAergic neuronal morphology of animals that
226 were raised on an *E. coli* diet supplemented with 20 mM βHB throughout development, from egg
227 laying until the time of the assay (typically late L4s or young adults). We found that βHB
228 supplementation significantly reduced the hypersensitivity of *daf-18/PTEN* mutants to the

229 cholinergic drugs aldicarb and levamisole (Figures 4A and 4B). Moreover, β HB supplementation
230 rescued the post-prodding shortening in *daf-18/PTEN* mutants (Figure 4C). Accordingly, we found
231 that *daf-18/PTEN* mutants showed a significant increase in the proportion of closed omega turns
232 during their escape response compared to the naïve condition (Figure 4D). In contrast, β HB
233 exposure does not change the number of closed omega turns in *daf-16/FOXO* null mutants or the
234 double null mutant *daf-18; daf-16* (Figure 4D).

235 We subsequently analyzed the changes in body length induced by optogenetic activation of both
236 GABAergic and cholinergic neurons in animals exposed to a diet enriched with β HB. Interestingly,
237 we found that *daf-18/PTEN* mutants exposed to β HB, but not wild-type or *daf-16/FOXO* mutant
238 animals, exhibited increased elongation following optogenetic activation of GABAergic neurons
239 (Figures 4E, 4F, and 4G). Furthermore, we observed that the hypercontraction observed in *daf-18/PTEN*
240 mutants after the activation of cholinergic neurons is significantly reduced in animals
241 exposed to β HB (Figures 4H, 4I and 4J). These findings suggest that this ketone body can
242 rebalance excitatory and inhibitory signals in the neuromuscular system of *daf-18/PTEN C. elegans*
243 mutants.

244 We also evaluated the morphology of GABAergic motor neurons in *daf-18/PTEN* animals exposed
245 to β HB. We found that β HB supplementation reduced the frequency of defects in GABAergic
246 processes (Figure 4K). Consistently, β HB exposure did not significantly reduce the defects on
247 GABAergic neurons of either *daf-16/FOXO* null mutants or *daf-18; daf-16* double mutants. Taken
248 together, these results demonstrate that dietary β HB ameliorates the defects associated with
249 deficient GABAergic signaling in *daf-18/PTEN* mutants.

250 It is noteworthy that we did not observe any improvement in either neuronal outgrowth defects in
251 the AIY interneuron or the migration of the HSN motor neurons (Figure S6) in *daf-18/PTEN*
252 mutants exposed to β HB, even though these defects were shown to depend on the reduction of
253 DAF-16/FOXO activity^{52,53}. AIY neurite and HSN soma migration take place during

254 embryogenesis^{52,54}. It is therefore possible that βHB may not go through the impermeable chitin
255 eggshell of the embryo, as has been reported with other drugs (see below)⁵⁵.

256

257 *β-Hydroxybutyrate Exposure During Early L1 Development Sufficiently Mitigates GABAergic*
258 *System Defects*

259 In the above experiments, animals were exposed to βHB throughout development. We next asked
260 whether there is a critical period during development where the action of βHB is required. We
261 exposed *daf-18/PTEN* mutant animals to βHB-supplemented diets for 18-hour periods at different
262 developmental stages (Figure 5A). The earliest exposure occurred during the 18 hours following
263 egg laying, covering *ex-utero* embryonic development and the first 8-9 hours of the L1 stage. The
264 second exposure period encompassed the latter part of the L1 stage, the entire L2 stage, and
265 most of the L3 stage. The third exposure spanned the latter part of the L3 stage, the entire L4
266 stage, and the first 6-7 hours of the adult stage (Figure 5A). Interestingly, we found that the earliest
267 exposure to βHB was sufficient to increase the proportion of *daf-18/PTEN* mutant animals
268 executing a closed omega turn during the escape response. However, when the animals were
269 exposed to βHB at later juvenile stages, their ability to enhance the escape response of *daf-*
270 *18/PTEN* mutants declined (Figure 5B). Moreover, exposing animals to βHB for 18 hours starting
271 from egg laying was enough to reduce morphological defects in the GABAergic motor neurons of
272 these mutants (Figure 5C and Figure S7). Interestingly, βHB has no effect on GABAergic
273 commissures in either recently hatched L1s or L4 *daf-18/PTEN* mutants when exposure is limited
274 to the first 9 hours after egg laying (where *ex utero* embryonic development occurs), possibly due
275 to the impermeability of the chitinous eggshell (Figure S7). Thus, it is likely that βHB acts at early
276 L1 stages to mitigate neurological GABAergic defects in *daf-18/PTEN* mutants.

277 Taken together, our findings demonstrate that mutations in *daf-18*, the *C. elegans* orthologue of
278 *PTEN*, lead to specific defects in inhibitory GABAergic neurodevelopment without significantly
279 affecting cholinergic excitatory signals. These GABA-specific deficiencies manifest as altered

280 neuronal morphology, hypersensitivity to cholinergic stimulation, reduced responses to
281 optogenetic GABAergic neuronal activation, mild body shortening following touch stimuli, and
282 deficits in the execution of the omega turn. We have determined that these impairments in
283 GABAergic development result from reduced activity of the FOXO orthologue DAF-16 in *daf-*
284 *18/PTEN* mutants. Importantly, our study's pivotal finding is that a βHB- enriched diet during early
285 development, robustly mitigates the deleterious effects of *daf-18/PTEN* mutations in GABAergic
286 neurons. This protective effect is critically dependent on the induction of DAF-16/FOXO by this
287 ketone body.

288

289 **DISCUSSION**

290 Mutations in *daf-18/PTEN* are linked to neurodevelopmental defects from worms to
291 mammals^{11,52,53}. Moreover, decreased activity of PTEN produces E/I disequilibrium and the
292 development of seizures in mice⁷. The mechanisms underlying this imbalance are not clear. Our
293 results demonstrate that reduced DAF-18/PTEN activity in *C. elegans* generates guidance
294 defects, abnormal branching, incomplete commissural outgrowth and deficient function of
295 inhibitory GABAergic neurons, without affecting the excitatory cholinergic neurons.
296 *daf-18/PTEN* deficient mutants have a shorter lifespan⁵⁶. One possibility is that the defects in
297 GABAergic processes are due to neurodegeneration associated with premature aging rather than
298 developmental flaws. However, this idea is unlikely given that the neuronal defects in DD neurons
299 are already evident at the early L1 stage. In *C. elegans*, DD GABAergic motor neurons undergo
300 rearrangements during the L1-L2 stages^{48,49}. In newly hatched L1 larvae, each DD motor neuron
301 innervates ventral muscles and extends a commissure to the dorsal nerve cord to receive synaptic
302 inputs from cholinergic DA and DB neurons^{48,49,57}. In adults, the DD commissure morphology is
303 maintained, but the synaptic output shifts to dorsal muscles, and input is provided by VA and VB
304 cholinergic motor neurons in the ventral nerve cord⁵⁷. A plausible possibility is that this remodeling
305 process makes DD neurons specifically sensitive to PI3K pathway activity. However, since the

306 DD commissures are defective already at the very early L1 stage (prior to rewiring) and similar
307 defects are observed in VD neurons, which are born post-embryonically between late L1 and L2
308 stages and do not undergo this remodeling^{46,49,57}, we can infer that the deficits caused by *daf-*
309 *18/PTEN* deficiency affect the development of the entire GABAergic system, independently of the
310 synaptic rearrangement of DD neurons.

311 Strikingly, cholinergic neurons have no noticeable morphological or functional defects in *daf-*
312 *18/PTEN* mutants. Loss-of-function mutants in the neuronal integrin *ina-1*, ortholog of human
313 ITGA6, affect the guidance of GABAergic commissures, without affecting cholinergic neurons⁴⁴.
314 Similar to *PTEN*, mutations in neuronal integrins have been linked to neurodevelopmental defects
315⁵⁸. Interestingly, the PI3K/Akt/FOXO pathway and integrin signaling are interrelated in mammals
316⁵⁹. This observation opens the possibility that one of the mechanisms by which *daf-18/PTEN*
317 mutants have defects in GABAergic neurodevelopment involves integrin expression and/or
318 function. Interestingly, mutations in *eel-1*, the *C. elegans* ortholog of HUWE1, or in subunits of the
319 Anaphase-Promoting Complex (APC), lead to developmental and functional alterations in
320 GABAergic neurons but not in cholinergic neurons^{19,60}, despite their expression in both neuronal
321 types. This observation suggests the existence of compensatory or redundant mechanisms in
322 cholinergic neurons that may not be present in GABAergic neurons.

323 In mammals, defects in *PTEN* mutants have been typically related to altered function of the mTOR
324 pathway^{8,61,62}. However, our results suggest that, in the *C. elegans* neuromuscular system,
325 decreased activity of DAF-18/PTEN affects GABAergic development due to a downregulation of
326 DAF-16/FOXO transcription factor activity. The FOXO family of transcription factors is conserved
327 throughout the animal kingdom⁶³. There is increasing evidence demonstrating the key role of this
328 transcription factors family in neurodevelopment⁶⁴⁻⁶⁶. Downregulation of FOXO activity early in
329 development reproduces neuropathological features found in ASD patients, i.e., increased brain
330 size and cortical thickness^{67,68}. The autonomic activity of DAF-18/PTEN and DAF-16/FOXO
331 coordinates axonal outgrowth in *C. elegans* AIY interneurons and rat cerebellar granule neurons

332 52. On the other hand, DAF-18/PTEN and DAF-16/FOXO in the hypodermis control neuronal
333 migration of the HSN neuron during development⁵³. Our rescue experiments strongly suggest that
334 the PI3K/Akt/DAF-16 pathway modulates the development of GABAergic motor neurons by acting
335 autonomously in these cells. Noteworthy, autonomic DAF-16/FOXO activity in GABAergic motor
336 neurons is also key for axonal growth during regeneration⁶⁹. These results further emphasize the
337 importance of DAF-16/FOXO in neuronal development and axonal growth.

338 In many patients suffering from epilepsy, ketogenic diets can control seizures^{27,28}. Furthermore,
339 they can reduce behavioral abnormalities in individuals with ASD²⁹. While the mechanisms
340 underlying the clinical effects of ketogenic diets remain unclear, it has been shown that these
341 diets correlate with increased GABA signaling⁷⁰⁻⁷². We demonstrate here that dietary
342 supplementation of the ketone body βHB ameliorates morphological and functional defects in
343 GABAergic motor neurons of *daf-18/PTEN* mutants. Although ketone bodies were historically
344 viewed as simple carriers of energy to peripheral tissues during prolonged fasting or exercise, our
345 findings confirm more recent reports showing that βHB also possesses a variety of important
346 signaling functions⁷³. We can hypothesize several distinct, non-mutually exclusive models by
347 which βHB can induce DAF-16/FOXO-dependent signaling. βHB directly inhibits mammalian
348 histone deacetylases HDAC1 and HDAC2, increasing histone acetylation at the FOXO3a
349 promoter and inducing the expression of this gene⁷⁴. HDAC1 and HDAC2 play an important role
350 as redundant regulators of neuronal development⁷⁵. Interestingly, in *C. elegans* βHB inhibits the
351 class I HDACs to extend worm lifespan in a DAF-16/FOXO-dependent manner⁵⁰. Therefore, βHB-
352 mediated HDAC inhibition may upregulate transcription of DAF-16/FOXO counterbalancing
353 hyperactivation of the PI3K pathway in *daf-18/PTEN* mutants. Another potential mechanism for
354 the effect of βHB involves the inhibition of the insulin signaling pathway. In mammals, the
355 administration of βHB downregulates the insulin signaling in muscle⁷⁶. Moreover, several reports
356 have shown that βHB administration reduces phosphorylation and activity of Akt/protein kinase
357 downstream of the insulin receptor^{77,78}. In *C. elegans*, inhibition of AKT-1 activates DAF-16/FOXO

358 ⁷⁹. Although understanding the mechanism behind β HB's action will require further studies, our
359 results demonstrate that this ketone body positively modulates DAF-16/FOXO during neuronal
360 development.

361 Multiple reports, from *C. elegans* to mammals, suggest that there is a sensitive period, typically
362 early in development, where pharmacological or genetic interventions are more effective in
363 ameliorating the consequences of neurodevelopmental defects⁸⁰. However, recent evidence
364 shows that phenotypes associated with certain neurodevelopmental defects can be ameliorated
365 by interventions during adulthood⁸¹. Our results show that β HB can ameliorate the phenotypic
366 defects of *daf-18/PTEN* mutants only when exposure occurs during an early critical period. The
367 inefficacy of β HB at later stages suggests that the role of DAF-16/FOXO in the maintenance of
368 GABAergic neurons is not as relevant as its role in development.

369 Our experiments do not allow us to distinguish whether the effect of β HB is preventive, reversible,
370 or both. Our results suggest that the improvement is not due to prevention in DDs because the
371 defects are present in newly hatched larvae regardless of the presence or absence of β HB, and
372 DD post-embryonic growth does not add new errors. Unlike in early L1 stages, the protective
373 effect of β HB becomes evident when analyzing the commissures of L4 animals. In this late larval
374 stage, not only the DDs but also the VD neurons are present. This leads us to speculate that β HB
375 may have a preventive action on the neurodevelopment of VD neurons. We also cannot rule out
376 that this improvement may be due, at least partially, to a reversal of defects in DD neurons. It is
377 intriguing how exposure to β HB during early L1 could ameliorate defects in neurons that mainly
378 emerge in late L1s (VDs). We can hypothesize that residual β HB or a metabolite from the previous
379 exposure may prevent these defects in VD neurons. β HB, in particular, has been shown to
380 generate long-lasting effects through epigenetic modifications⁸². Further investigations are
381 needed to elucidate the underlying fundamental mechanisms regarding the ameliorating effects
382 of β HB supplementation on deficits in GABAergic neurodevelopment associated with mutations
383 in *daf-18/PTEN*.

384 Across the animal kingdom, food signals increase insulin levels leading to the activation of
385 Akt/PI3K pathway⁸³⁻⁸⁵. In *C. elegans*, the L1 larval stage is particularly sensitive to nutritional
386 status. *C. elegans* adjusts its development based on food availability, potentially arresting in L1
387 in the absence of food⁸⁶. Strong loss-of-function alleles in the insulin signaling pathway exhibit
388 constitutive L1 arrest⁸⁷, highlighting the critical importance of this pathway during this larval stage.
389 Hence, it is not surprising that dietary interventions targeting the PI3K pathway at these critical
390 early L1 stages can modulate developmental processes. Our pharmacological experiments
391 showed that mutants associated with an exacerbation of the PI3K pathway, which typically inhibits
392 the nuclear translocation and activity of the transcription factor DAF-16/FOXO, lead to E/I
393 imbalances that manifest as hypersensitivity to cholinergic drugs. We demonstrated that these
394 imbalances arise from defects that occur specifically in the neurodevelopment of GABAergic
395 motor neurons. Interestingly, mutants inhibiting the PI3K pathway do not show differences in their
396 sensitivity to cholinergic drugs compared to wild-type animals. This observation can be explained
397 by a critical period during neurodevelopment when the Insulin/Akt/PI3K pathway must be
398 maintained at very low activity (or even deactivated). These low activity levels of the
399 Insulin/Akt/PI3K pathway would allow for a high level of DAF-16/FOXO activity, which, according
400 to our results, appears to be key for the proper development of GABAergic neurons.
401 The fine regulation of insulin and insulin-like signaling during early development is a conserved
402 process in animals^{88,89}. In mammals, for instance, conditions like Gestational Diabetes Mellitus
403 (GDM), characterized by fetal hyperinsulinemia and high levels of IGF-1⁹⁰, are associated with
404 neurodevelopmental defects⁹¹. Our results lead to the intriguing idea that dietary interventions
405 that increase DAF-16/FOXO activity, such as βHB supplementation, could constitute a potential
406 therapeutic strategy for these pathologies. Future studies using mammalian models are crucial to
407 shed light on the potential of this hypothesis.
408
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410 **References**

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634

635

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653

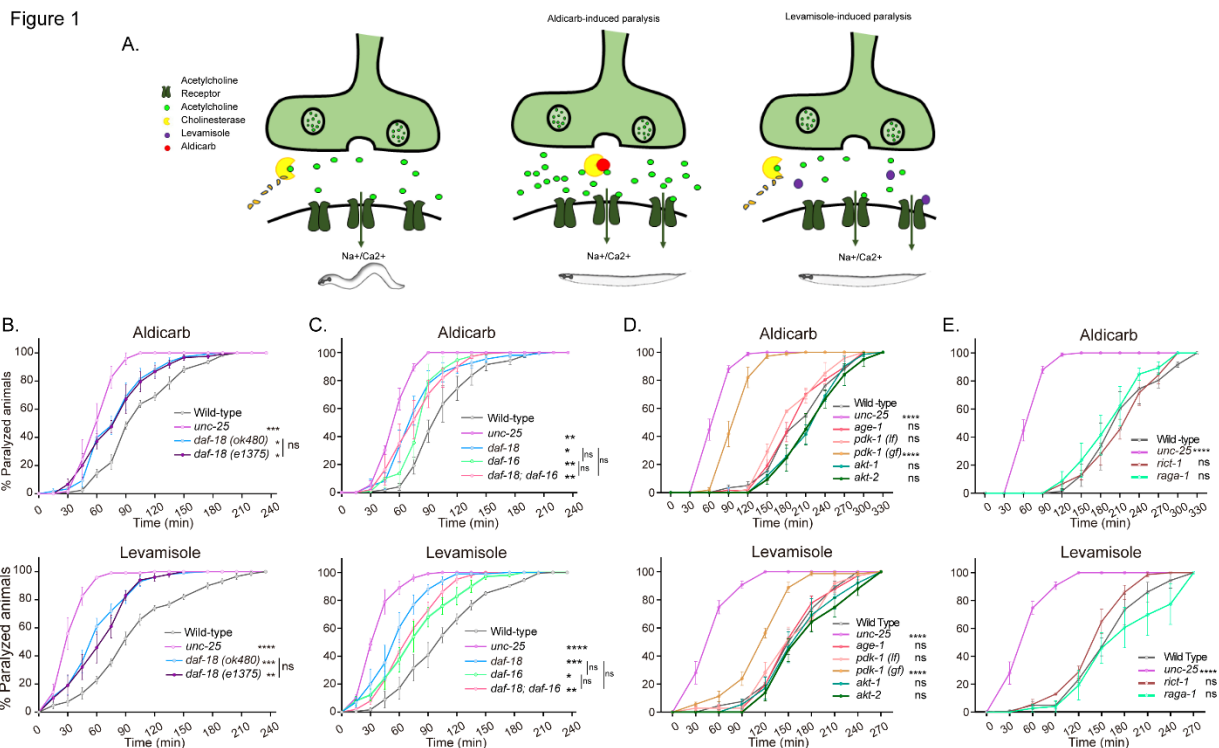
654 **Conflict of Interest**

655 The authors declare that the research was conducted in the absence of any commercial or
656 financial relationships that could be construed as a potential conflict of interest.

657

658 **Main Figures**

659 **Figure 1**



660

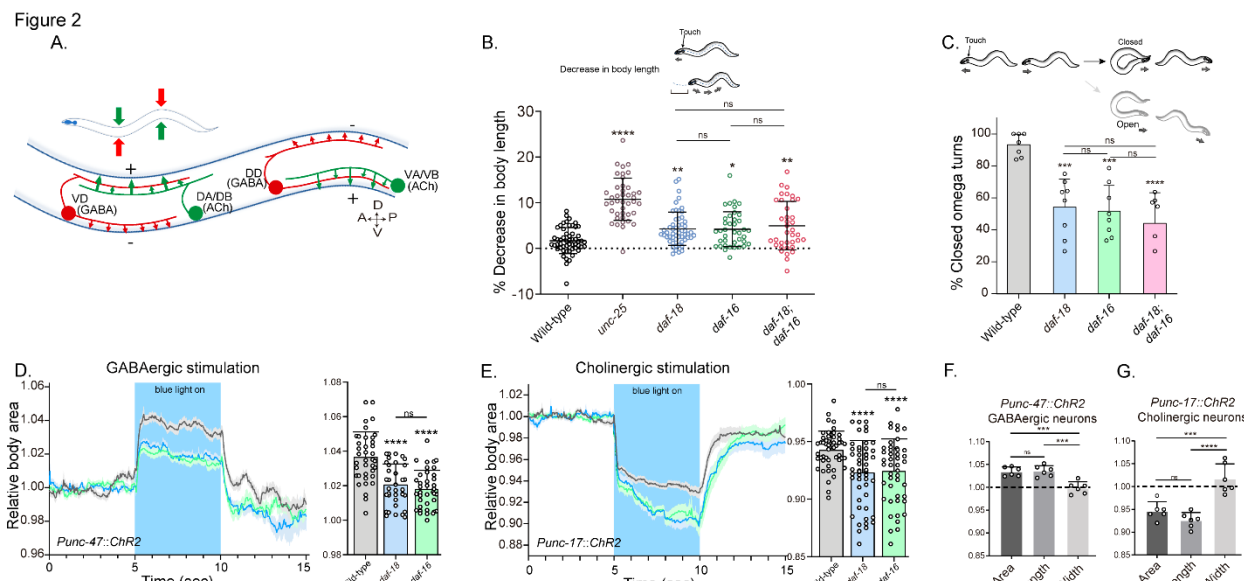
661 **Figure 1. *daf-18/PTEN* mutants are hypersensitive to cholinergic drugs.**

662 **A.** Schematic representation of paralysis induced by aldicarb and levamisole. Aldicarb acts by
663 inhibiting acetylcholinesterase, leading to an accumulation of acetylcholine at the
664 neuromuscular junction, resulting in continuous stimulation of muscles and worm paralysis.
665 Levamisole functions as an agonist at nicotinic acetylcholine receptors, causing prolonged
666 depolarization and, also, muscle paralysis. **B-E.** Quantification of paralysis induced by aldicarb
667 (Top) and levamisole (Bottom). The assays were performed in NGM plates containing 2 mM
668 aldicarb or 0.5 mM levamisole. Strains tested: N2 (wild-type) (B-E), CB1375 *daf-18(e1375)* (B),
669 OAR144 *daf-18(ok480)* (B-C), GR1310 *akt-1(mg144)* (D), TJ1052 *age-1(hx546)* (D), VC204 *akt-*
670 *2(ok393)* (D), VC222 *raga-1(ok386)* (E) and KQ1366 *rict-1(ft7)* (E). All of these strains carry
671 loss-of-function mutations. Furthermore, the strains denoted as "*pdk-1 (lf)*" and "*(gf)*" correspond

672 to JT9609 *pdk-1*(sa680), which possesses a loss-of-function mutation, and GR1318 *pdk-*
673 *1*(mg142), which harbors a gain-of-function mutation in the *pdk-1* gene, respectively. The strain
674 CB156 *unc-25*(e156) was included as a strong GABA-deficient control (B-D). At least four
675 independent trials for each condition were performed (n= 25-30 animals per trial). One-way ANOVA was used to test
676 statistical differences in the area under the curve (AUC) among different strains. Post-hoc analysis after One-
677 Way ANOVA was performed using Tukey's multiple comparisons test (B and C) and Dunnet's to
678 compare against the wild-type strain (D and E) (ns p > 0.05; * p ≤ 0.05; ** p ≤ 0.01; *** p ≤
679 0.001; **** p ≤ 0.0001).

680

681 **Figure 2**



682

683

684 **Figure 2. *daf-18/PTEN* mutants exhibit phenotypes typical of GABA-deficient animals.**

685 **A.** Schematic of *C. elegans* adult neuromuscular circuit. Red indicates GABAergic motor
686 neurons (DD/VD) and green indicates cholinergic motor neurons (VA/VB and DA/DB). The VA
687 and VB cholinergic motor neurons send synaptic inputs to the ventral body wall muscles and the
688 DD GABAergic motor neurons. The release of ACh from VA/VB neurons leads to the contraction
689 of the ventral body wall muscles and the activation of DD GABAergic motor neurons that
690 release GABA on the opposite side of the worm, causing relaxation of the dorsal body wall
691 muscles. Conversely, activation of the DA and DB cholinergic motor neurons produces
692 contraction of the dorsal body wall muscles and activates the VD GABAergic motor neurons.
693 The VD GABAergic motor neurons release GABA, causing relaxation of the ventral body wall
694 muscles, and thus contralateral inhibition. **B.** Quantification of body shortening in response to
695 anterior touch. Data are represented as mean ± SD. n= 50-70 animals per genotype distributed
696 across four independent experiments. Kruskal-Wallis analysis with Dunn's post-test for multiple

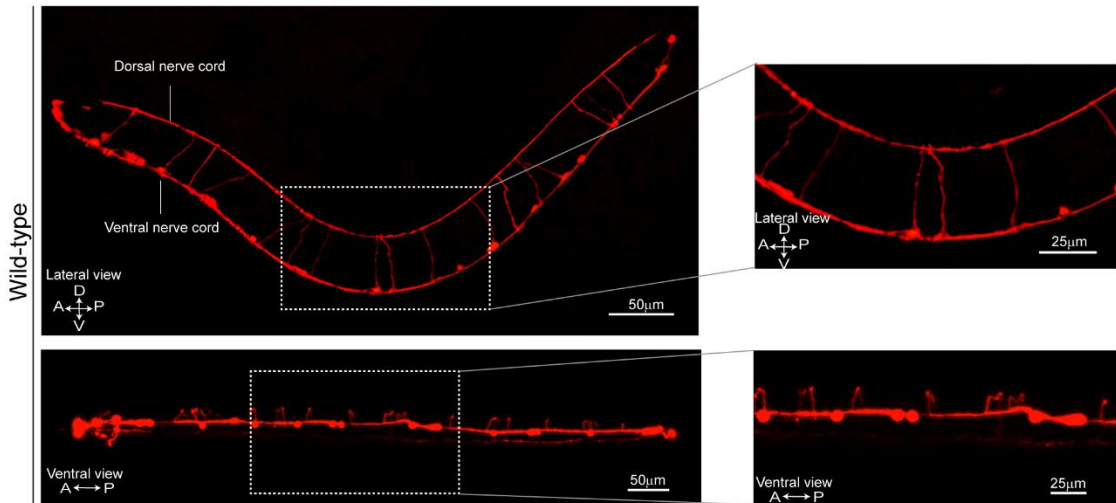
697 comparisons was performed. **C.** (Top) Scheme of *C. elegans* escape response in NGM agar.
698 After eliciting the escape response by an anterior gentle touch, the omega turns were classified
699 as closed (head and tail are in contact) or open (no contact between head and tail). (Bottom)
700 Quantification of % closed omega turns/ total omega turns. At least six independent trials for
701 each condition were performed (n= 20-25 animals per genotype/trial). Data are represented as
702 mean \pm SD. One-way ANOVA with Tukey's post-test for multiple comparisons was performed.
703 **D-E.** Light-evoked elongation/contraction of animals expressing Channelrhodopsin (ChR2) in
704 GABAergic (D) and Cholinergic (E) motorneurons. Animals were filmed before, during, and after
705 a 5-second pulse of 470 nm light stimulus (15 frames/s). The body area in each frame was
706 automatically tracked using a custom FIJI-Image J macro. The averaged area of each animal
707 during the first 125 frames (0-5 s) established a baseline for normalizing light-induced body area
708 changes. To compare the changes induced by optogenetic activity between different strains, the
709 body area measurements for each animal were averaged from second 6 (1 second after the
710 blue light was turned on) to second 9 (1 second before the light was turned off). These mean \pm
711 SD values are depicted in the bar graph shown to the right of each trace representation (n=40-
712 55 animals per genotype). Tukey's multiple comparisons method following One-Way ANOVA
713 was performed for D, while Dunn's multiple comparisons test after Kruskal-Wallis analysis was
714 used in E. **F-G.** Manual Measurement of Body length and width upon Optogenetic Stimulation of
715 GABAergic (F) and Cholinergic (G) neurons. At the 2.5-second time point of light stimulation, we
716 manually measured both the width and length of multiple animals and compared these
717 measurements with the corresponding areas obtained from automated analysis (see Materials
718 and Methods). The width of the worms remained relatively constant, highlighting that the
719 alterations in body area primarily stem from changes in the animal's length. One-way ANOVA
720 with Tukey's post-test for multiple comparisons was performed. Data are shown as mean \pm SD.
721 (ns p > 0.05; * p \leq 0.05; ** p \leq 0.01; *** p \leq 0.001; **** p \leq 0.0001).

722

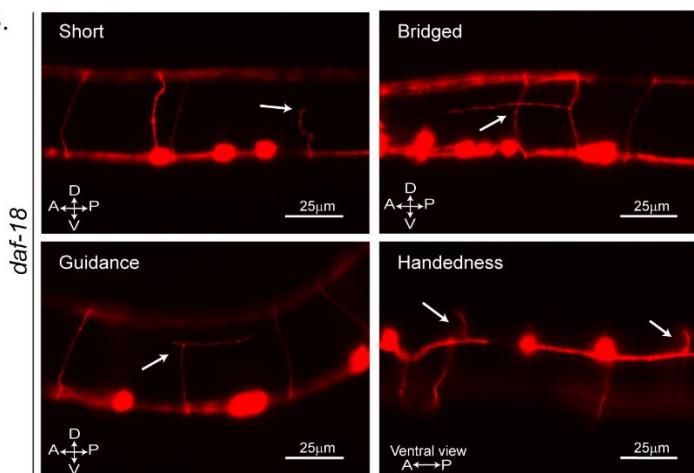
723 **Figure 3**

Figure 3

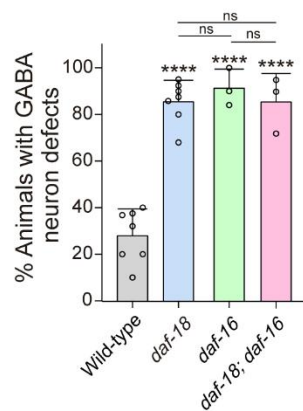
A. *Punc-47::mCherry* (GABAergic neurons)



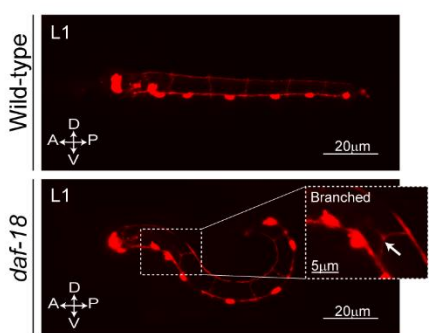
B.



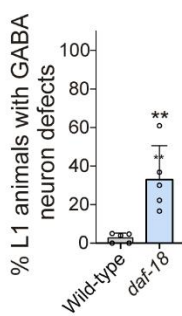
C.



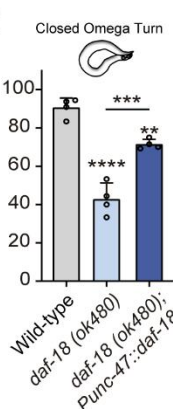
D.



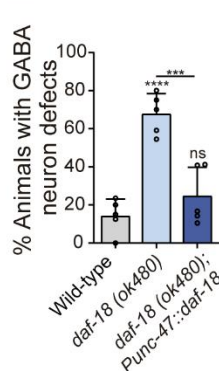
E.



F.



G.



725 **Figure 3. *daf-18/PTEN* mutants show neurodevelopmental defects in GABAergic motor -**
726 **neurons.**

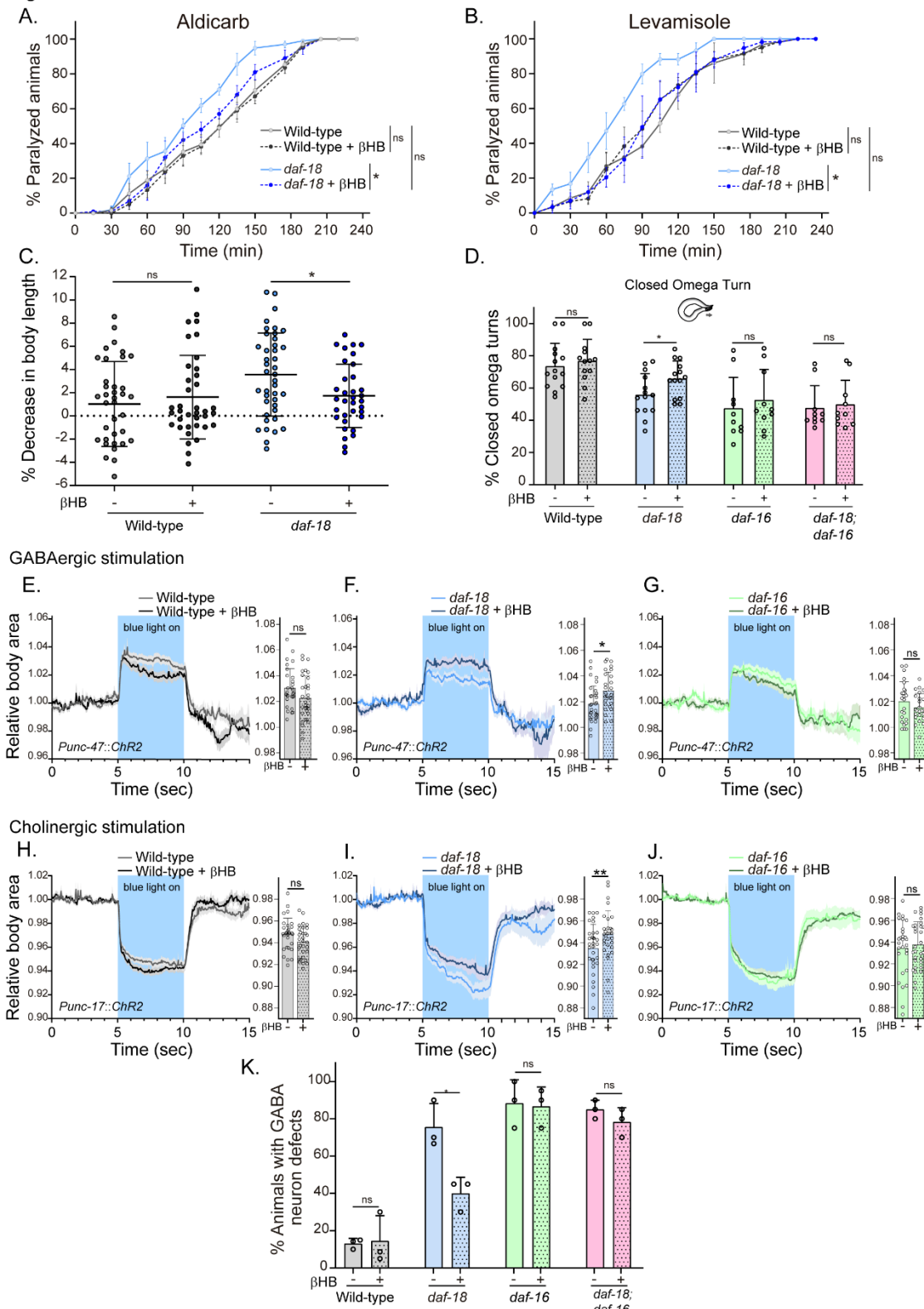
727 **A.** Representative images of wild-type animals expressing *mCherry* in the GABAergic motor
728 neurons are shown laterally (Top) and ventrally (Bottom). In the insets, commissures are
729 depicted at a higher resolution. Note that in the ventral view, all the processes travel through the
730 right side of the animal's body. **B.** Representative images of commissure defects observed in
731 *daf-18 (ok480)* mutants (arrows). The defects shown are: Short, commissure length less than
732 half of nematode width; Bridged, neighboring commissures linked by a neurite; Guidance,
733 commissures that do not reach dorsal nerve cord; and Handedness, commissure running along
734 the opposite side of the animal's body. **C.** Quantification of GABAergic system defects. Each bar
735 represents the mean \pm SD. One-way ANOVA and Tukey's multiple comparisons test were used
736 for statistics (ns p > 0.05; **** p \leq 0.0001). At least three independent trials for each condition
737 were performed (n: 20-25 animals per genotype/trial). **D.** Representative image of L1 animals (1
738 h post-thatch) expressing *Punc-47::mCherry* in wild-type (Top) and *daf-18(ok480)* mutant
739 (Bottom) backgrounds. In this larval stage, only six GABAergic DD motor neurons are born. The
740 inset shows a typical defective (branched) commissure. **E.** Quantification of GABAergic system
741 defects in L1s. Each bar represents the mean \pm SD. Two-tailed unpaired Student's t test. (**p \leq
742 0.01). At least five independent trials for each condition were performed (n: ~20 animals per genotype/trial). **F-G.**
743 Quantification of closed omega turns/total omega turns and commissure defects in GABAergic neurons of
744 animals expressing *daf-18/PTEN* solely in GABAergic neurons. One-way ANOVA and Tukey's
745 multiple comparisons test were used for statistics (ns p > 0.05; **p \leq 0.01; *** p \leq 0.001; **** p \leq
746 0.0001). At least four independent trials for each condition were performed (n: 15-20 animals
747 per genotype/trial)

748 A-Anterior; P-Posterior; D-Dorsal; V-Ventral.

749

750 **Figure 4**

Figure 4



751

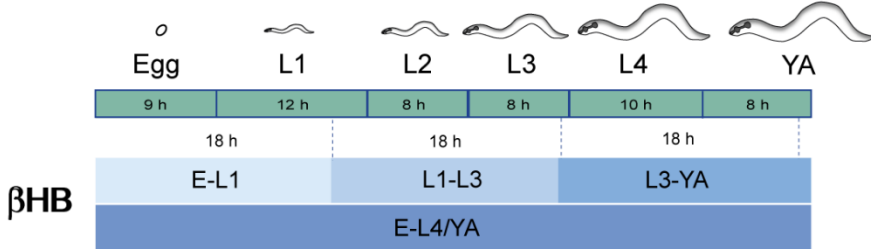
752 **Figure 4. Dietary βHB supplementation ameliorates GABAergic deficits in *daf-18/PTEN***
753 **mutants.**

754 Animals were exposed to βHB (20 mM) throughout development (from embryo to L4/young
755 adults). **A and B.** Quantification of paralysis induced by cholinergic drugs. At least four
756 independent trials for each condition were performed (n: 20-25 animals per genotype/trial). Two-
757 tailed unpaired Student's t-test (ns p > 0.05; *p ≤ 0.05; ** p ≤ 0.01) was used to compare βHB
758 treated and untreated animals. **C.** Measurement of body length in response to anterior touch.
759 n= 30-40 animals per genotype distributed across three independent experiments. Two-tailed
760 unpaired Student's t-test was used to compare βHB treated and untreated animals (ns p > 0.05;
761 * p ≤ 0.05). **D.** Quantification of closed omega turns/total omega turns during the escape
762 response. At least eight independent trials for each condition were performed (n= 20 animals
763 per genotype/trial). Results are presented as mean ± SD. Two-tailed unpaired Student's t test
764 (ns p > 0.05; *p ≤ 0.05). **E-J.** Light-evoked elongation/contraction of animals expressing ChR2 in
765 GABAergic (E-G) and cholinergic (H-J) motorneurons. The mean body area (mean ± SD) during
766 3 seconds of the light pulse is depicted in the bar graph shown to the right of each trace
767 representation (see figure 2) (n=25-35 animals per condition). Two-tailed unpaired Student's t-
768 test (ns p > 0.05; *p ≤ 0.05; ** p ≤ 0.01). **K.** Quantification of commissure defects in GABAergic
769 neurons. Results are presented as mean ± SD. Two-tailed unpaired Student's t-test (ns p >
770 0.05; *p ≤ 0.05). At least three independent trials for each condition were performed (n= ~20
771 animals per genotype/trial).

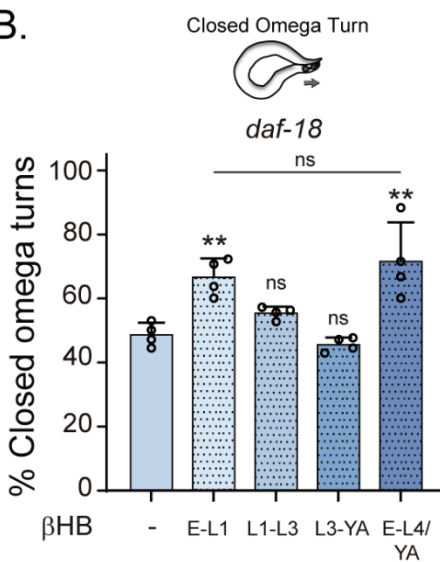
772

Figure 5

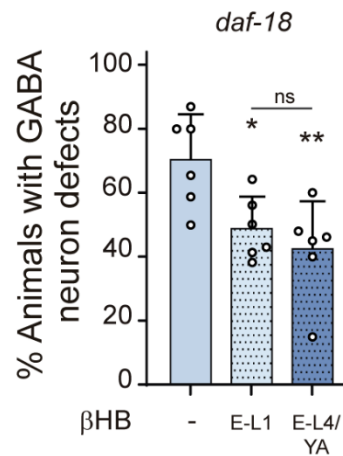
A.



B.



C.



773

774 **Figure 5. Early developmental stages are critical for β HB-modulation of GABAergic
775 signaling.**

776 **A.** Animals were exposed to β HB-enriched diet for 18-h periods at different developmental
777 stages: i) E-L1 covered *ex-utero* embryonic development (~ 9 h) and the first 8-9 h of the L1
778 stage; ii) L1-L3 covered the latter part of the L1 stage (~3-4 h), the entire L2 stage (~8 h), and
779 most of the L3 stage (~6-7 h), iii) L3-YA(Young Adult) spanned the latter part of the L3 stage (~
780 1-2 h), the entire L4 stage (~10 h), and the first 6-7 h as adults, and iv) E-L4/YA implies
781 exposure throughout development (from embryo to Young Adult). **B-C.** Quantification of closed
782 omega turns/total omega turns in *daf-18/PTEN* (B) and GABAergic commissure defects (C) in *daf-18/PTEN*
783 mutants exposed to β HB at different developmental intervals. Four and six independent trials for each

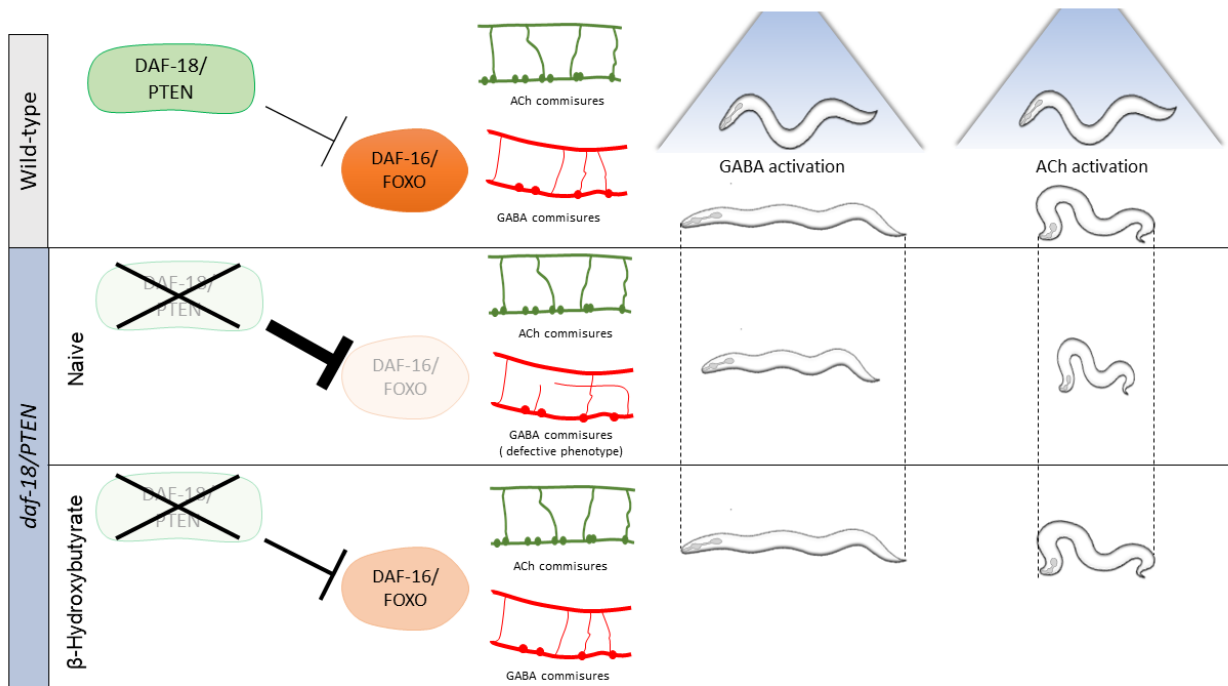
784 condition were performed in B and C, respectively (n=: 20-25 animals per genotype/trial).

785 Results are presented as mean \pm SD. One-way ANOVA with Tukey's post-test for multiple

786 comparisons was performed (ns p > 0.05; *p \leq 0.05; ** p \leq 0.01).

787

788 **Graphical Abstract**



789

790

791 **Materials and methods**

792 **C. elegans culture and maintenance.** All *C. elegans* strains were grown at room temperature
793 (22°C) on nematode growth media (NGM) agar plates with OP50 *Escherichia coli* as a food
794 source. The wild-type reference strain used in this study is N2 Bristol. Some of the strains were
795 obtained through the *Caenorhabditis* Genetics Center (CGC, University of Minnesota). Worm
796 population density was maintained low throughout their development and during the assays. All
797 experiments were conducted on age-synchronized animals. This was achieved by placing
798 gravid worms on NGM plates and removing them after two hours. The assays were performed
799 on the animals hatched from the eggs laid in these two hours.

800 Transgenic strains were generated by microinjection of plasmid DNA containing the construct
801 *Punc-47::daf-18cDNA* (kindly provided by Alexandra Byrne, UMASS Chan Medical School) at
802 20 ng/μL into the germ line of (*daf-18(ok480)*; *lin-15(n765ts)*) double mutants with the co-
803 injection marker *lin-15* rescuing plasmid pL15EK (80 ng/μL). At least three independent
804 transgenic lines were obtained. Data are shown from a single representative line.

805 The strains used in this manuscript were:

806 CB156 *unc-25(e156)* III
807 MT6201 *unc-47(n2409)* III
808 CB1375 *daf-18(e1375)* IV
809 OAR144 *daf-18(ok480)* IV
810 GR1307 *daf-16(mgdf50)* I
811 OAR115 *daf-16(mgDf50)* I; *daf-18(ok480)* IV
812 OAR161 *daf-18(ok480)*; *wpEx173[Punc-47::daf-18 + myo-2::GFP]*
813 LX929 *vsIs48[Punc-17::gfp]*
814 IZ629 *ufls34[Punc-47::mCherry]*
815 OAR117 *ufls34[Punc-47::mCherry; daf-18(ok480)]*
816 OAR118 *vsIs48[Punc-17::GFP]; daf-18(ok480)]*

817 OAR142 *ufis34*[*Punc-47::mCherry*; *daf-16(mgDf50)*]
818 OAR143 *ufis34* [*Punc-47::mCherry*; *daf-16(mgDf50)*; *daf-18(ok480)*]
819 CF1553 *muls84*[(*pAD76*) *Psod-3::gfp* + *rol-6(su1006)*]
820 OAR140 *muls84*[(*pAD76*) *Psod-3::gfp* + *rol-6*]; *daf-18(ok480)*
821 OAR141 *muls84*[(*pAD76*) *Psod-3::gfp* + *rol-6*]; *daf-16(mgDf50)*
822 OH99 *mgls18*[*Ptx-3::gfp*]
823 OAR83 *daf-18(ok480)*; *mgls18*[*Ptx-3::gfp*]
824 MT13471 *nls121*[*Ptph-1::gfp*]
825 OAR112 *nls121*[*Ptph-1::gfp*; *daf-18(ok480)*
826 IZ805 *ufls53*[*Punc-17::ChR2*]
827 ZM3266 *zxls3*[*Punc-47::ChR2::YFP*]
828 OAR177 *ufls53*[*Punc-17::ChR2::YFP*; *daf-18(ok480)*
829 OAR178 *ufls53*[*Punc-17::ChR2::YFP*]; *daf-16(mgDf50)*
830 OAR179 *zxls3*[*Punc-47::ChR2::YFP*]; *daf-18(ok480)*
831 OAR180 *zxls3*[*Punc-47::ChR2::YFP*]; *daf-16(mgDf50)*
832 TJ1052 *age-1(hx546)*
833 GR1310 *akt-1(mg144)*
834 GR1318 *pdk-1(mg142)*
835 JT9609 *pdk-1(sa680)*
836 VC204 *akt-2(ok393)*
837 VC222 *raga-1(ok386)*
838 KQ1366 *rict-1(ft7)*
839
840 **Paralysis assays.** Paralysis assays were carried out in standard NGM plates with 2 mM
841 aldicarb (Sigma-Aldrich) or 0.5 mM levamisole (Alfa Aesar). 25-30 L4 worms were transferred to

842 each plate and paralyzed animals were counted every 15 or 30 minutes. An animal was
843 considered paralyzed when it did not respond after prodding three times with a platinum wire on
844 the head and tail¹. At least four independent trials with 25-30 animals for each condition were
845 performed. The area under the curve (AUC) for each condition in each experiment was used for
846 statistical comparisons

847 **Escape response.** Escape response assays were performed on NGM agar plates seeded with
848 a thin bacterial lawn of OP50 *E. coli*. To maintain tight control of growth and moisture, 120 µL of
849 bacteria were seeded 24 hours before the assay and grown overnight at 37° C. The day of the
850 assay, young adult worms were transferred to the plates and allowed to acclimate for at least 5
851 min. Omega turns were induced by gentle anterior touch with fine eyebrow hair and were
852 classified as closed when the worm touched the tail with its head as previously described².
853 Between 4 and 7 independent trials with ~20 animals for each condition were performed.

854 **Body length assays.** Body length measurements were performed in standard NGM agar plates
855 without bacteria. Young adult synchronized worms were transferred into the plates and allowed
856 to acclimate for at least 5 min. Worms were recorded with an Amscope Mu300 camera. Animal
857 body length, before and after touching with a platinum pick, was measured using FIJI Image J
858 software. Quantification of body shortening after touching was calculated as the decrease of
859 body length related to the length of the animal before being touched.

860 **Commissure analysis.** Synchronized L1 or L4 animals carrying the fluorescence reporters
861 *vsIs48* (*Punc-17::GFP*, cholinergic neurons) or *ufls34* (*Punc-47::mCherry*, GABAergic neurons)
862 were immobilized with sodium azide (0.25 M) on 2% agarose pads. Commissures of GABAergic
863 and cholinergic neurons were scored with a Nikon Eclipse TE 2000 fluorescence microscope. A
864 commissure is generally composed of a single process, and occasionally two neurites that
865 extend together dorsally. Defects on commissures, including guidance defects, abnormal
866 branching, and incomplete commissures were classified similarly to previous reports³. The

867 percentage of animals with at least one commissure defect was calculated for each neuronal
868 class (e.g., cholinergic or GABAergic). At least three trials (~20 animals per condition in each
869 trial) were analyzed for each individual experiment. Representative images shown in the figures
870 were collected using laser confocal microscopy (ZEISS LSM 900 with AirScan II) with 20 \times and
871 63 \times objectives.

872 **β -hydroxybutyrate assays.** Worms were exposed to 20 mM DL-3-hydroxybutyric acid sodium
873 salt (Acros Organics) on NGM agar plates seeded with *E. coli* OP50. We synchronized the
874 animals similarly to other experiments by placing gravid animals in NGM plates containing 20
875 mM of β HB and removing them after two hours.

876 For experiments involving exposure at different developmental stages, animals were transferred
877 between plates with and without β HB as needed. For the earliest exposure, eggs were laid on
878 plates containing 20 mM of β HB and then transferred to drug-free plates to complete their
879 development. Conversely, for later exposures, animals were born on β HB-free plates and
880 subsequently transferred to β HB-containing plates at the specified time (See Figure 5).

881 ***sod-3* expression.** *sod-3* expression levels were analyzed in transgenic strains containing the
882 transcriptional reporter *mul84*, as described previously^{4,5}. Synchronized L4 animals were
883 anesthetized with Sodium Azide (0.25 M) and mounted on 2% agarose pads. Images were
884 collected using a Nikon Eclipse TE 2000 fluorescence microscope. GFP fluorescence intensity
885 was quantified in same-sized ROIs at the head of the animal using Image J FIJI software.
886 Results were normalized to control conditions (wild-type individuals without β HB). ~35-60
887 animals for each genotype/condition were analyzed.

888 **Optogenetic assays.** We examined young adult animals (6-8 hours post-L4 stage) that express
889 Channelrhodopsin (ChR2) in either GABAergic (*Punc-47::ChR2*) or cholinergic neurons (*Punc-*
890 *17::ChR2*). We transferred these animals to a NGM 6mm agar plate without food, let them

891 acclimate for 5 minutes, and recorded each animal at 15 frames per second using an Allied
892 Vision Alvium 1800 U-500m camera. To stimulate neuronal activity, we exposed the animals to
893 470 nm light pulses for 5 seconds. These light pulses were delivered using a custom Python
894 script (VIMBA Peron) to an Arduino Uno microcontroller, which operated a Mightex compact
895 universal LED controller (Mightex SLC-MA02-U). The light emission was achieved through a
896 Mightex High-Power LED Collimator Source (LCS-0470-03-11). To precisely track the changes
897 in the worm's body, we continuously monitored its area from 5 seconds before the light stimulus,
898 during the light stimulus and until 5 seconds afterward. We developed a FIJI-Image J macro
899 capable of automatically tracking the body area in each frame, capitalizing on the clear contrast
900 between the worm's body and the background. As demonstrated in Movies 3 and 4, changes in
901 body area directly corresponded to alterations in the animal's length. To compare the changes
902 induced by optogenetic activity between different strains, the body area measurements for each
903 animal were averaged from second 6 (1 second after the blue light was turned on) to second 9
904 (1 second before the light was turned off). These average values were used for the statistical
905 comparisons detailed in each figure legend.

906 To validate our measurement system, we manually measured the width of 6-8 animals at the
907 2.5-second point of light stimulation and compared them to the body area and length. Our
908 observations consistently showed that, regardless of whether the area increased or decreased
909 (depending on the activation of GABAergic or cholinergic neurons), the width remained mostly
910 unchanged (Figure 2F and 2G). Therefore, the observed changes in the animal's area
911 measured by our FIJI-Image J macro indeed represent alterations in the animal's length.

912 **AIY and HSN analysis.** Synchronized L4 or Young Adult worms carrying the fluorescence
913 reporters *Ptx-3::gfp* (AIY interneurons expressing GFP) and *Pth-1::gfp* (HSN expressing GFP)
914 were immobilized with sodium azide (0.25 M) on 2% agarose pads and analyzed with a Nikon
915 Eclipse TE 2000 fluorescence microscope. AIY neurons morphology were sorted in qualitative

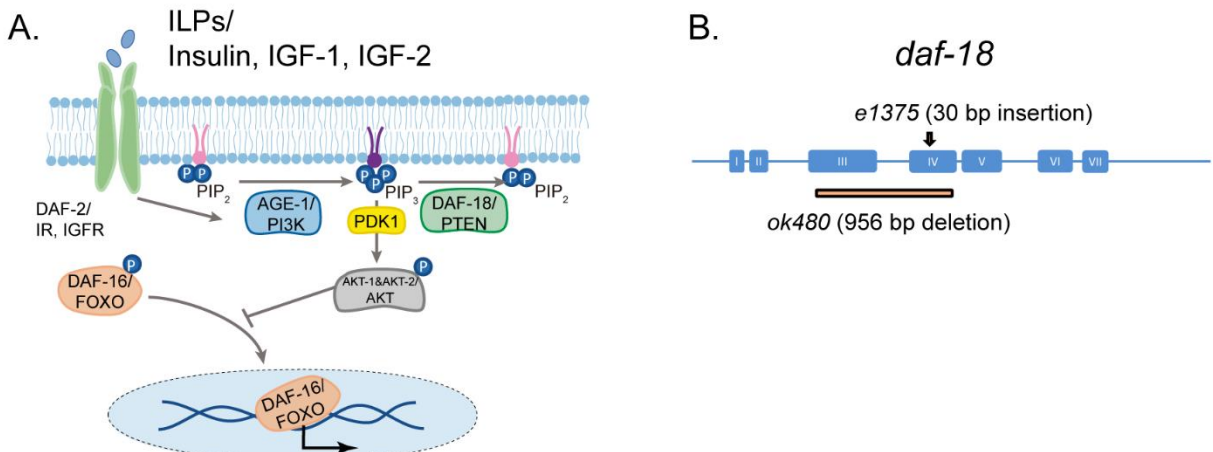
916 categories (see figure legend) while the migration of HSN was classified in quantitative
917 categories using ImageJ software.

918 **Statistical Analysis.** The results presented in each figure are the average of at least three
919 independent assays. Bars represent mean \pm SD. Typically, one-way ANOVA was employed for
920 analyzing multiple parametric samples, and Tukey's post hoc test was used for pairwise
921 comparisons among all groups. For comparisons against a control group, Dunnett's post hoc
922 test was used. For multiple non-parametric samples, the Kruskal-Wallis test was applied
923 followed by Dunn's post hoc test, which was also utilized for comparisons against the control
924 group. In cases where comparisons were made between two independent conditions, a t-test
925 was utilized for parametric data, while the Mann-Whitney U test was employed for non-
926 parametric data. We used the software GraphPad Prism version 6.01 to perform statistics. The
927 statistical information is indicated in the figure legends. For all assays, the scoring was done
928 blinded. All raw data are available in Open Science Framework
929 (https://osf.io/mdpgc/?view_only=3edb6edf2298421e94982268d9802050).

930

931

Supplementary Figure 1



932

933

934 **Supplementary Figure 1. *daf-18/PTEN* is the negative modulator of the PI3K/AKT pathway.**

935 **A.** *daf-18/PTEN* encodes a lipid and protein phosphatase that hydrolyzes phosphatidylinositol

936 (3,4,5)-trisphosphate (PIP₃) to phosphatidylinositol-4,5-bisphosphate (PIP₂). It is the main

937 negative modulator of PDK and AKT activity. In *daf-18/PTEN* mutants, AKT is overactivated

938 leading to high levels of DAF-16/FOXO phosphorylation that prevents the translocation of this

939 transcription factor to the nucleus. **B.** Gene structure of *daf-18*. Coding sequences are

940 represented by blue boxes. The *daf-18(e1375)* mutant allele inserts a 30 bp sequence in exon

941 IV. This insertion occurs downstream of the phosphatase catalytic domain and causes a

942 frameshift that leads to premature truncation of the protein. This *e1375* mutation partially

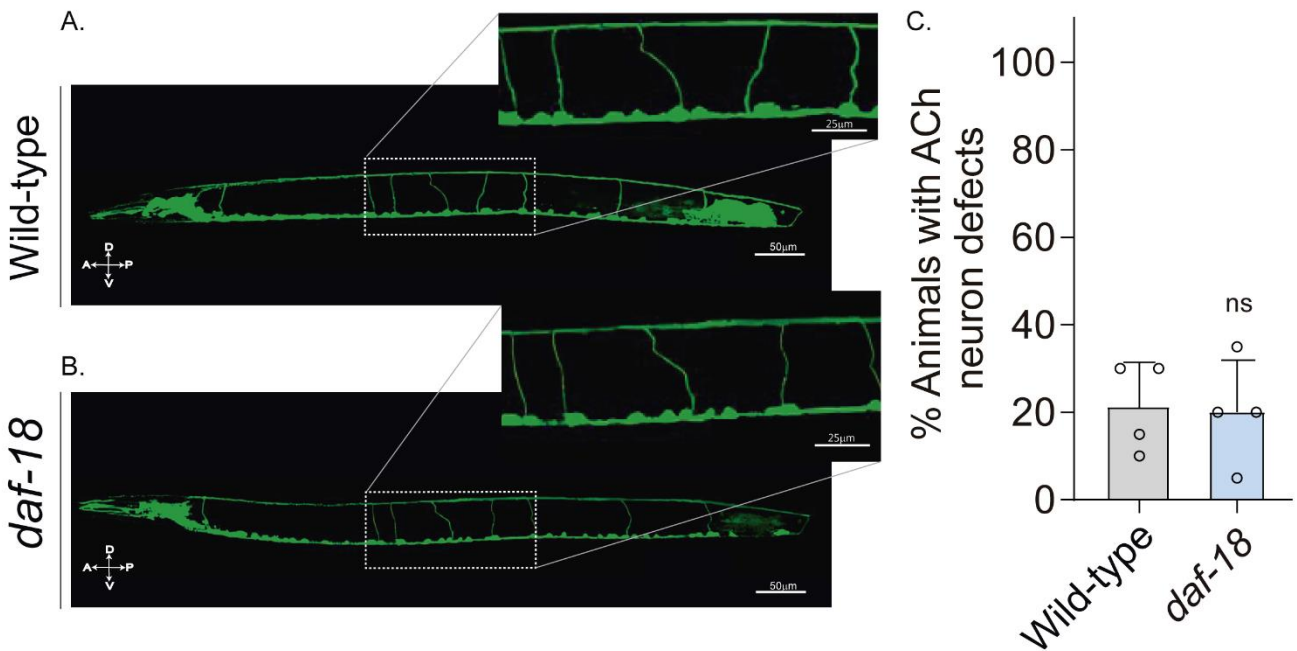
943 reduces DAF-18 function. The *daf-18(ok480)* allele contains a 956 bp deletion that removes

944 most of exon 3 and exon 4 and is generally considered to be a null allele.

945

Supplementary Figure 2

Punc-17::GFP (Cholinergic neurons)



946

947 **Supplementary Figure 2. *daf-18/PTEN* mutations do not affect excitatory cholinergic
948 motor-neuron morphology**

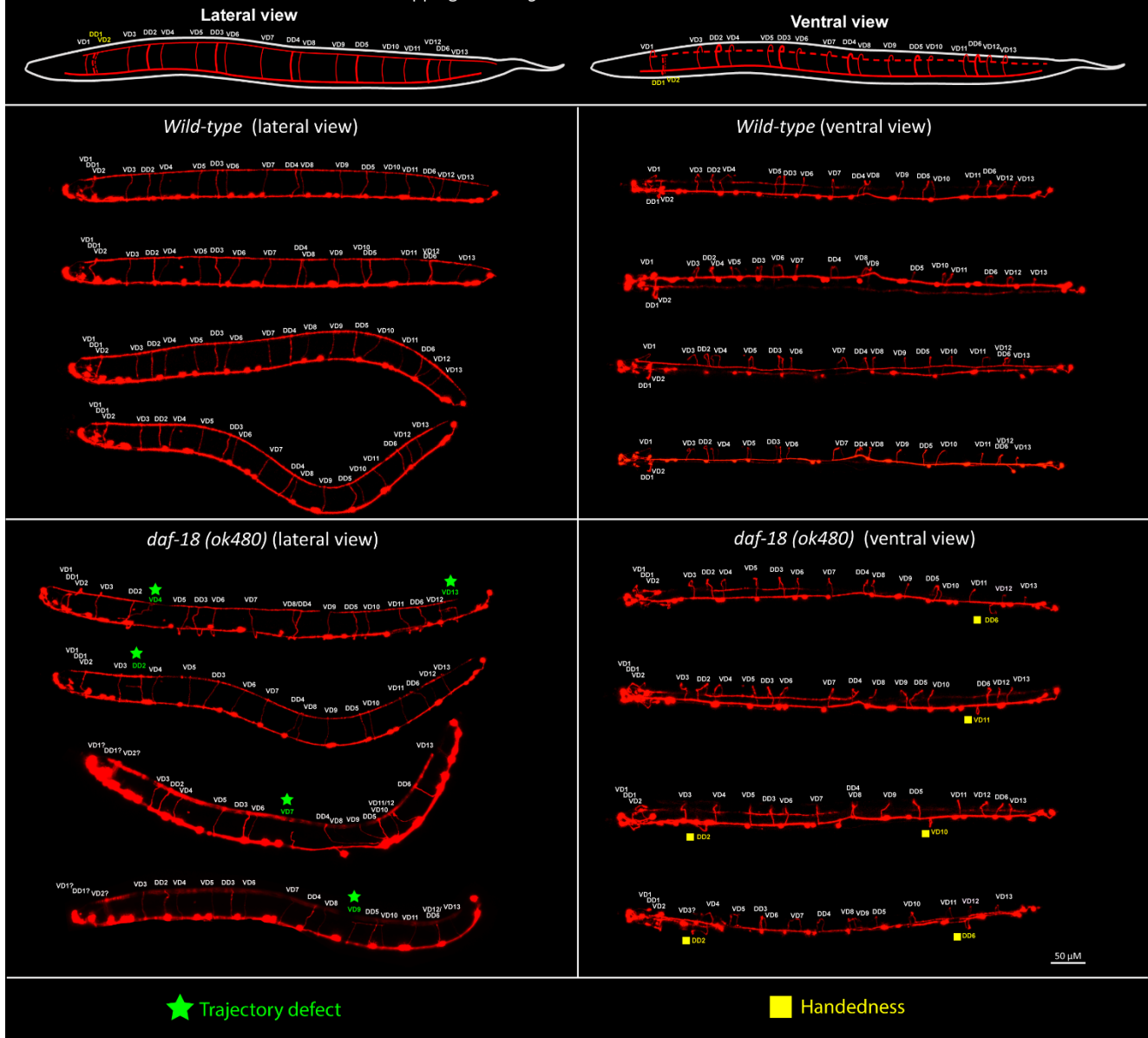
949 **A.** Representative images of animals expressing GFP in the cholinergic neurons. In the insets,
950 the commissural processes can be appreciated with higher resolution. **B.** Quantification of
951 cholinergic system defects. Each bar represents the mean \pm SD for at least four trials (~ 20
952 animals per trial). Statistical significance between the strains was determined by two-tailed
953 unpaired Student's t-test. (ns $p > 0.05$).

954 A anterior; P Posterior; D Dorsal; V Ventral.

955

Supplementary Figure 3

Canonical Mapping of *C. elegans* VDs and DDs commissures



956

957

958 **Supplementary Figure 3. *daf-18/PTEN* deficiencies affect DDs and VDs GABAergic**

959 **neurons**

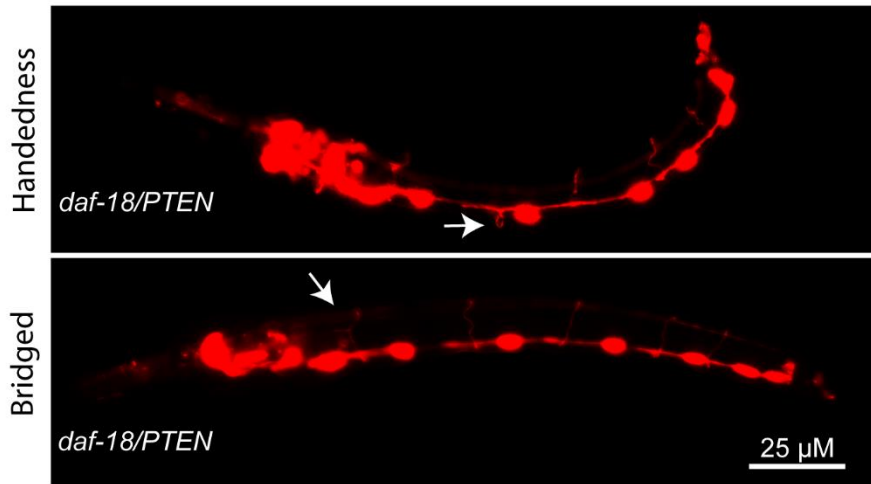
960 **Top**, schematic representation of the location of the commissures belonging to the two types of
 961 GABAergic neurons in lateral and ventral views (based on^{6,7}). **Below**, representative images of
 962 wild-type and *daf-18(ok 480)* worms viewed laterally (left) and ventrally (right). Note that in both
 963 lateral and ventral views (handedness errors) defects appear in both DDs and VDs neurons.

964 Errors in the first 3 commissures were not considered due to the difficulty of identifying the
965 commissures corresponding to VD1, DD1, and VD2 neurons.

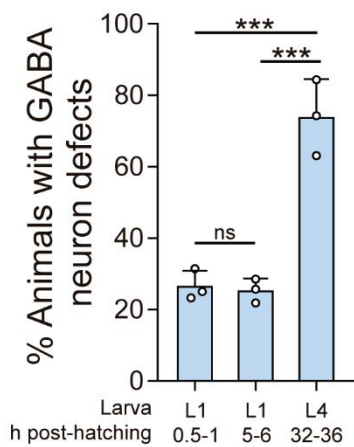
966

Supplementary Figure 4

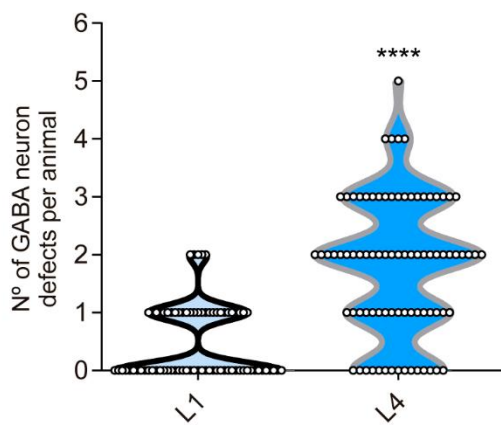
A.



B.



C.



967

968 **Supplementary Figure 4. DD-GABAergic neurons show defects in recently hatched L1**
969 **animals of *daf-18/PTEN* mutants**

970

971 **A.** Representative images of the most common errors observed in L1 animals (1 h post-hatch)
972 of *daf-18* mutants. These types of defects, plus others such as short commissures and guidance
973 defects, are also observed in L4 animals (Figure 3 and S3). **B.** GABAergic commissure defects
974 were quantified in *daf-18/PTEN* mutants at various developmental stages: 0.5-1 h post-hatching
975 (early L1 larva), 5-6 h post-hatching (mid-L1 larva), and L4 stage (32-36 h post-hatching). Three

976 independent trials for each condition were performed, with at least 30 animals per condition/trial.

977 Results are presented as mean \pm SD. A one-way ANOVA with Tukey's post-hoc test was used

978 (ns $p > 0.05$; ** $p \leq 0.01$; *** $p \leq 0.001$; **** $p \leq 0.0001$). **C.** Quantification of the number of errors

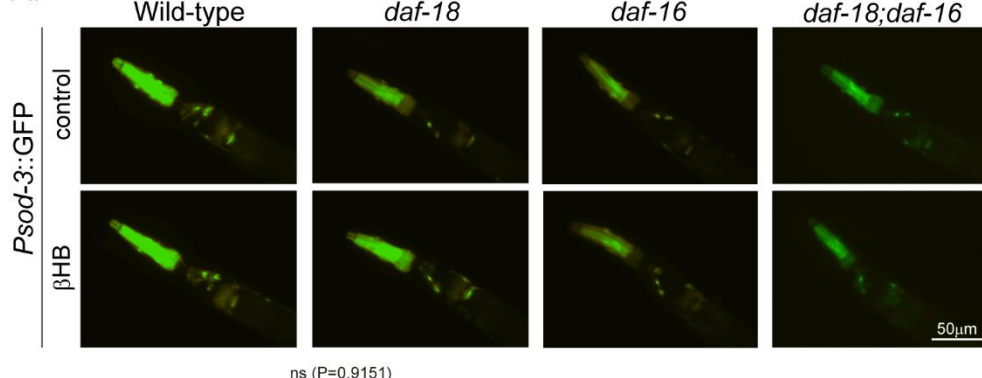
979 per animal in L1 and L4 larvae. Statistical significance was determined by Mann Whitney test

980 (**** $p \leq 0.0001$; n=75-80)

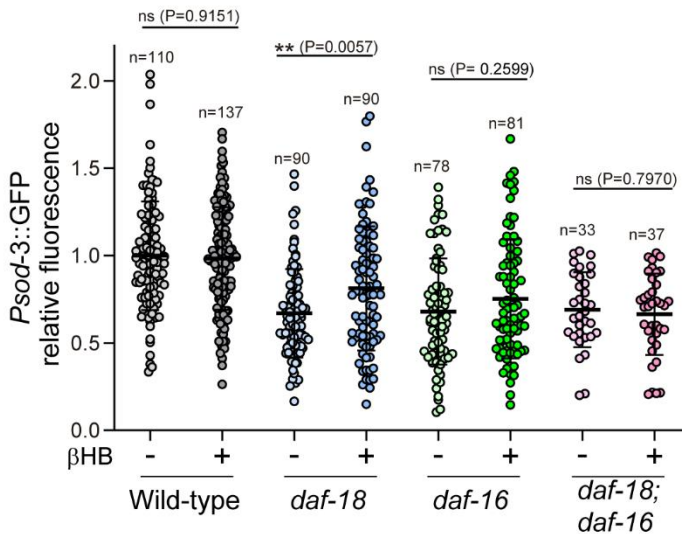
981

Supplementary Figure 5

A.



B.



982

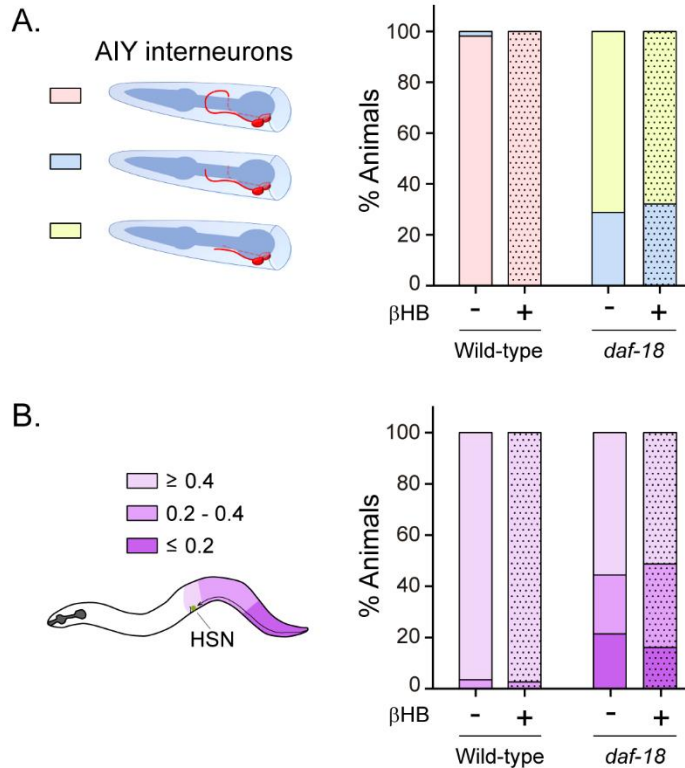
983 **Supplementary Figure 5. Exposure to β HB induces *sod-3* expression in *daf-18/PTEN*, but**

984 not in *daf-16/FOXO* mutants.

985 **A.** Representative fluorescence images (20X magnification) of worms expressing *Psod-3::GFP*
986 in different genetic backgrounds (wild-type, *daf-18(ok480)*, *daf-16(mgDf50)* and *daf-18(ok480)*;
987 *daf-16(mgDf50)*) upon exposure to β HB (20mM). **B.** Corresponding quantification of the
988 fluorescence intensity per animal in the head. Scatter dot plot (line at the median) with the
989 relative expression of *Psod-3::GFP* normalized to naïve wild-type animals. Statistical
990 significance between the treatment and the corresponding control was determined by Mann
991 Whitney test (ns $p > 0.05$; ** $p \leq 0.01$; **** $p \leq 0.0001$, $n=40-90$).

992

Supplementary Figure 6



993

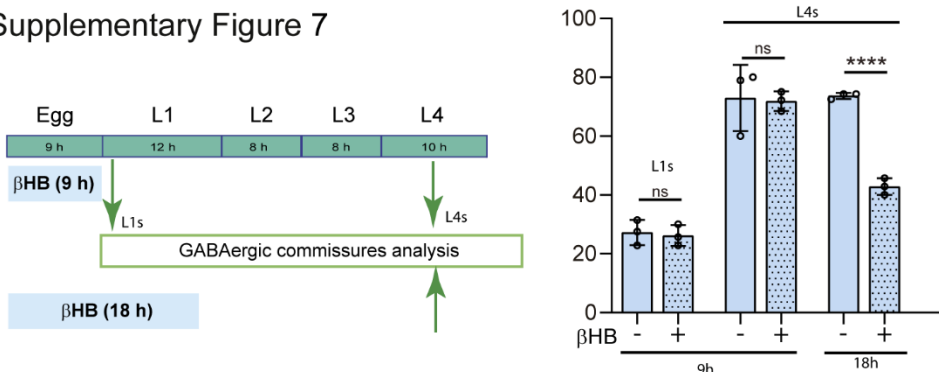
994 **Supplementary Figure 6. β HB does not prevent neurodevelopmental defects in AIY and**
995 **HSN neurons.**

996 **A.** AIY processes were visualized in transgenic animals expressing cytoplasmic *GFP* in AIY
997 neurons (*Ptx-3b::GFP*) in wild-type and *daf-18(ok480)* mutant backgrounds. AIY neuronal
998 growth defects were quantified as described before (Christensen *et al.*, 2011). Left: Scheme of
999 AIY morphology and location in the nematode nerve ring. Blue, pharynx; red, AIY interneurons.
1000 Pink: wild-type AIY morphology. The two interneurons meet at the dorsal midline. Light blue and
1001 yellow: denote different levels of AIY neurite truncation. Right: Percentage of animals with
1002 truncated neurites in wild-type and *daf-18(ok480)* mutants under exposure (or not) to β HB (20
1003 mM). **B.** HSN were visualized in transgenic animals expressing *GFP* in serotonergic neurons
1004 (*Pth-1::GFP*) in wild-type and *daf-18(ok480)* mutant backgrounds. HSN under-migration
1005 defects were identified as described before (Kennedy *et al.*, 2013). Left: Schematic
1006 representation of the HSN migratory route during embryogenesis and the corresponding

1007 location of the HSN (green circle) in a young adult animal. Only one of two bilaterally symmetric
1008 HSNs is illustrated. Colors show information about the position of HSNs: Light purple: complete
1009 migration (≥ 0.4), middle purple: intermediated migration ($>0.2 < 0.4$), dark purple: unmigrated
1010 (≤ 0.2). Right: Quantification of the percentage of animals with different HSN migration positions
1011 (the most under-migrated neuron of each animal is considered). Bars represent the mean
1012 values of at least three independent experiments. Note that there is no significant effect with
1013 β HB treatment compared to controls.

1014

Supplementary Figure 7



1015

1016

1017 **Supplementary Figure 7. βHB does not prevent neurodevelopmental defects in**
1018 **GABAergic neurons when applied exclusively during ex-utero embryonic development.**

1019

1020 Quantification of GABAergic commissure defects in L4-stage of *daf-18/PTEN* mutant animals
1021 exposed to βHB during the first 9 h post-egg laying (just before hatching) and 18 hours post-egg
1022 laying. The animals were then transferred to control plates without βHB and maintained until
1023 GABAergic commissures analysis. Scoring was performed in 0.5-1 h post-hatching (early L1
1024 larva) and L4 animals (green arrows). A two-tailed unpaired Student's t-test was used for
1025 statistical analysis. Data represent three independent trials with at least 20 worms per trial.
1026 Results are presented as mean ± SD.

1027 **References**

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