Abstract. CAP23 is a major cortical cytoskeleton–associated and calmodulin binding protein that is widely and abundantly expressed during development, maintained in selected brain structures in the adult, and reinduced during nerve regeneration. Overexpression of CAP23 in adult neurons of transgenic mice promotes nerve sprouting, but the role of this protein in process outgrowth was not clear. Here, we show that CAP23 is functionally related to GAP43, and plays a critical role to regulate nerve sprouting and the actin cytoskeleton. Knockout mice lacking CAP23 exhibited a pronounced and complex phenotype, including a defect to produce stimulus-induced nerve sprouting at the adult neuromuscular junction. This sprouting deficit was rescued by transgenic overexpression of either CAP23 or GAP43 in adult motoneurons. Knockin mice expressing GAP43 instead of CAP23 were essentially normal, indicating that although these proteins do not share homologous sequences, GAP43 can functionally substitute for CAP23 in vivo. Cultured sensory neurons lacking CAP23 exhibited striking alterations in neurite outgrowth that were phenocopied by low doses of cytochalasin D. A detailed analysis of such cultures revealed common and unique functions of CAP23 and GAP43 on the actin cytoskeleton and neurite outgrowth. The results provide compelling experimental evidence for the notion that CAP23 and GAP43 are functionally related intrinsic determinants of anatomical plasticity, and suggest that these proteins function by locally promoting subplasmalemmal actin cytoskeleton accumulation.

Key words: neurite outgrowth • neuromuscular junction • synaptic plasticity • actin dynamics • nerve sprouting

Introduction

The plasticity of the nervous system in the adult involves alterations in the functional properties of preexisting circuitry as well as physical rearrangements of the synaptic circuits. In the latter case, synapses are formed or dismantled, and nerves sprout or retract. Such anatomical changes are thought to provide a basis for persisting alterations in circuit function, which is important for learning, memory, and repair. The molecular mechanisms that regulate anatomical plasticity in the adult are poorly understood, but important constraints that shape the plasticity of neural systems derive from specific intrinsic properties of their neuronal components (Bisby and Tetzlaff, 1992; Caroni, 1997, 1999; Sengpiel et al., 1998; Atoninii et al., 1999; Bravin et al., 1999). Consequently, identifying relevant intrinsic neuronal components and pathways that control growth in neurons should provide an entry point to dissect the mechanisms that control anatomical plasticity.

The neural growth–associated protein GAP43 has properties of a general intrinsic determinant of anatomical plasticity. First, its expression in neurons correlates closely with axonal elongation, synaptogenesis, and nerve sprouting during development and in the adult (Skene, 1989; Bisby and Tetzlaff, 1992). Second, overexpression of GAP43 promotes nerve sprouting in transgenic mice and process outgrowth in cultured cells (Yankner et al., 1990; Aigner et al., 1995). However, gene inactivation studies in mice did not reveal defects that would directly support the notion that GAP43 plays a general role in promoting axon elongation or nerve sprouting. Thus, although GAP43−/− mice exhibited high postnatal mortality, defects in the formation of axonal projections were restricted to growth cone guidance at specific choice points, and to the formation of specific topographic maps in the target region (Strittmatter et al., 1995; Sretawan and Kruger, 1998; Mair et al., 1999; Zhu and Julien, 1999). These results suggested that GAP43 is involved in translating signals important for growth cone guidance. However, mainly because of the fact that the molecular mechanisms through which GAP43 affects growth cone activity have remained unclear, the relation between this critical role of GAP43 and a hypothetical, more general function in promoting neurite outgrowth has been difficult to assess.
GA P43 may be related to the protein kinase C (PKC) substrates neurogranin, MARCKS, MacMARCKS, and CA P23 (A derem, 1995; Benowitz and R. outtenberg, 1997; Wiederkehr et al., 1997). GA P43 and neurogranin are expressed exclusively in the nervous system, whereas MARCKS and MacMARCKS are also expressed at high levels in many types of differentiating and motile nonneural cells. CA P23 is a neural protein, except for a widespread expression pattern during early embryonic development (Widmer and Caroni, 1990). Although GA P43, CA P43, and CA P23 have no sequence homologies, they share a number of characteristic properties (see Fig. 1 A). These include the following: (1) regulated, abundant expression related to contact-mediated differentiation, cell-surface activity, motility, and process outgrowth; (2) membrane association mediated by palmitoylation or myristoylation; (3) highly hydrophilic, with markedly acidic isoelectric points, rodlike structures, and a unique amino acid composition; (4) the presence of one unique stretch of 10–28 basic and hydrophobic residues, the effector domain (ED); this domain binds acidic phospholipids, calmodulin (Cam), and actin filaments in a mutually exclusive manner, and contains the PK C phosphorylation site(s); (5) colocalization at characteristic patches at the cell membrane; and (6) induction of dynamic actin structures at the surface of transfected cells (A derem, 1995; Benowitz and R. outtenberg, 1997; Wiederkehr et al., 1997). Like GA P43, expression of CA P23 is enhanced when axons elongate, and its overexpression in adult neurons of transgenic mice promotes nerve sprouting (Caroni et al., 1997). These observations have raised the possibility that CA P23 and GA P43 may be closely related functionally, possibly acting through a common mechanism to mediate morphogenetic processes, including neurite outgrowth.

Here, we tested the notion that GA P43 and CA P23 may be related functionally in vivo, and that they may play a critical role for nerve sprouting in the adult. CA P23-deficient mice exhibited high postnatal mortality, and pronounced abnormalities at the neuromuscular junction (nmj), including a nearly complete absence of stimulus-induced nerve sprouting (Celio et al., 1997). Sensory neurons was carried out according to the protocol described in Wiederkehr et al. (1997). The following antibodies were used: rabbit antiserum to COOH-terminal peptide sequences from CA P23, GA P43, and MARCKS (Wiederkehr et al., 1997; Laut et al., 2000); rabbit antiserum to rat CAPR was from Genosys Biotechnologies Inc.; mA b 34hL to rat substance P was from A. nava Trading; rabbit antiserum to parvalbumin was from SWant AG; rabbit antiserum to parvalbumin was a gift from H. Celi (University of Fribourg, Switzerland); FITC-choleratoxin B subunit was from Sigma Chemical Co. For immunocytochemistry of muscle tissue, mice were perfused with 4% paraformaldehyde-containing PBS, and cryostat sections were incubated with antibodies as described in Caroni et al. (1997). RITC- α-bungarotoxin and A lexα-label antibody were from Molecular Probes Inc. Transmission electron microscopy of mouse medial gastrocnemius muscle was according to standard procedures.

Materials and Methods

Targeting of Mouse CA P23 Gene

A genomic clone for the generation of a targeting construct was derived from a mouse 129/Sv BAC library (Genomys), using mouse CA P23 cDNA as a probe. The targeting strategies are outlined in Fig. 2 A (knockout mice) and Fig. 4 A (knockin mice). NLS lacZ corresponds to lacZ cDNA with a nuclear targeting sequence. IRES corresponds to an internal ribosomal entry signal sequence for bicistronic mRNA production. Embryonic stem cells and mice were screened with genomic Southern blots, using probes corresponding to the regions indicated in Fig. 2 A and Fig. 4 A. Northern blots were carried out with total RNA samples from adult mice brain according to standard procedures.

Sensory Neuron Cultures

Sensory neurons were derived from dorsal root ganglia of P0-P6 mice. Briefly ganglia freed of surrounding membranes were digested for 20 min at 37°C in the presence of 0.05% trypsin and 0.04% collagenase H. The digestion was stopped, and neurons were dissociated by four subsequent triturations steps. A filter washing, neurons were plated onto poly-L-lysine/ laminin (40 μg/ml)-coated glass coverslips at a density of ~2,000 cells/cm², in the presence of L 15, with antibiotics, 10% FCS, 30 mM NaHCO₃, and 50 ng/ml of either NGF or NT3 (Sigma Chemical Co.). Cells were analyzed 15 h after plating. Where indicated, cytochalasin D (Sigma Chemical Co.) was added to the culture medium 2.5 h after plating, at a final concentration of 2–20 nM. For quantitative analysis of neurite outgrowth patterns, type-identified neurons (expression pattern and morphology) from three independent cultures (total of 15 neurons [n]; 2 neurites analyzed per neuron) were scored for the following parameters: neurite length (average length of two longest neurites); blebbing (average number of blebs [swellings] per neurite length); hairs (average number of filopodial side-branches per neurite length); and branching (average number of neurite branch points per neurite length). To obtain a measure for neurite winding (winding factor), we subdivided neurites in 10-μm segments, and determined the number of times that the neurite crossed a straight line between the beginning and the end of the 10-μm segment. Total numbers of crossing points were normalized to neurite length, and the average values for 15 neurons were determined.

Results

Distinct Expression Patterns of CA P23 and GAP43 in the Adult Nervous System, and Expression of CA P23 at the Adult Neuromuscular Junction

During mouse nervous system development, CA P23, GA P43,
and MARCKS mRNAs were coexpressed at high levels in most, if not all types of neurons (not shown). GAP43 and CA P23 were downregulated with comparable kinetics that correlated with the time, when axonal arborization and synaptogenesis in the target region decline (Caroni and Becker, 1992; Caroni et al., 1997; Benowitz and Ruttenberg, 1997; Frey, D., and P. Caroni, unpublished results). In the adult, expression of these proteins was maintained in distinct, partially overlapping sets of neurons (Fig. 1 B; see McNamara and Lenox, 1997, for a detailed analysis of MARCKS and GAP43 expression patterns). First, there were significant differences in the expression patterns of CA P23 and GAP43 throughout major brain structures in the adult (Fig. 1 B). CA P23 expression was particularly high throughout the adult neocortex, where GAP43 was mainly expressed in layer V projection neurons. In contrast, CA P23 was expressed at very low levels in thalamic and brain stem structures, and expression was not detectable in the cerebellar cortex. Strong CA P23 expression was detected in the hippocampal formation including dentate gyrus granule cells, which do not express GAP43 in the adult. Second, at a higher resolution, the analysis revealed differences in the expression patterns of subsets of neurons. Thus, for example, apparently all DRG neurons expressed high levels of CA P23 mRNA (Fig. 1 B) and protein (see Fig. 6), whereas the expression of GAP43 was restricted to a subset of DRG neurons in the adult (Fig. 1 B).

Because of its unique anatomical arrangement, the neuromuscular junction provides a convenient experimental system to investigate stimulus-induced synaptic sprouting in the adult (Brown, 1984; Caroni, 1997). The expression of GAP43 in adult, unlesioned spinal motoneurons is undetectably low (Caroni and Becker, 1992), and botulinum toxin A (BotA)–induced sprouting at the adult nmj of GAP432/2 mice was not impaired (data not shown). Likewise, we could not detect MARCKS immunoreactivity at the adult nmj (not shown). In contrast, although it was significantly lower than during development, CA P23 mRNA was clearly detectable at clustered, large ventral horn neurons, presumably motoneurons, in the adult, and the corresponding protein could be detected at the adult nmj (Fig. 1 C). In addition, an antibody to the reporter gene NLS-lacZ visualized tSC immunoreactivity at nonlesioned, adult nmjs of CAP232/2 mice (Fig. 1 C).

**Critical Role of Motoneuron CAP23 for Stimulus-induced Nerve Sprouting at the Adult Neuromuscular Junction**

To investigate the role of CA P23 in neurite outgrowth and anatomical plasticity, we generated mice with an inactivated CA P23 allele (Fig. 2 A). The targeting construct included a nuclear lacZ gene to monitor CA P23 gene activity in the mutant mice (Fig. 2 A). CA P232/2 and CA P232/2 were born with Mendelian frequencies, but in the absence

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**Figure 1.** Expression patterns of mouse CA P23 and GAP43 in the adult. (A) Schematic representation of the structures of GAP43 and CA P23. The NH2-terminal with the acylation domains is to the left; (blue box) basic ED with PKC phosphorylation sites; and (white rectangles) predominantly acidic and highly hydrophilic regions with similar amino acid compositions. The relative dimensions of the proteins and the EDs are in scale. (B) CAP23 mRNA and GAP43 mRNA. (C) CAP23 immunoreactivity at the adult nmj. Double labeling for postsynaptic acetylcholine receptors (bottom, α8gtx) and CAP23 or reporter gene lacZ (top). Note the persistence of CAP23 signal at denervated (i.e., nerve-free) nmj, and synaptic, plus extrasynaptic (arrow) accumulation of reporter gene signal in homozygous CA P232/2 mice. Bar, 50 μm.
of CAP23, only ~10% of the mice survived to adulthood. Adult CAP23−/− mice weighed 50–70% of their wild-type littermates and were distinctly hyperactive, but they exhibited no elevated mortality. The brain of CAP23−/− mice exhibited enlarged ventricles, and pronounced axonal and synaptic ultrastructural abnormalities were detected in the hippocampus and neocortex (Frey, D., L. Xu, and P. Caroni, unpublished observations). In spinal motoneurons, GAP43 and CAP23 are expressed at high levels during axonal growth and target innervation, and are down-regulated to undetectable (GAP43) or low (CAP23) levels between postnatal day 8 and P15, when neuromuscular synapses initiate their final maturation (Caroni and Becker, 1992; Caroni et al., 1997). In the absence of CAP23, axon numbers in the peripheral nerves were normal, but both the nerve terminal and the tSC exhibited characteristic and unique abnormalities (Fig. 2, B and C) that became apparent from P12-15. In the medial gastrocnemius muscle, nerve terminal subdomains were markedly heterogeneous in size, with frequent, grossly oversized terminal regions and occasional swollen endings that were completely wrapped by tSCs (Fig. 2, B and C). The extent of these abnormalities had muscle region- and muscle type-specific features, and was least pronounced at slow-type neuromuscular synapses, e.g., in the soleus muscle (not shown). tSC processes were loosely packed and distinctly hypertrophic, with excess growth into the extracellular space (Fig. 2 C). These ultrastructural features and the corresponding overall appearance of these synapses are suggestive of instability with extensive ongoing remodeling (Bernstein and Lichtman, 1999).

tSC process outgrowth at the adult neuromuscular junction is known to promote synaptic sprouting (Son and Thompson, 1995). However, even when sprouting was induced by a local paralyzing injection of BotA, which induces strong sprouting in the soleus muscle of wild-type...
mice, CA P23<sup>−/−</sup> mice exhibited little or no nerve sprouting in this muscle (Fig. 3). To determine whether it was the absence of CA P23 in the motor nerve that impaired nerve sprouting, we crossed Thy1CA P23 mice that overexpress CA P23 from P7-9 on specifically and constitutively in neurons (Caroni et al., 1997) into the CA P23<sup>−/−</sup> background. This restored BotA-induced sprouting in the soleus (Fig. 3). Similar results were obtained when CA P23<sup>−/−</sup> mice were crossed into a Thy1GAP43 (Aigner et al., 1995) background (Fig. 3). Therefore, reexpression of CA P23 or GAP43 in adult motor nerves was sufficient to rescue nerve sprouting at the neuromuscular synapse.

GAP43 Can Functionally Substitute for CA P23 In Vivo

To determine whether, and to what extent, GAP43 can substitute for CA P23, we generated CA P23<sup>gap43/gap43</sup> knockin mice. The additional, bicistronic GAP43 mRNA species was larger than that of the endogenous GAP43 transcript, thus, allowing its independent detection on Northern blots. Heterozygous CA P23<sup>1/gap43</sup> mice expressed reduced levels of CA P23, and two different GAP43 transcripts, whereas homozygous CA P23<sup>gap43/gap43</sup> mice expressed no CA P23, and higher levels of the extra GAP43 transcript (Fig. 4 A). In situ hybridization analysis confirmed the additional expression of GAP43 in the characteristic CA P23 pattern, including strong signals throughout the neocortex and in dentate gyrus granule cells (not shown). Unlike CA P23<sup>−/−</sup> mice, CA P23<sup>gap43/gap43</sup> mice exhibited no elevated mortality (Fig. 4 B), and were normal in size. Ultrastructural analysis revealed no gross abnormalities at the neuromuscular junction, in the neocortex, and in the hippocampus (not shown). At the light microscopic level, nmjs of CA P23<sup>gap43/gap43</sup> mice were slightly broader than normal, but exhibited no signs of instability and no nerve terminal expansions (Fig. 4 C). Upon BotA-induced paralysis, nmjs in the soleus exhibited robust ultraterminal sprouting, although at close examination sprouts were more branched and shorter than in wild-type mice (Fig. 4 D). Therefore, GAP43 can effectively substitute for CA P23 in vivo, providing compelling experimental evidence for the notion that these proteins are related functionally, and exert partially redundant functions in vivo.

A Neurite Outgrowth Phenotype in the Absence of CA P23 Is Phenocopied by Cytochalasin D, a Drug that Inhibits Actin Filament Polymerization

To investigate more directly cell autonomous roles of CA P23 in neurite outgrowth, we analyzed dissociated DRG neuron cultures, which in a wild-type background express substantial levels of CA P23. For these experiments, neurons were plated on a laminin substratum (40 μg/ml) and cultured in the presence of NGF or NT3. When compared to wild-type neurons, neonatal sensory neurons from CA P23<sup>−/−</sup> mice extended axons that were thin and strikingly twisted, with frequent varicosities and characteristic bulbous endings (Fig. 5). Time-lapse video microscopic analysis suggested that, in the absence of CA P23, growth cones are less stable, failing to maintain a spread configuration, frequently changing growth direction, and exhibiting a tendency to extend, forward or backward, along neighboring neurites in the culture (not shown). Some of these features are strikingly reminiscent of chick DRG neurons depleted of GAP43 by an antisense approach, where the phenotype included reduced accumulation of filamentous actin in growth cones (Aigner and Caroni, 1995). Therefore, we determined whether the abnormal appearance of CA P23-deficient neurons was due to a defect in the actin cytoskeleton. As shown in Fig. 5, the neurite outgrowth phenotype could be reproduced to a
large extent by growing wild-type neurons in the presence of low concentrations of cytochalasin D, which inhibits actin polymerization and induces loss of growth cone actin structures (Fig. 5). As discussed below, these findings are consistent with the notion that the absence of CAP23 leads to selective deficits in the stability of growth cone actin structures.

Shared and Unique Roles of CAP23 and GAP43 in Neurite Outgrowth

To investigate the effects of CA P23 depletion and GAP43 replacement on defined subtypes of DRG neurons, we classified cultured neurons by their morphology and expression patterns, using CAP23, GAP43, MARCKS, parvalbumin, substance P, CGRP, calbindin,
and cholera toxin B as markers (Lawson, 1992). In addition, we analyzed cultures in the presence of either NGF or the related neurotrophin NT3, which supports a different subpopulation of DRG neurons (Friedel et al., 1997). The expression of CAP23 and GAP43 was restricted to neurons, whereas MARCKS was also expressed at high levels in glial satellite cells (not shown). All neurofilament-positive neurons in wild-type cultures expressed substantial levels of CAP23, whereas GAP43 and MARCKS were only expressed in subsets of neurons. For the purpose of this analysis, we focused on three operationally defined DRG subtypes: type A expressed high levels of CAP23, GAP43, and MARCKS, whereas types-B and -C expressed high levels of CAP23, but no detectable GAP43 or MARCKS (Fig. 6 A). Types-A and -B neurons grew several thin processes, with small growth cones, whereas type C neurons extended comparatively broad processes, with large, well spread growth cones.

In the absence of CAP23, type A neurons grew twisted processes (winding), which failed to consistently extend away from the cell body (Figs. 6 B and 7). There also was an increase in varicosities and blobby growth cones (blebbing) (Fig. 7). These defects were not noticeably alleviated when these neurons expressed GAP43 instead of CAP23 (Figs. 6 B and 7). In the absence of CAP23, type B neurites (no GAP43, no MARCKS) grew in a strikingly thin and meandering pattern (winding), and growth cones were blobby. In these neurons, expressing GAP43 instead of CAP23 rescued the thin axon and the winding phenotype (Fig. 7). In addition, when compared to wild-type type B neurons, knockin neurites expressing GAP43 instead of CAP23 exhibited frequent filopodia side branches (hairs) and growth cones spread more (Figs. 6 B and 7). Finally, in the absence of CAP23 type C neurons grew shorter, broad processes, that lacked a well defined transition between growth cone and neurite shaft region, and branched frequently (Figs. 6 B and 7). In the presence of GAP43 instead of CAP23, neurite length was restored, but branching remained elevated (Figs. 6 B and 7).

**Discussion**

We have provided evidence that CAP23 in motoneurons is critical for stimulus-induced nerve sprouting at the NMJ in the adult, and that GAP43 can, to a large extent, functionally substitute for CAP23 in mice. Abnormal neurite outgrowth in the absence of CAP23 can be phenocopied by cytochalasin D, which interferes with actin filament polymerization. Representative phase-contrast photographs of 15-h DRG cultures on laminin, in the presence of NGF. Note the thin, meandering neurites, varicosities, and bulbous growth cones (arrows, bottom row) in the absence of CAP23 or in the presence of cytochalasin D. Bars: 50 μm.
In the companion article (Laux et al., 2000), we show that GAP43, CAP23, and MARCKS are closely related mechanistically: they accumulate at subplasmalemmal rafts, where they modulate PI(4,5)P₂ and regulate actin dynamics. In the following sections, we discuss the roles of CAP23 at the nmj in the adult, and the implications of the finding that CAP23 and GAP43 have common as well as unique functions.

Roles of CAP23 at the Neuromuscular Junction

In the soleus muscle, blockade of neuromuscular transmission with BoTA induces ultraterminal nerve sprouting, which can be detected at >90% of the neuromuscular synapses, starting 3–5 d after toxin application (Brown, 1984; Frey et al., 2000). Sprouts grow in length and complexity for ~4 wk, and form ectopic synapses that restore neuromuscular transmission (DePaiva et al., 1999). In CA P23⁻/⁻ mice, <5% of the synapses exhibited any detectable sprouting 7 d after toxin application. This was not due to a reduced effectiveness of the toxin, as supported by the absence of muscle twitch upon nerve stimulation, pronounced muscle atrophy, and expansion of nerve branches at the nmj (see below). The absence of nerve sprouting...
was particularly striking, when considering the pronounced hypertrophy and process extension by synapse-associated tSCs in these mice, which can promote nerve sprouting in wild-type mice (Son and Thompson, 1995). A t 30 d after paralysis, when in wild-type mice sprouting and synaptogenesis peaked, sprouting in the absence of CAP23 was still restricted to less than half of the synapses in the soleus, where sprouts were very short, with no evidence of ectopic synapses. Reintroduction of either CAP23 or GAP43 in motoneurons after birth effectively rescued the sprouting defect, providing experimental evidence that intrinsic CAP23 or GAP43 is necessary for paralysis-induced ultraterminal sprouting at this synapse. At close examination, sprouts in the presence of GAP43 were shorter and more branched than in the presence of CAP23, a finding consistent with the effects of these proteins on nerve sprouting in the presence of wild-type CAP23 alleles (Caroni et al., 1997), and with the observation that CAP23 and GAP43 have shared, as well as unique functions in neurite outgrowth (see below). To our knowledge, these results provide the first experimental evidence for the notion that CAP23 (or GAP43) are intrinsic determinants promoting anatomical plasticity in the adult. These findings suggest that the expression levels of these proteins in a particular neuron may be one factor determining its tendency to exhibit anatomical plasticity in the target region (Caroni, 1997). In analogy to the results with PC12 cells (Laux et al., 2000), this pathway is likely to involve stimulus-induced formation of dynamic peripheral actin structures that are required for sprout formation and growth.

In addition to motoneurons, CAP23 was also expressed in tSCs at the nmj. In the absence of CA23, the nmj exhibited striking anatomical alterations, including highly irregular nerve profiles, large bloblike expansions at the end of terminal nerve branches, and tSC hypertrophy. These abnormal features became apparent during the third postnatal week, when nmjs expand in size, and when GAP43 cannot be detected anymore at this synapse. Although they recovered sprouting competence and nerve terminal branches were much more regular in shape, CA23−/−× Thy1CA23 mice still exhibited abnormally expanded synaptic regions, sprout endings (Fig. 3), and Schwann cell processes (not shown), suggesting that the absence of CA23 in tSCs led to a defect in synaptic growth control. Analysis of nerve–muscle preparations from CA23 mutant mice revealed that these neuromuscular synapses had average quantal contents about twice as high as controls, and that this phenotype was not affected by the reintroduction of CA23 in motoneurons (Frey, D., L. Xu, P. Larou, unpublished results). One possible interpretation of these findings is that tSCs control synaptic growth and synaptic strength at the adult neuromuscular junction through a mechanism that requires CAP23 in the tSC.

**CAP23 and GAP43 Are Closely Related Functionally**

Replacing the coding sequence of CAP23 with that of GAP43, thus, generating knockin mice that expressed no CA23, but instead GAP43, largely rescued the phenotypes caused by the absence of CA23. Rescued features included early mortality, body weight, nmj morphology, synaptic strength at the nmj (not shown), and stimulus-induced nerve sprouting at the nmj. Therefore, although they do not share homologous sequences, CAP23 and GAP43 are closely related functionally in vivo, suggesting that they may be components of the same regulatory pathway. In the companion paper, Laux et al. (2000), we show that GAP43, CA23, and MARCKS are closely related mechanistically, as central components of a novel pathway, to modulate the accessibility of PI(4,5)P2 at plasma-membrane rafts, and regulate actin dynamics and process outgrowth.

Although characteristic defects specifically because of the absence of CA23 could be detected in vitro (see below), nmj morphologies and sprouting patterns in knockin mice were slightly different than in wild-type mice, and homozygous knockin mice were sterile (not shown), the efficient functional replacement of CA23 by GAP43 is striking, and was not anticipated to this extent. Thus, although
the two proteins have similar effects in transfected cells (Wiederkehr et al., 1997), and both induce nerve sprouting at the adult nmj of transgenic mice (Caroni et al., 1997), they not only lack sequence homology, but their EDs have significantly different regulatory properties, and their association with the plasma membrane is also regulated differently (Fig. 8). Major differences include the following: (1) binding of calmodulin by the ED, which is calcium-dependent and of high affinity for CAP23, but calcium-independent and of low affinity for GAP43 (Skene, 1989; Maekawa et al., 1994; Benowitz and Ruttenberg, 1997); (2) GAP43 is effectively phosphorylated by protein kinase C in situ, whereas under comparable experimental conditions, only a small fraction of CAP23 is phosphorylated (Widmer and Caroni, 1990); and (3) GAP43 tightly associates with the plasma membrane via palmitoylation, whereas membrane association by CAP23 not only depends on myristoylation, but also on the presence of the ED (not shown). One interpretation of these findings is that certain cellular functions depend critically on the expression of a minimal amount of many of these related proteins. These minimal requirements may not depend critically on the specific regulatory properties of CAP23 or GAP43 (Fig. 8). This interpretation is consistent with the observation that CAP23−/− mice, which expressed a third of wild-type CAP23 levels, exhibited a partial phenotype, with characteristic, but weaker, deformations and bloblike expansions at the nmj, average weights that were ~80% of normal, slightly elevated mortality, but no alterations in nmj synaptic strength. This haplo-insufficiency of CAP23 supports the notion that CAP23 levels can be critically important, and that GAP43 may rescue these deficits via a quantitative mechanism. The expression of GAP43 and CAP23 (and MARCKS; Laux et al., 2000) is highly regulated with respect to cell type, stimulus responsiveness, and quantity, and there is a general tendency for markedly reduced levels of these proteins in adult neurons. We suggest that the quantitative decline in neuronal CAP23 and GAP43 levels may be one factor restricting anatomical plasticity in the adult nervous system.

**Unique Properties of CAP23 and GAP43 in Neurite Outgrowth**

The analysis of dissociated DRG neurons in culture revealed abnormalities in neurite outgrowth that were specifically due to the absence of CAP23, and were not compensated by GAP43. These were evident in the global effects of CAP23 depletion on DRG cultures (Fig. 5) as well as in the effects on type A neurons, which expressed CAP23, GAP43, and MARCKS in a wild-type background, and no CAP23, but substantially higher levels of GAP43 immunoreactivity in a knockin background (Figs. 6 and 7). Deficits specifically due to the absence of CAP23 were thinner neurites, a characteristic winding pattern of neurite outgrowth, and a prevalence of pronoucedly bulbous growth cones (Figs. 5–7). These defects could be phenocopied by low doses of cytochalasin D (Fig. 5), suggesting that they were specifically due to deficits in the actin cytoskeleton in the absence of CAP23. Interestingly, a striking excess of winding neurites, with large bloblike endings was also detected during the reinnervation of skeletal muscle by regenerating peripheral nerves of CAP23−/− mice (not shown). Time-lapse analysis revealed that in the absence of CAP23, growth cones frequently changed extension direction, indicating that the winding pattern was due to an alteration in growth cone behavior. A closer inspection, growth cones that did exhibit a spread configuration were abnormally phase-dense, and labeling for microtubules revealed unusually high densities and disordered arrangements of this cytoskeletal element in the growth cone (not shown). Combined with the similar effects of cytochalasin D on neurites and growth cones, these observations suggest that in the absence of CAP23, growth cone actin barriers that restrict microtubule invasion (Forscher and Smith, 1988) are impaired. The frequent and abnormal occurrence of motile activity from the shaft region of the growth cone, which was again phenocopied by cytochalasin D (not shown), suggests that CAP23 may play a critical role to form an actin-based cortical cytoskeleton at the transition zone between growth cone and neurite. Such a role may also be required to prevent the unusual deformations and the striking bloblike expansions at the nmj of CAP23−/− mice.

The analysis of type B and type C DRG neurons that expressed high levels of CAP23, but no detectable GAP43 nor MARCKS provided further valuable information about common and shared properties of CAP23, GAP43, and MARCKS (Laux et al., 2000) in neurite outgrowth. First, although they exhibited anomalous growth patterns, neurites still extended in the absence of all three proteins (Fig. 6). This is in apparent contrast to the observation that the dominant-negative ΔED mutants suppressed neurite outgrowth in PC12 cells, and delayed peripheral nerve regeneration in transgenic mice (Laux et al., 2000). Possibly, DRG neurons express further functionally related proteins, such as MacMARCKS (Aderem, 1995) or the more remotely related protein paralemmin (Kutzleb et al., 1998). Alternatively, explanations include that DRG neurons may extend neurites more vigorously than PC12 cells, thus not critically depending on the function of these proteins. Second, GAP43 partially restored neurite outgrowth properties in type B (neurite diameter and winding) and
type C (neurite length) neurons, whereas neurites of type A neurons were comparable in knockout and knockin cultures. This observation further supports the notion that CA P23 and GAP43 are closely related functionally. Third, the expression of GAP43 instead of CA P23 in type B and type C neurons led to new neurite outgrowth features in these neurons, such as enhanced growth cone spreading and filopodia. These specific effects of GAP43 on neurite outgrowth in vitro are reminiscent of its effects on nerve sprouting at the NMJ (Caroni et al., 1997), and suggest that GAP43 may predominantly affect different aspects of growth cone actin dynamics than CA P23 (Aigner and Caroni, 1995).

What mechanisms could be responsible for the differential effects of CA P23 and GAP43 on neurite outgrowth? As summarized in Fig. 8, CA P23 and GAP43 are regulated differently, with respect to ED interactions and retention at the cell membrane. In addition, the ED is positioned differently relative to the NH2-terminal and the acylation site of these proteins. These differences could affect the retention of CA P23 and GAP43 at growth cone subregions, resulting in different contributions of the two proteins to growth cone actin dynamics. In addition, the different responsiveness of the EDs of CA P23 and GAP43 to calcium/calmodulin and PKC may lead to a predominant effectiveness of each protein at different points in the growth cone regulation cycle. A detailed elucidation of the mechanisms involved in differential regulation of actin dynamics and growth cone activity by CA P23 and GAP43 should shed light on their specific functions in the nervous system, during development and in the adult.

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