Characterization of *Xenopus laevis* Guanine Deaminase Reveals New Insights for its Expression and Function in the Embryonic Kidney

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**Background:** The mammalian guanine deaminase (GDA), called cypin, is important for proper neural development, by regulating dendritic arborization through modulation of microtubule (MT) dynamics. Additionally, cypin can promote MT assembly in vitro. However, it has never been tested whether cypin (or other GDA orthologs) binds to MTs or modulates MT dynamics. Here, we address these questions and characterize *Xenopus laevis* GDA (Gda) for the first time during embryonic development. **Results:** We find that exogenously expressed human cypin and Gda both display a cytosolic distribution in primary embryonic cells. Furthermore, while expression of human cypin can promote MT polymerization, *Xenopus* Gda has no effect. Additionally, we find that the tubulin-binding collapsin response mediator protein (CRMP) homology domain is only partially conserved between cypin and Gda. This likely explains the divergence in function, as we discovered that the cypin region containing the CRMP homology and PDZ-binding domain is necessary for regulating MT dynamics. Finally, we observed that gda is strongly expressed in the kidneys during late embryonic development, although it does not appear to be critical for kidney development. **Conclusions:** Together, these results suggest that GDA has diverged in function between mammals and amphibians, and that mammalian GDA plays an indirect role in regulating MT dynamics. *Developmental Dynamics* 2019. © 2019 Wiley Periodicals, Inc.

**Key words:** cypin; GDA; microtubule dynamics; development

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**Introduction**

The proper establishment of neuronal morphology is essential for the correct communication between neurons, and the cytoskeleton plays a key role in determining neuronal morphology during development. In particular, microtubules (MTs) orchestrate formation of neuronal polarity and ensure proper axon and dendrite morphology. MT dynamics are widely regulated by proteins that bind to the plus-end of the MT, called plus-end tracking proteins (+TIPS), in addition to other types of MT-binding proteins (Bearce et al., 2015; Cammarata et al., 2016; Lasser et al., 2018). While many studies have begun to investigate MT plus-end dynamics in the developing nervous system, it is clear that much more information is needed to reveal a comprehensive picture for how MT dynamics regulate neuronal development.

One important MT regulator in the developing nervous system is the mammalian guanine deaminase (GDA), called cypin (cytosolic PSD-95 interactor) (Akum et al., 2004). Cypin has been linked to autism spectrum disorders and has been described as a risk biomarker for these disorders (Braunschweig et al., 2013). In addition, cypin levels increase in rat striatal neurons in response to nigro-striatal degeneration, which models the degeneration of neurons in Parkinson’s disease (Fuller et al., 2014). Multiple neurodevelopmental disorders are associated with defects in axon guidance and dendritogenesis. Furthermore, it has been shown that cypin plays a key role in establishing neuronal morphology by antagonizing PSD-95 function and increasing dendrite branching (Firestein et al., 1999; Riefler et al., 2003; Akum et al., 2004). Cypin contains a domain with homology to the collapsin response mediator protein (CRMP), which binds tubulin and has a suggested role in regulating MT assembly in the growth cone (Schmidt and Strittmatter, 2007). In addition, cypin can bind tubulin heterodimers and regulate MT assembly in vitro (Akum et al., 2004). Moreover, the binding partner of cypin, PSD-95, binds to the known +TIP, EB3, in neurons and hence regulates MT dynamics (Sweet et al., 2011). However, while cypin has previously been proposed to regulate MT dynamics, it has not yet been determined whether...
cypin can actually localize to MT plus-ends or alter MT dynamics in living cells.

Multiple +TIPs have been described and characterized using *Xenopus laevis* (Lee et al., 2004; Lowery et al., 2013; Marx et al., 2013; Nwagbara et al., 2014; Lucaj et al., 2015; Rutherford et al., 2016; Erdogan et al., 2017). *Xenopus* embryos can be easily manipulated, and their primary embryonic neural cells are facile to obtain, culture, and image, as they display large growth cones (10 microns or more), which are useful for imaging cytoskeletal dynamics (Erdogan et al., 2016; Slater et al., 2017). Live cell imaging is particularly important, as some +TIPs only bind to growing MTs and, thus, their localization dynamics cannot be visualized using immunohistochemistry of fixed cells (Nwagbara et al., 2014).

In this work, we investigated whether human GDA (cypin) and/or *Xenopus laevis* GDA (Gda) can localize to MT plus-ends, perhaps functioning as a +TIP, and tested whether it has a role in regulating MT dynamics. Additionally, we examined *gda* expression in *Xenopus laevis* during embryonic development, as its expression and function had not been well characterized. We visualized exogenously-expressed cypin and Gda for the first time in living primary embryonic neural growth cones using fluorescent confocal microscopy and found that neither of them colocalize with MT plus-ends, suggesting that GDA does not function as a +TIP. Despite the lack of MT binding, we found that expression of cypin, but not Gda, promotes MT assembly in neuronal growth cones. Moreover, we found that the CRMP homology domain, which has been described as important for binding tubulin and regulating MT assembly, is not fully conserved in *Xenopus*. By generating a chimera containing the N-terminal region from Gda plus the cypin domain containing the CRMP homology and PDZ-binding domain, we found that this cypin region is necessary for regulating MT dynamics. Finally, using whole-mount in situ hybridization, we observed that *gda* is highly expressed in the kidneys and may be important for proper kidney function. Our results suggest that the function of GDA to regulate MT dynamics is not conserved in *Xenopus*.

### Results

**CRMP Homology Domain is not Conserved Between Human and *Xenopus* GDA**

While a previous phylogeny study reported that the guanine deaminase (GDA) DNA sequence has been conserved from prokaryotes to higher eukaryotes (Fernandez et al., 2009), this early study did not examine the entire amino acid sequence nor did it look specifically at the CRMP homology domain. *X. laevis*, *X. tropicalis*, *Danio rerio*, *Gallus*, *Homo sapiens*, *Mus musculus*, and *Rattus norvegicus* form part of the same related clusters of GDA sequences (Fernandez et al., 2009). Here, we compared the GDA protein sequences of all the aforementioned species. *X. laevis* is an allotetraploid organism and has two GDA homeologs, S and L; thus, we considered both homeologs for protein sequence comparison. We found the following sequence homologies of each organism with human cypin:
**X. laevis** GDA S-homeolog 60%; **X. laevis** GDA L-homeolog 61%; **X. tropicalis** 61%; zebrafish 59%; chicken 72%; rat 91%; and mouse 91% (Fig. 1).

In addition, while all GDAs were previously described to have well-conserved zinc-binding domains, the CRMP homology and PDZ-binding domains are not completely conserved. *Gallus*, *Homo sapiens*, *Mus musculus*, and *Rattus norvegicus* all have a PDZ-binding domain, which is absent from **X. laevis**, **X. tropicalis**, and *Danio rerio* (Fig. 1). Moreover, even though the CRMP homology domain was present in all the organisms, it is not well conserved. While chicken, rat, and mouse have strong conservation with human (82%, 91%, 91%, respectively), the CRMP homology domains in **X. laevis** GDA S and L, **X. tropicalis**, and zebrafish are not as similar (60%, 53%, 53%, 58% conservation with human GDA, respectively) (Fig. 1). Notably, neither the PDZ-binding domain nor the CRMP domains are conserved between amniotes and amphibians/fish.

**GDA Has a Cytosolic Distribution and Does not Accumulate at MT Plus-Ends**

To explore the possibility that GDA may function as a +TIP and thereby bind to the plus-ends of MTs to regulate MT dynamics...
(Bearce et al., 2015), we first examined whether human (cypin) and/or *X. laevis* (Gda) GDA localize to MT plus-ends or not. We co-injected mRNAs coding for green fluorescent protein (GFP)-tagged N- or C-terminus cypin and Gda along with mKate2-MACF43, a known marker of growing MTs that localizes to the MT plus-ends (Honnappa et al., 2009), into *X. laevis* embryos at the 2-cell stage. We prepared spinal cord explants from stages 20–24 *Xenopus* embryos, and we used live imaging to visualize the exogenously-expressed GDAs. We found that neither cypin nor Gda localized to the MT plus-end in neuronal growth cones (Fig. 2A), nor in neural-derived mesenchymal cells (Fig. 2B), as GDAs did not colocalize with mKate2-MACF43. Rather, in all cases, GDA was diffused throughout the cytosol and did not appear to localize to MTs at all, consistent with what has been previously described for cypin localization (Firestein et al., 1999; Akum et al., 2004; Chen and Firestein, 2007; Sweet et al., 2011). These data suggest that neither human nor *Xenopus* GDA accumulate on MTs plus-ends, and thus, GDA does not act as a +TIP in *Xenopus* embryonic cells.

**Exogenous Expression of Cypin, but not Gda, Promotes MT Polymerization in Living Cells**

Considering that some proteins regulate MT dynamics through indirect mechanisms, for example, by binding tubulin heterodimers or other proteins that bind to MTs (Andersen, 2000; Akhmanova and Steinmetz, 2008), we sought to determine whether overexpression of cypin or Gda (Fig. 2C) could affect MT dynamics indirectly in living cells. We measured MT behavior in cultured primary embryonic neuronal growth cones obtained from *Xenopus* stages 20–24, by quantifying the MT growth track velocity, lifetime, and length of mKate2-MACF43 comets after GDA overexpression (Supplementary Movie S1, which is available online). We observed that axonal growth cones overexpressing cypin showed an 11% increase in MT growth track velocity and a 17% increase in MT growth track length compared with those of control cells, while no change was observed in mKate2-MACF43 comet lifetime (Fig. 2D–F). Similar results were observed for both N- and C-terminal tagged GFP-cypin (data not shown). However,
we observed no changes in MT dynamics in cells overexpressing Gda compared with control cells (Fig. 2D–F). These data show that cypin overexpression increases MT polymerization rate in living Xenopus embryonic cells, and that human and Xenopus GDAs are different in their ability to modulate MT dynamics.

Additionally, considering the differential regulation of MT dynamics by cypin and Gda, and the striking differences between the cypin and Gda CRMP homology and PDZ-binding domain, we decided to evaluate if this cypin region is responsible for the effect on MT dynamics. We generated a chimera GDA containing the N-terminal half from Gda and the CRMP homology plus PDZ-binding domain from cypin (Fig. 2C) and examined the effect of the chimera GDA on MT dynamics. We found that the chimera GDA was able to increase MT growth track velocity and MT growth track length compared with those of control, similar to cypin (Fig. 2D–F). Our results suggest that the cypin region containing both CRMP homology and PDZ-binding domains plays an important role in promoting MT polymerization.

**Xenopus GDA**

Xenopus GDA is Strongly Expressed in the Embryonic Kidney and Thus may Play a Role in Kidney Function

Given that our results suggested that Xenopus GDA did not function as a promoter of MT polymerization, we wondered if it was still expressed in the developing brain, like cypin. To characterize *gda* expression during Xenopus embryonic development, we used reverse transcription-polymerase chain reaction (RT-PCR) and whole-mount in situ hybridization at different stages of development. First, we performed RT-PCR using cDNA from embryos at various stages including two-cell, blastula, neurula, 3 and 6 days post fertilization (dpf), as well as cDNA from the adult brain. We found that strong *gda* expression was not apparent until 3–6 dpf.
and persisted, although to a lesser extent, through adulthood within the brain (Fig. 3A). Whole-mount in situ hybridization showed no staining in early stages including two-cell, blastula, and 1 dpf, confirming the RT-PCR results (Fig. 3B). However, specific accumulation of gda expression in two bilateral regions behind the embryonic eye became apparent at 3 dpf and remained at a high level to 6 dpf (Fig. 3B). Following the stages of *Xenopus laevis* development (Nieuwkoop and Faber, 1994), and considering the localization and the tubular structures (Fig. 3B), we identified these areas to be the developing kidneys. Thus, we used the proximal tubule kidney marker, xSGLT1K (Zhou and Vize, 2004) to perform in situ hybridization in 4.5 dpf embryos. Our results showed that both gda and xSGLT1K are expressed in the same location (Fig. 3C), confirming the expression of gda in the kidneys.

We were also interested in determining a possible function of Gda in *Xenopus*. Because our overexpression analysis suggested that Gda does not play a role in modulating MT dynamics, and our expression analysis showed that gda is not present in most embryonic cells during early neural development, we reasoned that Gda cannot be playing a role to promote MT polymerization in the early nervous system. Given its strong expression in the kidneys beginning at 3 dpf, we wondered if Gda might play a role in kidney development or function. However, pronephros development in *Xenopus* begins at stage 12.5, and kidneys become initial functional by stage 38, which corresponds to 2 dpf (Vize et al., 1995; Wessely and Tran, 2011). Thus, it did not seem likely that Gda was required for initial kidney development.

Still, we wondered whether Gda might be important for kidney function. It has been demonstrated that an altered osmoregulatory function of the kidney leads to edema formation due to water retention (Wessely and Tran, 2011; Krneta-Stankic et al., 2017) and evaluation of edema has been widely used to assess kidney function (Getwan and Lienkamp, 2017; Krneta-Stankic et al., 2017). Thus, we used an antisense oligonucleotide to knock down gda at 4 dpf (Vize et al., 1995; Wessely and Tran, 2011). Thus, it did not seem likely that Gda was required for initial kidney development.

CRMP Homology Domain is not Conserved Between Human and *Xenopus* GDA

A GDA evolutionary study had determined that the *Rattus norvegicus*, *Xenopus laevis*, and *Homo sapiens* GDA proteins are part of the same sequence cluster, meaning that their sequences are evolutionarily similar (Fernandez et al., 2009). While the GDA enzymatic activity of guanine deamination into xanthine and ammonia has been conserved across many species (Fernandez et al., 2009), few studies have examined GDA function during embryonic development. One of the most prominent studies of GDA function has examined the role of the mammalian ortholog, cypin, during neuronal development (Akum et al., 2004; Chen et al., 2005; Charych et al., 2006; Chen and Firestein, 2007). Importantly, GDAs contain a PDZ-binding domain that is important for interaction and clustering with other proteins (Sheng and Sala, 2001). The cypin PDZ-binding domain is necessary for binding to PSD-95 protein (Firestein et al., 1999), which has been described to increase dendritic branching (Firestein et al., 1999) and indirectly affect MT dynamics (Sweet et al., 2011). Of interest, while the enzymatic active site and divalent cation binding domain are conserved among these species, the PDZ-binding domain is not present in either of the *Xenopus* homeologs (Fernandez et al., 2009).

Additionally, GDAs also have a CRMP homology domain that is known to bind tubulin heterodimers (Firestein et al., 1999; Fukata et al., 2002). The CRMP homology domain is necessary for inducing MT polymerization in vitro (Akum et al., 2004). Our work is the first to evaluate the sequence homology of the CRMP homology domain between cypin and Gda, showing that the CRMP homology domain is not fully conserved. These differences in the PDZ-binding and CRMP homology domains between cypin and Gda likely explain the divergence in function between the proteins as it relates to their ability to regulate MT dynamics.

GDA Does not Localize to the MT Plus-End

Our results show that exogenous expression of GFP-tagged cypin and Gda present a cytosolic distribution in living cells and are not enriched at MT plus-ends, where the majority of the proteins that regulate MT plus-end dynamics are localized.
Nevertheless, our observations are consistent with previous immunocytochemistry studies showing that cypin has a cytosolic localization in rat primary hippocampal neurons (Firestein et al., 1999; Akum et al., 2004; Chen and Firestein, 2007; Sweet et al., 2011). Additionally, while GDA is obtained primarily in the soluble fractions after subcellular fractionation, it is also present in other intracellular compartments, where it exists in isoenzymic forms (Kumar et al., 1965, 1967). Considering this evidence, even though GDA localization is not enriched on MTs, we cannot discard a possible GDA activity near MT lattice or plus ends, which could still allow cypin to have an indirect effect on MT dynamics.

**Overexpression of Cypin, but not Gda, Affects MT Dynamics In Vivo**

Using live imaging, we showed that cypin overexpression increases the MT growth velocity and MT track length in neuronal growth cones. These results are in accordance with previous studies showing that rat cypin promotes MT assembly and polymerization using in vitro assays (Akum et al., 2004). Additionally, it is well known that cypin increases dendritic arbor length (Akum et al., 2004) and that it interacts with PSD-95 (Firestein et al., 1999). Cypin interaction with PSD-95 leads to decreased dendrite branching, potentially caused by the disruption of the MT cytoskeleton (Charych et al., 2006). Of interest, PSD-95 was found to bind and reduce the MT binding of the known +TIP EB3, as well as reduce EB3 comet lifetime and velocity (Sweet et al., 2011). Thus, given that cypin has been shown to oppose PSD-95 function, and that PSD-95 reduces MT polymerization, it is possible that cypin expression in our study perturbs the ability of PSD-95 to inhibit MT polymerization.

However, we observed that expression of mammalian cypin, but not Xenopus Gda, promotes MT polymerization. This result is consistent with the fact that the CRMP homology domain, which has been shown to bind tubulin and regulate MT assembly (Schmidt and Strittmatter, 2007), is not fully conserved between cypin and Gda. Additionally, the cypin PDZ-binding domain that has been described to bind PSD-95 is responsible for regulating MT dynamics (Sweet et al., 2011) is absent from Gda. Finally, we found that by adding the cypin CRMP homology and PDZ-binding domains to the N terminal region of Xenopus Gda, we observed MT dynamics regulation similar to that of cypin. This result further supports the importance of the CRMP and PDZ-binding domain in regulating MT dynamics.

**Xenopus Gda is Strongly Expressed in the Embryonic Kidney**

We found that *gda* is highly expressed at 3 dpf, which corresponds to stages 41–44, a relatively late time point in Xenopus embryonic development. By this point, significant dendrite branching and elongation in Xenopus retinal ganglion cells has already occurred (stages 32–34) and the specialization into various dendritic subtypes is in process (stages 34–46) (McFarlane and Lom, 2012). While we cannot overlook the possibility of Gda being involved in the second phase of dendrite branching and differentiation, this hypothesis is unlikely due to the fact that whole-mount in situ hybridization showed minimal *gda* expression in the brain or neural tube in Xenopus. While the lack of *gda* staining in the neural tube is unsurprising given that cypin is absent from the mammalian spinal cord (Firestein et al., 1999), the lack of *gda* expression in the brain is striking, as cypin is highly expressed in the developing brain in mammals. This suggests that cypin may have evolved to serve a different function in mammals than that observed in Xenopus.

In the present work, we demonstrated by in situ hybridization that *gda* is expressed primarily in the kidney. This expression pattern is more restricted than that observed for cypin, which is expressed in the human nervous system, placenta, liver, and kidney, and absent from the heart, lung, muscle, and pancreas (Firestein et al., 1999). Moreover, cypin protein is expressed in rat kidney, liver, lung, brain, and spleen (Firestein et al., 1999). Thus, rat cypin expression pattern contrasts with that of human cypin, which is not expressed in the lung, suggesting that even within mammals, cypin has a slightly different expression pattern.

To date, most studies relating to GDA function have focused on its role in maintenance of homeostasis of triphosphate nucleotides, its role as a nitrogen source, its signal transduction pathway, and, lately, on its role in neuronal development. Although serum GDA function is often altered in kidney diseases (Caraway, 1966; Prodanov and Astrug, 1971; Kebejian and al-Khalidi, 1973; Kuzmits et al., 1980), whether GDA functions during kidney development or maintenance had not been addressed. Xenopus begin developing a primitive kidney called pronephroi by stage 12.5, which becomes fully functional by stage 38, approximately 2 days postfertilization. Later in development, the pronephroi is replaced by a more complex kidney structure, the mesonephroi (Vize et al., 1995; Wessely and Tran, 2011). In the present work, we showed that the peak *gda* expression is at 3 dpf, which is temporally coincident with a fully functional primitive kidney. Thus, a GDA effect on kidney functioning or maturation to mesonephroi might be expected. Moreover, knockdown of Xenopus GDAs resulted in edema formation, which has been shown to be indicative of kidney malfunction (Wessely and Tran, 2011; Kreneta-Stankic et al., 2017). Nevertheless, at the Xenopus developmental time point that we performed our studies, the pronephroi has still not matured to mesonephroi; thus, it remains to be determined if GDA has an effect on kidney maturation to mesonephroi, or if the effect is restricted to kidney function. Additionally, given that edema also results from cardiac and other developmental issues, it is also possible that other organ systems are affected by reduced GDA function.

**Conclusion**

The present study shows that human cypin and Xenopus GDA have different protein sequences and functions. In addition, our MT dynamics analysis suggests that the role of GDA in regulating MT dynamics is not conserved among species. Furthermore, *gda* is expressed with high specificity in the kidneys during Xenopus development. Thus, our study suggests a novel function for GDA and warrants further study of its potential role in kidney maturation and function.

**Experimental Procedures**

**Culture of Xenopus Embryonic Explants**

Eggs obtained from female X. laevis frogs (NASCO, Fort Atkinson, WI) were fertilized in vitro, dejellied, and cultured following standard methods (Sive et al., 2010). Embryos were staged
according to Nieuwkoop and Faber (1994), and embryonic explants were dissected at stage 22–24 and cultured as described (Lowery et al., 2012). All experiments were approved by the Boston College Institutional Animal Care and Use Committee and performed according to national regulatory standards.

**Guanine Deaminase Constructs and Sequence Alignment**

The *X. laevis* GDA L homeolog (GDA) sequence was designed based on the GDA annotated sequence from the laevis genome, version 9.1 (www.xenbase.org) using the transcript model of chr1L:128,874,669..128,903,678 (+ strand), which was predicted to contain the *X. laevis* GDA genomic sequence. The GDA sequence was acquired from Biomatik (Biomatik, Atlanta, GA) in pBluescript, and then subcloned into the GFP-pCS2+ vector to add a GFP tag to the N-terminus of GDA. The human cypin sequence, based on NCBI reference sequence NM_004293.4, was obtained from Origen (Origen, Rockville, MD) and subcloned into a GFP-pCS2+ to generate GFP-cypin and cypin-GFP. The cypin-chimera was obtained by removing the GDA region, beginning with the CRMP domain, and replacing it with the human cypin CRMP homology and PDZ-binding domain. The NCBI Reference Sequences for the GDA protein sequence comparison were the following: *Xenopus laevis* S homeolog [XP_018099342.1]; *Xenopus laevis* L homeolog [NP_001083074.1]; *Homo sapiens* [NP_001229434.1]; *Xenopus tropicalis* [XP_004910825.1]; *Danio rerio* [NP_001018510.1]; *Rattus norvegicus* [NP_113964.2]; *Mus musculus* [NP_034396.1]; *Gallus* [XP_424835.3]. The GDA protein sequences were compared using Clustal Omega (www.ebi.ac.uk/Tools/msa/clustalo/; RRID: SCR_001591).

**RNA and Morpholino**

Capped mRNA was transcribed in vitro using the SP6 mMessage mMACHINE Kit (Thermo Fisher Scientific, Waltham, MA). RNA was purified with LiCl precipitation and re-suspended in nuclease-free water. Constructs coding for GFP, GFP-GDA, GFP-cypin and cypin-GFP, cypin-chimera, and mKate2-MACF43 (a gift from Hogenraad Lab sub-cloned into pCS2+) were used as templates for RNA in vitro transcription. Embryos were injected at the two- to four-cell stage in each dorsal blastomere (in 0.1×MMR containing 5% Ficoll) with total mRNA amount per embryo of 1,000 pg of cypin, *gda*, and 200 pg of mKate2-MACF43.

A morpholino (MO) was designed specific to the *Xenopus GDA* L homeolog with the sequence 5’ TCCCCAACACAAAGTCCTTA CCACA 3’ and targeted to the exon 3 – intron 3 splice junction (Fig. 4C) (Gene Tools, LLC, Philomath OR). A standard control MO (5’-ccttacctacgattataa-3’) (Gene Tools, LLC, Philomath OR) was used as control MO. The GDA MO was injected at concentrations of 10, 20, and 50 ng per embryo to assess knock down of GDA (Fig. 4C). The GDA transcript levels were measured by RT-PCR, and the PCR product was sent for sequencing to ensure the introduction of a stop codon. A 50% KD was obtained when injecting 20 ng of the MO (Fig. 4A), which was used for the rest of the experiments to generate a partial KD phenotype.

**RT-PCR**

Whole RNA was extracted from embryos at various stages and adult *Xenopus* brain using Trizol reagent (Invitrogen, LifeTechnologies) and reverse transcribed using SuperScript IV reverse transcriptase (Invitrogen, LifeTechnologies). PCR was then performed with HotStarTaq Plus DNA Polymerase (Qiagen, Hilden, Germany) and primers specific to *Xenopus gda*: forward 5’CCCTT GTCCTGCCGATATTAC’ and reverse 5’ GTAGCAGCCATGAGCATCAC 3’ PCR was performed on the same samples of cDNA, also using primers specific to the housekeeping gene ODC1: forward 5’GCCATTTGAAAGACCTCTCCATC’ and reverse 5’T GGTTGATTCTCCTGCCAC’ as a RT-PCR control.

**Microscopy**

Live images of cultured cells were obtained with a CSU-X1M 5000 spinning-disk confocal (Yokogawa, Tokyo, Japan) on a Zeiss Axio Observer inverted motorized microscope with a Zeiss 63× Plan Apo.1.4 numerical aperture lens (Zeiss, Thornwood, NY). Images were acquired with an ORCA R2 charge-coupled device camera (Hamamatsu, Hamamatsu, Japan) controlled with Zen software. For time-lapse, images were collected every 2 sec for 1 min. For two-color images, the red and green channels were imaged sequentially. The images were deconvolved using the “Iterative Deconvolve 3D” plugin within ImageJ.

Lateral images of embryos at 4 dpf, for quantification of edema formation, in addition to in situ hybridization images, were obtained with a SteREO Discovery.V8 microscope using a Zeiss 1× objective for edema formation experiments and 5× or 8× objectives for in situ hybridization, and Axiocam 512 color camera (Zeiss, Thornwood, NY).

Images of immunostained fluorescent kidneys at 4 dpf were obtained with a Leica TCS SP5 scanning confocal inverted microscope (Leica, Buffalo Grove, IL) using a 10× objective. Lateral images of embryos were obtained in a BABB solution on a MatTek glass bottom dish.

**plusTipTracker Software Analysis**

MT dynamics were analyzed from mKate2-MACF43 movies from embryonic mesenchymal cells using plusTipTracker, as previously described (Applegate et al., 2011; Lowery et al., 2013; Stout et al., 2014). The same parameters were used for all movies: maximum gap length, eight frames; minimum track length, three frames; search radius range, 5–12 pixels; maximum forward angle, 50°; maximum backward angle, 10°; maximum shrinkage factor, 0.8; fluctuation radius, 2.5 pixels; and time interval 2 sec. Only cells with a minimum number of 10 MT tracks in a 1-min time lapse were included for analysis. All raw data of automated tracking of mKate2-MACF43 comets were normalized to the same-day control means. Dot plots were made using GraphPad Prism (GraphPad Software, La Jolla, CA). To determine statistical differences, unpaired two-tailed *t*-tests were used for comparing two conditions, after confirming normalized data distribution (GraphPad). A total of 83–154 growth cones were analyzed for each condition, from three independent experiments.

**Whole-Mount In Situ Hybridization**

Embryos at various developmental stages from two-cell to 6 dpf were fixed overnight at 4°C in a solution of 4% paraformaldehyde in phosphate-buffered saline (PBS), gradually dehydrated using ascending concentrations of methanol in PBS, and stored in methanol at –20°C before in situ hybridization, which was performed as previously described (Sive et al., 2007). Embryos
were treated with proteinase K for 1–2 min in the case of 1 dpf embryos and 3–5 min for 3–6 dpf embryos, and then bleached under a fluorescent light in 1.8X saline–sodium citrate, 1.5% H2O2, and 5% (vol/vol) formamide for 45 min before prehybridization. Probes for both GDA and the proximal tubule kidney marker, xSGLT1K (sodium glucose transporter), were used at a concentration of 0.5 μg/ml and hybridized overnight. Sglt1K construct pCMV-SPORT6, Sali/T7 was a gift from the Wessely Lab (Cleveland Clinic, Lerner Research Institute, Cleveland, OH). The full-length antisense digoxigenin-labeled hybridization RNA probes were transcribed in vitro using the T7 MAXIscript kit (Thermo Fisher Scientific, Waltham, MA). The probes were purified using ammonium acetate precipitation and resuspended in nuclease-free water.

Quantification of the Edema Formation
Embryos were injected with 20 ng of the control MO, Xenopus GDA MO, or Xenopus GDA MO plus 1,000 pg GDA mRNA, for rescue experiments. From lateral images of 4 dpf injected tadpoles, the formation of edema was analyzed by measuring the length from the top of the face to the bottom of the chest (face–chest), and as the size of the head can vary from tadpole to tadpole, the length from the middle of the cement gland to the middle of the eye (cement gland–eye) was used as an internal reference (Fig. 4C). The ratios of the former over the latter measurement were compared among the different conditions, to normalize the data for differences in tadpole size. The presence of edema in the embryo was defined as when the above ratio was 2× greater than standard deviation of ratios from the control MO injected embryos. Dot plots were produced using GraphPad Prism (GraphPad Software, La Jolla, CA).

Whole-Mount Immunostaining
Embryos were injected at the four-cell stage in the left ventral blastomere to target the developing kidney on the left side of the embryo, leaving the right side as an uninjected internal control. Embryos were injected with 20 ng of control MO with 300 pg of GFP mRNA or 20 ng of Xenopus GDA MO with 300 pg of GFP mRNA. At 4 dpf, the embryos were fixed overnight at 4°C in a solution of 4% paraformaldehyde in PBS. After 24 hr, the 4% paraformaldehyde solution was removed and embryos were dehydrated in multiple methanol washes, and subsequently rehydrated in a methanol/PBS series. The eyes and surrounding cartilage were removed from each embryo for ease of mounting when imaging. Embryos were bleached under a fluorescent lamp in a 1.8X saline-sodium citrate, 1.5% H2O2, and 5% (vol/vol) formamide solution for 30 min. After incubating the embryos in blocking solution (3% bovine serum albumin, 1% Triton X–100 in PBS) for 2 hr, embryos were incubated overnight at 4°C in 2% 4A6 and 2% 3G8 monoclonal antibodies (European Xenopus Resource Center) in blocking solution. After multiple rinses, embryos were incubated with 0.1% anti-mouse Alexa 488 overnight at 4°C. Embryos were then cleared overnight in BABB solution (33% benzyl alcohol, 66% benzyl benzoate).

Quantification of Pronephric Tubule Diameter
The embryonic kidneys were immunostained using both 3G8 and 4A6 antibodies, specific markers of pronephric tubules and nephro stomes, respectively, to obtain a full picture of the kidney. The diameter of each tubule at its widest point was measured at four layers through the immunostained embryonic kidney: 35%, 45%, 55%, and 65% through the sample. The average of the widest diameter was obtained for each sample, then a ratio was calculated between the average tubule diameter of the injected and uninjected sides. The Xenopus GDA knockdown condition was compared with the control MO condition, and the results were presented with a dot plot produced with GraphPad Prism (GraphPad Software, La Jolla, CA).

Statistical Analysis
Statistical analyses were performed using GraphPad Prism (GraphPad Software Inc). The D’Agostino and Pearson normality test was used to determine if the data was distributed normally. For MT dynamics analyses, the unpaired Student’s t-test was performed when comparing two conditions. To assess statistical significance when quantifying the edema formation, the Mann Whitney test was used, as GDA KD population did not pass the normality test. At least three independent experiments were carried out for each condition. The alpha value was set at 0.05 for all statistical tests, and the P values are represented as follows: *P < 0.05, **P < 0.01, ***P < 0.001 and ****P < 0.0001. Values are expressed as mean ± standard error of the mean.

Abbreviations
+TIPs plus-end-tracking proteins
CRMP collapsin response mediator protein
cypin cytosolic PSD-95 interactor
dpf days post fertilization
F-actin actin filament
cRNA guanine deaminase
KD knock down
MO morpholino
MT microtubule

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