Zebrafish Cerebellum as a Genetic Model for Spinocerebellar ataxia

The cerebellum is an essential brain compartment that controls motor coordination and learning among other functions to allow us to move smoothly and skilfully. In addition, mounting evidence involves the cerebellum in higher functions such as cognition and processing of emotions.1 The cerebellum is found in all jawed vertebrates, gnathostomes,2 and is highly conserved in its cytoarchitecture, connectivity, and function as well as the underlying cellular and genetic mechanisms of differentiation and homeostasis. In zebrafish, the cerebellum develops, differentiates, and reaches functional maturity during embryonic and larval stages during which zebrafish are almost transparent, making them a powerful tool for bioimaging. Moreover, as zebrafish are genetically tractable, they allow for combining molecular genetic studies with in vivo imaging of cell biological and physiological processes making use of the numerous available fluorescent proteins and biosensors.3,4 As zebrafish are becoming an increasingly popular model in the biomedical field, the stage is set to expand cerebellar research in zebrafish to causal investigations of human cerebellar diseases. Spinocerebellar ataxias (SCAs) represent a group of inherited neurodegenerative diseases of the cerebellum that commonly lead to motor discoordination due to affected cerebellar Purkinje cells (PCs). Towards modeling SCAs in zebrafish, we identified a small-sized PC-specific regulatory element that was used to develop coexpression vectors with tunable expression strength. These vectors allow for in vivo imaging of SCA-affected PCs by high-resolution fluorescence imaging. Next, zebrafish with SCA type 13 (SCA13) transgene expression were established, revealing that SCA13-induced cell-autonomous PC degeneration results in eye movement deficits. Thus, SCA13 zebrafish mimic the neuropathology of an SCA-affected brain as well as the involved loss of motor control and hence provide a powerful approach to unravel SCA13-induced cell biological pathogenic and cytotoxic mechanisms.

SCAs are inherited in an autosomal dominant manner, which prompted us to establish a transgenic zebrafish model affected by a gain of toxicity of a pathogenic SCA allele. As cell type, we chose Purkinje cells (PCs) as these are most commonly affected by SCAs and represent the central neuronal population and sole output of the cerebellum. The PCs integrate external sensory stimuli and motor actions, which are processed into output signals relayed onto efferent neurons usually of the motor system, explaining the difficulties in motor control upon PC degeneration.

PCs in zebrafish develop fast and emerge already at 56 hours postfertilization (hpf) with increasing cells numbers until 7 days postfertilization (dpf).6 By this stage, PC production reaches a plateau with a population size of about 420 PCs,6 which is far less than the over 200 000 PCs estimated to populate the mouse cerebellum. At 7 dpf, the cerebellar circuitry in zebrafish is established and PCs like in mammals receive synaptic input from parallel fibres – the granule cell axons – and climbing fibres – axonal connections from the inferior olive. The PC axons form collaterals to other PCs and project either directly out of the cerebellum to terminate on neurons of the octaval nuclei corresponding to vestibular neurons, or they connect to nearby eurydendroid cells, which represent the equivalents to deep cerebellar nuclei neurons in mammals3,7 (Figure 1). Also, the electrophysiological properties of PCs similar to their mammalian counterparts are established at 7 dpf, allowing them to control visual stimuli driven eye or swimming movements.
movements like the optokinetic response (OKR) or the optomotor reflex (OMR) and being involved in motor learning. Optogenetic silencing of PCs consequently impairs OKR and OMR reminiscent of loss of motor control in SCA patients. These findings support the view that zebrafish PCs are well suited to model cerebellar neurodegenerative diseases, as they will allow for continuously monitoring and thus for interconnecting pathogenic molecular processes with anatomical, physiological, and eventual behavioural deficits that mimic clinical signs of SCAs.

Establishment of a PC-Specific Coexpression System in Zebrafish

For this approach, a prerequisite is the possibility to genetically target PCs in a cell-type-specific manner. We therefore first invested into characterizing a regulatory element upstream of zebrafish carbonic anhydrase VIII (ca8), which is predominantly expressed in differentiating and mature PCs in zebrafish. By transient and stable transgenesis, we first revealed that a 7.54-kb region upstream of the ca8 transcriptional start site was able to drive enhanced green fluorescent protein (EGFP) expression in zebrafish PCs besides expression in a few other tissues such as in the notochord. Subsequently, we narrowed it down to a PC exclusive regulatory element of a size of 258 bp, which we named ca8-derived Purkinje-cell-specific enhancer element (cpce) (Figure 2A to C). Intriguingly, this regulatory element from zebrafish also mediates PC-specific expression in organotypic slice cultures of the mouse cerebellum (Figure 2D), suggesting that regulatory mechanisms of PC-specific expression are conserved among teleosts and mammals. The small size of the cpce further allowed for tandem orientation of several copies of this element, resulting in PC-specific bidirectional expression vectors of different expression strengths when 2, 4, or 8 copies of the cpce were used. The SCA-causing pathogenic alleles can therefore be coexpressed with fluorescent protein reporters at varying concentrations to allow for investigating dose-dependent effects and monitoring disease progression by in vivo time-lapse imaging. Copy numbers of 4 and more cpce elements resulted in increasing ectopic transgene expression in tectal and retinal cells, which could be restricted to PCs again by an miRNA-mediated gene silencing strategy. miRNA181a is predominantly expressed in tectal and retinal cells. Incorporating the miRNA181a target site into the 3′UTR of both transgenes eliminated successfully their expression outside of the cerebellum, while maintaining it at high levels in PCs. This strategy can be easily adopted for other regulatory elements with unwanted expression profiles. Finally, we established a series of PC-specific expression vectors containing 1, 2, and 4 cpce elements, respectively, in bidirectional orientation with different multiple cloning sites and miRNA181a target sites inserted into the corresponding 3′UTR sequences followed by polyA sequences where necessary (4× cpce) (Figure 2E). This vector set will enable fast, dose-independent, and comparable PC-specific disease modelling in zebrafish accessible for disease monitoring by fluorescent in vivo imaging for SCAs, but also for other diseases affecting PCs such as various types of dystonia, ataxia telangiec-tasia, Niemann-Pick disease, and autism among others, when making use of PC-specific expression of recombinases, transcriptional activators, and repressors for example.

Neurodegenerative Disease Modelling for Spinocerebellar Ataxia Type 13 (SCA13) in Zebrafish

Using these vectors, we set out to establish a genetic model of SCA13 in zebrafish. This neurological disease is inherited in
an autosomal dominant manner leading to cerebellar atrophy.\textsuperscript{8} SCA13 is caused by mutations in the \textit{KCNC3} allele encoding the voltage-gated potassium channel \textit{KCNC3/Kcnc3}. Its rodent homologue is expressed in neurons with high-frequency firing rate with prominent expression in cerebellar PCs.\textsuperscript{9} It is therefore likely that cerebellar atrophy in SCA13 patients is caused by degenerating PCs as primarily affected neuronal cell type, yet there is currently no in vivo model of SCA13 established, which shows clear signs of neuronal degeneration followed by loss of motor control. Hence, causal analysis of SCA13 is hampered. We initiated SCA13 modelling in zebrafish by analysing the spatiotemporal expression of the zebrafish \textit{ KCNC3 } homologues \textit{kcnca} and \textit{kncb}. Although \textit{kncb} was strongly expressed in larval PCs, \textit{kncb} expression was barely detected in this cell type. Human, rodent, and zebrafish \textit{kncb} alleles generate a number of splice isoforms terminating in different C-terminal domains of the potassium channel. Splice-isoform-specific mRNA in situ hybridization as well as fluorescence-assisted cell sorting (FACS)-mediated single PC reverse transcription polymerase chain reaction (RT-PCR) revealed that \textit{kncb-}X12 is expressed at highest abundance in zebrafish PCs. This splice isoform contains the shortest C-terminus of all \textit{kncb} splice isoforms. We therefore generated a \textit{kncb-}X12 \textit{R335H} zebrafish mutant allele (named hereafter \textit{kncb} \textit{R335H}) (Figure 2F) mimicking a mid-age onset \textit{SCA13} variant causing progressive cerebellar atrophy in humans.\textsuperscript{10}

By introducing either zebrafish wild type \textit{kncb} (\textit{kncb} \textit{WT}) or \textit{kncb} \textit{R335H} into PC-specific expression vectors coexpressing two fluorescent reporters, membrane-targeted \textit{TagRFP-T} and nuclear-localized EGFP (Figure 2G), transgenic zebrafish were generated by microinjection into one-cell stage embryos. This allowed for monitoring transgene expressing PCs in the differentiating cerebellum using in vivo confocal microscopy. \textit{kncb} \textit{WT}-expressing PCs differentiated indistinguishably from wild-type PCs, increased in number, and developed an elaborated dendritic tree at 7 dpf. In contrast, \textit{kncb} \textit{R335H}-expressing PCs displayed progressively increasing signs of axonal and dendritic fragmentation and diminished in cell numbers indicative of progressive neuronal degeneration, which was confirmed by PC counts making use of nuclear fluorescent protein expression (Figure 2H to J). Moreover, when analysed for PC-mediated behaviour, larvae with PC-specific \textit{kncb} \textit{WT} expression displayed a normal performance of the OKR, a

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**Figure 2.** Zebrafish disease modelling of SCA13 using a Purkinje-cell-specific coexpression system. CCe indicates corpus cerebelli; Val, lateral division of the valuza cerebelli; EGFP, enhanced green fluorescent protein; PC, Purkinje cell.

(A) Schematic map of ca8-derived Purkinje-cell-specific enhancer (cpce) and drawing of construct carrying the cpce: E1b promoter, driving EGFP reporter expression.

(B) EGFP expression is restricted to cerebellar PCs (green) in 4 dpf stable transgenic larval fish \textit{Tg(cpce-E1b:GFP)bz13} carrying the reporter transgene shown in (A).

(C) Schematic drawing of the zebrafish \textit{Kcnc3a} subunit reveals the S1-S6 transmembrane segments (4× sites (4 mir181aT) immediately upstream to both polyA sequences to eliminate ectopic expressions outside of cerebellum. Two multiple cloning sites (MCS1 and MCS2) can be used for the insertion of different genes of interest.

(F) Schematic drawing of the zebrafish \textit{Kcnc3a} subunit reveals the S1-S6 transmembrane segments homologous to those in mammals. Positively charged arginine residues (indicated as +) located in the S4 segment, essential for voltage sensing, are also conserved in zebrafish \textit{Kcnc3a}. The third arginine in the S4 segment is substituted by histidine (R335H), which mimics \textit{SCA13R420H} plus homologous to those in mammals.

(G) Schematic drawing of a PC-specific transgene to express \textit{Kcnc3awt} and \textit{Kcnc3aR335H} generated by alternative splicing is indicated in blue.

(H) Each image show PCs expressing \textit{Kcnc3awt} (upper, or lower rows, respectively), monitored at 4, 7, 11 dpf zebrafish. Cerebellar PCs expressing \textit{Kcnc3aR335H} (I), whereas those expressing \textit{Kcnc3awt} (J) show degenerative changes with fragmented red labelled puncta from dendritic or axonal structures (J).
reflexive eye movement driven by visual stimuli. Consistent with progressive PC degeneration larvae with kcn3aR335H expressing PCs instead were impaired in proper OKR performance, which is reminiscent of oculomotor disturbance, a clinical hallmark of SCA patients. Taken together, our results strongly suggest that KCNC3R420H in analogy to its zebrafish homologue kcn3aR335H induces PC degeneration in a cell-autonomous manner, which appears to be the main cause for the atrophy of the cerebellum leading to ataxic behavior observed in patients carrying this mutated pathogenic allele.

### Molecular Mechanism Underlying SCA13

**Pathogenic Mutants induced PC Degeneration**

Although the main neuropathological features such as cerebellar atrophy and uncoordinated motor behaviour are shared among SCA13 patients, the clinical symptoms of SCA13 are somewhat heterogeneous as well as the age of disease onset and the course of disease progression. This is explained by allelic heterogeneity as currently more than 10 KCNC3-mutated alleles have been reported so far, ranging from different missense mutations to a C-terminal deletion in the encoded potassium channel. This suggests that the cause of SCA13 is also somewhat heterogeneous influencing the strength of cytotoxic signals and mechanisms in PCs. Electrophysiological studies support this hypothesis, as they indicate that different SCA13 alleles could affect the excitability of neurons to a different extent linked to the disease phenotype.

For example, the adult onset KCNC3R420H mutant allele encodes a non-functional channel subunit mimicked by its zebrafish counterpart kcn3aR335H. The heterotetrameric wild-type/R420H mutant channels do not result in altering the kinetic behaviour as Kv3.3 channels. This mutant subunit as well as another SCA13 variant R423H, however, mediate a dominant negative effect likely by being trapped along the secretory pathway and mostly in the Golgi together with wild-type subunits – is misrouted. Because overexpression of Kv3.1-Kv3.4) depends on unusually positively shifted potassium channels belonging to the family of KCNC/Kv3 isoform expressing neurons. This argues for alternative cell biological mechanisms mediating cytotoxicity in SCA13-affected PCs either acting alone or together with the altered electrophysiological properties.

Further insight was provided by recent cell biological and biochemical studies showing that KCNC3R420H and KCNC3R423H are aberrantly glycosylated. In addition, these channel subunits are inefficiently transported to the plasma membrane and are retained to a large extent in the Golgi apparatus. At least in cells expressing KCNC3R420H, the epidermal growth factor receptor (EGFR) – in addition to KCNC3 wild-type subunits – is misrouted. Because overexpression of EGFR in Drosophila eyes could rescue KCNC3R423H induced eye malformation, retention of crucial signalling molecules in the secretory pathway could represent an additional cell biological mechanism of cytotoxicity in SCA13-affected neurons. It would be of interest if KCNC3R423H affects intracellular membrane transport of EGFR as well, to reveal if this is an allele-specific or common cell biological defect in SCA13-affected cells. The PC-specific coexpression system established by us will allow for testing the involvement of such mis-trafficked molecules in PC degeneration. It will further allow for discriminating whether mis-trafficking alone or in conjunction with altered neuronal excitability leads to progressive cytotoxicity (Figure 3). In addition, recent evidence involved another SCA13 allele KCNC3G592R in the regulation of actin dynamics in axonal growth cones. As this research was performed on cultured cells, in vivo confirmation could be obtained easily in
zebrafish, followed by comparison to implications of KCNC3R420H and KCNC3R423H on actin regulation.

Clearly, the molecular mechanisms causing the SCA13 disease are far from being understood, but distinct research avenues have been opened that await unravelling. Our genetic tools and models for deciphering cell biological and physiological causes of PC degeneration in a cell-autonomous manner directly in vivo may make use of coexpression of fluorescent reporters and biosensors will eventually enable clarification of the pathogenic mechanisms underlying PC degeneration in and reporting of spinocerebellar ataxia type 13 (SCA13). Moreover, modelling SCA13 in zebrafish with the possibility to monitor behaviour from single cells to the entire animal offers powerful approaches for in vivo screening and validation of compounds towards counteracting pathogenic molecular and physiological disease mechanisms. Thus, this little fish could make a big splash.

**Author Contributions**

KN and RWK wrote this commentary. KN and AD prepared figures. All authors edited the manuscript.

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**Figure 3.** Schematic representation of SCA13-induced PC degeneration. KCNC3R420H and KCNC3R423H variants show dominant negative effects on WT KCNC3 by being retained in the Golgi apparatus together with WT channel subunits, thereby reducing PC excitability. A possible PC death-associated effector activated by reduced PC excitability could be exerted via increased intracellular Ca\(^{2+}\) levels likely caused by broadening action potentials excessively activating P/Q-type voltage-gated Ca\(^{2+}\) channels (VGCC). In addition, the Golgi-retained KCNC3R423H mutant was shown to impair EGFR trafficking, inducing loss of EGFR trophic support, which may be needed for PC survival. EGFR indicates epidermal growth factor receptors; PC, Purkinje cell.

**Figure 5.** Schematic representation of SCA13-induced PC degeneration. KCNC3R420H and KCNC3R423H variants show dominant negative effects on WT KCNC3 by being retained in the Golgi apparatus together with WT channel subunits, thereby reducing PC excitability. A possible PC death-associated effector activated by reduced PC excitability could be exerted via increased intracellular Ca\(^{2+}\) levels likely caused by broadening action potentials excessively activating P/Q-type voltage-gated Ca\(^{2+}\) channels (VGCC). In addition, the Golgi-retained KCNC3R423H mutant was shown to impair EGFR trafficking, inducing loss of EGFR trophic support, which may be needed for PC survival. EGFR indicates epidermal growth factor receptors; PC, Purkinje cell.

**Supplementary Figure 1.** Schematic representation of SCA13-induced PC degeneration. KCNC3R420H and KCNC3R423H variants show dominant negative effects on WT KCNC3 by being retained in the Golgi apparatus together with WT channel subunits, thereby reducing PC excitability. A possible PC death-associated effector activated by reduced PC excitability could be exerted via increased intracellular Ca\(^{2+}\) levels likely caused by broadening action potentials excessively activating P/Q-type voltage-gated Ca\(^{2+}\) channels (VGCC). In addition, the Golgi-retained KCNC3R423H mutant was shown to impair EGFR trafficking, inducing loss of EGFR trophic support, which may be needed for PC survival. EGFR indicates epidermal growth factor receptors; PC, Purkinje cell.