Optical Silencing of *C. elegans* Cells with Arch Proton Pump

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**Abstract**

*Background:* Optogenetic techniques using light-driven ion channels or ion pumps for controlling excitable cells have greatly facilitated the investigation of nervous systems *in vivo*. A model organism, *C. elegans*, with its small transparent body and well-characterized neural circuits, is especially suitable for optogenetic analyses.

*Methodology/Principal Findings:* We describe the application of archaerhodopsin-3 (Arch), a recently reported optical neuronal silencer, to *C. elegans*. Arch::GFP expressed either in all neurons or body wall muscles of the entire body by means of transgenes were localized, at least partially, to the cell membrane without adverse effects, and caused locomotory paralysis of worms when illuminated by green light (550 nm). Pan-neuronal expression of Arch endowed worms with quick and sustained responsiveness to such light. Worms reliably responded to repeated periods of illumination and non-illumination, and remained paralyzed under continuous illumination for 30 seconds. Worms expressing Arch in different subsets of motor neurons exhibited distinct defects in the locomotory behavior under green light: selective silencing of A-type motor neurons affected backward movement while silencing of B-type motor neurons affected forward movement more severely. Our experiments using a heat-shock-mediated induction system also indicate that Arch becomes fully functional only 12 hours after induction and remains functional for more than 24 hour.

*Conclusions/Significance:* Arch can be used for silencing neurons and muscles, and may be a useful alternative to currently widely used halorhodopsin (NpHR) in optogenetic studies of *C. elegans*.

**Introduction**

Optogenetics is a novel technique for controlling excitable cells such as neurons, muscles and neuroendocrine cells. By exploiting light-driven ion channels or ion pumps derived from microbes and lower eukaryotes, optogenetics enables repression and activation of cells when illuminated with a particular wavelength of light. By enabling temporal manipulation of neuronal activity through an easy-to-induce and non-invasive treatment, this technique has the potential to revolutionize research on the subject of excitable cells *in vivo* [1–3].

A nematode, *Caenorhabditis elegans* (*C. elegans*) has a well-characterized neural anatomy: the complete structure as well as the wiring patterns of each neuron of this organism have been determined by reconstruction from serial electron micrograph sections [4]. Together with the availability of various easy-to-handle genetic tools, these characteristics make this model organism an attractive one for studying the functions of single targeted cells in neural networks [3]. On the other hand, the small size of *C. elegans* neurons and their poor accessibility with respect to the application of microelectrodes has hindered conventional electrophysiological approaches. Methods for investigating neural functions in *C. elegans* have thus been almost entirely limited to laser killing of neurons, administration of neuroactive drugs, or use of neural mutants. As neither temporary activation nor repression of neuronal activity can be achieved with traditional techniques such as laser killing or drug treatment, optogenetic methods will be of great benefit to the study of *C. elegans* neurons. Optogenetics allows the investigation of neuronal functions in freely moving animals, increasing the prospects for behavioral studies using *C. elegans*. Optogenetic approaches applied to *C. elegans*, using channelrhodopsin-2 for activation of cells, and halorhodopsin (NpHR) for silencing of cells, have been reported [6–8].

NpHR is a light-driven inward chloride pump present in *Natronomonas pharaonis* [9], which inhibits electrical excitation of genetically targeted cells under illumination by yellow light (580 nm) with millisecond temporal resolution. NpHR, however, is capable of only modest hyperpolarizing currents in response to light, and once activated, NpHR enters long-lasting inactive states [10]. Recently, an optogenetical application of another light-driven pump, Archaerhodopsin-3 (Arch), derived from halobacteria *halorubrum sodomense*, was reported [11]. Arch is an outward proton pump driven by green light (550 nm), and generates a neural silencing current at a lower level of illumination *in vitro* compared with NpHR. Additionally, Arch recovers spontaneously from light-dependent inactivation. It has been reported that Arch...
enables the silencing of neurons in the awake brain of a mouse [11]. Therefore, Arch may serve as another useful optogenetic tool for neural silencing experiments in vivo.

In this study, we examined the applicability of Arch for optogenetic manipulation of C. elegans cells in vivo. We found that worms forced to express Arch::GFP in neurons or muscle cells of the entire body stopped their locomotion and became paralyzed when illuminated with green light. Worms expressing Arch::GFP in different subsets of motor neurons exhibited distinct behavioral defects under green light illumination. We also examined the acquisition of Arch function and its stability in C. elegans cells by using a heat-shock gene induction system. We found that Arch becomes fully functional 12 hours after induction of expression, and remains functional for at least 24 hours.

Results

Arch::GFP Expression in C. elegans Cells

To examine whether Arch, a light-driven proton pump, can be used to repress the electrical activity of neurons and muscle cells in Caenorhabditis elegans (C. elegans), we first tried to express C-terminally GFP-tagged Arch (Arch::GFP) in worms by using the myo-3 gene promoter, which encodes a myosin heavy chain expressed specifically in the body wall and vulva muscles of the organism, and the promoters of the F25B3.3 and aex-3 genes, both of which drive neuronal-expression. Then, we analyzed the expression pattern of GFP and the locomotion behavior of transgenic worms under green light illumination.

In worms carrying the transgene myo-3p::Arch::gfp, ncEx301Ex[myo-3p::Arch::gfp], an intense GFP signal was observed in granular or vesicular structures close to the cell membrane and in myofibril-like structures of the body wall muscles. The signal was also detected at muscle cell boundaries, suggesting localization of Arch to the cell membrane. A weak GFP signal was seen in the cytoplasm, but not in the nucleus (Fig. 1A, B).

In worms carrying the transgene F25B3.3p::Arch::gfp, ncEx3034[F25B3.3p::Arch::gfp], and in worms carrying the transgene aex-3p::Arch::gfp, ncEx3026[aex-3p::Arch::gfp], GFP signals were observed in the soma and the axon of many neurons in the head, mid body and tail (Fig. 1C, E). The signal often demarcated the cell soma. An intense granular or vesicular GFP signal was sometimes detected in the soma and in axons (Fig. 1D, F).

All transgenic strains carrying Arch::gfp stopped locomotion and became paralyzed under green light illumination to the whole body. C. elegans worms do not have inherent rhodopsin genes [12], and it is presumed that the level of endogenous retinoids, if these even exist, is insufficient for the proper functioning of rhodopsins and it is presumed that the level of endogenous retinoids, if these even exist, is insufficient for the proper functioning of rhodopsins. Hence, we expressed Arch pan-neurally in C. elegans to examine the effect of Arch on the organism, and the promoters of the myo-3 and aex-3 genes, both of which drive neuronal-expression. We used a heat-shock gene induction system. We found that Arch becomes fully functional 12 hours after induction of expression, and remains functional for at least 24 hours.

Light Intensity Dependence of Arch Neuronal Silencing Activity

In a manner similar to that for the observed Arch neuronal silencing activity, the locomotion of ncEx3031[myo-3p::Arch::gfp] animals expressing Arch in body wall muscle cells was profoundly affected by green light illumination. With ncEx3031[myo-3p::Arch::gfp] animals, however, illumination did not lead to immediate arrest of locomotion; worm locomotion first slowed down, and paralysis gradually occurred. At an illumination intensity of 14.7 mW/mm², complete arrest of locomotion required illumination for 8 seconds (n = 9). This is in contrast to worms expressing Arch in neurons, which stopped locomotion immediately after the onset of illumination. Therefore, we evaluated the effect of illumination on the locomotion of ncEx3031[myo-3p::Arch::gfp] animals over 30 seconds of illumination. At 0.7 mW/mm², only a fraction of the animals had stopped locomotion. At 8.3 mW/mm², approximately half of the animals stopped locomotion, whereas all animals appeared to be paralyzed when subjected to 14.7 mW/mm² illumination (Fig. 2).

When continuously exposed to green light at an intensity of 14.7 mW/mm² for a period of 30 seconds, some ncEx3031[myo-3p::Arch::gfp] animals remained paralyzed during illumination (48% n = 55), but others recovered partially and started to move while still under illumination, albeit at a much slower rate than usual. With an illumination period longer than 30 seconds, or when subjected to intermittent illumination cycles consisting of 1 second of illumination followed by 1 second of non-illumination, repeated 30 times, locomotion recovery required more time. Some worms failed to resume normal fast locomotion even after a 10-second period of non-illumination. Given a sufficient time following light cessation, however, all worms eventually resumed normal locomotion.

Light-mediated Silencing of Subsets of Ventrall Cord Motor Neurons

In order to utilize Arch for analyzing the function of neuron-circuits of worms, we then tried to express Arch in subsets of neurons. Freely moving C. elegans worms crawl forward on agar in a sinusoidal motion, which was occasionally interrupted by brief
backward movements. Worms can also be directed to move backward or forward by tactile stimulation. We have examined how locomotory behaviors are affected by silencing different subsets of motor neurons in the ventral nerve cord. The locomotion is executed by body wall muscles whose contraction and relaxation is regulated by the cholinergic A- and B-type motor neurons, and the GABAergic D-type motor neurons, respectively [13,14,15]. Previous studies also showed that A-type motor neurons (VA, DA) direct backward locomotion while B-type motor neurons (VB, DB) direct forward locomotion [13,16].

First, to achieve silencing of D-type motor neurons (VD, DD) selectively, Arch::GFP was expressed under the control of the promoter of the unc-47 gene encoding a vesicular GABA transporter [17] (Fig. S1A). Without illumination, *ncEx2351[unc-47p::Arch::gfp] animals moved in a normal sinusoidal motion. When *ncEx2351 animals moving freely were illuminated with green light, they exhibited loopy sinusoidal forward movement (Fig. 3A), and movement of some animals appeared to slow down in either direction. In wild type animals, gentle touch to the anterior part of the body elicits backward movement, whereas posterior touch elicits forward movement. Under green light illumination, the extent of backward movement elicited by gentle anterior touch was smaller in *ncEx2351 worms (Fig. 3C). Likewise, forward movement elicited by gentle posterior touch was affected to various extents. We observed that the onset of touch elicited movement was delayed about 1 second in 2 out of 30 animals.
Arch::GFP in VA and VB motor neurons under the control of 
and it often stopped halfway. Posterior gentle touch to 
the anterior part of the body connected to the head moved.

minutes. The posterior half of the body was often motionless, while 
 locomotion was arrested under continuous illumination over 20
acr-5
shorter distance and their movement was loopy.

cases animals responded to anterior gentle touch, they retreated a

illumination to 
Green light illumination retarded or arrested the locomotion, 
in an abnormal loopy foraging behavior [14,18]. Thus, green light 
elicited backward movement, or did it only incompletely (Fig. 3C). In

animals was normal, but backward movement was abnormal:

animals often moved

ncEx2365[del-1p::Arch::gfp] (diamonds) exhibited higher responsiveness at lower light intensities than nc3031Ex[myo-
3p::Arch::gfp] (triangles) animals did. (mean±SEM; n = 3. Five animals were examined for each trial). doi:10.1371/journal.pone.0035370.g002

suggesting that the GABAergic system might play a role in 
switching promptly the direction of movement. We also examined the 
effect of silencing of GABAergic neurons in the background of 
Roller phenotype in ncEx2311[unc-47p::Arch::gfp; rol-6d] animals. Green light illumination retarded or arrested the locomotion, 
which lasted over 60 seconds of illumination (Fig. S2).

Slow locomotion and defective responses to gentle touch are the 
phenotypes observed in mutants with defects in GABA neurons 
(Fig. 3B, C). A previous study also showed that killing the four 
GABAergic RME neurons in the head of wild type worms resulted 
in an abnormal loopy foraging behavior [14,18]. Thus, green light 
illumination to ncEx2351 animals appears to lead to phenotypes 
expected from silencing GABAergic neurons.

Next, to achieve silencing of A-type motor neurons specifically, 
Arch::GFP was expressed under the control of the promoter of the 
unc-4 gene [19,20] (Fig. S1B). When illuminated with green light, forward movement of freely moving ncEx3068[unc-4p::Arch::gfp] animals was normal, but backward movement was abnormal: 
Whereas normal worms move backward a distance of about one 
body length, ncEx3068[unc-4p::Arch::gfp] animals often moved 
backward only as far as a half the body length, and sometimes 
did not move at all (Fig. 3B). Under green light illumination, anterior gentle touch to ncEx3068[unc-4p::Arch::gfp] worms did 
not elicit backward movement, or did it only incompletely (Fig. 3C). In 
contrast, the posterior touch elicited normal forward movement. 
Sometimes worms stopped moving for a while after a harsh touch.

To achieve silencing of B-type motor neurons specifically, the 
promoter of the acr-5 gene was used [20] (Fig. S1C). When illuminated with green light, forward movement of ncEx2371[acr-
5p::Arch::gfp] animals slowed down, and about half of the animals 
stopped locomotion completely within 10 seconds (Fig. 3B). The 
locomotion was arrested under continuous illumination over 20 
minutes. The posterior half of the body was often motionless, while 
the anterior part of the body connected to the head moved. 
Backward movement was also affected, though to a lesser extent, 
and it often stopped halfway. Posterior gentle touch to ncEx2371[acr-
5p::Arch::gfp] animals under green light illumination failed to elicit 
forward movement almost completely (Fig. 3C). Although in most 
cases animals responded to anterior gentle touch, they retreated a 
shorter distance and their movement was loopy.

When ncEx2365[del-1p::Arch::gfp] animals, which expressed 
Arch::GFP in VA and VB motor neurons under the control of the 
del-1 gene promoter [20,21] (Fig. S1D), were illuminated with 
green light, locomotion appeared to slow down in either direction 
(Fig. 3B). Response to gentle touch was sometimes affected in either 
direction (Fig. 3C). Backward movement was often incomplete.

Arch::GFP Expressed by Heat-shock Promoter

Next, to investigate the timing of the acquisition of the silencing 
function and the stability of Arch following its expression in C. elegans 
cells, we exploited a gene induction system using the 
promoter of the heat-shock protein gene hsp-16.2 [22].

In ncEx3003[hsp-16.2p::Arch::gfp] animals, GFP expression was 
detected 3 hours after heat-shock. The level of GFP expression 
then increased gradually, reached a stable plateau 9–12 hours after 
heat shock, remained constant for as long as 24 hours and started 
to decay slowly thereafter. The level of GFP expression in most 
animals 48 hours after heat shock was lower compared with the 
level 24 hours after heat shock, but higher compared with the level 
3 hours after heat shock. In some animals, GFP expression was 
detected even 72 hours after heat shock.

The GFP signal was observed in the entire region of the body 
(Fig. 1G). In neurons, the GFP signal was localized to the cell 
membrane and in the cytoplasm, but not in the nucleus (Fig. 1H). 
In muscle cells, the GFP signal was high at the cell surface (Fig. 1I).

An intense GFP signal was sometimes detected in granular or 
vesicular structures in the cytoplasm. To help clarify the 
localization of Arch in cells, we compared Arch expression with 
the GFP signals in ncEx17[hsp-16.2p::gfp] transgenic worms, which 
only express GFP under the control of the same heat-shock 
promoter [23]. The GFP signal in ncEx17[hsp-16.2p::gfp] worms 
was distributed diffusely in the cytoplasm, without visualizing 
granular or vesicular structures or demarcating cell boundaries. 
These differences further support the conclusion that Arch::GFP is 
localized to intracellular vesicles and the cell membrane.

Next, we investigated the function of Arch expressed by heat 
shock (Fig. 4A(i), B). Three hours after heat shock, green light 
illumination at 19.6 mW/mm² had no effect on the locomotion of heat-shock-induced ncEx3003[hsp-16.2p::Arch::GFP] worms, 
although a GFP signal was clearly observed. Six hours after heat 
shock, 5% of the animals stopped locomotion in response to green 
light illumination. In response to green light illumination, half of 
the animals stopped locomotion 9 hours after heat shock, and 90% 
of the animals stopped 12 hours after heat-shock. Up to 48 hours 
after heat shock, green light illumination still caused locomotion 
arrest in most animals. Some animals stopped locomotion in 
response to green light illumination even 72 hours after heat shock, 
although the fraction of paralyzed worms (20%, n = 10) was much 
smaller than the fraction 48 hours after heat shock.

To summarize the temporal changes of the Arch silencing 
function and expression, the worms became fully responsive to 
green light illumination 12 hours after heat shock, while Arch 
expression monitored with GFP reached the maximal level 9–12 
hours after heat shock. Thereafter, the responsiveness of worms 
to green light illumination appeared to correlate with the expression 
level of Arch::GFP, which sometimes persisted for longer than 48 
hours. Since heat-shock-mediated expression of GFP is induced 
only temporarily, the long-lasting perdurance of GFP expression, 
as well as the responsiveness to light observed in some worms, 
indicate that Arch can be stably retained in a functional form for a 
long period.

We also investigated the light sensitivity of Ex3003[hsp-
16.2p::Arch::GFP] worms 24 hours after heat shock. A small 
fraction of animals stopped locomotion when illuminated at 
2.2 mW/mm², half of the animals stopped at 4.9 mW/mm², and 
all the animals stopped when illuminated at 11.1 mW/mm².
Uptake and Stability of ATR in Arch

The heat-shock induction experiments described above suggested that Arch acquire silencing function fully only 12 hours after heat shock induction. The unexpectedly long period that is needed before Arch acquires silencing function prompted us to examine the possibility that Arch may need to undergo maturation processes that might require several hours to complete. To investigate whether ATR, an essential cofactor of Arch, might be involved in the posited maturation processes, we examined the incorporation and stability of ATR in Arch.

We examined whether the time required for incorporation of ATR into Arch caused the observed delay of several hours before Arch becomes functional following its expression (Fig. 4). ncEx3003[ksp-16.2p::Arch::gfp] worms raised in the absence of ATR were heat shocked and were transferred to a plate containing ATR 24 hours later (Fig. 4A(ii)). The worms raised without ATR expressed GFP similarly to those raised with ATR. About half of the animals stopped locomotion in response to green light illumination 1.5 hours after transfer, and more than 80% of the animals stopped locomotion under green light 3 hours after transfer (Fig. 4B). Therefore, ATR seems to be incorporated into the preexisting Arch that was synthesized in the absence of ATR, and function there. The results also indicate that Arch is stable in the absence of ATR. Importantly, the relatively short 1.5-hour period that worms required before becoming responsive to green light illumination following transfer to ATR-containing plates indicates that incorporation of ATR into Arch occurs rather rapidly, and this alone probably cannot explain the observed delay of several hours.

We also examined how long ATR keeps functioning after being incorporated into Arch. First, we estimated how long the ATR
that worms have ingested with food remains available for Arch when worms are transferred to plates without ATR. \( ncEx3003[hsp-16.2p::Arch::gfp] \) worms raised in the presence of ATR were heat shocked and transferred to a plate without ATR at various time points. When transferred immediately or 3 hours after heat shock, worm locomotion was not interrupted in response to green light for at least 48 hours after transfer. Given that the expression level of Arch 9 hours after heat shock is sufficiently high for half the worms to show a response to illumination when ATR is present, this result indicates that the level of ATR in the body of worms after transfer to an ATR-free plate seems to be reduced to a level insufficient to be incorporated into Arch within 6 hours. We then examined the stability of incorporated ATR. \( ncEx3003[hsp-16.2p::Arch::gfp] \) worms raised in the presence of ATR were heat shocked and transferred to a plate without ATR 24 hours later (Fig. 4A(iii)). When exposed to green light illumination, the worms stopped locomotion even 24 hours after transfer (93%, \( n = 15 \)), when free ATR should be no longer available for Arch in the body of the worms. Therefore, ATR incorporated into Arch appears to be stable and keep functioning for at least 24 hours.

Effects of GFP Segment in Arch::GFP

Another possible rate-limiting step in the process whereby Arch acquires a silencing function is translocation to the plasma membrane. Although we did not notice any obvious change in the localization pattern of Arch::GFP around 9 hours after heat shock, a slight change in localization, such as a shift to the plasma membrane from nearby cytoplasm, might nevertheless lead to significant functional differences. To investigate whether removing the GFP segment, whose hydrophilic nature can potentially affect incorporation into membranes, from Arch::GFP might facilitate translocation of Arch to the plasma membrane, we generated transgenes expressing Arch alone via a heat-shock promoter, \( Ex[hsp-16.2p::Arch] \).

The dependence of responsiveness to light intensity was almost identical for \( Ex[hsp-16.2p::Arch] \) and \( Ex[hsp-16.2p::Arch::gfp] \) animals, with slightly higher responsiveness for the former at lower light

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**Figure 4. Locomotion assay using heat shock-mediated induction of Arch.** (A) Scheme showing the schedule of transferring worms to plates with or without all-trans-retinal (ATR) in heat shock induction experiments. i: Animals were cultivated in the presence of ATR throughout the experiments. (Figs. 3B, 4B) ii: Animals were transferred to ATR-supplemented plates 24 h after heat shock (Fig. 4B). iii: Animals were cultivated in the presence of ATR and transferred to ATR-free plates 24 h after heat shock. (B) Time course of light-elicited locomotory paralysis after heat-shock induction of Arch::GFP. Heat-shocked \( ncEx3003[hsp-16.2p::Arch::gfp] \) animals cultivated in the presence of ATR throughout the experiment (circles) (Fig. 4A(ii)) or transferred from ATR-free to ATR-supplemented plates 24 h after heat shock (triangles) (Fig. 4A(iii)) were examined at each time point. When ATR was present throughout the experiment (circles), paralysis of worms was first noticed 6 h after heat shock. The paralysis rate reached a plateau 12 h after heat shock, and remained constant for 48 h. (mean ± SEM; \( n = 3 \), Five animals were examined for each trial.) Worms cultivated in the absence of ATR throughout the experiment did not respond to illumination at any time point. When animals were grown and heat shocked in the absence of ATR and then transferred to ATR-supplemented plates 24 h later (triangles), half of them were paralyzed by illumination 1.5 h after transfer, and the paralysis rate reached a plateau within 3 h. (mean ± SEM; \( n = 4 \), Five animals were examined for each trial.).

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intensities (Fig. 5A) (p<0.05 Student’s t-test). With Ex[hsp16-2p::Arch], 2% of the animals became paralyzed at an illumination intensity of 0.5 mW/mm², half of the animals at 3.0 mW/mm², and all the animals at 16.1 mW/mm². With Ex[hsp16-2p::Arch::gfp], 2% of the animals became paralyzed at 0.5 mW/mm², half of the animals at 1.8 mW/mm², and all the animals at 16.1 mW/mm².

The responsiveness of Ex[hsp16.2p::Arch::gfp] animals to light also changed over time in a manner similar to that of Ex[hsp16-2p::Arch::gfp] animals (Fig. 5B). With Ex[hsp16-2p::Arch::gfp] animals, paralysis by green light illumination was first noticed 6 hours after heat shock and the percentage of paralyzed animals increased thereafter. The worms exhibited maximum responsiveness around 12 hours after heat shock, with 95% of the animals stopping locomotion. Thus, deletion of GFP from Arch::GFP does not appear to accelerate the acquisition of the silencing function. The responsiveness to light after 12 hours decreased slightly faster in Ex[hsp16-2p::Arch::gfp] animals (p<0.01 Student’s t-test; 40% of Ex[hsp16-2p::Arch::gfp] animals were paralyzed by green light illumination 48 hours after heat shock, while more than 60% of Ex[hsp16-2p::Arch::gfp] animals were paralyzed. It appears that the GFP segment might inhibit the degradation of Arch.

**Discussion**

In this paper we report for the first time that Arch derived from archaeabacteria can be expressed in *C. elegans* cells without apparent adverse effects, and can be used to silence activity of neurons and muscles, thereby modifying worm’s behaviors.

Arch expressed in neurons and muscles affects worm behavior differently.

We found that illumination at a similar level of light intensity affected worms differently depending on whether Arch was expressed in neurons or muscles. An apparently stronger GFP signal in neurons compared with muscle cells in our transgenic lines suggests that the concentration of Arch::GFP is higher in neurons. Since the amplitude of change in membrane potential driven by Arch-mediated proton current is probably proportional to the density of Arch on the cell membrane, illumination with the identical light intensity would promote larger hyperpolarizing effects in neurons than in muscle.

Neighboring muscle cells are known to be coupled electrically through gap junctions [24]. With their larger cell size, mosaic expression of Arch and electric coupling of cells in muscle tissues would weaken the polarizing effect of the Arch-mediated proton current. Ex[myo-3p::Arch::GFP] transgenic lines so far established express Arch::GFP in highly mosaic patterns, and we have noticed the presence of dead embryos expressing GFP very intensely in line cultures. Although we have not noticed any morphological damage in Arch-expressing muscle cells, highly abundant expression of Arch or the presence of excessive amounts of the promoter sequence of the *myo-3* gene in muscle cells might have a harmful effect on the development or survival of worms.

The heat-shock-mediated Arch expression in *C. elegans* cells with Arch::GFP occurs ubiquitously. While Arch expressed both in neurons and in muscle cells may affect locomotion,
the finding that Ex[hsp-16.2p::Arch::GFP] worms exhibited higher sensitivity to green light illumination than Ex[myo-3::Arch::GFP] worms suggest that locomotory paralysis in Ex[hsp-16.2p::Arch::GFP] worms is mainly attributed to the silencing of neurons rather than muscle cells.

Arch-expressing C. elegans Responds to Light Precisely

By employing various illumination patterns, we showed that worms expressing Arch respond to light quickly and precisely. This finding is consistent with a previous report showing that Arch-mediated neuronal silencing occurs within a millisecond time range [11]. We also showed that continuous illumination for as long as 30 seconds leads to sustained paralysis in most worms expressing Arch in neurons, which is also consistent with a previous finding that prolonged activation does not significantly affect Arch activity.

Since Arch is an outward proton pump [11], prolonged activation of Arch may cause crucial changes in intracellular and extracellular pH levels. The failure of Arch-expressing worms to immediately recovery locomotion after the cessation of illumination following prolonged illumination might be due to disturbances of cellular pH conditions. Nevertheless, the absence of long-lasting effects on neurons or muscle cells is notable, since worms eventually resumed locomotion at normal rates in the absence of illumination.

Silencing of different subsets of neurons elicited distinct behavioral defects.

To investigate the neuron-circuits governing the locomotory behavior of worms, we have used several promoters that drive expression of Arch in selected subsets of motor neurons of the ventral nerve cord. We found that worms expressing Arch by different promoters exhibited distinct behavioral phenotypes specific for each promoter when illuminated with green light.

Responses of worms expressing Arch in GABAergic neurons including D-type motor neurons generally mimic the phenotype of genetic mutants defective in functioning of GABAergic neurons, supporting the notion that specific silencing of GABAergic neurons is achieved successfully. We have, however, observed several differences in the effects between acute silencing and chronic inactivation in mutants. Under a freely moving condition, animals expressing Arch in GABAergic neurons moved more actively than unc-47 mutants. In unc-47 mutants, movement of the head and tail are usually repressed, while green light-mediated silencing of GABAergic neurons elicited enhanced head movement, which resulted in loopy forward movement. We also noticed that light-mediated silencing of GABAergic neurons affected touch responses more strongly than a mutation in the unc-47 gene. The differences may reflect in part incomplete silencing of the GABAergic system due to the mosaic expression of Arch, which is characteristic for transgenes carried by an extrachromosomal array. It may be also due to the properties of the particular mutant allele used in this study. Alternatively, although the finding that acute silencing and chronic inactivation generally has similar effects suggests the absence of developmental mechanisms acting to compensate for locomotory defects of the mutants, neuron-circuits relevant to certain behaviors might be modified through chronic inactivation of GABAergic neurons.

We found that silencing subsets of cholinergic motor neurons had neuron subset-specific effects. At hatching, only three types of muscle cells are present, DA, DB and DD. A classic study using the laser ablation method showed that killing most of DA in newly hatched larvae resulted in animals that could not move backward but could move forward normally, whereas killing most of DB neurons resulted in defective forward movement without affecting backward movement [16]. Light-mediated activation of Arch expressed in DA and VA motor neurons by using the unc-4 promoter caused the direction-specific defect in locomotion, confirming the essential role of A-type neurons in backward movement. The results of the Arch expression experiment using the arc-5 promoter, which drives gene expression in DB and VB motor neurons, but not in A-type motor neurons, also confirmed the essential role of B-type motor neurons in forward movement. Interestingly, we have found that backward movement was often performed incompletely by silencing B-type neurons. While B-type neurons may not play a significant role in triggering backward movement, they might be critically required during the later phase of backward movement. By silencing both VA and VB neurons in Ex[hsp-16.2p::Arch::GFP] animals, we found that movement in either direction was affected. Accordingly, a previous study showed that unc-4(gf)-mediated genetic cell ablation under the control of the del-1 promoter resulted in uncoordinated movement, in which either the forward or backward was irregular and appeared to be executed with difficulty [21].

To summarize, behavioral phenotypes elicited by silencing each subset of motor neurons are generally consistent with known functions of the corresponding neuron subset. It should be noted that we have used transgenes carried by extrachromosomal arrays, which lead to mosaic expression of Arch among target neurons, indicating that complete silencing of a targeted neuron subset would result in a stronger and clearer phenotype. It is also noted that the behavioral changes were observed throughout the period of continuous illumination, which was sometimes longer than 20 minutes, further confirming the utility of Arch as an optogenetic tool in C. elegans.

Time Lag Between Induction and Functioning of Arch

The induction of Arch by heat shock seems to occur only transiently. It was reported that the level of hsp16-2 mRNA reached a maximum 1 hour after onset of heat shock, and decreased to the normal level within 2 hours after heat shock was terminated [22]. We found that the expression level of Arch::GFP did not increase markedly later than 9 hours after heat shock, indicating that synthesis of Arch persisted only for a limited period. This transient induction system enabled us to reveal hitherto unnoticed properties of Arch in vivo.

We found that Arch remains functional for a relatively long time in C. elegans cells, sometimes for several days, which is one advantage that Arch has as an experimental tool. An unexpected finding is that Arch becomes fully functional only 12 hours after its induction. We also found that the expression of Arch::GFP first detected 3 hours after heat shock and the expression level increased up to 9 hours after heat shock. This relatively slow onset and gradual increase in Arch::GFP expression is likely to account for a major part of the observed delay in acquiring a silencing function after Arch induction. It is, however, possible that other processes, such as translocation to the cell membrane and molecular modification, may also partly contribute to the delay.

To affect membrane potential effectively, Arch must be recruited to the cell membrane. We found that Arch::GFP is localized to the cell membrane, but a very intense GFP signal also occurs in intracellular vesicular structures. Translocation of Arch from such vesicles to the plasma membrane might be involved in creating the delay in acquiring the silencing function. Although the GFP segment, by promoting cytoplasmic localization, might affect incorporation of tagged-proteins into the plasma membrane, we found that the removal of GFP from Arch::GFP did not drastically alter either the time course or the dependence on light intensity concerning the responsiveness of Arch-induced worms.
Another possibility that would account for the delay is that newly synthesized Arch might undergo a conformation change, or post-translational modifications before becoming effective. Our finding that ATR can be incorporated into preexisting Arch within 1.5 hours argues against the idea that the delay represents a period needed for incorporation of ATR. A previous in vitro study showing that ATR does not affect the rate-limiting step of bacteriorhodopsin folding [25] also supports this notion. We have also shown that Arch, once having incorporated ATR, remains functional in the absence of ambient ATR. This is consistent with the fact that retinoids keep binding to bacteriorhodopsin after light-driven isomerization [26].

Comparison with NpHR

NpHR, an inward chloride pump, is currently the most widely used optogenetic tool for silencing neurons. A previous study showed that NpHR can silence neurons at a lower light intensity than NpHR [11]. In a previous study, the green light with an intensity of 4.4 mW/mm² or 10.2 mW/mm² were used for silencing of C. elegans [5]. To make Arch a useful alternative to NpHR in C. elegans dependence of silencing effects of these two pumps directly, Arch is considered as a potential tool for optogenetic studies of C. elegans.

Materials and Methods

C. elegans Culture Conditions

Worm strains expressing Arch were grown on Nematode Growth Medium (NGM) plates [27] seeded with solution of Escherichia coli OP50 and 500 μM all-trans-retinal (ATR) (Sigma-Aldrich, USA). Animals were maintained at 20°C in the dark unless indicated otherwise. C. elegans wild-type strain N2 and CB307 unc-47(e307) III were used.

Plasmid Construction

Arch::gfp cDNA was amplified from AAV-FLEX-Arch-GFP [11] (Addgene, Cambridge, MA, USA) by PCR using primers 5′ TTGGTACCTTACTTGTACAGCTCGTCCA3′ and 5′ TTATGCTACATTATGAGAATTCACTGCG3′ as primers, and was cloned into pBluescriptII (pBS, Stratégene, USA) with restriction enzymes KpnI/Sall. To generate his-16.2p::Arch::gfp (pOKA020), Arch::gfp cDNA cut out from pBS cDNA was cloned into the KpnI/Sall site of pPD49.78 containing the his-16.2 promoter [28].

The other plasmids were made using the TOPO cloning kit and the Gateway system (Invitrogen, USA). To generate the entry clone pENTR/D carrying Arch::gfp (pOKA002), Arch::gfp plasmid was PCR-amplified by using 5′GGTACCTTACTTGTACAGCTCGTCCA3′ and 5′TTGGTACCTTACTTGTACAGCTCGTCCA3′ as primers, and was cloned into pBluescriptII (pBS, Stratégene, USA) with restriction enzymes KpnI/Sall. To generate his-16.2p::Arch::gfp (pOKA020), Arch::gfp cDNA cut out from pBS cDNA was cloned into the KpnI/Sall site of pPD49.78 containing the his-16.2 promoter [28].

To TOPO reaction, pOKA003 was recombined into pDEST-hsp-16.2p [29] by LR reaction (gift from Hidetoh Kuroyanagi). Destination vectors containing neuron-subset-specific promoters were constructed by inserting the PCR-amplified genomic fragments into the SphI site of pDEST-PL (gift from Hidetoh Kuroyanagi) using the following primers:

- for pDEST-del-1p, 5′AAGCATGCGTGTGTAATGAGAAATGGACC-GCTGTCG3′ and 5′AAGCATGCGTCTGGGTAAAACCTATCATACGAAACTGC3′;
- for pDEST-unc-4p, 5′AAGCATGCGTCTGGGTAAAACCTATCATACGAAACTGC3′ and 5′AAGCATGCGTCTGGGTAAAACCTATCATACGAAACTGC3′. To generate del-1p::Arch::gfp, unc-47p::Arch::gfp, acr-5p::Arch::gfp, and unc-4p::Arch::gfp, pOKA020 was recombined into pDEST-del-1p, pDEST-unc-4p, pDEST-acr-5p, and pDEST-unc-4p (gift from Hidetoh Kuroyanagi) by LR reaction respectively.

Transgenic Strains

Transgenic animals were generated by microinjection of DNA into the distal arms of gonads of N2 hermaphrodites as previously described [30]. pOKA049 (myo-3p::Arch::gfp, 70 ng/μl) and pRF4 (rol-6d, 400 ng/μl), which is a transgenic marker conferring the Roller phenotype, were injected into N2 worms to create the line nEx3031[pOKA049 (myo-3p::Arch::gfp), pRF4 (rol-6d)]; pOKA052 (pF25B3.3p::Arch::gfp, 170 ng/μl) and pRF4 (600 ng/μl) to create nEx3034[pOKA052(F25B3.3p::Arch::gfp), pRF4 (rol-6d)]; pOKA048 (unc-4p::Arch::gfp, 160 ng/μl) and pRF4 (600 ng/μl) to create nEx3026[pOKA048 (unc-4p::Arch::gfp), pRF4 (rol-6d)]; pOKA020 (his-16.2p::Arch::gfp, 75 ng/μl) and pRF4 (125 ng/μl) to create nEx3002, nEx3003 and nEx3004[pOKA020 (his-16.2p::Arch::gfp), pRF4 (rol-6d)]; pOKA056 (his-16.2p::Arch, 110.8 ng/μl), pPD49.78 GFP (his-16.2p::Arch, 140.0 ng/μl) and pRF4 (230 ng/μl) to create nEx3019, nEx3040 and nEx3041[pOKA056 (his-16.2p::Arch), pPD49.78 GFP (his-16.2p::Arch::gfp), pRF4 (rol-6d)].

Non-roller transgenic animals were generated by using pCFJ90 (myo-2p::mCherry) which drives expression of mCherry in the pharyngeal muscles, as a transgenic marker. We injected into N2 worms pOKA073 (del-1p::Arch::gfp, 200 ng/μl), pCFJ90 (10 ng/μl) and pBluescript (500 ng/μl) to create nEx3235[pOKA073(del-1p::Arch::gfp), pCFJ90 (myo-2p::mCherry)]; pOKA074 (unc-47p::Arch::gfp, 200 ng/μl), pCFJ90 (10 ng/μl) and pBluescript (500 ng/μl) to create nEx3251[pOKA074(unc-47p::Arch::gfp), pCFJ90 (myo-2p::mCherry)]; pOKA075 (acr-5p::Arch::gfp, 200 ng/μl), pCFJ90 (10 ng/μl) and pBluescript (500 ng/μl) to create nEx3257[pOKA075(acr-5p::Arch::gfp), pCFJ90 (myo-2p::mCherry)]; pOKA068 (unc-4p::Arch::gfp, 200 ng/μl), pCFJ90 (10 ng/μl) and pBluescript (500 ng/μl) to create nEx3063[pOKA068(unc-4p::Arch::gfp), pCFJ90 (myo-2p::mCherry)].
all neurons of an *ncEx3034* (F25B3.3p::Arch::gfp) animal or an *ncEx3026* (nx-3p::Arch::gfp) animal.

**Heat-shock Induction of Arch**

All worms tested were *F*<sub>1* roller progeny of *P*<sub>0 roller adults picked onto NGM plates with or without ATR three days before experiments. Late L4 larvae were transferred to NGM plates with or without ATR. The culture plates containing worms were placed in a bacteria culture incubator at 37°C for 30 minutes, and then were returned to the worm incubator at 20°C. The room was maintained at 25°C.

**Microscopic Observation of GFP-expressing Worms**

For observation of GFP fluorescence, a fluorescence stereo microscope SZX12 (Olympus, Tokyo, Japan) with a 100W HBO mercury lamp (Osram, Munich, Germany), and a fluorescence microscope Axiosplan 2 (Zeiss, Oberkochen, Germany) with filter set #10 for GFP with a 100W HBO mercury lamp (Osram, Munich, Germany) were used. Images were acquired with a digital CCD camera Cool SNAP HQ² (Photometrics, Tucson, AZ, USA) using PM Capture Pro software (Nippon Roper, Japan), and processed using ImageJ public domain software.

**Locomotion Assay Under Illumination**

For locomotion assays to examine dependence on light intensity, 5 animals were transferred to a new 30mm NGM plate supplemented with ATR. Worms exhibiting the Roller phenotype or expressing mCherry- in the pharynx were chosen as transgenic animals. With strains in which the Arch::GFP expression was driven by tissue-specific promoters, animals with intense GFP signals were further selected for the assay. Animals were observed under a fluorescence stereo microscope (SZX12), and were illuminated with a 100W HBO mercury lamp through an excitation filter for Red Fluorescent Protein DsRed (545±10 nm, Olympus, Tokyo, Japan). Images of moving animals were recorded using a digital video camera G2-HM570B (JVC, Yokohama, Japan) mounted on the microscope. In most experiments, worms were used only once for a single series of illumination experiments with varying light intensities (from 0.1 to 17 mW/mm²). and were then discarded unless otherwise indicated.

For locomotion assays of animals expressing Arch::GFP in motor neuron subsets, animals were observed under a fluorescence stereo microscope (MVX10), and were illuminated with a 100W HBO mercury lamp through the filter set U-MRFPHQ (Olympus, Tokyo, Japan) with the light intensity of 14.7 mW/mm² at 550 nm. Locomotory defects of freely moving animals were scored through observation of each animal for 5 min. At least 30 animals were examined for each transgenic line. The response to gentle touch was assayed as described by Hart [31]. Briefly, for examining worm’s responses to gentle touch, animals were touched by stroking a hair across the worm’s body at different positions: just behind the pharynx for anterior touch, and just anterior to the anus for posterior touch. For examining the response to harsh touch, animals were prodded with a platinum wire. At least 30 animals were examined for each transgenic line, and each animal was examined 5 times.

In time course analyses of silencing activity of *ncEx3003* after heat-shock induction, animals were used only once for each time point. In experiments comparing the time course for heat-shock-induced Arch::GFP in *ncEx3002*, *ncEx3003* and *ncEx3004* animals, and Arch in *ncEx3039*, *ncEx3040* and *ncEx3041* animals, worms were used for a single series of illumination experiments at various time points (3, 6, 9, 12, 24 and 48 hours after heat shock). After observation at each time point, plates were returned to the worm incubator at 20°C, and before the next observation, plates were left for 30 minutes at 25°C.

**Measurement of Light Intensity**

Light intensity at 550 nm, corresponding to the Arch absorption maximum, was measured at the object plane using an optical power meter (#3664, Hioke, Ueda, Japan) with an optical sensor (#9742, Hioke, Ueda, Japan). Various intensities of illumination were implemented by changing the magnification of the objective lens of the microscope. To calculate the intensity of illumination, we measured the radius of the illuminated area, calculated the area, and then divided the total light intensity by the area.

**Supporting Information**

**Figure S1 Expression of Arch::GFP in *C. elegans* driven by various promoters.** (A) A fluorescent micrograph of an *nc2351Ex[u-47p::Arch::gfp]* animal. Arch::GFP is expressed in D-type motor neurons (VD, DD) (arrow). (B) Expression of Arch::GFP in an *nc3060Ex[u-4p::Arch::gfp]* animal. Arch::GFP is expressed in A-type motor neurons (arrow). (C) Expression of Arch::GFP in an *nc2371Ex[u-5p::Arch::gfp]* animal. Arch::GFP is expressed in B-type motor neurons (arrow). (D) Expression of Arch::GFP in an *nc2363Ex[del-1p::Arch::gfp]* animal. Arch::GFP is expressed in VA and VB motor neurons (arrow). The fluorescence of mCherry expressed in the pharynx is also detected. Scale bar = 100 μm. Anterior is toward the right and dorsal is up. (TIF)

**Figure S2 Defects in the locomotory behavior caused by silencing subsets of motor neurons in strains carrying the *rol-6d* gene.** (A-C) Locomotory behavior of worms expressing Arch in motor neurons subsets. Arch::GFP was expressed in D-type (VD, DD) and VA + VB motor neurons in *ncEx2311[unc-47p::Arch::gfp; rol-6d]* and *ncEx2322[del-1p::Arch::gfp; rol-6d]* animals exhibiting the Roller phenotype. To evaluate differences of animal’s behavior statistically, we used Fisher’s exact test for the locomotion assay of freely moving animals (A), and Student’s t-test for the touch response assay (B, C). For all transgenic strains, animals behaviors under green light illumination (ON = open box) and those without illumination (OFF = filled box) differed significantly (p < 0.001). Error bars indicate ±SEM. (A) Forward (F) and backward movement (B) was scored in worms moving freely. Percentage of animals exhibiting the “Class 3 (severe),” “Class2 (mild)” and “Class1 (no)” phenotype in locomotory behaviors when they were illuminated with green light is shown. (B) Forward movement (F) to gentle posterior touch and backward (B) movement to gentle anterior touch were scored. Responses out of five touches are shown. (C) Forward (F) and backward movement (B) to harsh touch was scored, and was shown additively in each bar. Responses out of five touches are shown. (TIF)

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**Author Contributions**

Conceived and designed the experiments: AO YS ST. Performed the experiments: AO. Analyzed the data: AO ST. Contributed reagents/materials/analysis tools: YS ST. Wrote the paper: AO ST.
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