A dominant mutation in a neuronal acetylcholine receptor subunit leads to motor neuron degeneration in Caenorhabditis elegans

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A Dominant Mutation in a Neuronal Acetylcholine Receptor Subunit Leads to Motor Neuron Degeneration in Caenorhabditis elegans

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Inappropriate or excessive activation of ionotropic receptors can have dramatic consequences for neuronal function and, in many instances, leads to cell death. In Caenorhabditis elegans, nicotinic acetylcholine receptor (nAChR) subunits are highly expressed in a neural circuit that controls movement. Here, we show that heteromeric nAChRs containing the acr-2 subunit are diffusely localized in the processes of excitatory motor neurons and act to modulate motor neuron activity. Excessive signaling through these receptors leads to cell-autonomous degeneration of cholinergic motor neurons and paralysis. C. elegans double mutants lacking calreticulin and calnexin—two genes previously implicated in the cellular events leading to necrotic-like cell death (Xu et al. 2001)—are resistant to nAChR-mediated toxicity and possess normal numbers of motor neuron cell bodies. Nonetheless, excess nAChR activation leads to progressive destabilization of the motor neuron processes and, ultimately, paralysis in these animals. Our results provide new evidence that chronic activation of ionotropic receptors can have devastating degenerative effects in neurons and reveal that ion channel-mediated toxicity may have distinct consequences in neuronal cell bodies and processes.

Introduction
Roles for ionotropic receptor-mediated signaling in the nervous system extend far beyond a well characterized participation in cell–cell communication at synapses. Ionotropic receptor activation is one of several key factors that influence cell survival in developing and mature nervous systems. For example, signaling through nicotinic acetylcholine receptors (nAChRs) promotes the elimination of neurons in the developing avian autonomic nervous system (Hruska and Nishi, 2007; Hruska et al. 2009), and inappropriate pharmacological activation of nAChRs in the Caenorhabditis elegans nervous system leads to developmental arrest (Ruaud and Bessereau, 2006). In the mature nervous system, inappropriate or excessive ion channel activation can have dramatic consequences. In mammals, hypoxic events, such as stroke, lead to excitotoxic cell death as a consequence of excess glutamate release and hyperexcitation of ionotropic glutamate receptors (iGluRs) (Sattler and Tymianski, 2001). Likewise, mutations that cause prolonged activation of nAChRs or iGluRs can lead to neurodegeneration and cell death in organisms ranging from nematodes to mammals (Treinin and Chalfie, 1995; Zuo et al. 1997; Heintz and Zoghbi, 2000; Orr-Urtreger et al. 2000; Labarca et al. 2001; Orb et al. 2004; Miwa et al. 2006). Excess ion channel activation is also a contributing factor in neurodegenerative diseases. For example, the selective vulnerability of motor neurons to cell death in amyotrophic lateral sclerosis (ALS) is believed to arise, at least in part, from hyperactivation of calcium-permeable AMPA-type iGluRs (Kwak and Weiss, 2006; Grosskreutz et al. 2010). Interestingly, in various mouse models of motor neuron diseases, including ALS, genetic manipulations that prevent the death of motor neuron cell bodies are not successful in halting disease progression (Sagot et al. 1995; Gould et al. 2006). This result implies that ion channel hyperactivation may contribute to degenerative events that persist even under conditions when neuronal cell death is attenuated. Shared features of ion channel-mediated degeneration across these diverse receptor types and systems suggest that aspects of this process may be widely conserved across organisms (Driscoll and Gerstbrein, 2003). In most cases, however, a cohesive picture of the cellular events that influence the progression toward cell death as a consequence of ion channel hyperactivation remains unclear.

Ionotropic receptor signaling and its contribution to neurodegeneration can be dissected in detail in the compact nervous system of the nematode Caenorhabditis elegans. Here, we provide evidence that the non-α nAChR subunit ACR-2 contributes to a heteromeric receptor that is important for regulating the activity of excitatory motor neurons. A pore modification in ACR-2 leads to loss of motor neurons and paralysis of the animal. Genetic ablation of nAChR subunits that coassemble into a heteromeric receptor complex with ACR-2 suppresses ACR-2(L/S) (where L/S
is leucine-to-serine substitution) toxicity. In addition, *C. elegans* double mutants lacking two genes previously implicated in calcium homeostasis and necrotic cell death (crt-1/calreticulin and cnx-1/calnexin) are resistant to nAChR-mediated toxicity and possess normal numbers of motor neuron cell bodies. Nonetheless, we observe a progressive degeneration of the motor neuron processes that leads to paralysis in these animals. Thus, ion channel hyperactivation has distinct consequences for neuronal cell bodies and processes.

**Materials and Methods**

**Molecular biology**

*nAChR subunit constructs*. The full-length ACR-2::GFP transgene (pDM1232) was generated by introducing the green fluorescent protein (GFP) coding sequence in-frame into the sequence of an acr-2 genomic fragment (−3353 to +7776 bp relative to the translational start site) encoding the intracellular loop between transmembrane domains TM3 and TM4. The full-length ACR-2(L/S) construct (pBB9) was generated by PCR-based site-directed mutagenesis using mutant primers and pDM1232 as the template. The *Pacr-2::ACR-12 cDNA (pHP3)* and *Pacr-4::ACR-12 cDNA (pBB25)* constructs were generated by amplifying the acr-12 cDNA from the expressed sequence tag yk1093d12 (gift from Yuji Kohara, National Institute of Genetics, Mishima, Japan) using sequence-specific primers designed to the start and stop of *acr-12* and subcloning it into the Nhel/Sacl sites of a plasmid containing an −3.3 kb promoter for the *acr-2* gene or into a plasmid containing a 1.3 kb promoter for the *unc-47* gene.

**Transcriptional reporters.** The *Punc-47::mCherry construct (pPRB5)* was generated by subcloning an *Aeg1/AattI* fragment that contained the full-length mCherry coding sequence into a vector containing a 1.3 kb promoter for the *unc-47* gene. The *Pacr-2::GFP (pPRB19)* construct was generated by subcloning an *AattI/BamHI* fragment that contained an 7.7 bp promoter for the *acr-2* gene or into a plasmid containing a 1.3 kb promoter for the *unc-47* gene. The *Pacr-2::ACR-12 cDNA (pHP3)* and *Pacr-4::ACR-12 cDNA (pBB25)* constructs were generated by amplifying the *acr-12* cDNA from the expressed sequence tag yk1093d12 (gift from Yuji Kohara, National Institute of Genetics, Mishima, Japan) using sequence-specific primers designed to the start and stop of *acr-12* and subcloning it into the Nhel/Sacl sites of a plasmid containing an −3.3 kb promoter for the *acr-2* gene or into a plasmid containing a 1.3 kb promoter for the *unc-47* gene.

**C. elegans strains**

*C. elegans* strains were grown on nematode growth medium (NGM) plates with the OP50 strain of *Escherichia coli* at 22°C using standard laboratory procedures. Wild-type animals are the N2 Bristol strain. All transgenic strains were obtained by microinjection of plasmid DNA into the germ line and data presented are from a single representative transgenic line unless noted otherwise. In all cases, lin-15(n7655) mutants were injected with the *lin-15* rescuing plasmid (pL15E; 30 ng/μl) and one or more of the following plasmids (30 ng/μl): pBB9, pBB25, pDM1232, pHP3, pPRB4, pPRB5, pPRB14, pPRB19. Multiple independent extragenic lines were obtained for each transgenic strain. Stably integrated lines were generated as necessary by x-ray integration and outcrossed at least four times to wild type. The transgenic strain expressing the integrated *ACR-2(L/S) transgene (ufIs25)* was outcrossed 10 times to wild type. The following strains were used in this study: RB1559 (acr-2(ok1887)), IZ421 (acr-12(ok3767)), RB2071 (ced-3(ok7374)), VC1801 (cox-1(ok2234)), RB1021 (ctx-1(ok4984)), IB274 unc-29(ok229), CB904 unc-38(e264), CB306 unc-50(c306), VC731 unc-63(ok1075), CB883 unc-74(e883), IZ380 afs-331, IZ814 afs-325, IZ625 afs-25;afs-31, IZ790 afs-49, IZ627 afs-42, LS949 var-148, IZ924 afs-25;var-48, IZ950 ced-3(ok2734); IZ625;var-48, IZ877 ctx-1(ok2234);IZ625;var-48, IZ926 ctx-1(ok4984); IZ625;var-48, IZ976 ctx-1(ok1048);ctx-1(ok2234);IZ625;var-48, IZ971 ctx-1(ok4984);ctx-1(ok2234);IZ625;var-48, IZ927 ctx-1(ok4984);ctx-1(ok2234);IZ625;var-48, IZ412, IZ787 unc-29(c29);I2525, IZ928 arc-3(ok2049);I2525, IZ299 arc-5(ok180);I2525, IZ390 arc-9(ok333);I2525, IZ2931 arc-19(ok676);I2525, IZ2932 arc-23(ko2840);I2525, IZ446 unc-38(e264);I2525, IZ659 unc-50(c306);I2525, IZ291 unc-63(ok1075);I2525, IZ490 unc-68(e540);I2525, IZ604 unc-74(e883);I2525, IZ673 arc-12(ok367);I2525, IZ676 unc-12(ok367);I2525, IZ681 arc-12(ok367);I2525.

**Genetic screen and identification of suppressors of ACR-2(L/S)-induced paralysis**

Paralyzed animals expressing the integrated ACR-2(L/S) array (ufIs25) were mutagenized with 50 μM ethyl methanesulfonate (Brenner, 1974). Young adult F2 progeny of ～20,000 mutagenized animals were washed twice with M9 and transferred to fresh plates. After allowing time for the animals to disperse, moving animals were picked to single plates. Eighty-one candidate suppressors were isolated. A secondary screen showed that fifty-one of these were resistant to the paralyzing effects of levamisole. For genetic complementation tests, males carrying a mutation in candidate levamisole-resistance genes were crossed with hermaphrodites carrying the *acr-2(L/S) transgene* and a suppressor mutation. F1 progeny were scored for paralysis. A cross was performed in parallel using N2 males to identify X-linked suppressor mutations and determine dominance/recessivity. The mapping of *acr-12* alleles was performed in the presence of the *ACR-2(L/S) transgene*. A strain carrying the integrated *ACR-2(L/S) transgene (ufIs25)* on linkage group I was backcrossed 7× to the CB4856 Hawaiian strain. *acr-12* alleles were mapped to a region the right of +8 on the X chromosome using the SNP mapping procedure described as previously (Wicks et al. 2001; Davis et al. 2005).

**Behavioral assays**

All behavioral analysis was performed with young adult animals (24 h post-L4) at room temperature (22°C–24°C); different genotypes were scored in parallel, with the researcher blinded to the genotype.

**Aldicarb and levamisole assays.** Staged populations of adult animals (~10) were transferred to NGM plates containing 1 mM aldicarb (Chem-Serv, and movement was assessed every 15 min for 2 h. Data represent the mean ± SEM of at least four assays. For levamisole assays, staged populations of adult animals were scored for paralysis after 120 min on plates containing 200 μM levamisole.

**Body bend analysis.** Body bends were scored on unseeded NGM agar. Animals were transferred from their culture plate to an unseeded plate and allowed to crawl away from any food that might have been transferred. The animals were then gently transferred without food to another unseeded plate and allowed to recover for 1 min. After the recovery period, the animals were filmed for 5 min using an ImagingSource DMK 21F94 FireWire camera and iMovie software.

**Microscopy**

Confocal microscopy was performed using a Zeiss Axioskop 2 microscope system and LSM Pascal 5 imaging software (Zeiss). Images were processed using ImageJ software (open source). Epifluorescent imaging was performed using a Zeiss Axiosimager M1 microscope and Axiosvision software (Zeiss). Movies and still images for behavioral analyses were obtained using an Olympus SZ61 upright microscope equipped with a FireWire camera (ImagingSource). For the developmental timeline, synchronized populations were obtained by bleaching gravid animals on NGM plates seeded with OP50. The resulting progeny were allowed to mature at room temperature. Animals were imaged at 16, 28, 38, and 48 h after bleaching using wide-field epifluorescent microscopy, and the number of surviving cell bodies were counted manually with the researcher blinded to genotype.

**Results**

**acr-2 encodes a nicotinic receptor subunit expressed in cholinergic motor neurons**

The *C. elegans* genome encodes 29 nAChR subunits that contribute to the formation of distinct classes of homopentameric and heteropentameric receptors (Jones et al. 2007; Rand, 2007). Two classes of nAChRs formed from six of these subunits are expressed in body wall muscle cells and required for neuromuscular signaling (Richmond and Jorgensen, 1999; Francis et al. 2005; Touroultine et al. 2005). Previous studies have suggested that the expression of several nicotinic acetylcholine receptor subunits, including the non-α subunit acr-2, is enriched in motor neurons of the ventral nerve cord (Hallam et al. 2000; Cinar et al. 2005; Jospin et al. 2009). To determine which ventral cord motor neurons express acr-2, we examined transgenic strains that expressed GFP (Chalfie et al. 1994) under the control of *acr-2* regulatory sequences. Expression of *Pacr-2::GFP* was limited to the nervous system.
system and largely restricted to neurons located in the ventral nerve cord. Expression of Pacr-2::GFP did not overlap with expression of a GABAergic mCherry reporter (Punc-47::mCherry), indicating that expression of acr-2 was limited to cholinergic motor neurons (DA, VA, DB, VB) in the ventral nerve cord of adult animals (Fig. 1A). To determine the subcellular localization of ACR-2, we generated transgenic strains that expressed a full-length ACR-2::GFP fusion protein in which GFP was inserted in frame in the intracellular loop region located between transmembrane domains three and four (Fig. S1, available at www.jneurosci.org as supplemental material). Expression of the GFP reporter construct could be observed at all larval stages and in the adult. We noted the onset of expression in late embryogenesis by the threefold stage (~550 min after fertilization). In first larval stage (L1) animals, when DA and DB motor neurons are the sole excitatory motor neurons present, ACR-2 expression was clearly visible in ventral nerve cord processes. DA and DB motor neuron dendrites receive synaptic input in the ventral cord, and these neurons extend commissural axons to the dorsal cord where they form neuromuscular synapses with the dorsal musculature. In adult animals, we found that the fusion protein was diffusely localized to neuronal processes of the dorsal and ventral nerve cords. Our analysis suggests that ACR-2 is diffusely localized with enriched expression in the dendritic compartment of cholinergic motor neurons.

The locomotory control interneurons provide synaptic input to the excitatory motor neurons; however, the role of acetylcholine (ACH) in this signaling remains unclear. To evaluate whether the locomotory control interneurons are cholinergic, we examined transgenic strains that coexpressed GFP under control of regulatory elements for the gene encoding the ACh vesicular transporter (Punc-17::GFP) together with the red fluorescent protein mCherry expressed under the control of the regulatory elements for the nmr-1 gene (Pnmr-1::mCherry) (Figure S2, available at www.jneurosci.org as supplemental material). nmr-1 is expressed in the AVA, AVD, AVE, and PVC locomotory control interneurons, as well as the RIM and AVG neurons (Brockie et al. 2001). We noted no overlap in the pattern of the red and green fluorescent signals with the possible exception of the interneuron AVE, indicating that these two reporters labeled almost completely independent cell populations. These data are consistent with the idea that the locomotory control interneurons are not primarily cholinergic. The enriched expression of ACR-2 in the dendritic compartment of motor neurons may reflect the involvement of ACR-2 receptor complexes at synapses between AVE and cholinergic motor neurons; however, the lack of Punc-17::GFP expression in a majority of locomotory control interneurons and the diffuse distribution of ACR-2 are inconsistent with an exclusive role at synapses.
acr-2 mutants have motor deficits

The cholinergic motor neurons in the ventral nerve cord make synaptic contacts onto the body wall musculature that drives nematode locomotion. To evaluate the contribution of ACR-2 to cholinergic motor neuron excitability and motor output, we obtained a strain carrying a deletion mutation (ok1887) in the acr-2 genomic locus. The acr-2(ok1887) eliminates ~2.8 kb of chromosomal DNA, including the transcriptional start, and is likely to be a null. Animals homozygous for the ok1887 allele are healthy and viable. acr-2(ok1887) mutants are not obviously uncoordinated, though closer inspection revealed a modest decrease in locomotion rate (Fig. 1B). Expression of the full-length ACR-2::GFP fusion protein in acr-2(ok1887) mutants was sufficient to restore normal movement.

The acetylcholinesterase inhibitor aldicarb has proven to be a useful tool for detecting alterations in neurotransmitter release from cholinergic motor neurons. To test whether ACR-2 receptor complexes may be important for regulating activity of the cholinergic motor neurons, we examined whether acr-2 mutant worms exhibit altered sensitivity to the paralyzing effects of aldicarb. acr-2 mutant animals were slightly resistant to paralysis by aldicarb, and this effect was normalized by expression of ACR-2::GFP (Fig. 1C). These data are consistent with the notion that ACR-2 plays a role in modulating the activity of cholinergic motor neurons but suggest that ACR-2 is not absolutely required for motor neuron depolarization.

Introducing a dominant mutation in ACR-2 leads to profound motor deficits

The second transmembrane domains of Cys-loop family ligand-gated ion channel subunits are well known to line the ion channel pore and play a critical role in channel gating. In particular, a highly conserved nonpolar residue (typically leucine) in the M2 region has been shown to have profound effects on receptor activation properties (Fig. S3, available at www.jneurosci.org as supplemental material). Substitution of a polar amino acid (e.g., serine) for the leucine at this position produces a gain-of-function effect, resulting in increased receptor activation and very slow inactivation (Revah et al., 1991; Labarca et al. 1995).

We engineered the homologous leucine-to-serine point mutation into the sequence encoding the M2 9–12 neurons (arrowheads). C, C1, Wide-field epifluorescent images of a transgenic animal expressing Punc-47::mCherry (C) and a transgenic animal coexpressing Punc-47::mCherry with ACR-2(L/S) (C1). The full complement of Punc-47::mCherry-labeled motor neurons remains in transgenic ACR-2(L/S) animals and is indicated. D, Quantification of the total number of motor neurons in wild-type (gray) and ACR-2(L/S) animals (black); *p < 0.01. For all images, animals are positioned with the head on the left side of the image.
of a gain-of-function receptor. Our results suggest that motor output to head muscles is unaffected, while control of body wall musculature is dramatically impaired in these animals.

**ACR-2(L/S)-induced motor neuron degeneration**

Examination of the ventral nerve cord region of transgenic ACR-2(L/S) animals by differential interference contrast (DIC) microscopy showed that a subset of the ventral nerve cord neurons that normally express *acr-2* swelled beyond their normal diameter and eventually disappeared, presumably as a result of cell death (Figs. 3, 4). These results suggest that enhanced cholinergic signaling mediated by receptors incorporating ACR-2(L/S) leads to motor neuron toxicity. To characterize this in more detail, we examined the effects of ACR-2(L/S) expression in strains carrying fluorescent reporters that label populations of cholinergic neurons. We observed only dim P*acr-2*::GFP fluorescence in the ventral nerve cord of ACR-2(L/S) animals, suggesting that many of the neurons labeled by this reporter were lost (data not shown).

To evaluate the specificity of this effect for neurons that expressed *acr-2*, we examined a P*unc-17*::GFP reporter that is expressed in all cholinergic neurons (Fig. 3) (Chase et al. 2004). While ACR-2(L/S) expression did not produce obvious differences in the number of head neurons labeled by P*unc-17*::GFP, we observed a dramatic decrease in the number of ventral nerve cord motor neuron cell bodies; yet, several motor neuron cell bodies remained present. The surviving neurons included the six VC motor neurons that do not normally express *acr-2* and a more variable group of 10–12 additional excitatory motor neuron cell bodies (Figs. 3; Fig. S4, available at www.jneurosci.org as supplemental material). Based on the position and number of cell bodies and commissural processes, the additional surviving neurons included both DA and DB motor neurons that normally express *acr-2*, as well as AS motor neurons that do not. Similar to our observations for ACR-2(L/S)-induced paralysis, the effects of ACR-2(L/S) on motor neurons were dominant. To evaluate the effects of ACR-2(L/S) on GABA motor neurons, we coexpressed ACR-2(L/S) together with a mCherry transcriptional reporter that labeled GABA neurons (P*unc-47*::mCherry) (Fig. 3C,D). We observed that the full complement of GABA neurons was present and morphologically normal. These data suggest that ACR-2(L/S) acts cell autonomously to promote degeneration of motor neurons and that specific neurons are differentially susceptible to the effects of ACR-2(L/S) expression.

**Motor neuron loss occurs soon after onset of ACR-2(L/S) expression**

As noted above, we observed clear ACR-2::GFP fluorescence in late embryogenesis. We found that threefold embryos coexpressing ACR-2(L/S) with the P*acr-2*::GFP transcriptional reporter possessed normal numbers of GFP-positive neurons, suggesting that ACR-2(L/S) toxicity occurred after hatch (Fig. S4, available at www.jneurosci.org as supplemental material). To precisely determine the onset of motor neuron cell death, we imaged transgenic ACR-2(L/S) animals that coexpressed the P*unc-17*::GFP transcriptional reporter at various time points ranging from newly hatched larvae to adults (Fig. 4). We observed significant motor neuron loss in newly hatched larvae. Roughly 40% of the 16 cholinergic motor neurons present in L1 animals were lost within 16 h after hatch. During the transition from the first larval stage to the second larval stage (L2) the number of ventral nerve cord motor neurons increases substantially, with the addition of >50 motor neuron cell bodies (Sulston and Horvitz, 1977). In transgenic ACR-2(L/S) animals we observed only a slight increase in the number of cell bodies labeled by P*unc-17*::GFP over the course of development. Even by the time transgenic ACR-2(L/S) animals had clearly reached adulthood, the number of
Punc-17::GFP labeled motor neuron cell bodies was roughly comparable to that of a wild-type L1 animal. The VC motor neurons do not express acr-2 and are clearly present in transgenic ACR-2(L/S) animals (Fig. 3; Fig. S5, available at www.jneurosci.org as supplemental material). Therefore, the small developmental increase in the number of motor neuron cell bodies that we observe in transgenic ACR-2(L/S) animals likely represents the postembryonic addition of VC neurons. These results suggest that the other classes of motor neurons that are born postembryonically and normally express acr-2 (e.g., VA, VB) are almost completely absent in adult transgenic ACR-2(L/S) animals.

**Mutations that suppress paralysis define the subunits of a neuronal nAChR**

To identify genes required for the toxic effects of transgenic ACR-2(L/S) expression, we conducted a forward genetic screen for suppressors of ACR-2(L/S)-induced paralysis. We screened the F2 progeny of mutagenized hermaphrodites that expressed ACR-2(L/S) and selected animals that exhibited improved movement. A close examination of the mutants isolated from the screen showed that two phenotypic classes were easily distinguishable. One class of animals phenocopied wild-type animals, whereas the other showed a deleterious phenotype. The latter group was quantified over a 5 min period. Data represent the mean ± SEM of at least 10 animals for each genotype.

We found that members of this class were strongly levamisole-resistant, and we isolated alleles of the levamisole resistance genes unc-38, unc-63, unc-74, and unc-50 from these animals (supplemental Table S1, available at www.jneurosci.org as supplemental material). We also crossed available strains carrying single loss-of-function mutations in known levamisole resistance genes with transgenic ACR-2(L/S) transgene. Animals were placed on a food-free agar plate, and the average number of body bends per minute was quantified over a 5 min period. Data represent the mean ± SEM of at least 10 animals for each genotype.

### Table 1. Several nAChR subunits are required for ACR-2(L/S)-induced paralysis

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**Figure 5.** Mutations in nicotinic acetylcholine receptor subunits suppress ACR-2(L/S)-induced paralysis. **A.** Quantification of the average number of body bends per minute for wild-type animals, acr-12(ok367), unc-63(ok1075), unc-38(e264), unc-74(e883), and unc-50(e306) mutants in the absence (gray bars) or presence (black bars) of the ACR-2(L/S) transgene. Animals were placed on a food-free agar plate, and the average number of body bends per minute was quantified over a 5 min period. Data represent the mean ± SEM of at least 10 animals for each genotype. **B.** Schematic of the membrane topology of ACR-12 with approximate location, allele names, and molecular nature of loss-of-function mutations that suppress ACR-2(L/S) toxicity indicated. **C.** Quantification of the average number of body bends per minute for the following genotypes: wild type, acr-12(ok367), transgenic ACR-2(L/S), acr-12 mutants expressing ACR-2(L/S), acr-12 mutants expressing ACR-2(L/S) together with an extrachromosomal array containing Punc-47::ACR-12, and acr-12 mutants expressing ACR-2(L/S) together with an extrachromosomal array containing the Pacr-2::ACR-12 cDNA. Data represent the mean ± SEM, for 5–10 animals. **D–H.** Still images of adult animals on NGM plates without food for the genotypes indicated. “GABA” and “ACh” refer to specific expression of the acr-12::GABA or cholinergic neurons using the unc-47 or acr-2 promoters, respectively.

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<td>—</td>
<td>nAChR α subunit</td>
</tr>
<tr>
<td>acr-23(ok2840)</td>
<td>—</td>
<td>nAChR α subunit</td>
</tr>
<tr>
<td>unc-63(e37)</td>
<td>+</td>
<td>nAChR α subunit</td>
</tr>
<tr>
<td>unc-38(e264)</td>
<td>+</td>
<td>nAChR α subunit</td>
</tr>
<tr>
<td>unc-29(e229)</td>
<td>—</td>
<td>nAChR non-α subunit</td>
</tr>
<tr>
<td>unc-50(e306)</td>
<td>+</td>
<td>Golgi integral membrane protein</td>
</tr>
<tr>
<td>unc-74(e883)</td>
<td>+</td>
<td>Function unknown</td>
</tr>
</tbody>
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muscles and neurons. Our analysis suggests that each of these gene products may contribute in a cooperative fashion to the generation of functional ACR-2(L/S) receptors and subsequent toxicity.

A smaller number of animals isolated from the screen exhibited uncoordinated movement with deep body bends and showed normal sensitivity to the paralyzing effects of levamisole. We determined that the suppressor mutations in this second phenotypic class represented one complementation group and were X linked. Using single-nucleotide polymorphism mapping, we mapped one allele to the right of +8 on the X chromosome. The nAChR subunit gene acr-12 is located in this genomic region. Sequence analysis revealed four nonsense mutations and two missense mutations in the acr-12 coding sequence among our second class of suppressors (Fig. 5B). We also found that a dele-
tion mutation (ok367) in the acr-12 gene suppressed paralysis in transgenic ACR-2(L/S) animals and prevented the loss of motor neuron cell bodies. Expression of a full-length acr-12 rescuing construct in acr-12(ok367);ACR-2(L/S) animals restored paralysis, verifying that acr-12 is required (Fig. 5C,F). In contrast, we found that several other nAChR subunits with restricted expression to the nervous system were not required for ACR-2(L/S)-induced paralysis (Table 1). acr-12 encodes a nicotinic receptor α subunit that is broadly expressed in the nervous system, including many ventral cord motor neurons, but is not expressed in body wall muscles (Cinar et al. 2005; Gottschalk et al. 2005). acr-12(ok367) mutants have grossly normal movement and show normal sensitivity to the paralyzing effects of levamisole. To test whether acr-12 expression solely in cholinergic motor neurons is sufficient for ACR-2(L/S)-induced toxicity, we specifically restored expression of acr-12 to either ACh or GABA motor neurons of acr-12(ok367) mutants that carried the ACR-2(L/S) transgene (Fig. 5G–H). We found that specific expression of the acr-12 cDNA in cholinergic motor neurons of transgenic acr-12(ok367); ACR-2(L/S) mutants led to ACR-2(L/S) toxicity and paralysis. In contrast, specific expression in GABA neurons had no effect. Our results indicate that acr-12 expression in cholinergic motor neurons is specifically required for ACR-2(L/S)-induced cell death. Furthermore, our results suggest that coassembly of ACR-2(L/S) into a heteromeric receptor complex with UNC-38, UNC-63, and ACR-12 is required to produce toxicity.

**ACR-2(L/S)-induced motor neuron loss is suppressed in cnx-1; crt-1 double mutants**

At least two mechanistically distinct types of cell death have been described. Programmed cell death or apoptosis is a form of cell death common in development and tissue homeostasis and occurs by a genetic program that is broadly conserved across metazoans (Danial and Korsmeyer, 2004). Necrosis generally occurs following cellular injury and is often characterized by swelling of the dying cell (Festjens et al. 2006; Golstein and Kroemer, 2007). Our forward genetic screen did not identify genes previously implicated in the execution of either of these forms of cell death. To determine how ACR-2(L/S) expression leads to cell death, we introduced the ACR-2(L/S) transgene into genetic backgrounds deficient for genes essential for either apoptotic cell death or necrotic cell death (Mattson et al. 2000; Rao et al. 2004; Szydlowska and Tymianski, 2010). Calreticulin/CRT-1 and calnexin/CNX-1 are endoplasmic reticulum (ER) resident proteins that serve dual functions as 

Calnexin and calreticulin perform similar cellular functions, we generated a strain carrying loss-of-function mutations in both genes to test whether they may act redundantly in ACR-2(L/S)-induced toxicity. cnx-1; crt-1 double mutants were viable although smaller in size than wild-type animals, had grossly nor-
that the full complement of 16 cholinergic motor neuron cell bodies was present in L1 cnx-1;crt-1;ACR-2(L/S) animals, and we did not detect obvious defects in the connectivity of the cholinergic motor neurons (Fig. 7C). These results indicate that embryonic born motor neurons developed normally and made appropriate synaptic contacts onto their partner muscle cells, suggesting that the combined loss of crt-1 and cnx-1 is strongly neuroprotective against ACR-2(L/S) toxicity in L1 animals. We observed that adult cnx-1;crt-1 double mutants expressing ACR-2(L/S) also possessed normal numbers of cholinergic motor neuron cell bodies, providing additional evidence that the presence of either calnexin or calreticulin is required for the cellular events that lead to ACR-2(L/S)-induced cell deaths (Figs. 6F; G, 7D). However, larvae that had progressed beyond L1 and adult cnx-1;crt-1;ACR-2(L/S) animals were unable to propagate sinusoidal body bends and move effectively. This observation suggested that ACR-2(L/S) expression in cnx-1;crt-1 double mutants led to a progressive disruption of motor function even when motor neuron death was attenuated.

To determine whether altered motor neuron connectivity may underlie the paralysis of adult cnx-1;crt-1;ACR-2(L/S) animals, we made a close examination of the cholinergic motor neuron processes (Fig. 7E–7J). We found that defects in the motor neuron processes of control cnx-1;crt-1 double mutants occurred only rarely (Fig. S6, available at www.jneurosci.org as supplemental material), and these animals exhibited grossly normal movement across all stages of development. In cnx-1;crt-1 double mutants that expressed ACR-2(L/S), we observed defasciculation of the ventral nerve cord neuronal processes (in 96% of animals scored; n = 30), as well as defects in the morphology of commissural axons (in 75% of axons) (Fig. 7). We often observed several classes of defects within individual animals and, in some instances, individual commissural axons contained multiple defects. The defects were of several types. First, we observed abnormal axon branches that often terminated in growth cone-like structures (54% of animals). Second, we observed ectopic sprouting with no clear single axonal process present (71% of animals). Finally, we observed axons with abnormal trajectories and wandering growth (83% of animals). These results suggest that the muscle targets of cholinergic motor neuron processes are not appropriately innervated in adult animals. We observed that the frequency of these defects increased dramatically after the first larval stage (Fig. 7I), suggesting that a progressive deficiency

Figure 7. Progressive destabilization of motor neuron processes in cnx-1;crt-1;ACR-2(L/S) animals. A, B, Frames showing the movement of L1 transgenic ACR-2(L/S) (A) or cnx-1;crt-1;ACR-2(L/S) (B) animals 30 s (t = 0), 60 s (+30), or 90 s (+60) after transfer to an agar plate. The black arrow marks the starting positions of the worms in each frame. The transgenic ACR-2(L/S) animal does not move from where it was placed on the plate. The dashed line shows the movement path of the L1 cnx-1;crt-1;ACR-2(L/S) animal over the course of 60 s. The additional tracks on the plate show the path of the animal in an ~30 s period before imaging began. C, D, Confocal images of a first larval stage (C) or adult cnx-1;crt-1;ACR-2(L/S) animals expressing an integrated Punc-17:GFP (D). Images show Z-projections of 15 confocal planes (0.5 μm/slice) (C) or 16 confocal planes (0.5 μm/slice) (D). Arrows (D) indicate positions of commissures. Dashed box (D) indicates area with multiple neuronal defects. E–G, Confocal images of cnx-1;crt-1;ACR-2(L/S) animals taken at hatch (E), 24 h posthatch (F), and adulthood (G). In each case, a region immediately posterior of the vulva was imaged and the ventral nerve cord is positioned at the top. Arrows indicate commissural processes; arrowheads indicate the ventral nerve cord. Images show Z-projections of 12 confocal planes (E), 23 confocal planes (F), or 33 confocal planes (G) (0.5 μm/slice). Scale bars represent 10 μm. H, Quantification of percentage of animals moving at hatch in ACR-2(L/S) and cnx-1;crt-1;ACR-2(L/S) animals. Data represent the mean number of animals ± SEM making more than two consecutive body bends during a 3 min period. I, Quantification of the percentage of cnx-1;crt-1 and cnx-1;crt-1;ACR-2(L/S) animals with defective neuronal process morphology. Observed defects include defasciculation of the ventral nerve cord, wandering commissural processes, or ectopic branching.
in the stabilization or maintenance of appropriate neuromuscular connectivity underlies the paralysis we observed in mature animals.

To better understand the mechanisms underlying the suppression of ACR-2(L/S)-mediated cell death in cnx-1;crt-1 animals, we directly evaluated the role of intracellular calcium. We found that culturing transgenic ACR-2(L/S) animals in the presence of dantrolene, an inhibitor of ER calcium release, or the calcium chelator EGTA (Fig. 8A,B) led to a modest but significant increase in the number of surviving motor neuron cell bodies in both the cell bodies (F) and ventral nerve cord processes (G) of wild-type and ACR-2(L/S) animals transgenically expressing ACR-2::GFP. Data represent the mean ± SEM, for at least 10 animals. **p < 0.01, compared to ACR-2(L/S) animals.

Figure 8. Perturbation of intracellular calcium and reduced ACR-2::GFP levels in cnx-1;crt-1 double mutants suggest dual mechanisms for cell death suppression. A–C, Quantification of the average number of Punc-17::GFP-labeled cell bodies present in the ventral nerve cord in adult animals treated with dantrolene (A), L1 animals treated with EGTA (B), and adult unc-68(e540) mutant animals (C). Data represent the mean ± SEM, for at least eight animals. **p < 0.01, ***p < 0.001, compared to untreated ACR-2(L/S) animals. D, E, Confocal images of adult wild type (D) or cnx-1;crt-1 (E) animals expressing ACR-2::GFP. Animals are taken immediately posterior to the vulva and show z-projections of seven confocal planes (0.5 μm/slice). F, G, Quantification of fluorescence levels in cell bodies (F) and ventral nerve cord processes (G) of wild-type and ACR-2(L/S) animals transgenically expressing ACR-2::GFP. Data represent the mean ± SEM, for at least 10 animals. **p < 0.01, compared to ACR-2(L/S) animals.

Discussion

Our analysis of ACR-2-containing nicotinic receptors in C. elegans neurons has revealed common features between the function of these receptors in the C. elegans nervous system and roles for heteromeric nAChRs in the vertebrate brain (Dani and Bertrand, 2007). First, heteromeric nAChRs in the mammalian brain are not primarily concentrated at postsynaptic sites; instead, they are more variably localized to presynaptic, preterminal, and nonsynaptic sites. Similarly, we find that ACR-2-containing nAChRs appear diffusely localized to the processes of excitatory motor neurons, suggesting that these receptors may function at extrasynaptic sites. Second, heteromeric brain nAChRs primarily function to modulate neurotransmitter release and neuronal excitability. Our studies of acr-2 loss-of-function mutants indicate that heteromeric nAChRs containing ACR-2 modulate the excitability of cholinergic motor neurons but are not absolutely required for motor neuron depolarization or ACh release at neuromuscular synapses. Third, mouse studies have shown that knock-in expression of a heteromeric brain nAChR subunit bearing a L/S pore modification homologous to the one we describe here causes dramatic neuron loss and perinatal lethality (Labarca et al. 2001). Likewise, transgenic expression of ACR-2(L/S) leads to cell-autonomous neurodegeneration.

Importantly, our transgenic approach also enabled us to identify genes required for ACR-2(L/S) toxicity. Mutations that suppressed both the paralysis and neurodegeneration caused by ACR-2(L/S) expression defined the constituent subunits of a putative multimeric ACR-2 receptor complex as well as genes required for receptor trafficking and assembly. Additionally, a single gene mutation in crt-1, previously shown to suppress other forms of ion channel-mediated cell death in C. elegans, partially suppressed ACR-2(L/S) toxicity. We found that the loss of motor neurons caused by ACR-2(L/S) expression was completely suppressed in adult cnx-1;crt-1 double mutants; yet, these animals remained paralyzed. Interestingly, suppression of ACR-2(L/S)-induced cell death uncovered a secondary consequence of ACR-2(L/S) expression: the accumulation of morphological defects in the processes of surviving motor neurons. These axonal defects resemble outgrowth errors typically associated with secondary regrowth of axons (Knobel et al. 2001; Hammarlund et al. 2007). Therefore, the severe morphological defects we observed in adult animals may reflect inappropriate regrowth subsequent to destabilization. We propose that the necrotic-like cell death and destabilization of neuronal processes observed in our studies may represent genetically separable events and suggest that our transgenic approach may afford a powerful system to tease apart the molecular pathways that differentially contribute to these two processes.

ACR-2 is part of a heteromeric nAChR in cholinergic motor neurons

We have demonstrated that acr-2 shows restricted expression to cholinergic motor neurons of the ventral nerve cord and appears diffusely localized in neuronal processes. These results suggest that the ACR-2 receptor complex may modulate motor neuron excitability by mediating signaling at extrasynaptic sites. Consistent with this notion, acr-2 loss-of-function mutants are not grossly uncoordinated and show only modest resistance to the paralyzing effects of the ACh esterase inhibitor aldicarb. Aldicarb-induced paralysis arises as a consequence of the prolonged action of ACh in the synaptic cleft; our analysis suggests that ACh release from motor neurons is decreased in acr-2 mutants. Another recent study reached a similar conclusion based on
electrophysiological analysis of acr-2 loss-of-function mutants (Jospin et al. 2009). We also show that the locomotory control interneurons (with the possible exception of AVE)—the major source of synaptic inputs to excitatory motor neurons—do not express a reporter that labels cholinergic neurons, suggesting these neurons are unlikely to be cholinergic. Therefore, what is the source of ACh for activation of ACR-2 receptor complexes? The presynaptic ACh release sites of en passant neuromuscular synapses are highly intermingled and densely packed because of the intercalation of neuronal processes in the nerve cord. Thus, one possibility is that these receptors are activated by spillover of ACh from release sites at nearby neuromuscular synapses.

Our genetic analysis showed that mutations in three genes encoding AChR subunits can suppress the neurotoxic effects associated with expression of pore-modified ACR-2(L/S) receptors. unc-38 and unc-63 are highly expressed in ventral cord motor neurons and also contribute to a heteromeric receptor complex that mediates excitatory neurotransmission at the NMJ (Fleming et al. 1997; Culetto et al. 2004). acr-12 is broadly expressed in the nervous system but is not expressed in body wall muscles (Cinar et al. 2005; Gottschalk et al. 2005). Our data are consistent with the notion that UNC-38, UNC-63, and ACR-12 coassemble with the ACR-2 subunit [either native ACR-2 or transgenic ACR-2(L/S)] to form heteromeric nAChRs in cholinergic motor neurons (Jospin et al. 2009). Loss-of-function mutations in any of these genes impair assembly or function of ACR-2 receptor complexes in cholinergic motor neurons and suppress ACR-2(L/S)-induced cell death. Several pieces of evidence support this idea. First, mutations in unc-29 and acr-16—genes that contribute to nAChRs at the NMJ and are essential for normal excitatory neurotransmission at neuromuscular synapses (Richmond and Jorgensen, 1999; Francis et al. 2005; Touroutine et al. 2005)—do not suppress ACR-2(L/S) neurotoxicity, indicating that reduced excitatory neuromuscular signaling alone is insufficient to suppress ACR-2(L/S)-induced toxicity. Second, specific expression of acr-12 in the cholinergic motor neurons of transgenic acr-12 mutants expressing ACR-2(L/S) was sufficient to produce paralysis, whereas specific expression of acr-12 in other neuron classes was without effect. Third, it has recently been shown that coexpression of five subunits—ACR-2, ACR-12, UNC-38, UNC-63, and ACR-3—was required for reconstitution of ACR-2 receptor complexes in a heterologous system (Jospin et al. 2009).

Pore-modified ACR-2(L/S) receptors cause ion channel-mediated neurotoxicity

We have shown that expression of the ACR-2(L/S) transgene leads to degeneration of the cholinergic motor neurons and paralysis, reinforcing the importance of these neurons in generating sinusoidal movement. The GABA motor neurons develop noradrenergic mechanisms of the central nervous system. Annu Rev Pharmacol Toxicol 47:699–729.

References


