Neurobiology of Disease

Spinocerebellar Ataxia Type 6 Protein Aggregates Cause Deficits in Motor Learning and Cerebellar Plasticity

In both human and mouse, numerous missense mutations in the P/Q-type Ca\(^{2+}\) channel sequence have been identified and linked to neurological defects (Pietrobon, 2005). In humans, mutations of the P/Q-type Ca\(^{2+}\) channel sequence lead to dominantly inherited neurological diseases that manifest as migraines in familial hemiplegic migraine-1, episodic ataxia, and epilepsy in episodic ataxia type 2, and to pure cerebellar ataxia in spinocerebellar ataxia type 6 (SCA6; Bidaud et al., 2006; Pietrobon, 2010). SCA6 has been identified as a poly-glutamine (polyQ) disease in which 21–33 residues are found in exon 47 of the P/Q-type channel C terminus (CT; Zhuchenko et al., 1997; Zoghbi and Orr, 2000; Mantuano et al., 2003). In humans, exon 47 undergoes alternative splicing, leading to CT-terminal isoforms containing either exon 47 alone or with the polyQ repeats (Zhuchenko et al., 1997). Protein aggregates containing the CT are present in the nucleus, cell bodies, and dendrites of Purkinje cells (PCs) SCA6 patients (Ishikawa et al., 1999a, 2001; Ishiguro et al., 2010).

Two main disease-related hypotheses have been suggested for CT-containing protein fragments found in SCA6 patients. First, the CT-containing proteins may constitute a degradation product of the channel that is more stable and therefore enriched in the disease form (Bidaud et al., 2006). This hypothesis is supported by the fact that CT domains of the Ca\(_{\alpha}2.1\) subunit are susceptible to proteolytic processing (Kubodera et al., 2003; Kordasiewicz et al., 2006). Indeed, when Ca\(_{\alpha}2.1\) subunits are expressed in HEK293 cells, the presence of 60–75 kDa peptides can be recognized by an antibody to the CT domain (Kubodera et al., 2003), and subunits carrying polyQ residues with different lengths (11 and 27) show differences in the stability of the proteolytic products (Kubodera et al., 2003). Second, the CACNA1A gene encoding the \(\alpha1\)A subunit may contain a transcription factor with an altered function in the disease form (Du et al., 2013). This hypothesis has been derived from the observation that the CT-containing polyQ stretch can induce a form of cell death that depends on its nuclear localization (Kordasiewicz et al., 2003).
2006). Interestingly, this CT-containing protein peptide can be generated by a bicistronic mRNA of the Cav2.1 α subunit that functions as a transcription factor to control expression of PC genes but reduces viability of PCs and induces cerebellar atrophy in the disease form (Du et al., 2013).

Cerebellar syndromes in SCA6 patients, such as ataxia of gait and stance, dysmetria, and oculomotor deficits, most likely develop as a result of changes in cerebellar output (De Zeeuw et al., 2011). The deficits in cerebellum function in SCA6 patients are suggestive for impairments in eyeblink conditioning (EBC), a form of associative motor learning that depends on the integrity of cerebellar synaptic plasticity (Kim and Thompson, 1997; Boele et al., 2010).

To analyze the physiological and behavioral consequences of CT polypeptides as predicted and found in SCA6 patients, i.e., CT-short (ending at exon 46) and CT-long polyQ (ending at exon 47 and containing 27 polyQ residues), we exogenously expressed the human CT polypeptides in mouse cerebellar PCs using viral and transgenic approaches. Our findings reveal that, as in SCA6 patients, a clustering of the CTs in nuclear inclusions and cytoplasmic aggregates occurred in infected PCs. Postnatal expression of the disease peptide causes SCA6 symptoms, including ataxia and PC degeneration, that are correlated with altered PC firing in vivo. Most importantly, expression of the SCA6 peptide causes a virtually complete loss of EBC that is correlated with the loss of cerebellar plasticity (i.e., LTP and LTD) at the parallel fiber (PF)-to-PC synapse. Thus, our results describe for the first time the physiological consequences of the human polyQ CT peptide as found in SCA6 patients on cerebellar plasticity. In addition, this study identifies an alteration in EBC as a new diagnostic strategy to detect Ca2+-channel-mediated diseases at an early disease state.

Materials and Methods

The constructs for the human Caα2.1,2.1 subunit corresponding to short or long isoforms were a generous gift from Dr. Cheng Chi Lee (University of Texas-Houston Medical School, Houston, TX). The Caα2.1,2.1 short ends at exon 46, whereas the long isoforms contain an additional pentanucleotide sequence before the stop codon (GCCGAC), inducing a frame shift. Therefore, the long isoforms express the additional exon 47. Of the murine Cav2.1, the long isoforms ends at exon 46, whereas the long isoforms contain an additional pentanucleotide sequence before the stop codon (GGCAG), inducing a frame shift. Therefore, the long isoforms express the additional exon 47.

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Electrophysiology

For recordings of EPSCs in PCs, whole-cell voltage-clamp recordings were made at 30°C from cerebellar slices as described previously (Womack and Khodakhah, 2003; Mark et al., 2011; Maejima et al., 2013). Parasagittal cerebellar slices (250 μm thick) were prepared on a Leica VT 1000S vibratome from at least four 21- to 32-old male mice. PCs were identified visually under an upright microscope (BX51 WI; Olympus) equipped with infrared illumination. During experiments, slices were perfused continuously with an artificial CSF (aCSF) containing the following (in mM): 125 NaCl, 2.5 KCl, 1.25 NaH2PO4, 30 NaHCO3, 1.25 NaHPO4, and 20 glucose (equilibrated with 95% O2 and 5% CO2). Borosilicate glass electrodes (2–4 MΩ) were filled with an internal solution containing the following (in mM): 60 CsCl, 10 MgCl2, 20 NaH2PO4, and 20 Na-acetate. Microscopy and image analysis

The distribution of the YFP-tagged Caα2.1,2.1 CT proteins in cerebellar slices was analyzed using a Leica TCS SP5 confocal laser scanning microscope (DMI6000 B; Leica) interfaced to a personal computer, running Leica Application Suite Advanced Fluorescence software (LAS AF 2.6). A 10×/0.3 numerical aperture (NA), 20×/0.7 NA, or 40×/1.1 NA objective was used to capture all photomicrographs. Sequential z-stacks were made for each section, and crosstalk of the fluorophores was eliminated automatically with LAS AF software. YFP fluorescence was detected by excitation with an argon laser (line 514 nm). Slow-fade with DAPI (Invitrogen) was used as mounting medium to label the nuclei of HEK293 cells and was detected by excitation with a diode laser at 405 nm. HEK293 cells were fixed for 10 min with a 4% formaldehyde solution (Sigma) in PBS, followed by PBS washes. Coverslips were mounted with Slow-Fade DAPI and left at room temperature overnight. For long-term storage, sealed coverslips were kept at 4°C. DAPI and CT YFP fluorescence was also detected on the upright Leica DMSFA using a 63× water objective (1.2 NA) coupled to a D5G light source (Sutter Instruments). The following Chroma filter sets were used (excitation, dichroic, emission, respectively): DAPI: D350, 50×, 400DCLP, D460/50M; YFP: HQ500/20s, Q515LP, HQ535/30m. Image analysis and deconvolution of images were performed using Volocity software.
and holding potential of 0 mV. All measurements of intrinsic firing properties were made at 35°C bath temperature. A combination of EPC10/2 amplifier and PatchMaster software (HEKA Elektronik) was used to control membrane voltage, stimulation timing, and data acquisition. Series resistance was compensated routinely by 70–80%. Membrane currents were filtered at 3 kHz and digitized at 20 kHz.

**Extracellular in vivo recordings of PCs**

Recordings have been performed as described previously (Kruse et al., 2014). Briefly, adult mice (one male and two females for floxed (fl)-CT-short; three males for CT-shortPC; three males for fl-CT-longQ27; and three males for CT-longQ27PC) were anesthetized with an intraperitoneal injection of ketamine (10 mg/kg)/xylazine (20 mg/kg), with supplemental ketamine doses during the experiment. After placing the mice in a stereotactic frame (Narishige), a sagittal incision was performed along the midline to expose the cranium and a window 6 mm caudal from bregma to give access to the vermis. A multielectrode device (Eckhorn microdrive; Thomas Recording) was used to monitor single PCs.

Single-cell activity was sampled with high temporal resolution (32 kHz) and analyzed offline for spike detection. Alternatively, signals of multiple electrodes were routed through an online spike sorter (Plexon), and time markers for each detected action potential was stored. Throughout the experiments, PCs were identified by the combined occurrence of simple and complex spikes. Complex spikes can only be detected in PCs and represent the climbing fiber input into PCs (Schmolesky et al., 2002).

**Figure 1.** CT-short predominantly targets to the nucleus, whereas CT-longQ27 is found in both the cytoplasm and the nucleus in HEK293 cells and cerebellar PCs. A, Schematic of the human Cav/H925112.1 isoforms and the derived C-terminal constructs as predicted and observed in SCA6 patients. Top, Full-length (FL) cDNAs of the human Cav/H925112.1 isoforms reveal the different splice variants. FL-short ends after exon 46, whereas FL-longQ27 contains the coding sequence of the additional exon 47. The polyQ stretch (27 polyQs) is located within exon 47. Bottom, The different CTs were fused to YFP. B, CT-short is targeted to the nucleus, whereas CT-longQ27 is targeted to the cytoplasm and nucleus in HEK293 cells. HEK293 cells were transfected with CT-short or CT-longQ27 and incubated for 14–20 h at 37°C. B1, Phase-contrast and DAPI and YFP fluorescence (bottom) images of transfected HEK293 cells. B2, xyz images of the cells shown in B1 demonstrates that CT-short formed nuclear aggregates that are larger in size compared with nuclear and cytoplasmic CT-longQ27 formed aggregates. B2, Bar graph of the fluorescence distribution of CTs between the nuclear and non-nuclear region of the cell. Cell borders were identified by the phase-contrast image, whereas nuclear regions were defined by the DAPI staining. The diagram shows that >90% of the YFP fluorescence of CT-short is found in the nucleus, whereas 50–60% of the CT-longQ27 YFP fluorescence is outside of the nuclear areas. B3, Diagram of the size of CT-containing aggregates reveal that CT-short forms larger aggregates than CT-longQ27. Statistical significance was evaluated with ANOVA (*p < 0.05, **p < 0.01). Error bars indicate SEM. C, CT-short is targeted predominantly to the nucleus, whereas CT-longQ27 is found predominantly in cytoplasmic aggregates in cerebellar PCs. AAV2-mediated expression of CT-short and CT-longQ27 in cerebellar PCs 3–4 weeks after injection into 2- to 3-week-old mice. C1, CT-short is found in large nuclear aggregates and distributed diffusely in some of the proximal dendrites. CT-longQ27 is found in small aggregates in the soma and in particular in the dendrites of PCs. C2, Number of PCs expressing CT-short (C2) or CT-longQ27 (C3) that reveal a punctate distribution in the cytoplasm or a diffuse or punctate distribution in dendrites. CT-longQ27 is found in small aggregates in the soma and in particular in the dendrites of PCs. Statistical significance was evaluated with ANOVA (**p < 0.01). Error bars indicate SEM.
Creation of transgenic animals and transgene detection
For stable integration of Ca\(^{2+}\)/H\(_{11001}\) channel constructs into the mouse genome, the human CT-short and CT-long\(\text{Q27}\) was cloned into pCZW (Braz et al., 2002), which contains a CMV and \(\beta\)-actin promoter and a lacZ expression cassette that is flanked with loxP sequences. After Cre-mediated recombination, the lacZ expression cassette is excised out, resulting in mice (CT-short\(\text{PC}\) and CT-long\(\text{Q27}\)\(\text{PC}\)) expressing the CTs only in the cell types containing the Cre recombinase. PolyA, Polyadenylation. Western blot analysis of cerebellar homogenates from adult mouse brains expressing YFP–CT-short and YFP–CT-long\(\text{Q27}\) specifically in PCs. YFP–CT-short and YFP–CT-long\(\text{Q27}\) were detected using a GFP antibody, and \(\alpha\)-tubulin was used as a loading control. No differences were detected in the expression levels between YFP–CT-short and YFP–CT-long\(\text{Q27}\) expressing mice. CT expression in fl-CT-short and fl-CT-long\(\text{Q27}\) mice and Cre recombinase-induced expression of CT-short\(\text{PC}\) and CT-long\(\text{Q27}\)\(\text{PC}\) in PCs (arrows show PC layer) resulted in a loss of \(\beta\)-gal staining in PCs but not in hippocampus (inset; top). PC-specific expression of Cre recombinase in CT-short\(\text{PC}\) and CT-long\(\text{Q27}\)\(\text{PC}\) mouse lines resulted in a loss of \(\beta\)-gal staining in PCs (bottom) but not in hippocampus (inset; top) and a gain of YFP–CT-short and YFP–CT-long\(\text{Q27}\) expression in PCs (D). YFP was visualized with a GFP antibody in cerebellar sagittal sections from adult mice.

Figure 2. Floxed CT peptide expressing mice for cell-type-specific Cre recombinase-mediated expression and characterization of CT-short and CT-long\(\text{Q27}\) in PCs of the cerebellum. A, Schematic description of the constructs used to create the transgenic animals expressing CT-short (fl-CT-short) or CT-long\(\text{Q27}\) (fl-CT-long\(\text{Q27}\)). CT-short and CT-long\(\text{Q27}\) were both cloned into pCZW (Braz et al., 2002), which contains a CMV and \(\beta\)-actin promoter and a lacZ expression cassette that is flanked with loxP sequences. After Cre-mediated recombination, the lacZ expression cassette is excised out, resulting in mice (CT-short\(\text{PC}\) and CT-long\(\text{Q27}\)\(\text{PC}\)) expressing the CTs only in the cell types containing the Cre recombinase. PolyA, Polyadenylation. B, Western blot analysis of cerebellar homogenates from adult mouse brains expressing YFP–CT-short and YFP–CT-long\(\text{Q27}\) specifically in PCs. YFP–CT-short and YFP–CT-long\(\text{Q27}\) were detected using a GFP antibody, and \(\alpha\)-tubulin was used as a loading control. No differences were detected in the expression levels between YFP–CT-short and YFP–CT-long\(\text{Q27}\) expressing mice. C, LacZ expression in fl-CT-short and fl-CT-long\(\text{Q27}\) mice and Cre recombinase-induced expression of CT-short\(\text{PC}\) and CT-long\(\text{Q27}\)\(\text{PC}\) in PCs (arrows show PC layer) of the cerebellum. Cerebellar sagittal sections from fl-CT-short and fl-CT-long\(\text{Q27}\) adult mice reveal \(\beta\)-gal staining in cerebellar PCs (bottom) and the hippocampus (inset; top). PC-specific expression of Cre recombinase in CT-short\(\text{PC}\) and CT-long\(\text{Q27}\)\(\text{PC}\) mouse lines resulted in a loss of \(\beta\)-gal staining in PCs (bottom) but not in hippocampus (inset; top) and a gain of YFP–CT-short and YFP–CT-long\(\text{Q27}\) expression in PCs (D). YFP was visualized with a GFP antibody in cerebellar sagittal sections from adult mice.
nase under the PC-specific Pcp2/L7 promoter. The Pcp2 promoter is a widely used promoter for driving PC-specific expression. Expression of transgenes under this promoter can be first detected at postnatal day 6 and full expression after postnatal week 2–3 (Barski et al., 2000). Because the promoter is not active early in development, we did not see any developmental problems related to the transgene expression in previous studies (Mark et al., 2011).

Western blots from three adult male age-matched fl-CT-short, fl-CT-longQ27, CT-shortPC, and CT-longQ27PC mouse brain homogenates were performed according to the methods of Han et al. (2006). The cerebellum from at least three adult male mice per genotype were analyzed. Detection of YFP constructs with anti-GFP (1:1000; Frontier Institute), followed by a secondary horseradish peroxidase antibody (1:5000; Pierce) was used. The blots were stripped with a mild stripping buffer (Abcam) and reincu-

Figure 3. Behavior analyses of CT-shortPC and CT-longQ27PC mice. Comparisons between CT-shortPC and CT-longQ27PC and fl-CT-short and fl-CT-longQ27 control mice on rotarod (A), beam-walk (B), and incline screen (C) tests. Statistical significance in all behavior experiments was evaluated with ANOVA. *p ≤ 0.05, **p ≤ 0.01. Data are reported as mean ± SEM. The number in parentheses indicates the number of animals tested.
bated with an anti-α-tubulin antibody (1:1000; Thermo Fisher Scientific) as a loading control. Blots were developed according to the SuperSignal West Dura Extended Duration Substrate protocols (Pierce).

Analysis of ataxic behavior
Analysis of ataxic behavior has been described in detail previously (Mark et al., 2011; Maejima et al., 2013) and briefly below.

Rotarod. Mice (four males and four females for fl-CT-short; six males and five females for CT-shortPC; and four males and four females for fl-CT-longQ27; and four males and four females for CT-longQ27PC) were placed on a rotarod (Columbus Instruments) rotating at 4 rpm for 1 min of acclimation. The rod was then accelerated at 0.1 rpm/s up to 40.0 rpm. The test continued until all mice had fallen off the rod. Latency to fall and rotations per minute at the time of fall were recorded for each mouse. Data were averaged over three trials per mouse.

Incline screen test. Two wire mesh screens, one at 60° angle and the other at 90° angle, were used for this test. Mice (two males and two females for fl-CT-short; six males and five females for CT-shortPC; and six males and five females for fl-CT-longQ27; and six males and five females for CT-longQ27PC) were placed facing downward in the middle of the screen. Latency to turn around and climb to the top was recorded. Maximum latency to complete the task was set at 60 s. If the mouse fell, a time of 60 s and a fall notation were recorded.

Beam walk. Mice (two males and three females for fl-CT-short; 10 males and eight females for CT-shortPC; and six males and four females for fl-CT-longQ27; and six males and four females for CT-longQ27PC) were analyzed for their capability to cross a narrow beam to an enclosed goal box. The horizontally placed, 70-cm-long beam was 10 mm wide and 60 cm above the table surface. One end of the beam was mounted to a small, illuminated platform, and the other end was fastened to an enclosed (15 cm²) goal box. Mice underwent training for 2 d (six trials per day) before data collection. Briefly, the mouse was placed at an illuminated end of the beam, and the time required to traverse the beam to the goal house was recorded. Mice were given a maximum of 120 s to transverse the beam, and a fall was also recorded as 120 s. In addition to recording the latency to transverse the beam, time spent immobile (idle), left and right paw slips, and falls were noted (Quinn et al., 2007). Data were averaged over three trials per mouse.

EBC
EBC was performed as described previously (Koekkoek et al., 2002; Boele et al., 2013). Briefly, at least four agematched (6 – 8 months old) male mice underwent a 3 d habituation (one session per day), followed by a 6 d training and 3 d extinction period. Animals were housed in their home cage until the start of each session. EBC experiments were performed in an isolated training box (Blink2.0; Neurasmus). Animals were placed inside of a slow rotating wheel to keep the mice alert during the sessions. Before the start of the habituation session, the minimal tone intensity that did not induce a startle response was determined (−70 dB). A sinuisoidal tone with a duration of 380 ms (rise/fall time 25 ms) and a frequency of 5 kHz were used as the conditioning stimuli (CS). The unconditioned stimuli (US) consisted of a 30-ms-long mild air puff to the cornea of the eye. In the training session, the CS was paired with the US with an interstimulus interval of 350 ms. During each session, animals received 64 trials in eight blocks. Each block consisted of one US, followed by six paired CS–US, followed by one CS, with each stimuli separated by a random intertrial interval of 30 s (± 10 s). Eyelid movements were monitored with the magnetic distance measurement technique (MDMT; Koekkoek et al., 2002). A magnetosensitive chip placed above the upper eyelid detects the movement of a magnet that has been placed on the lower eyelid of the mice before the recording session. During the habituation and extinction sessions, the blocks consisted of randomly paired US-only and CS-only trials. All experiments were approved by the Institutional Animal Research Facility.

Results
CT-short forms mainly nuclear inclusions, whereas CT-long forms mainly cytoplasmic aggregates in HEK293 cells and cerebellar PCs
PolyQ diseases are characterized by protein aggregations within the nucleus (nuclear inclusions) and cytoplasm (cytoplasmic ag-

Figure 4. Degeneration of PCs with age in CT-shortPC and CT-longQ27PC mice. A, Representative images of the PC layer in cerebellar slices from 18-month-old CT-shortPC and CT-longQ27PC mice. Arrows show missing PCs. B, Quantification of the number of PCs per area in cerebellar slices for 4- and 18-month-old CT-shortPC and CT-longQ27PC mice. Statistical significance was evaluated with ANOVA (**p < 0.01). Error bars indicate SEM.
SCA6 patients, we created two transgenic mouse lines (fl-CT-short and fl-CT-longQ27) for cerebellar expression of these constructs specifically in PCs using the Cre-lox system (Fig. 2A). Crossing the floxed CT-expressing mice with a mouse line that expresses Cre recombinase specifically in PCs (1gCPCP2-Cre, Barski et al., 2000) resulted in PC-specific expression of CT peptides beginning 1–3 weeks after birth (Fig. 2B–D; Mark et al., 2011). These mice expressing the CTs specifically in PCs of the cerebellum will be referred to as CT-shortPC and CT-longQ27PC. No differences in the amount of CT protein expression was found between the CT-shortPC- and CT-longQ27PC-expressing mice (Fig. 2B, D).

Postnatal expression of CT-longQ27PC but not CT-shortPC causes ataxia and alteration in the firing properties of PCs in vivo

SCA6 patients are characterized by late-onset ataxia and degeneration of cerebellar PCs. Therefore, we analyzed motor behavior of CT-shortPC- and CT-longQ27PC-expressing mice. We found that CT-longQ27PC-expressing mice develop signs of ataxia beginning at 8 months of age when tested on the rotarod (Fig. 3A). Rotarod performance declined further with age (Fig. 3A). Impairment of motor performance became also obvious in the incline screen test and the beam-walk test (Fig. 3B, C). In contrast, CT-shortPC-expressing mice developed late signs of ataxia only after 18 month of age and only on the rotarod but not on the incline screen or beam-walk tests (Fig. 3A–C). Both mouse lines showed an age-dependent reduction in the number of PCs. However, reduction in PC number occurred much earlier in CT-longQ27PC (after 4 months) compared with CT-shortPC (after 18 months)-expressing mice (Fig. 4).

Ataxia is most likely caused by an alteration in the information processing and output signals of the cerebellar cortex because of either neuronal degeneration (see above) or changes in the information processing of PCs before neuronal death. Therefore, we analyzed what extent expression of CT peptides interferes with the firing properties of PCs in vivo.

Indeed, simultaneous multiple electrode recordings revealed more irregular simple spike firing in CT-longQ27PC compared with CT-shortPC and control mice. The intermittent firing was not a ubiquitous and/or continuous feature of all PCs in CT-longQ27PC mice, because we sometimes recorded intermittently firing PCs together with regularly firing PCs. When multiple PCs with irregular firing were recorded in parallel, pauses in simple spike firing appeared independently at different cells with no indication of temporal coupling of pauses in different PCs (Fig. 5A, bottom). The mean frequency of simple spike firing was reduced significantly in CT-longQ27PC (Fig. 5B, left), which is probably a direct consequence of occurrence of pauses in simple spike trains. The simple spike irregularity, quantified by the coefficient of variation (CV) of the interspike interval distribution, was increased in both CT-shortPC and CT-longQ27PC when compared with control mice (CT-short, CV = 1.20 vs 0.74, p < 0.05; CT-longQ27PC, CV = 1.62 vs 0.61, p < 0.01; Fig. 4B, right).

The results on simple spike firing frequency and CV suggest that CT-longQ27PC mice have a higher incidence of irregular firing cells compared with CT-shortPC and control mice. To distinguish regularly from intermittently firing PCs, we plotted the simple spike rate versus the CV values. Regularly firing PCs might form a cluster with CVs < 1 and higher rates compared with irregularly firing PCs that are characterized by CVs > 1 and lower rates (Yartsev et al., 2009). The scatter plot revealed that CT-shortPC and CT-longQ27PC show two subpopulations of PCs...
when CV = 1 is used as a threshold (Fig. 5C). Accordingly, the incidence of intermittent firing cells (CV >1) was highest in CT-longQ27PC (23 of 51 PCs = 45.1%, three mice, 17.1–28.9 months) compared with 23.5% (8 of 34 cells) in CT-shortPC mice (n = 3, 16.1–33.9 months) and 9.1% (3 of 33) in fl-CT-longQ27 (control, three mice, 13.5–16.7 months) and 17.0% (8 of 47) in fl-CT-short mice (control, n = 3, 16.1–32.6 month). No differences were observed in the rates and regularity of complex spikes (Fig. 5D). Thus, the data suggest that PCs expressing CT-longQ27PC mice reveal a higher incidence of irregular firing. The CV for adjacent intervals (CV2) did not reveal significant differences between CT-short and CT-longQ27PC. This indicates that the variability in the simple spike firing in CT-longQ27PC is generated by slow modulations and longer pauses in firing that have a significant influence on the CV measure but weak influence on CV2.

Next we wanted to investigate the effects AAV2-expressing CTs (CT-short and CT-longQ27) on the spontaneous firing of PCs in the presence of synaptic transmission blockers. To accomplish this, we stereotactically injected AAV2-expressing CTs in 3- to 4-week-old mice and recorded 10–14 d later from positively expressing PCs in cerebellar slices. Spontaneously active PCs in cerebellar slices revealed either a tonic or a trimodal firing pattern, possibly reflecting enhanced bistability or tristability (Schonewille et al., 2006). PCs in the trimodal firing state cycled continuously among a silent, tonically active and bursting phase (Womack and Khodakhah, 2002; Fig. 6C1). We first analyzed tonically firing PCs and found that expression of CT-short and CT-longQ27 revealed a reduction in the firing frequency (Fig. 6A2). PCs expressing CT-longQ27-expressing PCs fire more irregularly compared with CT-short-expressing PCs as suggested by the higher CV (CT-short, CV = 0.10 ± 0.02; CT-longQ27, CV = 0.203 ± 0.049; Fig. 6A3). We observed more PCs expressing CT-short in the trimodal state compared with control and CT-longQ27-expressing PCs, whereas more PCs expressing CT-longQ27 appeared in the silent mode (putative downstate;
**A** Experimental Procedure

![Diagram of experimental procedure]

**B** Conditioning Day 2

![Graph showing CS and US for conditioning day 2]

**C** Conditioning Day 6

![Graph showing CS and US for conditioning day 6]

**D** Relative Conditioned Response

![Graph showing relative conditioned response]

**E** UR peak amplitude vs UR peak velocity

![Scatter plots showing UR peak amplitude vs UR peak velocity]

Figure 7. EBC in CT-short^Q27PC and CT-longQ27^PC mice. A, Schematic diagram of the experimental procedure used for analyzing the EBC in mice. Left, Animals underwent a 12-d behavioral assay with three sessions of habituation, six sessions of conditional learning, and three sessions of extinction with one session per day. The US was a 30 ms mild corneal air puff; the CS was a 380-ms-long auditory tone. Right, For habituation, a US was elicited randomly before the CS. For conditioning, the US was elicited at the end of the CS. For extinction the US (Figure legend continues.)
Schonewille et al., 2006). Additionally, PCs expressing CT-short and CT-longQ27 showed a reduced mean duration of the tonic state from 173.6 ± 24.5 s for CT-short and 49.7 ± 2.3 s for CT-longQ27 in the trimodal state (Fig. 6C). The drastic reduction in the mean duration of the tonic phase in CT-longQ27-expressing PCs was accompanied by an increase in the maximal firing rate (Fig. 6C). An increase in trimodal cycle time and maximal firing frequency has also been observed when SK channels, i.e., Ca^2+–activated K+ channels, are blocked in PCs (Womack and Khodakhah, 2003). Therefore, we compared the trimodal firing pattern and firing frequency between CT-longQ27-expressing PCs and PCs in which SK channels are blocked. Indeed, SK channel block leads to a similar, but more pronounced, change in trimodal firing as observed for CT-longQ27-expressing PCs (Fig. 6C), suggesting that CT-longQ27 may interfere with the interaction between P/Q-type channels and SK channels (Womack et al., 2004; Indriati et al., 2013).

Postnatal expression of CT-longQ27PC, but not CT-shortPC, impairs EBC and cerebellar plasticity at the PF–PC synapse. EBC is a form of associative learning that requires precise information processing within the cerebellar cortex (Kim and Thompson, 1997; De Zeeuw et al., 2011). For example, lesions in defined cerebellar areas in human patients lead to deficits in timing, acquisition, and extinction of eyeblink responses (Gerwig et al., 2007), and changes in synaptic plasticity of the cerebellar cortex impairs EBC (Koekkoek et al., 2003; Schonewille et al., 2010, 2011). To examine changes in EBC caused by the expression of CTs, we analyzed EBC responses in CT-shortPC, CT-longQ27PC, and control (fl-CT-short, fl-CT-longQ27) mice using an MDMT (Koekkoek et al., 2002). Closing of the eyelid is induced by a corneal air puff (the US) and is associated timely during a 6 d training session with a CS, i.e., an auditory tone (Fig. 7A). As shown in Figure 7B, we found that in CT-longQ27PC mice EBC responses are suppressed almost completely during the 6 d training period, whereas EBC responses in CT-shortPC mice are comparable with control mice.

Because changes in synaptic plasticity at the PF–PC synapse may impair EBC responses (Koekkoek et al., 2003; Schonewille et al., 2010, 2011), we analyzed whether LTD and LTP were affected in cerebellar slices in the presence or absence of CT peptides using whole-cell patch-clamp recordings after AAV2 expression. Indeed, we found that virally expressed CT-longQ27, but not CT-short, suppresses both LTD (Fig. 8A,B) and LTP (Fig. 8C,D) at the PF–PC synapse.

Discussion

PolyQ extension within the P/Q-type channel CT has only minor effects on gating and Ca^2+ permeability (Saegusa et al., 2007; Watase et al., 2008). Thus, a change in the biophysical properties of the channel may have only a small contribution to the disease phenotypes (Orr, 2012). After detection of the P/Q-type channel CTS in PCs from SCA6 patients (Ishiguro et al., 2010), it has been suggested that a CT peptide fragment might be the cause of the disease. The CT peptide fragments seem to be produced by either proteolytic processing (Kubodera et al., 2003; Kordasiewicz et al., 2006) or translation from a bicistronic mRNA (Du et al., 2013). Indeed, because they are localized in the cytoplasm and nuclei of PCs, they may interfere with the function of proteins in PCs and/or change their gene expression. Regardless of whether CT-longQ(disease) is produced via proteolytic processing or via its own transcriptional start site, CT-longQ(disease), has been shown to be toxic to cells in cultures and to cause cell death for PCs in vivo (Kordasiewicz et al., 2006; Du et al., 2013; Takahashi et al., 2013). Therefore, we were interested in establishing an animal model and viral techniques to analyze the physiological function of the two CT fragments, i.e., CT-short and CT-longQ27 for disease phenotypes. These two CT variants are present in the human cerebellum by the identified P/Q-type channel splice variants. Here, 65% of P/Q-type channel mRNA consists of the long form, whereas 35% consist of the short form (Soong et al., 2002). The ratio of the long (toxic) form of Ca2.1 mRNA compared with total Ca2.1 mRNA in cerebellar PCs of SCA6 patients (0.98 ± 0.06) is increased compared with control individuals (0.51 ± 0.05), suggesting a higher concentration of CT-longQ(disease) peptide fragments in the diseased state (Tsunemi et al., 2008; Tsu et al., 2011).

In our study, we found that long-term expression of CT-longQ27(disease) in cerebellar PCs leads to ataxia, associated with changes in PC firing and PC degeneration. These phenotypes are similar to phenotypes observed in other SCA6 knock-in mouse models, in which the length of the polyQ stretch had to be increased to 84 and 118 polyQs to cause disease symptoms (Watase et al., 2008; Unno et al., 2012). Compared with other mouse models, we now describe that CT-longQ27(disease), but not CT-short, interferes with synaptic plasticity at the PF–PC synapse and causes the loss of EBC, an associative motor learning paradigm. The EBC is a withdrawal response to protect the eye from noxious stimuli and is mediated by a very defined cerebellar circuit with the requirement of functional PC plasticity (Kim and Thompson, 1997; De Zeeuw and Yeo, 2005; Gao et al., 2012). Indeed, loss of LTP at PF–PC synapses may be associated with changes in motor learning (Schonewille et al., 2010, 2011; Galliano et al., 2013). Conversely, LTD might also contribute (Kishimoto et al., 2001, 2002; Koekkoek et al., 2003), but this form of plasticity is probably less critical for EBC (Schonewille et al., 2011). Human subjects with cerebellar lesions and brain imaging studies also reveal an important role of the human cerebellum in EBC (Gerwig et al., 2007). Thus, the important aspect of this finding is that deficits in EBC in humans may also be associated with SCA6 as observed in our animal model and serve as a diagnostic clinical tool for early detection of SCA6 and other diseases associated with cerebellar dysfunction.

Functional consequences of CT peptides expressed in PCs before cell death

Because CT-long is localized to the nucleus and the cytoplasm and has been identified as a transcription factor and a proteolytic product, changes in gene transcription and/or dominant-negative effects of the CT peptide might lead to functional alteration in PCs before cell death. In our experiments, the changes in synaptic plasticity and firing of PCs can be explained by
dominant-negative effects of the CT domain on P/Q-type channel interacting proteins. The CT domain of the P/Q-type Ca$^{2+}$ channel consists of binding domains for proteins involved in modulation, targeting, and structural organization of the channel complex in subcellular domains (Herlitze et al., 2003; Spafford et al., 2003). Therefore, a CT peptide expressed in the soma, axon, or dendrites of PCs would be expected to induce dominant-negative effects on these parameters before PC degeneration. For example, P/Q-type channels are tightly coupled and colocalized to Ca$^{2+}$-activated K$^+$ channels in PCs (Womack et al., 2004; Indriati et al., 2013). The functional interaction between these channels determines the mode and firing frequency of PCs.
Therefore, disruption of this interaction causes changes in the firing patterns of PCs similar to the alteration we observed with CT-longQ27. Indeed, chlorzoxazone, an activator of Ca\(^{2+}\)-activated K\(^+\) channels restores the precision of firing of PCs and improves motor behavior in both tottering and Cacna1a(-/-) mice, which are both ataxic P/Q-type channel mouse mutants (Alvina and Khodakhah, 2010). Furthermore, Berkelfeld et al. (2006) demonstrated the formation of macromolecular complexes with Ca\(^{2+}\)-activated K\(^+\) channels and P/Q-type channels in rat brains. One can postulate that the CT peptide may be disrupting the formation of these Ca\(^{2+}\)-activated K\(^+\) channel–P/Q-type channel complexes by blocking critical binding sites on the Ca\(^{2+}\)-activated K\(^+\) channels or P/Q-type channels, thus leading to a disruption of these “Ca\(^{2+}\) nanodomains” necessary for K\(^+\) channel activation. In addition, the CT binds proteins involved in membrane transport and synapse assembly, such as the Ca\(_{\alpha\beta}\) subunit (Herlitze et al., 2003). The multiple interaction sites within the CT domain of the P/Q-type Ca\(^{2+}\) channels make it a critical domain for the regulation of channel targeting to the membrane and synapses. Decreased clustering and current density of the channels have been demonstrated by the use of peptides that out-compete the interactions with Mint-1, CASK, and tctex-1 (Spaﬀord et al., 2003; Lai et al., 2005). The CT also contains binding sites for proteins involved in synaptic transmitter release, including Rim (Ribino et al., 2002; Shapira et al., 2003) and other proteins associated with Piccolo–Bassoon transport vesicles (Shapira et al., 2003). Because these proteins are involved in preforming synapses, CT peptides may interact with the clustering process required for synaptic targeting. Furthermore, the P/Q-type channel CT binds to proteins involved in synaptic plasticity, such as Ca\(^{2+}\)/calmodulin (Lee et al., 1999) and Ca\(^{2+}\) influx through dendritic P/Q-type channel, and is one of the major sources of dendritic Ca\(^{2+}\) necessary to induce LTD and LTP (Han et al., 2007). Therefore, changes in P/Q-type channel function in PCs in dendritic domains will most likely interfere with the expression of PC plasticity.

Dominant-negative effects on P/Q-type channel interacting proteins can be expected for both the CT-long and CT-short, because the described interacting proteins bind to both isoforms. However, we observed mainly behavioral and physiological effects for CT-longQ27 and not CT-short expressing PCs. The differences are most likely attributable to the different subcellular localization of the CTs within PCs. CT-short localizes mainly in the nucleus and at high expression levels diffuses throughout the cytoplasm. In contrast, CT-longQ27 is localized to cytoplasmic aggregations compatible with synaptic sites, in which it may compete with interacting proteins in subcellular domains.

In summary, we established a new SCA6 mouse model that will serve to understand the developmental onset of the disease and can be used for therapeutic drug testing. This mouse model develops late-onset ataxia associated with changes in PC firing and PC degeneration as observed in SCA6 patients. In addition, the mouse model identiﬁed the loss of EBC as a new diagnostic strategy for early detection of SCA6 in humans that is associated with changes in cerebellar plasticity.

References

Mark, Krause et al. Effects of P/Q-Type Channel Underlying SCA6

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